

TBEV-transmission and natural cycles

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Key points

- The natural cycle of the TBE virus is dependent on vector ticks and reservoir hosts.
- There are differing transmission cycles in varying environments, from cold northern coniferous forests to temperate central European forests.
- Within a natural transmission cycle there are different ways of transmission - tick-to-tick (transovarial, sexual), host-to-tick (viremic), and also tick-to-tick and host-to-host.
- The complexity of natural transmission cycles is inadequately explored and poorly understood.

Introduction

Ticks play a critical role in the transmission of a wide variety of viral, bacterial, and protozoan pathogens to humans and animals.^{1,2} In the case of humans, infection is accidental as these transmission cycles are invariably enzootic with the natural hosts most frequently being wild birds and mammals.¹ In order to be tangentially affected by such cycles, humans must be bitten by a vector tick species found in habitats visited by humans, as well as the tick's usual hosts, as the dispersal of ticks not attached to hosts covers only very short distances. In addition, the tick has to accept humans as a suitable host, meaning that the species involved usually have a broad host spectrum.

Nevertheless, these tick species may only be part of the transmission cycle, with eco-epidemiologically significant sub-cycles involving tick species not commonly in contact with humans.^{3,4} Thus, the transmission of tick-borne pathogens often comprises a complex network of interactions involving several tick and host species. Below, we provide background to the biology of ticks and how this can influence, specifically, the eco-epidemiological cycle of tick-borne encephalitis virus (TBEV).

Structure and morphology

Ticks are a group of hematophagous ectoparasites with about 989 living species.⁵ They belong to the phylum Arthropoda, the class Arachnida, the superorder Acarina, and the order Ixodida, and they are exclusively parasitic. The Ixodida contain 3 families: the Ixodidae with 15 genera (hard ticks), the Argasidae with 15 genera (soft ticks), and the Nuttalliellidae, represented by only one species, *Nuttalliella namaqua*.⁶⁻⁹ A summary of tick taxonomy and the two groups is presented in [Figure 1](#). All the tick species

involved in the eco-epidemiological cycle of TBEV belong to the Ixodidae. Details of tick biology generally can be found in a variety of publications, for example in Nicholson et al.,⁹ Petney et al.,¹⁰ and Sonenshine and Roe,¹¹ and a list of valid species names in Guglielmone and Nava.¹² The following genera of ticks contain species known to transmit TBEV or in their species TBEV was detected.

Ixodes is the largest tick genus, with 266 described species worldwide.⁵ *Ixodes* species are characterized by a distinct groove that encircles the anus anteriorly and a lack of eyes. Males have seven sclerotized ventral plates that are absent in the males of other genera. The genus *Ixodes* has been subdivided in roughly 15 subgenera (e.g., *Ixodes*, *Pholeoixodes*) on the basis of morphology.^{13,14} The genus has a worldwide distribution, including parts of Antarctica.⁹ Some species are particularly important as vectors of TBEV: *Ixodes ricinus* the castor bean tick or sheep tick in Europe and middle Asia, *Ixodes persulcatus* the taiga tick in northeastern Europe and northern Asia, and *Ixodes ovatus* in the forest belt of middle Asia and Japan.

The genus *Dermacentor* has 44 species worldwide.⁹ The basis capitulum appears rectangular when viewed dorsally. A pair of medially directed spurs occurs on the first pair of coxae. The palps are short and thick. The scutum is almost always ornamented, eyes present. *Dermacentor* species are found mostly in Europe, Asia, and North America.⁹ In Europe, TBEV has been recovered from 2 species, *Dermacentor reticulatus* (the ornate dog tick), *Dermacentor marginatus* (the ornate sheep tick), and in Asia from *Dermacentor nuttalli*.

Haemaphysalis is the second largest (176 species) tick genus.⁵ This eyeless genus can, in most cases, be identified by a pronounced lateral projection of palpal segment 2, which extends well beyond the basis capitulum. In Europe,

Figure 1: The *Ixodidae* family

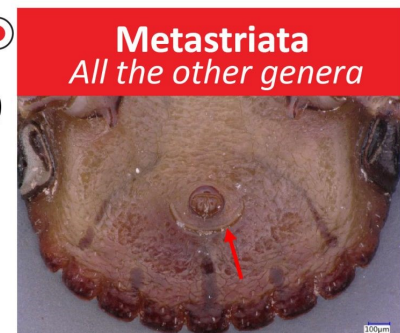
- **Phylum – Arthropoda**
- **Class – Arachnida**
- **Superorder – Acarina**
- **Order – Ixodida**
- **Suborder – Ixodina**

‣ **Family *Ixodidae***

- | | |
|--------------------------|--|
| • <i>Ixodes</i> | • <i>Dermacentor</i> |
| • <i>Amblyomma</i> | • <i>Haemaphysalis</i> |
| • <i>Anomalohimalaya</i> | • <i>Hyalomma</i> |
| • <i>Bothriocroton</i> | • <i>Margaropus</i> |
| • <i>Cosmiomma</i> | • <i>Nosomma</i> |
| • <i>Rhipicentor</i> | • <i>Rhipicephalus</i> (including <i>Boophilus</i>) |
| • <i>Africanella</i> | • <i>Robertsicus</i> |

‣ **Family *Argasidae***

‣ **Family *Nutalliellidae***



The *Ixodidae* family is divided in two groups: *Prostriata*, which includes only the genus *Ixodes* and which is characterized by an anal groove encircling the anus anteriorly (blue arrow); and *Metastriata*, including 14 genera, which all have an anal groove behind the anus (red arrow).

TBEV has been recovered from *Haemaphysalis punctata* (the red sheep tick), *Haemaphysalis concinna* in Europe and Asia, and from *Haemaphysalis longicornis* in Asia.⁹

The genus *Hyalomma* is relatively small with 27 species of small- to large-sized ticks.¹⁴ They are characterized by their elongated palps, which are at least twice as long as wide. The distinct eyes are located in sockets adjacent to the postero-lateral edges of the scutum that is unornamented. The distribution of *Hyalomma* species is limited to the Old World, primarily to arid or semiarid habitats. *Hyalomma marginatum* (the Mediterranean *Hyalomma*) is the only member of this genus from which TBEV has been recovered.^{15,16}

The biology of hard ticks

All the species known to transmit TBEV have a 3-host life-cycle (Figure 2). Each postembryonic life stage requires a blood meal from a suitable host, after which the tick detaches and molts in the leaf litter. The arrows with broken lines in the figure show the potential transmission paths to humans. The line from larvae to humans indicates that transovarial transmission from an infected female can

happen which results in infective larvae. Infection of the tick can occur when larvae, nymphs, or females feed on an infective host (see below).

The larva, nymph, and adult (female or male – Figures 3a, 3b, 3c, and 3d) are active stages that require a host (this is not the case for males of the genus *Ixodes*, which can mate off-host without feeding).¹⁷ Larvae are easily recognizable by the presence of only 3 pairs of legs, and absent spiracular and genital apertures (Figures 4a and 4b). Nymphs have 4 pairs of legs and spiracles (Figures 5a and 5b). Adult females have 4 pairs of legs, and spiracles, a genital aperture, and porose areas on the dorsal surface of the basis capituli (Figures 3a and 3b). Adult males have 4 pairs of legs, the scutum covers the entire dorsal surface, and seven hard sclerotized plates cover the ventral body surface of some species (Figures 3c and 3d).

