EVALUATION OF DIFFERENCE IN BACTERIAL CONTAMINATION OF TOOTHBRUSHES BETWEEN PATIENTS WITH GINGIVITIS AND PATIENTS WITH HEALTHY GINGIVA-A PILOT STUDY.

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Received: 25.06.19, Revised: 05.04.19, Accepted: 05.06.19

ABSTRACT

To Evaluate the difference in bacterial contamination of toothbrushes between patients with gingivitis and patients with healthy gingiva. To determine the bacterial contamination in terms of CFU/ml after brushing for a period of two weeks in patients with healthy gingiva. To determine the bacterial contamination in terms of CFU/ml after brushing for a period of two weeks in patients with gingivitis. To compare the difference in bacterial contamination in terms of CFU/ml between patients with healthy gingival and patients with gingivitis. The most commonly used method to maintain oral hygiene is toothbrush. Its main goal is to remove plaque, debris and stains which are responsible for gingivitis, periodontitis, tooth decay and halitosis. While removing, toothbrush becomes contaminated with blood, saliva, bacteria and soft debris. The toothbrush itself can act as a foci of infection and retard the disease prognosis and treatment outcomes.

Keywords: Toothbrush, Decontamination, bacterial colonization, brushing, gingivitis, healthy gingiva.

INTRODUCTION

The human oral cavity is invaded by a more number of bacteria flora than any other anatomic area in the body .It has been found that more than 700 species of bacteria out of which 400 species were found in periodontal pocket adiacent teeth⁽¹⁾.Maintaining good health is very important for a good quality of life.The impact of oral health on general health has been proved time and again by many studies. (2,3,4,5,6) The mouth serves as a "window" to the rest of the body, providing signals of general health disorders. Bacteria from the mouth can cause infection in other parts of the body when the immune system has been compromised by disease or medical treatments (e.g., infective endocarditis). Systemic conditions and their treatment are also known to impact on oral health (e.g., reduced saliva flow, altered balance of oral microorganisms). Periodontal disease has an impact on cardiovascular system, this statement was proved by many stuides. In 2006, Holmlund et al., periodontal disease and number of remaining teeth related to a past history of heart attack and high blood pressure or hypertension .Other study showed that both periodontal disease and overall tooth loss from any cause are closely related to cardiovascular disease. Almanet al (2011) have shown a signicant positive association between loss ofbone supporting teeth due to periodontal disease and CVD(25-28).Periodontal disease is often

considered the 'sixth complication' of diabetes⁽²⁹⁾. Poorly controlled diabetics are especially at risk

because they are more likely to developperiodontal disease than well-controlled diabetics . other manifestations includes, mouth lesions may be the first signs of HIV infection, aphthous ulcers are occasionally a manifestation of Coeliac disease or Crohn's disease, pale and bleeding gums can be a marker for blood disorders, bone loss in the lower jaw can be an early indicator of skeletal osteoporosis, and changes in tooth appearance can indicate bulimia or anorexia. Oral Prophylaxis is a premise for oral health and daily plaque and debris removal is considered important for health.Improper maintenance of oral hygiene leads to the accumulation of plaque around the tooth which is a primary cause for gingivitis and periodontitis. Thus, removal of plaque plays a main key role in maintaining oral hygiene. Tooth brushing is the most commonly used, easiest and effective method of oral hygiene practice performed around the world. (7) Toothbrush plays an important role in maintaining personal oral hygiene and it is effective toolfor removing the plaque. Not only the proper selection but also care should be taken in maintaining the toothbrush which is essential for good oral hygiene because the toothbrush also gets contaminated by bacteria. Toothbrushes must have the following requirements to remove the plaque; stiff bristles which is enough to remove plaque without causing trauma to the teeth and gums and small head with soft bristles. Organisms are not only associated with oral cavity but also seen in tooth which includes Streptococcus mutans, Staphylococcus aureus, Pseudomonas, Lactobacillus, Klebsiella, Candida species⁽¹⁾.Toothbrushes also has a significant role in disease transmission and increase the risk of infection since they can serve as a reservoir for microorganisms in healthy, oraldiseased and in immunocompromised people. Contamination is the state of retention and survival of infectious organisms that occur on animate or inanimate objects. (8) Contaminated toothbrushes may play a role in both systemic and localized diseases. This toothbrush contamination is associated with transmission of severe health problems which includes cardiovascular diseases, respiratory disorders, gastrointestinal diseases, arthritis, bacteremia , renal problems and stroke.^(7,8)Toothbrushes can become contaminated from the oral cavity, environment, hands, aerosol contamination, and storage containers and the bacteria which attach to the toothbrush gets accumulated and survive on toothbrushes will helps in transmitting the diseases. In 1920 Cobb reported that toothbrush is the cause of repeated infections in the oral cavity (9). Contaminated tooth brush acts as an environment for microbial transport, retention and growth. Toothbrush heads between the bristle tufts is a favourable medium for the growth of microorganisms. This can be the cause of reinfection person with pathogenic (autoinoculation) or it can acts as a significant risk of dissemination of infection for certain patients such as immunosuppressed,cardiopathic ,organ transplant recipients⁽¹¹⁾. Various factors such as inadequate storage, the toothbrush without decontamination, using the same toothbrush for a longer period of time without changing with new ones, survival of microorganisms for a long time leads to autoinoculation which may results in repeated entry of prospective pathogens and infection in the oral cavity especially in children and in immunocompromised patients. So, disinfection of the toothbrushes is an essential part to prevent various diseases.This condition is specifically important for children ,immunocompromised patients and those are undergoing organ transplantation or chemotheraphy.Actualizing the toothbrushes contamination, the problem of choice of proper tools and methods for their disinfection and the patients' education are important issues which should bring into the focus of dentists in everyday practice, because of the need of prevention the potential influence to oral andsystemic health.The toothbrush environment is also influenced by its design, in the mean of the filaments (number, position,

