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 Original Research Article

 2
 AN IN VITRO STUDY TO EVALUATE THE TRANSVERSE STRENGTH OF HEAT

 3
 CURED PMMA RESIN REINFORCED BY DIFFERENT CONCENTRATIONS OF TWO

 4
 NANOPARTICLES

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6 ABSTRACT

Purpose: The purpose of the study was to evaluate transverse strength of PMMA resin reinforced with
 various nanoparticles of different concentrations.

9 Method: One hundred samples of PMMA resin were made and divided into five groups (20 samples for 10 each group). The test specimens were divided into five groups depending on the concentration of 11 reinforcing nanoparticles as Group 1,2,3,4 and 5; Group 1: PMMA unreinforced (control group), Group 2: 12 PMMA reinforced with 2.5% nanozirconia, Group 3: PMMA reinforced with 5% nanozirconia, Group 4: PMMA reinforced with 2.5% titanium dioxide nanoparticles, and Group 5: PMMA reinforced with 5 % 13 titanium dioxide nanoparticles. Universal testing machine was used to conduct a three-point bending test 14 15 and evaluate the transverse strength of samples. Comparison of mean transverse strength for various 16 groups was carried out was done employing one way analysis of variance and Bonferroni post hoc tests. 17 Results: The highest and lowest mean transverse strength were of Group 3 and 1, respectively. 18 Bonferroni post hoc honestly significant difference multiple comparison for mean transverse strength 19 increase in strength to be statistically significant between all the groups (P < 0.05) except between the 20 samples of group G1 and G5 and G2 and G3.

21 Conclusion: Addition of nanoparticles in all concentrations significantly increased transverse strength of 22 heat cure PMMA resin as compared to control group. The best result was obtained after adding 5% of 23 nanozirconia particles to the conventional heat polymerized acrylic resin.

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25 **KEYWORDS**

26 PMMA, nanozirconia, titanium dioxide nanoparticles, transverse strength

27 1. INTRODUCTION

Edentulism has been a matter of great concern to a majority of people, be it partial or complete.
Replacement of teeth by artificial substitutes plays a vital role in leading a normal life.^[1]

30 Since 1930, acrylic resin polymethyl methacrylate (PMMA) has been the most popular material for 31 denture fabrication. Not only it provides an accurate fit, good esthetics, stability in the oral environment 32 but also is easy to handle in laboratory and clinics. Despite many advantages this material exhibits certain 33 limitations which render failure to fulfill mechanical requirements for dental applications. These include low fracture resistance and plaque accumulation^[2], high coefficient of thermal expansion and relatively 34 35 low modulus of elasticity. Fracture of maxillary dentures is twice more common than that of mandibular 36 dentures. Fractures caused outside the mouth are usually a result of heavy impact forces or a high stress 37 rate while those caused within the mouth are usually caused in function and can be attributed to a fatigue 38 phenomenon i.e. low and repetitive stress rate which commonly occurs over a period of time. This type of fracture is typically seen in midline of maxillary dentures than in mandibular dentures.^[3] 39

40 Release of PMMA from the dentures has reported to cause irritation to mucosa.^[5] Also being a 41 radiolucent material it cannot be imaged using standard radiographic techniques hence in cases of 42 accidental ingestion of prosthesis, aspiration or traumatic impaction of dental appliance, their detection 43 can become painstaking and invasive procedures may need to be carried out.^[4]

There have been several attempts to improve the mechanical properties of acrylic resins such as chemical modification or reinforcement with glass fibers, metal oxides and nanoparticles.

Recently, incorporation of nanofillers has been suggested to improve mechanical properties of PMMA. Fine particle size enables the homogenous distribution of nanofillers in PMMA and has reportedly improved the thermal properties of PMMA by increasing its thermal stability compared with pure PMMA.

