German *Culex pipiens* biotype *molestus* and *Culex torrentium* are vector-competent for Usutu virus

Cora M. Holicki¹, Dorothee E. Scheuch¹, Ute Ziegler¹, Julia Lettow²,³, Helge Kampen², Doreen Werner⁴ and Martin H. Groschup¹*

Abstract

**Background:** Usutu virus (USUV) is a rapidly spreading zoonotic arbovirus (arthropod-borne virus) and a considerable threat to the global avifauna and in isolated cases to human health. It is maintained in an enzootic cycle involving ornithophilic mosquitoes as vectors and birds as reservoir hosts. Despite massive die-offs in wild bird populations and the detection of severe neurological symptoms in infected humans, little is known about which mosquito species are involved in the propagation of USUV.

**Methods:** In the present study, the vector competence of a German (i.e. “Central European”) and a Serbian (i.e. “Southern European”) *Culex pipiens* biotype *molestus* laboratory colony was experimentally evaluated. For comparative purposes, *Culex torrentium*, a frequent species in Northern Europe, and *Aedes aegypti*, a primarily tropical species, were also tested. Adult female mosquitoes were exposed to bovine blood spiked with USUV Africa 2 and subsequently incubated at 25 °C. After 2 to 3 weeks saliva was collected from each individual mosquito to assess the ability of a mosquito species to transmit USUV.

**Results:** *Culex pipiens* biotype *molestus* mosquitoes originating from Germany and the Republic of Serbia and *Cx. torrentium* mosquitoes from Germany proved competent for USUV, as indicated by harboring viable virus in their saliva 21 days post infection. By contrast, *Ae. aegypti* mosquitoes were relatively refractory to an USUV infection, exhibiting low infection rates and lacking virus in their saliva.

**Conclusions:** Consistent with the high prevalences and abundances of *Cx. pipiens* biotype *molestus* and *Cx. torrentium* in Central and Northern Europe, these two species have most likely played a historic role in the spread, maintenance, and introduction of USUV into Germany. Identification of the key USUV vectors enables the establishment and implementation of rigor entomological surveillance programs and the development of effective, evidence-based vector control interventions.

**Keywords:** *Culex pipiens* biotype *molestus*, *Culex torrentium*, *Aedes aegypti*, Vector competence, Usutu virus
cause encephalitis or meningoencephalitis, with a possible lethal outcome in infected humans [3], and dengue and chikungunya viruses can lead to hemorrhagic and thereby sometimes fatal illnesses [4, 5].

Usutu virus (USUV; *Flaviviridae, Flavivirus*) is currently considered an emerging zoonotic arbovirus in Europe with an increase in registered cases and observed disease severity in both humans and animals. This flavivirus belongs to the Japanese encephalitis virus serocomplex with birds, mainly Eurasian blackbirds (*Turdus merula*), Eurasian magpies (*Pica pica*), house sparrows (*Passer domesticus*), and great grey owls (*Strix nebulosa*), as amplifying hosts [6, 7]. Susceptible avian species show mild to severe clinical signs including encephalitis, myocardial degeneration, and necrosis of the liver and spleen [8, 9]. Although USUV primarily causes a self-limiting, febrile illness with seroconversion in incidental hosts such as humans and horses, a neuroinvasive disease in humans, primarily in immunocompromised patients, should receive more attention. Recent studies even indicate that human USUV infections are not as sporadic as previously presumed [6, 10–12], with so far 49 reported acute infections over the years (1981–2018) in the Central African Republic, Burkina Faso, Italy, Croatia, Germany, France, and Austria [6].

USUV was first detected by McIntosh in 1959 in a *Culex neavei* mosquito (originally described as *Cx. univittatus*) near the Usutu River, Natal, South Africa [13]. Phylogenetic analyses suggest a minimum of three unique USUV dispersions from Africa to Europe (in the 1950s, 1980s, and 1990s). Presumably, the virus arrived along the eastern Atlantic and Black Sea/Mediterranean routes [6] with migratory birds as long-distance dispersal vehicles [14]. The first recorded outbreak of USUV in Europe occurred in 2001 in Austria among Eurasian blackbirds and great grey owls [15]. Thereupon, USUV ribonucleic acid (RNA) was retrospectively isolated from tissue samples of Eurasian blackbirds from 1996 in Italy [16]. Since then, USUV continued to colonize new ecological niches throughout Europe. In Germany, the first documented isolation of USUV was in 2010 from a pool of *Cx. pipiens* biotype *pipiens* mosquitoes collected in the city of Weinheim in the southwest of the country [17]. In the following 2 years, several epidemics in the Upper Rhine Valley resulted in substantial mortalities among the wild avifauna, primarily of the Eurasian blackbird [18]. To date, USUV has spread to the north and east of Germany, covering all German federal states, with major epidemics in 2016 and 2018 [19].

USUV is currently classified into eight genetically distinct lineages [20] depending on their geographical origin of isolation [14], of which five circulate in Germany: “Africa 2,” “Africa 3,” “Europe 2,” “Europe 3,” and “Europe 5” [8, 18–20]. The Africa 2 clade, for example, originated from South Africa and was isolated in Germany in 2015 from two carcasses of juvenile great grey owls from the Zoological Garden Berlin [8]. Till now, it is uncertain which vectors play an important role in the maintenance and spread of USUV in Germany.

USUV has been isolated from numerous mosquito species yet the primary vectors belong to the genus *Culex* [6]. This genus contains species that already are known to be fundamental in the spread of multiple other arboviruses including Sindbis virus, St. Louis encephalitis virus, Rift Valley fever phlebovirus, and WNV [21]. So far *Cx. neavei* from Senegal [22], *Cx. pipiens* and *Cx. quinquefasciatus* from North America [23], and *Cx. pipiens* from The Netherlands [24] have proved to be highly competent for USUV under laboratory conditions. In Europe, *Cx. pipiens* is highly abundant [25] and can be subdivided into two distinct biotypes: biotype *pipiens* (Linnaeus 1758) and biotype *molestus* (Forskål 1775), which vary greatly in their physiological and behavioral characteristics [26]. In Germany, both biotypes occur in sympathy or as hybrids, and their relevance for the spread of USUV strains must be further clarified [27]. *Culex torrentium*, another common species of temperate regions (e.g. Northern Europe), exhibits similar feeding-preferences as biotype *pipiens*, with both preferring to feed on birds, unlike the mammalophilic biotype *molestus* [26]. Primarily due to difficulties in the establishment of *Cx. torrentium* laboratory colonies, research on this species is very limited. Furthermore, there are only a few studies on the role of *Aedes* species such as *Ae. aegypti* or the invasive species *Ae. japonicus* and *Ae. albopictus* in the USUV transmission cycle [23, 28, 29]. To date, both an Italian and a North American *Ae. albopictus* population have been found to be refractory to an USUV infection [23, 28] while an *Ae. japonicus* population from The Netherlands could transmit USUV in its saliva [29].