Types of hard ticks

Ixodid ticks fall into two behavioral groups. Exophilic or non-nidicolous ixodid ticks occur in the open environment and are associated, with forests, savannahs, second-growth areas of scrub and brush, grassy meadows, semi-desert, or desert areas. These species are usually not very host-

Table 1: Tick species, tick habitats, and involved hosts in relation to the TBEV subtype and distribution

Tick species (subgenus)	Main habitats	Hosts	Virus subtype	Vector role	References **
<i>Ixodes</i> (<i>Ixodes ricinus</i>)	deciduous and mixed forests	reptiles, birds, mammals, human	ES, SS	principal vector in Europe	18-26
<i>Ixodes</i> (<i>Pholeoixodes</i>) <i>arboricola</i>	nidicolous, nests and burrows	birds	ES	persistence and transmission to white mice; considered to be a secondary amplifying vector of TBE virus in wild populations	27,28
<i>Ixodes</i> (<i>Pholeoixodes</i>) <i>lividus</i>	nests	birds	SS		19
<i>Ixodes</i> (<i>Pholeoixodes</i>) <i>hexagonus</i>	nidicolous, nests, burrows, caves, rock shelters, dog kennels and also buildings	hedgehogs, wild carnivores, dogs, rarely human	ES	transstadial and transovarial transmission; TBE virus isolates. Isolated from female and nymph infesting a hedgehog; a pool of 3 females from red fox	29-32
<i>Ixodes</i> (<i>Pholeoixodes</i>) <i>canisuga</i>	nidicolous, nests, burrows	hedgehogs, wild carnivores, dogs	?	little is known about the vector competence	
<i>Ixodes</i> (<i>Scaphixodes</i>) <i>frontalis</i>	nests	birds	ES	detection of TBEV; vector competence and importance in transmission cycle unknown	33-35
<i>Ixodes</i> (<i>Exopalgiger</i>) <i>trianguliceps</i>	endophilic, shady mixed and deciduous forests	small mammals (ca 50 species), birds, and a viviparous lizard	ES	vector and reservoir of TBE virus among the small mammals	30,36
<i>Ixodes</i> (<i>Ixodes</i>) <i>persulcatus</i>	exophilic, deciduous and mixed forests	polyxenic reptiles, birds, mammals, human	ES, SS, FES	principal vector for the Siberian and Far Eastern subtypes from north-eastern Europe to Russian Far East, China and Japan	18-20

ES, European subtype (TBEV-EU); FES, Far Eastern subtype (TBEV-FE); SS, Siberian subtype (TBEV-Sib)

* Reference for tick habitat and host: Nowak-Chmura and Siuda, 2012;³⁶ Petney et al., 2012;¹⁷ Guglielmone et al., 2014¹²

** Reference for tick species involved in TBE virus transmission

specific. Nidicolous or endophilic ixodid ticks live in or near the nests of their hosts, are adapted to highly specialized environments (crevices or other shelters used by their hosts), and tend to be more host-specific.⁹ Many *Ixodes* species are nidicolous.⁹ The main vectors of TBEV, *I. ricinus* and *I. persulcatus* are exophilic and exceptional both in terms of their large variety of hosts they use as well as the habitats they occupy.³⁷

Host-finding behavior

Ixodid ticks' host-seeking behavior is under the control of different abiotic factors that differ according to the region. In temperate and sub-polar regions, seasonal activity is

mainly regulated by ambient temperature, changing photoperiod, and incident solar energy, and in the more temperate regions, tick activity is often controlled by saturation deficit and relative humidity, with long-term dry conditions being adverse for survival.³⁸⁻⁴¹ Those species involved in the transmission of TBEV tend to quest passively or ambush their hosts by climbing onto weeds, grasses, or other lower vegetation to wait for a host to pass nearby.

Ixodes ricinus adults can climb as high as 1.5 m on brushy vegetation. The immature stages are found lower, up to 70 cm for larvae (O. Kahl, personal communication) and less than 1 m for nymphs.⁹ Ticks are able to sense a host with their Haller's organ, which is located on the tarsus I. Haller's

organ possesses chemo-, mechano-, and thermoreceptors that also ensures (together with the receptors on the palps) selection of a suitable feeding site on the host body. The most important stimuli are carbon dioxide (CO₂), vibration produced by moving potential hosts, and host temperature. For some species, visual images, host smell, and even noise can stimulate the tick.^{9,42,43}

Feeding behavior

Feeding behavior, even on preferred hosts, is not a uniform process. An ixodid tick may crawl on the host for several hours in search of a suitable feeding site. After attachment, many ixodid ticks secrete cement during the first 1–2 days to secure themselves at the wound site.⁴² The feeding tick begins salivating into the developing hematoma and sucking blood; phases of salivation and blood sucking alternate.⁹ Saliva not only plays an important role in the feeding tick's osmoregulation⁴⁴ but also has a variety of pharmacological effects. There is an extensive array of antihemostatic, anti-inflammatory, and immunomodulatory proteins and lipids in the tick saliva that suppress the host's ability to reject the feeding tick.^{9,45-47} Anticoagulant effects, inhibiting factor Xa, were first shown in *I. ricinus* in 1898-1899.⁴⁸ In addition, many tick species produce proteins that inhibit thrombin directly or inhibit the conversion of prothrombin to thrombin by inhibiting factor V. Other proteins prevent

platelet aggregation or bind, antagonize or degrade important host mediators of pain, itching and inflammation, particularly the host's own histamine, serotonin, and bradykinin.^{9,47}

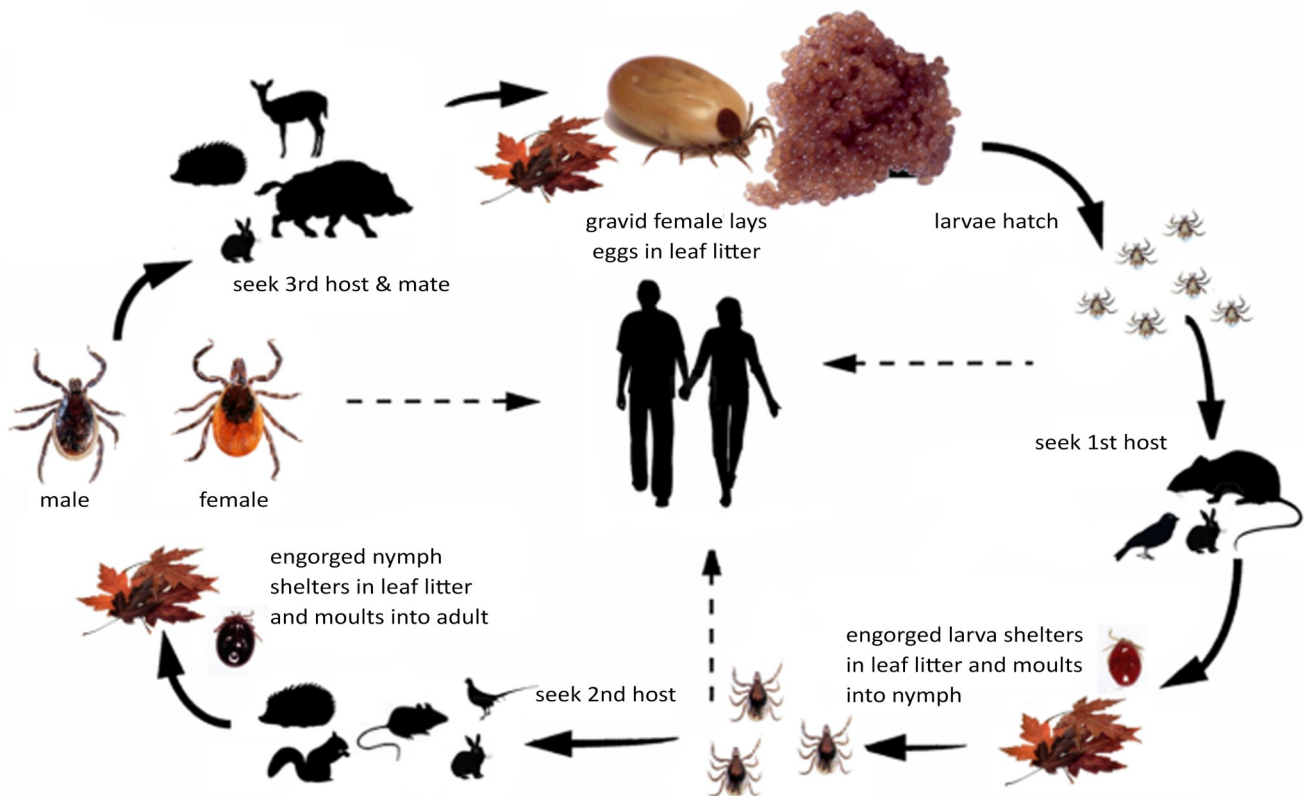
Ixodid ticks feed gradually because they must first produce new cuticle to accommodate the massive blood meal.⁹ Typical attachment periods range from as few as 2 days for larvae to as long as 13 days for females.⁴⁹

