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CA 2547073 C 2013/04/02

(11)(21) **2 547 073**

(12) **BREVET CANADIEN
CANADIAN PATENT**

(13) **C**

(86) Date de dépôt PCT/PCT Filing Date: 2004/11/23
(87) Date publication PCT/PCT Publication Date: 2005/06/16
(45) Date de délivrance/Issue Date: 2013/04/02
(85) Entrée phase nationale/National Entry: 2006/05/25
(86) N° demande PCT/PCT Application No.: US 2004/039555
(87) N° publication PCT/PCT Publication No.: 2005/053621
(30) Priorité/Priority: 2003/11/26 (US10/722,830)

(51) Cl.Int./Int.Cl. *A61K 6/083* (2006.01),
A61C 13/00 (2006.01), *A61K 6/087* (2006.01)
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(54) Titre : METHODE DE FABRICATION DE PROTEGE-DENTS ET D'ECLISSES DENTAIRES POUR LA NUIT
(54) Title: METHOD OF MAKING DENTAL NIGHT GUARDS AND SPLINTS

(57) **Abrégé/Abstract:**

The invention provides a method of shaping a dental product, comprising: shaping a curable polymerizable composition to form a shape of a dental product. The polymerizable composition is adapted to cure to form polymeric material, which at 37°C has a flexural modulus of less than 250,000 psi and a flexural strength of less than 7,000 psi.



(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property
Organization
International Bureau



(43) International Publication Date
16 June 2005 (16.06.2005)

PCT

(10) International Publication Number
WO 2005/053621 A1

(51) International Patent Classification⁷: **A61K 6/083**,
A61C 13/00, A61K 6/087

(21) International Application Number:
PCT/US2004/039555

(22) International Filing Date:
23 November 2004 (23.11.2004)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
10/722,830 26 November 2003 (26.11.2003) US

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(81) Designated States (unless otherwise indicated, for every
kind of national protection available): AE, AG, AL, AM,
AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN,
CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI,
GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE,
KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD,
MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG,
PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM,
ZW.

(84) Designated States (unless otherwise indicated, for every
kind of regional protection available): ARIPO (BW, GH,
GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM,
ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM),
European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI,
FR, GB, GR, HU, IE, IS, IT, LU, MC, NL, PL, PT, RO, SE,
SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ,
GW, ML, MR, NE, SN, TD, TG).

Published:
— with international search report
— before the expiration of the time limit for amending the
claims and to be republished in the event of receipt of
amendments

For two-letter codes and other abbreviations, refer to the "Guid-
ance Notes on Codes and Abbreviations" appearing at the begin-
ning of each regular issue of the PCT Gazette.

(54) Title: METHOD OF FORMING A DENTAL PRODUCT

(57) Abstract: The invention provides a method of shaping a dental product, comprising: shaping a curable polymerizable compo-
sition to form a shape of a dental product. The polymerizable composition is adapted to cure to form polymeric material, which at
37°C has a flexural modulus of less than 250,000 psi and a flexural strength of less than 7,000 psi.

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METHOD OF MAKING DENTAL NIGHT GUARDS AND SPLINTS

[0001] Intentionally left blank

[0002] The invention relates to method of making dental night guards and splints. The invention provides an improved method of making a dental night guard and a dental splint, which are light curable.

[0003] The invention uses dental compositions, which are preformed and/or to be formed into tooth restorative fillings, adhesives, cements, sealants, crowns, bridges, full dentures, partial dentures, denture liners, custom trays, artificial teeth, repairs for natural teeth, veneers, denture repairs, denture reline, night guards, splints, retainers, orthodontic components, provisional dental devices, inlays, onlays, orthodontic appliances, oral orthopedic appliances, temporary dentures, temporary partial dentures, maxillofacial prostheses, obturators, and ocular prostheses.

