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Comparative evaluation of addition of carbon nano fillers and graphene and its effect on impact strength and flexural strength of heat cure denture base resin. - An in vitro study

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Abstract

Context: Polymethylmethacrylate(PMMA) is widely used for fabrication of removable dental prosthesis, but it is still insufficient to fulfil the mechanical requirements for dental applications.

Aims: Comparative evaluation of addition of carbon nano fillers and graphene and its effect on impact and flexural strength of heat cure denture base resin - an in vitro study **Methods and Material:** Characterisation of carbon nanofillers (MWCNTs) and graphene were done to check the purity, diameter and size using FT RAMAN & FESEM. Monomer without microaddition acted as control group. Nanofillers by weight were added in monomer in different percentages 0.50% by weight for group b and

group c and combination of 0.25% by weight of graphene and 0.25% by weight of carbon for group d.Nanofillers were subjected to probe sonification apparatus for uniform dispersion. Polymer and monomer containing nanofillers were mixed according to the groups and packed in the mold and processed according to conventional method.80 specimens were fabricated and divided into four groups according to the test. The retrieved specimens were kept in artificial saliva before testing. Universal maching and izod impact testing machine was used to check impact and flexural strength

Statistical analysis used: Kruskall Wallis ANOVA and Mann Whitney test.

Results: 0.5% by weight of carbon nanofillers showed highest flexural and impact strength of PMMA resin followed by graphene and by combination of 0.25% by weight of carbon and 0.25% by weight of graphene nanofillers.

Conclusions: Carbon and graphene nanofillers increases the flexural and impact strength of PMMA resin so that ideal and best material can be determined for clinical success.

Keywords: Polymethylmethacrylate.multiwall carbon nanofillers,graphene,flexural strength,impact strength.

Introduction

Polymethyl methacrylate(PMMA) is widely used as denture base material. PMMA has mechanical properties such as hardness, rigidity, biological properties, aesthetic properties. One of the most common complications of denture base prosthesis is fracture of denture either from long-term fatigue failure caused by repeated masticatory force or from extra-oral high impact force resulting from accidental dropping of the prosthesis. Risk of fracture due to accidental fall of dentures, so high impact strength is a desirable property. ²

Flexural and impact strength are important properties which are essential for strength and to increase the longevity of prosthesis. Flexural fatigue occurs after repeated flexing of a material.^{3,4} The midline fracture in dentures is often the result of flexural fatigue.³ When the patient exhibits parafunctional habits such as bruxism and clenching, the flexural strength is an essential property. In order to overcome these problems and increase the longevity and durability of the prosthesis, several attempts were made to modify and improve the strength of the PMMA like zirconia, glass fibre, alumina, tin, and copper or addition of whisker to resin like tin oxide(TiO2),zinc oxide (ZnO2),aluminium oxide(AlO2). ⁵⁻⁸Recently, much attention has been directed toward the incorporation of

inorganic nanoparticles into PMMA to improve its properties like ZrO2, TiO2, and carbon nanotube (CNT.9-¹²MWCNTs (Multiwall Carbon Nanotubes)can successfully reinforce the fracture lines by strengthening the fibrils and bridging voids to enhance the fatigue performance of the polymer. ¹⁰Graphene has attracted great interest due to its exceptional physical, chemical, thermal and electrical properties. 13,18 Combination of graphene and MWCNTs has never been used in literature. Hence, aim of this in vitro study was comparitive evaluation of effect of MWCNTs(Multiwall Carbon Nanotubes) and graphene on flexural strength and impact strength of heat cure denture base resin.

Subjects and Methods

Group A	Heat cure denture base resin						
(Control group)	without incorporation of nano						
	fillers						
GROUP B	Incorporation of 0.5% by weigh						
	of carbon Nano fillers in heat						
	cure denture base resin.						
GROUP C	Incorporation of 0.5% by weigh						
	of graphene in heat cure denture						
	base resin.						
GROUP D	Incorporation of 0.25% of						
	carbon Nano fillers and 0.25% of						
	graphene in acrylic resin.						

