Review

Global impact of Torque teno virus infection in wild and domesticated animals

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Abstract

Infection with Torque teno viruses (TTVs) is not restricted to humans. Different domestic and wild animal species are naturally infected with species-specific TTVs worldwide. Due to the global spread of the infection, it is likely that essentially all animals are naturally infected with species-specific TTVs, and that co-evolution of TTVs with their hosts probably occurred. Although TTVs are potentially related to many diseases, the evidence of the widespread infection in healthy human and nonhuman hosts raised doubts about their pathogenic potential. Nonetheless, their role as superimposed agents of other diseases or as triggers for impairment of immune surveillance is currently under debate. The possible contribution of animal TT viruses to interspecies transmission and their role as zoonotic agents are currently topics of discussion.

Key words: TT virus; anellovirus; animals; ssDNA; epidemiology.

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Introduction

Anelloviruses (AVs) are a large group of small, non-enveloped viruses with a circular, single-stranded DNA genome of variable size, ranging from 2.1 to 3.8 kb [1-3]. Because of their high degree of genetic heterogeneity, AVs are divided into at least 11 genera, each of them including a various number of species [4]. AVs are ubiquitous and highly prevalent worldwide [5,6]. They infect their hosts with multiple and often divergent strains [7,8] and are transmitted through multiple routes [9-15]. Infection with AVs probably occurs early in life [16] and leads to a progressive persistent infection of the natural host [17]. Human Torque teno virus (TTV) is the prototype of the Anelloviridae family, but a number of related viruses have been described in other mammalian hosts. Human TTV was originally reported in 1997 in a Japanese patient with post-transfusion, non-A-G hepatitis [18]. Since its discovery, a series of related human AVs have been characterized [19]. Furthermore, a series of closely related TTV-like viruses, and two distinct genera comprising viruses with smaller genome size and similar genetic organization, but only partial sequence homology, designated Torque teno midi virus (TTMDV) and Torque teno mini virus (TTMV), have been characterized [20-22]. A large number of AV species has also been reported in wild and domestic animals, including non-human primates, wild boars, badgers, pine martens, tupaias, rodents, bats, sea turtles, sea lions, livestock (pigs, sheep, cattle, camels, and poultry), and companion animals (cats and dogs) [3,5,23-29]. Indeed, since to date the presence of TTVs infecting mammalian species has been poorly investigated, it would not be surprising if all mammals were infected with species-specific TTVs. Human TTVs and TTVs infecting mammalian species have similar genomic and transcriptional organization, suggesting that co-evolution of TTVs with their hosts probably occurred [30]. However, AVs from different species share a low sequence homology, with genetic diversity varying between 35% (within genus) and 56% (among genera) [19]. In particular, AVs infecting animal species are highly divergent from those detected in humans, both in sequence composition and genome length, although phylogenetically some animal TTVs cluster with the human TTVs [31]. Despite over 17 years of investigation, little is known about the pathogenesis and possible disease associations with AV infections, in part due to the lack of a robust cell culture system for viral replication or a suitable small-animal model.
Discovery and characterization of TTV

