Molecular approach for identification of mosquito species (Diptera: Culicidae) in Province of Alessandria, Piedmont, Italy

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Abstract. The aim of the present work was to identify cryptic species in the Anopheles maculipennis and Culex pipiens complexes and to study the genetic structure of the dominant mosquito species Ochlerotatus caspius (Diptera: Culicidae) in the Province of Alessandria close to the vast area untreated rice fields in Piedmont, NW Italy. With the help of PCR-RFLP analysis, four members of the Anopheles maculipennis complex were identified: A. messeae, A. maculipennis, A. sacharovi and A. atroparvus. Only C. pipiens f. molestus was identified in 11 habitats studied in Piedmont. Partial sequences of the cytochrome c oxidase subunit 1 (COI) mitochondrial gene and the second internal transcribed spacer (ITS2) of nuclear ribosomal RNA genes for Italian O. caspius are reported here for the first time. The results indicate that this species diverged from Iranian representatives of this species about one million years ago. The great diversity of mosquito species in Piedmont considerably increases the risk of vector-borne diseases.

INTRODUCTION

Exact identification of mosquitoes is essential for controlling the species that are vectors of human and animal diseases. It is particularly difficult to identify representatives of species complexes, such as Anopheles maculipennis and Culex pipiens (Diptera: Culicidae). The morphology of larvae and adults of species within these complexes are similar or identical. However, they do show considerable differences in ecological and physiological features, including food preferences. Their epidemiological significance is also different.

Importantly, the causative agents of many diseases are transferred between hosts by certain species of bloodsucking Culicidae (e.g. Turell et al., 2001; Becker et al., 2003; McAbee et al., 2008). In a recent study, the following dangerous viruses were detected in pooled mosquito samples in Italy: West Nile virus in C. pipiens, two orthobunyavirus and Taha virus in Ochlerotatus caspius and Batai virus in A. maculipennis (Cazolari et al., 2010). Since different species within these complexes can sustain arboviral outbreaks (Talbalaghi et al., 2010), specific diagnosis of the vectors is essential for developing control strategies.

Unfortunately, morphological features are often insufficient for identifying species of bloodsucking mosquitoes as some are very variable and consequently unreliable [for example, siphonal index of C. pipiens larvae (Fedorova & Shaikevich, 2007)], while many diagnostic features (especially paleoae and chaetae) are not present on collected material. This problem can be solved by using molecular-genetic analysis and developing DNA markers for specifically identifying species and subspecies of these insects.

One of the basic molecular-genetic markers used to study species composition of mosquitoes is the second internal transcribed spacer (ITS2) of nuclear ribosomal RNA genes (rRNA) genes. rRNA genes of mosquitoes are represented in the genome as a set of tandem repeats. Each transcriptional unit consists of the genes encoding three ribosomal RNAs (18S, 5.8S, and 28S) separated by internal transcribed spacers ITS1 and ITS2. The coding 28S gene sequence is followed by an external transcribed spacer. Transcriptional units are separated by un-transcribed spacers. The rRNA genes are conserved, whereas internal transcribed spacers are variable and useful for comparing species. It is known that the structure of ITS2 varies among different mosquito species (Porter & Collins, 1991), the length being from 421 bp in A. maculipennis to 442 bp in A. sacharovi, however A. messeae, A. melanoon, A. atroparvus, and A. labranchia have a fragment of similar length (Marinucci et al., 1999).

Recently, a complex of PCR-RFLP methods was developed to identify some cryptic species in the Anopheles maculipennis complex (Nicolescu et al., 2004; Ejob et al., 2008).

Another approach is to analyze the 5’ end of the cytochrome c oxidase subunit 1 mitochondrial gene (COI). This region, known as the DNA barcode (Barcode of Life Initiative, 2003), was used previously to study the Iranian O. caspius (Azari-Hamidian et al., 2009). For O. caspius in Italy, microsatellite and allozyme polymorphism and mitochondrial COI gene structure are reported (Porretta et al., 2007). Although the nucleotide structure of the ITS2 area of C. pipiens is variable it cannot be used to distinguish between C. pipiens and C. pipiens f. molestus, however, a method based on the analysis of the 5’end
of the mitochondrial COI gene can be successfully used to identify mosquitoes of the C. pipiens complex (Shaikevich, 2007).

In this study, simple PCR-based assays (amplification of the ITS2 within nuclear RNA genes of the A. maculipennis complex and amplification of the COI gene within mtDNA of C. pipiens complex followed by endonuclease digestion) were used for the molecular identification of cryptic species.