color, grouping, fixation), as well as by the design of its holder. Caudry et al. found that bacteria are strongly adhered to the toothbrush filaments and the retention of moisture, epithelial and oral debris in the filament bundles raise the survival. The usage bacterial of the toothbrush in an extended period of time makes it a reservoir of microorganisms despite the fact that it is used to lower the present flora in the contaminating dental plaque, so the microorganisms may be imported in the mouth again(autoinoculation). So,the contamination of toothbrush can be prevented by immersing it in disinfectant solutions like 0.1% Chlorhexidinegluconate and 1% Sodium hypochlorite and replacing in a regular time period. So far many studies have evaluated the contamination risk of tooth brushes, within the bias of literature search, it was inferred that, none of the studies has focussed on the difference in contamination between a patient with gingivitis against health gingiva. This difference is studied and found to be true significance, it could help in patient education and help in better treatment outcomes. Hence this study was done to investigate and compare the bacterial load on toothbrushes used by patients with healthy gingiva and gingivitis.

Materials and methods

Study design

A non randomized clinical trial.

Study setting

Approximately1000 patients are visiting saveetha dental college daily. Among them, 90% of the patients are diagnosed with poor oral hygiene and they were given a demo of modified bass brushing method followed by health education to improve their oral health.

Study Population

18 to 45 years who visited the OP of saveetha dental college were selected based on the study criteria.

Eligibility Criteria Inclusion Criteria

Patients with age group between 18-45 years Group-A(gingivitis)-Based on gingival index by Loe and Silness.Group-B(Healthy gingiva)-gingiva which is firm inconsistency, with pink colour and scalloped margins were included in this study.

Exclusion Criteria: Patients with a history of systemic disease (Myocardial infarction, ischaemic heart disease, COPD, Bronchial asthma, Hyperthyroidism, Hyperthyroidism, Hyperthyroidism, Hypertholestrolemia, hypertension, diabetes mellitus, renal disorders, blood disorders, Parkinsons disease, cushing syndrome), patients who had periodontitis and who are not willing to participate were excluded from this study.

Informed consent

Prior to start the study written informed consent was obtained from all the participants.Institution ethical

committe approval was also obtained prior to the study.

Sample size

Based on the study by Taji.et.al, the sample size of this present study was 10%.

Sampling

A non probability type of sampling was used. Selective/judgemental. Patients visiting the OP was choosen based on the inclusion and exclusion criteria until the sample size was achieved in each group.

Armamentarium

The following equipments/materials were used for the study

- Steriled mouth mirror
- Surgical gloves
- Steriled containers
- Normal saline
- Cuvettes
- Micropipette
- Petri dish
- Nutrient agar
- Spirit lamp
- Metal loop
- Incubator

Method

All the gingivitis patients were selected based on gingival index given by Loe H and Silness P (1963). For assessing the severity of gingivitis, and its location by examining qualitative changes of gingival tissues. The severity of gingivitis is scored on the selected index teeth (16,36,12,32,24,44). Tissues surrounding each tooth divided into 4 gingival scoring units which are Disto-facial papilla, Facial margin, Mesio-facial papilla and Lingual gingival margin.

