UNITARY MOLDED GOLF BALL

FIG. 1.

HOMOGENEOUS MOLDED GOLF BALL SUITABLE FOR OFFICIAL PLAY.

FIG. 2.

RANGE BALL (WHITE THROUGHOUT)

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UNITARY MOLDED GOLF BALL


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16 Claims. (Cl. 273—218)

This invention relates to a novel molded golf ball and is a continuation-in-part of my co-pending application Ser. No. 252,588 filed Jan. 21, 1963, and now abandoned. The invention more particularly relates to a molded golf ball having all the desirable characteristics of the best conventionally wound golf balls while avoiding many of the disadvantages of the wound balls.

Golf balls have evolved from their early form which simply consisted of a stuffed cover to a highly sophisticated article which must have many very precise characteristics in order to satisfy players and be acceptable for tournament use. The only golf balls which have thus far been found acceptable for these purposes are wound balls which are made by winding a tensioned rubber thread about a suitable core, covering the wound article with a tough cover, such as of gutta-percha compounds or the like and finally painting or covering ball.

In order to comply with the rules of the game, as for example as set forth by the United States Golf Association (U.S.G.A.), golf balls must be greater than 1.680" in diameter, must weigh less than 1.620 ounces and cannot have an initial velocity of more than 255 feet per second as measured on a standard U.S.G.A. golf ball testing machine. In addition to these required characteristics, the balls must be as round as possible, and retain this roundness even after use, must be well balanced so that they fly and roll true and make a good sound when struck by the golf club, generally referred to as "click" and be able to stand the rigors of play without adverse physical effect.

The characteristics of the wound ball may be controlled by suitable selection and control of the core, the winding tension and the cover. The manufacture of wound balls is, however, relatively slow involving the fabrication of the core and the precise winding of the thread on the core by means of winding machinery which have limited capacity, the fabrication of the cover, and the application of the cover to the winding. The most expensive premium balls are generally so constructed that the ball flies at an initial velocity approaching the 255 feet per second limit. For this purpose, however, a very thin cover must be used due to its relatively lesser resilience and the same is very prone to cutting, when hit with the edge of a golf club which may render the ball no longer usable. Even the thicker covers on the sub-premium balls are prone to cutting and damage and are often rendered unsatisfactory for further use due to such cutting or damage. Throughout the evolution of the golf ball, there have been many attempts and suggestions to produce golf balls by molding as, for example, from elastomers, such as vulcanized rubber. These attempts and suggestions have included completely molded balls and balls having molded interiors and provided with a separate cover. While it can be readily appreciated that a molded golf ball could be produced much more cheaply than a wound golf ball and could eliminate many of the disadvantages of a wound ball, nevertheless, and in spite of constant and continual efforts to produce a satisfactory molded ball, none have been successfully produced, and the only acceptable balls available are of the wound type.

It has never proven feasible nor possible to mold a golf ball from an elastomer which would have the required characteristics as set forth above. Ordinary elastomers, such as rubber or the like, even when vulcanized do not have the characteristics to allow the production of a ball with sufficient hardness, compression, click, and the like. If it is attempted to increase the degree of hardness by increasing the cross-link density, caused by the vulcanization, as for example, with sulfur, the structure becomes too brittle and too inflexible and this cannot be remedied by the use of plasticizers, etc. Furthermore, in order to obtain the desired density, thick fillers and additives have to be used and all in all, it was simply not possible to obtain a ball of the required density which would still have the other necessary characteristics for a successful golf ball.

In accordance with the invention, I have surprisingly discovered that if an elastomer is cured in the presence of a relatively large amount of a monomer which is capable of acting as a cross-linking agent for the elastomer and under conditions which will additionally result in polymerization of the monomer, a structure is formed which consists of the elastomer cross-linked into a three dimensional network with a high density of long, flexible cross-links and that this structure is capable of being produced in the form of a molded golf ball having all of the required golf ball characteristics and yet overcoming the disadvantages of the wound ball.

In the accompanying drawing:

FIG. 1 shows a perspective view partially in section of a golf ball suitable for official play under the U.S.G.A. rules produced in accordance with the invention, and FIG. 2 shows a perspective view partially in section of a range ball produced in accordance with the invention.

