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Detection and characterization of the Dezidougou virus (genus *Negevirus*) in mosquitoes (*Ochlerotatus caspius*) collected in the Republic of Sakha (Yakutia)

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Abstract

Introduction. Monitoring and research on arthropod-borne microorganisms is important. Recently, with the development of next-generation sequencing methods, many previously unknown viruses have been identified in insects.

Aim of the study. Isolation of viruses from mosquitoes sampled in the Republic of Sakha (Yakutia), followed by the study of a new for Russia negevirus isolated from mosquitoes of the species *Ochlerotatus caspius*, including determination of its complete nucleotide sequence, phylogenetic and virological characteristics.

Materials and methods. Dezidougou virus isolation was performed on C6/36 (*Aedes albopictus*) cell culture. Electron microscopy was performed using a JEM 1400 electron microscope. Nucleotide sequence screening was performed by NGS on a high-throughput sequencer MiSeq, Illumina (USA). Full genome nucleotide sequence was determined by Sanger sequencing. Phylogenetic analysis was performed using GenBank database, using Vector NTI Advance 11 and MEGA 11 programs.

Results. The virus isolated from mosquitoes replicated efficiently in C6/36 cells, causing their death. However, it did not replicate in the mammalian cell cultures used. The isolated virus did not cause pathologic manifestations in suckling mice when infected intracerebrally. Electron microscopic examination of the purified virus-containing suspension showed the presence of spherical viral particles with a diameter of 45–55 nm. The results of full genome sequencing identified it as belonging to Dezidougou virus, first isolated in Côte d'Ivoire. The nucleotide sequence of the genome of Yakutsk 2023 strain of Dezidougou virus was deposited in GenBank (PP975071.1).

Conclusion. Dezidougou virus of genus *Negevirus* was isolated and characterized for the first time in the Russian Federation. Further studies on the prevalence of negeviruses, their virological features, potential importance for public health and their impact on vector competence of vectors are important and promising.

Keywords: *Dezidougou virus; negeviruses; insect specific viruses; Republic of Sakha (Yakutia)*

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Ethical Statement. The authors confirm compliance with institutional and national standards for the use of laboratory animals according to the «Consensus author guidelines for animal use» 2010. The study protocol was approved by the Bioethics Committee of the State Scientific Center of Virology and Biotechnology «Vector» of Rosпотребнадзор (Protocol No. 02 of April 03, 2023).

ОРИГИНАЛЬНОЕ ИССЛЕДОВАНИЕ

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Выявление и характеристика вируса *Dezidougou* (род *Negevirus*) в комарах (*Ochlerotatus caspius*), собранных на территории Республики Саха (Якутия)

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Резюме

Введение. Мониторинг и исследование микроорганизмов, переносимых членистоногими, имеют важное значение. В последнее время с развитием методов секвенирования нового поколения (NGS) у насекомых идентифицировано множество ранее неизвестных вирусов.

Цель исследования. Изоляция вирусов из комаров отобранных в Республике Саха (Якутия), с последующим исследованием нового для России негевируса, выделенного из комаров вида *Ochlerotatus caspius*, включая определение его полной нуклеотидной последовательности, филогенетическую и вирусологические характеристики.

Материалы и методы. Изоляцию вируса *Dezidougou* проводили на культуре клеток C6/36 (*Aedes albopictus*). Электронную микроскопию осуществляли с использованием электронного микроскопа JEM 1400. Скрининговое определение нуклеотидных последовательностей выполняли с применением метода NGS на высокопроизводительном секвенаторе MiSeq, Illumina (США). Определение полногеномной нуклеотидной последовательности проводили секвенированием по методу Сэнгера. Филогенетический анализ выполняли с использованием базы данных GenBank и программ Vector NTI Advance 11, MEGA 11.

Результаты. Выделенный из комаров вирус эффективно реплицировался в клетках C6/36, вызывая их гибель. При этом он не размножался в использованных клеточных культурах млекопитающих. Выделенный вирус при интрацеребральном инфицировании мышей-сосунков не вызывал у них патологических проявлений. При электронно-микроскопическом исследовании очищенной вирусосодержащей суспензии было показано наличие сферических вирусных частиц диаметром 45–55 нм. Результаты полногеномного секвенирования идентифицировали его принадлежность к вирусу *Dezidougou*, впервые выделенному в Кот д'Ивуаре. Нуклеотидная последовательность генома штамма Yakutsk 2023 вируса *Dezidougou* была депонирована в базе данных GenBank (PP975071.1).