[0004] The invention uses dental compositions, which are adapted to be light cured to form dental night guards and dental splints in patients' mouths. The flexibility and softness of these dental night guards and dental splints enables them to be readily removable from a patient's mouth without discomfort to the patient. Light curing of the polymerizable materials used to form these dental night guards and dental splints enables them to be readily shaped to the form of a patient's mouth. Preferably they are at least partially cured before removing them from the mouth to better retain their shape. After being removed from the mouth they may be further shaped and further cured.

[0005] Beneficially, light curing polymerizable material allows the user to, at least partially, cure the material in the mouth, when it has the most desired shape. Thus, the user has selective control of the curing of the material. Advantageously, selected separate portions (amounts, zones, regions or areas) of the material may be effectively partially or completely polymerized without effectively partially or completely polymerizing (or curing) the remainder of the material to be formed into a dental night guard or dental splint. Thus, the portions polymerized retain their form, while the

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remainder of the material may be further shaped and formed to improve the fit of the dental night guard or dental splint to the patient.

[0006] Dental compositions useful in accordance with the invention may include organic or inorganic filler particles or fibers. Dental compositions useful in accordance with the invention preferably include more than 5 percent by weight of one or more semicrystalline polymerizable compounds. Dental compositions useful in accordance with the invention preferably are dimensionally stable below 36°C and effectively fluid above 40°C.

[0007] “Flexural strength”, and “flexural modulus” as used herein refers to results of testing according to ASTM D790 (1997). “Compliance” as used herein refers to the inverse of flexural modulus.

[0008] Dental compositions useful in accordance with the invention may include from 0 to about 95 percent by weight filler particles. In a preferred embodiment of the invention polymerizable dental compositions useful in accordance with the invention include from about 5 to about 90 percent by weight filler. More preferably, these polymerizable dental materials include from about 20 to about 85 percent by weight filler. Most preferably, these polymerizable dental materials include from about 40 to about 80 percent by weight filler.

[0009] The filler particles preferably include organic and/or inorganic particles, and preferably reduce polymerization shrinkage, improve wear resistance and modify the mechanical and physical properties. Preferred fillers are polymers, glasses formed from or including, barium, calcium, strontium, lanthanum, tantalum, and/or tungsten silicates and aluminates and/or aluminosilicates, silica, quartz, ceramics, nanoparticles. Preferably the filler particles have a range of particle sizes of from 0.01 micrometers to 100 micrometers.

[0009a] Specific aspects of the invention include:

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a method of shaping a flexible dental product selected from the group consisting of nightguards and splints, comprising the steps of: shaping a first light curable polymerizable composition to form a dimensionally shape-stable, uncured layer over a cast model of a patient's teeth, the first light curable polymerizable
5 composition comprising a semicrystalline material; irradiating said first light curable polymerizable composition to form a partially cured shaped layer over the cast model; applying a second light curable polymerizable composition to the partially cured shaped layer; shaping the second light curable polymerizable composition to form a dimensionally shape-stable, partially cured dental product over the cast model of the
10 patient's teeth; irradiating the second light curable polymerizable composition and the partially cured shaped layer to form a cured flexible dental polymeric product, said product having a flexural modulus of less than 250,000 psi and a flexural strength of less than 9,000 psi at 23° C., and removing the flexible dental product from the cast model; and

15 a method of shaping a flexible dental product selected from the group consisting of nightguards and splints, comprising the steps of: shaping a first light curable polymerizable composition to form a portion of a dental product over a cast model of a patient's teeth, irradiating said first light curable polymerizable composition to form a partially cured flexible portion of a dental polymeric product; shaping a
20 second light curable polymerizable composition over the partially cured flexible portion of the dental polymeric product; irradiating the partially cured flexible portion of the dental polymeric product and the second light curable polymerizable composition to form a cured flexible dental polymeric product, said cured flexible dental polymeric product at 37° C. has a flexural modulus of less than 80,000 psi and
25 a flexural strength of less than 7,000 psi, and removing the flexible dental product from the cast model; wherein the first light curable polymerizable composition is different from the second light curable polymerizable composition.