Grading of the gingivitis

Score-0,gingival statusis normal gingivaand the criteria isnatural coral pink gingiva;Score-1,gingival status is mild inflammation and the criteria isslight changes in colour, slight edema. No bleeding on probing; Score-2 ,gingival status is moderate inflammation and the criteria is Redness, edema glazing and it bleeds on probing and score-3,gingival status is severe inflammation and the criteria is marked redness and edema/ ulceration/ tendency to bleed spontaneously.All the examinees who met the criteria were informed about the study.Both were each given a new toothbrush with same brand of fluoridated tooth paste. Each subjects were given a demo of modified bass brushing method and they were requested to follow twice daily for a period of 2 weeks, since it is effecting in cleaning proximal and gingival sulcus. At the end of 2 weeks, brushes were collected in a sterile bag and processed. Each toothbrush was then transferred into the container containing 10ml of steriled normal saline and mixed vigorously for 1 minute.After

mixing,50 μ L of saline was transferred into the cuvette which is incubated at 37°C for 1hr by placing in the incubator. 50 μ L of saline was then spread onto the plates of nutrient agar for the growth of an aerobic bacteria. Each sample was processed 3 times and incubated to minimise the manual and laboratory errors. The nutrient medium was incubated aerobically for 24hrs at 37°C. Then total bacterial count was done. The results are tabulated which are as follows,



Figure-1 depicts the bacteria in agar plate of gingivitis patients.



Figure-2 depicts the bacteria in agar plate of patients with healthy gingiva.

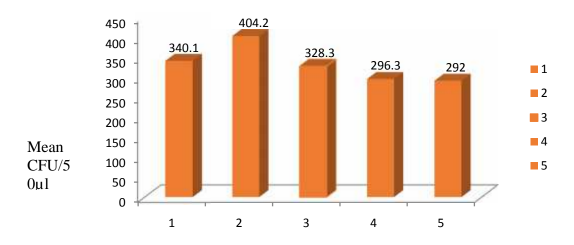
Limitations of the study

A first limitation was the time constraint. A second limitation was the small sample size.

Direction of the future research

It is the follow up of this present study, assessment of progression and prognosis of a disease by using decontaminant solutions.

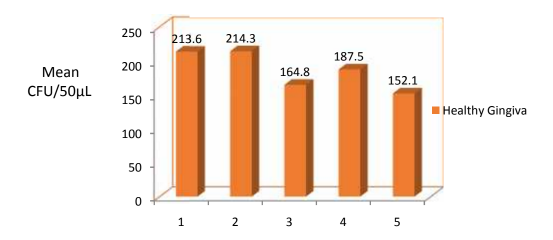
Result
Graph-1 shows the mean of bacterial colony counts of 5 samples in gingivitis patients.



No: of samples

Graph-1,depicts the mean of bacterial colony count in terms of CFU/50 μ L of all the 5 samples in gingivitis patients. Sample-1 has a mean of 340.1 cfu/50 μ L, sample-2 has a mean of 404.2 cfu/50 μ L, sample-3 has a mean of 328.3 cfu/50 μ L, sample-4 has a mean of 296.3 cfu/50 μ L and sample-5 has a mean of 292cfu/50 μ L.

Graph-2 shows the mean of bacterial colony counts of 5 samples in patients with healthy gingiva.



Graph-2,depicts the mean of bacterial colony count in terms of CFU/50 μ L of all the 5 samples in patients with healthy gingiva. Sample-1 has a mean of 213,6 cfu /50 μ L, sample-2 has a mean of 214,3cfu/50 μ L, sample-3 has a mean of 164.8 cfu/50 μ L, sample-4 has a mean of 187.5 cfu/50 μ L and sample-5 has a mean of 152.1 cfu/50 μ L.

