***MECHANICAL TRANSMISSION OF LUMPY SKIN DISEASE VIRUS BY STOMOXYS SPP (STOMOXYS CALSITRANS, STOMOXYS SITIENS, STOMOXYS INDICA), DIPTERA: MUSCIDAE.***

***Arman Issimov\*1,Lespek Kutumbetov2, Mukhit B. Orynbayev2, Berik Khairullin2, Balzhan Myrzakhmetova2, Kulyaisan Sultankulova2, Peter J. White1.***

***1. Sydney School of Veterinary Science, Faculty of Science, University of Sydney***

***2. RGE “Research Institute for Biological Safety Problems” Committee of Science, The Ministry of Education and Science of the Republic of Kazakhstan***

***\* Corresponding Author***

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**Summary**

Lumpy skin disease (LSD) is an emerging disease in Kazakhstan and currently the means of transmission is uncertain. In the current study, mechanical transmission of Lumpy Skin Disease Virus (LSDV) by *Stomoxys* species from infected to naive animals has been demonstrated under laboratory conditions. Flies partially fed on LSDV infected cattle were placed onto recipient animals within a 1-hour time period to complete their feeding process. In addition to this, virus was isolated from all three Stomoxys species immediately and 6h post feeding on LSDV infected animal while virus DNA was detectable up to 48h post-feeding by PCR.

Samples collected for PCR from recipient animals tested positive in 5 out of 6 cases, while the virus was isolated from 4 of 6 animals. The clinical signs exhibited by recipient animals were mostly moderate in nature with only one severe case. To our knowledge, this is the first time that transmission of LSDV by three *Stomoxys* species has been demonstrated and their role as mechanical vectors of LSDV is indicated.

**Introduction**

Lumpy skin disease virus (LSDV) belonging to genera Capriposviruses, Poxviridae family, is a highly contagious infectious disease of cattle mainly characterised by multiple skin lesions, fever, enlargement of superficial lymph nodes, profuse salivation, lacrimation and nasal discharge as well as oedema and swelling of the joints (Davies, 1991). The disease was recorded in Zambia for the first time in 1929. Subsequently, LSDV has become endemic throughout the African continent and Near East and continues to spread north posing a threat to Europe and Central Asian regions. Recently, an LSD outbreak has been documented in Greece (Tasioudi et al., 2016) and within a month the disease was clinically confirmed and reported in Azerbaijan, the Russian Federation and the Republic of Kazakhstan (OIE, 2016; Sprygin et al., 2018; Zeynalova, Asadov, Guliyev, Vatani, & Aliyev, 2016).

 The World Organization for Animal Health (OIE) classified LSD as a notifiable disease due to its significant economic impact (E. S. M. Tuppurainen & Oura, 2012). In addition, it has a detrimental effect on the development of animal production. Economic damage results from a sharp decline in milk yield, milk quality and hide damage, body weight reduction, abortion, infertility and in some cases death of the animal (Babiuk, Bowden, Boyle, Wallace, & Kitching, 2008). Morbidity rates may vary significantly during LSD outbreaks and reach up to 100% whereas the mortality rate is usually low (less than 5%) reaching 20% on some occasions (OIE, 2010; E. S. M. Tuppurainen & Oura, 2012; Woods, 1988).

It is thought that a variety of blood-feeding insects may play a significant role in LSDV transmission by acting as mechanical vectors. This assumption is based on the seasonality of outbreaks of LSD, occuring during hot and wet summer seasons (Weiss, 1968). Studies on using the basic reproduction numbers to evaluate the risk of LSDV transmission by blood feeding insects suggests that *S. calcitrans* and *Aedes aegypti* are the most competent vectors of LSD whereas *Culicoides nubeculosus, Anopheles stephensi, and Culex quinquefasciatus* are unlikely to be competent at transmitting LSDV (S. Gubbins, 2019). Previous studies have reported the capability of stable flies (*Stomoxys calcitrans*) to transmit sheep poxand goat pox virus(Kitching & Mellor, 1986; Sohier et al., 2019) and LSDV by mosquitoes (Aedes aegypti) (Chihota, Rennie, Kitching, & Mellor, 2001). In more recent studies, stable flies (*Stomoxys calcitrans)* have been shown to transmit LSDV from experimentally infected animals to naïve cattle (Sohier et al., 2019). It is anecdotally believed that stable flies are most likely involved in LSD epidemiology since the virus had been recovered from individual flies caught on LSDV infected animals in the field (Weiss, 1968). The aim of this study was to determine the vector capability of three *Stomoxys* spp in the transmission of LSD virus.