Table.1 Grouping of Specimens

A total 80 specimens were fabricated for testing two parameters i.e. impact strength & flexural strength and were divided into 4 groups (Table.1).10 samples for each group (n=10) for flexural strength .10 samples for each group (n=10) for impact strength

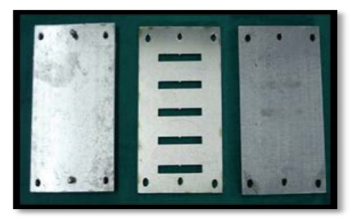
Preparation of Stainless Steel Mold

The molds were fabricated for flexural strength consisted of stainless steel plates of 2mm thickness with ten slots of dimension 60x40x2 mm (length x width x thickness)

according to $(ISO20795)^{17}(Fig.1 a)$. Mold for fabrication of samples for impact strength consisting of 5 slots were made up of dimension 80 X 10 X 4mm (length x width x thickness) $(ISO20795)^{11}(Fig.1.b)$



Fig.1 a)Fabricated molds for flexural strength



b) Impact strengthCharacterisation of Nanoparticles



Fig. 2: Graphene and Multiwall Carbon nanotubes solution-United Nanotech Innovations Pvt.Ltd Bangalore



Fig.3: FT RAMAN (Bruker's MultiRAM)



Fig.4: FE-SEM (JEOL JSM 5800; JOEl Ltd, Tokyo, Japan).

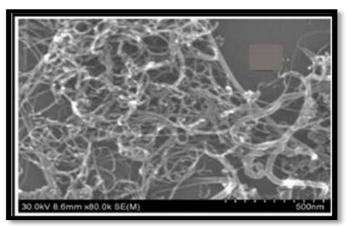


Fig. 5: FE-SEM images of MWCNT's after characterisation

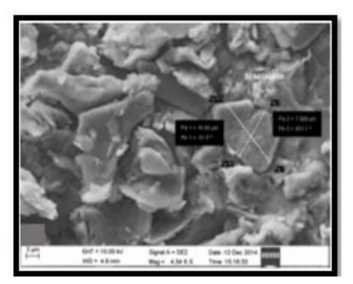
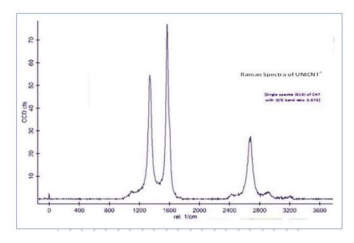
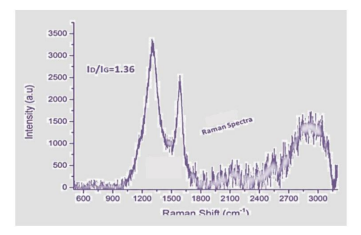


Fig. 6: FE-SEM images of graphene after characterisation Graphene and carbon nanofiller (MWCNTs) solution (Fig 2) used in this study were in ethanol dispersed form.MWCNT and graphene nanoparticles were characterized with the help of Fourier Transform Raman (FT RAMAN) & Field Emission Scanning Electron Microscopy (FE-SEM)(fig3 &4). Graphs of MWCNTs (Graph.1) and Graphene (graph.2) were verified for purity of the nanofillers.FE-SEM image of MWCNTs (Fig.5) and graphene (Fig.6) verified the size of nanoparticles .FE-SEM aids in examining small area of contamination spots.FT-RAMAN provides a fingerprint by which the molecule can be identified and also the purity of the nanofillers.



Graph1: FT RAMAN spectroscopy of MWCNT's after characterization



Graph 2: FT Raman spectroscopy of graphene after characterization

Preparation of Acrylic Denture Base Resin Specimens

Percentage of Na	Amount	Amount	Amount
fillers	Nano fillers	PMMA	monomer
	in µl		
0.5% Carbon Na	0.083 μ1	5gm	2ml
fillers			
0.5%Graphene	0.0108 μ1	5gm	2ml
Nano fillers			
0.25%Carbon Na	0.042 μ1	5gm	2ml
fillers			
0.25% Graphene	0.054 μ1	5gm	2ml
Nano fillers			

Table 2: Percentages and amounts of polymer, monomer and carbon (MWCNTs) and graphene Nano filler used in the study