In this research, vector competence studies were performed with German mosquito populations to better comprehend and predict the spread of enzootic USUV strains throughout Germany and of strains newly introduced into the country. Infection experiments were carried out with German *Cx. pipiens* biotype *molestus* (i.e. a mammalophilic species, possibly transmitting USUV to humans) and *Cx. torrentium* (i.e. an ornithophilic species, possibly essential for sustaining the enzootic transmission cycle between birds) to investigate their susceptibility for the USUV Africa 2 strain, “USUV-Berlin” [8]. For comparison purposes, the vector competence of “Southern European,” Serbian *Cx. pipiens* biotype *molestus*, and Malaysian *Ae. aegypti* colonies were also examined.
Methods

Mosquito collection, rearing, and identification
For the infection experiments, the following laboratory colonies were used: Cx. pipiens biotype molestus from the “Wendland,” Lower Saxony, Germany (established in 2012); Cx. pipiens biotype molestus from Novi Sad, the Republic of Serbia (established in 2012); Ae. aegypti originally from Malaysia (Bayer CropScience, Langenfeld, Germany). All the tested mosquito colonies were reared at 24 °C ± 1 °C with a relative humidity of approximately 60–70% and a 16 h light/8 h dark photocycle. The adult mosquitoes were provided with a 5–6% sugar solution ad libitum and for egg production with bovine or chicken (EDTA or heparin) whole blood, depending on the feeding preference of the species/biotype. Culex torrentium females were produced from egg rafts collected in the field from different populations near Berlin and Bonn, Germany, after species identification of two to three larvae hatching from each raft by means of a real-time polymerase chain reaction (PCR) [27]. To ensure that all tested mosquito populations were free from flaviviruses prior to the experiments, individual non-engorged females per population were examined via an USUV-specific reverse transcription quantitative real-time PCR (RT-qPCR) [17] and a SYBR® Green-based quantitative real-time pan-flavivirus assay [30].

Virus strain and cultivation
An USUV Africa 2 strain was used for all infection experiments. The strain was isolated from the brain of a succumbed great grey owl in Berlin, Germany, in 2015 (GenBank accession no. KU664608) [8]. The virus was passaged twice on confluent monolayers of Vero cells (Collection of Cell Lines in Veterinary Medicine, Friedrich-Loeffler-Institut (FLI), Greifswald-Insel Riems, Germany), and cell culture supernatant was harvested 4 days post infection (dpi). The virus stock was aliquoted and stored in cryovials at −70 °C until further use. For virus cultivation, Vero cells were maintained in minimum essential medium (MEM), supplemented with 2% fetal calf serum (FCS) and 1% antibiotics (penicillin and streptomycin; Merck, St. Louis, MO, USA). Virus was quantified by means of an endpoint dilution assay, and the virus titer was calculated with the Spearman-Karber algorithm [31]. The used stock had a titer of approximately 10^8.1 TCID50/ml per ml.

Oral infection with USUV
Adult mosquitoes (3–14 days old) were sorted into groups of ten individuals under 100% carbon dioxide (CO2) anesthesia 1 day prior to infection. After sugar deprivation for 24–48 h, Culex mosquitoes were fed artificially overnight. They were exposed to cotton swab ends soaked in an infectious blood meal. Aedes mosquitoes were allowed to feed for 1 h on virus-spiked blood offered via a Hemotek PS5 feeder (Hemotek Ltd, Lancashire, UK), using hog gut and parafilm to seal-off the reservoirs. The blood meal consisted of 10% virus stock, 60% bovine EDTA blood, and 30% sugar solution with a stock concentration of 5–6%. To confirm that the used blood was free from WNV-specific antibodies, serum samples from the cows were examined with the ID Screen® WN competition enzyme-linked immunosorbent assay (ELISA) (IDVet, Grabels, France). After the preparation of the blood meal, a remainder of it was titrated on Vero cells to calculate the exact virus titer in TCID50/ml, which varied between the experiments. In two of the experiments, the virus titer of the blood meal had dropped by about two logs for unknown reasons. The mosquitoes were, therefore, split into two groups, one was exposed to a high virus titer (10^7.4 TCID50/ml) and the other to a low virus titer (10^5.1 TCID50/ml).

Engorged mosquitoes were transferred under 100% CO2 sedation into chambers and kept in an incubator (climate test cabinet MKKL, Flohr Instruments, Nieuwegein, The Netherlands; dehumidifier MG50, Munters, Bedfordshire, UK) under 25 °C ± 1 °C and a relative humidity of 80–85% and offered cotton pads soaked with 5–6% sugar solution ad libitum. A minimum of one engorged female per species/population was frozen at −70 °C to confirm virus ingestion and as a baseline for virus replication in the mosquitoes.

Forced salivation assay
By means of forced salivation, saliva was collected after an incubation period of both 14 and 16 (14/16) or 21 days, respectively. After immobilization of the mosquitoes by removing their legs and wings under 100% CO2 anesthesia, their proboscises were inserted into cut 10-µl filter tips filled with 10 µl of phosphate-buffered saline (PBS). Saliva was collected for half an hour, and the tip content was subsequently transferred into a 1.5-ml tube containing another 10 µl of PBS [32]. Directly thereafter, the mosquito saliva samples were inoculated onto a 96-well plate cell monolayer of Vero cells (i.e. 20 µl of each saliva solution was diluted in 150 µl of MEM + 2% FCS + 1% antibiotics). After 7 days, the cells were examined under the light microscope, fixed with 7.5% neutral buffered formalin (Carl Roth, Karlsruhe, Germany), and stained with crystal violet (Carl Roth). If the saliva appeared to contain viable and replicable virus after the 7 days (i.e. represented by a distinct cytopathogenic effect in the Vero cells), 140 µl of the cell culture supernatant was collected before fixing, and the RNA was extracted using the QIAamp Viral RNA Mini Kit (QIAGEN, Hilden, Germany) according to manufacturer’s instructions.
RNA extracts, eluted in 50 µl of the elution buffer, were tested with an USUV-specific RT-qPCR assay (5 µl at a time) [17] with a standard curve running in parallel [33]. All RT-qPCR assays were completed with the AgPath-ID One-Step RT-PCR Reagents (ThermoFisher Scientific, Darmstadt, Germany) and the CFX96™ Real-Time PCR Detection System (Bio-Rad Laboratories, Feldkirchen, Germany). To confirm correct species association, mosquitoes which contained USUV in their saliva were retroactively once again identified to species and biotype level. For the sake of completeness this was not only done with the field-derived mosquitoes (i.e. positive Cx. torrentium) but also with the mosquitoes originating from the laboratory colonies. For molecular identification a real-time PCR assay [27] was used for the Cx. pipiens complex and cytochrome c oxidase subunit 1 barcoding [34, 35] for Ae. aegypti.