49 However, size, shape, type and concentration of nanoparticles added affect the properties of resin.^[5]

50 The few studies conducted on the effect of nanoparticles on the transverse strength have been more or 51 less conclusive and unclear.^[6]

52 Titanium dioxide and zirconium oxide nanoparticles have become popular as reinforcement nanofillers 53 recently. Titanium is used since it increases the surface hydrophobicity, reduces the adherence of 54 biomolecules, aids in colouring, has antimicrobial properties and improves mechanical properties of 55 PMMA resins. Spherical particles of titanium dioxide have been used to improve the flexural strength as 56 spherical particles increase the polishability and mechanical properties. Other structures such as 57 nanotubes and fibers which have been recently discovered show much better properties.^[7]

58 Zirconia has exhibited excellent biocompatibility and being white in colour it is less likely to interfere with

59 esthetics.^[4]

60 Zirconium oxide nanoparticles mechanically reinforce the polymers and allows for high impact strength,

61 fracture toughness, hardness and density of the reinforced PMMA matrix.^[8]

62 Modifying nanozirconia powder by coating with a layer of trimethoxysilypropylmethacrylate (TMSPM)

63 renders more radiopacity as it decreases the radiographic density and allows more absorption of 64 radiation.^[4]

Hence, the purpose of this study was to evaluate and compare transverse strength of heat cured PMMA
 resin after its reinforcement with zirconium oxide and titanium dioxide nanoparticles in concentrations 2.5

67 wt % and 5wt % each.

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70 2. MATERIAL AND METHODS

- 71 2.1 The study proceeded as follows:-
- 72 i. Fabrication of metal dies-

Three stainless steel metal dies of dimensions 65mm x 10mm x 3 mm were fabricated. The
 selection for dimensions of the dies was based on ADA specification no. 12.

75 ii. Fabrication of test samples-

76 Preparation of molds for fabrication of wax pattern:

a. The stainless steel metal dies were impressed upon putty material (Affinis, New Delhi, India) so
as to create a mold space. Molten wax (No.2, Rolex, Ashoosons Dental Care Pvt. Limited, Delhi,
India) was then poured onto the mold spaces so created and left to cool. The wax patterns of
dimension 65 x 10 x 3mm as per ISO 1567 standardization were obtained after cooling. These

81 patterns were lubricated with a thin layer of petroleum jelly (Loba Chemie, India) and invested in 82 Type II dental plaster (Dentex, India). After the investing materials had set, the flasks were 83 placed for dewaxing in a conventional water bath. The molds so created were thoroughly flushed with hot water. The flasks were left to cool followed by application of a layer of separating media 84 85 (DPI Cold Mould Seal, Bombay Burmah Trading Corporation Ltd., Mumbai, India) to prepare the flasks for packing. The appropriate amount of heat cure acrylic resin required was prepared from 86 87 a mixture of polymer and monomer in the ratio of 3:1 by volume i.e. 3 parts of polymer and one 88 part of monomer with the help of electronic weighing machine (Jeejex Digital Electronics SF-400, 89 Jiya Sales, India).

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91 iii. Division of samples into various groups (Fig.1):

92 G1 - Control (DPI Heat Cure, Bombay Burmah Trading Corporation Ltd., Mumbai, India)

93 G2- DPI Reinforced with 2.5 wt% nanozirconia particles (ZrO₂, Purity 99.9%, Average particle

94 size: 30-50nm, NanoResearch Lab, Jamshedpur, Jharkhand, India)

95 G3 - DPI Heat Cure with 5 wt% nanozirconia particles

96 G4 - DPI Heat Cure with 2.5% titanium dioxide nanoparticles (TiO₂, Anatase, Purity: 99.9 %,

97 Average particle size : 10-20nm, NanoResearch Lab, Jamshedpur, Jharkhand, India)

98 G5 - DPI Heat Cure with 5 wt% titanium dioxide nanoparticles

G1 (Control)	G2 (2.5 NZR)	G3 (5 NZR)	G4 (2.5 NTO)	G5 (5 NTO)
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102 Fig. 1 - Specimens for each group

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a. Preweighed nanoparticle powder was separately added to the acrylic resin powder to form the
 desired formulation and thoroughly mixed using a mortar and pestle (Uniteck Scientific &
 Electronic Industry, Chandigarh, India).