Fig.7: Electron precision balance (Hindustan Scientific Linkers)



Fig.8: Nanofillers measured using Micropipette

Electronic precision balance (Fig.7) was used to weigh 50 gm of heat cure acrylic resin - DPI Heat Cure- Dental Products of India Ltd Mumbai, India,20ml of monomer was measured using calibrated beaker.Monomer without micro additions of nanoparticles were mixed with polymer (PMMA) in standard 1:2 ratio by weight ⁽³⁾ acted as group A (control group). For fabrication of samples of group B and C 0.5% by weight of MWCNTs(multiwall carbon nanotubes) i.e. 0.083 μ1 and 0.5% by weight graphene i.e.0.0108 μ1 were measured using micropipette (Fig.8) and 0.25% by weight of MWCNTs(multiwall carbon nanotubes) i.e. 0.042 μ1 & 0.25% by weight of graphene i.e. 0.054 μ1 respectively for group D as mentioned in (Table.2).

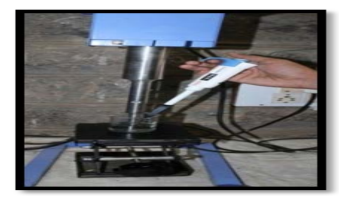


Fig.9: Nanofillers added to the beaker



Fig. 10: Ultra probe sonication



Fig.11: Mixing of the mixture

Measured nanofillers were added to a beaker containing monomer (Fig.9)Graphene particles and MWCNTs (multiwall carbon nanotubes) would be subjected to ultra probe sonication at 140W, 40 KHz for 3mins according to the groups (Fig.10) Probe sonication apparatus (Ultrasonicator Apparatus, Pci Analytics Pvt. Ltd) is used for breaking them into individual nano crystal and for uniform dispersion in the monomer (methyl methacrylate) to prevent agglomeration Monomer containing MWCNTs (multiwall carbon nanotubes) and graphene of different concentration by weight of respective groups were mixed with polymer (PMMA) in standard 1:2 ratio by weight (3). (Fig.11)The jar was kept closed till the mix attained the dough stage. Petroleum jelly was applied to the porcelain mixing jar and molds. The resin dough was removed from the mixing jar and loaded in the mold. The mold was reassembled and bench pressed incrementally pressure was applied 1500 psi until metal to metal contact of the mold was achieved. The mold was clamped and kept overnight for bench curing. The samples were processed

by conventional curing method as per American Dental Association (ADA)specification no 12.



Fig. 12: Finishing of the specimens



Fig.13: Samples kept in artificial saliva

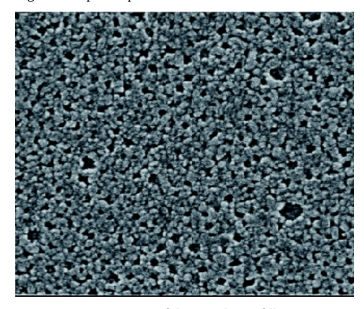


Fig.14: FESEM images of dispersed nanofillers in resin After curing this assembly was bench cooled, the specimens were retrieved and any excess flash was trimmed off and finishing of samples was done (Fig.12). Artificial saliva (Spinco Biotech Pvt.Ltd,Chennai) (Fig

1.c) was used to remove any impurity from the samples and to standardize the procedure. Samples were kept in artificial saliva for 48hrs before testing (Fig.13). Before testing the samples for impact and flexural strength, the samples were evaluated for uniform dispersion of nano fillers in PMMA resin with the help of FE-SEM. (Fig.14)

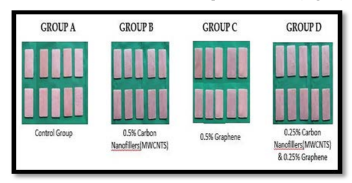


Fig.15:Samples Of Different Group for flexural strength

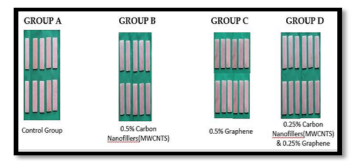


Fig.16: Samples of Different Group for impact strength

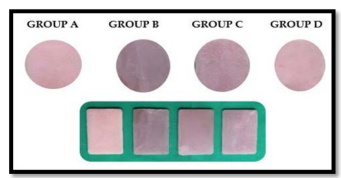


Fig.17: Samples Of Different Group showing color variation

Four study groups each contained about 20 specimens, with different concentration of nanoparticle in specimens and control group for flexural strength and impact strength. (Fig.15,16)



Fig.18: Universal testing machine for Three-point flexural strength test.