In 1997, Nishizawa et al., using representational difference analysis, isolated a sequence of 500 nucleotides from the blood of a patient suffering from a form of post-transfusion hepatitis of unknown etiology. The representative clone (N22) was found to originate from the genome of a non-enveloped, single-stranded DNA virus, provisionally named TTV, the initials of the index patient [1,18]. Subsequent cloning and sequencing experiments conducted by Okamoto et al. led to the characterization of a linear genome of 3,739 kb, called TA278 [1]. In 1999, the presence of a GC-rich sequence of approximately 120 nt was reported, allowing the recognition of the circular nature of the TTV genome with negative polarity [32]. The genomic full-length of the TA278 isolate was finally determined to be 3,853 nt, and the presence of the unique stem-and-loop structure in the GC-rich region was demonstrated [1,2]. Based on this and other structural and molecular characteristics that this virus shared with members of the family Circoviridae, and in particular with chicken anemia virus (CAV), TTV was initially recognized as the first human circovirus [32]. However, further studies showed no significant sequence identity between TTV and circovirus, and the virus was considered as the only representative of the new family Circinoviridae, from Latin circinatio, "circle" [2]. Shortly after the discovery, viral isolates were identified with genetic and structural characteristics similar to TTV, but with considerable genetic variability; among these, isolates PMV, SANBAN, and YONBAN [21,33] showed a nucleotide divergence greater than 50% compared to the prototype TA278. Subsequently, shorter sequences were isolated with genomic features intermediate between TTV and CAV; their characterization led to the identification of a new group of viruses named TTV-like mini virus (TTLMV, TTMV) [21]. Finally, in 2005, the International Committee on Taxonomy of Viruses (ICTV) proposed to classify TTMV and TTV into the new genus Anellovirus, and to assign to the acronym the meaning of Torque teno virus, from the Latin torque (necklace) and tenuis (narrow), referring to the characteristics of their genome [34]. TTV became the prototype of a large group of viral agents with similar genomic organization but a low sequence homology, isolated both in humans and in animals. In fact, in the following years, similar viruses were isolated in other species of vertebrates, including non-human primates (chimpanzees, macaques, tamarin monkeys, and douroucouli), pets (dogs and cats), livestock (pigs, cattle, sheep, camels, and poultry), and wild boars, badgers, pine martens, tupaias, rodents, bats, sea turtles, and sea lions [3,5,23-29,35]. The natural infection of pigs was first described by Leary et al. in 1999 [5]; the first complete genome of Torque teno sus virus (TTSuV) was described by Okamoto in 2002 [3]. In 17 years since discovery, over 200 complete or partial sequences of viral genomes like TTV were characterized in humans and more than 10 species of vertebrates. In 2009, the ICTV established the new family Anelloviridae, divided it into 11 genera, each one with a different number of species. Three other genera, Mutorquevirus, Nutorquevirus, and Xitorquevirus, which include newly discovered isolates, are currently under consideration by the ICTV for inclusion within the family (Table 1). Human TTVs belong to the genus Alphatorquevirus, which includes 29 species. In addition to TTV, in humans there are two other TTV-like species: TTMV and TTMDV, classified within the family Anelloviridae and belonging to the genera Betatorquevirus and Gammatorquevirus, respectively. TTVs infecting pigs (Sus domesticus) include four species described so far: Torque teno sus virus 1a and 1b (TTSuV1a and TTSuV1b), belonging to the genus Iotatorquevirus,

| Genus            | Species | Host                  |
|------------------|---------|-----------------------|
| Alphatorquevirus | 29      | Humans, chimpanzees   |
| Betatorquevirus  | 12      | Humans, nonhuman primates |
| Deltatorquevirus | 1       | Tupaia                |
| Epsilontorquevirus | 1    | Tamarin               |
| Etatorquevirus   | 2       | Cats                  |
| Gammatorquevirus | 15      | Humans, chimpanzees   |
| Iotatorquevirus  | 2       | Swine                 |
| Kappatorquevirus | 2       | Swine                 |
| Lambdatorquevirus | 1     | Sea lions             |
| Thetatorquevirus | 1       | Dogs                  |
| Zetatorquevirus  | 1       | Douroucouli           |
and *Torque teno sus virus* k2a and k2b (TTSuVk2a and TTSuVk2b), the latter being the most recently discovered species, belonging to the genus *Kappatorquevirus* [36].

**Structure and genomic organization of TTV**

The virus appears as a spherical particle with icosahedral symmetry and a diameter of 30–50 nm. The buoyant density in cesium chloride was found to be 1.31–1.33 g/cm³ for TTV in serum and 1.33–1.35 g/cm³ for TTV in feces [2]. Resistance to treatment with detergents, solvents, and dry heat confirmed the absence of a lipid envelope [37,38]. TTV proteins have not been well characterized yet; however, similar to CAV, it is believed that the capsid is made up of a single structural protein. Sensitivity to DNase I, mung bean nuclease, and RNase A, and resistance to restriction enzyme NdeI have proven the single-stranded nature of the viral DNA [2,37]. Hybridization with positive-sense RNA molecules and subsequent digestion has demonstrated the negative polarity of the viral DNA [2]. DNA size varies from 2.1 to 3.9 kb, depending on the host species [2,7,18,32]; in general, the sizes of the animal AVs are smaller than those of human AVs. TTVs infecting humans and chimps have a genome size ranging from 3.7 to 3.9 kb; the genome size of TTSuV (approximately 2.8 kb) is intermediate between the size of human TTV and the size of the smallest AV identified so far (< 2.1 kb), isolated in the cat [3]. Despite the variability in sequences and size, the various species of animal and human TTVs show similar genomic organization [3,24,39]. The genome consists of a coding region that contains at least three open reading frames (ORFs), and a non-translated region (UTR) of about 1.2 kb (0.8 kb in TTSuV), containing a GC-rich tract [40]. The highly conserved UTR region contains the TATA box and the polyA tail sequences and it is used as a transcription factor (Figure 1). The messenger RNAs (mRNAs) analyzed were transcribed from a plasmid containing the complete genome construct of TTV in COS1 cells [41]. Three spliced mRNAs of 3.0 kb, 1.2 kb, and 1.0 kb with common 5'-and 3'-termini were recovered, and it was demonstrated that the splicing sites link distant ORFs to create two new ORFs capable of encoding 286 and 289 amino acids, respectively [41]. ORF1 encodes a product of about 700–770 amino acids, which is considered the viral capsid protein [3]. ORF2 codes for products of about 200 amino acids and in some genotypes it is divided into smaller ORF2a and ORF2b, the latter containing a CAV-like conserved amino acid motif, similar to protein-

