Molecular Identification, Phylogenetic Status, and Geographic Distribution of Culicoides oxystoma (Diptera: Ceratopogonidae) in Israel

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Abstract

Culicoides oxystoma (Diptera: Ceratopogonidae) is an important vector species, reported mainly from Asia, with high potential to transmit viral diseases affecting livestock. In Japan, many arboviruses have been isolated from C. oxystoma, suggesting it as a key player in the epidemiology of several Culicoides-borne diseases. Over the years, C. oxystoma has also been reported in the Middle East region, including Israel. In this region, however, C. oxystoma cannot be easily distinguished morphologically from its sibling species included in the Culicoides schultzei complex. We therefore used genomic data for species identification and phylogeny resolution. Phylogenetic analyses based on internal transcribed spacer 1 (ITS-1) of ribosomal DNA and the mitochondrial gene encoding cytochrome oxidase subunit I (COI) showed that C. oxystoma from Israel is closely related to C. oxystoma from Japan. Using differential probing PCR, we showed that C. oxystoma is distributed all over the country, especially in Mediterranean climate regions. *Culicoides oxystoma* is less common or even absent in arid regions, while the other genetic cluster of C. schultzei complex was found only in the east of the country (mostly arid and semiarid regions). The molecular finding of C. oxystoma in wide geographical regions, together with its high proportion in the general Culicoides population and its vectoring potential, imply that it may be an important vector species in the Middle East.

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Introduction

Culicoides biting midges (Diptera: Ceratopogonidae) are among the smallest hematophagous flies. They occur throughout the world, except New Zealand and the Antarctic. They play an essential role in the epidemiology of several major veterinary diseases as vectors of arboviruses which affect ruminants and equines [1]. In livestock, diseases caused by bluetongue virus (BTV) and epizootic hemorrhagic disease virus (EHDV), among others, are of international significance, and have an important impact on the economy and animal welfare [2,3]. Several *Culicoides* species have been associated with the transmission of BTV and EHDV, hence restricting these diseases to areas where specific competent vector species occur and to their seasonal activity [4–6]. Since the intimate relation between a given virus and a vector determines the latter’s competence [7], the taxonomic status of *Culicoides* vector species is essential for defying their role in disease epidemiology. Taxonomic identification of *Culicoides* species is based on morphological parameters that include wing pattern, measurements and proportions of antennal and palpal segments, counts of their various sensilae and genitalia [8]. This routinely used assessment can be time-consuming, particularly among *Culicoides* complexes, and is not always reliable for differentiating closely related species [9,10]. Few taxonomists are able to distinguish roughly between sibling species, thus classification may result in misidentification [11]. It is therefore most efficient to use molecular and morphological methods in parallel, since specimen mounting is usually not efficient for downstream molecular procedures and vice versa.

Common and dominant *Culicoides* species recorded in Israel, in the eastern Mediterranean region and in Africa are *Culicoides imicola* and midges belonging to *Culicoides schultzei* complex [1,12]. *Culicoides imicola* sensu stricto, the most important proven vector of BTV [1,13], is quite easily identified by its typical wing pattern. Some members of the *C. schultzei* complex are also considered as potential vectors of BTV and EHDV [14]. This complex consists of several species described in the past (C. oxystoma, *C. schultzei*, C. subschultzei, C. kingi, *C. rhizophorensis*, *C. enderleini*, *C. nevilli* and *C. nosschultzei*) [15,16]. Although the taxonomic status of *C. schultzei* complex has been studied [13,15–20], the literature reveals confusing and inconsistent reports. Callot et al. [18] first reported on samples identified as *C. schultzei* Enderleini found in Israel which, several years later, were named kingi-schultzei gp. in light of the great morphological variability found among individuals [17]. Later, Cornet [19] described the discrepancies in this group and recorded six African species in a preliminary note. These species were then named and described in detail [15]. At that time, the taxonomic status of *Culicoides* BTV vectors was discussed by Wirth and Dyce [20], who referred to *C. schultzei* as a complex of species which requires further study. In that report, the identity of...
Culicoides oxystoma, a common species in Asia closely related to C. schultzei, was also stated as unclear. In Israel, C. oxystoma was reported as a member of the C. schultzei complex [12,21]. Furthermore, three different morphological types of C. oxystoma were recorded, one of which was similar to the identified C. oxystoma sensu Arnaud from Japan [15]. Boorman [16] noted that the exact identity of this species is in doubt and evaluated that most of the records of C. schultzei group from north of the Sahara refer to C. oxystoma.