**Pathogen screening in mosquito bodies and legs plus wings**

After saliva collection, the mosquito bodies (thorax and abdomen) and legs plus wings were stored separately in 2-ml screw cap tubes with two 3-mm steel beads, 560 µl of AVL viral lysis buffer, and carrier RNA (QIAGEN) at −70 °C. For RNA extraction, mosquito bodies and legs plus wings were homogenized for 2 min at 30 Hz (TissueLyser II; QIAGEN). Next, insoluble debris was pelleted by centrifugation for 1 min at 13,000 rpm (5430R centrifuge; Eppendorf, Hamburg, Germany), and RNA was extracted from the supernatant with a BioSprint 96 (QIAGEN) or KingFisher Flex Purification System (ThermoFisher) using the NucleoMag VET kit (MACHEREY-NAGEL, Düren, Germany) and following the manufacturer’s instructions. RNA extracts were eluted in 100 µl elution buffer and stored at −70 °C. RNA solutions (5 µl at a time) were tested for viral RNA, using the same USUV-specific RT-qPCR assay as above [17] with the standard curve for quantification [33] and the CFX96™ Real-Time PCR Detection System (Bio-Rad Laboratories).

**Interpretation of results**

Feeding rate refers to the number of engorged females compared to the overall number of females exposed to the infectious blood meal. Survival rate at a given time point is defined as the number of live mosquitoes out of the total number of engorged females subjected to the experiment (minus the day-0 samples). Mosquito bodies, legs plus wings, and saliva samples were considered USUV-positive if they contained USUV-specific RNA (cycle threshold < 36). A minimum of two technical replicates were completed for each positive sample in the RT-qPCR assay [17]. The infection rate describes the number of infected mosquitoes (i.e. USUV-positive bodies) out of the total number of mosquitoes analyzed. To verify that USUV infection and replication in the respective species were, however, not limited to the midgut and that viral dissemination to secondary tissues took place, legs plus wings were also examined. The dissemination rate is defined as the number of USUV-positive legs plus wings samples among the infected (i.e. USUV-positive bodies) mosquitoes. Nonetheless, a mosquito species was only considered competent for USUV when viral RNA was found in its saliva. Transmission rate is measured as the number of USUV-positive saliva samples out of the number of mosquitoes with a disseminated infection (i.e., USUV-positive bodies and legs plus wings). Transmission efficiency describes the total number of USUV-positive saliva samples out of the total number of mosquitoes analyzed.

**Data analyses**

Statistical analyses and graphical displays were completed with the R version 3.6.0. (26 April 2019) [36] and the additional package “lsmeans” [37]. Generalized binomial regression models (GLM) were used to investigate the effect of mosquito species, the days post infection, and the virus titer in the blood meal including their interactions (explanatory variables) on the feeding, survival, infection, dissemination, and transmission rates and transmission efficiencies (response variables). Least-squares means (LSM) [38] were used for testing linear contrasts among predictions with Tukey’s adjustment for P-values [39]. Results were considered statistically relevant when the P-values (summarized in Additional file 1: Tables S1–S3) were < 0.05. In the case of highly unbalanced mosquito numbers, a Fisher’s exact test with Bonferroni correction was implemented instead.

Individual samples where the extremities were virus-positive but not the bodies were not included in the analyses. Two mosquitoes had USUV-positive bodies and saliva even though USUV-specific RNA quantities in their legs plus wings were close to the detection limit. However, before virus could be secreted with the saliva, the mosquitoes must have developed a disseminated infection. Therefore, the two samples were included when counting USUV-positive legs plus wings samples, and the low viral RNA quantities in the legs plus wings were attributed to errors in the extraction process.

**Results**

**Feeding and survival rates**

Of the 284 German and the 101 Serbian Cx. pipiens biotype molestus, which were offered an infectious blood meal, 72.9% and 100%, respectively, were fully engorged after exposure (Fig. 1). Similarly high was the feeding rate...
in the *Ae. aegyti* with 80.7% of the 197 exposed females. By contrast, of the F0 generation of the field-collected *Cx. torrentium*, only 24.7% of the 89 females took a blood meal, significantly less than of the *Ae. aegyti* and both the *Cx. piriens* biotype *molestus* laboratory colonies from Germany and Serbia (Fisher’s exact test: *P* < 0.001 for all). The few engorged *Cx. torrentium* females, however, showed high survival rates of 94.1% and 100% from 0 to 14/16 dpi and from 14/16 to 21 dpi (Fig. 1; only including the survival rates from 0 to 14/16 dpi), respectively, while survival rates of the laboratory colonies after feeding from the infectious blood meal were lower. However, differences were only significant for groups of mosquitoes surviving from 0 to 14/16 dpi but not from 14/16 to 21 dpi (summarized in Additional file 1: Table S3). For example, from 0 to 14/16 dpi, the survival rate of the German *Cx. piriens* biotype *molestus* was 45.6%, which was significantly lower than for *Ae. aegyti* with 67.1% (GLM and LSM: *df* = infinity, *Z*-ratio = 4.0, *P* = 1.4 × 10−3).

Similarly, the survival rate of the Serbian *Cx. piriens* biotype *molestus* was 27.3%, which was significantly lower than the survival rates of both *Ae. aegyti* (GLM and LSM: *df* = infinity, *Z*-ratio = 6.0, *P* < 0.001) and *Cx. torrentium* (GLM and LSM: *df* = infinity, *Z*-ratio = −3.6, *P* = 9.0 × 10−3).