The ball as shown in FIG. 2 differs from that shown in FIG. 1 in that the elastomer of which it is composed contains sufficient white pigment, such as titanium dioxide, so that the ball is uniformly white throughout. With this white pigment the ball may be over weight and may not be as lively as desirable for official play, but is excellently suited for driving range use.

For the production of the homogeneous molded golf balls in accordance with the invention, the elastomer is cured while being molded under heat and pressure in the form of the golf ball in the presence of the monomer, a material promoting polymerization of the monomer, as for example a free radical type polymerization initiator, such as a peroxide catalyst and preferably a filler in order to adjust the ultimate density to the desired value.

As monomers which are capable of acting as cross-linking agents for elastomers will generally polymerize or cure per se into a hard, brittle form, it was completely unexpected that the curing of the monomer within the elastomer would result in a material having the necessary toughness, resilience, etc. for forming a satisfactory molded golf ball.

The elastomer is molded into the form of the golf ball, in intimate mixture with the monomer, and preferably a filler material, in the presence of a polymerization initiator for the monomer and under sufficient heat and pressure to cure the composition.

The elastomer may be any elastomer which is capable of being cross-linked and is preferably a rubber such as a polymer, copolymer, or terpolymer of butadiene, isoprene, or chloroprene. Most preferably the elastomer is a cis-butadiene rubber containing at least 40% cis. Examples of other elastomers include chlorinated and sulfochlorinated polyethylene, amorphous polyolefins, such as polyethylene and polypropylene copolymers and terpolymers with unsaturated monomers; silicone rubbers; polypropylene oxide elastomers; butyl rubber; poly-sulfides; acetaldehyde copolymers; urethane elastomers; ethylene-vinyl acetate copolymers; styrene copolymers with acrylic and methacrylate esters of alcohols containing...
at least four carbon atoms and the like. The elastomers should initially have a sufficient molecular weight to constitute a viscous liquid.

The monomer may be any monomer which is capable of cross-linking the elastomer and undergoing further polymerization. In order to serve as cross-linking agents, the monomers generally must be in the form of difunctional compounds, i.e., compounds having two readily polymerizing groups or sites. The term "monomer" as used herein is used in its broadest sense and is not restricted to monomeric units but includes partial polymers capable of further polymerization. Examples of these monomers which are referred to herein and in the claims as cross-linking monomers include: vinyl, allyl, methallyl, furfuryl, crotyl and cyaninyl esters of the following acids: oxalic, maleic, succinic, glutaric, adipic, pimelic, suberic, azelaic, sebacic, maleic, itaconic, citric, mesaconic, furmaric, acetic, phthalic, isophthalic, terephthalic, naphthalene dicarboxylic, mellitic, pyromellitic, trimetric, acrylic, methacyrylic, cinnamic, and crotonic. Also suitable are polyamides and imides of the following acids: maleic, itaconic, acrylic, methacrylic crotonic, citraconic, acetic and cinnamic; polyol esters and anhydrides of acrylic, methacrylic, crotonic and cinnamic. Other suitable materials include di- and triallyl cyanurate; di- and triallyl isocyanurate; diallyl amine; allyl ether; allyl glycolate; di-, tri- and tetravinyl and allyl silanes. Still further suitable materials are low molecular weight reactive polymers such as polymers of butadiene, isoprene, chloroprene and epoxidized derivatives of these materials. The diacrylates and dimethacrylates, as for example butylene glycol dimethacrylate, have been proven preferable though divinyl compounds, such as divinyl benzene have also proven highly suitable.

In order to vary the density so that the finished ball will have the desired weight and will not exceed the maximum allowable weight, a filler may be required which may also serve to reinforce the composition. Any known or conventional filler may be used which should be in finely divided form, as for example in a form between about 20 and 525 and preferably about 60 and 325 mesh U.S. Standard screen size. Most preferably the filler material is in the form of a silica or silicate as fillers in this form actually act as an adjunct to the cross-linking and thus aid the same and become a more integral part of the composition additionally and a minor amount of cork should preferably be used. In addition to the silica and silicate fillers, such as fire retardants, such as fire retardant compositions containing halogenated silicas, such as calcium silicate, and the like, and other fillers in addition or in place of the silica, or silicates, such as carbon black, cork, titania, cotton flock, cellulose flock, leather fibre, plastic fibre, plastic film, leather flour, fibrous fillers, such as asbestos, glass, and synthetic fibers, metal oxides, carbonates, and talc can be used.