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Detailed Description of the Invention

[0010] Preformed curable dental devices and sheets for making curable dental devices may be stored in packages or below solidification temperatures. For example, preformed curable dental devices and sheets for making curable dental devices may be stored at solidification temperatures at and below 35°C.

[0011] Flexible dental polymeric material useful in accordance with the invention has a flexural modulus in order of increasing preference of less than 350,000; 300,000; 250,000; 200,000; 170,000 psi, and a flexural strength in order of increasing

preference of less than 9,000; 8,000; 7,000; or 6,000 psi at 23°C. Flexible dental polymeric material useful in accordance with the invention has a flexural modulus in order of increasing preference of less than 250,000; 200,000; 150,000; 100,000; 80,000; 50,000 or 40,000 psi, and a flexural strength in order of increasing preference of less than 7,000; 6,000; 5,000; 4,000; 3,000 or 2,000 psi at 37°C.

[0012] Flexible dental polymeric material useful in accordance with the invention has a flexural modulus in order of increasing preference of at least 10,000; 20,000; or 50,000 psi, and a flexural strength in order of increasing preference of at least 100; 200; or 500 psi at 23°C. Flexible dental polymeric material useful in accordance with the invention has a flexural modulus in order of increasing preference of at least 5,000; 10,000; or 25,000 psi, and a flexural strength in order of increasing preference of at least 20; 50; or 100 psi at 37°C.

[0013] Dental polymerizable material useful in accordance with the invention is shaped and polymerized to form a night guard and/or a splint. Preferably polymerizable material useful in accordance with the invention is dimensionally stable and moldable under finger pressure at 23°C-37°C. Preferably while it is at 45°C it is flowable. Preferably it cools sufficiently from 45°C within 0.1 to 20 minutes in an air atmosphere at 23°C to become dimensionally stable.

[0014] The invention provides a method of shaping a polymerizable composition to form a shape of a dental product. The polymerizable composition is adapted to form polymeric material, which at 37°C has a flexural modulus of less than 250,000 psi and a flexural strength of less than 8,000 psi. Preferably the polymerizable composition effectively does not flow without application of more than 0.001 psi of pressure on the composition to force the composition to flow, and the composition is moldable by a pressure of at least 1 psi. Preferably the flexural modulus of the polymeric material and the flexural strength of the polymeric material are at least 20 percent less at 37°C than at 23°C. More preferably the flexural modulus of polymeric material and the flexural strength of the polymeric material are at least 50 percent less at 37°C than at 23°C. Most preferably the flexural modulus of the polymeric material and the flexural strength of the polymeric material are at least 70 percent less at 37°C than at 23°C.

[0015] Preferably polymeric material useful in accordance with the invention has a flexural modulus at 37°C of less than 100,000 psi and a flexural strength at 37°C of

less than 5,000 psi. More preferably the polymeric material at 37°C has a flexural modulus of less than 60,000 psi and a flexural strength of less than 3,000 psi. Most preferably the polymeric material at 37°C has a flexural modulus of less than 40,000 psi and a flexural strength of less than 2,000 psi.

[0016] Dental polymeric material useful in accordance with the invention has a compliance in order of increasing preference of at least 4×10^{-6} in²/pound; 1×10^{-5} in²/pound; or 2×10^{-5} in²/pound at 37°C.

[0017] The invention provides a method of shaping a polymerizable composition to form a shape of a polymerizable dental product. The polymerizable composition is adapted to form polymeric material, which at 23°C preferably has a flexural modulus of less than 250,000 psi and a flexural strength of less than 9,000 psi.

[0018] The polymerizable composition is adapted to form polymeric material, which at 37°C preferably has a flexural modulus of less than 250,000 psi and a flexural strength of less than 8,000 psi.

[0019] In the following examples, unless otherwise indicated, all parts and percentages are by weight; Lucirin™ TPO refers to 2,4,6-trimethylbenzoyldiphenylphosphine oxide made by BASF.