**Infection, dissemination, and transmission rates**

The infection, dissemination, and transmission rates of the four mosquito infection experiments conducted in this study are summarized in Table 1. The virus titers of the blood meals were distinguished as being either high with 107.4 TCID50/ml or low with 105.1 TCID50/ml. After uptake via the infectious blood meal, USUV was able to establish an infection in all of the tested species. However, only *Cx. piriens* biotype *molestus* and *Cx. torrentium* proved competent for USUV. After oral exposure to a high virus titer (107.4 TCID50/ml) in the blood meals, viral RNA could be detected in the bodies, legs plus wings, and saliva samples of multiple mosquitoes from both the German and Serbian *Cx. piriens* biotype *molestus* colonies. The saliva secreted by six of the German and six of the Serbian *Cx. piriens* biotype *molestus* mosquitoes contained virus particles that successfully infected mammalian Vero cells with substantial virus replication 7 dpi. Significant differences, however, were not observed among the vector competence indices (infection, dissemination, and transmission rates) between either the mosquito species or the collection dates (i.e. 14/16 and 21 dpi). This, however, may be a result of the small sample sizes per species and time point. By contrast, a low virus titer (105.1 TCID50/ml) in the blood meals yielded a very low infection rate of 5.6% (2/36) 14 dpi and 5.3% (1/19) 21 dpi in the German *Cx. piriens* biotype *molestus* with no virus replication in secondary tissue. Nonetheless, the low titer still inflicted one out of a total of eight *Cx. torrentium* specimens 21 dpi, with subsequent virus dissemination and transmission. Extensive virus replication took place with RNA viral loads of 3.9 × 106 per µl of total RNA in the body and 4.7 × 104 per µl of total RNA in the legs plus wings. In comparison, the day-0 *Cx. torrentium* mosquitoes contained on average 3.5 × 104 per µl of total RNA. Furthermore, the virus titer in the blood meals not only affected the infection, dissemination, and transmission rates but also the viral loads in the individual mosquito samples. For example, after oral inoculation with a high titer, 6.3 × 105 virus copies per µl of total RNA were found on average in the bodies and 1.2 × 104 virus copies per µl of total RNA in the legs plus wings. The day-0 *Cx. torrentium* mosquitoes contained on average 3.5 × 104 per µl of total RNA. Furthermore, the virus titer in the blood meals not only affected the infection, dissemination, and transmission rates but also the viral loads in the individual mosquito samples. For example, after oral inoculation with a high titer, 6.3 × 105 virus copies per µl of total RNA were found on average in the bodies and 1.2 × 104 virus copies per µl of total RNA in the legs plus wings. The day-0 *Cx. torrentium* mosquitoes contained on average 3.5 × 104 per µl of total RNA. Furthermore, the virus titer in the blood meals not only affected the infection, dissemination, and transmission rates but also the viral loads in the individual mosquito samples. For example, after oral inoculation with a high titer, 6.3 × 105 virus copies per µl of total RNA were found on average in the bodies and 1.2 × 104 virus copies per µl of total RNA in the legs plus wings.
in a decline in the number of virus copies to $8.7 \times 10^3$ per μl of total RNA in the bodies and no virus copies in the legs plus wings of the German Cx. pipiens biotype molestus 14/16 and 21 dpi. Even when exposed to a high virus titer, Ae. aegypti had an infection rate of 0% 14/16 dpi and only 18.2% 21 dpi, with the detection of virus dissemination in only one mosquito. USUV could not be detected in the saliva of Ae. aegypti mosquitoes.

### Discussion

The goal of this study was to identify German mosquito species that are vector-competent for USUV to establish evidence-based recommendations regarding the implementation of mosquito surveillance and control strategies. The study proved for the first time that Serbian and German Cx. pipiens biotype molestus and German Cx. torrentium mosquito populations are competent in replicating and transmitting the German USUV Africa 2 strain from Berlin. The vector competence of Culex mosquitoes for zoonotic viruses like USUV is, nevertheless, not a new discovery. Several infection experiments have been performed in the past with Culex species [6].

This genus includes species which are not only abundant in Europe but can also function as bridge vectors because of the high variability in their host-feeding patterns, transmitting zoonotic pathogens from birds to mammals [40]. A Cx. neavei population from Senegal, a species which is ubiquitous in Africa, was verified to be highly competent for the Africa 2 USUV strain SAAR-1776 (GenBank accession no. AY453412) and is therefore a likely endemic vector in the USUV transmission cycle in Africa. Fourteen days post exposure to a blood meal with a virus titer of $10^6.5$ plaque forming units (PFU) per ml, its infection rate was 91%, its dissemination rate 40%, and its transmission rate 81% [22]. These values are very similar to the rates found in this report for the German Cx. pipiens biotype molestus. Recently performed infection experiments with a Cx. pipiens population from The Netherlands [24] demonstrated for the first time the ability of European mosquitoes to transmit a Europe 2 USUV strain from Italy (Bologna/09; GenBank accession no. HM569263). Similarly to the African Cx. neavei and the German Cx. pipiens biotype molestus from this study, the infection rates were high (80%), with virus...
dissemination and accumulation rates in the saliva of 69% [24]. However, contrary to the infection of the European mosquitoes with the Italian USUV strain, two *Cx. pipiens* populations from the UK [41] showed a low susceptibility to the African SAAR-1776 strain. Infection rates ranged from 0 to 14.2%, and only 1 out of 48 *Cx. pipiens* was positive for USUV in its saliva 14 dpi [41]. The variation in susceptibility could be due to the genetic variability of the virus strains with in situ evolution and host-specific mutations in European but not African lineages [14]. Therefore, it is especially interesting that *Cx. pipiens* and *Cx. quinquefasciatus* colonies from North America [23], where USUV has not been detected yet, proved highly susceptible to the very same Africa 2 strain (SAAR-1776). *Culex pipiens* showed an infection rate of 59%, and of these, 24% were positive in their saliva. *Culex quinquefasciatus* yielded similar results [23]. It is, therefore, highly likely that varying blood meal virus titers (10<sup>7.5</sup> compared to 10<sup>6.0</sup> TCID<sub>50</sub>/ml) or disparities in the vector competence of geographically distinct mosquito populations of the same species also influenced the results.

So far, no vector competence studies have been performed with *Cx. torrentium* and USUV. This is, therefore, the first confirmation of USUV transmission by a *Cx. torrentium* specimen. Due to the limited availability of *Cx. torrentium* females for this study the vector competence of this species was only examined after feeding from a blood meal with a low virus titer. It would be interesting to also test the species’ vector competence after feeding from a high virus titer and compare it to that of other species. Only recently was a German *Cx. torrentium* population described to be highly competent for the closely related flavivirus WNV with transmission rates of up to 90% [42]. *Culex torrentium* even transmitted WNV to a greater degree than *Cx. pipiens* biotype molestus [42].

In addition to the results from the vector competence experiments, the role of *Culex* mosquitoes in the transmission of USUV is reinforced through the findings of USUV in native mosquitoes in Africa and Europe. For example, USUV was detected in *Cx. antennatus*, *Cx. modestus*, *Cx. neavei*, *Cx. perexiguus*, *Cx. perfuscus*, *Cx. pipiens*, and *Cx. quinquefasciatus* [6]. Since the first isolation of USUV in Germany in 2010, multiple detections in *Cx. pipiens* sensu lato (s.l.) have been described: in 2014 near Freiburg, Baden-Wuerttemberg (Europe 3) [43], in 2015 near Leipzig, Saxony-Anhalt (Africa 2) [44], and in 2016 in Emsdetten, North Rhine-Westphalia (Africa 3) [43]. To date, USUV-positive *Cx. pipiens* s.l. have also been collected in Austria, France, Italy, Serbia, Spain, and Switzerland [6]. It must, however, be kept in mind that the mere isolation of USUV from field-collected, homogenized, whole mosquitoes cannot give a precise indication on the vector status of a specific mosquito species.