An *I. ricinus* female can reach approximately 450 mg at the end of feeding from approximately 2 mg at the beginning of feeding.⁵⁰

Drop-off

The controlled timing of drop-off from the host offers important ecological advantages. For non-nidicolous ticks, such drop-off rhythms are synchronized with host behavioral patterns. This tends to disperse fed ticks in optimal habitats where they can develop and reproduce. Photoperiod appears to be the dominant abiotic exogenous factor affecting drop-off patterns. The daily light:dark cycle induces a regular rhythm of feeding and dropping off. Detachment may occur while hosts are inactive in their nests or burrows or, alternatively, it may be coordinated with the period of high host activity.⁵⁰

Figure 2



The life-cycle of *Ixodes ricinus*. The dotted arrows indicate potential transmission to humans. ©Nina Littwin

Figure 3a

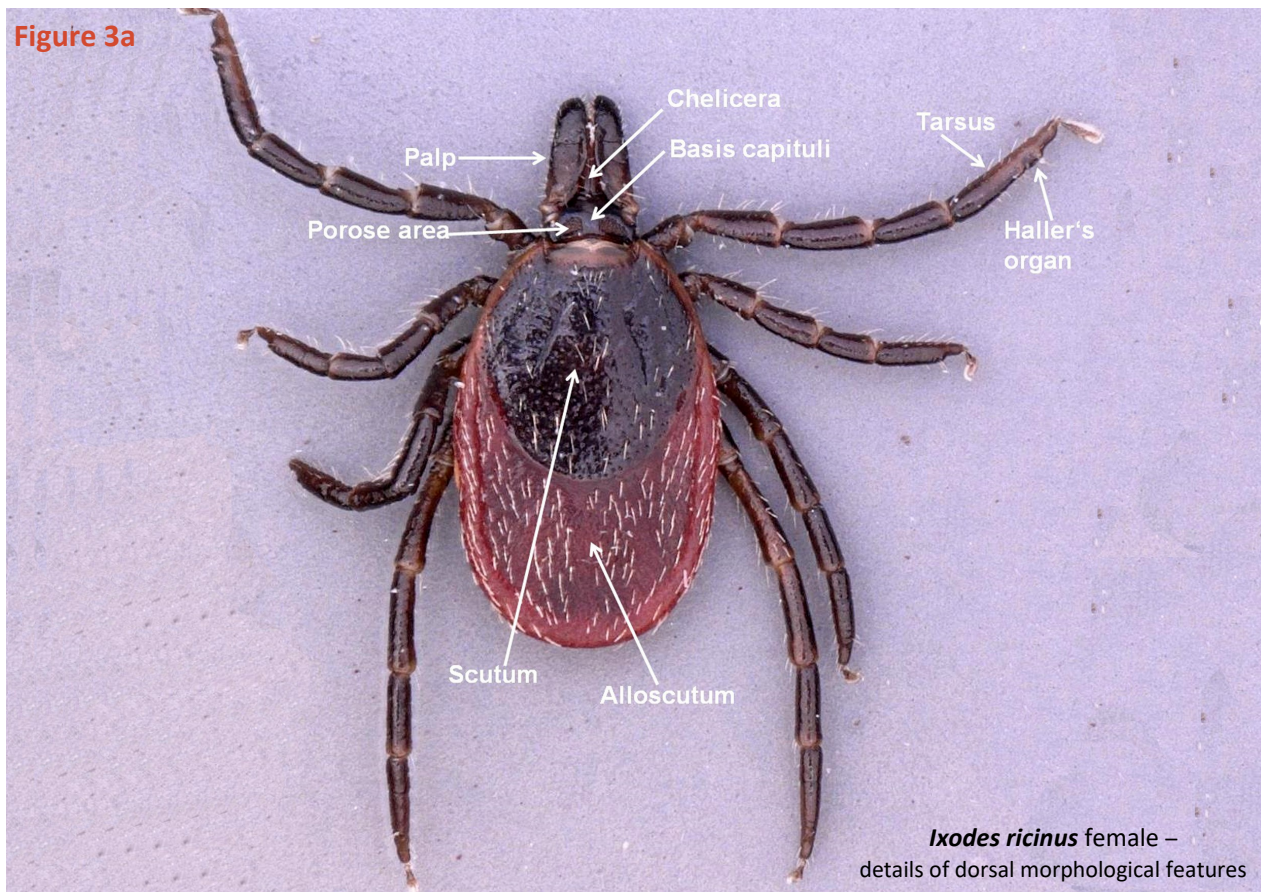


Figure 3b

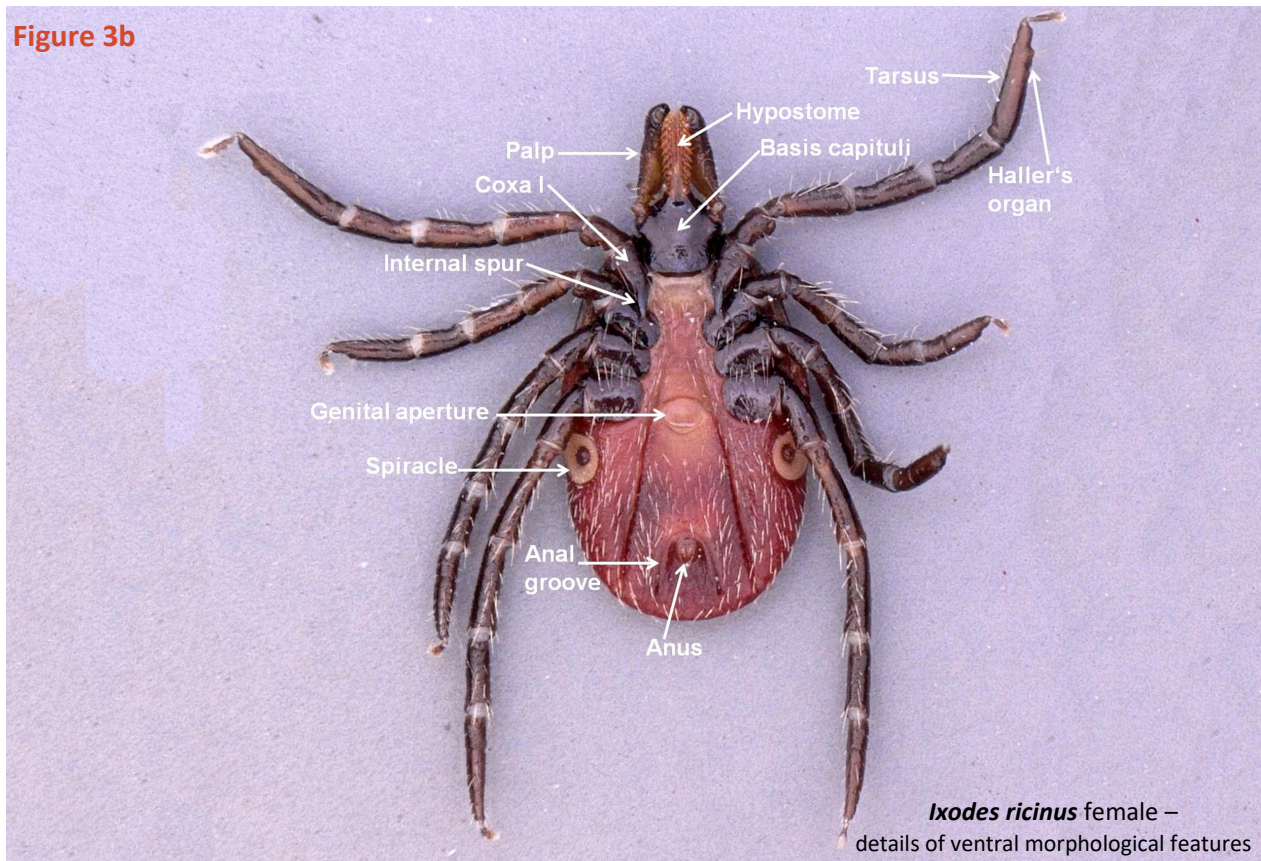


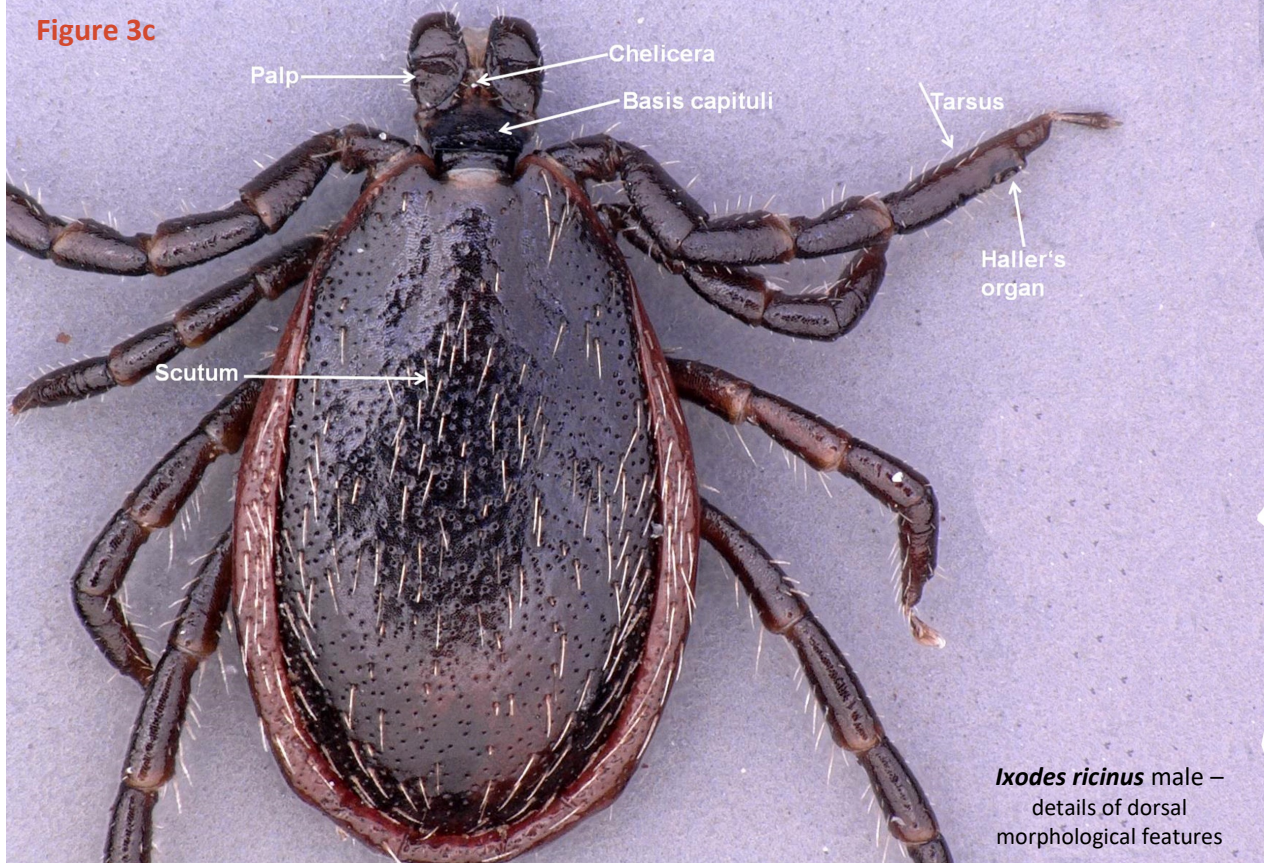
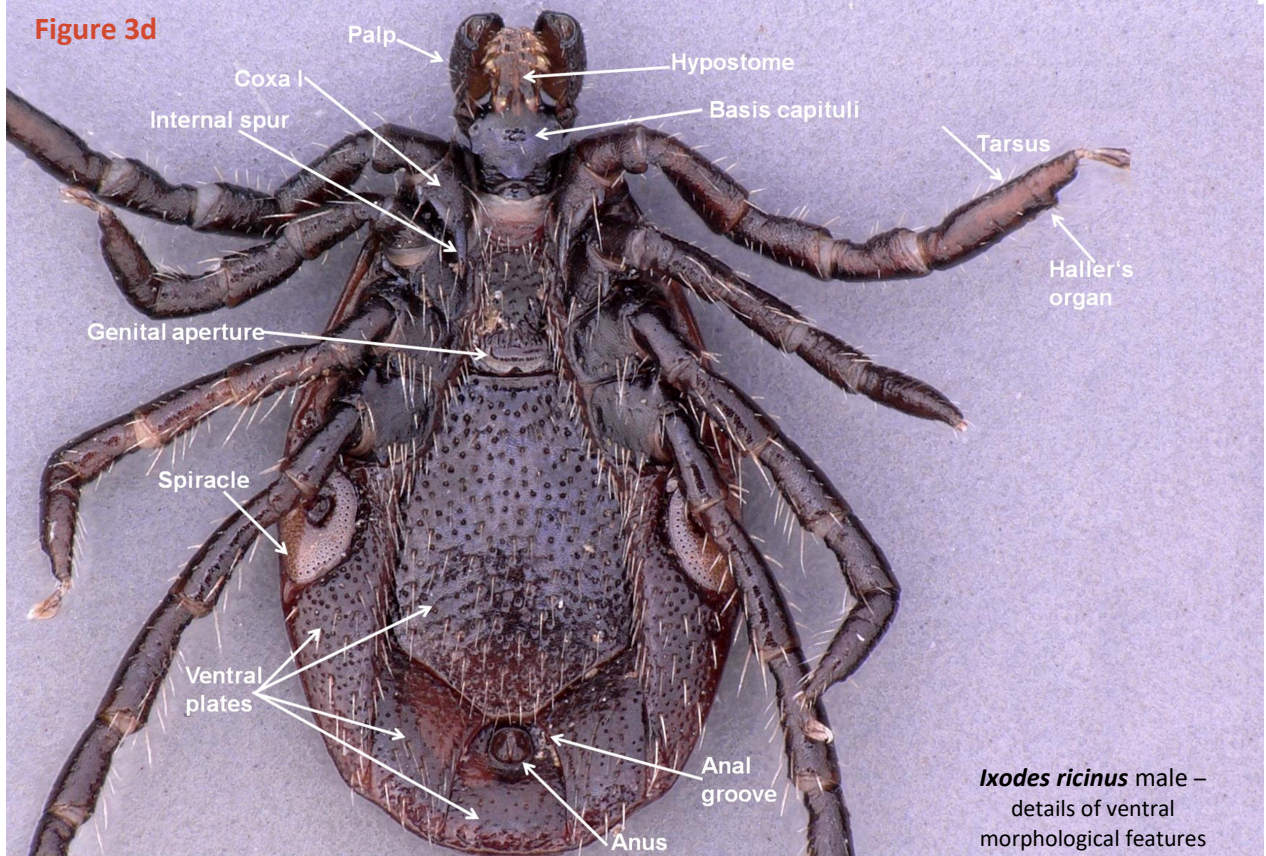
Figure 3c**Figure 3d**

