Prevalence of Oral *Trichomonas tenax* in Periodontal Lesions of Down Syndrome in Tabriz, Iran

**ABSTRACT**

**Background:** It was presented that *Trichomonas tenax* is one of the parasites which is responsible for oral infection. This study was considered to estimate the prevalence of this parasite in oral cavity of Down syndrome patients with periodontal lesions and in healthy population from welfare organization in Tabriz, Iran.

**Materials and Methods:** In the case-control study, 52 patients with periodontal disease (case group) and 52 individuals with healthy gingiva (control group) selected for the study. Examination was done by dental mirror and periodontal probe. After using plaque detector tablets, sampling was done by entering sterile paper into periodontal pocket for 20 seconds.

The amount of plaque was measured by plaque index. Finally samples sent to laboratory for prepared PCR reaction.

**Results:** In the case group, 14 patients were infected and in the control group 5 individuals. Percentage of infection in case group was 18.8% and in the control group was 3% that difference was statically significant. Plaque index in the case group was 72 ± 10.2 and in the control group was 68 ± 11.4 and difference between two groups was not significant.

**Conclusion:** Parasitic infections in Down syndrome were higher than healthy children while plaque index was not significantly different between the two groups. Therefore follow-up of orders are necessary in control of parasitic infection in Down syndrome that have intrinsic defect of immune systems.

**Keywords:** Anaerobic parasite, Gingivitis, Periodontics

**INTRODUCTION**

The oral cavity of human is colonized by specific bacteria, fungi and protozoa. Among these microorganisms, *Trichomonas tenax* (*T. tenax*) is known as common anaerobic parasite of oral cavity and also found in sub maxillary glands [1].

*T. tenax* is commonly found in oral cavity and patients with poor oral hygiene and periodontal disease involved. The occurrence of *T. tenax* is dependant on host ‘age. Different factors are responsible for transmission of parasite that includes incidence by saliva through kissing, or application of polluted dishes and drinking water [2,3]. Depending on oral health status, level of contamination is reported between 0 to 25% [2].

Several studies were undertaken in relation to prevalence of periodontal disease in syndrome down (DS) patients (trisomy 21) [4-7]. It has been shown that defective neutrophil chemotaxis influences the progression of periodontal disease in DS patients [8]. Also, the studies display that the level of prostaglandin E2 (PGE2) detected in gingival crevicular fluid (GCF) from DS patients is increased, a fact that may be of importance in the pathogenesis of the periodontal disease frequently seen in these patients [9]. Moreover, pulmonary trichomoniasis is usually caused by aspirated *T. tenax* [2,10]. Afterwards, health of DS patients could be affected because of this infections and intrinsic defect of the immune system in DS [11]. As correlation to this parasite and periodontal diseases in DS patients, pulmonary trichomoniasis and due to public health importance, this study was performed to determine the prevalence of *T. tenax* in oral cavity of DS patients with periodontal disease and in healthy population in Tabriz, Iran.

**MATERIALS AND METHODS**

The study comprised 104 individuals that were selected from Welfare Organization of Tabriz, Iran. A total of 52 DS patients (5-12 years old) (24 females and 28 males) with periodontal disease were choice as case group and a total of 52 individuals aged between 5-12 years (31 females and 21 males) without any periodontal disease and healthy gingiva were also picked as control group. Each participant was examined using a dental mirror and a periodontal probe. Then learn how to use plaque disclosing tablets. Individual's plaque was measured by O'Leary's index and gingival condition was obtained based on Gingival Index [12]. It scores the marginal and interproximal tissues distinctly on the basis of 0 to 3. The criteria are:

0= Normal gingiva;
1= Mild inflammation – slight change in color and slight edema but no bleeding on probing;
2= Moderate inflammation – redness, edema and glazing, bleeding on probing;
3= Severe inflammation – marked redness and edema, ulceration with tendency to spontaneous bleeding.

A sterile, absorbent paper point was quietly inserted into the periodontal pocket (sulcus). After 20 seconds, the papers were put in a tube having 100µl of NTE buffer [100 mM NaCl, 10 mM Tris-HCl (pH 7.5), 1mM EDTA], PCR of 18S rRNA gene were used for detection of *T. tenax*. Total genomic DNA extracted by phenol-chloroform method according to Sambrook. The 18S rRNA gene was amplified directly from dental plaque with the forward and reverse primers TGBK-F and TGBK-R (5′-AGGAGCTCGGTAAATCCAG-3′ and 5′-CTTGTACATTCTCCTTC-3′), respectively [13].

A written consent was provided from each patient and the study was approved in the university Ethics Committee. A questionnaire documented the history of patients’ general health, any antibiotic consumption (in the three last months), periodontal disease, oral and maxillofacial disease. The statistical analysis was done via SPSS software (Version 11.5) by t-test and Chi-square test to study the correlation between the kind of oral disease, age and sex with the existence of *T. tenax*.

**RESULTS**

In the case group, 14 DS patients (26.9%) and in the control group 5 individuals (9.6%) were infected to *T. tenax* and were positive in the PCR [Table/Fig-1,2]. Amplification was approximately 1000 bp that was compatible with published reports [13].
Comparing gingivitis index show that difference between these two groups (case and control) is statistically significant ($p<0.001$, $X^2=23.76$). But comparing gingivitis index between female and male of the case group ($p=0.05$, $X^2=23.76$) statistically non-significant. Also, in the control group difference between female and male was statistically non-significant [Table/Fig-3].