The inconclusive taxon status of C. schultzei complex, together with the expertise required to morphologically differentiate the members of this group, raise the need for less subjective identification methods that are based on genomic data. Although genomic records are unavailable for C. schultzei complex, previous studies have successfully initiated reliable molecular methods for the identification and definition of phylogenetic status of other Culicoides species. These methods are based on several genomic data including internal transcribed spacer 1 (ITS-1) and 2 (ITS-2) regions of ribosomal DNA [22–29], as well as the mitochondrial gene cytochrome oxidase subunit I (COI) and subunit II (COII) [10,30–33] that are performed using regular PCR, real-time PCR, and micro-array. Recently, identification method based on mass-spectrometry was developed [34].

In this work, we applied molecular methods to simplify the identification of the commonly found C. schultzei complex, showing that some midges morphologically classified as C. schultzei complex from Israel are genetically closely related to C. oxystoma from Japan. As this species has the potential to transmit important arboviruses [16,35], we analyzed its phylogenetic status and its geographical distribution to evaluate its vectoring potential in the Middle East region.

Results

Morphological identification of Culicoides species

A total of 217 specimens were classified based on wing patterns as members of the C. schultzei complex. These specimens could be easily morphologically differentiate from C. imicola found in the same collection batch. Although wing characterization was unique for each species, slight variations in wing pattern among individuals within a species was frequently observed (Figure 1). Due to the inconclusive taxon status of C. schultzei complex, together with the expertise required to morphologically differentiate its members, no attempt was made to distinguish between the species of this complex.

Molecular identification and phylogenetic analyses of Culicoides species based on ITS-1, and COI gene

Midges identified morphologically as C. schultzei complex (n = 10, two collection site: Newe-Ur, located in the Jordan Valley in a semi-arid climate region, and Bet-Elazari, located in the center of the country in a Mediterranean climate region) were subjected to specific PCR using the ITS-1 primers. This PCR resulted in two different band sizes (Figure 2A) that were sequenced. The obtained sequences [GenBank: JN408469–JN408478] were closely related to the published sequence of C. oxystoma [GenBank: AB462279] [36]. The upper bands consisted of 420 to 423 bp and showed 87% similarity to C. oxystoma, whereas lower bands were 400 to 408 bp and showed 90% similarity to C. oxystoma (Figure 2A). According to sequence differences, species specific PCR for ITS-1 region was developed to differentiate between the first (Figure 2B) and the second (Figure 2C) genetic groups of the C. schultzei complex.

Phylogenetic trees generated with the two different methods: maximum likelihood (Figures 3A, 4), and neighbor joining (data not shown) revealed that the 10 individual midges identified morphologically as C. schultzei complex split significantly into two different genetic groups with an average of 83% similarity between the groups. One genetic group consisted of three midges collected in Neve-Ur (99.05% sequence homologies within the group) while the second genetic group consisted of seven midges collected from both locations (Neve-Ur and Beit-Elazari) (97.94% sequence homologies within the group) together with the available sequence of C. oxystoma.

Parallel PCR amplification of COI gene from the same specimens showed uniformity in band sizes (data not shown), yet their sequences [GenBank: JN545045–JN545054] revealed variation, showing between 90 and 98% similarity to the published sequence of C. oxystoma [GenBank: AB360979]. Consistent with the ITS-1 data, phylogenetic analysis based on the COI gene of the sequences obtained in this study and 8 C. oxystoma sequences from Japan [37], showed that midges (10 individuals) identified morphologically as C. schultzei complex in our study are...
significantly separated into two different genetic populations (Figure 3B) with an average of 91% similarity between the two groups. The first group which consisted of 3 individuals (99.3% sequence homologies within the group) showed 90% similarity to the published sequence of *C. oxystoma* and branched in the same clade of other species belonging to subgenus *Avantis*: *C. brevitarsis, C. obsolutes* [28] (Figure 4). COI gene sequences obtained from the same *C. imicola* specimens [GenBank: JN545055, JN545056] showed 100% nucleotide sequence identity to each other and to several published *C. imicola* sequences from Israel and Europe [30], and these constituted a phylogenetically distinct population positioned in a separate clade from the members of the *C. schultzei* complex (Figure 3B).