**Replication of TTV**

The distribution of TTV in the tissues of the natural hosts and their detection in a wide variety of biological samples suggest that AVs are able to replicate in different cell types, recognizing receptors distributed across different tissues [45-47]. For DNA replication, similar to circovirus and in analogy to the plant and bacterial viruses with circular single-stranded DNA, a rolling circle mechanism of replication has been proposed, leading to the formation of a circular, double-stranded intermediate [2]. As a DNA polymerase-coding sequence has never been demonstrated, it is believed that AVs, like most small DNA viruses, replicate their DNA using the DNA polymerase and the replication machinery of the host cell [48]. Replicating forms of the TTV genome have been detected in a variety of tissues, including lung, stimulated peripheral blood mononuclear cells, bone marrow, spleen, liver, pancreas, kidney, thyroid gland, and lymph nodes [45,49,50]. Furthermore, TTV was detected in the serum of most infected hosts. In bone marrow-derived cells, three types of mRNAs of TTV

![Genomic organization of TTV](Image 1)
(2.9 kb, 1.2 kb, and 1.0 kb) have been detected and produced by alternative splicing, as well as by transfecting African green monkey cells with whole TTV genome; in this case, at least six proteins derived from three distinct mRNAs, expressed from two different start codons, have been produced.

**Genetic variability of TTV**

Using conserved UTR-specific primers for detection of TTV DNA, several variants with high genetic diversity have been detected [7,21,32,35,51,52]. Human TTVs belonging to the genus *Alphatorquevirus* are divided into at least 29 species and more than 40 genotypes and 70 subtypes, with nucleotide differences exceeding 50% among species, more than 30% between genotypes and between 15% and 29% among subtypes. The sequence homology between human and swine TTV is below 45% [3,39,53]. Sequence homology between the TTSuV1 and TTSuV2 species was estimated to be around 56%, nucleotide divergence being higher among isolates of TTSuV1 (30%) compared to TTSuV2 isolates (15%). Genetic variability is greater in ORF1, where at least three hypervariable regions characterized by several insertions and deletions and amino acid diversity among TTV isolates higher than 70% have been described. In TTV isolates belonging to the same genotype and in isolates belonging to different genotypes, at least 19 recombination sites have been identified, 13 of them located in the UTR [54]. Co-infection with strains of different genotypes is very frequent in humans [8,55], as is co-infection with viral subtypes and multiple species of TTSuV in swine [56]. Finally, it is possible that the interspecies transmission of TTVs between humans and livestock or pet animals may contribute to the genetic variability of the virus [30].