The polymerization initiator may be any known or conventional initiator capable of causing the cross-linking monomer to further polymerize and cross-link. Generally, the initiators are of the free radical type, such as a peroxide, perfluorosulphonic acid compounds, hydroxines, amine oxides, ionizing radiation and the like. Peroxides, such as dicumyl peroxide and other commercially available peroxides conventionally used as polymerization catalysts may be most conveniently used.

The amount of the cross-linking monomer must be within about 20 and 90% by weight of the elastomer or in order to obtain the necessary degree of cross-linking to form the required three dimensional molecular network and to give the ball its characteristics. Amounts of the cross-linking polymer, as high as 90% by weight of the elastomer, may be used; but amounts of about 40 and 75% by weight of the elastomer are preferred.

The amount of the filler material is dictated by its type and the type of the other constituents in order to obtain the required density and may vary between about 20 and 90% by weight of the elastomer, and preferably 30 and 70% of the elastomer.

The polymerization initiator need only be present in the catalytic amount required for this function and may be in general used in the amounts that the particular agent is generally used as a polymerization catalyst. In connection with peroxides, the same, for example, may be used in amounts of about 0.2-10% by weight of the elastomer.

When using the preferred components, the best results are obtained with compositions having 100 parts by weight of the cis-butadiene rubber and approximately 40-70 parts by weight of the methacrylate ester and 20-60 parts by weight fillers.

For the production of the golf balls, the ingredients should initially be intimately mixed using, for example, the usual rubber mixing rolls or a Banbury mixer until the composition is reasonably uniform. The mixing is basically done in a manner which is common in the elastomer art and there is practically no danger of over mixing. The temperature of the mixing is not critical but should, of course, be below curing temperature, and the same is generally effected at room temperature although with frequent ingredients may be slightly warmed. This again follows rubber milling practice and involves a new technique, the heating up being relatively low.

The molding is effected in mating precision hemisphere molds or dies whose molding surface is covered with multiple regular projections to give the molded ball conventional dimpled or waffled surface appearance in order to improve its aerodynamic characteristics. The molding is a simple, straight-forward operation effected in the conventional manner used in precision molding. The material, after being thoroughly mixed may be formed into slugs in the customary manner and is then molded, which may be cylindrical or any other desired shape which will facilitate the insertion in the mold should be proportioned so that the mold is fully filled. The mating halves of the mold are then closed so that the mold cavity is entirely filled. The mold halves may be held together with pressures between about 100 and 3,000 p.s.i. preferably 1000 to 2000 p.s.i. The actual pressure actually exerted on the material being molded during the molding is probably between about 50 and 1,000 and preferably 100 and 500 p.s.i. Molding temperatures may vary depending on the particular composition used and may, for example, vary between 200-400°F. When utilizing the low density polyurethane compositions, the temperature preferably should not be below about 290°F. Optimum results are obtained in the neighborhood of 300°F, with a molding time of about 10 minutes. The molding should be effected until the composition is substantially completely cured, and there is no real upper limitation on the length of the curing time except as is dictated by practical considerations. In general, curing times between about 1 and 60 minutes, and preferably 4 and 30 minutes will generally suffice. Higher temperatures may be used up to about 400°F, but a problem arises. For optimum properties, the molded ball must be homogeneously cured, that is to say the heat must have reached the center of the ball before the outside layers become completely cured. With a cold mold, this presents no problem and temperatures may reach an end point as high as 400°F. However, normally in successive molding operations when a mold is used, it is difficult to mold optimum balls at temperatures above 320°F. Since, however, there is no advantage in high temperatures, it is preferred to operate at or near 300°F, with a molding time of approximately ten minutes at this temperature. The time is not critical but nothing is gained in longer times after the curing is complete.
however, using 12.5 parts of the cork, then the compression increases to 95 and the hardness to 95b.

