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 ² Comparative evaluation of salivary leptin levels in
 ³ healthy and chronic periodontitis patients with or
 ⁴ without Diabetes Mellitus
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6 Running title: Salivary leptin levels in healthy and chronic periodontitis patients with or without Diabetes

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- 9 ABSTRACT

INTRODUCTION: Leptin is a hormone-like protein, or some call it as an
 acute phase inflammatory protein, which recently has gained attention due
 to its role in regulating metabolism of the human body as well as affecting
 the body's defense mechanisms, including macrophages.

MATERIALS AND METHODS: Patients visiting Saveetha Dental College 14 from July to November 2020 were examined. Thirty patients (15 males and 15 15 females) were included in this study and subdivided into three groups. 16 Group A has participants with healthy periodontium. Group B, periodontitis 17 and diabetes mellitus. Group C has periodontitis patients only. 18 **RESULTS:** Salivary leptin was observed to be the highest in periodontitis 19 patients with diabetes mellitus followed by patients with periodontitis only 20 and minimal levels in periodontal health patients. 21 **CONCLUSION:** The results indicate that salivary leptin levels are raised in

CONCLUSION: The results indicate that salivary leptin levels are raised in
 the saliva of patients with periodontitis with diabetes mellitus than healthy
 controls and hence can be an important biomarker for periodontal
 therapeutic purposes.

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KEYWORDS: Leptin, diabetes, periodontitis, innovative technology,
inflammation.

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30 **1. INTRODUCTION**

The disease of the periodontium is a chronic inflammatory condition and its 31 severe form is characterized by periodontal ligament loss and resorption 32 and destruction of surrounding alveolar bone [1]. It has now become one of 33 the most common oral diseases in the world that affects many individuals 34 eventually leading to the loss of the teeth. Previous literature has reported 35 the prevalence of periodontitis in developing countries higher than the 36 developed countries. Overall, the disease is prevalent with around 20-50% 37 of the population around the world [2]. The most important risk factors that 38 are known to be associated with periodontitis are poor oral hygiene, 39 Diabetes, tobacco use, alcohol consumption and stress [3]. 40

There has been consistent evidence in previous literatures regarding 41 periodontal diseases which states diabetes mellitus is one of the major 42 systemic risk factors which can play a role in the initiation and progression 43 of the disease and subsequently leading to the destruction of the 44 periodontium[4]. The Gingival crevicular fluids (GCF) and saliva of diabetic 45 patients with periodontitis have higher levels of various types of cytokines 46 which are inflammatory mediators as compared to non diabetic individuals 47 with periodontitis^[5]. Epidemiological studies have proved that diabetes 48 49 mellitus is an important risk factor for periodontitis, and the risk of the disease is greater if glycemic control is poor. People with poorly controlled 50 or uncontrolled diabetes are at a higher risk of periodontitis[6]. 51

52 Leptin is a hormone-like protein that is derived from adipocytes that plays a

53 major role in mechanisms like in metabolism, regulating weight and

function of reproductive organs. It is also known to play a significant role in 54 certain inflammatory conditions through its direct effect on innate and 55 adaptive immune cells [7]. Leptin is also known to have other functions 56 such as stimulating energy expenditure, and modulation of lipid and bone 57 metabolism, function of pancreatic beta cells, insulin sensitivity[8]. Previous 58 literature has shown the presence of leptin in healthy gingival tissues and in 59 60 the gingiva with inflammatory disease which is reduced with the progression of inflammation and is increased in pocket depth [9]. Overall, it 61 62 seems that the expression of the leptin gene has an important role in the modulation of inflammatory processes such as periodontitis[10]. The 63 present study was carried out to compare the salivary leptin levels in 64 healthy individuals and patients with periodontitis with or without diabetes 65 mellitus. 66

67 2.MATERIALS AND METHOD

68 2.1.Study design

This study was designed as a retrospective study. Patients aged 20 to 50 years, visiting Saveetha Dental hospital from July to November 2020 were examined. Thirty patients (15 males and 15 females) were included in this study and subdivided into three groups. (n for each group is not clear) Group A consisted of participants with clinically healthy periodontium.

Group B with periodontitis and diabetes mellitus (type 1 or 2). Group C with
 periodontitis patients only.