Of the invasive mosquito species in Europe, USUV nucleic acid was isolated from the Asian bush mosquito *Ae. japonicus* in Graz, Austria [45], and the Asian tiger mosquito *Ae. albopictus* in Emilia-Romagna region, Italy [46–48]. Two vector competence studies have described *Ae. albopictus* as being refractory to an USUV infection, where neither an African (SAAR-1776) nor various Italian strains (GenBank accession nos. KF055442, KF055441, and KF055440) resulted in virus accumulation in the mosquitoes’ saliva [23, 28]. In comparison, this study also investigated the USUV vector competence of the yellow fever mosquito *Ae. aegypti*. This species is thermophilic and endemic in tropical and subtropical regions of the world. Multiple introductions into Europe have been described, such as into a German household [49] and a Dutch airport [50], by means of passive mosquito dispersal. Similarly to *Ae. albopictus*, the results of this study show that *Ae. aegypti* mosquitoes are relatively refractory to USUV. The species is, therefore, unlikely to be involved in the transmission of USUV in Europe despite virus replication and dissemination in individual specimens. The infection rates of both the German and Serbian *Cx. pipiens* biotype molestus colonies were higher than those of the tested *Ae. aegypti*. None of the tested *Ae. aegypti* saliva samples contained viable virus. Nonetheless, due to the ongoing expansion of invasive species such as *Ae. albopictus* [51, 52] and *Ae. japonicus* [53, 54], the need to also investigate the role of these mosquito species in the transmission of USUV to incidental hosts (e.g. humans and equines) remains.

Multiple studies have demonstrated the virus-dose dependency of a mosquito species’ vector competence after an oral infection with USUV [22] and WNV [55]. This correlates with the results in this report where a two-log drop in the virus titer resulted in the absence of USUV-specific RNA in secondary tissues and the saliva of German *Cx. pipiens* biotype molestus. The generalized binomial regression model calculated a significant increase in the infection rates after exposure to a high rather than a low virus titer in the blood meals (GLM and LSM: df = infinity, Z-ratio = − 4.7, P < 0.001). Nonetheless, recurring USUV outbreaks among the German avifauna, such as in 2017 and 2018 [19], verify the probable presence of numerous wild bird species producing viral loads high enough to infect susceptible mosquito species. In the case of the closely related WNV, viremia levels in birds can reach 10<sup>12</sup> TCID<sub>50</sub>/ml, yet most birds do not reach titers > 10<sup>8</sup> TCID<sub>50</sub>/ml [55].

The virus titer is not the only variable to take into account, as there is also a strong temperature dependency of vector competence. For example, the Dutch *Cx. pipiens* infected with the Italian USUV strain had distinctly higher infection rates at 28°C than at 18°C (90%
versus 11%) [24]. This temperature dependency of vector competence is multifactorial and varies between mosquito species. The positive correlation between the vector competence for USUV and the ambient temperature is among others a result of the ectothermic nature of mosquitoes, where an increase in temperature directly results in an increase in the replication rate of the virus [56]. This, in turn, results in a reduction of the extrinsic incubation period [56–58]. Furthermore, a reduced effectiveness of the midgut barrier at higher temperatures, temperature-induced changes in the regulation of biotype-specific immune-responsive genes, and temperature-dependent activation of RNA interference pathways can all influence vector competence [58]. All experiments in this study were performed at 25 °C. This temperature resembles the average summer conditions around the Mediterranean Sea and southeastern European countries but is higher than the recorded long term mean temperatures of German summers. Yet, extreme heat waves such as in 2003, 2018, and 2019 have clearly demonstrated that an increase from the long-term mean temperatures in Germany is becoming more frequent [59, 60]. The summer of 2018 was the second warmest ever recorded in Germany, possibly shortening the extrinsic incubation period and allowing a more rapid autochthonous virus transmission [61]. Therefore, the occurrence of temperatures high enough to support the USUV vector-host transmission cycle in Germany is also conceivable and can help explain the increasing frequency of avian infections [18, 19, 62].

In this experiment, numerous other factors may have had an effect on the results. Aside from the Cx. torrentium specimens, all mosquitoes originated from established colonies with multiple generations in the laboratory. This may have had an impact on the genetic and phenotypic variations of the mosquito strains. The susceptibility of a mosquito to a virus cannot only be generation- but also time- and population-specific, with geographic variations in the vector competence [55, 63]. Furthermore, to obtain optimal feeding rates a sugar solution was applied to the blood meals. Even though a generation- but also time- and population-specific, with geographic variations in the vector competence [55, 63]. Furthermore, to obtain optimal feeding rates a sugar solution was applied to the blood meals. Even though a reduction of the extrinsic incubation period [56–58]. Furthermore, a reduced effectiveness of the midgut barrier at higher temperatures, temperature-induced changes in the regulation of biotype-specific immune-responsive genes, and temperature-dependent activation of RNA interference pathways can all influence vector competence [58]. All experiments in this study were performed at 25 °C. This temperature resembles the average summer conditions around the Mediterranean Sea and southeastern European countries but is higher than the recorded long term mean temperatures of German summers. Yet, extreme heat waves such as in 2003, 2018, and 2019 have clearly demonstrated that an increase from the long-term mean temperatures in Germany is becoming more frequent [59, 60]. The summer of 2018 was the second warmest ever recorded in Germany, possibly shortening the extrinsic incubation period and allowing a more rapid autochthonous virus transmission [61]. Therefore, the occurrence of temperatures high enough to support the USUV vector-host transmission cycle in Germany is also conceivable and can help explain the increasing frequency of avian infections [18, 19, 62].

In this experiment, numerous other factors may have had an effect on the results. Aside from the Cx. torrentium specimens, all mosquitoes originated from established colonies with multiple generations in the laboratory. This may have had an impact on the genetic and phenotypic variations of the mosquito strains. The susceptibility of a mosquito to a virus cannot only be generation- but also time- and population-specific, with geographic variations in the vector competence [55, 63]. Furthermore, to obtain optimal feeding rates a sugar solution was applied to the blood meals. Even though a final sugar concentration of 1.5–1.8% in the blood meals is probably too low to influence a mosquito’s vector competence [64], its influence on the mosquito’s immune response and its gut microbiota [64, 65] and therefore also on its vector competence for USUV remains unclear. Not only can the gut microbiome or the presence of intracellular Wolbachia bacteria influence a mosquito’s vector competence but also coinfections with other viruses or pathogens. For example, the German and Serbian Cx. pipiens biotype molestus colonies tested competent for USUV in this study already proved their high transmission efficiencies for a German WNV isolate [66].