**Animal TTVs**

TTV infection is not restricted to humans. Using highly conserved primers derived from the UTR of the TTV genome, a variety of TTV-like viruses have been detected in non-human primates and tupaias (tree shrews, *Tupaia belangeri chinensis*) [5,35,57]. The entire nucleotide sequences of species-specific TTVs that infect nonhuman primates, such as the chimpanzee (*Pan troglodytes*), Japanese macaque (*Macaca fuscata*), cotton-top tamarin (*Saguinus oedipus*), and douroucouli (*Aotes trivirgatus*) have been determined [24,35]. Human and simian TTVs share closely related genome organization and presumed transcriptional profiles, showing nearly 85% sequence similarity. However, it is now well established that TTV variants in nonhuman primates are species-specific [5,23,24,35,58] and that TTVs from macaques and tamarins are increasingly divergent from TTV variants infecting humans and chimpanzees [35]. Recent evidence shows that simian TTV can infect humans, as it has been observed that approximately 10% of Japanese patients with liver diseases are infected with simian TTV, although the mode of transmission of the infection from animals to humans has not been demonstrated [59]. Furthermore, TTV DNA has been detected in serum samples obtained from wild and domesticated animals, including chickens, pigs, wild boars, camels, cats, dogs, pine martens, sea lions, cows, and sheep [3,5]. Even more recently, Nishiyama et al. reported the identification of diverse anelloviruses in several species of wild rodents; the viruses are highly prevalent in wood mice (*Apodemus sylvaticus*) and field voles (*Microtus agrestis*), detectable at a low frequency in bank voles (*Myodes glareolus*), but absent from house mice (*Mus musculus*) [28]. The presence of TTV in dogs (now named *Torque teno canis virus*) has been documented, and the full-length nucleotide sequence of Japanese and Chinese isolate strains has also been reported; results indicate that dogs are naturally infected with species-specific TTVs with small genomic sizes and suggest a global distribution of TTVs with extremely divergent genomic sequences and lengths in animals [3,60]. In Italian dogs, the presence of *Torque teno canis virus* has recently been checked for, and about 10% of the animals were found to be infected with highly divergent variants (unpublished data). Indeed, since TTV has been detected in different animals, it might be considered as a zoonotic infection. Hence, it is possible that the virus can be transmitted from humans to animals and *vice versa*. Many recent studies have focused on TTVs that infect pigs (TTSuV). The AVs that infect swine, domestic pigs, and wild boars are spread all over the world, regardless of the race, age, gender, and state of health of the animals [30,61-63]. A recent retrospective study has documented the circulation of the virus in Spanish farms since 1985 [64]. Globalization and international trade in live animals have been proven to play a crucial role in the spread of TTSuV [65]. In addition, the vast spread of the infection in swine and the absence of geographical clustering of the isolates have led some authors to speculate that the dissemination on a planetary scale may be partly due to the use of contaminated vaccines [13]. The global prevalence of infection varies
between 16.8% and 100% for TTSuV1 [66-68], and between 31% and 90% for TTSuV2 [63,66,69]. Data related to infection with TTSuVk2b, the most recently identified species, are still limited. A study on 244 pig serum samples from 17 different countries reported an overall 40% prevalence of infection [36], whereas Blois et al. reported an 11.5% prevalence of TTSuVk2b in 721 Italian pigs [70]. The high prevalence of TTSuV in the swine population indicates the existence of effective mechanisms of transmission. Although the fecal-oral route is considered the primary mode of transmission, other ways may contribute to the dissemination of the virus. TTSuVs have been found in the feces and in the nasal secretions of piglets in their first week of life; the prevalence of infection increases with age, and the virus is shed more through nasal secretions than through fecal elimination [71,72]. TTSuVs are also found in seminal fluid, indicating the possibility of sexual transmission of infection [73]. TTSuV DNA has also been found in colostrum and stillborn piglets, indicating that vertical and transplacental or intra-uterine transmission may contribute to the dissemination of the virus [14]. So, infection with TTSuV occurs early in life and leads to a progressive persistent infection, with viremia levels increasing with the age of the animals. Most tissues and pig organs, if not all (brain, lungs, kidneys, bone marrow, heart, spleen, and mesenteric and mediastinal lymph nodes), test positive at five weeks of age, reaching the highest prevalence at slaughter [46]. In particular, lymphoid tissues and T lymphocytes are hypothesized to be the target site and the target cells for TTSuV, respectively [74]. As in humans, TTV shows a high degree of genetic variability in the swine population, and the presence of co-infection with multiple species and viral subtypes is a frequent event [39,56,75,76]. The production of antibodies elicited against the viral capsid protein, which is associated with a reduction in viremia, has been demonstrated [77], but these antibodies seem unable to eradicate the infection, which in part could explain the persistence of the virus in the animal host. Finally, the intra-uterine infection of the fetus before the acquisition of mature immunocompetence could explain the establishment of a state of immunotolerance against TTSuV [46].