Discussion

The result of this study showed that the bacterial contamination was more in toothbrush used by gingivitis patients than the patients with healthy gingiva and the predominant microorganisms

isolated were S. aureus, and S.mutans. In the present study, microbial contamination was seen in all the 10 toothbrushes (100%) and this finding was consistent with some previous studies found microbes on all of the tested toothbrushes⁽¹²⁻¹⁵⁾.But in one of the

previous studies, microbial contamination was seen in 7 out of 10 toothbrushes (70%)⁽¹⁶⁾.Bunetel et al. found that toothbrushes used by patients with existing oral disease quickly became contaminated ⁽¹⁸⁾.Several of the studies found that toothbrushes were contaminated before use(17-20). Caudry et al. found that toothbrushes are heavily contaminated with normal use⁽⁸⁾.In the present study, Predominant microorganisms isolated were S. aureus, S.mutans and this finding was consistent with most similar studies^(12,13,14).In other study,Microbial growth was detected on almost all of the brushes tested in this study (>90%), with development of streptococci observed on the vast majority of the brushes, which shows that toothbrushes are an excellent means of transport for bacteria. Nearly half of the brushes showed growth of mutans streptococci, members of the oral microflora, that are currently considered to be major cariogenic agents (24). Other study reported that toothbrushes are heavily infected with escheriachia coli followed by klebsiella pneumonia, streptococcus pyogenes, staphylococcus auerus⁽³¹⁾.Glass found that toothbrushes from both healthy patients and patients with oral disease contained potentially pathogenic bacteria and viruses as Staphylococcus aureus, Pseudomonas, and herpes simplex virus⁽¹⁷⁾.Svanberg M. found that toothbrushes could be heavily infected mutans microorganisims especially streptococci (30). In the present study, the mean of bacterial colony count in gingivitis patients ranges from 10² to 10⁵ Colony forming units $/50\mu$ L and in patients with healthy gingiva the mean ranges from 10⁻¹ to 10⁻³ colony forming units/50µL.In one of the previous studies, the total microbial load per tooth- brush was found to be 10 4to 10 6colony forming units⁽¹⁵⁾.The American Dental Association recommends a routine change of toothbrushes every 3 months (7). According to the reports of Denny and glass^(23,24) healthy patients replace their toothbrush every two weeks. Patients who are sick should change their toothbrushes at the beginning of an illness, when they first feel better, and when they are completely well. Chemotherapy immuneor patients should change suppressed their toothbrushes every three days, and persons submitted to major surgery should change their toothbrushes every day. So,the replacement of toothbrush in regular time periods is very essential to prevent the continuation of reinfection of oral diseases.

Conclusion

The result of this study showed that the bacterial contamination was more in toothbrush used by gingivitis patients than the patients with healthy gingiva and the predominant microorganisms isolated were S. aureus, and S.mutans. Toothbrushes have an important role in transferring

microorganisms which increases the risk of infection. So, the dentist should be more responsible in order to aware the patients for the issue of choosing, keeping and maintaining the hygiene of the toothbrushes, as well as their replacement in regular period of time.

References

- Karibasappa GNI, Nagesh L, Sujatha BK, Assessment of microbial contamination of toothbrush head: an in vitro study, Indian J Dent Res. 2011 Jan-Feb;22(1):2-5.
- Aubrey Sheiham, Richard Geddie Watt, The Common Risk Factor Approach: a rational basis for promoting oral health, Community Dent Oral Epidemiol, 2000 Dec; 28(6):399-406.
- M.A. Awad, D. Locker, N. Korner-Bitensky, Measuring the Effect of Intra-oral Implant Rehabilitation on Health-related Quality of Life in a Randomized Controlled Clinical Trial, journal of dental health, J Dent Res. 2000 Sep;79(9):1659-63.
- B.LamsterDDS, MMScEvanthiaLallaDDS, MSTWencheS.BorgnakkeDDS, PhD2George W.TaylorDMD, DrPH3,The Relationship Between Oral Health and Diabetes Mellitus,journal of American dental association,Volume 139, Supplement 5, October 2008.
- Renata S.LeiteDDS, MSNicole M.MarlowMSPHJyotikaK.FernandesMDKathieHerma yerMD, MS(Guest Editor),,Oral Health and Type 2 Diabetes,The American Journal of the Medical Sciences,Volume 345, Issue 4, April 2013, Pages 271-273.
- 6. MikaelaKogawaabDanielaCorrêaGrisibDenisePinheir oFalcãoaIngridAquinoAmorimcdTaiaMaria BertoRezendebcdIzabel Cristina RodriguesdaSilvaeOsmarNascimentoSilvafOctávioLui zFrancodghRivadávioFernandesBatistadeAmorima,Im pact of glycemic control on oral health status in type 2 diabetes individuals and its association with salivary and plasma levels of chromograninA,oral biology ,Vol 62 Feb 2016.
- ADA.org: ADA statement on toothbrush care: cleaning, storage and replacement, 2009.
- Caudry SD, Klitorinos A, Chan EC. Contaminated toothbrushes and their disinfection. Journal (Canadian Dental Association) 1995;61(6):511–516.
- Carolus M. Cobb, M.D, The Tooth Brush as a Cause of Repeated Infections of the Mouth, Boston Med Surg J 1920; 183:263-264.
- AuxiliaHemamaliniTilak I, StanicaPrasath 2, Lakshmi.T 3,Evaluation of the contamination and effective disinfection of toothbrushes used by children,ijcrr,Vol03 issue 09.
- 11. White JC, Niven CF., Jr Streptococcus S. B. E.: A Streptoccocus associated with subacute bacterial endocarditis. J Bacteriol. 1946;51:717–22.
- SumaSogi HP, Subbareddy VV, Shashi Kiran ND. Contamination of toothbrush at different time intervals and effectiveness of various disinfecting solutions in reducing the contamination of toothbrush. J Ind Soc PedoPrev Dent 2002;20:81-85.