**Materials and methods**

*Animal Ethics*

Animals were maintained in the insect-free facility, with drinking water and hay provided ad libitium. The experiment was conducted in accordance with national and international laws based on the European Convention for the Protection of Vertebrate Animals used for Experimental and Other Scientific Purposes guidelines (Ausems, 1986). The protocol was approved by the Committee on the Ethics of Animal Experiments of the Research Institute for Biological Safety Problems of the Science Committee of Ministry of Education and Science of the Republic of Kazakhstan. (permit number: 0904/126).

*Experimental animals*

Eight Kazakh White-headed cattle approximately 12-months-old were purchased from a local farm (Korday District) where LSDV has never been recorded, and vaccination was not practiced. Prior to the experiment, animals were kept under quarantine conditions in the insect secured containment laboratory of the Research Institute for Biological Safety Issues (RIBSP) for 15 days. During the quarantine period, blood samples were tested for the presence of antibodies against capripox viruses using the SNT serum neutralization test (Beard et al., 2010).

After veterinarian examination had determined that the animals were healthy, they were divided into 4 groups and placed in isolated rooms of Vivarium No. 202 of the RIBSP.

*Experimental design*

Two donor animals (D1 and D2) were artificially infected with a virulent strain of LSDV and served as a source of infection for *Stomoxys Calcitrans, Stomoxys sitiens* and *Stomoxys indica*.

The titre of the virus inoculum was determined as 106 TCID50/mL. The virus suspension was inoculated in a volume of 2 ml at two sites on each side of the body, (neck and flank) and administered subcutaneously. Personnel participating in this experiment wore overalls (DuPont, Tyvek) and respirators (class N95). Plastic baits containing flies were disinfested externally using Lysoformin 3000 prior to moving between animal units. Five-day old unfed batches of Stomoxys calcitrans Stomoxys sitiens and Stomoxys indica containing approximately 100 flies each were allowed to feed to repletion on the donor animals 5 days following the development of LSDV lesions. Engorged flies were then tested for LSDV presence immediately after engorgement and at different time intervals; 6, 24 and 48 hours post-feeding. They were maintained in the environmental chamber at 250C and 80% humidity, and heparinized bovine blood soaked into cotton pads were offered.

The second batch containing approximately 400 flies each was placed onto an animal with LSDV by using a glass or plastic cylindrical bait containing 40 flies each, closed with a reticular membrane at both ends. Flies on lesions were fed for not more than one minute to prevent the complete engorgement of flies with blood, and then they were moved to a healthy animal to complete feeding. The feeding completion time was determined by the cessation of the flies’ attacking the skin. **Recipient animals were observed daily for clinical manifestation of LSDV.** Blood samples for PCR and VI were taken at intervals of 3,5,7,9,11,13,15,17,19,21,23 and 25 days post exposure. Serum samples were collected on day 7, 14, 21 and 28 post exposure.

In addition, *Stomoxys* spp were examined for the presence of LSDV in the fly proboscis following feeding on donor animals. To do so, 100 adult (*S. Calcitrans*), 100 (*S. sitiens*) and 150 (*S. indica*) were fed to repletion on lumpy skin disease infected cattle. Following blood-meal engorged flies were anesthetized and kept in wet ice. Flies were then washed three times in PBS and rinsed using distilled water to eliminate surface contamination. Proboscises were removed aseptically using entomological forceps and tested separately according to species. Removed proboscises were homogenized and tested by PCR and VI.

*Virus strain*

The LSDV strain used in this study was isolated from an infected cow during an outbreak of LSD in Atyrau, Kazakhstan in 2016. The virus was passaged five times on primary lamb testis (LT) and three times in vitro using calves. According to Plowright and Ferris (1958) pre-pubertal lambs were used to prepare primary LT cell cultures. Cell cultures showed 90% cytopathic effects (CPE), were freeze–thawed three times, and then centrifuged (2000 g for 20 min) and stored at -80C until needed.

*Virus amplification test*

A PCR assay was performed using the protocol published by E. S. Tuppurainen, Venter, and Coetzer (2005).