107 b. Proportionate amount of the monomer and the polymer was taken in the mixing jar and thoroughly mixed until dough stage was attained. The flask (Varsity Flask, No-7, S.S. Products, 108 109 India.) was closed and trial closure was carried out using hydraulic press (Unident Instruments 110 India Private Limited, India). After removal of flash, flask was then clamped and pressure maintained for 30 minutes to allow proper penetration of monomer into polymer. The flask was 111 then immersed in an acrylizer (Unident Instruments India Private Limited, India) at room 112 temperature. The temperature was raised to 74 C, held for 8 hours, then raised to 100 C for an 113 114 hour. After the completion of the curing cycle, the flask was removed from the water bath and 115 bench cooled for 30 minutes, immersed in cool tap water for 15 minutes prior to deflasking. Samples were then contoured using carbide bur, finished with sandpaper and polished using 116 slurry of coarse pumice. The width and thickness of each samples was measured using a digital 117 118 vernier calliper (PRECISE, Sudershan Pvt Ltd, Delhi, India) with a resolution of 0.01mm. Since

- width and thickness were factors assessed for determining transverse strength, only the resin
 samples with a slight variation in size up to + 0.3mm were included in the study.
- 121 In this manner, a total of 100 acrylic samples divided into five groups each containing 20
- samples of compression moulded heat cure acrylic denture base resin were fabricated.
- All the specimens were stored in distilled water at a temperature of 37° C for 48 hours prior to transverse strength testing.^[9]

125 **2.2 Calculation of Transverse Strength:**

To determine transverse strength, fracture load was measured using the three-point bending test according to ISO 178 on a universal testing machine (ASIAN Test Equipments, Micronix Intruments, India). Then specimens were placed on a 3-point flexure apparatus and the support span was 50 mm. Load was applied at the midpoint of the sample with a crosshead speed of 5mm/min until the specimen fractured and fracture load was recorded (Fig. 2).^[7] The transverse strength values of each specimen were derived using formula:

132 TS = <u>3WL</u>

133 $2bd^2$

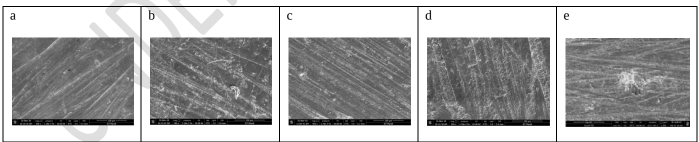
where TS is the transverse strength (in MPa), W is the fracture load (N), L is the distance between the two supports, b is the specimen width, and d is the specimen thickness. The data thus obtained was subjected to statistical analysis.



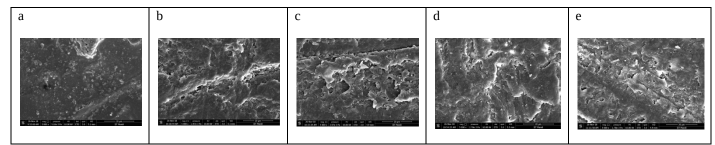
138 Fig. 2 - Testing the specimen on Universal Testing Machine

139 **2.3 SEM ANALYSIS:**

SEM (FEI Nova NanoSEM 450, USA) was used to examine the surface of fractured cross-section of the specimens. The acceleration voltage, used to perform SEM evaluation, was set at 10 kV and the working distance was 5.3 mm with a 3 spot size. A 10 nm gold-palladium coating was applied to provide conductivity to the material. Images were recorded at different magnifications to study distribution of particles (Fig. 3a-e and Fig. 4a-e).



