Abstract: Method of making an abrasive article (e.g., an abrasive wheel or a honing stick), a cutting tool or a cutting tool insert comprising glass-ceramic abrasive particles in a ceramic matrix.

(54) Title: METHOD OF MAKING ABRASIVE ARTICLES, CUTTING TOOLS, AND CUTTING TOOL INSERTS
METHOD OF MAKING ABRASIVE ARTICLES, CUTTING TOOLS, AND CUTTING TOOL INSERTS

Background

Ceramic composites are known in the art. Uses of known ceramic composites include abrasive articles (e.g., abrasive wheels and honing sticks), cutting tools, and cutting tool inserts.

The ceramic composite, abrasive, and cutting tool industries continue to desire new materials, abrasive articles, cutting tools and cutting tool inserts, as well as methods for making the same.

Summary

In one aspect, the present invention provides a method of making an abrasive article (e.g., an abrasive wheel or a honing stick) comprising glass-ceramic abrasive particles in a ceramic matrix each comprising a mixture of metal oxides, wherein the glass-ceramic abrasive particles (i.e., glass-ceramic particles having an average hardness of at least 15 GPa (in some embodiments, at least 16 GPa, at least 17 GPa, or even at least 18 GPa)) and the matrix each comprise at least 35 (in some embodiments, at least 40, 45, 50, 55, 60, 65, or even at least 70) percent by weight Al₂O₃, based on the total weight of the glass-ceramic abrasive particle and the matrix, respectively, the method comprising:

providing a mixture of at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 (in some embodiments, either greater than 1.5:1 or less than 1:1.5, or even either greater than 2:1 or less than 1:2);

heating the mixture so that the glass of the ceramic particles coalesces to provide a body; and

heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic (i.e., at least a portion of the glass crystallizes),

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to provide the abrasive article..

In one aspect, the present invention provides a method of making a cutting tool or cutting tool insert comprising glass-ceramic abrasive particles in a ceramic matrix each comprising a mixture of metal oxides, wherein the glass-ceramic abrasive particles and the matrix each comprise at least 35 (in some embodiments, at least 40, 45, 50, 55, 60, 65, or even at least 70) percent by weight Al$_2$O$_3$, based on the total weight of the glass-ceramic abrasive particle and the matrix, the method comprising:

- providing a mixture comprising at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 (in some embodiments, either greater than 1.5:1 or less than 1:1.5, or even either greater than 2:1 or less than 1:2);

- heating the mixture so that the glass of the ceramic particles coalesces to provide a body; and

- heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic (i.e., at least a portion of the glass crystallizes), to provide the cutting tool or cutting tool insert.

In some embodiments, the glass-ceramic abrasive particles have a first composition and a plurality of ceramic particles comprising glass, the ceramic particles having a second composition, wherein the first and second compositions each comprise a mixture of metal oxides, wherein the amount of any particular metal oxide by weight in the first composition is at least 90 (in some embodiments, at least 95, 97, 98, or even at least 99) percent of its weight percent in the second composition, and the combined amount of metal oxides that are different between the first and second compositions do not exceed 10% (in some embodiments, do not exceed 10%, 5%, 3%, 2%, or even do not exceed 1%) of the total weight of all first and second compositions metal oxides, wherein the first and second compositions each comprise at least 35 (in some embodiments, 40, 45, 50, 55, 60, 65, or even at least 70) percent by weight Al$_2$O$_3$, based on the total weight of the first and second compositions, respectively. Typically, the glass-ceramic abrasive particles have a
density at least 90 (in some embodiments, at least 93, 95, 96, 97, 98, or even at least 99) percent of theoretical density.

In some embodiments, the glass-ceramic abrasive particles and ceramic particles each comprise at least 35, 40, 45, 50, 55, 60, 65, or even at least 70 percent by weight \( \text{Al}_2\text{O}_3 \), based on the total weight of the glass-ceramic abrasive particles and the ceramic particles, respectively. In some embodiments, the glass-ceramic abrasive particles and the ceramic particles each contain not more than 10 (in some embodiments, not more than 9, 8, 7, 6, 5, 4, 3, 2, 1, or even zero) percent by weight collectively \( \text{AS}_2\text{O}_3, \text{Bi}_2\text{O}_3, \text{B}_2\text{O}_3, \text{GeO}_2, \text{P}_2\text{O}_5, \text{SiO}_2, \text{TeO}_2, \text{V}_2\text{O}_5 \), alkaline oxides, and alkaline earth oxides, based on the total weight of the glass-ceramic abrasive particles and the ceramic particles, respectively.

In some embodiments, the glass-ceramic abrasive particles provide in a range from 10 to 70 (in some embodiments, in a range from 20 to 50, 20 to 40, or even from 25 to 35) percent by volume of the abrasive article, cutting tool, or cutting tool insert, and the ceramic particles provide a range from 30 to 90 (in some embodiments, in a range from 50 to 80, 60 to 80, or even from 65 to 75) percent by volume of the abrasive article or cutting tool, based on the total volume of the abrasive article, cutting tool, or cutting tool insert, respectively.

In this application:

"amorphous material" refers to material derived from a melt and/or a vapor phase that lacks any long range crystal structure as determined by X-ray diffraction and/or has an exothermic peak corresponding to the crystallization of the amorphous material as determined by a DTA (differential thermal analysis) as determined by the test described herein entitled "Differential Thermal Analysis";

"cell" refers to the smallest structural unit of a ceramic material that forms at least two distinct crystalline phases if the ceramic material is crystallized. In an amorphous material, a cell is essentially uniform in composition and essentially free of microstructural features (e.g., grain boundaries). In the context of the present invention, an individual cell generally corresponds to the individual body (e.g., particle, glass body, etc.) that is coalesced with a plurality of other bodies to form a larger consolidated ceramic material.

The size of a cell can be determined using metallography and microscopic methods known in the art (e.g., SEM, TEM).
"ceramic" includes glass, crystalline ceramic, glass-ceramic, and combinations thereof;

"complex metal oxide" refers to a metal oxide comprising two or more different metal elements and oxygen (e.g., CeAl₅O₁₈, Dy₃Al₅O₁₂, MgAl₂O₄, and Y₃Al₅O₁₂);

"complex Al₂O₃-metal oxide" refers to a complex metal oxide comprising, on a theoretical oxide basis, Al₂O₃ and one or more metal elements other than Al (e.g., CeAl₅O₁₈, Dy₃Al₅O₁₂, MgAl₂O₄, and Y₃Al₅O₁₂);

"complex Al₂O₃-Y₂O₃" refers to a complex metal oxide comprising, on a theoretical oxide basis, Al₂O₃ and Y₂O₃ (e.g., Y₃Al₅O₁₂);

"complex Al₂O₃-REO" refers to a complex metal oxide comprising, on a theoretical oxide basis, Al₂O₃ and rare earth oxide (e.g., CeAl₅O₁₈ and Dy₃Al₅O₁₂);

"glass" refers to amorphous material exhibiting a glass transition temperature;

"glass-ceramic" refers to ceramic comprising crystals formed by heat-treating glass;

"Tg" refers to the glass transition temperature as determined by the test described herein entitled "Differential Thermal Analysis";

"Tₓ" refers to the crystallization temperature as determined by the test described herein entitled "Differential Thermal Analysis";

"rare earth oxides" refers to cerium oxide (e.g., CeO₂), dysprosium oxide (e.g., Dy₂O₃), erbium oxide (e.g., Er₂O₃), europium oxide (e.g., Eu₂O₃), gadolinium oxide (e.g., Gd₂O₃), holmium oxide (e.g., Ho₂O₃), lanthanum oxide (e.g., La₂O₃), lutetium oxide (e.g., Lu₂O₃), neodymium oxide (e.g., Nd₂O₃), praseodymium oxide (e.g., Pr₆O₁₇), samarium oxide (e.g., Sm₂O₃), terbium oxide (e.g., Tb₂O₃), thorium oxide (e.g., Th₄O₇), thulium oxide (e.g., Tm₂O₃) and ytterbium oxide (e.g., Yb₂O₃), and combinations thereof; and

"REO" refers to rare earth oxide(s).

Further, it is understood herein that unless it is stated that a metal oxide (e.g., Al₂O₃, complex Al₂O₃-metal oxide, etc.) is crystalline, for example, in a glass-ceramic, it may be glass, crystalline, or portions glass and portions crystalline. For example, if a glass-ceramic comprises Al₂O₃ and ZrO₂, the Al₂O₃ and ZrO₂ may each be in a glassy state, crystalline state, or portions in a glassy state and portions in a crystalline state, or even as a reaction product with another metal oxide(s) (e.g., unless it is stated that, for
example, Al₂O₃ is present as crystalline AI2O₃ or a specific crystalline phase of AI2O3 (e.g., alpha Al₂Os), it may be present as crystalline AI2O₃ and/or as part of one or more crystalline complex AI₂O₃-metal oxides).