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78 2.1.1Inclusion criteria

Healthy periodontium of similar age and gender who had less than 10% of
sites with bleeding on probing, no sites with Probing depth of 4 mm, no

clinical attachment loss >2 mm were included in group A. The inclusion
criteria for periodontitis in group B and C were as follows: no more than >2
teeth missing in each quadrant; <u>\$\$</u>30% of periodontal sites with PD \$\$4 mm;
<u>\$\$</u>20% of periodontal sites with interproximal clinical AL >2 mm; \$\$\$30% of
sites showing BOP.

86 2.1.2 Exclusion criteria

Patients with any other systemic diseases other than diabetes mellitus, smoking habit, history of periodontal treatment in the last 6 months, betel nut users, alcoholism.

90 2.1.3 Saliva collection

Participants were instructed to abstain from eating, drinking, and practicing oral hygiene procedures 12 hours before saliva collection. Whole unstimulated saliva was collected from all patients by spitting into saliva containers. The samples collected were transported to the laboratory and assessed for Leptin levels using ELISA method.

96 2.2 METHODOLOGY

Leptin Levels (Human LEP) in saliva samples were measured using EliKine 97 kit which is a commercially available kit that uses a quantitative sandwich 98 enzyme immunoassay technique which has an antibody specific for Human 99 LEP. The samples were diluted with the calibrator diluent provided with the 100 kit in the ratio of 1:100, and the assay was performed according to the 101 manufacturers' instructions. Standards were included in each run and all 102 103 results were reported within the linearity of the assay. The average of the duplicate readings for each standard, control, and sample were noted and 104 subtracted from the average zero standard optical density (O.D). The 105 concentrations read from the standard curve were multiplied by the dilution 106 107 factor.

108 The results were reported as concentration of Human LEP in picograms 109 per milliliter of sample.

110 **2.3 statistical analysis**

111 Obtained results are tabulated into excel sheets and imported to Statistical

- 112 Package of Social Sciences (SPSS, version 22). One way ANOVA test was
- used to provide statistical significance with P<0.05.

114 **3.RESULTS**

Among the 30 individuals, the salivary leptin levels were observed to be elevated in people with both periodontitis and diabetes mellitus than healthy periodontium individuals and people with periodontitis without diabetes mellitus.

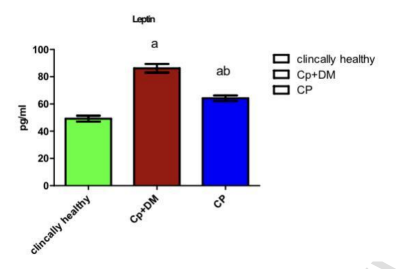
Table 1 shows the mean \pm SD salivary levels of leptin in the healthy individuals, individuals with periodontitis and diabetes and individuals with periodontitis only were 49.20 \pm 6.82, 86.20 \pm 9.93 and 64.20 \pm 6.57 pg/mL respectively. The *P* value for the three groups was found to be highly significant, *P*< 0.0001.

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INTERGROUP COMPARISON	CLINICALLY HEALTHY	CP WITH DIABETES	СР
MEAN	49.20	86.20	64.20
STD.DEVIATION	6.82	9.93	6.57
P VALUE	<0.0001		

126Table 1.Descriptive statistics of salivary leptin levels in healthy subjects, chronic127periodontitis (CP)with diabetes and Periodontitis.



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Fig. 1.Assessment of salivary Leptin concentration among periodontal health, periodontitis with diabetes mellitus and periodontitis only. The levels of salivary leptin were assessed by the Enzyme linked immunosorbent assay method. Significance at *P*<0.05(*P*=0.0001), a - compared with periodontal health group. b - compared with periodontitis with diabetes mellitus.

133 **4.DISCUSSION**

Leptin is a hormone-like protein or some call it as an acute phase
 inflammatory protein, which recently has gained attention due to its role in
 regulating metabolism of the human body as well as affecting the body's
 defense mechanisms, including macrophages [11,12]. Leptin is a member
 of the IL-6 family and prevents a reduction in the synthesis of mucin after
 the activation of lipopolysaccharides thereby having a direct effect on
 preventing the growth of bacteria and preserving health [13].

