Direct evidence for an expanded circulation area of the recently identified Balkan virus (Sandfly fever Naples virus species) in several countries of the Balkan archipelago

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Abstract

Background: Recently, Balkan virus (BALKV, family Phenuiviridae, genus Phlebovirus) was discovered in sand flies collected in Albania and genetically characterised as a member of the Sandfly fever Naples species complex. To gain knowledge concerning the geographical area where exposure to BALKV exists, entomological surveys were conducted in 2014 and 2015, in Croatia, Bosnia and Herzegovina (BH), Kosovo, Republic of Macedonia and Serbia.

Results: A total of 2830 sand flies were trapped during 2014 and 2015 campaigns, and organised as 263 pools. BALKV RNA was detected in four pools from Croatia and in one pool from BH. Phylogenetic relationships were examined using sequences in the S and L RNA segments. Study of the diversity between BALKV sequences from Albania, Croatia and BH showed that Albanian sequences were the most divergent (9–11% [NP]) from the others and that Croatian and BH sequences were grouped (0.9–5.4% [NP]; 0.7–5% [L]). The sand fly infection rate of BALKV was 0.26% in BH and 0.27% in Croatia. Identification of the species content of pools using cox1 and cytb partial regions showed that the five BALKV positive pools contained Phlebotomus neglectus DNA; in four pools, P. neglectus was the unique species, whereas P. tobbi DNA was also detected in one pool.

Conclusions: We report here (i) the first direct evidence that the Balkan virus initially described in coastal Albania has a much wider dissemination area than originally believed, (ii) two real-time RT-PCR assays that may be useful for further screening of patients presenting with fever of unknown origin that may be caused by Balkan virus infection, (iii) entomological results suggesting that Balkan virus is likely transmitted by Phlebotomus neglectus, and possibly other sand fly species of the subgenus Larroussius. So far, BALKV has been detected only in sand flies. Whether BALKV can cause disease in humans is unknown and remains to be investigated.

Keywords: Bunyaviridae, Phlebovirus, Arbovirus, Toscana virus, Meningitis, Fever, Sand fly, Phlebotomus, Phylogeny, Emergence

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Phleboviruses (family Phenuiviridae) are arthropod-borne viruses transmitted by mosquitoes, ticks and sand flies to vertebrate hosts [1]. Several phleboviruses belong to the Sandfly fever Naples species complex (which include at least two human pathogens, namely Toscana virus causing neurological infections and Naples virus causing incapacitating febrile illness) [2]. In the Old World, sand fly-borne phleboviruses are transmitted by Phlebotomus spp. and Sergentomyia spp. and show a wide distribution in all countries of the Mediterranean basin [2], http://ecdc.europa.eu/en/healthtopics/vectors/
vector-maps/Pages/VBORNET_maps_sandflies.aspx. During the last decade, several new phleboviruses were discovered in Mediterranean countries either in sand flies [3–9] or clinical samples [10]. Each was genetically related to any of the three following groups (based on antigenic relationships): Sandfly fever Naples species, Salehabad and Sandfly fever Sicilian/Corfou virus group. In the Balkans, the current knowledge on circulating phleboviruses is limited. Recently, the Balkan virus (BALKV) was discovered in sand flies collected in Albania and genetically characterised as a member of the Sandfly fever Naples species complex [11]. Two specific quantitative real-time RT-PCR assays were designed to screen entomological specimens collected in the surrounding countries, i.e. Croatia, Bosnia and Herzegovina (BH), Kosovo, Republic of Macedonia (RoM), and Serbia, to gain knowledge concerning the geographical area where exposure to BALKV exists.