Заключение. Впервые в Российской Федерации был выделен и охарактеризован вирус *Dezidougou* рода *Negevirus*. Проведение дальнейших исследований распространенности негевирусов, их вирусологических особенностей, потенциального значения для здравоохранения и влияния на векторную компетентность переносчиков является важным и перспективным.

Ключевые слова: вирус *Dezidougou*; негевирусы; insect specific viruses; Республика Саха (Якутия)

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Финансирование. Исследование было выполнено при поддержке Министерства науки и высшего образования Российской Федерации, соглашение № 075-15-2019-1665 и Государственного задания 9/21.

Конфликт интересов. Авторы декларируют отсутствие явных и потенциальных конфликтов интересов, связанных с публикацией настоящей статьи.

Этическое утверждение. Авторы подтверждают соблюдение институциональных и национальных стандартов по использованию лабораторных животных в соответствии с Consensus author guidelines for animal use 2010. Протокол исследования одобрен Комитетом по Биоэтике ФБУН «Государственный научный центр вирусологии и биотехнологии «Вектор» Роспотребнадзора (Протокол № 02 от 03.04.2023).

Introduction

Historically, there has been a special interest in arthropod-infecting viruses because of their involvement in the spread of viruses pathogenic to humans and animals. Initially, the term insect specific viruses (ISVs) referred to viruses of the genus *Ortoflavivirus* (family Flaviviridae) that could replicate only in insect cells but had a similar genome organization to orthoflaviviruses pathogenic to vertebrates [1, 2]. With the development of high-throughput sequencing methods, new ISVs have been identified [3, 4]. To date, the group of ISVs includes representatives of different families: Baculoviridae, Poxviridae, Iridoviridae, Ascoviridae, Polydnviridae, the genome of which is represented by double-stranded DNA; Parvoviridae (single-stranded DNA); Reoviridae, Tetraviridae, Dicrostoviridae, Nodaviridae, Picornaviridae, Flaviviridae (RNA(+)); Rhabdoviridae (RNA(-)) [1–5].

Representatives of the genus *Negevirus* are found in different parts of the world and infect a wide range of hematophages (mosquitoes of the genera *Culex*, *Aedes* and *Anopheles*, as well as mosquitoes of the genus *Lutzomyia*). At the same time, negeviruses are genetically close to plant viruses from the genera *Cilevirus*, *Higrevirus* and *Blunervirus* (family Kitaviridae), which allowed us to hypothesize the role of plants in the natural transmission cycle of negeviruses [6, 7]. The genus *Negevirus* is named after the first fully characterized isolate of *Negev virus* [6]. More than 36 species of viruses, not including 30 unclassified viruses (<https://www.ncbi.nlm.nih.gov/Taxonomy/Browse/>), are currently classified as negeviruses.

Dezidougou virus (DEZV) was first isolated at the Pasteur Institute (Dakar, Senegal) from a population of mosquitoes *Aedes aegypti* collected near the village of Dezidougou, Côte d'Ivoire, in 1987 [6, 7]. Later, DEZV was found in different parts of the world: Europe, Africa, Central and South America [6, 8, 9]. Virions of negeviruses have a spherical shape with a diameter of 45–50 nm [6]. The genome of negeviruses is represented by unsegmented single-stranded RNA with positive sense, 7–10 kb in size. Most negeviruses have three open reading frames (ORF) flanked by non-translated regions at the 5'- and 3'-ends. Each ORF is separated by short intergenic regions, the largest frame ORF1 encodes viral polymerase, ORF2 encodes glycoprotein, and ORF3 encodes membrane proteins [7]. At the end of the viral genome, a poly(A)-tail is present, ranging from 13 to 52 nt in length. [6].

The aim of this study was to isolate viruses from mosquitoes sampled in the Republic of Sakha (Yakutia) and to study DEZV virus, new for Russia, isolated from mosquitoes of species *Ochlerotatus caspius*, including determination of its complete nucleotide sequence, phylogenetic and virological characterization.

