Torque Teno Mini Virus Infection in Chronic Cervicitis and Cervical Tumors in Isfahan, Iran

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Key Words
Torque teno mini virus · Cervical cancer · Adenocarcinoma · Squamous cell carcinoma · Cervical intraepithelial neoplasia · Cervicitis · Nested polymerase chain reaction

Abstract
Objectives: Torque teno mini virus (TTMV) is classified as the Betatorquevirus genus of Anelloviridae. Little is known about the prevalence of TTMV in humans. Worldwide, cervical cancer is the second most common cancer affecting women. This study aimed to estimate the TTMV infection frequency in cervicitis cases and cervical tumors including intraepithelial neoplasia (CIN), squamous cell carcinoma (SCC) and adenocarcinoma, and the possible role of this virus in the etiology of them in an Isfahan population.

Methods: 79 cervicitis cases and 42 tumors were collected from histopathological files of Al-Zahra Hospital in Isfahan, Iran. DNA was extracted and subjected to nested polymerase chain reaction.

Results: Totally 62% of the tested samples were positive for TTMV. It was positive in 52.4% of adenocarcinoma, 68.4% of CIN and 100% SCC cases. In cervicitis, 48% of the cases were positive. In the phylogenetic construct two of the cervical tumor isolates and two of the cervicitis isolates were placed in the same cluster with already reported isolates from Japan (EF538880 and AB041962). Also, three of the cervical tumors isolated (JQ734980, JQ734981 and JQ734982) were placed in another cluster.

Conclusion: The presence of the virus in cervical tissues suggests possible sexual transmission of the virus.

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sexually transmitted pathogens also increase the risk of which are variably implicated in the pathogenesis.

The two main cervical cancers are adenocarcinoma and squamous cell carcinoma (SCC), of which SCC consists of 90–95% of all [10]. Adenocarcinoma is an epithelial gland cell tumor which progresses much faster than SCC [10]. The abnormal growth of precancerous cells in the cervix is called cervical intraepithelial neoplasia (CIN) [11].

Cervicitis is an inflammatory condition of the cervix which frequently is asymptomatic. It is considered to be associated with sexually transmissible pathogens with rates as high as 30–45% in populations affected with sexually transmitted infections [19]. While less than half of the cases are believed to be caused by Chlamydia and Neisseria gonorrhoeae, the etiology of the remainder which are referred to as nonchlamydial, nongonococcal cervicitis or nonspecific cervicitis are unknown [19]. Mycoplasma genitalium, Mycoplasma hominis, Ureaplasma urealyticum, herpes simplex virus, cytomegalovirus, adeno virus and Trichomonas vaginalis are other pathogens which are variably implicated in the pathogenesis [19].

Preliminary reports on the prevalence of TTMV in the human population indicate high rates of infection in French (76–77%) [20], Brazilian (72–77%) [8, 21] and in Norwegian (48%) [22] blood donors and also in French hemodialysis patients (95%) [23]. TTMV DNA has been detected in amniotic fluid, cord blood and breast milk [8, 24]. TTMV genome has also been detected in cervical swabs of healthy women (61%) [25], peripheral blood mononuclear cells, feces and saliva [6].

Considering this issue and the lack of information about the possible effects of TTMV in cervical tumors and cervicitis, the aim of this study was to determine the frequency of TTMV in cervical tumors and cervicitis cases and its possible correlation with them.

Seventy-nine cervicitis cases and 42 cervical tumors which were formalin-fixed and paraffin-embedded were collected from histopathological files of Al-Zahra Hospital in Isfahan, Iran. This study was approved by the local ethics committee. Of 42 tumors tested, 21, 19 and 2 cases were adenocarcinoma, CIN and SCC, respectively. Data on histopathological changes were collected by re-examination of standard hematoxylin- and eosin-stained sections. The tissues were stored in ambient conditions (10–20°) of the dry local climate. Before sectioning each sample, both the tissue blocks and the microtome were cleaned with absolute alcohol, and four 10-μm thick sections from each block were then collected [26, 27].

Sections of tissue were subjected to xylene treatment (1 ml) at 59° for 15 min in 1.5-ml Eppendorf tubes and centrifuged at 11,300 g for 10 min. The procedure was repeated three times, and the samples were then washed with 100% ethanol with three rounds of centrifugation at 9,660 g for 10 min. Finally, the samples were air-dried for 30 min [26, 28].