Figure 4a



Ixodes ricinus larva – dorsal view

Figure 4b



Ixodes ricinus larva – ventral view

Figure 5a



Ixodes ricinus nymph – dorsal view

Figure 5b



Ixodes ricinus nymph – ventral view

Host specificity

Tick species can be either opportunistic or specific with respect to the hosts they choose; both *I. ricinus* and *I. persulcatus* are opportunistic species, especially the immatures. For *I. ricinus*, more than 300 species of vertebrate hosts have been recorded.^{9,28} Larvae and nymphs of *I. ricinus* feed readily on lizards, birds, and small mammals, as well as on larger hosts including deer. Adults feed on medium-sized and large mammals, especially ungulates, as well as humans, as do the immature ticks.⁹ *Ixodes persulcatus* is more restricted to 46 species of hosts.^{51,52}

Questing height is also important. Ticks questing on or near the ground are exposed mostly to small animals, while those questing higher in the vegetation are more likely to encounter larger animals. The extent to which different hosts are utilized depends on host behavior and opportunities for contact, such as foraging range, time of day and time spent foraging, habitats visited, and other factors.¹⁴

Acceptance of a vertebrate animal is also dependent on physiological factors and the ability of the ticks to recognize it as a host. Host utilization may be influenced by the ability of ticks to evade or suppress host homeostatic systems and avoid rejection.⁴⁷

Hard tick ecology, environmental factors

Ticks occur in many terrestrial habitats ranging from cool, arboreal northern forests to hot, arid deserts. Each species, however, has become adapted to the specific types of habitat where it is generally found in highest abundance. All *I. ricinus* postembryonic stages are exophilic and depend entirely on a suitable combination of climatic variables, making them vulnerable to climate changes and especially to desiccation. Thus, they are mainly found in cool, moist forests.^{42,53,54}

Water balance is a critical determinant of a tick's ability to wait for hosts. Ticks may quest for weeks or even months while waiting for a host. When they have a body water deficit, they retreat to more sheltered, humid microenvironments, such as the rotting vegetation in a meadow or damp leaf litter on the forest floor. They secrete a hygroscopic salivary secretion onto their external mouthparts that collects atmospheric water at relative humidity = 80-85% (active water vapor sorption).⁴² Rehydrated ticks are able to resume host-seeking. Some ticks are able to remain in the questing position for many days without rehydration, while others must return to their humid microenvironments.⁵⁵ Dense ecotonal vegetation provides shade, increased moisture, protection from intense solar radiation, and plants that support the tick hosts.

There have been various studies showing the relationship between *I. ricinus* and vegetation type in central Europe^{56,57} and the capacity of this species to adapt to a large variety of biotopes with low temperature (e.g., Sweden) and high altitudes, up to 1500 m.⁵⁸⁻⁶⁰

Ixodes persulcatus is distributed in 14 countries, between 21° and 66° of northern latitude in Eurasia, mostly with a temperate continental climate.⁵¹ In a model predicting the suitable habitats for *I. persulcatus*, it was shown that temperature and humidity are the main factors in the distribution of this species.⁵¹ Vegetation also has an impact on the tick distribution, its requirement is wood and wet biotopes.⁵¹

Normally, temperature and relative humidity in a burrow, cave, or similar type of shelter are more uniform throughout the year than in the external macro-environment. The higher relative humidity in such microenvironments is due in part to the presence of hosts, their wastes, and the plant materials they use to construct or line their nests.⁶¹ Nidicolous ticks exhibit behavioral patterns that restrict their distribution to these sheltered locations. They avoid bright sunlight and low humidity, the type of conditions prevailing at the entrances of burrows or caves. Confined within these hidden, restricted locations, nidicolous ticks become active when hosts are present. However, when the hosts are absent, they may wait for up to several years for hosts to return, or until they die of starvation.

Diapause

An important physiological trait that enables ticks to survive adverse environmental conditions and conserve energy until conditions improve is diapause as a form of dormancy.⁶² Diapause is induced by an external cue before adverse conditions occur. It is not terminated by favorable external conditions – as it is the case with quiescence – but there is some diapause development before its termination. During diapause ticks become inactive, reduce their metabolic rates, and do not feed on hosts even when given the opportunity.⁵³ Diapause can occur in each life stage, whether it is unfed or engorged. This varies, however, between species and can also differ within a tick species in different geographic areas. As an example, oviposition can be delayed in *D. marginatus*. Engorged females that feed in late summer, early fall or in winter oviposit only in the following spring.⁵³

Life cycle and seasonal activity

Ixodes persulcatus inhabits mainly coniferous forests of Asia and Eastern Europe, while *I. ricinus* inhabits deciduous and mixed forests in the British Isles, in Continental Europe, and western Asia.^{51,53,63} *Ixodes persulcatus* adult females and eggs are unable to survive the winter, however, that *I.*