**Species distribution within three climate regions**

Based on ITS-1 band size which easily distinguished *C. oxystoma* from other *C. schultzei* complex species on an agarose gel, we determined the distribution of the two genetic groups in 15 settlements along three different climatic regions. *C. oxystoma* (*n = 156*) was shown to be widespread and prevalent in almost all geographical regions within the Mediterranean, semi-arid, and arid climates, whereas a separate genetic cluster of the *C. schultzei* complex (*n = 61*) was found in the eastern part of the country (Figure 5). Both genetic clusters were less common or absent in the arid climate regions.

**Discussion**

Despite much discussion of the taxonomic status of *C. oxystoma* as a member of the *C. schultzei* complex in Israel over the years, it remains unclear [12,15–20]. It has been proposed to be the Asian analog of the Afrotropical species *C. subschultzei* [8]. Here we suggest that the molecularly identified *C. oxystoma* from Israel is closely related to *C. oxystoma* from Japan rather than to the other morphologically comparable and indistinguishable members of the *C. schultzei* complex found in Israel. These results are in agreement

Cloning and sequencing of the ITS-1 rDNA of *C. imicola* (two individual midges from the two different collection sites) revealed 392 bp sequences [GenBank: JN408479, JN409480] which were 99% identical to each other and to the published sequence of *C. imicola* from the island of Corsica, France [20] [GenBank: AY96114]. These two *C. imicola* sequences (Figures 3A, 4) were positioned together with another published sequence of *C. imicola* and branched in the same clade of other species belonging to subgenus *Avantis*: *C. brevitarsis, C. obsolutes* [28] (Figure 4). COI gene sequences obtained from the same *C. imicola* specimens [GenBank: JN545055, JN545056] showed 100% nucleotide sequence identity to each other and to several published *C. imicola* sequences from Israel and Europe [30], and these constituted a phylogenetically distinct population positioned in a separate clade from the members of the *C. schultzei* complex (Figure 3B).
with the previously found morphological similarity of a *C. oxystoma* specimen from Israel to the identified *C. oxystoma* sensu Arnaud from Japan [15].

Variation between individual sequences in the same cluster and the phylogenetic analyses suggest that other genetic clusters within the *C. schultzei* complex are present that are not related to geographical location. Further analyses based on other genomic data may better resolve this complex. Our analyses can serve as a foundation to the future development of a simple identification method for other members of the *C. schultzei* complex.

Molecular identification of *C. oxystoma* is of great importance as this species is the most important potential vector of several arboviruses in Japan, with 85.9% of all viruses isolated from this species causing major livestock diseases, such as EHDV [35]. It would therefore be interesting to determine whether this important potential vector species plays a role in the epidemiology of *Culicoides*-borne diseases in Israel and the Middle East. In the past, members of the *C. schultzei* complex in Israel have been suspected as vectors for arboviruses [38-40]. Braverman et al. [21] evaluated the vectorial potential of three common species, including *C. oxystoma*, for BTV based on three criteria: the mean periods and survival rates between blood meals, and expectation of the species remaining infective throughout its life. Although *C. oxystoma* was present in the field for only a short period (ca. 2 weeks) during the BTV season, it showed the highest mean period between blood meals as well as the highest range of expectation to remain infective, suggesting that this species could serve as an important augmenting vector during that limited period. Although previous attempts to isolate arboviruses from pools of midges of the *C. schultzei* complex from Israel have been mainly unsuccessful [39], studies from other countries have demonstrated the isolation of arboviruses from this species complex [14,35]. In addition, BTV was shown to multiply in an Israeli *C. schultzei* gp. midges following intrathoracic inoculation [38]. Therefore, further attempts to
isolate viruses from recently field-collected *C. oxystoma* in Israel recent, as well as vector-competence experiments in the laboratory are required to determine the vectorial capacity of the *C. oxystoma* described in our study.

Other criteria which contribute to the vectorial capacity of Culicoides species are their prevalence, seasonality and geographical distribution. In this work, the molecularly identified *C. oxystoma* was dominantly found in almost all collection sites distributed in different geographical and climatic regions in Israel, suggesting this species' potential to serve as a vector.

In addition to *C. oxystoma*, we molecularly sequenced other species of the *C. schultzei* complex. To the best of our knowledge, these are the first reported sequences, other than those of *C. oxystoma*, representing the *C. schultzei* complex. Sequences of ITS-1 and COI in this other member of the species complex, showed 87 and 90% identity, respectively, to the available GenBank sequences of *C. oxystoma* from Japan. Phylogenetic analyses of this molecularly identified species place it in a separate clade and supports a different genetic population within the *C. schultzei* complex. Distribution studies of this distinct genetic group of the *C. schultzei* complex showed that it is mostly found in semiarid regions, suggesting an Afrotropical origin. Additional investigation of this species' genetic composition may further differentiate it into separate genetic groups and give a clue as to its origin.