For DNA extraction, a QIAamp DNA Kit (QIAGEN, USA) was used according to manufacturer’s instructions. For protein digestion, Proteinase K was pipetted in the amount of 20 µl into microcentrifuge tubes (Eppendorf, Germany) containing 200 µl of homogenized sample. Then, 200 µl of Buffer AL was added to the mixture, vortexed and incubated at 56°C for 10 min. The mixture was then transferred to 2 ml collection tubes and 500 µl of Buffer AW1 was added prior to centrifugation at 13000 rpm (Centrifuge 5417, Eppendorph international, Germany) at room temperature for 3 min. This process was repeated by adding 500 µl of Buffer AW2 in the second cycle. DNA containing columns were then transferred into 1.5 ml tubes and eluted with 200 µl of Buffer AE by centrifugation (8000 rpm) at room temperature for 1 min and extracted DNA stored at - 70°C until assayed.

For PCR assay, to produce 192 bp of amplified nucleotides reactions the forward 5’-TCC-GAG-CTC-TTT-CCT-GAT-TTT-TCT-TAC-TAT-3’ and reverse 5’-TAT-GGT-ACC-TAA-ATT-ATA-TAC-GTA-AAT-AAC-3’ primers were used (Ireland & Binepal, 1998). The conditions for DNA amplification in a Thermal Cycler (Eppendorf Mastercycler) were as follows: 95°C for 2 min, 95°C for 45 s, 50°C for 50 s, 72°C for 1 min (34 cycles) and 72°C for 2 min. PCR products obtained were loaded in 1.5% agarose-gel electrophoresis and the results visualized using Bio-imaging systems MiniBIS Pro (Israel).

*Serum neutralization test*

The serum neutralization test was carried out utilizing 96-well microtitre plates following protocols of the BSL-3 laboratory of the RIBSP based on OIE (2018) manuals. Blood samples were collected from experimental animals in vacutainers containing clot activator on days 7, 14, 21, and 28 following infected insect exposure. Serum was separated from clotted blood samples by centrifugation (1500 g for 10 min), diluted 1/5 in Eagle’s/HEPES (N2-hydroxyethylpiperazine, N-2-ethanesulphonic acid) and heat inactivated at 56°C for 30 min. Next, to each well of 96-well microtitre plates, 50 µl of diluted serum aliquots were added. Tenfold serial dilution of LSDV from 10-1 to 10-8 was diluted in Eagle’s/HEPES in bijoux bottles. Fifty µl of the virus dilution was introduced to each well in triplicates per dilution, incubated at 37°C for 1 hour before adding LT cells to each well of the plates at a concentration of 105 cells/ml. The monolayers were examined daily for specific CPE and by inverted microscopy on day 9 post inoculation and end-points calculated according to Karber (1931). To determine virus neutralization index, the difference between the titre of virus in the test serum and negative serum from the same animal was taken. A neutralization index showing ≥1.5 was considered positive.

*Virus isolation*

Virus isolation was conducted according to Standard operational procedures of the BSL-3 laboratory of the RIBSP, based on OIE (2018) manuals. Briefly, 1 ml buffy coat or supernatant were administered on to lamb testes cells in 25 cm2 cell culture flasks and allowed to incubate at 37°C for 1 hour. Following incubation culture media was rinsed with PBS and overlaid with Glasgow's Minimal Essential Medium containing 0.1% penicillin, 0.2% gentamycin and 2% fetal calf serum. The cell monolayer was examined daily for characteristic CPE. In the case no CPE was observed, the cell culture was freeze–thawed three times and second or third blind passages were carried out. The utilized culture media were kept at -80°C until required. Cell culture flasks showing CPE were tested with gel-based PCR to confirm that CPE was induced by LSDV

**Results**

**As shown in Table 1, PCR and virus isolation tests were positive for all three *Stomoxys* spp collected immediately and 6 hours post-feeding. No virus was isolated from species from 24 and 48 hours post-feeding, whereas viral nucleic acid was detected up to 48h post-feeding by PCR. Further tests failed to detect LSDV in all three species between days 3 and 10 post-feeding.**