146 Fig. 3 - SEM images at 500x



147 Fig. 4 - SEM images at 5000x

148 **3. RESULTS**

- 149 In the present study all experimental groups other than unreinforced PMMA have shown a definite
- 150 increase in mean transverse strength with reinforcements. Data was analysed using computer software
- 151 STATA and SPSS-20.0, IBM software, Chicago.
- 152 The mean transverse strength (MPa) obtained for different categories of nanoparticles as well as with

153 control have been summarized in Table 1. Maximum values were obtained with G3, followed by G2, G4,

- 154 **G5** and minimum strength was obtained for **G1**.
- 155 In the present study all experimental groups other than unreinforced PMMA have shown a definite 156 increase in mean transverse strength with reinforcements.
- 157 A multiple comparison test by one way ANOVA as shown in **Table 2** revealed a significant difference 158 among the means obtained for all the groups (P < 0.001).
- For comparison within the group i.e. multiple group comparison: **Posthoc Bonferroni** test was applied which shows that the increase in transverse strength was statistically significant for all experimental groups in comparison with samples of unreinforced PMMA group except between G1 and G5 (P= 1.000) and G2 and G3 (P=.74).
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170 Table 1- Descriptive statistics of transverse strength values (MPa) obtained for control and two

Groups	N	Mean	StandardD	Standard	
		(MPa)	eviation	Error Mean	
G1	20	94.800	9.83483	2.19913	
G2	20	134.82	12.66516	2.83202	
G3	20	148.96	28.65139	6.40664	
G4	20	119.26	6.91676	1.54663	
G5	20	101.98	14.43716	3.22825	

171 types of nanoparticles:

173 Table 2: Intergroup comparison of mean of transverse strength (MPa) among various studied

174 groups:

Transverse	Sum of	df	Mean	F	Sig.
strength (MPa)	Squares		Square		
Between Groups	40373.034	4	10093.258		
Within Groups	25351.795	95	266.861	37.822	0.000*
Total	65724.828	99			

181 4. DISCUSSION

182Polymethylmethacrylate (PMMA) is the most popular material used widely in various fields of Prosthodontics. 183Amongst various mechanical properties it is the impact strength and fatigue strength of this material that 184are of utmost importance and in denture base polymer they are not found to be entirely satisfactory. The 185ultimate flexural strength (also called transverse strength or modulus of rupture) of a material reflects the 186potential of material to resist fatal failure under a flexural load.^[7] Hence, high flexural strength can be 187regarded as an important determinant for the success of dentures. Compressive, tensile and shear strengths 188collectively form the flexural strength of a material. As the flexural strength increases, the force required to 189fracture the material also increases.^[10]

190Transverse strength represents the type of loading borne by a denture in the mouth. A high value of 191transverse strength of denture base acrylic is desirable as it provides a superior clinical performance by the 192dentures.^[1] Fatigue phenomenon is the low and repetitive stress rate which commonly occurs over a period of 193time. This fatigue failure is not dependent on strong biting forces. Relatively small stresses caused by 194 mastication over a period of time can contribute to formation of a small crack, which propagates through the 195denture thereby resulting in a fracture. The maximum bite forces in a patient wearing complete dentures have 196been noted up to 700 N, but these values are reduced drastically (100 - 150 N) with the removal of dentures. 197Fractures of denture occur essentially because of concentration of stresses and increased flexing.^[2] 198Recently, incorporation of nanofillers has been suggested to improve PMMA properties. The structure of 199material that has particles of a nanometer size possesses special properties. It can be rendered to the high 200ratio of surface area to volume. Amongst a variety of nanoparticles available like silver, copper, zinc, silicon, 201titanium and their oxides, titanium dioxide has gained importance recently because of its higher photocatalytic 202activity, high stability, low cost and safety towards both humans and the environment. On the other hand 203 some studies found that titanium dioxide nanoparticles did not improve the transverse strength of PMMA. This 204 could be attributed to clustering of the particles within the resin which weakened the resin. It was found that 205TiO₂ nanoparticles has an effect on thermal stability of PMMA resin and it caused a decrease in the thermal 206expansion coefficient and contraction. A decrease in elastic modulus, transverse strength and toughness was 207 reported. Addition of nanozirconia was suggested to improve the mechanical properties of PMMA. This 208 helped in increasing the impact strength, transverse strength, compressive strength, fatigue strength, as well