In some embodiments, the glass-ceramic abrasive particles and ceramic particles each independently comprise at least 35 (in some embodiments, at least 40, 45, 50, 55, 60, 65, 70, or even at least 75) percent by weight AI2O3, based on the total weight of the glass-ceramic abrasive particles and ceramic particles, respectively, wherein the glass-ceramic abrasive particles and ceramic particles each independently contain not more than 10 (in some embodiments, not more than 5, 4, 3, 2, 1, 0.5, 0.1, or even zero) percent by weight collectively As₂O₃, Bi₂O₃, B₂O₃, GeO₂, P₂O₅, SiO₂, TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides, based on the total weight of the glass-ceramic abrasive particles and ceramic particles, respectively. In some embodiments, the glass-ceramic abrasive particles and ceramic particles, respectively, further comprise at least one metal oxide other than Al₂O₃ (e.g., a metal oxide selected from the group consisting of BaO, CaO, CeO₂, CuO, Dy₂O₃, Er₂O₃, Eu₂O₃, Gd₂O₃, Ho₂O₃, La₂O₃, Lu₂O₃, MgO, Nd₂O₃, Pr₆O₁₇, Sm₂O₃, Sc₂O₃, SrO, Tb₂O₃, Th₄O₁₇, TiO₂, Tm₂O₃, Yb₂O₃, Y₂O₃, ZrO₂, and combinations thereof).

In some embodiments, the first and second compositions each independently comprise AI₂O₃, at least one of REO or Y₂O₃, and at least one of ZrO₂ or HfO₂, wherein at least 80 (in some embodiments, at least 85, 90, 95, 97, 98, 99, or even 100) percent by weight of the glass-ceramic abrasive particles and ceramic particles each independently collectively comprise the AI₂O₃, the at least one of REO or Y₂O₃, and the at least one of ZrO₂ or HfO₂, based on the total weight of the glass-ceramic abrasive particles and ceramic particles, respectively. In some embodiments, the glass-ceramic abrasive particles and ceramic particles each independently comprise at least 35, 40, 45, 50, 55, 60, 65, 70, or even at least 75 percent by weight AI₂O₃, based on the total weight of the glass-ceramic abrasive particles and ceramic particles, respectively. In some embodiments, the glass-ceramic abrasive particles and ceramic particles each independently further comprise at least one metal oxide other than AI₂O₃, REO, Y₂O₃, HfO₂, and ZrO₂ (e.g., a metal oxide selected from the group consisting of BaO, CaO, MgO, SrO, TiO₂, and combinations thereof).
Typically, the glass-ceramic abrasive particles are comprised of at least 50 (in some embodiments, 55, 60, 65, 70, 75, 80, 85, 90, 95, or even 99) percent by volume crystalline ceramic, based on the total volume of the glass-ceramic abrasive particles. In some embodiments, the ceramic particles comprises at least 1, 2, 3, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 97, 98, 99, or even 100 percent by volume crystalline ceramic, based on the total volume of the ceramic particles.

**Brief Description of the Drawings**

FIG. 1 is a side view of an exemplary embodiment of an apparatus including a powder feeder assembly for a flame-melting apparatus.

FIG. 2 is a section view of the apparatus of FIG. 1.

FIG. 3 is an exploded section view of the apparatus of FIG. 1.

FIG. 4 is a side view of a portion of the powder feeder assembly of FIG. 1.

FIG. 5 is a perspective view of a portion of the powder feeder assembly of FIG. 1.

FIG. 6 is a cross-sectional view of a portion of the powder feeder assembly of FIG. 1.

FIG. 7 is a perspective view of an exemplary abrasive wheel.

FIG. 8 is a perspective view of an exemplary honing stick.

FIG. 9 is a perspective view of an exemplary cutting tool.

FIGS. 10-14 are SEM micrographs of Examples 1-5, respectively.

**Description**

The present invention pertains to methods of making abrasive articles (e.g., wheels and honing sticks), cutting tools and cutting tools inserts. The glass-ceramic abrasive particles and the ceramic particles, as applicable, are prepared by selecting the necessary raw materials and processing techniques.

Typically, the glass-ceramic abrasive particles and the ceramic particles comprise Al₂O₃ and at least one BaO, CaO₅Cr₂O₃, CoOₓCuO, Fe₂O₃, GeO₂, HfO₂, Li₂O, MgO, MnO, NiO, Na₂O, REO, Sc₂O₃, SrO, TiO₂, Y₂O₃, ZnO, or ZrO₂. The glass-ceramic abrasive particles and the ceramic particles typically each comprise Al₂O₃.
Sources, including commercial sources, for the metal oxides include the oxides themselves, metal powders, complex oxides, ores, carbonates, acetates, nitrates, chlorides, hydroxides, etc.

For example, sources, including commercial sources, of (on a theoretical oxide basis) Al₂O₃ include bauxite (including both natural occurring bauxite and synthetically produced bauxite), calcined bauxite, hydrated aluminas (e.g., boehmite, and gibbsite), aluminum, Bayer process alumina, aluminum ore, gamma alumina, alpha alumina, aluminum salts, aluminum nitrates, and combinations thereof. The Al₂O₃ source may contain, or only provide, Al₂O₃. The Al₂O₃ source may contain, or provide Al₂O₃, as well as one or more metal oxides other than Al₂O₃ (including materials of or containing complex Al₂O₃-metal oxides (e.g., DyaAlsO₁₂, Y₃Al₅O₁₂, CeAl₁O₁₈, etc.)).

Sources, including commercial sources, of rare earth oxides include rare earth oxide powders, rare earth metals, rare earth-containing ores (e.g., bastnasite and monazite), rare earth salts, rare earth nitrates, and rare earth carbonates. The rare earth oxide(s) source may contain, or only provide, rare earth oxide(s). The rare earth oxide(s) source may contain, or provide rare earth oxide(s), as well as one or more metal oxides other than rare earth oxide(s) (including materials of or containing complex rare earth oxide-other metal oxides (e.g., Dy₃Al₅O₁₂, CeAl₁O₁₈, etc.)).

Sources, including commercial sources, of (on a theoretical oxide basis) Y₂O₃ include yttrium oxide powders, yttrium, yttrium-containing ores, and yttrium salts (e.g., yttrium carbonates, nitrates, chlorides, hydroxides, and combinations thereof). The Y₂O₃ source may contain, or only provide, Y₂O₃. The Y₂O₃ source may contain, or provide Y₂O₃, as well as one or more metal oxides other than Y₂O₃ (including materials of or containing complex Y₂O₃-metal oxides (e.g., Y₃Al₁O₁₈)).

For embodiments comprising ZrO₂ and HfO₂, the weight ratio of ZrO₂:HfO₂ may be in a range of 1:zero (i.e., all ZrO₂; no HfO₂) to zero:1, as well as, for example, at least about 99, 98, 97, 96, 95, 90, 85, 80, 75, 70, 65, 60, 55, 50, 45, 40, 35, 30, 25, 20, 15, 10, and 5 parts (by weight) ZrO₂ and a corresponding amount OfHfO₂ (e.g., at least about 99 parts (by weight) ZrO₂ and not greater than about 1 part HfO₂) and at least about 99, 98, 97, 96, 95, 90, 85, 80, 75, 70, 65, 60, 55, 50, 45, 40, 35, 30, 25, 20, 15, 10, and 5 parts HfO₂ and a corresponding amount OfZrO₂.
Sources, including commercial sources, of (on a theoretical oxide basis) ZrO₂ include zirconium oxide powders, zircon sand, zirconium, zirconium-containing ores, and zirconium salts (e.g., zirconium carbonates, acetates, nitrates, chlorides, hydroxides, and combinations thereof). In addition, or alternatively, the ZrO₂ source may contain, or provide ZrO₂, as well as other metal oxides such as hafnia. Sources, including commercial sources, of (on a theoretical oxide basis) HfO₂ include hafnium oxide powders, hafnium, hafnium-containing ores, and hafnium salts. In addition, or alternatively, the HfO₂ source may contain, or provide HfO₂, as well as other metal oxides such as ZrO₂.

In some embodiments, it may be advantageous for at least a portion of a metal oxide source (in some embodiments, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, or even 100 percent by weight) to be obtained by adding particulate, metallic material comprising at least one of a metal (e.g., Al, Ca, Cu, Cr, Fe, Li, Mg, Ni, Ag, Ti, Zr, and combinations thereof), M, that has a negative enthalpy of oxide formation or an alloy thereof to the melt, or otherwise combining them with the other raw materials. Although not wanting to be bound by theory, it is believed that the heat resulting from the exothermic reaction associated with the oxidation of the metal is beneficial in the formation of a homogeneous melt and resulting glass. For example, it is believed that the additional heat generated by the oxidation reaction within the raw material eliminates, minimizes, or at least reduces insufficient heat transfer, and hence facilitates formation and homogeneity of the melt. It is also believed that the availability of the additional heat aids in driving various chemical reactions and physical processes (e.g., densification, and spherodization) to completion. Further, it is believed for some embodiments, the presence of the additional heat generated by the oxidation reaction actually enables the formation of a melt, which otherwise is difficult or not practical due to high melting point of the materials. Further, the presence of the additional heat generated by the oxidation reaction actually enables the formation of glass that otherwise could not be made, or could not be made in the desired size range. Another advantage of the invention includes, in forming the glasses, that many of the chemical and physical processes such as melting, densification and spherodizing can be achieved in a short time, so that very high quench rates may be achieved. For additional details, see U.S. Publication No. US2003-0110709A1, published June 16, 2003.
In some embodiments, for example, the raw materials are fed independently to form the molten mixture. In some embodiments, for example, certain raw materials are mixed together, while other raw materials are added independently into the molten mixture. In some embodiments, for example, the raw materials are combined or mixed together prior to melting. The raw materials may be combined, for example, in any suitable and known manner to form a substantially homogeneous mixture. These combining techniques include ball milling, mixing, tumbling, and the like. The milling media in the ball mill may be metal balls, ceramic balls, and the like. The ceramic milling media may be, for example, alumina, zirconia, silica, magnesia and the like. The ball milling may occur dry, in an aqueous environment, or in a solvent-based (e.g., isopropyl alcohol) environment. If the raw material batch contains metal powders, then it is generally desired to use a solvent during milling. This solvent may be any suitable material with the appropriate flash point and ability to disperse the raw materials. The milling time may be from a few minutes to a few days, generally between a few hours to 24 hours. In a wet or solvent based milling system, the liquid medium is removed, typically by drying, so that the resulting mixture is typically homogeneous and substantially devoid of the water and/or solvent. If a solvent based milling system is used, during drying, a solvent recovery system may be employed to recycle the solvent. After drying, the resulting mixture may be in the form of a "dried cake". This cake-like mixture may then be broken up or crushed into the desired particle size prior to melting. Alternatively, for example, spray-drying techniques may be used. The latter typically provides spherical particulates of a desired mixture. The precursor material may also be prepared by wet chemical methods including precipitation and sol-gel. Such methods will be beneficial if extremely high levels of homogeneity are desired.