MATERIAL AND METHODS

Study area and sampling sites

The adults and larvae of the mosquitoes analyzed were collected in the Province of Alessandria close to the vast area of untreated rice fields in Piedmont, NW Italy. The Province of Alessandria is 10 km from the biggest mosquito breeding site in untreated rice fields in Piedmont, NW Italy. The Province of Alessandria is 10 km from the biggest mosquito breeding site in Europe, approximately 220,000 ha of rice fields, which produce 50% of the rice grown in Europe. These rice fields are unique in Europe, and the adults and larvae of the mosquitoes analyzed were collected weekly by a monitoring network of CO$_2$ traps (see Table 1). The genera and species complexes were identified morphologically.

DNA isolation and analysis

Mosquito DNA was extracted using DI Atom™ DNA Prep kit (Isogen, Moscow, Russia). DNA samples were obtained from individual larvae and adults, either dry or preserved in 96% ethanol. Prior to DNA isolation from the ethanol-preserved material, ethanol was evaporated off by heating at 65°C for 30–40 min.

PCR-RFLP assay

The mtDNA gene COI was amplified using primers LCO and HCO (Folmer et al., 1994) for studying C. pipiens and Culex-

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<th>Municipality</th>
<th>Sample</th>
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PCR amplification reactions were run in a final volume of 25 µl with PCR buffer (Isogen), 20 µM of each dNTP, 2.5 mM MgCl$_2$, one unit of Taq DNA polymerase, 0.2 µM of each primer and 0.1 µg of the isolated DNA. Thermocycler conditions were as recommended for each primer pair.

For the Anopheles maculipennis complex, the improved PCR-RFLP method was used (Efjov et al., 2008). The PCR-RFLP was run using CfoI (Promega, Madison, WI, USA) and BstACI (Sibenzyme, Moscow, Russia) endonucleases. Restriction volumes were 20 µl. The digestion mixtures consisted of (i) 3 µl of the ITS2 PCR-product, 0.5 µl (5 U) of CfoI, 2 µl of the manufacturer’s incubation buffer, 0.2 µl BSA (Promega) and 14.3 µl ddH$_2$O, and (ii) 3 µl of the ITS2 PCR-product, 0.5 µl (5 U) of BstACI, 2 µl of the manufacturers incubation buffer and 14.5 µl ddH$_2$O (BSA is included in all Sibenzyme’s incubation buffers). Incubation time was not less then 2 h at 37°C.

For the precise identification of C. pipiens mosquitoes using the PCR-RFLP assay amplicons were digested with HaeIII (Promega) restriction endonuclease. Restriction enzyme digestions were performed in 20 µl volume. Digest master mix consisted of 5 µl of the COI PCR product, 0.5 µl (5 U) of HaeIII, 2 µl buffer C (Promega), 0.2 µl BSA (Promega) and 12.3 µl ddH$_2$O. The HaeIII restriction reaction was incubated for 1 h at 37°C. The digested products were visualized on a 2% agarose gel.

Sequence analysis

PCR products were identified by electrophoresis, using a 1% agarose gel (Sigma, St. Louis, MO, USA). Amplified DNA fragments were isolated from the gel using JetQuiCk Gel Extraction Spin Kit (Genomed, Loehne, Germany) and then directly sequenced on an ABI PRISM 310 using the ABI PRISM BigDye Terminator Cycle Sequencing kit (Applied Bio-
systems, Foster City, CA, USA) according to the instructions of
the manufacturer. The obtained sequences of
*O. caspius* DNA
are available in GenBank under accession numbers
HM140413–HM140418 for
*COI* and HM140419–HM140424
for *ITS2*. The DNA sequences were aligned using ClustalW and
analyzed using MEGA version 4 (Tamura et al., 2007).

**RESULTS AND DISCUSSION**

*Anopheles* spp.
The PCR with primers 5.8S and 28S produced ampli-
cons with lengths typical for mosquitoes of the complex
*A. maculipennis* (420–440 bp). An improved PCR-RFLP
method was used to identify species of the *A. maculipennis*
complex (Ejov et al., 2008). Identification was
based on species-specific restriction sites in the *ITS2*
locus. The recognition site for
*Cfo* I is the sequence
GCGC. After digestion with
*Cfo* I, variable fragments
were obtained (Fig. 1). Two fragments (56 and 42 bp) are
characteristic for all members of the
*A. maculipennis*
complex. Because of their low molecular weight these
fragments merged into one another on agarose gels (about
50 bp).

