Blood donation screening for West Nile virus (WNV) was mandatory in the Lazio region in 2017 and 2018 (June-November) according to the national surveillance plan. In these years, all five donations reactive in WNV nucleic acid amplification tests harboured instead Usutu virus (USUV). Clade ‘Europe 2’ was identified in four blood donations and a 2018 mosquito pool. The cocirculation of WNV and USUV in Lazio warrants increased laboratory support and awareness of possible virus misidentification.

In the summer and autumn seasons of 2017 and 2018, following serological identification of West Nile virus (WNV) infection in horses, mandatory WNV screening with nucleic acid amplification test (NAT) was established for blood donations in some provinces of the Lazio Region in central Italy: Viterbo (starting from 18 August 2017), Latina (12 September 2018) and Rome (10 October 2018). During this period, five blood donations were reactive in WNV NAT screening. In none of these samples could WNV positivity be established by further tests conducted at the Regional Reference Laboratory.

In the present study, we describe serological, detailed molecular and phylogenetic analyses undertaken to better characterise the WNV positivity observed during the screening.

Processing of blood samples

For the Lazio Region, centralised blood screening for WNV is performed at the Biological Qualification Center (CQB) for NAT blood screening at the Sandro Pertini Hospital in Rome, using NAT (Cobas WNV NAT screening test, Roche, Mannheim, Germany). Blood donations reactive in WNV NAT screening and confirmed twice by the same method at the CQB, are considered not suitable for the transfusion service and are discarded. Plasma aliquots of these donations are sent to the Regional Reference Laboratory for Arbovirosis (Laboratory of Virology, National Institute for Infectious Diseases L. Spallanzani, Rome) for confirmatory testing and further diagnostic investigation. Donors who provided WNV NAT-positive donations are recalled 2–3 weeks after donation, to complete serological investigation.

All five donors reactive in repeated WNV NAT were asymptomatic and fulfilled the eligibility criteria concerning WNV risk. According to Ministerial Decree of 2 November 2015, in order to guarantee self-sufficiency in blood components during the summer period, it is recommended to use the WNV NAT test as an alternative to the provision of 28 days suspension of donors who have spent at least one night in areas at risk of WNV infection [1]. The areas with a documented WNV circulation are established by an appropriate epidemiological surveillance system. The main demographic data of these donors are shown in the Table (Donors #1 to #5).  

Virological characterisation of WNV NAT-reactive samples

Plasma samples were concentrated by ultracentrifugation and tested with a nested pan-flavivirus-RT-PCR targeting the NS5 gene (modified from [2], amplicon size: 210 nt), followed by amplicon sequencing. New sequences described in this report have been registered in GenBank under accession numbers MK015649,
The sequencing results indicated that all five blood donations harboured Usutu virus (USUV) and not WNV. These results were further confirmed with a USUV-specific real-time RT-PCR, modified from Nikolay et al. [3], and by a USUV-specific nested RT-PCR targeting NS5, followed by sequencing. The phylogenetic tree, built with both neighbour-joining and Bayesian maximum clade credibility approaches, indicated that the USUV strains detected in four of the five blood donors from the Lazio region belonged to the clade ‘Europe 2’, while the strain detected in donor #1 belonged to the clade ‘Europe 3’ (Figure 1 and 2).

Phylogenetic tree based on the partial NS5 gene (position 9004–9406 of HM569263), built using the neighbour-joining method. The evolutionary distances were computed using the Kimura 2-parameter method. The rate variation among sites was modelled with a gamma distribution (shape parameter=0.25). Each record consists of accession number, place and year of detection/isolation. Phylogenetic analysis was conducted in the MEGA7 software package.

The serological investigation revealed that four donors did not have WNV- or USUV-specific antibodies at donation, while donor #1 had both IgG and IgM specific for WNV. In follow-up samples, obtained 2–3 weeks apart, USUV seroconversion was observed for donors #2 and #5, while the USUV-specific antibody titre exceeded the WNV-specific antibody titre in donor #1 (Table).

MK024375, MK006173, MK006174, MK006175, MK060108, MK060109, MK060110 and MK060111.