Particulate raw materials are typically selected to have particle sizes such that the formation of a homogeneous melt can be achieved rapidly. Typically, raw materials with relatively small average particle sizes and narrow distributions are used for this purpose. In some methods (e.g., flame forming and plasma spraying), particularly desirable particulate raw materials are those having an average particle size in a range from about 5 nm to about 50 micrometers (in some embodiments, in a range from about 10 nm to about 20 micrometers, or even about 15 nm to about 1 micrometer), wherein at least 90 (in some
embodiments, 95, or even 100) percent by weight of the particulate is the raw material, although sizes outside of the sizes and ranges may also be useful. Particulate less than about 5 nm in size tends to be difficult to handle (e.g., the flow properties of the feed particles tended to be undesirable as they tend to have poor flow properties). Use of particulate larger than about 50 micrometers in typical flame forming or plasma spraying processes tend to make it more difficult to obtain homogenous melts and glasses and/or the desired composition.

Furthermore, in some cases, for example, when particulate material is fed in to a flame or thermal or plasma spray apparatus, to form the melt, it may be desirable for the particulate raw materials to be provided in a range of particle sizes, including agglomerated forms.

Glasses for the ceramic particles, and/or for making the glass-ceramic abrasive particles and the ceramic particles, can be made, for example, by heating (including in a flame or plasma) the appropriate metal oxide sources to form a melt, desirably a homogenous melt, and then rapidly cooling the melt to provide glass. Some embodiments of glasses can be made, for example, by melting the metal oxide sources in any suitable furnace (e.g., an inductively or resistively heated furnace, a gas-fired furnace, or an electric arc furnace).

The glass is typically obtained by relatively rapidly cooling the molten material (i.e., the melt). The quench rate (i.e., the cooling time) to obtain the glass depends upon many factors, including the chemical composition of the melt, the glass-forming ability of the components, the thermal properties of the melt and the resulting glass, the processing technique(s), the dimensions and mass of the resulting glass, and the cooling technique. In general, relatively higher quench rates are required to form glasses comprising higher amounts of Al₂O₃, especially in the absence of known glass formers such as As₂O₃, B₂O₃, B₂O₅. GeO₂, P₂O₅, SiO₂, TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides. Similarly, it is more difficult to cool melts into glasses in larger dimensions, as it is more difficult to remove heat fast enough.

In some embodiments of the invention, the raw materials are heated into a molten state in a particulate form and subsequently cooled into glass particles. Typically, the
particles have a particle size greater than 25 micrometers (in some embodiments, greater
than 50, 100, 150, or even 200 micrometers).

The quench rates achieved in making glasses according to the methods of the
present invention are believed to be higher than \(10^2, 10^3, 10^4, 10^5\) or even \(10^{60}\)C/sec (i.e., a
temperature drop of \(1000^\circ\)C from a molten state in less than 10 seconds, less than a
second, less than a tenth of a second, less than a hundredth of a second or even less than a
thousandth of a second, respectively). Techniques for cooling the melt include discharging
the melt into a cooling media (e.g., high velocity air jets, liquids (e.g., cold water), metal
plates (including chilled metal plates), metal rolls (including chilled metal rolls), metal
balls (including chilled metal balls), and the like). Other cooling techniques known in the
art include roll-chilling. Roll-chilling can be carried out, for example, by melting the
metal oxide sources at a temperature typically 20-200\(^{\circ}\)C higher than the melting point, and
cooling/quenching the melt by spraying it under high pressure (e.g., using a gas such as air,
argon, nitrogen or the like) onto a high-speed rotary roll(s). Typically, the rolls are made
of metal and are water-cooled. Metal book molds may also be useful for
cooling/quenching the melt.

The cooling rate is believed to affect the properties of the quenched glass. For
instance, glass transition temperature, density and other properties of glass typically
change with cooling rates.

Rapid cooling may also be conducted under controlled atmospheres, such as a
reducing, neutral, or oxidizing environment to maintain and/or influence the desired
oxidation states, etc. during cooling. The atmosphere can also influence glass formation
by influencing crystallization kinetics from undercooled liquid. For example, larger
undercooling of Al\(\text{2}O_3\) melts without crystallization has been reported in argon atmosphere
as compared to that in air.

In one method, glasses for the ceramic particles, and/or for making the glass-
ceramic abrasive particles and the ceramic particles, can be made utilizing flame fusion as
reported, for example, in U.S. Pat. No. 6,254,981 (Castle). In this method, the metal oxide
source(s) are fed (e.g., in the form of particles, sometimes referred to as "feed particles")
directly into a burner (e.g., a methane-air burner, an acetylene-oxygen burner, a hydrogen-
oxygen burner, and the like), and then quenched, for example, in water, cooling oil, air, or
the like. Feed particles can be formed, for example, by grinding, agglomerating (e.g., spray-drying), melting, or sintering the metal oxide sources. The size of feed particles fed into the flame generally determines the size of the resulting particles comprising glass.

Some embodiments of glasses can also be obtained by other techniques, such as: laser spining melt with free fall cooling, Taylor wire technique, plasmatron technique, hammer and anvil technique, centrifugal quenching, air gun splat cooling, single roller and twin roller quenching, roller-plate quenching, and pendant drop melt extraction (see, e.g., Rapid Solidification of Ceramics, Brockway et al., Metals And Ceramics Information Center, A Department of Defense Information Analysis Center, Columbus, OH, January, 1984). Some embodiments of glasses may also be obtained by other techniques, such as: thermal (including flame or laser or plasma-assisted) pyrolysis of suitable precursors, physical vapor synthesis (PVS) of metal precursors and mechanochemical processing. Other techniques for forming melts, cooling/quenching melts, and/or otherwise forming glass include melt-extraction, and gas or centrifugal atomization.

Gas atomization involves heating feed particles to convert them to melt. A thin stream of such melt is atomized through contact with a disruptive air jet (i.e., the stream is divided into fine droplets). The resulting substantially discrete, generally ellipsoidal glass particles (e.g., beads) are then recovered. Examples of bead sizes include those having a diameter in a range of about 5 micrometers to about 3 mm. Melt-extraction can be carried out, for example, as reported in U.S. Pat. 5,605,870 (Strom-Olsen et al.). Container-less glass forming techniques utilizing laser beam heating as reported, for example, in U.S. Pat. No. 6,482,758 (Weber), may also be useful in making glasses useful in the present invention.

An exemplary powder feeder apparatus is illustrated in FIGS. 1-6. The powder feeder assembly 1000 holds and delivers powder 1110 to a flame-melting device 1500. Flame-melting device 1500 includes powder receiving section 1510 for receiving powder 1110 for melting and transforming into another material(s), such as those disclosed herein. Powder 1110 is delivered into powder receiving section 1510 through discharge opening 1130 of powder feeder assembly 1000. Connecting tube 1900 is positioned between discharge opening 1130 and powder receiving section 1510. Also, funnel 1300 is
positioned proximate to discharge opening 1130 for receiving and directing powder 1110 flow after it leaves discharge opening 1130.

Powder feeder assembly 1000 includes hopper 1100 for holding powder 1110. Typically, hopper 1100 includes body 1120 defined by a cylindrical wall, though other body shapes are possible. Also, hopper 1100 can be made from a unitary piece or multiple pieces. Hopper 1100 in the example embodiment illustrated also includes cover section 1200. Cover section 1200 includes opening 1710 for feeding powder 1110 into hopper 1100. Any commercially available delivery means can be used for filling hopper 1100 with powder 1110, such as a screw feeder, vibratory feeder, or brush feeder. Cover section 1200 can also include a section having shaft receiving opening 1422 (as illustrated in FIG. 6).

Brush assembly 1400 is disposed within hopper 1100 body 1120. Brush assembly 1400 is connected via shaft 1420 to means for rotating brush assembly 1400, such as motor 1800. Motor 1800 can also be connected to means for adjusting the speed of motor 1800, such as motor speed controller 1850. The brush assembly can use, for example, a Nylon Strip Brush (e.g., 1 inch (2.5 cm) overall height, 5/16 inch (0.8 cm) bristle length and 0.02 inch (5 millimeter) diameter), part# 74715T61, available from McMaster-Carr, Chicago, IL). The brush assembly is coupled to a shaft, which in turn is coupled to and driven by a DC Gear Motor (130 Volt, Ratio 60:1, Torque 22 lb.in.) (available, for example, from Bodine Electric Company, Chicago, IL). The speed of the motor can be controlled using an adjustable motor control (e.g., a Type-FPM Adjustable Speed PM Motor Control, Model # 818, also available from Bodine.