Materials and methods

During the field period of 2023, 500 mosquitoes were captured on the territory of the Republic of Sakha (Yakutia) in the Saisarsky (62.029955/129.668761) and Central districts (62.009133/129.744127). The mosquitoes were transported in cooler bags on wet cloths at +4 °C and

stored at –18–24 °C. Mosquitoes were sorted according to phenotypic traits and pooled into pools of 10 individuals. The morphological keys for genus determination were the size of the individual, color of scales, length of legs and structure of the mouth apparatus. A fragment of 16S rRNA and a fragment of the *COI* gene of the mitochondrial genome were sequenced to determine the mosquito species. A total of 51 mosquito pools were examined.

Before homogenization, all mosquitoes were washed in 70% ethanol and then twice with water to remove potential surface microorganisms. The resulting pools of *Ochlerotatus* sp. mosquitoes were homogenized in 300 µl of saline solution on a TissueLyser LT homogenizer (Qia-gen, The Netherlands).

Virus isolation was performed on the C6/36 (*Aedes albopictus*) cell culture, highly sensitive for ISVs [10–12]. The monolayer was grown to 80–90% confluency in a 24-well plate (Greiner, Austria) in DMEM F12 medium (Gibco, USA) containing 10% fetal bovine serum (Gibco, USA), 100 IU/mL penicillin and 100 µg/mL streptomycin (Gibco, USA) in an atmosphere with 5% CO₂ at 28 °C. The culture was infected with filtered mosquito homogenates and incubated for 7 days, evaluating possible cytopathic effects (CPE). After that, the plates with cell monolayers were subjected to three freeze/thaw cycles, the obtained suspensions were clarified from cell debris by centrifugation at 8000g at 4 °C for 5 min and used for passaging (infection of fresh C6/36 cell monolayers similar to the above described). After detection of pronounced virus-specific CPE, the viral suspensions were used to infect monolayers of C6/36 cells grown in culture vials (Greiner, Austria). Determination of the infectious titer of the obtained virus-containing suspension was performed according to the standard method on C6/36 culture by micromethod with registration of the results by microscopy and MTT test [13, 14]. The titer was calculated using the Spearman–Kerker method [15].

Light microscopy was performed using an Olympus CKX53 inverted microscope (Olympus, Japan), with fixation with an Olympus SC50 digital camera (Olympus, Japan) at ×200 magnification and digital processing in the Cellsens Standard program.

For electron microscopy, virus-containing culture medium was clarified from cells by centrifugation at 8000g for 10 min to remove residual cellular debris. The virus was concentrated using a VivaSpin centrifuge concentrator (Sartorius, Germany) for 30 min at 6000g. The concentrate was resuspended, fixed with formalin and applied as suspensions on copper grids coated with formvar substrate film and stabilized with carbon. The obtained drugs were stained with 1% aqueous uranyl acetate solution according to the generally accepted technique. The samples were examined using a JEM 1400 electron microscope (Jeol, Japan) at an accelerating voltage of 80 kV. Image analysis and processing were performed using iTEM software package (SIS, Germany).

The ability of DEZV to infect mammalian cells was investigated on transplantable cultures: HEK-293A (human embryonic kidney cells), Vero E6 (green monkey kidney cells) and SPEV (pig embryonic kidney cells).

A monolayer of cells in T-25 culture vials was infected with 0.1 mL of inoculum and incubated in DMEM F12 maintenance medium with 2% FBS (fetal bovine serum) for 10 days. The resulting cell lysates were tested for the presence of infectious viral progeny by titration on C6/36 cell culture.

The susceptibility of animals to DEZV was tested on 2–3-day-old suckling mice. Animals infected intracerebrally with 0.02 mL of inoculum were monitored for the 21 days.