Species distribution in this study revealed that both genetic clusters of the *C. schultzei* complex were less common or even absent in comparison with *C. imicola* in the sampled settlements located in the southern parts of the country (Negev Desert). This suggests that arid environmental conditions might be less suitable for this species complex's breeding and survival. On the other hand, midges from the *C. schultzei* complex are prevalent in Africa [15] and have been found in the Sinai Desert [41,42]. Thus, it is possible that the data in our study represent genetic populations that differ from those found in Africa, or that different trapping methods reveal different species assembles.

The wide distribution of *C. oxystoma* described herein places this potential vector species as an important Culicoides species in Israel. The molecular findings for *C. oxystoma* in this study may serve in future studies of vectoring capacity and epidemiology of Culicoides-borne diseases and their prevention.

**Methods**

**Ethics statement**

No specific ethics permits were required for the described studies. The collection of midges were coordinated and permitted by the farm owners. Collections were not performed in national parks or other protected area of land, and did not involve endangered or protected species.

**Collection of adult Culicoides species**

Midges were collected from 15 settlements located in different geographical and climatic regions throughout Israel during the summers of 2009–2010 (Table 1) using UV-light traps with a suction fan (John W. Hock Company, Gainesville, FL, USA) placed 1.5 to 2 m above the ground on cattle or horse farms. Insects were collected overnight into a plastic beaker containing absolute ethanol and kept at 4°C until analysis.

Two of the 15 settlements were used as collection sites for phylogenetic analysis. The first site, Neve-Ur, is located in the northern Jordan valley (south of the Sea of Galilee, in a semiarid region), and the second site, Beit-Elazari, is located in the center of the country (Mediterranean region). To study the geographical distribution of the molecularly defined species, we sampled midges from all settlements.

**Morphological identification**

Initial identification of Culicoides species was based on wing-spot pattern and other morphological keys [8]. While *C. imicola* wings are pale with darker markings, *C. schultzei* complex wings are characterized by two pale spots. Wings, head and abdomen of individual midges were mounted in a mixture of Canada balsam and phenol on microscope slides for morphological identification [43], leaving thorax and legs for subsequent DNA extraction [29].

**Genomic DNA extraction and PCR amplification**

Individual midges were washed with sterile double-distilled water and ground using a motorized hand homogenizer (pellet-pestle; Sigma, St. Louis, MO, USA) in a microcentrifuge tube containing 50 μL ice-cold lysis buffer as described by Frohlich et al. [44].

For PCR amplification of approximately 400 bp of the ITS-1 region of Culicoides, genus-specific primers PanCulF 5’-GTAGGTT-GAACCTGCGGAAGG-3’ and PanCulR 5’-TCCGCTCCTTCATCGAACCCT-3’ were synthesized (Metabion, Martinsried, Germany) and used according to the published cycling profile as previously described [22]. Amplification of the COI gene was carried out using primers C1-J-1718m 5’-GAGGATTTGGAATTTGATTAGT-3’ and C1-N-2191m 5’-TGCGGTCTTC-TGCGGTCTTGTCACGACA-3’ synthesized (Metabion, Martinsried, Germany) and used according to the published cycling profile as previously described [22]. Amplification of the COI gene was carried out using primers C1-J-1718m 5’-GAGGATTTGGAATTTGATTAGT-3’ and C1-N-2191m 5’-TGCGGTCTTC-TGCGGTCTTGTCACGACA-3’ synthesized (Metabion, Martinsried, Germany) and used according to the published cycling profile as previously described [22].

Table 1. Geographic trap locations used in this study.