Brush assembly 1400 includes bristle element 1410 having distal 1411 and proximate end 1412. When powder 1110 is placed into hopper 1100 for delivery to flame-melting device 1500, brush assembly 1400 is rotated within hopper 1100. When brush assembly 1400 is rotated, bristle element(s) 1410 urges powder 1110 in hopper 1100 through screening member 1600. By adjusting the rotational speed of brush assembly 1400, the feed rate of powder 1110 through screening member 1600 can be controlled.

Brush assembly 1400 cooperates with screening member 1600 to deliver powder 1110 having desired properties from discharge opening 1130 to powder receiving section 1510 of flame-melting device 1500. Distal end 1411 of bristle 1410 is located in close
proximity to screening member 1600. While a small gap between distal end 1411 of bristles 1410 and screening member 1600 can be used, it is typical to keep the gap on the same order of magnitude as the particle size of the powder, however, one of ordinary skill in the art will appreciate that the gap can be much larger, depending on the particular properties of the powder being handled. Also, distal end 1411 of bristle 1410 can be positioned flush with screening member 1600 or positioned to protrude into and extend through mesh openings 1610 in screening member 1600. For bristles 1410 to protrude through openings 1610, at least some of bristles 1410 need to have a diameter smaller than the mesh size. Bristle elements 1410 can include a combination of bristles with different diameters and lengths, and any particular combination will depend on the operating conditions desired.

Extending bristle 1400 end 1411 into and through openings 1610 allows bristles 1410 to break up any particles forming bridges across openings 1610. Also bristles 1410 will tend to break-up other types of blockages that can occur typical to powder feeding. Bristle element 1410 can be a unitary piece, or can also be formed from a plurality of bristle segments. Also, if it is desired that the bristle elements extend into and/or through the mesh openings, then bristle 1410 size selected needs to be smaller than smallest mesh opening 1610.

Referring to FIG. 3, in the exemplary embodiment illustrated, hopper 1100 can include a wall defining a cylindrical body 1120. This shape conveniently provides for symmetry that allows for a more controlled flow rate of powder from discharge opening 1130. Also, the cylindrical shape is well suited for using with rotating brush assembly 1400, since bristle element 1410 can extend to the wall, leaving little or no area on the screening member that can accumulate powder. However, other geometries are possible, as the particular conditions of use dictate.

Hopper 1100 also includes cover section 1200. Cover section 1200 has opening 1710 for receiving powder 1110 from hopper feeder assembly 1700. Cover section 1200 cooperates with body 1120 to form powder chamber 1160. Opening 1710 on cover 1200 can also be omitted or sealable so that a gas, such as nitrogen, argon, or helium can be input into gas input line 1150 on hopper 1100 for neutralizing the atmosphere or assisting in delivering the powder or particles to the flame-melting device. Also, gas can be used in
the system for controlling the atmosphere surrounding the powder or particles. Also, gas input line 1910 can be placed after discharge opening 1130, for example, on connecting tube 1900.

Entire powder feeder assembly 1000 can be vibrated to further assist in powder transport. Optionally, the screening member can be vibrated to assist powder transport through powder feeder assembly 1000. One of ordinary skill in the art will recognize that other possible vibrating means can be used, and there are abundant commercial vibrating systems and devices that are available depending on the particular conditions of use.

Referring to FIGS. 1-6, when hopper 1100 includes cover 1200 and body 1120, removable cover 1200 allows easy access to powder chamber 1160 for cleaning or changing screening member 1600. Also, brush assembly 1400 can be positioned to form the desired engagement between bristle elements 1410 and screening member 1600. When brush assembly 1400 is attached to rotating shaft 1420, shaft 1420 can protrude outside opening 1422 in cover 1200 to be driven, for example, by motor 1800. The speed of brush assembly 1400 can be controlled by means such as speed controller 1850. Further details regarding this exemplary powder feeding apparatus can be found in see U.S. Publication No. US2005-0133974A1, published June 23, 2005.

The addition of certain metal oxides may alter the properties and/or crystalline structure or microstructure of ceramics utilized in the present invention, as well as the processing of the raw materials and intermediates in making the ceramic. For example, oxide additions such as MgO, CaO, Li2O, and Na2O have been observed to alter both the $T_g$ and $T_x$ (wherein $T_x$ is the crystallization temperature) of glass. Although not wishing to be bound by theory, it is believed that such additions influence glass formation. Further, for example, such oxide additions may decrease the melting temperature of the overall system (i.e., drive the system toward lower melting eutectic), and ease glass formation.

Compositions based upon complex eutectics in multi-component systems (quaternary, etc.) may have better glass-forming ability. The viscosity of the liquid melt and viscosity of the glass in its’ working range may also be affected by the addition of metal oxides other than the particular required oxide(s).

Crystallization of glasses and ceramics comprising the glass to form glass-ceramics may also be affected by the additions of materials. For example, certain metals, metal
oxides (e.g., titanates and zirconates), and fluorides may act as nucleation agents resulting in beneficial heterogeneous nucleation of crystals. Also, addition of some oxides may change the nature of metastable phases devitrifying from the glass upon reheating. In another aspect, for ceramics according to the present invention comprising crystalline ZrO₂, it may be desirable to add metal oxides (e.g., Y₂O₃, TiO₂, CeO₂, CaO, and MgO) that are known to stabilize the tetragonal/cubic form of ZrO₂.

The particular selection of metal oxide sources and other additives for making ceramics utilized in the present invention typically takes into account, for example, the desired composition, the microstructure, the degree of crystallinity, the physical properties (e.g., hardness or toughness), the presence of undesirable impurities, visual appearance (e.g., coloration), and the desired or required characteristics of the particular process (including equipment and any purification of the raw materials before and/or during fusion and/or solidification) being used to prepare the ceramics.

In some instances, it may be preferred to incorporate limited amounts of metal oxides selected from the group consisting of: B₂O₃, B₁₂O₃, Na₂O, P₂O₅, SiO₂, TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides, and combinations thereof. Sources, including commercial sources, include the oxides themselves, complex oxides, elemental (e.g., Si) powders, ores, carbonates, acetates, nitrates, chlorides, hydroxides, etc. These metal oxides may be added, for example, to modify a physical property of the resulting glass-ceramic and/or improve processing. These metal oxides, when used, are typically added from greater than 0 to 20% by weight collectively (in some embodiments, greater than 0 to 5% by weight collectively, or even greater than 0 to 2% by weight collectively) of the glass-ceramic depending, for example, upon the desired property.

Useful formulations include those at or near a eutectic composition(s) (e.g., ternary eutectic compositions). In addition to compositions disclosed herein, other such compositions, including quaternary and other higher order eutectic compositions, may be apparent to those skilled in the art after reviewing the present disclosure.

The microstructure or phase composition (glassy/crystalline) of a material can be determined in a number of ways, including optical microscopy, electron microscopy, differential thermal analysis (DTA), and x-ray diffraction (XRD).
Using optical microscopy, amorphous material is typically predominantly transparent due to the lack of light scattering centers such as crystal boundaries, while crystalline material shows a crystalline structure and is opaque due to light scattering effects. A percent amorphous (or glass) yield can be calculated for particles (e.g., beads), etc. using a -100+120 mesh size fraction (i.e., the fraction collected between 150-micrometer opening size and 125-micrometer opening size screens). The measurements are done in the following manner. A single layer of particles, beads, etc. is spread out upon a glass slide. The particles, beads, etc. are observed using an optical microscope. Using the crosshairs in the optical microscope eyepiece as a guide, particles, beads, etc. that lay along a straight line are counted either amorphous or crystalline depending on their optical clarity (i.e., amorphous if they were clear). A total of 500 particles, beads, etc. are typically counted, although fewer particles, beads, etc. may be used and a percent amorphous yield is determined by the amount of amorphous particles, beads, etc. divided by total particles, beads, etc. counted. Embodiments of methods for making the amorphous (or glass) may have yields of at least 50, 60, 70, 75, 80, 85, 90, 95, or even 100 percent.

If it is desired for all the particles to be amorphous (or glass), and the resulting yield is less than 100%, the amorphous (or glass) particles may be separated from the non-amorphous (or non-glass) particles. Such separation may be done, for example, by any conventional techniques, including separating based upon density or optical clarity.

Using DTA, the material is classified as amorphous if the corresponding DTA trace of the material contains an exothermic crystallization event ($T_x$). If the same trace also contains an endothermic event ($T_g$) at a temperature lower than $T_x$, it is considered to consist of a glass phase. If the DTA trace of the material contains no such events, it is considered to contain crystalline phases.

Differential thermal analysis (DTA) can be conducted using the following method. DTA runs can be made (using an instrument such as that obtained from Netzsch Instruments, Selb, Germany under the trade designation "NETZSCH STA 409 DTA/TGA") using a -140+170 mesh size fraction (i.e., the fraction collected between 105-micrometer opening size and 90-micrometer opening size screens). An amount of each
screened sample (typically about 400 milligrams (mg)) is placed in a 100-microliter Al₂O₃ sample holder. Each sample is heated in static air at a rate of 10°C/minute from room temperature (about 25°C) to 1100°C.

Using powder x-ray diffraction, XRD₃ (using an x-ray diffractometer such as that obtained under the trade designation "PHILLIPS XRG 3100" from Phillips, Mahwah, NJ, with copper Kα1 radiation of 1.54050 Angstrom) the phases present in a material can be determined by comparing the peaks present in the XRD trace of the crystallized material to XRD patterns of crystalline phases provided in JCPDS (Joint Committee on Powder Diffraction Standards) databases, published by International Center for Diffraction Data.