209as its fracture toughness and hardness. An antifungal effect has also been reported which may play a 210preventive role in patients susceptible to fungal infections.^[11]

211Use of nanoparticles is based on the principle that reduction of filler size increases the mechanical properties 212of resins. Spherical particles of titanium dioxide have been reported to improve the transverse strength as 213they increase polishability and mechanical properties. It also increased the surface hydrophobicity, reduced 214the adherence of biomolecules, aided in colouring and exhibited antimicrobial properties.^[7]

215 The quest has been on for the most suitable concentration of different nanoparticles which can be added to 216the acrylic resin so that transverse strength is improved manifolds. However, it was found that concentrations 217above 5% have led to massive changes in the colour of acrylic. Therefore, two concentrations 2.5% and 5% 218were selected.^[12]

219The increase in transverse strength at 5% concentration of nanozirconia can be attributed to the high 220interfacial shear strength between nanofiller and matrix due to formation of cross-links or supra molecular 221bonding which covers or shields the nanofillers. This in turn prevents propagation of crack. Also the crack 222propagation can be changed by improving the bonding between nanofiller and resin matrix. An increase in 223transverse strength that occurred with addition of 2.5wt% zirconium dioxide nanoparticles can be attributed to 224the uniform distribution of the very fine size of nanoparticles that allows them to enter between linear 225macromolecular chains of the polymer.^[4]

226 The SEM micrograph studies have shown good surface characteristics with different nanozirconia 227concentrations. Moreover, a uniform distribution of particles was assumed as no big agglomeration was 228found. The SEM analytical studies have revealed that as the concentration increased, the polymer matrix got 229filled with nanoparticles that stopped crack propagation, resulting in stronger material. Uniform dispersion of 230the nanoparticles into the resin matrix filled the inter-polymeric chain spaces, which shows the importance of 231the additive content of the nanoparticles.^[13]

232Increase in transverse strength on addition of titanium dioxide nanoparticles in concentration 2.5% in PMMA 233matrix can be attributed to uniform dispersion of the small sized filler particles. This is responsible for the 234improved fracture resistance of PMMA. Addition of titanium dioxide nanoparticles up to 2.5% increased the 235strength, above which the strength decreased This can be explained by adversely affected degree of 236conversion which in turn leads to increase in the level of residual unreacted monomer that acts as plasticizer. 237Incorporation of nanoparticles causes these particles to agglomerate and aggregate. The agglomerated 238compounds can act as stress concentrating center in the matrix and adversely affect mechanical properties of 239the polymerized material. It is easily noted that the content of nano additives is of critical importance.^[14]

240By using scanning electron microscopy it has been observed that, at a concentration of 2.5%, nanoparticles of 241titanium dioxide were well distributed in specimen. The particles maintained their original size and had an 242active role in improving the mechanical properties. However, when the concentration was increased to 5.0%, 243the SEM images have demonstrated that the nano-oxides had agglomerated to a different extent, which 244resulted in a decrease in the mechanical properties of the resin.

245In the *in vitro* present study only one mechanical property i.e. transverse strength was taken into 246consideration. So, further studies may be carried out *in vivo* conditions to understand its effect in oral 247environment.

248 5. CONCLUSION

249 Within the limitations of this study, following conclusions were drawn:

- Addition of titanium dioxide nanoparticles beyond the concentration of 2.5 % decreased the
 transverse strength of conventional heat polymerized acrylic resin.
- The best result was obtained after adding 5% of nanozirconia particles to the conventional heat polymerized acrylic resin.

According to the results of this *in vitro* study, it could be concluded that nanoparticles may be considered as a new approach for denture base reinforcement. The reinforcement resulted in significantly higher transverse strength as compared to unreinforced resin.

- However, only one mechanical property i.e., transverse strength was taken into consideration in this *in vitro* study. Further studies considering other mechanical, esthetic and biological properties can be carried out.
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263 COMPETING INTEREST

- 264 Authors have no competing interest
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