Table 2. Animal hosts from which TBEV* has been recovered

Order/Family	Species	Virus type
Mammalia: Rodentia		
Muridae	<i>Apodemus agrarius</i> ^{86,87}	FES
	<i>Apodemus flavicollis</i> ^{81,88}	ES
	<i>Apodemus sylvaticus</i> ^{81,88}	ES
	<i>Apodemus speciosus</i> ⁸⁹	FES
	<i>Apodemus argenteus</i> ⁸⁹	FES
	<i>Myodes rufocanus</i> ⁸⁹	FES
	<i>Rattus norvegicus</i> ⁸⁹	FES
	<i>Microtus agrestis</i> ⁸⁷	ES
Cricetidae	<i>Microtus arvalis</i> ^{81,87}	ES
	<i>Myodes glareolus</i> ^{87,90,91}	ES
	<i>Myodes rufocanus</i> ⁸⁶	
	<i>Myodes rutilus</i> ⁸⁶	
Sciuridae	<i>Sciurus vulgaris</i> ^{81,92}	ES
Dipodidae	<i>Sicista betulina</i> ⁸⁶	
Eulipotyphlya		
Erinaceidae	<i>Erinaceus concolor</i> ⁹²	
	<i>Erinaceus roumanicus</i> ⁸¹	ES
Talpidae	<i>Talpa europaea</i> ⁹²	
Soricidae	<i>Sorex Araneus</i> ^{81,86}	ES
Goats	<i>Capra</i> sp. ⁹³⁻⁹⁵	
Sheep	<i>Ovis aries</i> ⁹³	
Bovidae	<i>Bos taurus</i> ⁹⁶	
Bison	<i>Bison bonasus</i> ⁹⁷	FES
Carnivora		
Canidae	<i>Vulpes vulpes</i> ⁹⁸⁻¹⁰⁰	
	<i>Canis familiaris</i> ¹⁰¹	FES
Mustelidae	<i>Mustela putorius</i> ¹⁰²	ES
Artiodactyla		
Cervidae	<i>Cervus elaphus</i> ^{103,104}	
	<i>Capreolus capreolus</i> ^{103,105}	
	<i>Alces alces</i> ¹⁰³	
Aves (families)**	Virus isolation ^{92, 106-108} : Passeriformes: Acrocephalidae, Bombycillidae, Corvidae, Emberizidae, Frigillidae, Hirundinidae, Laniidae, Motacillidae, Muscicapidae, Paridae, Passeridae, Psylloscopidae, Sittidae, Sturnidae, Sylviidae, Turdidae.	
	Others: Anatidae, Phasianidae, Picidae, Rallidae, Scolopacidae Transovarial transmission ⁵⁹ : Accipitridae, Charadriidae, Columbidae, Emberizidae, Laniidae, Troglodytidae, Turdidae	

ES, European subtype (TBEV-EU); FES, Far-Eastern subtype (TBEV-FE); SS, Siberian subtype (TBEV-Sib)

*Selected references; **Less information available

persulcatus larvae and nymphs, whether unfed or engorged, are able to overwinter. In contrast, eggs as well as unfed and satiated females of *I. ricinus* are capable of overwintering, a principal difference between the life-cycles of the two tick species. Vector tick activity is well correlated with the seasonal pattern of TBE occurrence. In such a focus, it is common for 2–3% of the ticks to be virus-infected.⁶³ In Northern and Central Europe, the seasonal activity of *I. ricinus* often has 2 peaks, one in spring (May–June) and the other one at the end of summer (September–October).

For *I. persulcatus* adults four types of seasonal dynamics throughout their distribution area were described, differing in the duration of the active period.^{63,64} In the north-western area of distribution, *I. persulcatus* becomes active immediately after the melting of the snow cover with a rapid increase in abundance in May, followed by a sharp decline in mid-summer. In Karelia (a middle taiga subzone), adult activity lasted on average 74 days. Between 2012 and 2023, the relative abundance of ticks increased significantly in comparison with the 1980s monitoring period, showing a tendency towards an earlier start of the tick activity, as in the 1980s.⁶³

Unfed *Dermacentor reticulatus* adults are mostly active in spring and autumn, occasionally in winter, but usually not in summer (June to early August).^{65–67} During periods of snow cover and the driest and hottest weeks of the year *Dermacentor reticulatus* is inactive.^{68–69} The larvae feed for 3–6 days, nymphs for 5–12 days, and females for 7–16 days, while males may remain in the host for a long time, even in the absence of females.^{51,52} Adults can overwinter unfed or engorged^{53,54} and are able to survive 2.5 years of starvation.⁵⁵ Interestingly, this tick can spend the whole winter on hosts.⁴⁹ *Dermacentor reticulatus* eggs can survive under water for several months and may be spread by floods into new areas.⁶⁶ *Dermacentor reticulatus* in immature life stages is assumed to be nidicolous and therefore cannot be collected from vegetation. Nevertheless, Schmuck et al.⁷⁰ collected *D. reticulatus* immatures (47 questing larvae and two nymphs) by flagging in June and July in 2018 and 2019, in two different locations close to the city of Leipzig, Germany. To understand under which circumstances *D. reticulatus* immatures were found outside the burrows of their hosts and can be collected from vegetation needs further investigation.⁷⁰

Tick species involved in TBEV transmission

Of the 54 species of ixodid ticks known from the Western Palearctic,⁵¹ eight species from three genera are known to be able to transmit TBEV, and the virus has been isolated from at least 14 other species (Table 1). *Ixodes ricinus*, the most commonly encountered European tick species, is considered to be the principal vector of TBEV there.⁷¹ Lichard and Kožuch²⁷ were able to show TBEV persistence

and transmission to white mice by *Ixodes arboricola*, which is considered a secondary amplifying vector of TBEV.⁷² *Ixodes persulcatus* is also known to transmit TBEV.^{73–75} It is the adult female *I. persulcatus*, which infects humans with TBEV and other zoonotic pathogens. Neither the larval nor the nymphal stage often attaches to humans.⁹ Both *D. marginatus* and *D. reticulatus* are also vectors of TBEV.^{24,65,76,77}

Haemaphysalis concinna is a known vector of TBEV as well.^{78,79} Evidence for the vectorial capacity of *Haemaphysalis inermis* for TBEV is available from Nosek et al.⁸⁰ The virus has been isolated in the Czech Republic from female and nymphal *I. hexagonus* infesting a hedgehog.^{29,81,82} TBEV also has been detected in *Haemaphysalis punctata*.⁸³

The role of *Dermacentor* ticks (Table 1) in the circulation of TBEV in the environment is unclear and poorly studied.⁶⁵ *D. reticulatus* appears to be spreading and population density increasing during recent decades.^{66,84,85} In eastern Poland, the mean prevalence of infection with TBEV found in questing adult *D. reticulatus* was 10.8% (range 7.3–14.3% in infected areas): This is considerably higher than the prevalence found in questing adult *I. ricinus* (1.6%, range 0.7–4.3% in infected areas).²¹

Prevalence of TBEV in questing adult *D. reticulatus* ticks from Białowieża Primeval Forest was similar (1.58%)²⁴ to that in questing *I. ricinus* (1.30%),⁷⁷ as was the case in Moldova (adult *I. ricinus* 3.8%, adult *D. reticulatus* 3.9%, but adult *Haemaphysalis punctata* 8.8%).⁸⁰ The natural occurrence of TBEV in a *D. reticulatus* tick population was also proven for Germany during 2016 to 2018 by isolation of several TBEV strains from this tick species in a natural focus.⁶⁵

The differences in TBEV prevalence in the various vector species remain puzzling. Questing *I. ricinus* usually have a very low prevalence of the virus, ranging from no virus in many areas to less than 1% in most others, and rarely reaching 2–5%, in unfed adults.^{111–114} Knap and Avsic-Zupanc¹¹⁵ showed that over a 4-year period, the prevalence was at the expected low level in the 8 areas studied, but that no area was consistently positive for the virus. This may be related to the frequently low sample sizes (14/30 samples had fewer than 300 specimens).