Furthermore, XRD can be used qualitatively to determine types of phases. The presence of a broad diffuse intensity peak is taken as an indication of the amorphous nature of a material. The existence of both a broad peak and well-defined peaks is taken as an indication of existence of crystalline matter within a glass matrix.

In some embodiments, the amorphous (glass) particle size is selected for the coalescing step. In some embodiments, the amorphous material has an average particle size of less than 3 micrometers (in some embodiments, less than 2, 1, 0.75, or even less than 0.5 micrometer), in order to produce glass-ceramic having an average cell size of less than 3 micrometers (in some embodiments, less than 2, 1, 0.75, or even less than 0.5 micrometer). Although not wanting to be bound by theory, it is believed that smaller cell sizes lead to improved mechanical properties for consolidated ceramics (e.g., higher strength and higher hardness).

The initially formed glass or ceramic (including glass prior to crystallization) may be larger in size than that desired. For example, it may be desirable for the ceramic particles comprising glass be smaller sized particles. If the glass is in a desired geometric shape and/or size, size reduction is typically not needed. The glass or ceramic can be converted into smaller pieces using crushing and/or comminuting techniques known in the art, including roll crushing, jaw crushing, hammer milling, ball milling, jet milling, impact crushing, and the like. In some instances, it is desired to have two or multiple crushing steps. For example, after the ceramic is formed (solidified or, in the case of the glass-ceramic, in some embodiments, after conversion to glass-ceramic), it may be in the form of larger than desired. The first crushing step may involve crushing these relatively large
masses or "chunks" to form smaller pieces. This crushing of these chunks may be accomplished, for example, with a hammer mill, impact crusher or jaw crusher. These smaller pieces may then be subsequently crushed to produce the desired particle size distribution. In order to produce the desired particle size distribution (sometimes referred to as grit size or grade), it may be necessary to perform multiple crushing steps. In general the crushing conditions are optimized to achieve the desired particle shape(s) and particle size distribution. Resulting particles that are not of the desired size may be re-crushed if they are too large, or "recycled" and used as a raw material for re-melting if they are too small.

In some embodiments, the initially formed amorphous material (including glass prior to crystallization) may be formed at the desired average particle size (e.g., less than 3 micrometers). Spray-pyrolysis, plasma processing, and condensation from vapors can be used to form smaller particle sizes.

The shape of particles can depend, for example, on the composition and/or microstructure of the ceramic, the geometry in which it was cooled, and the manner in which the ceramic is crushed (i.e., the crushing technique used). In general, where a "blocky" shape is preferred, more energy may be employed to achieve this shape. Conversely, where a "sharp" shape is preferred, less energy may be employed to achieve this shape. The crushing technique may also be changed to achieve different desired shapes. For some particles an average aspect ratio ranging from 1:1 to 5:1 is typically desired, and in some embodiments, 1.25:1 to 3:1, or even 1.5:1 to 2.5:1.

It is also within the scope of the present invention, for example, to directly form articles in desired shapes. For example, desired articles may be formed (including molded) by pouring or forming the melt into a mold. Also see, for example, the forming techniques described in application having U.S. Serial No. 10/358,772, filed February 5, 2003.

It may be desirable to grade or screen the ceramic particles to prove a desired size distribution. For example, the glass-ceramic abrasive particles are usually graded to a given particle size distribution before use. Such distributions typically have a range of particle sizes, from coarse particles to fine particles. In the abrasive art this range is sometimes referred to as a "coarse", "control" and "fine" fractions. Hence it may be desirable to grade glass-ceramic particles of the first composition according to an abrasive
industry accepted grading standard. Abrasive particles graded according to abrasive
industry accepted grading standards specify the particle size distribution for each nominal
grade within numerical limits. Such industry accepted grading standards (i.e., specified
nominal grades) include those known as the American National Standards Institute, Inc.
(ANSI) standards, Federation of European Producers of Abrasive Products (FEPA)
standards, and Japanese Industrial Standard (JIS) standards.

In some embodiments, the glass-ceramic abrasive particles have an aspect ratio
greater than 2:1, 2.5:1, or even greater than 3:1.

In some embodiments, glasses for the ceramic particles, and/or for making the
glass-ceramic abrasive particles and the ceramic particles, may be smaller than desired.
The size of such glasses can be increased, for example, via consolidation of the glass at
temperatures above glass transition temperature. This coalescing step in essence forms a
larger sized body from two or more smaller particles. For instance, the glass undergoes
glass transition (T_g) before significant crystallization occurs (T_x) as evidenced by the
existence of an endotherm (T'_g) at lower temperature than an exotherm (T'_x). The
temperature and pressure used for coalescing may depend, for example, upon composition
of the glass and the desired density of the resulting material. The temperature should be
greater than the glass transition temperature. In certain embodiments, the heating is
conducted at at least one temperature in a range from about 800°C to 1200°C (in some
embodiments, 800°C to 1000°C, 850°C to 1100°C, or even 900°C to 1000°C).

Typically, the glass is under pressure (e.g., greater than zero to 1 GPa or more)
during coalescence to aid the coalescence of the glass. Typically, the pressure is less than
100MPa (15,000 psi). In one embodiment, a charge of the particles, etc. is placed into a
die and hot-pressing is performed at temperatures above glass transition where viscous
flow of glass leads to coalescence into a relatively large part. Examples of typical
coalescing techniques include hot pressing, hot isostatic pressing, hot extrusion, hot
forging and the like (e.g., sintering, plasma assisted sintering). For example, particles
comprising glass (obtained, for example, by crushing) (including beads and
microspheres), fibers, etc. may formed into a larger particle size. Coalescing may also
result in a body shaped into a desired form (e.g., a geometric shape). Coalescing of the
glass may also be accomplished by a variety of methods, including pressure-less or pressure sintering, forging, hot extrusion, etc.).

In some embodiments, coalescing of the glass can be conducted in a gaseous atmosphere (e.g., nitrogen) at a pressure greater than 1.1 atm. (in some embodiments, at a pressure greater than 1.25 atm., 1.5 atm., 2 atm., 5 atm., or even greater than 10 atm.) sufficient to increase the rate of densification of the glass as compared to the same glass heated in the same manner except the pressure during the later heating is conducted in an atmosphere at a pressure of 1.0 atm., and wherein the gaseous atmosphere at a pressure greater than 1.1 atm. (in some embodiments, at a pressure greater than 1.25 atm., 1.5 atm., 2 atm., 5 atm., or even greater than 10 atm.) is in direct contact with at least a portion of the outer surface of at least a portion the glass being consolidated (see, for example, publication No. US 2006-00223 85-AL

In general, heat-treatment of the glasses for the ceramic particles, and/or for making the glass-ceramic abrasive particles and the ceramic particles, can be carried out in any of a variety of ways, including those known in the art for heat-treating glass to provide glass-ceramics. For example, heat-treatment can be conducted in batches, for example, using resistive, inductively or gas heated furnaces. Alternatively, for example, heat-treatment (or a portion thereof) of particles, for example, can be conducted continuously, for example, using a rotary kiln, fluidized bed furnaces, or pendulum kiln. In the case of a rotary kiln or a pendulum kiln, the material is typically fed directly into the kiln operating at the elevated temperature. In the case of a fluidized bed furnace, the glass to be heat-treated is typically suspended in a gas (e.g., air, inert, or reducing gasses).

The time at the elevated temperature may range from a few seconds (in some embodiments, even less than 5 seconds) to a few minutes to several hours. The temperature typically ranges from the $T_x$ of the glass to 1600°C, more typically from 900°C to 1600°C, and in some embodiments, from 1200°C to 1500°C. It is also within the scope of the present invention to perform some of the heat-treatment in multiple steps (e.g., one for nucleation, and another for crystal growth; wherein densification also typically occurs during the crystal growth step). When a multiple step heat-treatment is carried out, it is typically desired to control either or both the nucleation and the crystal growth rates. In general, during most ceramic processing operations, it is desired to obtain
maximum densification without significant crystal growth. Although not wanting to be bound by theory, in general, it is believed in the ceramic art that larger crystal sizes lead to reduced mechanical properties while finer average crystallite sizes lead to improved mechanical properties (e.g., higher strength and higher hardness). In particular, it is very desirable to form ceramics with densities of at least 90, 95, 97, 98, 99, or even at least 100 percent of theoretical density, wherein the average crystal sizes are less than 0.15 micrometer, or even less than 0.1 micrometer.

In some embodiments of the present invention, the glasses or ceramics comprising glass may be annealed prior to heat-treatment. In such cases annealing is typically done at a temperature less than the $T_x$ of the glass for a time from a few seconds to few hours or even days. Typically, the annealing is done for a period of less than 3 hours, or even less than an hour. Optionally, annealing may also be carried out in atmospheres other than air. Furthermore, different stages (i.e., the nucleation step and the crystal growth step) of the heat-treatment may be carried out under different atmospheres. It is believed that the $T_g$ and $T_x$, as well as the $T_x$-$T_g$ of glasses according to this invention may shift depending on the atmospheres used during the heat treatment.

One skilled in the art can determine the appropriate conditions from a Time-Temperature-Transformation (TTT) study of the glass using techniques known in the art. One skilled in the art, after reading the disclosure of the present invention, should be able to provide TTT curves for glasses used to make glass-ceramics utilized in the present invention, determine the appropriate nucleation and/or crystal growth conditions to provide glass-ceramics utilized in the present invention.