**Methods**

Sand flies were collected in the field in 2014; 10 stations in Kosovo and 8 stations in Serbia, in 2015; 5 stations in Croatia, 6 stations in BH, 5 stations in RoM, 1 station in Montenegro and 1 station in Serbia (Table 1) using a previously described method [11]. Traps were placed near animals with the consent of the owners. BALKV RNA was detected using 2 SYBR Green real-time RT-PCR specific assays targeting the polymerase gene (BALKV-L-F; 5′-CTD ATY AGY TGC TGC TAC AAT G-3′, BALKV-L-R; 5′-CCA TAA CCA AGA TAY TCA T-3′) and the nucleoprotein gene (BALKV-S-F; 5′-AGA GTR TCT GCA GCC TTT GTT CC-3′, BALKV-S-R; 5′-CAG CTA TCT CAT TAG GYT GT-3′). The cycling program consisted of 50 °C for 30 min and 95 °C for 15 min, followed by 40 cycles at 94 °C for 15 s, 60 °C for 30 s, and 72 °C for 45 s, with a final melting curve step at 95 °C for 1 min, 60 °C 30 s and 95 °C for 30 s. Melting curves for positives were at 75 °C for the polymerase assay and 79.5 °C for the nucleoprotein test.

Phylogenetic relationships were reconstructed using sequences of the S and L RNA segments. Positive samples were PCR-amplified targeting a portion of the polymerase [12] and the nucleoprotein genes [13, 14] (two systems producing overlapping sequences which were concatenated before analysis). Sand fly species identification within positive pools was performed using as previously described cytochrome c oxidase subunit 1 (cox1) and cytochrome b (cytb) barcoding gene regions followed by NGS sequencing.

**Fig. 2** Phylogeny of the Balkan virus and closely related phleboviruses using partial nucleotide sequences of the nucleoprotein gene (572 nt). Neighbor-joining analysis (Kimura 2-parameter model) was performed using MEGA6, with 1000 bootstrap replicates.
of the corresponding PCR products [11]. A 50 μl-volume of BALKV positive pools was inoculated onto Vero cells for attempting virus isolation [7, 9].

Results

In 2014 a total of 270 and 53 sand flies were collected from Kosovo and Serbia, respectively. In 2015, 453, 386, 37, 602 and 29 sand flies were trapped in Croatia, BH, Montenegro, RoM and Serbia, respectively (Table 1). BALKV RNA was detected in 4 pools from Croatia (3 collected in Vidonje [C41, C50, C51 at 42.98244N, 17.64294E (240 m)], 1 in Duba [C13 at 42.60032N, 18.33946E (475 m)]) and in 1 pool from BH in Sovici (B1 at 43.408240N, 17.329175E, 283 m) (Table 1, Fig. 1).

Although not quantitative, the low C_t values observed with the polymerase gene (C_t range 19.9–24.4) and the nucleoprotein gene (C_t range 19.8–32.8) SYBR Green real-time RT-PCR was indicative of high viral load in the positive pools. Phylogeny was reconstructed by using sequences in the S and L RNA segments that were 572 nt (Fig. 2) and 525 nt long, respectively (Fig. 3). Identical groupings were observed using both markers. BALKV formed a homogenous cluster with common ancestor supported by high bootstrap value. BALKV was included in the subgroup I of the Sandfly fever Naples species complex together with SFNV, Tehran, Zerdali and Fermo viruses.

For pool B1, failure to obtain a positive PCR with Nphlebo primers led us to sequence the 136 bp SYBR Green RT-qPCR product for genetic and phylogenetic analysis. Study of the diversity between BALKV sequences from Albania, Croatia and BH showed that (i) Albanian sequences were the most divergent (9–11% [NP]) from the others, and (ii) that Croatian and BH sequences were grouped (0.9–5.4% [NP]; 0.7–5% [L]) (GenBank: KY662227–KY662287).

Identification of the sand fly species contained in the BALKV-positive pools detected Phlebotomus neglectus sequences in all five pools; P. neglectus was the unique species in four pools, whereas P. tobbi DNA was present in 1 pool from Croatia (Table 2).