The serological investigation for WNV was performed with commercially available kits: indirect immunofluorescence assay (anti-West Nile virus IFA IgM and IgG, Euroimmun, Germany) for patients #1–#5 and ELISA (West Nile Virus IgM/IgG Capture DxSelect Focus Diagnostics, Diasorin, United States) for patients #6–#9.

The titles are expressed as highest serum dilution showing specific immunofluorescent staining. For WNV ELISA, the test results are expressed as pos/neg.

Anti-USUV IgM and IgG antibodies were determined by immunofluorescence assay using home-made glass slides, prepared with a mix of uninfected and USUV-infected Vero E6 cells, according to standard procedures established for flaviviruses [20].
Figure 1
Phylogenetic analysis of Usutu virus strains, neighbour-joining method, central and northern Italy, 2017–2018 (n = 50)

Phylogenetic tree based on the partial NS5 gene (position 9004–9406 of HM569263), built using the neighbour-joining method. The evolutionary distances were computed using the Kimura 2-parameter method. The rate variation among sites was modelled with a gamma distribution (shape parameter = 0.25). Each record consists of accession number, place and year of detection/isolation. Phylogenetic analysis was conducted in the MEGA7 software package.


Colour code of Usutu virus clades: red: Europe 1; blue: Europe 2; pink: Europe 3; green: Europe 4; yellow: Africa 3; black: Africa 2.
Vector investigation

According to the national integrated surveillance plan for WNV and USUV, the Istituto Zooprofilattico Sperimentale delle Regioni Lazio e Toscana performed the entomological investigation: adult mosquitoes were sampled at sites with known virus circulation (areas where horses positive for anti-WNV antibodies or blood donors reactive in WNV NAT had been identified) using traps of the model Italian Mosquito Trap (IMT; PeP, San Giuliano Milanese, Italy), BG Sentinel (BioGents, Regensburg, Germany) and Gravid (BioQuip Products, Rancho Dominguez, United States). The sampling protocol was defined depending on the number of WNV cases in the area and on the number of caught mosquitoes. After mosquito sorting and identification, those of the species *C. pipiens* were divided in pools of at most 100 specimens and analysed by RT-PCR for virus detection.

In the period between September and October 2018, 47 trapping exercises were performed at nine sites with known virus circulation in the provinces of Rome, Latina and Frosinone. A total of 2,443 specimens of *C. pipiens* were caught, divided into 38 pools and tested by RT-PCR for the presence of WNV and USUV RNA. Among the 38 tested pools, 14 were positive for USUV (minimum infectious rate: 1.3); they originated from two municipalities in the Latina province. Sequencing indicated that these viruses belonged to the ‘Europe 2’ clade and were very similar to those obtained from blood donors #2 to #5 (Figure 1).

During the same period, three horses with neurological signs and positive WNV serology were located in the province of Latina. One horse died, and WNV lineage 2 was detected by RT-PCR in its brain (cerebellum and medulla oblongata). These results were confirmed by the National Reference Centre for Foreign Animal Diseases (CESME), Istituto Zooprofilattico Sperimentale dell’Abruzzo e del Molise.

Characterisation of Usutu virus strains detected in blood donors from northern Italy

We also performed molecular analysis on four USUV-positive blood donations identified in 2017 by the
Turdus merula (Eurasian blackbird), reported in Europe in 2001, causing mass mortality of USUV was isolated in South Africa in 1959 and was first involved mosquitos as vectors and birds as amplifying clades so far recognised [5]. Its natural life cycle involves mosquitos as vectors and birds as amplifying hosts. Humans and horses are dead-end hosts, as for WNV.

USUV was isolated in South Africa in 1959 and was first reported in Europe in 2001, causing mass mortality of Eurasian blackbird (Turdus merula); however, a retrospective study showed that it has been present in Italy at least since 1996 [6]. Subsequently, USUV spread to other country, including Hungary (2003–2006), Switzerland (2006), Spain (2003–2006), Germany (2010), Czech Republic (2014), Belgium, France and the Netherlands (2016), where the virus was detected or isolated from mosquitos, birds and bats [7-10].