Prevalence of the virus in feeding ticks, although very variable, can be substantially higher.¹¹⁶ Waldenström et al.¹¹⁷ showed a low prevalence (0.5%) in nymphs and larvae feeding on migratory birds in Sweden, while Kazarina et al.¹⁰⁶ detected 14% nymphs and 7% larvae of *I. ricinus* on migratory birds infected in Latvia. Data for *I. persulcatus* are more variable. Korenberg and Kovalevskii¹¹⁸ reported a high TBEV prevalence in unfed adults, ranging from 10.9% to 38.7% over 6 years (mean 26.2%) in unfed adults in the Pre-

Ural Region, whereas the prevalence in the Primorski Region of the Russian Far-East ranged from a little over 1% to over 9% from 1970 to 1990, and in the Khabarovsk Region from 3.4% to 9.4% over 4 years.¹¹⁹ In the Novosibirsk Region, the prevalence of TBEV in unfed adult *I. persulcatus* was 3.6%, with 0.8% being pathogenic to laboratory mice.⁸⁶ In the same study, 3.3% of questing adult *I. pavlovskyi* were infected with the virus with 1.8% of the isolates being pathogenic. Information on less commonly encountered vectors is rarely available and sample sizes are usually low, making such data unreliable (e.g., Kim et al.)¹²⁰ Long-term studies and statistical analyses showed that higher average temperatures during the summer-autumn period may lead to higher levels of TBEV found in ticks and consequently increase the risk for humans to develop symptomatic TBE following an infected tick bite.¹²¹

Vertebrate hosts

The prevalence of antibodies to TBEV in hosts is quite variable.¹²² TBEV has been found in numerous mammal species from different families, as well as in a large number of passerine and non-passerine bird species (Table 2). Virus infection was demonstrated by antibodies to the virus or viral ribonucleic acid (RNA) detection in a wide variety of bird species,^{117,123} with virus isolation from *Turdus pilaris* (fieldfare) and *Acrocephalus dumetorum* (Blyth's reed warbler) opening the possibility of virus transfer to new foci during bird dispersal or migration. Viremia has been induced experimentally in birds, reaching levels sufficient to infect feeding ticks.^{123,124} Generally speaking, findings of TBEV in animals, whether indirect or direct, do not mean very much eco-epidemiologically. Only the demonstration of reservoir competence indicates an active role in the perpetuation of TBEV.

Red foxes (*Vulpes vulpes*) are known to be reservoir-competent for TBEV.^{99,125} Although *I. hexagonus* is a proven vector of TBEV, little is known about the vector competence of the fox tick *I. canisuga*.

In recent years, the detection of viral RNA in hosts has become possible. Tonteri et al.,¹²⁶ in Finland, detected the European (TBEV-EU) and Siberian (TBEV-Sib) subtypes in *M. glareolus*, TBEV-Sib in the shrew *Sorex araneus*, and TBEV-EU in *Microtus agrestis*. Achazi et al.⁸⁷ detected TBEV RNA in rodent brain tissue in prevalences up to 20% in TBE non-risk as well as in risk areas in east-German Federal States. In the Novosibirsk region of Siberia, where *I. persulcatus* and *I. pavlovskyi* are the main TBEV vectors, the prevalence of TBEV viral RNA in five small mammal species was extremely high.⁸⁶ It ranged from 35.3% for *A. agrarius* organs to 82.2% for *Myodes rutilus* blood, with a mean value for all species and tissues of 62.1%. All 3 virus subtypes were represented. In addition to small mammal hosts, larger wild and domestic animals frequently have high antibody prevalences. Because they feed large numbers of vector

ticks, they can be used as sentinels for the occurrence of TBEV in a given area.

TBEV transmission

Nuttall et al.¹²⁷ noted: "Reciprocal interactions of parasites transmitted by blood-sucking arthropod vectors have been studied primarily at the parasite-host and parasite-vector interface. The third component of this parasite triangle, the vector-host interface, has been largely ignored."

The adult female tick is considered to play only a minor role in virus circulation. Tick males, which either do not feed or feed for only a short time, might also be involved in virus transmission.¹²⁸ TBEV invades all tick tissues, including the salivary glands and ovaries,¹⁶ thus it may be transmitted by ticks in the following ways: 1) via saliva, 2) transovarially (vertically), and 3) sexually.¹²⁸⁻¹³⁰

TBEV transmission from vector ticks to hosts via saliva

Certain species of ticks are vectors and reservoirs of TBEV, and they can transmit the virus already when they start feeding^{131,132} with viral particles contained in the saliva, which the ticks release into the host tissues.¹³³ Tick-borne virus transmission occurs within 24 h of tick attachment, possibly within the first hour of feeding (Ebel et al. 2004). After attachment, many ixodid ticks secrete cement to fix themselves in the skin, but in a study Alekseev et al. (1996) showed that cement plug may act as bolus of infection for TBEV and *Borrelia burgdorferi* s.l. When an infected tick had been removed, the remaining cement cone embedded in the skin resulted in TBEV infection within one hour of detachment.¹⁸

TBEV is present in the alveolar cells of the salivary glands of *D. marginatus* and *H. inermis* females in as few as five days after their feeding on viremic white mice.¹³⁵ Also certain vertebrates, so-called reservoir hosts, are important for the amplification of the virus and are together with vector ticks the basis for the heteroxenous TBEV perpetuation.¹³⁶

Viremic transmission from hosts to feeding ticks

Ticks become infected with TBEV while they feed on a viremic host.^{63,137} Nosek et al.^{138,139} proved that a viremia in a host lower than 10^1 mouse LD₅₀/0.03 ml was insufficient to cause infection in ticks. In individual engorged *I. ricinus* ticks, the virus titer was 10^1 - 10^4 mouse LD₅₀/0.03 ml. Viremic white mice served as virus donors. Grešíková and Nosek⁸⁰ demonstrated the persistence of TBEV in *H. inermis* (from larva to nymph) and then the transmission from *H. inermis* nymphs to white mice. Viremia surpassing the threshold values of infectivity for tick vectors was also found in some juvenile and adult *Myodes rufocanus*, *M. rutilus*, and *Micromys minutus*. The viremia level depends

on the rodent species and age, and exhibits individual variability.¹⁴⁰

Co-feeding transmission

TBEV transmission is also possible from infected to non-infected ticks during feeding close to each other on a non-viremic host.^{141,142} Cellular infiltration of tick feeding sites, and the migration of cells from such sites, can provide a vehicle for transmission between co-feeding ticks that is independent of host viremia.¹⁴³ The non-viremic route of transmission between co-feeding ticks can even occur in rodents that are already immune to TBEV.¹⁴⁴ The degree of co-feeding virus transmission may be influenced by local climatic factors that affect the seasonal timing of tick host-seeking activity and, as such, can be used to predict the focal distribution of TBEV.^{141,142}

Transovarial transmission

Another possible way for ticks to transmit TBEV involves transovarial transmission and transstadial persistence (see below), which were described for the first time as early as 1940.¹⁴⁵ However, only some eggs in the batch of a TBEV-infected vector tick female become infected.¹³⁰ In addition, virus can partly be lost during transition from stage to stage,¹⁴⁶ and not all tick individuals reach the next life stage irrespective of the presence or absence of the pathogen. Danielova and Holubova¹³⁰ found that only 0.23% of larvae coming from infected females were TBEV-positive. Other studies showed that 0.58% to 0.75% of the larvae were transovarially infected. Thus, the rate of transovarial transmission remains below 1%. Nuttall et al.¹⁴⁷ suggest that transovarial transmission is important for the maintenance of a natural focus even if it occurs at a very low rate.

