Role Of Dermatoglyphics As A Potential Diagnostic Marker For Periodontitis: A Clinical Study

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Abstract:

Background: Periodontitis is a chronic inflammatory disease affecting the connective tissue attachment and supporting bone around the teeth. Over the years, the retrospective analysis of the possible etiological factors responsible for periodontal infections has magnified to its brink. The necessity lies in segregating the evidences and acknowledging their potential role in disease pathogenesis. Genetic predisposition might explain such an individual variability that can be predicted by using various cytogenetic markers. However, these studies are far more costly and complicated. The present study was conducted to study the role of palmar dermatoglyphics as a diagnostic tool for periodontal infections.

Materials and methods: The present cross sectional study comprised of 120 subjects who were categorized as those suffering from chronic generalized periodontitis (n = 60), in whom variouspalmar patterns were evaluated as compared to healthy controls (n = 60), to find a "dermatoglyphic marker," if any.

Results:A statistically significant higher percentage of ulnar loops (80.5%) was seen in CGP patients as compared to a majority of whorls (80.17%) in the healthy controls (p < 0.001). The percentage of presence of palmar accessory tri-radii was similar in both groups $\chi 2 = 0.963$ (p = 0.326). Mean a-t-d angle variation was statistically significant in the CGP group as compared to the healthy group $\chi 2 = 83.796$ (p = 0.000).

Conclusion: The results of the study revealed that dermatoglyphics holds promising results for determining the genetic susceptibility of individuals to develop periodontitis.

Keywords: Dermatoglyphics, chronic generalized periodontitis, palmar patterns.

I. Introduction

The study of definite ridge patterns on the fingers, palms and soles is known as 'Dermatoglyphics'. The concept emerged by the name 'epidermal ridge configurations' and later sculpted into the terminology 'skin casings'. In 1926, Dr. Harold Cummins, father of American fingerprint analysis and Mildo, replaced the existing terms by the word dermatoglyphics.¹

The initiation of ridge pattern formation occurs in the 6-7th week of the embryonic period and terminates around the 20^{th} week of gestation. In this span of foetal development any abnormal in-utero genetic or non-genetic factors can disturb the patterns. Hence, these are rightly considered as, "the window of intrauterine and congenital abnormalities".²

Ridge patterns remain unchanged for life and no two individuals share the same patterns which show a genetic association of dermatoglyphics. The most commonly encountered process of recording the patterns is rapid, non- invasive, economical and thus a convenient technique for the clinician as well as the patients.

In dentistry, assessment of dermatoglyphics has been linked with several disorders like, hereditary gingival fibromatosis,³ cleft lip/ palate,⁴ bruxism,⁵ dental caries⁶etc. Periodontitis is the destruction of the periodontium and has several etiological factors which are accredited with disease onset and progression. Genetics has been associated with periodontitis since ancient times. However, the techniques used to confirm the genetic basis of periodontitis are unavailable at most times and are expensive as well. Various epidemiological studies have been conducted to confirm the role of genetic alterations in the pathogenesis of an antenatal disturbance like periodontitis.^{7,8} In order to alleviate this predicament, research has recently brought dermatoglyphics into the limelight.²The primitive studies by Yilmaz⁷ and Atasu⁸were successful to acknowledge ridge pattern variations in different forms of periodontitis. Genetic correlation of aggressive periodontitis with ridge patterns has also been determined.¹

The present study was conducted assuming the hypothesis that an antenatal disturbance like periodontitis should manifest in a prenatal event such as dermal ridge formation. Hence the aim of the present study was to assess the role of dermatoglyphics as a diagnostic tool for chronic generalized periodontitis (CGP) and to compare these patterns with those of periodontally healthy subjects.

II. Materials And Methods

The study comprised of subjects referred to the Department of Periodontology, Swargiya Dadasaheb Kalmegh Smruti Dental College and Hospital, Nagpur. The study received ethical committee approval of the institution. The participants were explained about the study and an informed consent was obtained.