The first human USUV infection was reported in the Central African Republic in 1981 [11]; the first neuroinvasive human infection in Europe was reported in Italy in an immunocompromised host in 2009 [12]; several cases of disease in humans were described thereafter [13-15].

Recently, USUV circulation has been increasing in several Europe regions, overlapping with areas endemic for WNV that exploits the same main mosquito vector [16]. Molecular and serological evidence of USUV circulation has been reported in northern Italy, involving the ‘Europe 2’ and the ‘Europe 4’ clades [4] and infecting birds, mosquitos and humans; human infections have been both asymptomatic and symptomatic [13].

An annual surveillance plan is issued by the Italian Ministry of Health with the aim of monitoring WNV and USUV circulation and triggering appropriate public health measures. The purpose is the limitation of virus spread, control of the vector population and elimination of transmission through blood, blood components, tissues and organ donation during the period of increased vector activity in the summer and autumn season (June–November) [1]. In 2017 and 2018, respectively two and three blood donations from the Lazio region were positive in WNV NAT screening. Molecular characterisation revealed that the WNV NAT positivity was indeed due to the presence of USUV RNA. Intensified entomological surveillance for arboviruses detected 14 mosquito pools that were positive for USUV RNA.

Partial molecular characterisation indicated close genetic relationships between human and mosquito USUV strains detected in Lazio region; all belonged to the ‘Europe 2’ clade. The only USUV strain of clade ‘Europe 3’, detected in donor #1, was presumably acquired in Switzerland, according to the travel history of the donor; in addition, this donor’s place of residence, Argentina, accounted for the serological pattern observed at donation, that was consistent with previous exposure to other related flaviviruses.

The presence of WNV lineage 2 detected by molecular assay in the brain of a symptomatic horse in the province of Latina is evidence of the cocirculation of WNV and USUV in this area. Data from blood donors in Lombardy confirmed the cocirculation, in northern Italy, of different USUV strains (belonging to clades ‘Europe 2’ and ‘Europe 4’). This finding is in line with previous reports [13] and indicates autochthonous circulation after multiple importation events.

This is the first report showing circulation of USUV virus in the Lazio region, both in mosquito vectors and in human hosts, while so far, USUV has not been detected in other animal hosts (neither equids nor birds).

This study corroborates the hypothesis that USUV can cause clinically asymptomatic infection in humans [17]; this, in turn, may be of concern regarding the safety of the blood supply in areas where the virus is active. Although there is no evidence of human-to-human transmission of USUV in the transfusion (or transplant) setting, asymptomatic infected donors may donate USUV-infectious blood, which may cause severe disease in immunocompromised patients [18].

Public health authorities, blood transfusion services and clinicians should be aware of the expanding risk of USUV infection in humans, especially during the summer, and advanced laboratory support is required to address the problem of possible virus misidentification by screening tests.

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Authors’ contributions

Conceptualisation: FC, MRC; Methodology: FC, FR; Software: FC, FM; Formal analysis: FC, FR; Investigation: SM, CC, TC, FR, EP, FC, IR, CDL, FR, LB, EL; Resources: IR, CDL, FR, CC; Conceptualisation: FC, MRC; Methodology: FC, FR; Software: FC, FM; Formal analysis: FC, FR; Investigation: SM, CC, TC, FR, EP, FC, IR, CDL, FR, LB, EL; Resources: IR, CDL, FR, CC; Writing – original draft: FC; Writing – review and editing: MRC, FC, GI, AS; Resources: IR, CDL, FR, CC; Formal analysis: FC, FR; Investigation: SM, CC, TC, FR, LB, EL; Resources: IR, CDL, FR, CC; Conceptualisation: FC, MRC; Methodology: FC, FR; Software: FC, FM; Formal analysis: FC, FR; Investigation: SM, CC, TC, FR, EP, FC, IR, CDL, FR, LB, EL; Resources: IR, CDL, FR, CC; Writing – original draft: FC; Writing – review and editing: MRC, FC, GI, AS.

Conflict of interest

None declared.

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