Danielova et al.¹⁴⁸ detected TBEV in 2 out of 647 flagged larvae of *I. ricinus*, which indicates transovarial transmission.

Transstadial persistence

TBEV was not detectable in *I. ricinus* nymphs 14 days after molting from larvae that had engorged on viremic *A. flavicollis*, but TBEV was present in these ticks two months post ecdysis. Many nymphs contained the virus, indicating that the latter undergoes an eclipse phase during metamorphosis.¹⁴⁹

Sexual transmission in ticks

Transmission of TBEV from males to females¹²⁸ is successful in only 10% of copulations in infected *I. persulcatus*, but it may provide notable support for the transfer of the virus to the following generation of ticks if transovarial transmission follows. A mathematical model of sexual transmission of

the virus¹⁵⁰ was developed long before determining that such a sort of transmission occurs. Virus exchange between a non-engorged female and an infected male of *I. persulcatus* that ‘feeds’ on (i.e., attaches to) the female before or after copulation is quite probable, and it has been proven that the saliva of starved males contains a fairly large amount of virus, sufficient for infecting not only animals but also humans. The feeding of *I. persulcatus* males on females with which they later copulate can be observed in 2–10% of cases.^{151,152}

Vertical TBEV transmission in vertebrates

TBEV transmission from mother to her offspring in small rodents, e.g., red voles (*M. rutilus*), was shown for naturally infected reservoir hosts as well as after experimental infection with different sublethal doses of the virus.¹⁵³ TBEV RNA was detected in up to 90% of the newborn rodents, 240–280 days after experimental infection of their parents, by real-time polymerase chain reaction (RT-PCR), enzyme-linked immunosorbent assay (ELISA), and bioassays. The small amounts of TBEV RNA detected in the embryos, placenta, and blood serve as evidence of prenatal transmission. Postnatal transfer of the virus might occur through the rodent’s milk. Vertical virus transmission may occur before, during, and/or after birth of the baby rodents with a high frequency. In natural foci, this could ensure long-term persistence of TBEV in mammal hosts¹⁴⁹ without involving any arthropod vectors.⁸⁶ Divé et al.¹⁵⁴ reported detailed investigation of pre- and postnatal health assessment of three children in the context of severe maternal TBEV infection during pregnancy. The clinical and virological data strongly suggest that fetal TBEV infection did not occur, despite severe manifestations in the pregnant females. Non-reservoir hosts do not directly participate in virus transmission, but can play an important role in the maintenance of natural foci. The density of reservoir-incompetent hosts may have either a positive effect on virus transmission, by amplifying the tick population, or a negative (‘dilution’) effect, as tick bites on a non-reservoir host cannot lead to virus transmission.^{155,156}

Alimentary route of transmission

Humans mostly become infected with TBEV via tick bites, but viral transmission is also possible via the consumption of unpasteurized goat, cow and sheep milk.¹⁵⁷ Approximately 1% of all TBEV infections in humans are probably acquired by consuming infected unpasteurized milk and milk products from infected livestock, particularly goats.¹⁵⁸

Outbreaks due to alimentary virus transmission are known from Eastern, Central and Southern Europe, and have to be considered particularly in cases of local epidemics.¹⁶²⁻¹⁶⁴ Ličková et al.¹⁶⁵ summarize the history and recent

alimentary TBEV infections in Europe. In an alimentary outbreak in Germany, due to consumption of a fresh goat cheese, the virus could be for the first time isolated from naturally infected cheese.¹⁶⁴

TBEV interhuman transmission of TBEV by breast milk has not been confirmed or ruled out. Kerlik et al.¹⁶⁶ reported a case of probable transmission of TBEV from an unvaccinated mother to an infant through breast-feeding.

The natural cycle

The natural cycle of TBEV is highly complex, and many details remain obscure. The three prevailing TBEV subtypes overlap in some areas, they all have multiple mammalian reservoir hosts and various tick vectors, and in some areas these subtypes occur sympatrically. Humans are not included in these natural cycles, but may enter those transmission cycles inadvertently.

Small mammals as a reservoir and vector ticks play a central role in the natural cycle of TBEV, but non-reservoir hosts such as birds and large vertebrates, such as wild ungulate species, or foxes, may also indirectly contribute to the spread and maintenance of TBEV. Additionally, changing climatic patterns, as well as changes in ecosystems, may not only affect the spatial distribution of TBEV, but also the maintenance of small natural TBEV foci.^{167,168} Small rodents such as *A. flavicollis* are important hosts for the larvae of *I. ricinus*, the probably most important TBEV amplifying host in Central Europe. *Apodemus flavicollis* temporarily develops high virus titers necessary to infect ticks. Detailed studies by Radda et al.,^{98,149} who trapped small rodents and collected the engorged ticks in a natural TBE focus for 2 years, showed that given certain prerequisites are fulfilled (high numbers of rodents, vector tick larvae and nymphs feeding on these rodents), such a natural TBEV focus is able to sustain itself without any significant input of other hosts. This may explain why many of these natural foci are stable, but restricted to small areas, and why they harbor TBEV-positive ticks over a long period of time. Forest structure, especially deforestation and reforestation, are known to have a huge impact on ticks and vertebrate reservoir hosts for many tick-borne pathogens.^{159,160}

Experimental transstadial maintenance of TBEV in *D. marginatus* and *D. reticulatus* ticks emphasizes the role of both species. TBEV infection and transmission rates in *Dermacentor* species to hosts are somewhat lower than in species of the genera *Ixodes* and *Haemaphysalis*.¹⁶¹ Feeding larvae and nymphs of *I. persulcatus* may become infected with TBEV if the virus titer in the host blood reaches at least 3.0 log₁₀ LD₅₀/0.03 mL.¹⁵¹ Such levels of viremia occur only in small rodents and are a critical factor in the virus circulation between vertebrates and ticks in natural foci. In small rodents, the infection is asymptomatic.⁹⁰

TBEV has been isolated from a wide range of birds from many different families, including migratory species, which may be important for the distribution of the virus. A common strategy for migratory birds is to rest at certain stopover sites along their routes. At these sites, the birds can be infested with ticks or engorged ticks can detach after engorgement. Sándor et al.¹⁶⁹ detected 4 different tick species on 11 different bird species in the Danube Delta, including larvae, nymphs, and females of *I. ricinus*.

A high variability is found between areas and years with respect to viral prevalence in both vertebrate hosts and vector tick populations, while consistent differences between vectors. For example the generally higher TBEV prevalences in *I. persulcatus* compared with those in *I. ricinus* may relate to the ecology/biology of the individual vectors. The complexity is well defined by the various mathematical models aimed at exploring the dynamics of TBEV ecology.^{156,170,171} Hartemink et al.¹⁷¹ list 19 parameters based on field data to define the basic reproduction number (R_0) of tick-borne infections, while Rosà et al.¹⁵⁶ list 32 parameters in a more comprehensive model. Unfortunately, no single study has been able to comprehensively measure all the parameters needed to test these models, although approximations are available.

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