120 subjects were divided into two groups (Group A& Group B). Group A comprised of 60 subjects who were diagnosed with CGP after clinical and radiographic examination whereas group B consisted of 60 periodontally healthy subjects. The mean age of the participants were 30 ± 12 (Average age: 40). Exclusion criteria was subjects having congenital/ systemic diseases or those diagnosed with aggressive periodontitis.

All the participants were asked to wash their hands with soap in order to remove all the dirt. Then the fingertip and palmar patterns of each participant were recorded by using standard ink method, using blue duplicating ink (Kores India Limited, Mumbai), thick white printing paper (JK Copier, A4 size, 100 g/m2) and sponge pad.⁹The finger and palmar prints obtained using this method were studied with the help of magnifying glass. For each individual all ten finger patterns were recorded. The parameters were assessed by both the qualitative and quantitative analysis. The parameters assessed qualitatively were finger print patterns including arches, loops (ulnar loop, radial loop) & whorls and palmar patterns in which assessment of accessory tri-radii was evaluated as shown in Fig I. Palmar tri radii patterns are of two types: digital (a, b, c, d) and axial/carpal (t). Accessory tri radii which are considered as genetic variants are denoted as a', b', c', d' and t'.¹⁰

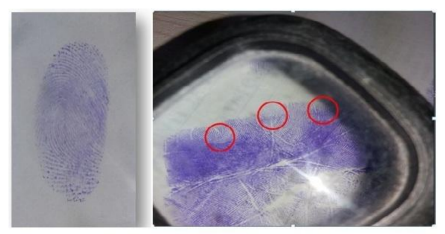


Fig I: Analysis of finger and palmar patterns

The finger and palmar prints were analyzed using Cummins, Mildo and Penrose method.¹¹According to this method the patterns showing maximum frequency in each individual was considered for evaluation. The a-t-d angle which is a palmar pattern is formed between the tri radii a-t-d.¹²It was measured using a protractor and assessed by quantitative analysis as shown in Fig II.

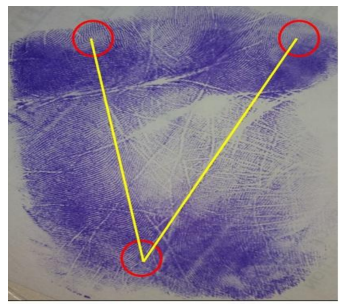


Fig II: Measurement of atd angle

III. Statistical Analysis

Calculation of percentage of fingertip pattern frequency on all 10 fingers of each subject of both groups followed by their comparative evaluation after statistical analysis by paired't' test (p < 0.001) was done.Calculation of percentage of palmar accessory tri- radii and 'a- t- d angle' variation, if any, on both palms of each subject of both groups followed by their comparative evaluation after statistical analysis by Chi Square test (p < 0.001) was done.All the above analyses were performed using statistical package SPSS software (version 20.0)

IV. Results

IV.1. Qualitative analysis: Table 1 depicts the demographic data of fingerprint patterns in the study groups. The results showed a statistically significant higher percentage of ulnar loops(80.5%) in CGP patients as compared to a majority of whorls (80.17%) in the healthy controls (p < 0.001). The percentage of arches was 10% in CGP group as compared to 5.67% in healthy individuals. The percentage of presence of palmar accessory tri-radii was similar in both groups $\chi^2 = 0.963$ (p = 0.326) as described in Table 2.

% of Pattern Frequency								
Group	Arch	Ulnarloop	Radial loop	Whorl				
CGP								
(in %)	10%	80.5%	12.33%	5.5%				
MEAN	2.22±1.07	8.18±1.30	1.55±0.88	1.94±0.96				
Healthy								
(in %)	5.67%	2%	10.33%	80.17%				
MEAN	2.13±1.00	1.2±0.42	2.21±1.16	8.15±1.21				
p value	0.005	<0.001	<0.001	<0.001				

Table 1

P value is set at <0.001

CGP: Chronic generalized periodontitis

% of Palmar accessory tri radii								
			Group		Total			
			CGP	Healthy				
	Absent	Count	47	51	98			
Palmar accessory		% within Group	79.7%	86.4%	83.1%			
tri radii		Count	13	9	22			
	Present	% within Group	20.3%	13.6%	16.9%			
		Count	60	60	120			
Total	% within Group	100.0%	100.0%	100.0%				

Table 2

P value is set at <0.001

CGP: Chronic generalized periodontitis

IV.2. Quantitative analysis:

The angle between triradii a, t and d (atd angle) was measured on right and left hand separately. Analysis of the percentage of mean a-t-d angle variationwas statistically significant in the CGP group as compared to the healthy group $\chi 2= 83.796$ (p= 0.000) as shown in Table 3.