Heat-treatment may occur, for example, by feeding the material directly into a furnace at the elevated temperature. Alternatively, for example, the material may be fed into a furnace at a much lower temperature (e.g., room temperature) and then heated to desired temperature at a predetermined heating rate. It is within the scope of the present invention to conduct heat-treatment in an atmosphere other than air. In some cases it might be even desirable to heat-treat in a reducing atmosphere(s). Also, for, example, it may be desirable to heat-treat under gas pressure as in, for example, hot-isostatic press, or in gas pressure furnace. Although not wanting to be bound by theory, it is believed that atmospheres may affect oxidation states of some of the components of the glasses and
glass-ceramics. Such variation in oxidation states can bring about varying coloration of glasses and glass-ceramics. In addition, nucleation and crystallization steps can be affected by atmospheres (e.g., the atmosphere may affect the atomic mobilities of some species of the glasses).

It is also within the scope of the present invention to conduct additional heat-treatment to further improve desirable properties of the material. For example, hot-isostatic pressing may be conducted (e.g., at temperatures from about 900°C to about 1400°C) to remove residual porosity, increasing the density of the material.

The ratio of the total volume of the crystalline particles to the total volume of glass beads in the abrasive articles is in a range of 1:5 to 5:1 (in some embodiments, 1:3 to 3:1, of even 1:1.5 to 1.5:1)).

Abrasive articles according to the present invention can be made by a method comprising:

- providing a mixture comprising at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, and wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 (in some embodiments, either greater than 1.5:1 or less than 1:1.5, or even either greater than 2:1 or less than 1:2);
- heating the mixture so that the glass of the ceramic particles coalesces to provide consolidating glass of the ceramic particles to provide a body; and
- heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic (i.e., at least a portion of the glass crystallizes),


to provide the abrasive article.

Cutting tools and cutting tool inserts according to the present invention can be made by a method comprising:

- providing a mixture comprising at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, and wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the
first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 (in some embodiments, either greater than 1.5:1 or less than 1:1.5, or even either greater than 2:1 or less than 1:2);

heating the mixture so that the glass of the ceramic particles coalesces to provide consolidating glass of the ceramic particles to provide a body; and

heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic (i.e., at least a portion of the glass crystallizes), to provide the cutting tool or cutting tool inserts.

The mixture comprising the glass-ceramic abrasive particles and the plurality of ceramic particles comprising glass can be placed, for example, in a mold generally having the shape of the desired composite article, and the glass of the ceramic particles consolidated, for example, as described to provide the abrasive article, cutting tool or cutting tool insert. Optionally, glass present in the abrasive article, cutting tool, or cutting tool insert can be heat-treated, for example, as describe herein to convert at least a portion of the glass to crystalline ceramic and provide glass-ceramic (i.e., at least a portion of the glass crystallizes).

Typically, glass-ceramics are stronger than the glasses from which they are formed. Hence, the strength of the material may be adjusted, for example, by the degree to which the glass is converted to crystalline ceramic phase(s). Alternatively, or in addition, the strength of the material may also be affected, for example, by the number of nucleation sites created, which may in turn be used to affect the number, and in turn the size of the crystals of the crystalline phase(s). For additional details regarding forming glass-ceramics, see, for example, Glass-Ceramics, P.W. McMillan, Academic Press, Inc., 2nd edition, 1979.

As compared to many other types of ceramic processing (e.g., sintering of a calcined material to a dense, sintered ceramic material), there is relatively little shrinkage (typically, less than 30 percent by volume; in some embodiments, less than 20 percent, 10 percent, 5 percent, or even less than 3 percent by volume) during crystallization of the glass to form the glass-ceramic. The actual amount of shrinkage depends, for example, on the composition of the glass, the heat-treatment time, the heat-treatment temperature, the heat-treatment pressure, the density of the glass being crystallized, the relative amount(s)
of the crystalline phases formed, and the degree of crystallization. The amount of shrinkage can be measured by conventional techniques known in the art, including by dilatometry, Archimedes method, or measuring the dimensions of the material before and after heat-treatment. In some cases, there may be some evolution of volatile species during heat-treatment.

In some embodiments, the relatively low shrinkage feature may be particularly advantageous. For example, articles may be formed in the glass phase to the desired shapes and dimensions (i.e., in near-net shape), followed by heat treatment to at least partially crystallize the glass. As a result, substantial cost savings associated with the manufacturing and machining of the crystallized material may be realized.

In another aspect, for example, during heat-treatment of some exemplary glasses for making glass-ceramics utilized in the present invention, formation of phases such as \( \text{La}_2\text{Zr}_2\text{O}_7 \) and/or cubic/tetragonal \( \text{ZrO}_2 \), in some cases monoclinic \( \text{Zr}\theta_2 \), may occur at temperatures above about 900\(^0\)C. Although not wanting to be bound by theory, it is believed that zirconia-related phases are the first phases to nucleate from the glass. Formation of \( \text{Al}_2\text{O}_3 \), \( \text{ReAlO}_3 \) (wherein Re is at least one rare earth cation), \( \text{ReAl}\pi\text{Ois} \), \( \text{Re}_3\text{Al}_5\text{O}_{12} \), \( \text{YsAlSiO}_{12} \), etc. phases are believed to generally occur at temperatures above about 925\(^0\)C. Typically, crystallite size during this nucleation step is on order of nanometers. For example, crystals as small as 10-15 nanometers have been observed. For at least some embodiments, heat-treatment at about 1300\(^0\)C for about 1 hour provides a full crystallization. In general, heat-treatment times for each of the nucleation and crystal growth steps may range from a few seconds (in some embodiments, even less than 5 seconds) to several minutes to an hour or more.

The average crystal size can be determined by the line intercept method according to the ASTM standard E 112-96 "Standard Test Methods for Determining Average Grain Size". The sample is mounted in mounting resin (such as that obtained under the trade designation "TRANSOPTIC POWDER" from Buehler, Lake Bluff, IL) typically in a cylinder of resin about 2.5 cm in diameter and about 1.9 cm high. The mounted section is prepared using conventional polishing techniques using a polisher (such as that obtained from Buehler, Lake Bluff, IL under the trade designation "EPOMET 3"). The sample is polished for about 3 minutes with a diamond wheel, followed by 5 minutes of polishing.
with each of 45, 30, 15, 9, 3, and 1-micrometer slurries. The mounted and polished sample is sputtered with a thin layer of gold-palladium and viewed using a scanning electron microscopy (such as Model JSM 840A from JEOL, Peabody, MA). A typical back-scattered electron (BSE) micrograph of the microstructure found in the sample is used to determine the average crystallite size as follows. The number of crystallites that intersect per unit length ($N_L$) of a random straight line drawn across the micrograph are counted. The average crystallite size is determined from this number using the following equation.

\[
\text{Average Crystal Size} = \frac{1.5}{N_L M},
\]

where $N_L$ is the number of crystallites intersected per unit length and $M$ is the magnification of the micrograph.

In another aspect, glass-ceramics utilized in the present invention may comprise at least $I_7$$2$, 3, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, 97, 98, 99, or even 100 percent by volume crystallites. The crystallites present in glass-ceramics utilized in the present invention may have an average size of less than 1 micrometer, less than 0.5 micrometer, less than 0.3 micrometer, less than less than 0.2 micrometer, or even less than less than 0.15 micrometer.

Examples of crystalline phases which may be present in ceramics according to the present invention include: alumina (e.g., alpha and transition aluminas), REO, $Y_2O_3$, $HfO_2$, $ZrO_2$ (e.g., cubic $ZrO_2$ and tetragonal $ZrO_2$), one or more other metal oxides such as BaO, CaO, $Cr_2O_3$, CoO, CuO, $Fe_2O_3$, GeO$_2$, Li$_2$O, MgO, MnO, NiO, Na$_2$O, P$_2$O$_5$, Sc$_2$O$_3$, SiO$_2$, $Bi_2O_3$, SrO, TeO$_2$, TiO$_2$, V$_2$O$_5$, ZnO$_5$ as well as "complex metal oxides" (including complex $Al_2\theta_3$-metal oxide (e.g., complex $Al_2O_3$-REO (e.g., $ReAlO_3$ (e.g., Gd$AlO_3$, La$AlO_3$), $ReAlI_{10}18$ (e.g., LaAlInO$_8$), and $Re_3Al_5O_{12}$ (e.g., Dy$_3$Al$_5$O$_{12}$)), complex $Al_2O_3$-$Y_2O_3$ (e.g., $Y_3Al_5O_{12}$), and complex $ZrO_2$-REO (e.g., $La_2Zr_2O_7$)) and combinations thereof. Typically, ceramics according to the present invention are free of eutectic microstructure features.

In some embodiments, ceramics according to the present invention further comprise $ZrO_2$ and/or $HfO_2$ up to 30 percent by weight (in some embodiments, in a range
from 15 to 30 percent by weight ZrO$_2$ and/or HfO$_2$, based on the total weight of the ceramic.