The deparaffinized tissue sections were treated with 900 μl of solution containing 50 μl of 5 M NaCl, 200 μl of 0.5 M EDTA and 650 μl of retrieval solution (1 M Tris, 0.5 M EDTA, 10% sodium dodecyl sulfate) and thermomixed at 450 rpm for 15 min at 59°. 90 μl of 0.5 mg/ml proteinase K was then added and thermomixed at 500 rpm for 3 h at 59°. An equal volume of phenol/chloroform/isoamyl alcohol (25:24:1) (Merck, Germany) was added and the samples were then centrifuged at 4,290 g for 10 min. The upper phase was collected and transferred to another microtube and 0.1 volume 3 M sodium acetate was added, and the samples were vortexed for 1 min. Then 2 volumes of cold 100% ethanol (Merck) were added and the sample was incubated at –20° overnight. The precipitated DNA was centrifuged at 9,660 g at 4°. The supernatant was discarded and the DNA precipitate was washed once with 75% ethanol. The pelleted DNA was dissolved in 50 μl of distilled water or TE solution (Tris-HCl buffer (10 m M , pH 8.0) containing 1 m M EDTA) after complete drying. Purity of extracted DNA was estimated as the ratio between spectrophotometric absorption at 260 and 280 nm (OD₂₆₀/OD₂₈₀) [28].

To confirm the quality of the extracted DNA, primers (D-141F: 5′-ACCAAGGAGCAAGAGACA-3′ and D-141R: 5′-GTCTTCCGGCCTAGTCTT-3′) were used for detection of the housekeeping gene GJB2 of human chromosome 13, with a product size of 125 bp [29]. Amplification of the GJB2 gene by polymerase chain reaction (PCR) was performed in 25 reaction volumes containing 2 μl of the tissue extract, 1 U of Smartaq DNA polymerase per microliter (Cinnagen, Iran), 0.4 μM of each primer, 240 mM of each dNTP, 20 mM of Tris-HCl, 3 mM MgCl₂, 50 mM KCl and 20 mM of ammonium sulfate. Thermal cycling consisted of an initial denaturing step at 94° for 5 min, followed by 34 cycles of denaturation at 94° for 60 s, 60 s of annealing at 63.2°, 1 min of elongation at 72°, and a post-elongation step of 10 min at 72°.
TTMV in Cervicitis and Cervical Tumors

The primer systems used for amplification of TTMV DNA (340 bp) included M1359/M1365 (sense: GTTTATSMCGYAGACGGAG, PCR product: 433 bp; antisense: TYTGCGAAWAGGCGSTCTAA, PCR product: 433 bp) and M1360/M1366 (sense: GAAGGTGAGTGAAACAC- CG, PCR product: 340 bp; antisense: AGGGCSTCTAA- WTCCTCCKTC, PCR product: 340 bp) (K: G or T, M: A or C, S: G or C, W: A or T, Y: C or T) primer sets in nested PCR [3].

In the first round of PCR, 3 μl of DNA, and in the second round, 1 μl of the PCR product was used in a 25-μl reaction mixture containing 1 U SmarTaq polymerase (Cinnagen), 0.4 μM each primer, 240 μM each dNTP, 20 mM Tris-HCl, 3 mM MgCl₂, 50 mM KCl and 20 mM ammonium sulfate. Thermal cycling conditions in the first round were as follows: initial denaturation at 94° for 5 min followed by 30 cycles of denaturation at 94° for 40 s, annealing at 60° for 40 s, and an extension step at 72° for 50 s. The amplification program was followed by a final extension step at 72° for 5 min. The thermal cycling conditions in the second round of PCR were the same as in the first round. PCR products (10 μl) from the second round were loaded onto a 1% agarose gel (Sigma) containing ethidium bromide and electrophoresed, and the DNA was viewed under UV light.

The PCR products (bands of about 340 bp) of seven randomly selected samples from cervicitis cases and tumor cases were subjected to agarose gel electrophoresis (1%) and DNA was extracted according to guidelines of the DNA Gel Extraction Kit K0513 (Fermentas, Germany). The DNAs were sequenced using an Applied Biosystem 3730 DNA Analyzer (Geneservice, UK). A nucleotide BLAST search of the determined sequences against the nucleotide sequence database (NCBI, National Center of Biotechnology Information) was performed.

A phylogenetic tree was constructed using the neighbor-joining method in MEGA4 (Molecular Evolutionary Genetics Analysis software version 4.1) [30] based on our sequenced amplicons with accession numbers of JQ734980, JQ734981, JQ734982, JQ734984 and JQ734986 of tumor cases and accession numbers of JQ734979 and JQ734985 of cervicitis cases against sequences obtained from GenBank. TTMV virus Isfahan isolates (GQ337059, GQ337060, GQ337061 and GQ337962), nine other TTMV isolates and two variants of TTV were obtained from GenBank database.

Fisher’s exact test was used for statistical analysis using GraphPad Instat software version 3.05 (GraphPad, USA).

In the gel electrophoresis, expected 340-bp bands for TTMV were observed (fig. 1). In the phylogenetic construct, two of the cervical tumor isolates (JQ734984 and JQ734986) and two of the cervicitis isolates (JQ734979 and JQ734985) were placed in the same cluster with the already reported isolates from Japan (EF538880 and AB041962). Also, three of the cervical tumors isolated (JQ734980, JQ734981 and JQ734982) were placed in another cluster. This cluster also included the already reported isolates (GQ337060 and GQ337062) from sera of the same region. However, they were also far from isolates (GQ337059 and GQ337061) from the same region which were placed in another cluster (fig. 2).