For individuals from population 36, 389-bp and 50-bp
fragments, common for *A. atroparvus*, were obtained.
DNA of the mosquito from population 33 was cut into
fragments of 207, 111, 78, and 50 bp, corresponding to *A.
sacharovi*. 50 bp is almost not visible in lane 33-1 on Fig.
1, but other visible fragments are specific only for *A.
sacharovi* (Nicolescu et al., 2004). For the individuals
from populations 25 and 67 (sample 67-2) restriction
fragments of 272, 102, and 50 bp were obtained. These
fragments correspond to *A. maculipennis*. DNA from
other individual from population 67 (sample 67-1) was
cut into fragments of 141, 135, 111 and 50 bp, as in both
*A. messeae* and *A. melanoon*. For exact identification of
the latter one an additional restriction reaction with
*BstAC* I was run, which indicated fragments of 292 and
150 bp, common for *ITS2* of *A. messeae* (not shown).

There is no site for
*BctAC* I in the ITS2 of the DNA of
*A. melanoon*.

The analysis of *Anopheles* spp. from various biotopes
using PCR-RFLP revealed four members of the *A. macu-
lipennis* complex: *A. messeae*, *A. maculipennis*, *A.
sacharovi*, and *A. atroparvus*.

Five species of the *A. maculipennis* complex are
common in Europe: *A. maculipennis*, *A. messeae*, *A.
melanoon*, *A. sacharovi*, and *A. atroparvus*. All are zoo-
and anthropophilic and attack humans and animals both
indoors and outdoors. After feeding on blood, *A. maculi-
ppennis*, *A. sacharovi*, *A. messeae*, and *A. atroparvus*
rest indoors (are endophilic), while *A. melanoon* rests out-
doors (is exophilic). *A. messeae* hibernates, while *A.
maculipennis*, *A. melanoon*, *A. sacharovi*, and *A.
atroparvus* can suck blood all year round, but do not lay
eggs in winter. Therefore, a correct identification of the
vectors is essential for the development of control strate-
gies. Since the mtDNA of *A. maculipennis* mosquitoes is
more polymorphic than the ITS regions of the rRNA tran-
scriptional units, using the *ITS2* sequences for the PCR-
RFLP analysis for identifying the species within this com-
plex is likely to prove more accurate. It has been possible
to identify species using these methods and these markers
are very promising for use in future studies.

**Culex pipiens**
The *C. pipiens* complex in Europe is represented by
*Culex pipiens* (Linnaeus) and *C. pipiens* f. *molestus*
(Forskal). Although very similar in morphology these two
forms are quite different in their physiology and behaviour. *C. pipiens* is rather ornithophilic while *C. pipiens f. molestus* is mostly anthropophilic. *C. pipiens* is anautogenous (unable to lay a batch of eggs until it has had blood meal) and eurygamous (unable to mate in confined spaces). In contrast, the *molestus* form is autogenous (can produce a first batch of eggs prior to feeding on a blood meal) and eugametic (mates in confined spaces). *C. pipiens* can hibernate, but not *C. pipiens f. molestus*. Although both forms are vectors there is no data on their relative abilities as vectors of disease.

Earlier, the DNA polymorphism in a 603-bp fragment of the 5' end of the mitochondrial *COI* gene using endonuclease *Hae*III restriction endonuclease to distinguish between anautogenous (*pipiens*) and autogenous (*molestus*) forms (Shaikievich, 2007). After restriction of amplificates of DNA of *COI* gene using endonuclease *Hae*III two fragments (206 and 397 bp) were recorded in anautogenous *C. pipiens*, whereas the 603 amplicon from autogenous *C. pipiens* remained unchanged (Fig. 2). This method was used in the analysis of mosquitoes from Piedmont. Some examples are presented in Fig. 2. After amplification and digestion with *Hae*III only the uncut 603-bp fragment was obtained for all the specimens of *C. pipiens* studied. Altogether 48 individuals of the *molestus* form were identified in Piedmont using restrictase *Hae*III. Almost all the *C. pipiens* studied were collected from cans, water containers and sewage water in an urban area. Homogeneous populations of the anthropophilic form *molestus* were recorded in 11 populations (see Table 1).

Mainly anthropophilic populations of *C. pipiens* are recorded in Egypt (Gad et al., 1999). In regions of southern Russia in summer, mosquitoes of the *molestus* form may temporarily establish homogeneous populations or coexist with the anautogenous form of *pipiens* in above-ground water bodies (barrels) (Vinogradova et al., 2007). Heterogeneous populations are also recorded in Germany, where some populations in water barrels, buckets, and garden ponds consist of up to 20% of the autogenous form (Becker et al., 1999).

These results confirm that *molestus* mosquitoes can survive if the temperature conditions are favourable in open habitats, although in northern zones these mosquitoes occur mostly underground and biotopically separated from the form *pipiens*.