**Table 1. PCR and virus isolation results of *Stomoxys spp* at different time intervals following feeding on LSDV infected donor animals.**

|  |  |  |  |
| --- | --- | --- | --- |
| **Time/day post feed**  | *Stomoxys Calcitrans* | *Stomoxys sitiens* | *Stomoxys indica* |
| PCR(no positive/no tested) | Virus isolation(TCID50/mL), mean | PCR(no positive/no tested) | Virus isolation(TCID50/mL), mean | PCR(no positive/no tested) | Virus isolation(TCID50/mL), mean |
| **0h** | **12/15** | **3.5** | **13/15** | **3.1** | **11/15** | **3.3** |
| **6h** | **8/15** | **1.8** | **11/15** | **2.0** | **8/15** | **1.9** |
| **1** | **5/15** | **-** | **4/15** | **-** | **3/15** | **-** |
| **2** | **2/15** | **-** | **3/15** | **-** | **2/15** | **-** |
| **4** | **0/15** | **-** | **0/15** | **-** | **0/15** | **-** |
| **7** | **0/15** | **-** | **0/15** | **-** | **0/15** | **-** |
| **14** | **0/15** | **-** | **0/15** | **-** | **0/15** | **-** |

**Table 2. Clinical manifestations of LSD for recipient animals infected through transmission of LSDV by *Stomoxys spp***

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| **Animal ID** | **Route of transmission** | **Numbers and species of flies fed** | **Body temperature** | **Lesions** | **Enlargement of lymph nodes** | **Additional clinical signs** |
| **R1** | **Mechanical** | 210 *Stomoxys Calcitrans* | Up to 40.1°C (on day 17 post exposure) | Small skin nodules developed on fly feeding side (Day 10 post exposure) | **Mild enlargement:**prescapular and precrural lymph nodes**(Day 11** post exposure**)** | Nasal and ocular discharges (**Day 11** post exposure) |
| **R2** | **Mechanical** | 210 *Stomoxys Calcitrans* | Up to 39.9°C (on day 6 post exposure) | **Small swellings at feeding sites (Day 14** post exposure**)** | **Mild enlargement:**prescapular and precrural lymph nodes**(Day 9** post exposure**)** | Nasal and ocular discharges (on day 6 post exposure) |
| **R3** | **Mechanical** | 200 *Stomoxys sitiens* | Up to 39.7℃ (on day 13 post exposure) | Large swellings at feeding sites (Day 7 post exposure) | **Mild enlargement:**prescapular lymph node**(Day 13** post exposure**)** | Edema of the hind limbs (Day 13 post exposure) |
| **R4** | **Mechanical** | 200 *Stomoxys sitiens* | Up to 39.5℃ (on day 9 post exposure) | **Skin lesions** at feeding sites (Day 14 post exposure) | **Mild enlargement:**prescapular and precrural lymph nodes**(Day 14** post exposure**)** | Nasal and oral discharges, rhinitis (Day 11 post exposure) |
| **R5** | **Mechanical** | **230** *Stomoxys indica* | Up to 39.8℃ (on day 12 post exposure) | **Small skin lesions** at feeding sites (Day 10 post exposure) | **Mild enlargement:**prescapular lymph node**(Day 11** post exposure**)** | **-** |
| **R6** | **Mechanical** | **230** *Stomoxys indica* | **-** | Small-sized swellings on the feeding site (Day 2 post exposure) | **-** | **-** |

Demonstration *of LSDV transmission by Stomoxys Calcitrans*

Recipient animals R1 and R2 showed characteristic clinical signs of LSD similar to those described by E. S. M. Tuppurainen and Oura (2012) and Coetzer (2004). On day 8 post infection the body temperature for R1 raised to 39.6°C and reached 40.1°C on day 17 (Table2). In addition to this prescapular and precrural lymph node enlargementand excessive nasal and ocular discharge was observed. Multiple small skin nodules developed on the fly feeding side of R2 and caused pain when palpated, indicated by animals twitching the skin, flicking the tail, kicking or stamping. Elevation in body temperature of R2 increased to 39.9 °C on day 6 post exposure and remained mostly beyond 39.0℃ up to day 28 post exposure (Table2).

Blood samples collected for PCR tested positive for R1 on days 7, 13, 17 and 21 post exposure. Samples for R2 were positive on days 11, 13 and 15 post exposure. Virus was isolated from blood samples of animal R1 on day 15 whereas samples from R2 exhibited CPE on days 9, 11,14, 21 and 28 post feeding (Table3). The index for virus neutralization was ≥log1.5.