PREPARATION 1 – Preparation of TBDMA Oligomer

[0020] A reactor is charged with 1176 grams of trimethyl-1,6-diisocyanatohexane (5.59 mol) and 1064 grams of bisphenol A propoxylate (3.09 mol) under dry nitrogen flow and heated to about 65°C under positive nitrogen pressure. To this reaction mixture, 10 drops of catalyst dibutyltin dilaurate are added. The temperature of the reaction mixture is maintained between 65°C and 140°C for about 70 minutes and followed by additional 10 drops of catalyst dibutyltin dilaurate. A viscous paste-like isocyanate end-capped intermediate product is formed and stirred for 100 minutes. To this intermediate product, 662 grams (5.09 mol) of 2-hydroxyethyl methacrylate and 7.0 grams of BHT as an inhibitor are added over a period of 70 minutes while the reaction temperature is maintained between 68°C and 90 ° C. After about five hours stirring under 70°C, the heat is turned off, and TBDMA oligomer is collected from the reactor as semi-translucent flexible solid and stored in a dry atmosphere.

PREPARATION 2 – Preparation of Monomer

[0021] A reaction flask is charged with 151.25 grams of octadecyl isocyanate and heated to about 70°C under a positive nitrogen pressure. To this reactor are added 125.3 grams of caprolactone 2-(methacryloyloxy)ethyl ester, 0.12 gram of catalyst dibutyltin dilaurate and 0.58 grams of butylated hydroxy toluene (BHT). The addition is slow and under dry nitrogen flow over a period of two hours. The temperature of the reaction mixture is maintained between 70°C and 85°C for another 2.5 hours, the reaction product is discharged as clear liquid into plastic containers and cooled to form a semi-opaque solid and stored in a dry atmosphere.

Example 1. Polymerizable Material

[0022] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 100.0 grams of TBDMA oligomer prepared by following the procedure of Preparation 1, 0.35 gram of 2,4,6- trimethylbenzoyldiphenylphosphine oxide, (Lucirin TPO made by BASF).

Example 2. Polymerizable Material

[0023] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 100.0 grams of TBDMA oligomer prepared by following the procedure of Preparation 1, 0.25 gram of 2,4,6- trimethylbenzoyldiphenylphosphine oxide (Lucirin TPO made by BASF) and 1.0 gram of visible light initiating solution. Visible light initiating solution contains 13.3% camphorquinone (CQ), 23.0% methacrylic acid (MAA), 1.3% butylated hydroxytoluene (BHT), 46% N, N-dimethylaminoethylneopentyl acrylate, 16.3% γ -methacryloxypropyltrimethoxysilane and 66.7% 1,6-hexanediol dimethacrylate (HDDMA).

Example 3. Polymerizable Material

[0024] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 94.0 grams of TBDMA oligomer prepared by following the procedure Preparation 1, 3.65 grams of octadecyl acrylate (ODA) and 2 grams of monomer prepared by following the procedure of Preparation 2, 0.25 gram of 2,4,6- trimethylbenzoyldiphenylphosphine oxide, (Lucirin TPO made by BASF) and 1.0 gram of visible light initiating solution. Visible light initiating solution contains

13.3% camphorquinone (CQ), 23.0% methacrylic acid (MAA), 1.3% butylated hydroxytoluene (BHT), 46% N, N-dimethylaminoethylneopentyl acrylate, 16.3% γ -methacryloxypropyltrimethoxysilane and 66.7% 1,6-hexanediol dimethacrylate (HDDMA).

Example 4. Polymerizable Material

[0025] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 65.0 grams of TBDMA oligomer prepared by following the procedure of Preparation 1, 35.0 grams of monomer prepared by following the procedure of Preparation 2, 4 grams of octadecyl acrylate (ODA), 0.35 gram of 2,4,6-trimethylbenzoyldiphenylphosphine oxide (Lucirin TPO, made by BASF), and 0.65 gram of visible light initiating solution. Visible light initiating solution contains 13.3% camphorquinone (CQ), 23.0% methacrylic acid (MAA), 1.3% butylated hydroxytoluene (BHT), 46% N, N-dimethylaminoethylneopentyl acrylate, 16.3% γ -methacryloxypropyltrimethoxysilane and 66.7% 1,6-hexanediol dimethacrylate (HDDMA).