USUV and WNV coinfections are not only highly conceivable in mosquitoes but have also been described in 34 avian species and in horses in Europe [7]. Future studies should, therefore, focus on what effect viral interferences will have on the antibody production in susceptible hosts and the immune response of mosquitoes [67].

**Conclusion**

Both the German (“Central European”) and Serbian (“Southern European”) Cx. pipiens biotype molestus laboratory colonies could transmit the USUV Africa 2 lineage at an extrinsic incubation temperature of 25 °C. A specimen obtained from field-collected egg rafts of Cx. torrentium, a highly prevalent species in Northern Europe, was also capable of transmitting USUV. The tropical mosquito species Ae. aegypti, on the other hand, showed reduced susceptibility with no detection of virus in its saliva.

### Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13071-020-04532-1.

**Abbreviations**

CI: Confidence interval; CO₂: Carbon dioxide; dpi: Days post infection; FCS: Fetal calf serum; FLI: Friedrich-Loeffler-Institut; GLM: Generalized binomial regression model; LMS: Least-squares mean; MEM: Minimum essential medium; NA: Not applicable; PBS: Phosphate-buffered saline; PCR: Polymerase chain reaction; PFU: Plaque forming units; RNA: Ribonucleic acid; RT-qPCR: Reverse transcription quantitative real-time PCR; s.l.: Sensu lato; TCD₅₀: Tissue culture infective dose; UK: The United Kingdom; USA: The United States of America; USUV: Usutu virus; WNV: West Nile virus.

**Acknowledgements**

A special thank you goes to Jana Schulz (FLI) for her support with the statistical analyses of the data. We are grateful to René Schottner and Cornelia Steffen (FLI) for excellent technical assistance in the laboratory. Furthermore, we thank René Klein, Marlene Hausner, and Juliane Horenk (FLI) for rearing the mosquitoes and their species identification. We would like to thank Dušan Petrič (Faculty of Agriculture, University of Novi Sad, Novi Sad, the Republic of Serbia) for supplying the Serbian Cx. pipiens biotype molestus colony and Cornelia Silaghi (FLI) for her theoretical support. We are thankful to Vinicius Pinho dos Reis (FLI) for providing us with the synthetic USUV-RNA for the calibration curve and Bärbel Hammerschmidt (FLI) for supplying the cattle blood for the infectious blood meals. We also thank Anna Heitmann and Stephanie Jansen (Bernhard-Nocht-Institut, Hamburg, Germany) for their instructions and theoretical advice.

**Authors’ contributions**

MHG, HK, CMH, and DES contributed to the study design. CMH, DES, and JL performed the experiments. CMH performed the analysis of results and wrote the manuscript. UZ, MHG, HK, and DW provided necessary resources such as the mosquitoes. DES, UZ, HK, DW, and MHG reviewed and edited the
manuscript. MHG and HK acquired funding. All authors read and approved the final manuscript.

Funding
Open Access funding enabled and organized by Projekt DEAL. This work was financially supported by the German Federal Ministry of Food and Agriculture (BMEL) through the Federal Office for Agriculture and Food (BLE), Grant numbers 2819104915, 2819104615, and 2818SE001. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Availability of data and materials
Data supporting the conclusions of this article are included within the article and its additional files. Raw data generated and/or analyzed during the present study are not publicly available, but are available from the corresponding author upon reasonable request.

Ethics approval and consent to participate
Blood for mosquito feeding was collected from animals kept at the FLI, Greifswald-Insel Riems, Germany. The animals were held and sampled according to national and European legislation (Directive 2010/63/EU on the protection of animals used for scientific purposes), and the procedures were approved by the competent authority of the Federal State of Mecklenburg-Western Pomerania, Germany (Reference No. 72213-2-041/17, Approved 12 February 2018).

Consent for publication
Not applicable.

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

Author details
1 Institute of Novel and Emerging Infectious Diseases, Friedrich-Loeffler-Institut, Federal Research Institute for Animal Health, Greifswald-Insel Riems, Germany. 2 Institute of Infectiology, Friedrich-Loeffler-Institut, Federal Research Institute for Animal Health, Greifswald-Insel Riems, Germany. 3 Present Address: Department of Molecular Genetics and Infection Biology, University of Greifswald, Greifswald, Germany. 4 Biodiversity of Aquatic and Semiaquatic Landscape Features, Leibniz Centre for Agricultural Landscape Research, Müncheberg, Germany.