*Demonstration of LSDV transmission by Stomoxys sitiens*

On day 6 post infection the body temperature for R3 was 39.3℃ and elevated up to 39.7 ℃ on day 13 post exposure (Table2). The body temperature for R4 was 39.5℃ on day 9 post exposure and remained chiefly above 39.0℃ up to day 17 post exposure (Table2). Recipient animals R3 and R4 developed large swellings on the fly feeding sites on day 7 post exposure (Table2). In addition, the enlargement of prescapular lymph nodes were detected when palpated. On day 14 post exposure R3 developed edema of the hind limbs extending from metatarsus to tarsal joints (Table2). R4 manifested nasal and oral discharge as well as rhinitis on day 11 post exposure. Virus DNA was detected from blood samples for R3 by conventional PCR on days 7, 9, 13 and 17 post exposure while R4 tested positive on days 11 and 15 post exposure. Virus isolation was conducted using a buffer coat, and CPE in the cell culture was seen on the second blind passage for blood samples collected on day 13 post exposure (Table3). Virus neutralization index demonstrated an index > log1.5 for both recipient animals on days 14 and 21 post exposure.

*Demonstration of LSDV transmission by Stomoxys indica*

Animal R5 manifested a mild elevation in body temperature of 39.1℃ on day 10 post exposure and 39.8 on day 12 post exposure (Table2). Mild enlargement of prescapular lymph nodes was also detected. Skin lesions, approximately 3 cm in diameter, was observed on feeding sites of flies on day 10 post exposure and disappeared on day 11 post exposure (Table2). Blood samples tested positive by PCR on day 9 and 11 post exposure. No CPE was seen in the blood samples collected. Seroconversion was detected on days 14 and 21 post exposure (Table3), showing an index > log1.5.

On the other hand, the recipient animal R6 demonstrated no clinical signs of LSD apart from small-sized swellings on the insect feeding site on day 2 post exposure (Table2), which gradually disappeared by day 4 post exposure. All samples tested were negative.

*Detection of LSDV in the proboscis of stable flies fed on LSDV infected cattle.*

LSDV DNA was detected using gel-agarose PCR in 10 out of 12 samples collected from *S. Calcitrans,* in 9 out of 12 samples for *S. sitiens and* in six of the 12 proboscis samples for *S. indica* respectively. The proboscis samples from *S. Calcitrans* demonstrated CPE in 14 of 20 (70%) flasks. The samples for *S. sitiens* demonstrated CPE in twelve out of eighteen (66%) flasks while the sample for *S. indica* demonstrated CPE in 7 of 22 (32%) flasks.

**Table 3.** Test results obtained from animals R1, R2, R3, R4, R5 and R6.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Animal ID | Transmission | Gel-based PCR | SNT | Virus isolation | Results |
| **R1** | **Mechanical** | Pos (days 7, 13, 17, 21 pe) | Pos(days 14, 21, 28 pe | Pos (day 15 pe) | Infected |
| **R2** | **Mechanical** | Pos (days 11, 13, 15pe) | Pos(days 14, 21, 28 pe | Pos (days 9, 11 pe) | Infected |
| **R3** | **Mechanical** | Pos (days 7, 9, 13, 17 pe) | Pos(days 14, 21 pe | Pos, 2nd passage(day 13 pe),  | Infected |
| **R4** | **Mechanical** | Pos(days 11, 15 pe) | Pos(days 14, 21 pe | Pos, 2nd passage(day 13 pe) | Infected |
| **R5** | **Mechanical** | Pos (day 9, 11 pe) | Pos(days 14, 21 pe | - | Infected |
| **R6** | **Mechanical** | - | - | - | Not Infected |

 **(**pe) – post exposure

**Discussion**

There are several studies investigating the role of stable flies as a mechanical vector for the transmission of disease (Foil, Meek, Adams, & Issel, 1983; Mellor, Kitching, & Wilkinson, 1987; Turell & Knudson, 1987). It has been assumed that the transmission of LSDV by blood-feeding insects provided a short-term mechanism of transmission (Chihota, Renniet, Kitchingt, & Mellort, 2003). The outcomes obtained in this study indicate a longer duration of possible transmission. In this study LSDV was shown to survive inside infected stable flies for at least 6 hours without a noticeable loss in titre. In other words, the virus could be localized within the insect vector, where stable flies play an intermediary role for virus transmission. In addition to that, harbored LSDV is protected from detrimental ambient conditions and this implies a more sophisticated means of transportation than just "occasional contact".