Example 4

The examples may be repeated using cis polyisoprene, SBR rubber, butadiene-acrylonitrile copolymer, butyl rubber and ter-polymer, in place of the cis butadiene; divinyl adipate, methylenebis-acrylamide, dialkyl phthalate, ethylene glycol-diacylate, divinyl terephthalate in place of the polybutylene glycol dimethacrylate or divinyl benzene; calcium silicate, aluminum silicate, calcium carbonate, carbon black in place of the silica; and t-butyl peroxide, cumene hydroperoxide, benzoyl peroxide, perbenzoic acid, azo bis-oxybutyronitrile, t-butyl perbenzoate, in place of the dicumyl peroxide, and in each case golf balls of excellent quality, meeting the requirements of the U.S.G.A. will be produced.

I have furthermore discovered that if the procedure described above is followed, but if a white pigment is incorporated in the mixture prior to molding, a golf ball is obtained which is excellently suited for use by golf driving ranges, i.e. practice ranges. The ball is heavier and is generally officially over-weight and is generally not as lively and does not have the initial velocity usually desired by players in ordinary and tournament play. Both these characteristics, however, enhance its desirability for driving range use. Furthermore, and of prime importance, the ball is perfectly white throughout and does not have to be painted. While it does not have as high a gloss as does a painted golf ball, its whiteness is permanent and its appearance is entirely adequate and the elimination of the need for painting further reduces driving range operational costs.

As a white pigment titanium dioxide is preferred but other white pigments, such as barium sulfate, zinc sulfide, barium carbonate, lithopone, titanium with barium base, titanium with calcium base, titanium with aluminum silicate, white lead, calcium carbonate, aluminum oxide may be used.

The amount of the white pigment should be sufficient to give the molded ball the desired white appearance, and in general amounts of 2 to 40% and preferably 5 to 10% of the total ball should be used.

The white pigment may be used in place of or in addition to a filler, such as the silica. The ball should preferably contain about 25% by weight of the elastomer filler of which 4 to 10% and preferably 12 to 20% is in the form of the white pigments, such as the titanium dioxide.

In all other respects the driving range balls are produced in the identical manner as described for the golf balls above and have the same rugged and durable characteristics.

The following examples are given by way of illustration and not limitation of the driving range golf balls:

Example 5

A mixture is prepared with 2.5 pounds of a mixture of 8 parts of the dimethacrylate of Example 1, 4 ounces of titanium dioxide and 8 pounds of the "Hysil" silica filler powder. The amount of dicumyl peroxide curing agent was 2 ounces. When molded, the ball was white with a hardness of about 80s. It is not satisfactory for tournament play, but is a useful ball for driving range use with a good click and a good feel.

Example 6

A mixture was made up of 16 ounces parts of cis butadiene polymer, 8.5 ounce parts of butylisocyanate dimethacrylate 6.5 ounce parts of "Hysil" silica, 1.5 ounce parts of dicumyl peroxide and 0.5 ounce parts of dicumyl peroxide curing agent. Molding was effected at 300° F. for about 10 minutes under a pressure of 185 p.s.i. A white ball was produced having good bounce and good click, excellently suited for driving range use.

Example 7

The procedure of Example 6 was followed, but the amount of titanium dioxide was reduced to 1 part. Molding was at 195 p.s.i. under the same conditions and balls were obtained with good bounce and click, having a hardness of 100 and a compression of 90.

All of the balls made in Examples 5 to 7 were white and could be used for driving range purposes without painting. The balls from all examples exhibit a slight sweetish odor, which to many people is not unpleasant and which is practically unnoticeable with painted balls. However, if it is desired a small amount of an odorant such as isobornyl acetate can be added.

White rubber has been described in detail with reference to certain specific embodiments, various changes and modifications which fall within the spirit of the invention and scope of the appended claims will become apparent to the skilled artisan. The invention, therefore, is only intended to be limited by the claims or their equivalents wherein I have endeavored to claim all inherent novelty.

What is claimed is:
1. A homogeneous molded golf ball of high durability and good click having the following specifications:

- Weight: 1.600-1.620 oz.
- Specific gravity: 1.04-1.12
- Hardness (Shore): 75-99
- Compression range: 40-150
- Initial velocity (USG.A.): 200-250 ft./sec.
- Size: sphere of 1.689 to 1.685" in diameter.