Screening for ISV nucleotide sequences was performed by high-throughput sequencing. Total RNA was extracted using the Extract RNA Reagent (Eurogen, Russia) according to the manufacturer's protocol. The aqueous phase obtained after addition of chloroform and subsequent centrifugation was collected and diluted 1 : 1 with freshly prepared 70% ethanol and purified on Cleanup Mini spin columns (Eurogen, Russia) and treated with Benzonase (Merck, Germany) [16]. Synthesis of the first strand of cDNA was performed using the NEBNext Ultra Directional module (New England Biolabs Inc., USA). Second strand DNA synthesis was performed using UMI Second Strand Synthesis Module for QuantSeq FWD Illumina (Lexogen, Austria). Prepared dsDNA libraries were analyzed on a MiSeq high-throughput sequencer (Illumina, USA). Cutadapt (version 1.18) and SAMtools (version 0.1.18) were used to remove Illumina adapters and re-read sequences. Contigs were assembled *de novo* using the MIRA assembler (version 4.9.6).

The complete nucleotide sequence of DEZV RNA was determined by reverse transcription polymerase chain reaction (RT-PCR) using primers (**Appendix 1**) complementary to the genome fragments of the virus under study. RT-PCR was performed in 15 μ L of the reaction mixture in a C1000 thermocycler (Bio-Rad, USA). The obtained amplicons were separated by gel electrophoresis in 2% agarose gel in Tris-acetate buffer (Eurogen, Russia) with 0.1% ethidium bromide (Sisco Research Laboratories, India).

Sanger sequencing reaction was performed using BigDye Terminator v. 3.1 Cycle Sequencing Kit (Applied Biosystems, USA), on a 3500xl Genetic Analyzer (Applied Biosystems, USA). The obtained nucleotide sequences were aligned to prototype sequences using the UniproUGENE v. 1.48 software product. Phylogenetic analysis was performed using the GenBank database. Phylogenetic trees were constructed using Vector NTI Advance 11 and MEGA 11 programs. Phylogenetic trees were built by maximum likelihood method using 500 bootstrap replicates.

The authors confirm compliance with institutional and national standards for the use of laboratory animals according to Consensus author guidelines for animal use 2010. The study protocol was approved by the Bioethics Committee of the State Scientific Center of Virology and Biotechnology «Vector» of Rospotrebnadzor (Protocol No. 02 of 03.04.2023).

Results

In an attempt to isolate viruses from homogenates of 51 mosquito pools, 3 consecutive passages were performed on C6/36 culture. When the 3rd passage was performed on a monolayer of C6/36 cells corresponding to one of the pools used for infection, cytopathic manifestations were recorded on the 5th day of incubation, which increased with further incubation. To confirm the presence of lytic-inducing virus, an additional 4th passage was performed on a confluent C6/36 monolayer. **Fig. 1** shows microphotographs illustrating the presence of pronounced virus-specific CPE leading to 100% cell death by day 6 post-infection. The infectious activity of the isolated viral isolate was determined by titration on C6/36 cell culture and amounted to 7.1 log₁₀ TCID₅₀/mL. Electron microscopic examination was performed to determine the shape and size of viral particles. The negatively stained virus suspension contained particles predominantly 48–52 nm in diameter, round in shape, with a more electron-dense area in the central part (**Fig. 2**).

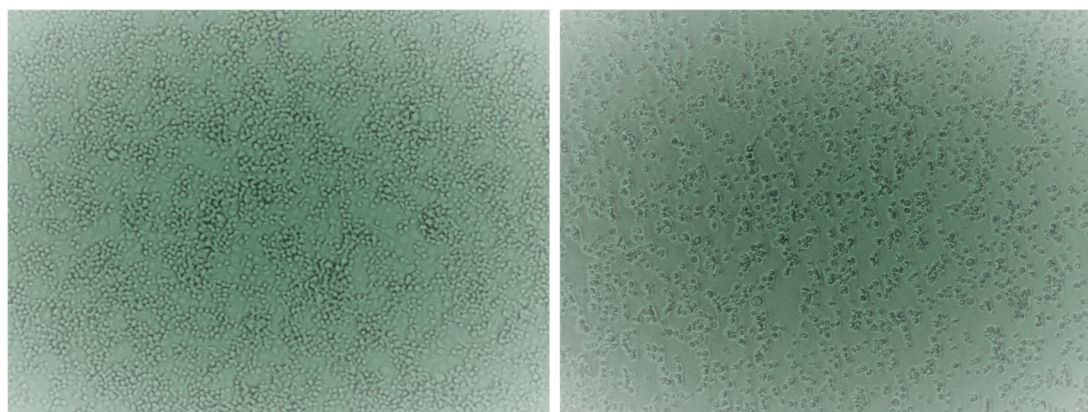


Fig. 1. Light microscopy ($\times 200$) of C6/36 cell culture infected with DEZV Yakutsk 2023 strain 120 hours after infection.