Example 5. Polymerizable Material

[0026] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 62.3 grams of TBDMA oligomer prepared by following the procedure of Preparation 1, 26.0 grams of monomer formed by following the procedure of Preparation 2, 2.0 grams of octadecyl acrylate (ODA), 9.0 grams of Genomer™ 4246 (urethane acrylate sold by Rahn AG), 0.35 gram of 2,4,6-trimethylbenzoyldiphenylphosphine oxide (Lucirin TPO, made by BASF), and 0.35 gram of visible light initiating solution. Visible light initiating solution contains 13.3% camphorquinone (CQ), 23.0% methacrylic acid (MAA), 1.3% butylated hydroxytoluene (BHT), 46% N, N-dimethylaminoethylneopentyl acrylate, 16.3% γ -methacryloxypropyltrimethoxysilane and 66.7% 1,6-hexanediol dimethacrylate (HDDMA).

Example 6. Polymerizable Material

[0027] A light curable polymerizable material is prepared by stirring at 85°C a liquid of 38.75 grams of TBDMA oligomer prepared by following the procedure of

Preparation 1, 40.0 grams of monomer prepared by following the procedure of Preparation 2, 20 grams of CN962 urethane acrylate (sold by Sartomer Company, Inc.), 0.25 gram of 2,4,6- trimethylbenzoyldiphenylphosphine oxide (Lucirin TPO, made by BASF), and 1.0 gram of visible light initiating solution. Visible light initiating solution contains 13.3% camphorquinone (CQ), 23.0% methacrylic acid (MAA), 1.3% butylated hydroxytoluene (BHT), 46% N, N-dimethylaminoethylneopentyl acrylate, 16.3% γ -methacryloxypropyltrimethoxysilane and 66.7% 1,6-hexanediol dimethacrylate (HDDMA).

Example 7. Preformed Polymerizable Laminated Splint with Flexible and rigid layers

[0028] 15 g of polymerizable material formed by following the procedure of Example 3 is formed into a 2 mm thick sheet, warmed to 37°C and then shaped in a splint shaped mold. Then 15 g of polymerizable material formed by following the procedure of Example 4 is formed into a 2mm thick sheet, warmed to 37°C and then applied onto the splint shaped sheet to form a preformed polymerizable laminated splint having flexible and rigid layers.

Example 8. Flexible Polymeric Night guard

[0029] A) A two part plaster cast model of a patient's teeth is formed and coated with a release agent. 15 grams of polymerizable composition formed by following the procedure of Example 5 is formed into a 2 mm thick sheet, applied over the cast two part model, and shaped using finger pressure and trimming to form a polymerizable night guard. The polymerizable composition is further adapted to the two part model as the model is articulated.

[0030] B) Air barrier coating (sold by Dentsply International under the trademark: Eclipse) is painted onto the polymerizable night guard and onto the cast. While on the cast the polymerizable night guard is then cured for 10 minutes in a light curing unit (sold by Dentsply International under the trademark: Eclipse). The polymeric night guard formed is clear. It is then washed with water to remove all traces of Air Barrier Coating. The night guard is then finished and polished.

Example 9. Laminated Polymeric Night Guard Having Flexible And Rigid Layers

[0031] A shaped rigid layer of a night guard is prepared by following the procedure of Example 8 part A, except that 10 g of the product composition formed by following the procedure of Example 4 is used in place of that of Example 5. Then the shaped rigid layer of a night guard is cured in a light unit (sold by Dentsply International under the trademark: Eclipse) for 4 minutes. 5 grams of the product composition of Example 1 is applied to the partially cured shaped rigid layer of a night guard while warm. The applied composition is shaped using finger pressure and trimming to form a night guard. Mold release agent is applied to the incisal and occlusal surface of the opposing cast model. The night guard is examined and adjusted to fit articulator. Then the two layer night guard is cured on the cast model by following the procedure of Example 8 part B.