| Climatic region | Altitude (m) | Geographic coordinates | Trap location | No. |
|-----------------|--------------|------------------------|---------------|-----|
| Mediterranean | 185 | 33°01’16"N, 35°34’37"E | Ayelot Hashahar | 1 |
| Mediterranean | 138 | 32°54’17"N, 35°32’59"E | Vered Hagall | 2 |
| Semiarid | -185 | 32°35’15"N, 33°33’13"E | Neve Ur | 3 |
| Mediterranean | 27 | 32°24’06"N, 34°55’46"E | Givat Haaim | 4 |
| Mediterranean | 22 | 32°22’49"N, 34°52’29"E | Kfar Vitkim | 5 |
| Mediterranean | 29 | 32°11’56"N, 34°49’36"E | Rishpon | 6 |
| Mediterranean | 52 | 32°02’32"N, 34°49’31"E | Ramat Gan | 7 |
| Arid | -215 | 32°00’00"N, 35°26’43"E | Gilgal | 8 |
| Mediterranean | 82 | 31°57’23"N, 34°55’31"E | Ben Shemen | 9 |
| Mediterranean | 77 | 31°55’20"N, 34°49’21"E | Netzer Sereni | 10 |
| Mediterranean | 54 | 31°50’45"N, 34°48’34"E | Beit Elazari | 11 |
| Mediterranean | 61 | 31°43’07"N, 34°43’45"E | Kfar Warburg | 12 |
| Arid/Semiarid | 97 | 31°18’14"N, 34°31’09"E | Urim | 13 |
| Arid/Semiarid | 185 | 31°16’50"N, 34°27’05"E | Ein HaBesor | 14 |
| Arid | 120 | 29°53’46"N, 35°03’34"E | Yotvata | 15 |

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30 cycles of 1 minute at 94°C, 30 seconds 53°C, 30 seconds 72°C and a final elongation of 10 minutes at 72°C. All amplifications were carried out in a total reaction volume of 25 μL using GoTaq colorless master mix (Promega, Madison, WI, USA), 2 μL DNA and 1 μL of 10 μM of each primer. Amplification was performed in a thermal cycler (Biometra, Goettingen, Germany). PCR products were stained with EZ-vision (Amresco, Solon, OH, USA), loaded onto a 1.5% agarose gel and visualized under UV light.

Cloning and sequencing

PCR products of each of the genes (ITS-1 and COI) of 12 individual midges (10 of C. schultzei complex, 2 of C. imicola) were cloned into the pCR2.1 TOPO TA Cloning System (Invitrogen, Carlsbad, CA, USA) and transformed into HIT Competent Escherichia coli DH5α (RBC Bioscience, Taipei, Taiwan). Transformed colonies grown on LB (Lysogeny broth) agar containing ampicillin, X-gal (5-bromo-4-chloro-indolyl-galactopyranoside) and IPTG (Isopropyl β-D-thiogalactopyranoside) were selected according to color differentiation. Clone inserts were reamplified using the above-described PCR. Inserts containing plasmids were isolated with the DNA-spin™ kit (Intron Biotechnology, Kyungki-Do, Korea). DNA sequencing was carried out utilizing the BigDye Terminator cycle sequencing chemistry from Applied Biosystems (ABI, Carlsbad, CA, USA), the 3700 DNA analyzer, and the ABI data-collection and sequence-analysis software (Center for Genomic Technologies, The Hebrew University of Jerusalem). Sequences were analyzed using DNAMAN (Lynnon Corporation, Pointe-Claire, Canada) and deposited in GenBank.

Phylogenetic analysis

A total of 30 sequences of Culicoides ITS-1 available in GenBank (chosen according to representative clades identified in [28]), were used for phylogenetic analysis, including an outgroup and the sequences from 12 individual midges obtained in this study. In addition, phylogenetic analysis of a total of 21 COI gene sequences including 8 GenBank published C. oxystoma, one C. imicola, and the 12 midges obtained in this study was performed.

Multiple alignments of the DNA fragments were generated using MUSCLE [45]. Phylogenetic trees obtained from the alignments were created using the maximum likelihood method based on the Tamura 3-parameter best fit model [46] with 1000 bootstraps. The trees with the highest log likelihood are shown. The percentage of trees in which the associated taxa clustered together is shown next to the branches. All analyses were conducted in MEGA 5.10 [47]. In addition, neighbor joining trees were built (Data not shown).

Species distribution

A total of 217 midges morphologically identified as members of the C. schultzei complex were sampled from the traps located in 15 settlements throughout Israel with an average of 15 individuals from each settlement (range: 2–39). In one of the settlements (Yotvata), C. schultzei complex midges were absent and only C. imicola midges were collected. Culicoides were then classified according to ITS-1 band size on the agarose gel into two groups: C. oxystoma and C. schultzei complex. Species distribution was demonstrated on a topographic-climatic map of Israel generated using ArcGIS™ 9.3. Climate regions are based on the Köppen classification [48].

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Author Contributions

Conceived and designed the experiments: YG NM. Performed the experiments: NM YB. Analyzed the data: NM YG EK. Contributed reagents/materials/analysis tools: NM YS. Wrote the paper: YG NM YB.

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