On the left is a control of C6/36 cell culture.

Рис. 1. Световая микроскопия ($\times 200$) культуры клеток C6/36, инфицированной штаммом DEZV Yakutsk 2023, через 120 ч после инфицирования.

Слева представлен контроль культуры клеток C6/36.

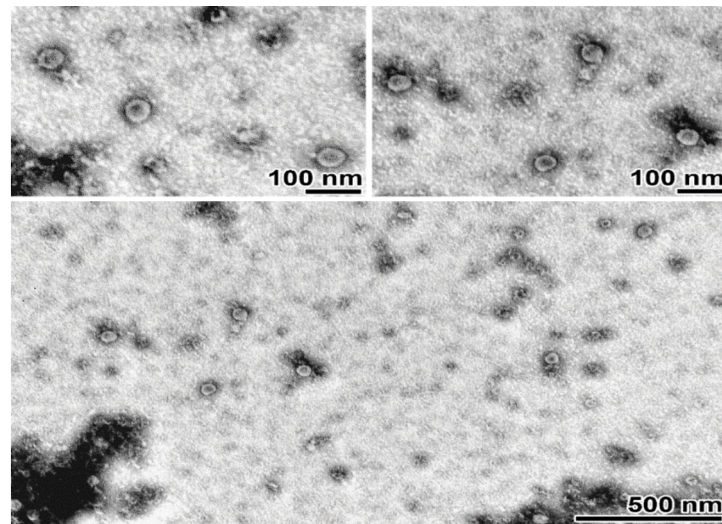


Fig. 2. Transmission electron microscopy of a purified virus suspension.

Rounded particles with a diameter of 45–55 nm and an electron-dense region in the central part. Contrasted with 2% uranyl acetate. The scale is indicated on images.

Рис. 2. Просвечивающая электронная микроскопия очищенной суспензии вируса.

Частицы округлой формы диаметром 45–55 нм и электронно-плотной областью в центральной части. Контрастирование 2% уранилацетатом. Бар указан на снимках.

To determine the potential ability of the viral isolate to replicate in mammalian cells, cell cultures Vero E6, HEK-293A and SPEV were infected at a multiplicity of 10 TCID₅₀/cell and incubated for 10 days. No virus-specific effects on cell cultures were detected. Cell lysates of the above cultures were titrated on C6/36 culture. Infectious virus was detected. The results obtained prove the inability of the isolated virus to replicate in the used mammalian cells.

To reveal possible pathogenic properties of the isolated virus, suckling mice were infected. It was shown that intracerebral infection with a dose of 10⁵ TCID₅₀/mouse did not cause any clinical manifestations of infection during the whole period of observation (21 days).

Nucleotide sequence screening of total RNA from infected C6/36 cells by next-generation sequencing (NGS) allowed us to identify a fragment of nucleotide sequence corresponding to ORF1 of the Dezidougou negevirus genome. The fragment was 471 bp in length with a read count of 1465 and a coverage of 293. Sanger sequencing was performed to determine the complete nucleotide sequence of DEZV negevirus. The genome is a non-segmented positive-sense single-stranded RNA 9010 nt in size, and has three open reading frames. The obtained sequence was deposited in the GenBank database under accession number: PP975071.1. The similarity of the nucleotide sequence of the isolated DEZV Yakutsk 2023 variant with known negevirus isolates is summarized in the following **Table**. The highest level of similarity (92%) in nucleotide sequence DEZV Yakutsk 2023 showed in comparison with the isolate from *Coquillettidia richiardii* from Germany (DEZV OP576003). The similarity rate in comparison with a DEZV variant isolated in Spain (MT096525) was 86%. The level of similarity with nucleotide

sequences of other negevirus was as follows: with Kustavi Negevirus (ON949944, Spain) – 69.6%, Utsjoki Negevirus 3 (ON955101, Finland) – 72.1%, Wallerfield virus (KX518839, Panama) and Uxmal virus (MH719095.1, Mexico) – 68%. The presence of a large number of amino acid substitutions indicates a significant genetic diversity of negevirus.