Example 10. Night guard having Flexible and rigid Layers Formed from a Polymerizable Laminate

[0032] A plaster cast of a patient's teeth is coated with a release agent. 20 grams of the laminated product composition formed by following the procedure of Example 7 is applied over the cast. The composition is shaped using finger pressure and trimming to form a night guard. When adapting resin to articulated models, mold release agent is applied to the incisal and occlusal surface of the opposing model. Then the two layer night guard is cured on the cast model by following the procedure of Example 8 part B.

Example 11. Flexible Night Guard Partially Cured in the Mouth

[0033] 15 grams of the product composition formed by following the procedure of Example 5 is shaped into an arch and applied over teeth arch in patient's mouth. The composition is shaped using finger pressure and trimming to form a night guard. After the night guard is examined and adjusted to fit inside the mouth, it is partially cured in the mouth for 2 minutes using a handheld light (sold by Dentsply International under the trademark: ProLite). The partially cured night guard is then removed from the mouth. Then the partially cured night guard is cured on the model by following the procedure of Example 8 part B.

Example 12. Flexible Night Guard Partially Cured in the Mouth

[0034] A night guard is cured on a model by following the procedure of Example 11, and pouring fast-set plaster into the back of partially cured night guard arch to form a model, just prior to curing on the model by following the procedure of Example 8 part B.

Example 13. Night Guard Cured Without a Model

[0035] The night guard of Example 12 is warmed in water at 70°C for 2 minute, and adapted and molded to the teeth arch in patient's mouth. Then it is removed from the patient's mouth.

Example 14. Laminated Polymeric Night Guard Having Flexible And Rigid Layers

[0036] A shaped rigid layer of a night guard is prepared by following the procedure of Example 8 part A. 5 grams of the product composition of Example 1 is applied to the shaped rigid layer of a night guard. The applied composition is shaped using finger pressure and trimming to form a night guard. Mold release agent is applied to the incisal and occlusal surface of the opposing cast model. The night guard is examined and adjusted to fit articulator. Then the two layer night guard is cured on the cast model by following the procedure of Example 8 part B.

Example 15. Night Guard Having Flexible And Rigid Layers Partially Cured In The Mouth

[0037] 15 grams of the product composition formed by following the procedure of Example 6 is shaped into an arch and applied over teeth arch in patient's mouth. Then partially cured night guard is cured 2 minutes with a hand held curing light. 15 g of clear composition prepared by following the procedure of Example 2 is molded over the surface of the partially cured night guard arch, which is then reseated back to patient's mouth and examined. The arch is adjusted to fit the teeth surface inside the mouth. Using the handheld light this layer is cured to fix its form. The partially cured night guard arch is then removed from the mouth, and cured by following the procedure of Example 8 part B, except without a model. The night guard is then warmed in boiling water and bench cooled for 1 minute, adapted and molded to the teeth arch in the patient's mouth. It is removed from the patient's mouth after it cools to 37°C.

[0038] Prior light curable night guards include those made from polymerizable compositions, such as Triad™ TranSheet, sold by Dentsply International Inc and Eclipse™ Clear Baseplate, sold by Dentsply International Inc. For any particular shape (or geometry) flexural modulus is representative of the force required to deform (or deflect) a material. As shown in Table, at 23°C the Flexural strength and the flexural modulus for a flexible night guard prepared by following the procedure of Example 5 is less than half of the flexural strength and the flexural modulus for night guards made from Eclipse Clear Baseplate and Triad TranSheet.