Comparing of means on percentage of plaque index was shown in [Table/Fig-4]. According to the result, variances of two groups (case and control) were not equal and this difference was statistically non-significant. While variances between female and male in case and control group were equal and statistically non-significant. Comparison of positive infection to $T.tenax$ in two groups by fisher test showed that difference was statically significant ($X^2=5.21$, $p<0.05$). But difference between female and male of two groups was non-significant [Table/Fig-5]. Comparison of positive infection to $T.tenax$ between female by fisher test revealed that difference was statically non-significant ($X^2=1.24$, $p>0.05$). But difference between female and was significant. ($X^2=4.73$, $p<0.05$) [Table/Fig-6].

**DISCUSSION**

The result of this study presented that percent of infection to $T.tenax$ and gingival index in the case group (DS patients) were significantly higher than control group although no significant difference was found in plaque index results between the groups. DS patients mostly would be involved in periodontal disease due to poor oral hygiene, irregular teeth and high frenum attachment. For this reason, it is expected that prevalence of $T.tenax$ in DS patients be higher than healthy children and the result of this study exhibited this subject. Percentage of infection to $T.tenax$ was 26.9%. In the previous study, range of infection to this parasite, among children aged 2-12-year-old, reported approximately 4% [14]. Also, showed that DS patients have inappropriate activity of matrix metalloproteinase that increased the risk of periodontal disease [15]. As well as, $T.tenax$ could produce cathepsin B-like proteinases that affect pathological process such as facilitate penetration of the host, digestion of host proteins and interference with the host immune system [16]. Also, in patients with marginal periodontitis or gingivitis, detection of $E.gingivalis$ and $T.tenax$ were examined. According to the results, *Entamoeba gingivalis* was more dominant among females, whereas $T.tenax$ was not found in both patients and control groups [17].

In the other study prevalence of $E.gingivalis$ and $T.tenax$ were investigated which patients have gingivitis and scale. Results showed that in 48 (21.8%) of the positive 58 specimens, $E.gingivalis$ was discovered alone while $T.tenax$ was present in only two (1%) specimens. In 8 (3.6%) cases, $E.gingivalis$ and $T.tenax$ were recognized together. In addition, other factor such as gum

**Table/Fig-1**: PCR products on the basis of *Trichomonas tenax* 18S rRNA gene. M: molecular size marker, 100bp.

**Table/Fig-2**: Prevalence of *Trichomonas tenax* according to the positive PCR bond.

**Table/Fig-3**: Comparison of infection to $T.tenax$ in two groups.

**Table/Fig-4**: Comparison of means on percentage of plaque index.

**Table/Fig-5**: Comparison of infection to $T.tenax$ in two groups.
problems, existence of tartar, smoking, brushing and control habits were establish to be statistically significantly associated with the oral protozoa [18].

In the other same study, prevalence of *E. gingivalis* and *T. tenax* in oral parasitic infection was evaluated. According to their results, nine infection (6 with *E. gingivalis* and 3 with *T. tenax*) in the case group were determined but in the control group just one infection by *E.gingivalis* was reported [6].

Another research on diagnosis of *E. gingivalis* and *T. tenax* was done between 176 specimens (8 to 19-year-old). Finding showed that 25 subjects (14.2%) had only *E. gingivalis*, 5 (2.9%) had an invasion of *E. gingivalis* and *T. tenax* together and 2 individuals (1.1%) had only *T. tenax*. Result established that oral protozoa was found in the groups (children and teenagers) with having cured or complete dentition. Occurrence rate of parasite was higher in 11 to 19-year-old persons than in the lower age groups. Also, both investigated protozoa can arise simultaneously such that their occurrence rate was deepened on age (increasing with age in rate of *E. gingivalis* and sex dependent (rate is higher in boys than girls) [19]. It is noted that DS patients have intrinsic defect of the immune system [11], therefore these results confirmed that DS patient could be more prone to *T. tenax* infection. Beside determination *T. tenax* in periodontal patients, it was showed that frequency of occurrence of parasite dependence on state of periodontium and hygiene of oral cavity [20]. It is thought that this parasite could enter into respiratory tract from oropharynx by aspiration. Either determined that it feed from bacteria and alone not able to make pulmonary disease and co-infection was accruing between parasite and bacterial flora of oropharynx [21]. On the other hand clinical and epidemiological studies showed close relationship between plaque and periodontal disease. Which in the poor removal of plaque, gingivitis was created and caused progressive periodontitis [22]. According to our results, it is necessary to following oral and general hygiene order to control of parasitic infestations, especially between DS patients which inhibit progress of periodontal disease and related infection such pulmonary trichomoniases in DS patients.

**CONCLUSION**

According to children dental health especially in Down syndrome, detection of parasites responsible to infection is very important. *T. tenax* was one of parasite determined in DS patients in this study because these children had not high oral hygiene. So for this reason were will be exposure to other high risk disease like as trichomoniases by infection of *T. tenax*. Furthermore it is being mentioned that oral health and existence of inflammation is affected *T. tenax* infection thus identification of this infection in DS patients help them to have healthy oral and teeth. In our study gingivitis and gingiva index was high in DS patients comparing control group. Finally, it can be concluded that DS children should receive dental care to have low infection to *T. tenax*, related pulmonary disease and gain high hygiene oral cavity.

**REFERENCES**