Fig. 3 shows the phylogenetic tree with our isolated DEZV Yakutsk 2023 and negevirus. The closest to DEZV Yakutsk 2023 is the prototypical Dezidougou virus strain 8345 (OP576003.1) from Germany with an identity level of 91.95%. Among other negevirus, the closest are viruses isolated in Finland (Utsjoki Negevirus 1, ON949947), Castlereas virus in Australia (KX886280) and Ying Kou virus in China (isolate NC 040636.1) with a similarity level of 72.1–72.4% in nucleotide sequences.

Discussion

Over the past decades, a large number of insect viruses have been discovered that belong to various families, including viruses belonging to genus *Negevirus*.

The virus isolated by us from pools of mosquitoes sampled in the Republic of Sakha (Yakutia) replicated efficiently in C6/36 (*Aedes albopictus*) cells, causing their death. At the same time, it was unable to infect and multiply in mammalian cell cultures (Vero E6, HEK-293A and SPEV). The isolated virus did not cause pathologic manifestations in suckling mice when infected intracerebrally with a high dose. The results of complete genome sequencing determined that it belonged to negevirus, the highest level of homology was noted with Dezidougou virus, first isolated in Côte d'Ivoire. Electron microscopic examination of purified virus suspension showed the presence of spherical viral particles

Table. Levels of homology (%) of nucleotide/derived amino acid sequences of strain DEZV Yakutsk 2023 in comparison with the closest negevirus strains

Таблица. Уровни сходства (%) нуклеотидной/выведенной аминокислотной последовательностей штамма DEZV Yakutsk 2023 в сравнении с наиболее близкими штаммами негевиров

Name of strain Название штамма	GenBank	Country Страна	Year Год	Nucleotide sequence, % Нуклеотидная последовательность, %	Amino acid sequence, % Аминокислотная последовательность, %
Dezidougou virus strain 8345	OP576003.1	Germany Германия	2014	91.95	98.40
Dezidougou virus strain ArA 20086	JQ675604.1	Côte d'Ivoire Кот-д'Ивуар	1984	85.22	96.27
Dezidougou virus strain DEZI/ Aedes africanus/SEN/ DAK-AR-41524/1984	KY968698.1	Senegal Сенегал	1984	86.24	96.18
Dezidougou virus isolate FTA2-3	MT096525.1	Spain Испания	2015	86.08	96.44
Kustavi Negevirus isolate FIN/VS-2018/100	ON949944.1	Finland Финляндия	2017	69.61	71.62
Agua Salud Negevirus isolate PA-2013-MP416-PP	MK959116.1	Panama Панама	2013	67.31	57.38
Wallerfield virus strain TR7904	NC_023440.1	Trinidad and Tobago Тринидад и Тобаго	2009	67.84	61.27
Wallerfield virus isolate PA-2013-MP416-PP	MK959117.1	Panama Панама	2013	68.16	61.45
Uxmal virus isolate UXMV-M985	MH719095.1	Mexico Мексика	2007	68.57	60.58
Utsjoki Negevirus 3 isolate FIN/L-2018/06	ON955101.1	Finland Финляндия	2015	72.14	82.15
Culex Biggie-like virus strain CBigVL/Kern	MH188028.1	USA США	2016	70.88	55.19
Culex negev-like virus 3 strain mos172X44875	NC_035129.1	Australia Австралия	2015	70.00	55.30
Tanay virus isolate 11-3, complete genome	KF425262.1	Philippines Филиппины	2005	70.02	52.82
Goutanap virus 16GH1	LC504569.1	Ghana Гана	2016	68.85	56.00

with a diameter of 45–55 nm, characteristic of negeviruses. In a number of studies about negevirus, it has been shown that these viruses reproduce only in arthropod cells and do not replicate in vertebrate cells. Thus, in [6], the authors infected C6/36, Vero, and BHK-21 cell cultures, as well as newborn mice (intracerebral inoculation) with Negev (NEGV), Piura (PIUV), Dezidougou (DEZV), Ngewotan (NWTV), Loreto (LORV), and Santana (SANV) viruses. All of the above viruses were only able to replicate in C6/36 cells and did not cause any disease in sucklings, which is consistent with the results of the present study.