It is also with in the scope of the present invention to substitute a portion of the aluminum cations in a complex Al$_2$O$_3$-metal oxide (e.g., complex Al$_2$O$_3$-REO and/or complex Al$_2$O$_3$Y$_2$O$_3$ (e.g., yttrium aluminate exhibiting a garnet crystal structure)). For example, a portion of the Al cations in a complex Al$_2$O$_3$Y$_2$O$_3$ may be substituted with at least one cation of an element selected from the group consisting of: Cr, Ti, Sc, Fe, Mg, Ca, Si, Co, and combinations thereof. For example, a portion of the Y cations in a complex Al$_2$O$_3$Y$_2$O$_3$ may be substituted with at least one cation of an element selected from the group consisting of: Ce, Dy, Er, Eu, Gd, Ho, La, Lu, Nd, Pr, Sm, Th, Tm, Yb, Fe, Ti, Mn, V, Cr, Co, Ni, Cu, Mg, Ca, Sr, and combinations thereof. Further, for example, a portion of the rare earth cations in a complex Al$_2$O$_3$-REO may be substituted with at least one cation of an element selected from the group consisting of: Y, Fe, Ti, Mn, V, Cr, Co, Ni, Cu, Mg, Ca, Sr, and combinations thereof. The substitution of cations as described above may affect the properties (e.g., hardness, toughness, strength, thermal conductivity, etc.) of the ceramic.

Crystals formed by heat-treating glass to provide embodiments of glass-ceramics utilized in the present invention may be, for example, acicular equiaxed, columnar, or flattened splat-like features.

Some embodiments of glasses and glass-ceramics utilized in the present invention, and some glasses used to make such glass-ceramics, comprise at least 75 percent (in some embodiments, at least 80, 85, or even at least 90; in some embodiments, in a range from 75 to 90) by weight Al$_2$O$_3$, at least 0.1 percent (in some embodiments, at least 1, at least 5, at least 10, at least 15, at least 20, or 23.9; in some embodiments, in a range from 10 to 23.9, or 15 to 23.9) by weight La$_2$O$_3$, at least 1 percent (in some embodiments, at least 5, at least 10, at least 15, at least 20, or even 24.8; in some embodiments, in a range from 10 to 24.8, 15 to 24.8) by weight Y.O$_3$, and at least 0.1 percent (in some embodiments, at least 1, at least 2, at least 3, at least 4, at least 5, at least 6, at least 7, or even 8; in some embodiments, in a range from 0.1 to 8 or 0.1 to 5, or 0.1 to 2) by weight MgO, based on the total weight of the glass or glass-ceramic, respectively.
Some embodiments of glasses and glass-ceramics utilized in the present invention, and some glasses used to make such glass-ceramics, comprise at least 75 percent (in some embodiments, at least 80, 85, or even at least 90; in some embodiments, in a range from 75 to 90) by weight $\text{Al}_2\text{O}_3$, and at least 1 percent (in some embodiments, at least 5, at least 10, at least 15, at least 20, or even 25; in some embodiments, in a range from 10 to 25, 15 to 25) by weight $\text{Y}_2\text{O}_3$, based on the total weight of the glass-ceramic or glass, respectively.

Some embodiments of glasses and glass-ceramics utilized in the present invention, and some glasses used to make such glass-ceramics, comprise at least 75 (in some embodiments, at least 80, 85, or even at least 90) percent by weight $\text{Al}_2\text{O}_3$, and at least 10 (in some embodiments, at least 15, 20, or even at least 25) percent by weight $\text{Y}_2\text{O}_3$ based on the total weight of the glass-ceramic or glass, respectively.

For some embodiments of glasses and glass-ceramics utilized in the present invention, and some glasses used to make such glass-ceramics comprising $\text{ZrO}_2$ and/or $\text{HfO}_2$, the amount of $\text{ZrO}_2$ and/or $\text{HfO}_2$ present may be at least 5, 10, 15, or even at least 20 percent by weight, based on the total weight of the glass-ceramic or glass, respectively.

Certain glasses according to the present invention may have, for example, a $T_g$ in a range of about 750°C to about 950°C.

The average hardness of the ceramics utilized in the present invention can be determined as follows. Sections of the material are mounted in mounting resin (obtained under the trade designation "TRANSOPTIC POWDER" from Buehler, Lake Bluff, IL) typically in a cylinder of resin about 2.5 cm in diameter and about 1.9 cm high. The mounted section is prepared using conventional polishing techniques using a polisher (such as that obtained from Buehler, Lake Bluff, IL under the trade designation "EPOMET 3"). The sample is polished for about 3 minutes with a diamond wheel containing 125-micrometer diamonds, followed by 5 minutes of polishing with each of 45, 30, 15, 9, 3, and 1-micrometer slurries. The microhardness measurements are made using a conventional microhardness tester (such as that obtained under the trade designation "MITUTOYO MVK-VL" from Mitutoyo Corporation, Tokyo, Japan) fitted with a Vickers indenter using a 100-gram indent load. The microhardness measurements are made according to the guidelines stated in ASTM Test Method E384 Test Methods for

Certain glasses utilized in the present invention may have, for example, an average hardness of at least 5 GPa (in some embodiments, at least 6 GPa, 7 GPa, 8 GPa, or 9 GPa; typically in a range of about 5 GPa to about 10 GPa), and glass-ceramics according to the present invention or ceramics utilized in the present invention comprising glass and crystalline ceramic at least 5 GPa (in some embodiments, at least 6 GPa, 7 GPa, 8 GPa, 9 GPa, 10 GPa, 11 GPa, 12 GPa, 13 GPa, 14 GPa, 15 GPa, 16 GPa, 17 GPa, or 18 GPa (or more); typically in a range of about 5 GPa to about 18 GPa).

Certain glasses utilized in the present invention may have, for example, a thermal expansion coefficient in a range of about $5 \times 10^{-6}$/K to about $H \times 10^{-6}$/K over a temperature range of at least 25$^\circ$C to about 900$^\circ$C.

Typically, the matrix of the abrasive article, cutting tool, or cutting tool insert have a (true) density, sometimes referred to as specific gravity, of at least 40% (in some embodiments, at least 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 95%, 90%, 95%, 96%, 97%, 98%, 99%, 99.5%, or even 100%) of theoretical density. For cutting tools and cutting tool inserts, the (true) density of the matrix is typically at least 90% (in some embodiments, at least 95%, 96%, 97%, 98%, 99%, 99.5%. or even 100%) of theoretical density. Typically, the (true) density of the glass-ceramic abrasive particles are typically at least 85% (in some embodiments, at least 90%, 95%, 96%, 97%, 98%, 99%, 99.5% or even 100%) of theoretical density.

Embodiments of abrasive articles according to the present invention have a porosity in the range from 25 to 75 (in some embodiments, in a range from 35 to 65, or even 45 to 55) percent by volume. The porosity is typically present as interconnected pores. The pores may be induced by techniques known in the art, including use of additions of porous substances such as alumina or glass bubbles, and/or fugitive materials such as organic materials. Embodiments of cutting tools and cutting tool inserts according to the present invention have a porosity in the range from 0 to 10 (in some embodiments, in a range from 0 to 5, 0 to 3, or even 0 to 2) percent by volume.
An exemplary abrasive wheel is shown in FIG. 7. Abrasive wheel 70 has glass-ceramic abrasive particles 72 (of a first composition) and pores 73 distributed in matrix 74 (of a second composition).

An exemplary honing stick is shown in FIG. 8. Honing stick 80 has glass-ceramic abrasive particles 82 (of a first composition) distributed in matrix 84 (of a second composition).

An exemplary cutting tool is shown in FIG. 9. Cutting tool 90 includes cutting tool inserts 91 having glass-ceramic abrasive particles 92 (of a first composition) distributed in matrix 94 (of a second composition).

Advantages and embodiments of this invention are further illustrated by the following non-limiting examples, but the particular materials and amounts thereof recited in these examples, as well as other conditions and details, should not be construed to unduly limit this invention. All parts and percentages are by weight unless otherwise indicated. Unless otherwise stated, all examples contained no significant amount of As₂O₃, Bi₂O₃, B₂O₃, GeO₂, P₂O₅, SiO₂, TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides.

**Feed Powder Preparation**

For preparing spray dried feed powders, the raw materials, including the solvent, binder and dispersant were batched according to Table 1, below.

**Table 1**

<table>
<thead>
<tr>
<th></th>
<th>Composition 1 (ALZ), grams</th>
<th>Composition 2 (AY), grams</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPA</td>
<td>895</td>
<td>895</td>
</tr>
<tr>
<td>Dispersant</td>
<td>33.8</td>
<td>33.8</td>
</tr>
<tr>
<td>Binder</td>
<td>67.5</td>
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</tr>
<tr>
<td>Al₂O₃</td>
<td>804</td>
<td>1121</td>
</tr>
<tr>
<td>La₂O₃</td>
<td>689</td>
<td>--</td>
</tr>
<tr>
<td>ZrO₂</td>
<td>387</td>
<td>--</td>
</tr>
</tbody>
</table>
The source of the raw materials listed in Table 1 are provided in Table 2, below.