**Innocent bystanders or something else?**

AVs have been known to exist for almost two decades and, although much is known about their epidemiology, they are orphan viruses still in search of a disease. The high prevalence of infection in the general population and the capability to establish persistent infections raised doubts about their actual pathogenic potential, and some researchers have considered TTV a component of the human microbiota [78]. Although TTV has been potentially related to many diseases, there are only a few reports supporting the disease-inducing potential of TTV. A series of studies suggest a possible involvement of human TTVs in the pathogenesis of certain diseases, such as hepatitis [79], hematological disorders [80], respiratory diseases [12], rheumatic autoimmune diseases [81], and various malignant disorders. Regarding AVs infecting animals, most studies to date have focused on the role of TTSuV in pig diseases. Although TTSuVs were found at particularly high frequency in healthy animals [70,72], they are currently receiving more attention due to the latest results on disease association. In fact, TTSuVs are considered non-pathogenic by themselves, but there is increasing evidence that points to their influence on the development of some diseases or suggests that these viruses even affect disease outcome [66]. In particular, the co-infection with porcine circovirus type 2 (PCV2) and the associated porcine circovirus diseases deserve special attention [61]. First evidence highlighted the high prevalence of TTSuV2 infection in postweaning multisystemic syndrome (PMWS), although in most cases, such co-infection does not lead to a disease; however, it is subclinical, suggesting that the disease association is possibly a matter of viral load [66]. Recent reports showed that porcine TTV partially contributed to inducing porcine reproductive and respiratory syndrome, porcine dermatitis and nephropathy syndrome, and hepatitis [82,83]. The latest research indicating that TTSuV2 but not TTSuV1 seems to benefit from the host’s disease status suggests that TTSuVs may act as triggers for diseases that cause immune system impairment. In this context, the level of TTSuV2 viremia may be associated with the level of immunocompetence of the animals, resulting in uncontrolled in PMWS animals, which are known to be immunocompromised [66]. Similarly, TTV DNA load in infected humans appears to correlate with the level of immune competence of the host [84,85]. Some authors have recently suggested that the expansion of members of AVs in the human microbiome and the increasing viral load in plasma during immunosuppressive therapy could be used to predict and monitor immune competence [86]. The association between the single infection with TTSuV or the co-infection with PCV2 and abortion in pigs has not been confirmed [16]. However, a recent
study carried out using pigs infected with the hepatitis E virus has shown a correlation between TTSuV and the increased risk of developing severe hepatitis in animals co-infected with PCV2 [87]. More recently, a high prevalence of TTSuV1, but not TTSuV2, in pigs suffering from porcine respiratory disease complex has been shown [88]. As for other porcine pathogens (PCV), it is therefore possible to speculate that the TTSuV species are distinct not only from the genetic point of view, but also for their different pathogenic potential.

Conclusions

Since their identification about 20 years ago, many studies have been conducted on human and animal TTVs, in order to assess the extent of dissemination of the infection and the pathogenic potential of the viruses [30,89]. It is likely that such viruses would be considered as components of the host microbiota and, as such, unable to cause disease directly, but rather available to be engaged in physiological processes and to modulate the response of the organism to other pathogens. It was recently suggested that the dynamics of replication of human TTVs can correlate with the immune competence of the host, and that the viral load can be considered as a possible biomarker of immune responsiveness [84]. It has been also proposed that animal AVs, especially TTSuVs, could be utilized as a model to evaluate the dynamics and the effects of global trade on viral heterogeneity, and to understand how live animal movement affects virus evolution [65]. Furthermore, it was hypothesized that TTV could be a more appropriate indicator of viral pathogens in drinking water than currently used indicator systems based solely on bacteria, thus providing an efficient indicator system for viral pathogen risk [90]. This indicator would have broad application to drinking water utilities and would provide a better means to assess viral risk and protect public health, especially in developing countries. Furthermore, TTSuV has been detected in commercial pig products including vaccines, enzymes for laboratory use, and human drugs containing components of porcine origin [13]. Such evidence would indicate the need to screen these products for the presence of these viruses. Finally, it has been reported that the prevalence of TTV is geographically variable. Could such a large variability influence the behavior of the infection at an individual level? Could the chronic exposure to this virus in a particular geographical area, especially in developing countries, be a source of repetitive pathogenicity for hosts? These and other epidemiological and biological aspects of animal AVs should be studied with large comparative and prospective studies.

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