- Bhat SS, Hedge KS, George RM. Microbial contamination of toothbrushes and their decontamination. J Ind Soc PedoPrev Dent 2003;21:108-112.
- 14. Filho PN, Macari S, Faria G, Assed S, Ito IY. Microbial contamination of toothbrushes and their decontamination. Paediatric Dent 2000;22:381-4.
- Taji SS, Rogers AH. The microbial contamination of toothbrushes. A pilot study. Aust Dent J 1998;43:128-30.
- Mehta A, Sequeira PS, Bhat G. Bacterial contamination and decontamination of toothbrushes after use. N Y State Dent J. 2007;73:20–2.
- Glass RT, Lare MM. Toothbrush contamination: a potential health risk? Quintessence International. 1986;17(1):39–42
- Bunetel L, Tricot-Doleux S, Agnani G, Bonnaure-Mallet M. In vitro evaluation of the retention of three species of pathogenic microorganisms by three different types of toothbrush. Oral Microbiology and Immunology. 2000;15(5):313–316.
- 19. Glass RT. The infected toothbrush, the infected denture, and transmission of disease: a review. Compendium. 1992;13(7):592–598.
- 20. Grewal N, Swaranjit K. A study of toothbrush contamination at different time intervals and comparative effectiveness of various disinfecting solutions in reducing toothbrush contamination. Journal of the Indian Society of Pedodontics and Preventive Dentistry. 1996;14(1):10–13.
- 21. Verran J, Leahy-Gilmartin AA. Investigations into the microbial contamination of toothbrushes. Microbios. 1996;85(345):231–238.

- 22. Denny FW: Risk of toothbrushes in the transmission of res- piratory infections. Pediat Infect Dis J 10:710–11, 1991.
- Glass RT, Jensen HG: More on the contaminated tooth- brush: the viral story. Quint Int 19:713–16, 1988
- 24. Napimoga MH, Hofling IF, Klein MI, Kamiya RU, Goncalves RB. Transmission, diversity and virulence factors of Streptococcus mutans genotypes. J Oral Sci 2005;47:59-64.
- 25. Walls AWG, Steele JG (2001) Geriatric oral health issues in the United Kingdom. International Dental Journal 51:183-187.
- Genco RJ, Glurich I, Haraszthy V, Zambon J, DeNardin E (2001) Overview of Risk Factors for Periodontal Disease and Implications for Diabetes and Cardiovascular Disease. Compendium of Continuing Education in Dentistry (Special Issue) 22(2):21-23.
- Cohen DW, Rose LF, Minsk L (2001) The periodontal-medical risk relationship. Compendium of Continuing Education in Dentistry (Special Issue) 22(2):7-11.
- 28. Holmlund A, Holm G, Lind L (2006) Severity of Periodontal Disease and Number of Remaining Teeth Are Related to the Prevalence of Myocardial Infarction and Hypertension in a Study Based on 4,254 Subjects. Journal of Periodontology 77(7):1173-1178.
- American Academy of Periodontology, Accessed 30 October 2011.
- 30. Svanberg.M Contamination of toothpaste and toothbrush by Streptococcus mutans, European journal of oral science, Volume 86, issue 5.