Fig 19: Izod Impact Testing Machine for Izod impact strength test.

For flexural strength, Universal Testing Machine, Star Testing System, India .Model No. STS 248(Fig.18) was used. The distance between the specimen supports was 40 mm and the loading force was applied to the specimens at a crosshead speed of 5 mm/min until the specimens fractured. The maximum load exerted on the specimens was recorded. Impact strength test was conducted following the procedure given by the (ISO20795)¹⁷ with Izod impact testing machine (Fig.19).

Data was subject to statistical analysis using Statistical package for social sciences (SPSS v 22.0, IBM). Comparison of differences in means of flexural and impact strengths between the 4 groups was done using Kruskall Wallis ANOVA, followed by Mann Whitney test for pair-wise comparisons. For all the statistical tests,

p<0.05 was considered to be statistically significant, keeping α error at 5% and β error at 20%, thus giving a power to the study as 80%.

Results

Table 3: Comparison of mean flexural strengths of four groups evaluated by Kruskal-Wallis test.

	GROUPS	N	Mean (Mpa)	Std. Deviation	p value by Kruskal-Wallis ANOVA		
	Group A(Control)	10	58.29	1.862			
FLEXURAL STRENGTH	Group B (0.5% MWCNTs)	10	68.86	.884	< 0.001**		
	Group C (0.5%Graphene)	10	64.96	.953	HS		
	Group D (0.25% MWCNTs + 0.25%Graphene)	10	62.45	.935			

Higher flexural strength were of samples of Group B (heat cure denture base resin incorporated with 0.5% of carbon nanofillers (MWCNTs) by weight) 68.86 Mpa followed by Group C (heat cure denture base resin incorporated with 0.5% of graphene by weight) 64.96 Mpa, Group D (heat cure denture base resin incorporated with0.25 % of carbon nanofillers (MWCNTs) by weight and 0.25% of graphene by weight) 62.45 Mpa Group A control group 58.29 Mpa. There is a highly significant difference between the means of different groups (p<0.001).

Table 4: Comparison of impact strength of four groups evaluated by Kruskal-Wallis test.

	GROUPS	N	Mean (kJ/m ²⁾	Std. Deviation	p value by Kruskal-Wallis ANOVA
	Group A (Control)	10	7.65	.461	
Impact Strength	Group B (0.5% MWCNTs)	10	10.62	.858	< 0.001** HS
	Group C (0.5%Graphene)	10	8.52	.433	113
	Group D (0.25% MWCNTs + 0.25%Graphene)	10	9.81	.624	

The mean impact strength of all group specimens as tabulated in (Table 4)demonstrate higher impact strength for Group B (heat cure denture base resin incorporated with 0.5% of carbon nanofillers (MWCNTs) by weight) 10.62 kJ/m² followed by Group D (heat cure denture base resin incorporated with 0.25 % of carbon nanofillers (

MWCNTs) by weight and 0.25% of graphene by weight) 9.81 kJ/m², Group C (heat cure denture base resin incorporated with 0.5% of graphene by weight) 8.52 kJ/m² and Group A, control group 7.65 kJ/m². There is a highly significant difference between the means of different groups (p<0.001).

Table 5: Comparison of differences in means of flexural and impact strengths between the four groups by Mann Whitney test for pair-wise comparisons.