Received: 9 July 2020 Accepted: 7 December 2020

Published online: 30 December 2020

References
1. Medlock JM, Hansford KM, Schaffner F, Versteirt V, Hendrickx G, Zeller H, et al. A review of the invasive mosquitoes in Europe: ecology, public health risks, and control options. Vector Borne Zoonotic Dis. 2012;12:435–47.
2. Pfeffer M, Dobler G. Emergence of zoonotic arboviruses by animal trade and migration. Parasites Vectors. 2010;3:35.
3. DeBais RL, Tyler KL. West Nile virus meningoencephalitis. Nat Clin Pract Neurol. 2006;2:264–75.
4. Venturi G, Di Luca M, Fortuna C, Remoli ME, Riccardo F, Severini F, et al. Detection of a chikungunya outbreak in Central Italy, August to September 2017. Euro Surveillance. 2017;22:170064.
5. Tomassello D, Schlagenhauf P. Chikungunya and dengue autochthonous cases in Europe, 2007–2012. Travel Med Infect Dis. 2013;11:274–84.
6. Clé M, Beck C, Salinara S, Leccolinet S, Gutierrez S, Van de Perre P, et al. USUtu virus: a new threat? Epidemiol Infect. 2019;147:e232.
7. Roehs F, Fajardo A, Moratorio G, Vignuzzi M. Usutu virus: an arbovirus on the rise. Viruses. 2019;11:640.
8. Ziegler U, Fast C, Eiden M, Bock S, Schube C, Høper D, et al. Evidence for an independent third Usutu virus introduction into Germany. Vet Microbiol. 2016;192:60–6.
9. Chvala S, Kolodziejek J, Nowotny N, Weissenböck H. Pathology and viral distribution in fatal Usutu virus infections of birds from the 2001 and 2002 outbreaks in Austria. J Comp Pathol. 2004;131:176–85.
10. Pecorari M, Longo G, Gennari W, Grotolli A, Sabbatini AMT, Tagliacozz S, et al. First human case of Usutu virus neuroinvasive infection, Italy, August–September 2009. Euro Surveillance. 2009;14:19446.
11. Grotolli A, Marcacci M, Tagliacozz S, Gennari W, Di Gennaro A, Orsini M, et al. Usutu virus infections in humans: a retrospective analysis in the municipality of Modena, Italy. Clin Microbiol Infect. 2017;23:33–7.
12. Santini M, Vilibić-Cavlek T, Barić B, Barbici L, Savic V, Stevanović V, et al. First cases of human Usutu virus neuroinvasive infection in Croatia, August–September 2013: clinical and laboratory features. J Neurovirol. 2015;21:92–7.
13. Williams MC, Simpson DIIH, Haddow AJ, Knight EM. The isolation of West Nile virus from man and of Usutu virus from the bird-biting mosquito Mansonia autes (Theobald) in the Entebbe area of Uganda. Ann Trop Med Parasitol. 1964;58:367–74.
14. Engel D, Jöst H, Wink M, Börstler J, Boschi S, Garigliani MM, et al. Reconstruction of the evolutionary history and dispersal of Usutu virus, a neglected emerging arbovirus in Europe and Africa. mbio. 2016;7:e01938-15.
15. Chvala S, Bakonyi T, Bukovsky C, Meister T, Brugger K, Rubel F, et al. Monitoring of Usutu virus activity and spreading by use of dead bird surveillance in Austria, 2003–2005. Vet Microbiol. 2007;122:237–45.
16. Weissenbch H, Bakonyi T, Rossi G, Mann P, Nowotny N. Usutu virus, Italy. 1996. Emerg Infect Dis. 2013;19:274–7.
17. Jöst H, Bialonski A, Maus D, Sambri V, Eiden M, Groschup MH, et al. Isolation of Usutu virus in Germany. Am J Trop Med Hyg. 2011;85:551–3.
18. Ziegler U, Jöst H, Müller K, Fischer D, Rinder M, Tietze DT, et al. Epidemic spread of Usutu virus in Southwest Germany in 2011 to 2013 and monitoring of wild birds for Usutu and West Nile viruses. Vector Borne Zoonotic Dis. 2015;15:481–8.
19. Michel F, Sieg M, Fischer D, Keller M, Eiden M, Reuschel M, et al. Evidence for West Nile virus and Usutu virus infections in wild and resident birds in Germany, 2017 and 2018. Viruses. 2019;11:674.
20. Cadar D, Lühken R, van der Jeugd H, Garigliani M, Ziegler U, Keller M, et al. Widespread activity of multiple lineages of Usutu virus, western Europe, 2016. Euro Surveillance. 2017;22:30452.
21. Turell MJ. Members of the Culex pinnipes complex as vectors of viruses. J Am Mosq Control Assoc. 2017;28:123–6.
22. Nikolay B, Di Lallo M, Faye O, Boye CS, Sall AA. Vector competence of Culex neurons (Diptera: Culicidae) for Usutu virus. Am J Trop Med Hyg. 2012;86:993–6.
23. Cook CL, Huang YS, Lyons AC, Alto BW, Unlu I, Higgs S, et al. North American Culex pinnipes and Culex quinquefasciatus are competent vectors for Usutu virus. PLoS Negl Trop Dis. 2018;12:e0006732.
24. Fros JJ, Miesen P, Vogels CB, Gaibani P, Sambri V, Martina BE, et al. Comparative Usutu and West Nile virus transmission potential by local Culex pinnipes mosquitoes in North-Western Europe. One Health. 2015;1:31–6.
25. Hesson JC, Rettich F, Merdic E, Vignевич G, Ostman O, Schäfer M, et al. The arbovirus vector Culex torrentium is more prevalent than Culex pinnipes in northern and central Europe. Med Vet Entomol. 2014;28:179–86.
26. Brugman VA, Hernández-Triana LM, Medlock JM, Fooks AR, Carpenter S, Johnson N. The role of Culex pinnipes L. (Diptera: Culicidae) in virus transmission in Europe. Int J Environ Res Public Health. 2018;15:389.
27. Rudolf M, Czajka C, Börstler J, Melaun C, Jöst H, von Thien H, et al. First nationwide surveillance of Culex pinnipes complex and Culex torrentium mosquitoes demonstrated the presence of Culex pinnipes biotype pinnipes/molestus hybrids in Germany. PLoS ONE. 2013;8:e71832.
28. Pugioli A, Bonilauri P, Calzolari M, Lelli D, Carrieri M, Urbanelli S, et al. Does Aedes albopictus (Diptera: Culicidae) play any role in Usutu virus transmission in Northern Italy? Experimental oral infection and field evidences. Acta Trop. 2017;172:192–6.
29. Abbo SR, Visser TM, Wang H, Goertz GP, Fros JJ, Abma-Henkens MHC, et al. The invasive Asian bush mosquito Aedes japonicus found in the Netherlands can experimentally transmit Zika virus and Usutu virus. PLoS Negl Trop Dis. 2020;14:e0008217.
30. Viña-Rodríguez A, Sachse K, Ziegler U, Chaintotius SC, Keller M, Groschup MH, et al. A novel pan-flavivirus detection and identification assay based on RT-qPCR and microarray. Biomed Res Int. 2017;2017:42487/56.
31. Mayr A, Bachmann PA, Bibrack B, Wittmann G. Quantitative Bestimmung der Virusinfektiosität (Virusinfektion). In: Mayr A, Bachmann PA, Bibrack B,
Wittmann G, editors. Virologische Arbeitsmethoden, Band I (Zellkulturen - Bebrütete Hühnererei - Versuchstiere). Jena: Gustav Fischer Verlag; 1974. p. 35–9.
32. Heitmann A, Jansen S, Lühekn R, Leggewie M, Schmidt-Chanasis J, Tannich E. Forced salivation as a method to analyze vector competence of mosquitoes. J Exp. Vis 2018;18:3857980.
33. Pinho dos Reis V. The role of integrins in flavivirus infection [Doctoral dissertation]. [Greifswald, Germany]: Ernst-Moritz-Arndt-Universität Greifswald, 2018.