The frequency of TTMV infection in different cervical tumors and cervicitis cases is shown in figures 3 and 4. In total, 62% of the tested tumor samples were positive for TTMV. It was positive in 52.4, 68.4 and 100% of adenocarcinoma, CIN and SCC cases, respectively. In cervicitis samples, 48% of the tested samples were positive for TTMV.

The differences observed in the frequency of infection in different tumors were not significant (p > 0.05). The difference in the frequency of infection in tumor cases and cervicitis cases was not significant (p > 0.05).
The tumor and cervicitis cases were categorized into different age groups (≤20–39, 40–59, and ≥60). The frequencies of the virus infection in different age groups are shown in Table 1. The difference among different age groups was not significant (p > 0.05).

This study reports the presence of TTMV in cervical tumors and cervicitis cases for the first time. Preliminary reports on the prevalence of TTMV in the human population indicate high rates of infection in French (76–77%) [20], Brazilian (72–77%) [8, 21] and Norwegian (48%) [22] blood donors and also in French hemodialysis patients (95%) [23].

The percentage of TTMV infection in cervical tumors in this study was 62%, which was similar to 61% of infection of cervical swabs from healthy women tested in Italy [25]. Considering this, Fornai et al. [25] suggested

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**Fig. 2.** Phylogenetic tree was constructed using the neighbor-joining method implemented in MEGA4.1 with 1,000 bootstrap replicates. Evolutionary distances were computed using the maximum composite likelihood model. The units are base substitutions per site. The numbers displayed at each branch separation are bootstrap values in percent.
possible active replication of the virus in the genital tract, for example by lymphoid cells or exudation from plasma. The results obtained in this study show that the cervical tissues should be considered as the source of the virus in the genital tract. Matsubara et al. [24] detected TTMV in infant cord blood suggesting like other viruses, such as human immunodeficiency virus 1, rubella virus, cytomegalovirus and parvoviruses which are transmitted through the intrauterine route, the mother-to-child transmission of TTMV could happen. The detection of TTMV in cervical tissues in this study is in favor of their suggestion.

The report by Fornai et al. [25] that detected TTMV in cervical swabs of healthy women suggests the possible sexual transmission of the virus that is in agreement with the results in this study. Ghazimorad and Bouzari [31] detected the virus in 17 out of 100 sera from healthy blood donors in Isfahan. The frequency of the virus detected in cervicitis and cervical tumors in this study in the same area is significantly higher than the sera (p < 0.01). This may indicate that the changes in tissues induced by cervicitis and cervical tumors may have led to more replication and infection of the virus or the infection by the virus may have led to higher frequency of cervicitis or cervical tumors. Confirmation of these needs more sophisticated investigations.

Neil and Lampe [21] reported a high variability in one of the most conserved regions of the TTMV genomes detected in the sera of blood donors in Brazil. Takahashi et al. [1] also showed a high degree of genomic divergence for this virus. In the present study, a great diversity was also observed among the samples sequenced, with a homology rate from 70 to 85%, which confirms the former reports. Phylogenetic analyses showed that some of the virus isolates are similar to those which were reported from other areas, especially Japan. On the other hand, some were similar to already reported isolates from the sera of the same region. This also confirms the variability of the virus.

It has been suggested that anelloviruses may be involved in various diseases such as pancreatic cancer,

![Fig. 3. Frequency of TTMV infection in cervical tumors and cervicitis cases tested.](image1)

![Fig. 4. Frequency of TTMV infection in different tumors tested.](image2)

<table>
<thead>
<tr>
<th>Age groups</th>
<th>Tumors Total</th>
<th>TTMV+ positivity</th>
<th>Cervicitis Total</th>
<th>TTMV+ positivity</th>
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</thead>
<tbody>
<tr>
<td>≤20–39</td>
<td>7</td>
<td>4</td>
<td>58%</td>
<td>26</td>
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<td>40–59</td>
<td>21</td>
<td>13</td>
<td>62%</td>
<td>33</td>
</tr>
<tr>
<td>≥60</td>
<td>13</td>
<td>9</td>
<td>69%</td>
<td>8</td>
</tr>
</tbody>
</table>

Table 1. Frequency of TTMV in different age groups of cervical tumors and cervicitis cases.
systemic lupus erythematosus, idiopathic inflammatory myopathies, or chromosomal translocation [4]. Although a statistical association between anellovirus infection (or viral load) and disease has been suggested, a causal relationship has not been proven [6]. In comparison of tumor and cervicitis cases, no significant differences were observed in the frequency of TTMV infection (p > 0.05). Moreover, the difference in the frequency of the virus in the different tumors tested was not significant (p > 0.05). Our results are in favor of the idea that TTMV as an anellovirus is not pathogenic.

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References