2. Predominantly composed of a filled elastomer highly cross-linked into a three-dimensional network with long, flexible cross links formed from a polymerized cross-linking monomer, said elastomer being one capable of being so cross-linked by said cross-linking monomer.

3. A golf ball of size, spherical shape and weight to comply with the standard rules for golfing, having a maximum diameter of about 1.69" predominantly composed of a filled elastomer highly cross-linked into a three-dimensional network with long, flexible cross links formed from a length of at least 10 atoms, said elastomer being one capable of being so cross-linked by said cross-linking monomer, said filler being present in amount ranging between about 20 and 90% by weight of said elastomer.

4. A golf ball of size, spherical shape and weight to comply with the standard rules for golfing, having a maximum diameter of about 1.69" predominantly composed of a filled rubber elastomer highly cross-linked by about 40-75% by weight of a cross-linking monomer into a three-dimensional network with long, flexible cross links formed from a length of at least 10 atoms, said elastomer being one capable of being so cross-linked by said cross-linking monomer, said filler being present in amount ranging between about 20 and 90% by weight of said elastomer.

5. A homogeneous molded golf ball of high durability and good click acceptable for official play under the U.S.G.A. rules and having the following specifications:

- Weight: 1.600-1.620 oz.
- Specific gravity: 1.11-1.12
- Hardness (Shore): 90-100
- Compression range: 70-120
- Size: sphere of 1.680-1.685" in diameter.

and predominantly composed of a filled elastomer highly cross-linked into a three-dimensional network with long, flexible cross-links having a size of at least 1 carbon atoms formed from a polymerized cross-linking monomer, said elastomer being one capable of being so cross-linked by said cross-linking monomer.
After the molding, the balls are removed from the mold and any mold mark where the molds mate may be removed, as for example, by buffing, and the ball painted and marked, is then ready for marketing and use. Painting may be effected in the conventional manner using the conventional paints used for golf balls, as for example, enameled, polyurethane, epoxy, acrylic, or vinyl paints.

The size of the mold should be such that the finished balls have a diameter greater than 1.680" and is preferably between 1.680 and 1.685" in diameter. With this size the weight of the ball may be controlled so that it is less than 1.620 ounces and preferably between 1.600 and 1.620 ounces. The ball may be molded and will retain a roundness within .01" and preferably within .001". The density of the painted ball will be less than 1.13 and will preferably be between 1.11 and 1.12. The hardness of the ball as measured on a Shore hardness test will be greater than 75a and less than 99c, and will preferably be between 90–100b. The compression, as measured on a standard golf ball compression test machine will be between 40 and 150 points and preferably between 70 and 120 points. The balls will preferably bounce between about 60 and 75% of the height from which they are dropped in accordance with the Standard Bureau test.

The initial velocity of the balls, as tested on a standard U.S.G.A. test machine will be between 200 feet per second and 255 feet per second and preferably between about 230 and 250 feet per second. In actual play, however, due to internal dampening, the balls will appear livelier and may be driven further than conventionally wound balls which show a similar initial velocity.

The balls have all the desirable play characteristics of the best conventionally wound balls, have good click, excellent feel, and as contrasted to the covered wound balls, are so highly resistant to cutting that the same may practically be considered cut-proof, and indestructible in play. Furthermore, the balls will have a perfect center of gravity, excellent aerodynamic properties, superior roll, and even when severely distorted will return to round.

As contrasted to the conventionally covered wound balls, the balls in accordance with the invention may be marked by simply stamping with a flat die, and are heat-resistant, solvent-resistant, abrasive-resistant, have an excellent shelf-life, will not water-log and have a superior texture and appearance. If the paint on the ball becomes worn or damaged, the balls may be very readily remarketed by removing the old paint cover with grit-blasting, such as sand-blasting and repainting the same. In contrast, thereto, the conventional balls seldom last long enough to allow repainting.

The golf balls in accordance with the invention may be manufactured much more easily and cheaply than the conventional wound balls, and the required raw materials are readily available and relatively inexpensive. Critical for the desired characteristics is the combination of the elastomer with the relatively large amount of the cross-linking monomer which polymerizes as it cross-links the elastomer forming the three-dimensional network with the long, flexible cross-links. The combination of the elastomer with this monomer produces a synergistic effect since it yields a tough impact-resistant material having the desired and necessary characteristics, whereas either of these materials cured alone will not produce such a material.