		N Mean		Std. Deviation	Std. Error	p value of Mann Whitney test	
	Groups						
	Group A (Control)	10	58.29	1.862	.854	Group A Group B	0.000**
Flexural Strength	Group B (0.5% MWCNTs)	10	68.86	.884	1.092	Group A Group C	0.000**
	Group C (0.5%Graphene)	10	64.96	.953	.533	Group A Group D	0.028*
	Group D (0.25%MWCNTs+ 0.25% Graphene)	10	10 62.45 .9.	.935	.889	Group B Group C	0.000**
						Group B Group D	0.000**
						Group C Group D	0.033*
	Group A (Control)	10	7.65	.461	.299	Group A Group B	0.000**
Impact Strength	Group B (0.5% MWCNTs)	10	10.62	.858	.178	Group A Group C	0.024*
	Group C (0.5%Graphene)	10	8.52	.433	.194	Group D	0.000**
	Group D (0.25%MWCNTs+ 0.25% Graphene)	10	9.81	.624	.224	Group B Group C	0.000**
						Group B Group D	0.001**
						Group C Group D	0.000**

^{**=}statistically highly significant difference (p<0.001)

Discussion:

PMMA has been established as principal material in denture base construction. Nevertheless it is generally recognized that despite fulfilling aesthetic requirements, the fracture strength of PMMA are not entirely satisfactory and this is reflected by the expenditure on a large number of denture repairs annually. Most fractures of dentures occur during function, primarily from denture resin fatigue. Because of the risk of fracture, if patients drop their dentures, high impact strength is a desirable property. Recently, much attention has been directed toward the incorporation of inorganic nanoparticles into PMMA to improve its properties. Nanomaterials have been developed promptly and some researches of nanomaterials have been carried out in prosthodontics. Many of the current dental materials are available through nano

crystallization to improve their original performance and play a key role in oral applications.

Nano composite denture base has higher interfacial shear between the resin bond strength matrix nanomaterial's, compared to the conventional resin matrix. It creates thick interface, which enhances the bond between the resin molecules and creates higher molecular weight polymers.⁹ Because of the interface and cross linking polymerisation there is less chances of leaching of residual monomer from the resin.CNTs are strong, resilient, lightweight, and usually form stable cylindrical structures. CNTs have high mechanical properties with reported strengths 10 to 100 times higher than steel at a fraction of the weight. 14CNTs that have a flawless structure are classified into 2 main types, namely single walled and multiwall CNTs .Single-walled CNTs (SWCNTs) consist of a single graphite sheet seamlessly wrapped into a cylindrical tube, and multiwall CNTs (MWCNTs) have an array of such nanotubes concentrically nested like the rings of a tree trunk. Singlewalled CNTs (SWCNTs) are costly and less available.

The addition of carbon fibres to a matrix not only gives strength and elasticity to the material but also improves toughness .The stress transfer efficiency can be 10 times higher than that of traditional additives. ¹⁴Multi-walled carbon nanotubes (MWCNTs) have unique atomic structure, and extra ordinary mechanical properties, making ideal reinforcing materials. ¹⁵ An efficient exploitation of the CNT properties in polymers is related to their homogenous dispersions in the matrix or an exfoliation of the agglomeration and a good wetting with the polymer. Various dispersions methods (stirring, extrusion, sonication, etc.) for the distributions of CNTs (carbon nanotubes) in polymers have been used. A common technique used in order to disperse CNTs is the sonication's technique. ^{12,13,14}

^{* =} statistically significant difference (p<0.05)

A pulsed ultrasound exfoliates agglomerates and disperses CNTs in the matrix effectively. To take full advantage of these unique mechanical properties, optimization of nanotube-polymer interface properties such as wet ability and adhesion is required. In pure PMMA fibres, polymer necking occurs under increasing tension, which results in failure at relatively small strains. However, adding CNTs to a polymer may dramatically improve the resistance of the polymer to mechanical failure. Incorporating MWCNTs to polymer matrices may effectively bridge cracks and reduce the extent of plastic deformation by a PMMA matrix.¹⁴

MWCNTs can successfully reinforce the fracture lines by strengthening the fibrils and bridging voids to enhance the fatigue performance of the polymer. In the last years, graphene has attracted great interest due to its exceptional physical, chemical, thermal and electrical properties. This material can be described as a single layer of pure carbon only one atom thick.