**Ochlerotatus caspius**

The 563-bp sequences of the mitochondrial gene *COI* (Table 1). Four unique haplotypes were identified. The individuals studied varied by less than 1%. The mosquitoes from populations 1, 2 and 3 were identical in this 563-bp DNA fragment. DNA from a mosquito of population 7 differed by only one nucleotide. The highest polymorphism was shown by a mosquito from population 10: six nucleotide substitutions. The highest polymorphism was shown by a mosquito from population 10: six nucleotide substitutions. Most substitutions were at the third codon position and did not affect the amino acid sequence. Only the transition G → A at position 37 at the first codon position results in the amino acid substitution valine-isoleucine (Val → Ile); this substitution was detected in two of the mosquitoes studied from populations 10 and 15 (Fig. 3).

GenBank search revealed that the *COI* sequences of *O. caspius* from Piedmont share a 98% similarity with specimens from Iran (Azari-Hamidian et al., 2009). The neighbour-joining dendrogram shows two clusters: the first is formed by Italian and the second – Iranian *O. caspius* (Fig. 4).

In addition, sequences of the variable ITS2 region were analyzed in the ribosomal RNA of *O. caspius*. Samples from the same populations: 1, 2, 3, 7, 10, and 15 (Table 1) were studied. The size of the obtained PCR fragments of the ITS2 region of *O. caspius* was about 370 bp (not shown). Primers were selected so that the ITS2 sequences

### COI

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<tr>
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### ITS2

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![Fig. 4. Dendrogram based on 563 bp *COI* gene mtDNA sequence data of the relationship of the haplotypes within *O. caspius*. Och1, Och2, Och3, Och7, Och10, and Och15 – *O. caspius* from Piedmont; the sample numbers correspond to those in Table 1. The tree was constructed using neighbour-joining method with Kimura 2-parameter distances. Numbers are bootstrap percentages (1000 replicates).](image-url)
were flanked with a 88-bp fragment of the 5.8S and 45-bp fragment of the 28S rRNA genes. Therefore, the size of the actual ITS2 of O. caspius is about 254 bp.

The ITS2 of the five individuals investigated was almost identical. Only one mosquito differed from the others by two nucleotide substitutions and one deletion. Four variable sites were detected in six sequences (Fig. 3). Pair-wise distances between samples from different populations varied from 0 to 1.1%. In comparison with all ITS2 sequences of O. caspius from Spain and Tunis available in GenBank, the overall average pair-wise distance is below 0.2%.

Unfortunately, only the mtDNA of Iranian mosquitoes was studied and only ITS2 of Tunisian and Spanish, mosquitoes. Consequently, it is difficult to compare the genetic structure of geographically remote populations of O. caspius using the same marker. The polymorphism in mtDNA of Italian and Iranian O. caspius is 10 times more than that of the ITS2 sequences of Italian, Spanish, and Tunisian O. caspius. O. caspius is thought to have originated in central Asia and later spread to Western Europe and North America (Ross, 1964; Minar, 1976 cited from Milankov et al., 2009). The nucleotide sequence divergence in insect mtDNA is thought to be approximately 2% for pair-wise divergence per million years (Guillemaud et al., 1997). If so, the differences in COI suggest that it is about one million years since the divergence of the geographically distinct populations in Iran and Italy. During this period, the spread in the distribution of insects was strongly affected by the availability of their habitats, and probably, the distinctions operated at the subspecies level. It would be interesting to compare the DNA of O. caspius from Italy and central Asia with other populations of this species using the same set of mitochondrial and nuclear markers.

CONCLUSIONS

By using molecular analysis it was possible to confirm the presence in the Province of Alessandria of several species of mosquito that are capable of transmitting infections.

The DNA analysis revealed that the Anopheles maculipennis complex in the area of Piedmont includes A. messeae, A. maculipennis, A. sacharovi, and A. atroparvus, and the Culex pipiens complex is represented by the anthropophilic form molestus. All these species are of high epidemiological significance in Mediterranean countries.

It is suggested that the level of genetic homogeneity in O. caspius in the region of Piedmont is high, even though only six individuals were studied. Our data accord with the conclusions reached using other markers such as allozymes and the gene COI, which are not significantly genetically diverse in O. caspius from the plain of the Po River in the north of Italy (Porretta et al., 2004). Probably, there are no genetic barriers preventing crossing between populations of O. caspius in northern Italy; this could increase the risk of rapid population growth in this favourable rice field habitat.

A change in the climate in the Piedmont region, together with the development of insecticide resistance, could result in a rapid increase in the abundance and geographic range of mosquito populations. This could complicate the development of an effective control of mosquitoes. The use of a DNA-based method to confirm the identity of mosquito species is likely to greatly reduce the risk of misidentification and results in a more efficient monitoring of control programs.

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**METAPATHOGEN**: http://www.MetaPathogen.com


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