The genus *Negevirus* consists of a diverse group of insect-specific viruses, with a genome represented by single-stranded (+)RNA, isolated from mosquitoes and phlebotomine mosquitoes in Brazil, Colombia, Peru,

Panama, USA, Germany, Spain and Nepal. These viruses were isolated from pools of mosquitoes collected in the field, suggesting that they are fairly widespread among mosquitoes in the wild. Most negevirus are characterized by high genetic variability and the presence of interspecies transmission. Negevirus are widely distributed geographically and have a diverse range of hosts among insects – mosquitoes of the genera *Culex*, *Aedes* and *Anopheles*, sand flies of the genus *Lutzomyia*, etc. [17, 18]. Our results on the isolation of DEZV in the Republic of Sakha (Yakutia) in Russia confirm the data on the wide geographical distribution of *Negevirus*.

The biological and potential public health significance of negevirus remains to be determined. Since negevirus were originally isolated from naturally selected mosquitoes of different genera that are vectors of arbo-

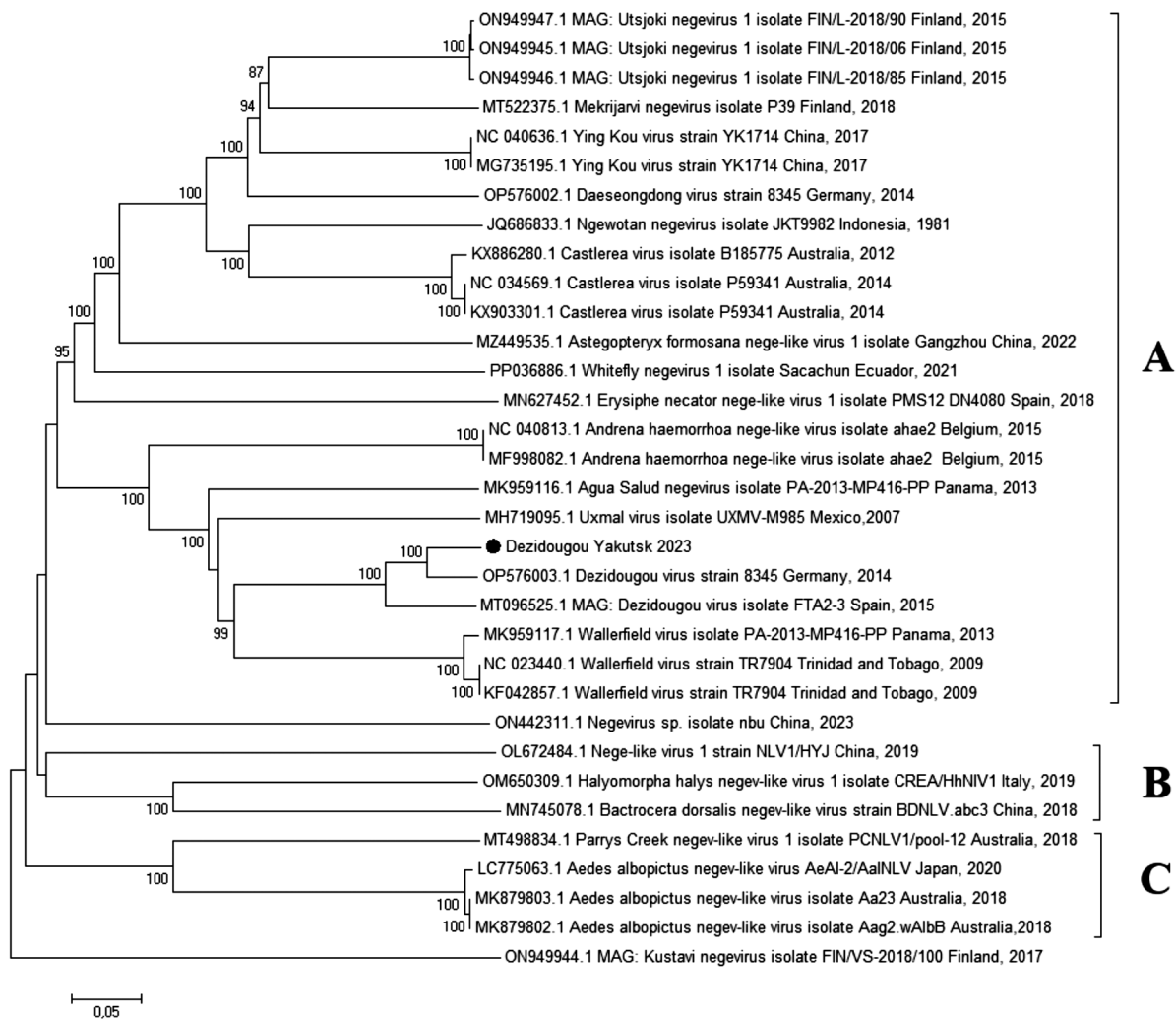


Fig. 3. Phylogenetic tree showing the maximum likelihood analysis of full-length viral sequences of DEZV and viruses of the genus Negevirus.

The sequence characterized in this study is highlighted with the symbol (●). A, B, and C are the main branches of negevirus.

Рис. 3. Филогенетическая диаграмма, отображающая анализ максимального правдоподобия полноразмерных вирусных последовательностей DEZV и других негевировусов.

Последовательность, охарактеризованная в этом исследовании, выделена символом (●). А, В и С – основные ветви негевировусов.

viruses, it is possible that negevirus infection may influence the susceptibility and vector competence of vectors to viral pathogens of vertebrates. For example, it has been shown experimentally on *Ae. aegypti* that infection of mosquitoes with certain strains of the bacterial endosymbiont *Wolbachia* prevents dengue virus replication and reduces vector competence [18, 19]. And if a bacterial endosymbiont can alter mosquito vector competence to arboviruses, it is likely that a viral symbiont could also have a similar effect. It is thought that ISVs could also potentially be used as biological control agents with the proposed elimination of superinfection with human pathogenic arboviruses and maintenance of the effect in nature through transovarial transmission [20]. In a recent study, the negevirus Piura has been shown to effectively inhibit Zika virus replication in insect cells. Co-infection