TABLE 1

LIGHT CURABLE NIGHT GUARDS	Flexural strength at 37°C (psi)	Flexural strength at 23°C (psi)	Flexural modulus at 37°C (psi)	Flexural modulus at 23°C (psi)
EXAMPLE 5	1,000	5,400	33,000	168,000
TRIAD TranSheet	11,000	12,500	330,000	360,000
ECLIPSE Clear Baseplate	14,100	16,400	366,000	432,000

[0039] As shown in Table 1, at 37°C the flexural strength of night guards made from Eclipse Clear Baseplate and Triad TranSheet is more than ten times the flexural strength of night guards prepared by following the procedure of Example 5. Also, as shown in Table 1, at 37 °C the Flexural modulus of night guards made from Eclipse Clear Baseplate and Triad TranSheet is about ten times the Flexural modulus of night guards prepared by following the procedure of Example 5. The low Flexural strength and low Flexural modulus of night guards made in accordance with the invention enable them to be at least partially cured in the mouth and then removed from the patient's mouth without hurting the patient.

[0040] It should be understood that while the present invention has been described in considerable detail with respect to certain specific embodiments thereof, it should not be considered limited to such embodiments but may be used in other ways without departure from the scope of the appended claims.

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CLAIMS:

1. A method of shaping a flexible dental product selected from the group consisting of nightguards and splints, comprising the steps of:

5 shaping a first light curable polymerizable composition to form a dimensionally shape-stable, uncured layer over a cast model of a patient's teeth, the first light curable polymerizable composition comprising a semicrystalline material;

 irradiating said first light curable polymerizable composition to form a partially cured shaped layer over the cast model;

10 applying a second light curable polymerizable composition to the partially cured shaped layer;

 shaping the second light curable polymerizable composition to form a dimensionally shape-stable, partially cured dental product over the cast model of the patient's teeth;

15 irradiating the second light curable polymerizable composition and the partially cured shaped layer to form a cured flexible dental polymeric product, said product having a flexural modulus of less than 250,000 psi and a flexural strength of less than 9,000 psi at 23° C., and

 removing the flexible dental product from the cast model.

20 2. The method of claim 1 wherein said flexural modulus of said polymeric product and said flexural strength of said polymeric product are at least 70 percent less at 37° C. than at 23° C.

3. The method of claim 1 wherein said polymeric product at 37° C. has a flexural modulus of less than 100,000 psi and a flexural strength of less than 3,000 psi.

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4. The method of claim 1 wherein said polymeric product at 37° C. has a flexural modulus of less than 80,000 psi and a flexural strength of less than 2,000 psi.

5. The method of claim 1, wherein said flexural modulus of said polymeric product and said flexural strength of said polymeric product are at least 20 percent less at 37° C. than at 23° C.

6. The method of claim 1, wherein said flexural modulus of said polymeric product and said flexural strength of said polymeric product are at least 50 percent less at 37° C. than at 23° C.

7. A method of shaping a flexible dental product selected from the group consisting of nightguards and splints, comprising the steps of:

shaping a first light curable polymerizable composition to form a portion of a dental product over a cast model of a patient's teeth,

irradiating said first light curable polymerizable composition to form a partially cured flexible portion of a dental polymeric product;

15 shaping a second light curable polymerizable composition over the partially cured flexible portion of the dental polymeric product;

irradiating the partially cured flexible portion of the dental polymeric product and the second light curable polymerizable composition to form a cured flexible dental polymeric product, said cured flexible dental polymeric product at 20 37° C. has a flexural modulus of less than 80,000 psi and a flexural strength of less than 7,000 psi, and

removing the flexible dental product from the cast model;

wherein the first light curable polymerizable composition is different from the second light curable polymerizable composition.

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8. The method of claim 7 wherein said flexural modulus of said polymeric product and said flexural strength of said polymeric product are at least 70 percent less at 37° C than at 23° C.

9. The method of claim 7 wherein said polymeric product at 37° C. has a
5 flexural modulus of less than 80,000 psi and a flexural strength of less than 2,000 psi.

10. The method of claim 7, wherein said flexural modulus of said polymeric product and said flexural strength of said polymeric product are at least 20 percent less at 37° C. than at 23° C.