Graphene is flexible, practically transparent, very strong and biocompatible. Graphene shows superior mechanical properties including high fracture resistance, excellent mechanical strength, high Young's Modulus (1TPa). 18 Furthermore, it is biocompatible, and very light, it has a large surface area, and the reinforcement of denture base material has been a subject of interest to the dental material community. The effects of CNT and graphene reinforcement on some mechanical properties of denture base materials have not been explored. 13,17 This investigation studied the effect of MWCNT reinforcement on the mechanical properties of a commonly used PMMA denture base material. The null hypothesis that the addition of multiwall carbon nanotubes (MWCNTs) by weight and graphene by weight would not improve the flexural strength and impact strength of the prosthesis was rejected.

Multi wall carbon nanotube and graphene was chosen because limited studies are there in literature and also it is known for its high mechanical properties, like flexural strength impact strength, hardness. Multiwall carbon nanotube and graphene was incorporating concentrations 0.5% by weight of MWCNTs and graphene by weight and combination of 0.25%MNCT and graphene. 0.5% was selected because studies have shown that concentration higher than 0.5% leads to more discoloration of PMMA resin and also it leads to decrease in strength of the resin.25 Higher concentrations of nano fillers will lead to flexural and impact strength deterioration of the resin material. This is attributed to higher filler content above saturation point at which the resin cannot incorporate further filler particles. Any attempt to add filler particles after reaching saturation of matrix leads to interruption in the resin matrix continuity and thus causing a decrease in the strength of reinforced specimens. These findings are consistent with those reported by Sodagar.A and Mars.B. 15,17

The increase in impact strength and flexural is due to the interfacial shear strength between nano-filler and matrix is high due to formation of cross-links or supra molecular bonding which cover or shield the Nano fillers that in turn prevent propagation of crack. Mars and Pienkowski ¹⁵ studied that CNT effectively bridge the cracks, also CNT is strong and stable because, carbon in nanotubes are arranged in hexagonal ring. This lead to a reduction in segmental motion thus increasing the impact strength and flexural strength

Nano fillers have tendency to agglomerate when incorporated with heat cure resin directly to prevent this ultrasonification prevent agglomeration of nanoparticles and leads to uniform dispersion of nanoparticles in monomer by using probe ultrasonification apparatus. Agglomeration causes cluster formation of

nanoparticles in PMMA resin leading to decrease in strength of the PMMA resin. The samples were stored in artificial saliva for 48hrs to simulate oral environment and to remove any impurity before testing. ²⁰

Incorporation of nano fillers increases flexural and impact strength of heat cure denture base resin. Highest impact and flexural strength was shown by incorporation of 0.5% by weight of carbon nano fillers (MWCNTs), but there was more colour change in the specimen compared to other group i.e. 0.5% graphene incorporated in heat cure denture base resin and combination of 0.25% of carbon nanofillers (MWCNTs) and 0.25% of graphene by weight incorporated in heat cure denture base resin (Fig.17). The colour change of group B specimen i.e heat cure denture base resin incorporated with 0.5% by weight of carbon nano fillers (MWCNTs) was more compared to other groups can be used in patients with darker complexion for characterisation. Heat cure denture base resin incorporated with 0.5% of graphene or combination of 0.25% of carbon nanofillers (MWCNTs) and 0.25% of graphene by weight improved flexural strength and impact strength as well as less colour change in the specimen as compared to control group, so they can be used in patients with fair complexion.

Conclusion

Within the limitations of this study, following conclusions can be drawn:

- 1. 0.5% by weight of nano fillers increases flexural and impact strength of heat cure denture base resin
- 2. Highest impact and flexural strength was shown by incorporation of 0.5% by weight of carbon nano fillers (MWCNTs), but there was more colour change in specimen compared to other group I.e. 0.5% graphene and combination of 0.25% of carbon nano fillers(MWCNTs) and 0.25% of graphene by weight.but it can be used in patients with darker complexion for characterisation.

3. 0.5% of Graphene or combination of 0.25% of carbon nano fillers(MWCNTs) and 0.25% of graphene by weight can be used to increase the flexural and impact strength of heat cure denture base resin as well as in patients with fair complexion.

Further in-vivo studies using these materials are recommended to substantiate these results and for the characterisation of the prosthesis in patient with darker complexion so that ideal and best material can be determined for clinical success.

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