The long, flexible cross-links thus formed will generally have a length of at least 10 carbon atoms and normally the length of the cross-links will be substantially in excess of this. Thus, for example, when using butylene-1,3-dimethacrylate a cross-link of 11 atoms minimum length is obtained, and the average length is a multiple of this value.

The following examples are given by way of illustration and not limitation:

**Example 1**

A composition of 100 parts of predominantly cis butadiene polymer (62.5 parts of butylene glycol dimethacrylate, 62.5 parts of fine silica filler sold under the trade name of "Hyasil," and 3.13 parts of dicumyl peroxide, are thoroughly mixed on rubber rolls at approximately room temperature until the resulting material is completely homogeneous. The material is made up into cylindrical slugs and placed in molds. Molding is effected at pressures from 125 to 300 p.s.i. at a mold temperature of 300° F. with a molding time of about 10 minutes.

The balls are removed from the mold, the thin finish from the mold is buffed off, and the ball is then painted and marked.

The balls have the following characteristics:

<table>
<thead>
<tr>
<th>Size</th>
<th>inches in diameter</th>
<th>1.680</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roundness</td>
<td>inches</td>
<td>.002</td>
</tr>
<tr>
<td>Hardness</td>
<td>96b</td>
<td></td>
</tr>
<tr>
<td>Compression</td>
<td>125</td>
<td></td>
</tr>
<tr>
<td>Initial velocity</td>
<td>238±2</td>
<td></td>
</tr>
</tbody>
</table>

Their play characteristics may be considered at least equivalent to the conventional premium-quality wound balls, and they have excellent click and other playing characteristics. The balls are practically cut-free and indestructible and their resistance to extreme stresses is shown by clamping the ball and a premium-wound ball in a vise and applying pressure until the wound ball explodes and removing the molded golf ball. It is slightly out of shape but in a few moments resumes its normal round shape.

**Example 2**

A composition of 100 parts of cis butadiene, 62.5 parts of divinyl benzene, 62.5 parts of fine silica of a particle size between .01 and 0.1 micron and 3.13 parts of dicumyl peroxide, are thoroughly mixed on rubber rolls at approximately room temperature until the composition is completely homogeneous. The material is made up into cylindrical slugs and placed in molds. Molding is effected at pressures from 125 to 130 p.s.i. at a mold temperature of 300° F. with a molding time of about 10 minutes. The balls are removed from the mold, the thin finish is buffed off and the balls are then painted white. The golf balls produced were of excellent quality, having the following characteristics:

<table>
<thead>
<tr>
<th>Size</th>
<th>inches in diameter</th>
<th>1.681</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
<td>oz</td>
<td>1.610</td>
</tr>
<tr>
<td>Roundness</td>
<td>inches</td>
<td>.002</td>
</tr>
<tr>
<td>Density</td>
<td>1.12</td>
<td></td>
</tr>
<tr>
<td>Hardness</td>
<td>94b</td>
<td></td>
</tr>
<tr>
<td>Compression</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td>Initial velocity</td>
<td>238±2</td>
<td></td>
</tr>
</tbody>
</table>

**Example 3**

A composition of 100 parts by weight of cis polybutadiene, 56.2 parts by weight of butylene glycol dimethacrylate, 37.5 parts by weight of fine silica sold under the trade name "Hyasil," 6.2 parts by weight of cork having a particle size below 60 mesh, and 3.13 parts by weight of dicumyl peroxide, are thoroughly mixed on rubber rolls at approximately room temperature until the resulting mixture is homogeneous. The material is made up into cylindrical slugs and molded, in the identical manner described in Example 1, into golf balls. The balls produced have the following characteristics:

<table>
<thead>
<tr>
<th>Size</th>
<th>inches in diameter</th>
<th>1.680</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weight</td>
<td>oz</td>
<td>1.58</td>
</tr>
<tr>
<td>Compression</td>
<td>85</td>
<td></td>
</tr>
<tr>
<td>Hardness</td>
<td>93b</td>
<td></td>
</tr>
<tr>
<td>Initial velocity</td>
<td>238±2</td>
<td></td>
</tr>
</tbody>
</table>

If the balls are made up in the identical manner except,
5. A golf ball of spherical shape having a maximum diameter of about 1.69", of high durability and good
click predominately composed of a filled diene-rubber
cross-linked with about 25–90% by weight of a cross-
linking monomer for said rubber into a three dimension-
al network with long, flexible cross-links having a length
of at least 10 carbon atoms, said diene-rubber being one
of capable being so cross-linked by said cross-linking
monomer, said filler being present in amount ranging
between about 20 and 90% by weight of said rubber.