% of a- t- d angle variation								
		Group		Total				
			CGP	Healthy				
atd angle variation		Count	11	60	71			
	Absent	% within Group	16.9%	100.0%	58.5%			
	Present	Count	49	0	49			
		% within Group	83.1%	0.0%	41.5%			
Total		Count	60	60	120			
		% within Group	100.0%	100.0%	100.0%			

Table 3

P value is set at <0.001

CGP: Chronic generalized periodontitis

V. Discussion

The role of dermatoglyphics in periodontal diseases had its inception in a study by Yilmaz et al 1993 in 70 individuals comprising of 36 early onset periodontitis cases, 20 adult periodontitis and 20 periodontally healthy patients and evaluated the quantitative and qualitative patterns of ridged skin. They confirmed the role of heredity in the etiopathogenesis of periodontal diseases.⁷

In a case control study Atasu et al 2005 performed qualitative analysis of fingertip, palms and sole patterns of 158 subjects which encompassed 36 Juvenile Periodontitis (JP), 45 Rapidly Progressing Periodontitis (RPP), 38 Adult Periodontitis (AP) cases compared to 39 healthy controls. A decreased frequency of ulnar loops on all digits and increased frequency of 'e' tri- radii on sole pattern was found in the JP group. An increased frequency of radial loops on second digit & IV and H loops and't, b' triradiion the palms in RPP group whereas increased frequency of concentric whorls and ulnar loops on all digits was encountered in AP group.⁸ In the present study an increased frequency of ulnar loop patterns was observed whereas no statistically significant difference was observed in the palmar accessory tri radii frequency.

Gupta and Karjodkar 2013 evaluated the variation in the atd angle amongst other dermatogylphic patterns in precancerous and cancerous lesions of the oral cavity.²This value is between 30 to 65 in healthy individuals where as variation in value ($< 30^{\circ}$ or $>65^{\circ}$)indicates delayed development of tri- radius 't' and a shift in its position thus altering the angle.¹² To our knowledge the present study is the first to evaluate this parameter in periodontitis cases. In this study we found that an acute atd angle was formed on the palmar patterns of the test group as compared to the healthy controls. This indicates that there is altered development of the tri radius 't' causing a shift in its position.

In another case control study by Devishree et al 2015 evaluating the qualitative parameters on finger tips in 15 aggresive periodontitis as compared to 15 periodontally healthy cases an increased frequency of ulnar loops was found in the test group.¹ However in our study a similar finding was observed in chronic periodontitis. It can thus be assumed that although an increased frequency of ulnar loops is a characteristic feature of periodontal diseases, a higher level of genetic analysis is required to categorize the type of periodontitis.

Babitha et al 2016 conducted an observational study to assess correlation between dermatoglyphics, ABO blood group, Rh factor in periodontally healthy (n=100) and diseased patients (n=100). An increased frequency of whorls and loops were encountered in chronic periodontitis patients. Also majority of the test group participants had 'O' positive blood group as compared to 'A' positive healthy controls.¹³The results of the present study can be correlated with the aforesaid study.

VI. Conclusion

Within the limits of the study, it can rightly be concluded that, dermatoglyphics can be a potential diagnostic aid in determining the genetic basis of CGP. The prevalence of a specific pattern predominantly in the latter, justifies that presence of ulnar loops and 'a-t-d angle' variation can be considered as a "dermatoglyphic marker" in the diagnosis of CGP. Further large scale qualitative and quantitative research will aid in proving the rightfulness of the present study.

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