34. Folmer O, Black M, Hoeh W, Lutz R, Vrijenhoek R. DNA primers for amplification of mitochondrial cytochrome c oxidase subunit I from diverse metazoan invertebrates. Mol Mar Biol Biotechnol. 1994;3:294–9.
35. Hebert PDN, Ratnasingham S, de Waard JR. Barcoding animal life: cytochrome c oxidase subunit 1 diversities among closely related species. Proc R Soc Lond B. 2003;270:596–9.
36. R Development Core Team. R: a language and environment for statistical computing. Vienna: R Foundation for Statistical Computing; 2019.
37. Lenth RV. Least-squares means: the R package lsmeans. J Stat Softw. 2016;69:1–33.
38. Harvey WR. Least-squares analysis of data with unequal subclass numbers. Washington, DC: USDA National Agricultural Library; 1960.
39. Tukey JW. The philosophy of multiple comparisons. Stat Sci. 1991;6:100–16.
40. Hanmer GJ, Kitron UD, Brawin JD, Loss SR, Ruiz MO, Goldberg TL, et al. Culicidae (Diptera: Culicidae): a bridge vector of West Nile virus to humans. J Med Entomol. 2008;45:125–8.
41. Hernández-Triana LM, de Marco MF, Mansfield NL, Thorne L, Lumley S, Marston D, et al. Assessment of vector competence of UK mosquitoes for Usutu virus of African origin. Parasites Vectors. 2018;11:381.
42. Jansen S, Heitmann A, Lühekn R, Leggewie M, Helms M, Badusche M, et al. Culxere Lorentzen: a potent vector for the transmission of West Nile virus in Central Europe. Viruses. 2019;11:492.
43. Scheuch DE, Schäfer M, Eiden M, Heym EC, Ziegler U, Walther D, et al. Detection of Usutu, Sindbis, and Batai viruses in mosquitoes (Diptera: Culicidae) collected in Germany, 2011–2016. Viruses. 2018;10:389.
44. Sieg M, Schmidt V, Ziegler U, Keller M, Höper D, Heenemann K, et al. Outbreak and co-circulation of three different Usutu virus strains in Eastern Germany. Vector Borne Zoonotic Dis. 2017;17:662–4.
45. Camp Jv, Kolodzieck J, Nowotny N. Targeted surveillance reveals native and invasive mosquito species infected with Usutu virus. Parasites Vectors. 2019;12:46.
46. Tambu M, Boninlau P, Bellini R, Calzolari M, Albien A, Sambri V, et al. Detection of Usutu virus within a West Nile virus surveillance program in Northern Italy. Vector Borne Zoonotic Dis. 2011;11:551–7.
47. Mancini G, Montarsi F, Calzolari M, Capelli G, Dottori M, Ravagnan S, et al. Mosquito species infected in the circulation of West Nile and Usutu viruses in Italy. Vet Ital. 2017;53:97–110.
48. Calzolari M, Boninlau P, Bellini R, Albien A, Delfilippo F, Maioli G, et al. Mosquitoes species involved in the circulation of West Nile and Usutu viruses in mosquitoes sampled in Emlia-Romagna region (Italy) in 2009. PLoS ONE. 2010;5:e14324.
49. Kampen H, Jansen S, Schmidt-Chanasis J, Walther D. Indoor development of Aedes aegypti in Germany, 2016. Euro Surveillance. 2016;21:30407.
50. Ibañez-Justicia A, Gloria-Soria A, den Hartog W, Dik M, Jacobs F, Stoor A. The first detected airborne introductions of yellow fever mosquitoes (Aedes aegypti) to Europe, at Schiphol International airport, the Netherlands. Parasites Vectors. 2017;10:603.
51. Kuhlisch C, Kampen H, Walther D. The Asian tiger mosquito Aedes albopictus (Diptera: Culicidae) in Central Germany: surveillance in its northernmost distribution area. Acta Trop. 2018;188:78–85.
52. Walther D, Scheuch DE, Kampen H. The invasive Asian tiger mosquito Aedes albopictus (Diptera: Culicidae) in Germany: local reproduction and overwintering. Acta Trop. 2017;166:186–92.
53. Kampen H, Werner D. Out of the bush: the Asian bush mosquito Aedes japonicus japonicus (Theobald, 1901) (Diptera, Culicidae) becomes invasive. Parasites Vectors. 2014;7:59.
54. Zielke DE, Walther D, Kampen H. Newly discovered population of Aedes japonicus japonicus (Diptera: Culicidae) in Upper Bavaria, Germany, and Salzburg, Austria, is closely related to the Austrian/Slovenian bush mosquito population. Parasites Vectors. 2016;9:163.
55. Vogels CB, Goertz GP, Pijpman JP, Koenraadt CJ. Vector competence of European mosquitoes for West Nile virus. Emerg Microbes Infect. 2017;6:e96.
56. Lambrechts L, Paphiatis M, Fauq S, Carrington LB, Kramer LD, Thomas MB, et al. Impact of daily temperature fluctuations on dengue virus transmission by Aedes aegypti. Proc Natl Acad Sci USA. 2011;108:7460–5.
57. Lühekn R, Jost H, Cadar D, Thomas SM, Bosch S, Tannich E, et al. Distribution of Usutu virus in Germany and its effect on breeding bird populations. Emerg Infect Dis. 2017;23:1994–2001.
58. Vogels CB, Fros JJ, Goertz GP, Pijpman JP, Koenraadt CJ. Vector competence of northern European Culex pipiens biotypes and hybrids for West Nile virus is differentially affected by temperature. Parasites Vectors. 2016;9:393.
59. Deutscher Wetterdienst (DWD). Deutschlands wetter in Sommer 2018: Außergewöhnlich warm, trocken und sonnig – viele neue regionale Rekorde. https://www.dwd.de/DE/presse/pressemittenliegen/DE/2018/20180830_deutschlandwetter_sommer_news.html. Accessed 30 Jan 2020.
60. Deutscher Wetterdienst (DWD). The weather in Germany in summer 2019. https://www.dwd.de/DE/press/press_release/EN/2019/20190830_the_weather_in_germany_in_summer_2019.html. Accessed 30 Jan 2020.
61. Ziegler U, Lühekn R, Keller M, Cadar D, van der Grinten E, Michel F, et al. West Nile virus epizootic in Germany. Antivir Res. 2019;162:39–43.
62. Michel F, Fischer D, Eiden M, Fast C, Reuschel M, Kuller K, et al. West Nile virus and Usutu virus monitoring of wild birds in Germany. Int J Environ Public Health. 2018;15:171.
63. Richards SL, Lord CC, Pesko KN, Tabachnick WJ. Environmental and biological factors influencing Culex pipiens quinquefasciatus (Diptera: Culicidae) vector competence for West Nile virus. Am J Trop Med Hyg. 2010;83:126–34.
64. Vaidyanathan R, Fleisher AE, Minnick SL, Simmons KA, Scott TW. Nutritional stress affects mosquito survival and vector competence for West Nile virus. Vector Borne Zoonotic Dis. 2008;8:727–32.
65. Muturi EJ, Dunlap C, Ramirez JL, Rooney AR, Kim CH. Host blood-meal source has a strong impact on gut microbiota of Aedes aegypti. FEMS Microbiol Ecol. 2019;95:fly213.
66. Holicki CM, Ziegler U, Ráileanu C, Kampen H, Werner D, Schulz J, et al. West Nile virus lineage 2 vector competence of indigenous Culex and Aedes mosquitoes from Germany at temperate climate conditions. Viruses. 2020;12:561.
67. Rizzoli A, Jiménez-Clavero MA, Barzon L, Cordioli P, Figuerola J, Koraka P, et al. The challenge of West Nile virus in Europe: knowledge gaps and research priorities. Euro Surveillance. 2015;20:21135.
Publisher's Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.