6. A golf ball according to claim 5 in which the diene-
rubber is cross-linked with a cross-linking monomer se-
lected from the group consisting of dimethacrylate esters
and divinyl compounds.

7. A golf ball according to claim 6 containing a filler
which is a member selected from the group consisting
of silica and silicates.

8. A homogeneous molded golf ball of high durability
and good click having a standard spherical shape of a
maximum diameter of about 1.69" predominately com-
posed of a diene elastomer highly cross-linked with about
25–90% by weight of a dimethacrylate ester cross-link-
ing monomer into a three dimensional network with long,
flexible cross-links and filled with about 25–90% by
weight of a finely divided filler, the principal constituent
of which is finely divided silica, said elastomer being one
capable of being so cross-linked by said cross-linking
monomer.

9. A golf ball according to claim 8 in which said
dimethacrylate ester is a glycol dimethacrylate.

10. A golf ball according to claim 9 in which said
glycol dimethacrylate is butylene glycol dimethacrylate.

11. A homogeneous molded golf ball of high durability
and good click having a standard spherical shape of max-
nimum diameter of about 1.69" predominately com-
posed of a cis butadiene elastomer highly cross-linked into a
three dimensional network with long, flexible cross links
by 25–90% by weight of a member selected from the
group consisting of butylene glycol dimethacrylate and
divinyl benzene and containing about 50–70 parts by
weight of finely divided silica filler.

12. A homogeneous molded golf ball of high durability
and good click acceptable for official play under the
U.S.G.A. rules and having the following specifications:

<table>
<thead>
<tr>
<th>Weight</th>
<th>1.600–1.620 oz</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity</td>
<td>1.11–1.12</td>
</tr>
<tr>
<td>Hardness (Shore)</td>
<td>90–100b</td>
</tr>
<tr>
<td>Compression range</td>
<td>40–120</td>
</tr>
<tr>
<td>Initial velocity (U.S.G.A.)</td>
<td>320–250 ft/sec</td>
</tr>
</tbody>
</table>

and predominately composed of cis butadiene rubber high-
ly cross-linked into a three dimensional network with long,
flexible cross links by about 56% by weight of butylene
glycol dimethacrylate and filled with 37% by weight of
finely divided silica and about 6 to 12% of cork.

13. A homogeneous molded golf ball for driving range
use comprising a molded sphere of the size and shape of
a golf ball predominately composed of a filled elastomer
highly cross-linked into a three dimensional network with
long, flexible cross links formed from about 25 to 90%
by weight based on said elastomer of a polymerized cross-
linking monomer, said filler being present in amount rang-
ing between about 20–90% by weight of said elastomer,
said elastomer being one capable of being so cross-linked
by said cross-linking monomer and containing sufficient
white pigment so that the ball is uniformly white through-
out.

14. A golf ball for driving range use according to
claim 13 in which said elastomer is a diene elastomer
cross-linked into said three dimensional network with
about 40–90% by weight of a cross-linking agent selected
from the group consisting of dimethacrylate esters, and
divinyl compound.

15. A golf ball for driving range use according to
claim 13 in which said pigment is titanium dioxide present
in amount of 2 to 40% by weight.

16. A highly durable homogeneous molded golf ball
for driving range use comprising a molded sphere of the
size and shape of a golf ball predominately composed of a
silica filled butadiene elastomer highly cross-linked into
a three dimensional network with long, flexible cross links
by about 25–90% by weight of a cross-linking monomer
selected from the group consisting of dimethacrylate
esters and divinyl compounds, and containing from 5 to
10% by weight of titanium dioxide to give the same a
uniform white color throughout, said elastomer contain-
ing a total amount of filler and titanium dioxide of about
20–90% by weight.

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DELBERT B. LOWE, Examiner.

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