In the present study the salivary leptin levels were determined in healthy 141 patients with periodontitis with or without diabetes 142 patients and mellitus[14,15]. Based on the results of the current study, there was an 143 increase in leptin salivary levels in periodontitis patients with diabetes 144 mellitus compared to patients with periodontitis without diabetes mellitus 145 and healthy subjects[16, 17]. A previous study conducted by Purwar et 146 al.[18] evaluated leptin levels in saliva and serum of patients with chronic 147

periodontitis and healthy controls where there was a significantly lower in
 healthy controls than those with periodontitis , consistent with the results of
 the current study[18].

In the current study, the salivary leptin levels are observed to be higher in periodontitis patients than healthy controls corroborating with Shimada et al. [19],Seteet al.[20],Purwaret al.[18],Mendoza et al.[21],Karthikeyanet al.[22],Kanoriyaet al.[23].

Salivary leptin has significant physiologic effects on oral keratinocytes 155 which contributes to the wound healing process in the oral mucosa. 156 Salivary leptin might have a role in the antimicrobial and anti-inflammatory 157 property of the saliva, in association with epidermal growth factor[24]. 158 Nokhbehsaimet al. [25] evaluated the serum levels of leptin in patients with 159 periodontitis with and without diabetic mellitus type II where it was 160 concluded that there was a positive correlation between serum leptin level 161 and clinical periodontal parameters [25]. 162

The limitation of this study was this being a unicentered cross-sectional study. Future interventional studies are needed to more strongly elucidate effect of salivary leptin levels in periodontitis patients with or without diabetes mellitus.

167 **5. CONCLUSION**

It can be concluded that diabetic patients have more periodontal tissue destruction and increased salivary leptin concentrations than non-diabetic with chronic periodontitis and patients with clinically healthy gingiva. Salivary leptin may be useful biomarkers of periodontal tissue destruction and helps in determining the pathogenesis of chronic periodontitis. In the

- 173 future, it can pave the way for developing novel treatment procedures in the
- therapeutic management of periodontal diseases.
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176 **REFERENCES**

- de Pablo P, Chapple IL, Buckley CD, Dietrich T. Periodontitis in systemic rheumatic diseases. Nat
 Rev Rheumatol. 2009;5(4):218-24.
- Mariano Sanz, Francesco D'Aiuto, John Deanfield, Francisco Fernandez-Avilés.European
 workshop in periodontal health and cardiovascular disease--scientific evidence on the association
 between periodontal and cardiovascular diseases: a review of the literature, Eur. Heart J. 2010;
 B3–B12.
- McGhee JR, Fujihashi K, Lue C, Beagley KW, Mestecky J, Kiyono H. Role of IL-6 in human
 antigen-specific and polyclonal IgA responses. AdvExp Med Biol. 1991; 310:113-21.
- Preshaw P. M., Bissett, S. M. Periodontitis: oral complication of diabetes. Endocrinol. Metab.
 Clin.North Am, 2013;42(4), 849–867.
- Chapple I. L. C., Genco, R. and working group 2 of the joint EFP/AAP workshop Diabetes and
 periodontal diseases: consensus report of the Joint EFP/AAP Workshop on Periodontitis and
 Systemic Diseases. J. Clin. Periodontol.2013;106–S112.
- 190

Soskolne W. A., Klinger A. The relationship between periodontal diseases and diabetes: an
 overview, Ann. Periodontol. 2001;6(1), 91–98.

- Fernández-Riejos P, Najib S, Santos-Alvarez J, Martín-Romero C, Pérez-Pérez A, González Yanes C, Sánchez-Margalet V. Role of leptin in the activation of immune cells. Mediators
 Inflamm. 2010;2010:568343.
- 196 8. Carbone F, La Rocca, C, Matarese, G. Immunological functions of leptin and adiponectin,
 197 Biochimie. 2012;94(10):2082–2088.
- 198 9. Johnson RB, Serio FG. Leptin within healthy and diseased human gingiva.J Periodontol.
 199 2001;72(9):1254-7.