Discussion

The Balkan Peninsula is the region where sand fly fever was first described at the end of the nineteenth century in BH [15, 16]. Subsequent studies provided direct and indirect evidence for the presence of viruses belonging to the SFNV in BH [17–21]. In Croatia, antibodies against SFNV were found in human populations, with highest rates (up to 53.9%) observed on islands and in coastal regions [18, 22–27]. BALKV belongs to the Sandfly fever Naples species complex where it is most closely to Fermo, SFNV YU 8–76, Zerdali and Tehran viruses isolated in Italy, Serbia, Turkey and Iran which are
grouped in the subgroup I [6, 9, 19, 28] (Figs. 2, 3). BALKV was first detected from two sand fly pools from Albania, Kruje region [11]. Here, we demonstrated that BALKV has a much larger circulation area that seems to be confined to the Adriatic coast of the Balkan Peninsula. This merits further confirmation through similar studies conducted north and south of the current study area (Fig. 1).

To our knowledge, BALKV is the first phlebovirus to be genetically identified in BH. Assuming that each positive pool contained one infected sand fly only, the sand fly infection rate of BALKV is 0.26% in BH and 0.27% in Croatia; which is higher than Zerdali virus (0.035%) and similar to Fermo virus (0.20%) [6, 9]. Identification of the species content of pools using *cox1* and *cytb* showed that *P. neglectus* is the only species to be found in all BALKV RNA positive pools; indicating that this species might be the vector of BALKV. Interestingly, *P. neglectus* belongs to subgenus *Larroussius*, similar to *P. tobbi* which seems to be a typical vector for Zerdali virus, another member of the *Sandfly fever Naples species* [9]. Together, these data support the hypothesis that *Larroussius* sand flies are typical vectors of the members of this virus group.

**Conclusions**

We report here (i) the first direct evidence that Balkan virus initially described in Coastal Albania has a much wider dissemination area than originally believed, (ii) two real-time RT-PCR assays that may be useful for further screening of patients presenting with fever of unknown origin that may be caused by Balkan virus infection, (iii) entomologic results suggesting that Balkan virus is likely transmitted by *Phlebotomus neglectus*, and possibly other sand fly species of the subgenus *Larroussius*. So far, BALKV has been detected only in sand flies. Whether BALKV can cause disease in humans is unknown and remains to be investigated.

**Abbreviations**

BALKV: Balkan virus; BH: Bosnia and Herzegovina; L: Large RNA segment; NGS: Next generation sequencing; NP: Nucleoprotein; RoM: Republic of Macedonia; RT-PCR: Reverse transcriptase polymerase chain reaction; S: Small RNA segment

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**Availability of data and materials**

Sequences generated in this study are available in the GenBank database under the accession numbers KY662276–KY662287.

**Authors’ contributions**

NA participated in field work, performed PCR and sequencing, and wrote the original MS; BA, VI, FM, JO, JS, DP, DB and PV organized and participated in the field work; VD set-up of PCR-based NGS identification of sand flies; SV participated to field work, performed PCR-based NGS identification; RNC analysed results, and coordinated the lab work. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

Traps were placed near animals with the consent of the owners.

### Table 2

Details of the Balkan virus positive pools with sandfly species identification using cytochrome b and cox1 sequences

| Trapping locality | Pool code | Sand fly species | Gene | Reads | No. of sand flies | Gender | Collection date | Altitude (m) |
|-------------------|-----------|------------------|------|-------|------------------|--------|----------------|--------------|
| Bosnia and Herzegovina | Sovici | P. neglectus | cytb | 1427 | 27 | male | 06/07/2015 | 283 |
| | | | cox1 | 4257 | | | | |
| Croatia | Duba | P. tobbi | cytb | 1211 | 20 | male | 13/07/2015 | 475 |
| | | | cox1 | 546 | | | | |
| | | P. neglectus | cytb | 967 | | | | |
| | | | cox1 | 7351 | | | | |
| | Vidonje | P. neglectus | cytb | 950 | 20 | female | 16/07/2015 | 240 |
| | | | cox1 | 8182 | | | | |
| | Vidonje | P. neglectus | cytb | 1834 | 20 | female (bf) | 16/07/2015 | 240 |
| | | | cox1 | 5302 | | | | |
| | Vidonje | P. neglectus | cytb | 3143 | 20 | female | 16/07/2015 | 240 |
| | | | cox1 | 22,867 | | | |
Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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