In the present study, the mechanical transmission of LSDV using *Stomoxys* species (*Stomoxys Calcitrans, Stomoxys sitiens* and *Stomoxys indica*) from infected to susceptible animals has been demonstrated under laboratory conditions. Five out of six cattle exposed to infected flies refeeding manifested mild to generalized LSD including viremia and fever. Animal R6 did not develop any clinical signs of LSD, apart from small-sized swellings on the insect feeding site on day 2 post exposure. This case could be potentially explained by the differences in the immune status of animals at the moment of exposure. Although the severity of clinical signs exhibited varies in animals, this aligns with records that less than 50% of the cattle infected experimentally or naturally with LSDV will manifest inapparent or generalized disease (Carn & Kitching, 1995; Prozesky & Barnard, 1982). All three species of flies demonstrated the capability to ingest and harbor virus particles and were able to transmit virus within a 1-hour time interval between the feeding processes. Moreover, LSDV was recovered from fly mouthparts within the same period with a virus titre 10-4TCID50/ml. Furthermore, LSDV can survive in *Stomoxys* species at least 6h following feeding on an infected animal, whereas viral nucleic acid was detected up to 48h post exposure.

In a previous study, stable flies failed to transmit LSDV from infected to naïve animals 24-hours after feeding on an infected animal. This implies that virus survival decreases over time in the flies’ gut environment. However, in the transmission experiments with shorter transmission periods, *S. calcitrans* demonstrated a vector competence of transmitting sheep pox and goat pox viruses (Webb, 1990) and LSDV (Sohier et al., 2019). Stable flies are known to be intrusive feeders, and due to aggressive attacks and painful bites, host animals take defensive actions resulting in interrupted feeding requiring flies to seek for a new host. Such feeding behavior thus requires taking 3 to 5 interrupted feeding sessions to achieve full repletion (Schofield & Torr, 2002). Blood-meal regurgitation by *S. calcitrans* prior taking another blood-meal has been experimentally recorded by Butler, Kloft, Dubose, and Kloft (1977). This implies that the mouthparts can be contaminated with the virus regurgitated during the second bloodmeal, which in turn will increase interrupted transmission rates of LSDV by stable flies.

In a recent study, the mechanical transmission of LSDV by hard ticks has been demonstrated (Lubinga et al., 2015; E. S. M. Tuppurainen et al., 2013). Moreover, they were capable of vertical transmission of LSDV (Lubinga, Tuppurainen, Coetzer, Stoltsz, & Venter, 2014; Eeva S. M. Tuppurainen et al., 2013). However, the mechanical role of the hard ticks in the mass dissemination of LSDV within a herd is highly restricted since, in most cases, the life cycle of the tick occurs on a single host (Koshy, Rajavelu, & Lalitha, 1982; Vatsya, Banerjee, Yadav, & Kumar, 2006). Taking into account the switch of hosts during stages and generally one meal per stage, their role as even putative vectors diminishes. After repletion, the female ticks drop to the ground, oviposite and die whereas male ticks remain on host animals for further feeding and mating with newly attached females (Cupp, 1991).

Under field conditions, a large number of flies that feed on the erupted lesions must carry or at least become contaminated with pathogen during the feeding process. Thus, it is highly likely that a single infected animal introduced into a herd will serve as virus source for a large number of resident stable flies. Given the fact that cattle can be easily infected with LSDV using an intradermal method of inoculation leads to the suggestion of insect vector involvement since insects introduce virus into their hosts in a similar manner. This factor will inevitably result in the fulminant spread of the disease within a herd.

It is reported that under experimental conditions stable flies range over large areas, with male flies travelling up to 28.9 km and females 21.9 km (Bailey, Whitfield, & Smittle, 1973). Moreover, the wind has a direct impact on insect distribution (Yeruham et al., 1995). In other studies, mathematical modelling used to calculate vector born dispersal of LSDV between herds located in close proximity revealed that most transmission is highly likely to occur over short distances, less than 5 kilometres (Simon Gubbins et al., 2018). Such a significant coverage range and vector capability of stable flies to carry pathogen may lead to LSDV escape from the initial outbreak foci and rapid dissemination over neighbouring farms. Given this fact, insect control programs must be considered during outbreaks of LSD as well as ring vaccination program within the flight range of hematophagous flies.

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**Conflict of Interest**

There is no conflict of interest.

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