Assessment of Distribution Patterns of Culex Disease Vectors by Molecular Assays

Abstract
Study of distribution patterns of Culex pipiens forms and their sibling species Culex torrentium in Berlin and Hannover, Germany was done. The mosquitoes of the Culex genus have epidemiological significance as vectors of arboviruses and other pathogens of humans and animals. Based on molecular genetics methods it was shown that the park in Berlin in 2009 was inhabited by Culex pipiens form pipiens, while in the water container in Hannover coexistence of Cx. pipiens form pipiens and Cx. torrentium was noted. All females carrying sucked human blood and caught in the flats belong to Cx. pipiens form molestus. Molecular differentiation revealed the coexistence of various Culex taxa in cities, which increases the probability of hybridization and the appearance of potential bridge vectors between birds and humans.

Keywords: Culex pipiens; Culex torrentium; PCR-RFLP; COI; ace-2; Wolbachia

Introduction

Culex pipiens complex taxa and sibling species Cx. torrentium are morphologically difficult to distinguish, but differ in behaviour and ecology and, accordingly, in epidemiology. With a view of their epidemiological importance, knowledge of the distribution of morphologically similar mosquitoes in nature is important for the understanding of their vector-pathogen dynamics and for their control. Mosquitoes of the genus Culex are vectors of many dangerous human and animal infections, such as filariasis, avian malaria and arboviruses - West Nile virus, Japanese encephalitis, St. Louis encephalitis, Sindbis virus and others. Besides this, people attacked by mosquitoes suffer from itching and allergic reactions. Mosquitoes of the Culex pipiens complex are of special interest. The complex includes closely related ubiquitous species. Common in Europe are Cx. pipiens Linnaeus, 1758 and sibling species Cx. torrentium Martini, 1925. The subspecies Cx. pipiens includes two forms, the nominate form and the molestus form. Within the temperate zone, these forms are biotopically isolated and have somewhat opposite biological features. C. pipiens form pipiens are unautogenous (able to produce of the first egg raft only after taking a blood meal), diapausing (able to enter reproductive diapause) and eurygamous (they need a large space for mating) mosquitoes, inhabit the open water bodies. While Cx. pipiens form molestus are autogenous (produce of the first egg raft without taking a blood meal), non-diapausing, stenogamous (able to mate in confined space, without swarming), and usually inhabit the underground water bodies. Cx. torrentium is morphologically and ecologically very similar to Cx. pipiens form pipiens, the mosquitoes are unautogenous, diapausing and eurygamous.

Identification of Cx. torrentium cause considerable problems and Cx. torrentium is often confused with Cx. pipiens. Unautogenous form pipiens and Cx. torrentium inhabit the same terrestrial water basins forming 'pure' or mixed populations [1,2]. In W Europe, Cx. torrentium is spread from Scandinavian Peninsula to Portugal, including Great Britain.

It is hard and often impossible to identify the species for a wild larva or imago of the Cx. pipiens complex; the morphological distinguishing traits are not obvious, the main traits being the genital structure in adults and siphonal index in larvae. However, these traits are variable, inducing problems in distinguishing between the two species [1,3]. Since the species of the complex show different physiology and behavior and are of different epidemiological significance, the methods for exact identification are being actively developed basing on DNA analysis (including PCR). Several methods have been developed to distinguish between Cx. pipiens and Cx. torrentium. One of the methods implies different lengths of the ITS2: 400 and 350 bp in Cx. pipiens and Cx. torrentium, respectively [4,5]. Another method suggested by Smith et al. [6] is based on polymorphism in intron 2 of the gene encoding acetylcholine esterase 2 (ace-2). Autogenous and anautogenous forms of Cx. pipiens differs in microsatellite signature (Bank & Fonseca, 2006) and with distinguished patterns of enzymes (Becker et al. [7]). The aim of this work was to study species composition of mosquitoes from the genus Culex in Berlin and Hannover, Germany with molecular-genetics methods.

Materials and Methods

Mosquito larvae were collected in September 2009 from a non-functional fountain in the park Turmstrasse, 10559 Berlin. In Hannover, mosquito larvae were collected from a container with rain water in a country house area, Prussweg, 30165 Hannover. The females carrying sucked human blood were caught in the flats of multi-storey buildings (Lubecker strasse, 10, 10559 Berlin and Prussweg 6, 30165 Hannover). The larvae and the imagos were partly dried and partly fixed with 96% alcohol for subsequent analysis. The collected larvae and imagos were morphologically...
characterized as *Cx. pipiens* s.l. by M.V. Fedorova and O.V. Bezzhonova.

DNA isolation, PCR and RFLP analyses were performed as described earlier [8]. The results were verified by analysis of the PCR products of the gene ace-2 [6] using the primers ACEpip, ACEtorr, and B1246s, suggested by the authors for Eurasia. *Wolbachia* infection was tested using PCR with primers wsp81F and wsp691R, complementary to a bacterial gene *wsp* [9].

The results of the PCR-RFLP assay were confirmed by selective sequencing of the gene COI. DNA sequencing was performed using the ABI PRISM 310 sequencer and the Applera (USA) reagents kit according to the instructions of the manufacturer. The obtained sequences were analyzed using software Chromas (http://www.technelysium.com.au), polymorphism analyses were conducted in MEGA6 [10]. The sequences of the COI gene amplification products were submitted to the GenBank (accession numbers HM008665-HM008672).