- 200 10. Bado A, Levasseur S, Attoub S, Kermorgant S, Laigneau JP, Bortoluzzi MN, Moizo L, Lehy T,
 201 Guerre-Millo M, Le Marchand-Brustel Y, Lewin MJ. The stomach is a source of leptin. Nature.
 202 1998;394(6695):790-3.
- 203 11. Friedman J. M. Leptin at 14 y of age: an ongoing story, AM. J. CLIN. NUTR. AM J CLIN NUTR,
 2009;973S–979S.
- 205 12.Tartaglia LA, Dembski M, Weng X, Deng N, Culpepper J, Devos R, Richards GJ, Campfield LA,
 206 Clark FT, Deeds J, Muir C, Sanker S, Moriarty A, Moore KJ, Smutko JS, Mays GG, Wool EA, Monroe
 207 CA, Tepper RI. Identification and expression cloning of a leptin receptor, OB-R.Cell.
 208 1995;29;83(7):1263-71.
- 13.Hsu A, Aronoff DM, Phipps J, Goel D, Mancuso P. Leptin improves pulmonary bacterial clearance
 and survival in ob/ob mice during pneumococcal pneumonia. ClinExp Immunol. 2007; 150(2):332-9.
- 14. Venkatesan J, Rekha P.D, Anil S, Bhatnagar I, Sudha P.N, Dechsakulwatana C, Kim S, Shim
 M.S. Hydroxyapatite from Cuttlefish Bone: Isolation, Characterizations, and
 Applications. Biotechnol. BioprocessEng.2018;23, 383-393.,383–393.
- 214
- 215 15. VellappallySajith, Al Kheraif, Abdulaziz A, Anil, SukumaranWahba, Ashraf A.IoT medical tooth
 216 mounted sensor for monitoring teeth and food level using bacterial optimization along with adaptive
 217 deep learning neural network. Measurement. 2019;672–677.
- 218 16. Del Fabbro M, Karanxha L, Panda S, Bucchi C, NadathurDoraiswamy J, Sankari M, Ramamoorthi
 219 S, Varghese S, Taschieri S. Autologous platelet concentrates for treating periodontal infrabony
 220 defects. Cochrane Database Syst Rev. 2018;26:11(11):CD011423
- 17. VellappallySajith, Al Kheraif, Abdulaziz A., Divakar, DarshanDevang., Basavarajappa, Santhosh
 Anil, Sukumaran; Fouad Hassan. Tooth implant prosthesis using ultra low power and low cost
 crystalline carbon bio-tooth sensor with hybridized data acquisition algorithm. Comput. Commun.
 2019;176–184.
- 18. Purwar P, Khan MA, Mahdi AA, Pandey S, Singh B, Dixit J, Sareen S. Salivary and serum leptin
 concentrations in patients with chronic periodontitis. J Periodontol. 2015;86(4):588-94.
- 19. Shimada Y, Komatsu Y, Ikezawa-Suzuki I, Tai H, Sugita N, Yoshie H. The effect of periodontal
 treatment on serum leptin, interleukin-6, and C-reactive protein.J Periodontol. 2010;81(8):1118-23.
- 229 20. Manuela RubimCamaraSete, Ronaldo Lira-Júnior, Ricardo Guimarães Fischer, Carlos Marcelo S
 230 Figueredo. Serum Adipokine Levels and their Relationship with Fatty Acids in Patients with Chronic

- 231 Periodontitis, Braz. Dent. J. 2015;169–174.
- 232 21. Mendoza-Azpur G, Castro C, Peña L, Guerrero ME, De La Rosa M, Mendes C, Chambrone L.
 233 Adiponectin, leptin and TNF-α serum levels in obese and normal weight Peruvian adults with and
 234 without chronic periodontitis. J ClinExp Dent. 2015;1;7(3):e380-6.
- 22. Karthikeyan B. V. and Pradeep, A. R. Gingival crevicular fluid and serum leptin: their relationship
 to periodontal health and disease, J. Clin. Periodontol.. 2007;467–472.
- 237 23. Kanoriya D, Pradeep AR, Mallika A, Singhal S, Garg V. Correlation of crevicular fluid and serum
 238 levels of retinol-binding protein 4 and leptin in chronic periodontitis and obesity. Clin Oral Investig.
 239 2017; 21(7):2319-2325.
- 24. Hart B. L., Korinek, E. and Brennan, P. Postcopulatory genital grooming in male rats: prevention
 of sexually transmitted infections, Physiol. Behav, 1987;41(4),321–325.
- 242 25. Nokhbehsaim M, Keser S, Nogueira AV, Jäger A, Jepsen S, Cirelli JA, Bourauel C, Eick S,
 243 Deschner J. Leptin effects on the regenerative capacity of human periodontal cells. Int J Endocrinol.
 244 2014;180304.

245