of C6/36 cells with PIUV resulted in a 10,000-fold reduction in the infectious titer of Zika virus compared to Zika virus mono-infected cells. At the same time, Zika virus was unable to inhibit PIUV replication [21]. E. Patterson et al. [22] demonstrated that Negev virus inhibited replication of Chikungunya arboviruses and Eastern equine encephalitis arboviruses belonging to the *Alphavirus* genus in coinfecting mosquito cells. Other researchers have shown that insect-specific *Culex* Flavivirus (CxFV) Izabal did not inhibit West Nile virus (WNV) replication in C6/36 cells and *Culex quinquefasciatus* mosquitoes. Most importantly, the WNV transmissible efficacy was enhanced in CxFV-infected mosquitoes [23].

It has previously been shown experimentally that negevirus are found in the salivary glands of insects and, therefore, there is the potential for transmission to the ver-

tebrate host during feeding [24, 25]. Consequently, humans and other vertebrates may have contact with negevirus, increasing the likelihood of some of them adapting and eventually evolving already as a conditional vertebrate pathogen. There is a reasonable assumption that many vertebrate viruses transmitted by arthropods were originally insect-specific [7]. The assumption that ISVs are ancestors of arboviruses makes these viruses a potential tool for studying the evolution of host-to-host transmission.

Conclusion

For the first time in the territory of the Russian Federation, a virus of genus *Negevirus* was identified and isolated in a pool of mosquitoes sampled in the Republic of Sakha (Yakutia). The isolated negevirus strain DEZV Yakutsk 2023 replicated efficiently in C6/36 cells (*Ae. albopictus*), causing their death. At the same time, it was unable to infect and multiply in the mammalian cell cultures used and did not cause pathologic manifestations in suckling mice upon intracerebral infection. The complete nucleotide sequence of the viral genome of DEZV Yakutsk 2023 was determined by NGS and Sanger sequencing and phylogenetic analysis was performed.

Further research on the prevalence of negevirus, their virological features, potential public health significance and impact on vector competence seems to be important and promising.

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