### Results and Discussion

As described earlier [8], after the PCR with primers CulexCOIF and CulexCOIR and subsequent restriction with *Hae*III of the PCR products two fragments, 206 bp and 397 bp were obtained for the *Cx. pipiens* form pipiens. The fragment of the *Cx. pipiens* form molestus COI gene remained unchanged (603 bp). For mosquitoes *Cx. torrentium* intraspecific variation of COI gene nucleotide sequence is characteristic and in recognition sequence of *Haell* endonucleases presence both adenine and guanine is possible: 60% -AGCC, 30% -GGCT (Shaikevich, 2009), therefore both variants of restriction are recorded. However, *Cx. torrentium* can be identified using *Bcl*I. After *Bcl*I restriction of the PCR products the DNA of *Cx. pipiens* of both forms is cut in three fragments: 406bp, 118bp and 79bp. The DNA of *Cx. torrentium* is cut into only two fragments: 524bp and 79bp.

Typical 603-bp amplification products were obtained after amplification with primers CulexCOIF and CulexCOIR for all 35 studied mosquitoes. The restriction analysis using *Bcl*I showed that all mosquitoes from Berlin belonged to *Cx. pipiens*, while 4 of 14 mosquitoes from Hannover were identified as *Cx. torrentium* (Table 1). The amplificates of 29 samples were cut by *Hae*III into two fragments, 206 and 397 bp. In 6 samples only uncut 603-bp PCR products were present, these were obtained from all the blood-sucked females collected in the flats and from one individual collected from water container in Hannover characterized as *Cx. torrentium* using *Bcl*I (Table 1). Additional analysis of the PCR products of the gene ace-2 confirmed that four individuals from Hannover population belong to *Cx. torrentium*. The other studied individuals belonged to *Cx. pipiens*.

### Table 1: The restriction analysis using *Bcl*I showed that all mosquitoes from Berlin belonged to *Cx. pipiens*, while 4 of 14 mosquitoes from Hannover were identified as *Cx. Torrentium.*

| Sample Site      | Sample Number | COI/HaeIII (bp) | COI/BclI (bp) | ACE (bp) | COI GenBank AC number |
|------------------|---------------|----------------|--------------|---------|-----------------------|
| Berlin, flat     | B1            | 603            | 406, 118 and 79 | 610     | HM008670              |
|                  | B2            | 603            | 406, 118 and 79 | 610     | HM008671              |
|                  | B3            | 603            | 406, 118 and 79 | 610     |                       |
|                  | B4            | 603            | 406, 118 and 79 | 610     |                       |
| Berlin, park     | B5            | 206 and 397    | 406, 118 and 79 | 610     | HM008668              |
|                  | B6            | 206 and 397    | 406, 118 and 79 | 610     | HM008669              |
|                  | B7            | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B8            | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B9            | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B10           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B11           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B12           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B13           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B14           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B15           | 206 and 397    | 406, 118 and 79 | 610     |                       |
|                  | B16           | 206 and 397    | 406, 118 and 79 | 610     |                       |
Selective sequencing of amplificates (Figure 1) has shown that the DNA of *Cx. pipiens* mosquitoes of form *pipiens* differs from DNA of form *molestus* by single nucleotide substitution: position 119 of the studied fragment of the gene COI. The DNA of mosquitoes of form *molestus* has a characteristic substitution of Guanine within the HaeIII site (GG'CC) with Adenine (AGCC) with a loss of the respective restriction site. The DNA of *Cx. torrentium* in the studied fragment differs from DNA of *Cx. pipiens* by 13 nucleotide substitutions (2.5%). These results are in agreement with earlier studies of the *Cx. pipiens* mosquitoes in Russia [2,8,9].

The results show that the anthropophylic females caught in multi-store buildings in Berlin and Hannover belong to *Cx. pipiens* form *molestus*. The non-functional park fountain in central Berlin is inhabited by *Cx. pipiens* form *pipiens*. Though the distance between the sampling sites is not long, populations of the form *pipiens* appear to exist separately, not mixing with individuals from multi-store buildings.

Of special interest is the mosquito population from the water container in Hannover, with coexistence of *Cx. pipiens* form *pipiens* and *Cx. torrentium*. Coexistence is not rare for these two species, earlier we recorded several mixed populations from Russia. *Cx. torrentium*, found in Russia was not infected by *Wolbachia*, contrary to *Cx. pipiens*, the populations of which contained up to 100% infected individuals (Vinogradova et al., 2007). *Wolbachia* infections in *Culex pipiens* complex mosquitoes in Germany also were detected [11]. The PCR with primers for the gene wsp of *Wolbachia* showed no positive results for all four individuals of *Cx. torrentium* from Hannover. Though the number of the tested *Cx. torrentium* mosquitoes is low, it can be suggested that this member of the *Cx. pipiens* complex is not infected by these symbiotic bacteria in Germany. Probably, bacterial infection play a key role in reproductive isolation of *Cx. pipiens* and *Cx. torrentium*, since it is known that *Wolbachia* induces cytoplasmic incompatibility in *Culex* mosquitoes.

In conclusion, the method of species identification used in this work proved efficient for *Culex pipiens* mosquitoes populations.
from N Germany. Only one amplification reaction and two restriction reactions are sufficient for exact identification of both Cx. pipiens forms, and also of Cx. torrentium. The analysis takes 2-3 days and is considerably less expensive than the direct sequencing of the PCR products. The DNA for the PCR-RELP analysis may be isolated from an individual or any part of an individual at any stage of development [12].

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