<table>
<thead>
<tr>
<th>Source</th>
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<tbody>
<tr>
<td>Al</td>
</tr>
<tr>
<td>Y₂O₃</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Raw Material</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alumina (Al₂O₃) particles</td>
<td>Obtained from Alcoa Industrial Chemicals, Bauxite, AR, under the trade designation “Al6SG”; average particle size 0.4 micrometer</td>
</tr>
<tr>
<td>Aluminum (Al) particles</td>
<td>Obtained from Alfa Aesar, Ward Hill, MA; -325 mesh particle size.</td>
</tr>
<tr>
<td>Isopropyl alcohol 99.0 % purity (IPA)</td>
<td>VWR International, West Chester, PA</td>
</tr>
<tr>
<td>Gadolinium oxide (Gd₂O₃) particles</td>
<td>Obtained from Molycorp Inc., Mountain Pass, CA</td>
</tr>
<tr>
<td>Lanthanum oxide (La₂O₃) particles</td>
<td>Obtained from Molycorp Inc., and calcined at 700°C for 6 hours prior to batch mixing</td>
</tr>
<tr>
<td>Yttrium oxide (Y₂O₃) particles</td>
<td>Obtained from H.C. Stark Newton, MA</td>
</tr>
<tr>
<td>Zirconium oxide (ZrO₂) particles</td>
<td>Obtained from Zirconia Sales, Inc. of Marietta, GA under the trade designation “DK-2”; average particle size 2 micrometer</td>
</tr>
</tbody>
</table>

The respective raw materials were charged into an alumina jar (27 cm diameter x 23 cm high; obtained from Coors, Golden, CO), that was partially filled with 2750 grams of alumina milling media (1.27 cm; obtained from Coors. The contents of the jar was milled for 3 hours at 33 rpm. After milling was completed, the resulting slurry was emptied and separated from the milling media. The slurry was then spray dried in an atmosphere controlled spray drier, using hot nitrogen as the drying gas and a gas-atomizing nozzle. The atomizing gas was also nitrogen. The spray-dried powders were recovered, and screened through a 70-mesh (212 micrometer opening size) ro-tap screen (obtained from Tyler, Mentor, OH).

Glass Beads Preparation
Glass beads were prepared using spray dried feed powders (prepared as described in the "Feed Powder Preparation" section (above). The spray dried feed powder was fed into a hydrogen/oxygen torch flame at about 10 grams/minute. More specifically, the powder was fed into the flame through a glass funnel that was attached to a vibratory powder feeder. The resulting melted raw materials were quenched in water contained in a 19-liter (5-gallon) rectangular container (41 centimeters (cm) by 53 cm by 18 cm height) to quench the molten droplets. The water was at about 20°C, and was a continuously circulating. The torch was obtained from Bethlehem Apparatus Co., Hellertown, PA under the trade designation "BETHLEHEM BENCH BURNER PM2D, MODEL B". The torch had a central feed port (0.475 cm (3/16 inch) inner diameter) through which the feed particles were introduced into the flame. The hydrogen and oxygen flow rates for the torch were 42 standard liters per minute (SLPM) and 18 SLPM, respectively. The angle at which the flame hit the water was about 90°. The distance between the flame and water surface was about 38 cm. The resulting (quenched) particles were collected in a pan and heated at 1100°C in an electrically heated furnace until dried (about 30 minutes).

A percent crystalline yield was calculated from the resulting flame formed beads. The measurements were done as follows. A single layer of beads was spread out upon a glass slide. The beads were observed at 32x using an optical microscope. Using the crosshairs in the optical microscope eyepiece as a guide, beads that lay horizontally coincident with crosshair along a straight line were counted either transparent or opaque (i.e., crystalline) depending on their optical clarity. A total of 500 beads were counted and a percent crystalline yield was determined by the number of opaque beads divided by total beads counted and multiplied by 100. The particles were predominantly transparent (>95% by number).

Differential Thermal Analysis (DTA) was conducted on beads using the following method. A DTA run was made (using an instrument obtained from Netzsch Instruments, Selb, Germany under the trade designation "NETZSCH STA 409 DTA/TGA") using a -140+170 mesh size fraction (i.e., the fraction collected between 105-micrometer opening size and 90-micrometer opening size screens). An amount of each screened sample was placed in a 100 microliter Al2O3 sample holder. Each sample was heated in static air at a rate of 10°C/minute from room temperature (about 25°C) to 1100°C.
If the DTA trace of the beads prepared exhibited an endothermic event as evidenced by a downward change in the curve of the trace. It is believed this event was due to the glass transition ($T_g$) of the beads. Thus, the material was determined to be glass. If the same material exhibited an exothermic event as evidenced by a sharp peak in the trace, it is believed that this event was due to the crystallization ($T_x$) of the material.

Powder X-ray diffraction, XRD, (obtained under the trade designation "PHILLIPS XRG 3100" from Phillips, Mahwah, NJ), with copper Kα1 radiation of 1.54050 Angstrom) was used to verify the absence of any discernible peaks corresponding to any crystalline phases (i.e., the beads were amorphous).

**Crystallization of Glass Beads**

For crystallization of the glass (including glass beads) a heat treatment schedule above the crystallization temperature, $T_x$, of the corresponding composition of the glass was conducted.

For the glass beads, a bed of glass beads (about 0.5 cm deep) was formed in a platinum crucible (9 cm diameter x 2.5 cm height). The platinum crucible was then placed in an electrically heated furnace (obtained from CM Furnaces, Bloomfield, NJ under the trade designation "RAPID TEMP FURNACE") and heated to 1350°C at a heating rate of 10°C/min. in air. The beads were then held at 1350°C for 30 minutes, after which the power was shut off, and the furnace was allowed to cool down to room temperature. After the heat treatment all of the beads were opaque indicating the beads were crystalline.

Powder X-ray diffraction, XRD, (obtained under the trade designation "PHILLIPS XRG 3100" from Phillips, Mahwah, NJ), with copper Kα1 radiation of 1.54050 Angstrom) was used to verify the crystallinity of the material.

**Abrasive Grains Preparation**

Abrasive grains for use in making examples were prepared by taking a quantity of the glass beads (having the desired composition) and encapsulating (i.e., canning) the glass beads in stainless steel foils, and sealed under vacuum. The encapsulated beads were then placed in a Hot Isostatic Press (HIP) (obtained form American Isostatic Presses, Inc., Columbus, OH under the trade designation "IPS EAGLE-6"). The HIPing was carried out.
at a peak temperature of 1000°C, and at about 3000 atm pressure of argon gas. The HIP furnace was first ramped up to 750°C at 10°C/minute, then from 750°C to 980°C at 25°C/minute. The temperature was maintained at 980°C for 20 minutes, and was then increased to 1000°C. After 10 minutes at 1000°C the power was turned off, and the furnace allowed to cool. The argon gas pressure was applied at a rate of 37.5 atm/minute. Argon gas pressure reached 3000 atm when the temperature of the furnace was 750°C. This pressure was maintained until the temperature of the furnace was allowed to cool down to about 750°C. The pressure was released at a rate of 30 atm/minutes. The resulting disks, about 10 cm in diameter and 4.5 cm in thickness, were crushed first by using a hammer into about 1 cm size pieces and then by using a "Chipmunk" jaw crusher (Type VD, manufactured by BICO Inc., Burbank, CA) into smaller particles and screened to provide a -20+80 mesh fraction corresponding to particle sizes ranging from 177 micrometer to 850 micrometer. The crushed and screened particles retained their transparency indicating that during HIPing of the beads, and crushing and screening of the discs no significant crystallization event took place.

A bed of—25+70 mesh glass beads particles, about 0.5 cm deep, was formed in a platinum crucible. (9 cm diameter x 2.5 cm height). The platinum crucible was then placed in an electrically heated furnace ("RAPID TEMP FURNACE") and heated to 1350°C at a heating rate of 10°C/min. in air. The beads were then held at 1350°C for 30 minutes, after which the power was shut off, and the furnace was allowed to cool down to room temperature. After the heat treatment all of the crushed glass particles were opaque as observed using an optical microscope (prior to heat-treatment, the particles were transparent) indicating the particles were crystalline.

Several crystallized particles from each heat-treatment were mounted in mounting resin (obtained under the trade designation "TRANSOPTIC POWDER" from Buehler, Lake Bluff, IL) in a cylinder of resin about 2.5 cm in diameter and about 1.9 cm high. The mounted section was prepared using conventional polishing techniques using a polisher (such as that obtained from Buehler, Lake Bluff, IL under the trade designation "ECOMET 3"). The sample was polished for about 3 minutes with a diamond wheel, followed by 5 minutes of polishing with each of 45, 30, 15, 9, 3, and 1-micrometer slurries. The microhardness measurements are made using a conventional microhardness tester.
(obtained under the trade designation "MITUTOYO MVK-VL" from Mitutoyo Corporation, Tokyo, Japan) fitted with a Vickers indenter using a 100-gram indent load. The microhardness measurements are made according to the guidelines stated in ASTM Test Method E384 Test Methods for Microhardness of Materials (1991).

The mounted, polished samples used for the hardness measurements were sputtered with a thin layer of gold-palladium and viewed using a scanning electron microscopy (SEM) (Model JSM 840A from JOEL, Peabody, MA).

Example 1

Example 1 composite article was prepared from a mixture of 15 grams of composition 1 (ALZ) crystalline beads (abrasive particles) (prepared as described in the sections entitled "Glass Beads Preparation" and "Crystallization of Glass Beads", above) and 15 grams of composition 1 (ALZ) glass beads (prepared as described in the sections entitled "Glass Beads Preparation", above). The crystalline beads were screened to obtain a -100+150 mesh size fraction (i.e., the fraction collected between 100-micrometer opening size and 150-micrometer opening size screens). The glass beads were screened to obtain a -50+70 mesh size fraction (i.e., the fraction collected between 300-micrometer opening size and 212-micrometer opening size screens). The ratio of the average size of the crystalline beads to the average size of the glass beads was 0.49:1.

The 30 gram mixture was placed in a graphite die and hot-pressed using uniaxial pressing apparatus (obtained under the trade designation "HP-50", Thermal Technology Inc., Brea, CA). The hot-pressing was carried out in a positive (about 0.07 MPa (10 psi)) nitrogen atmosphere and 5 MPa (0.75 ksi)) pressure. The sample was heated to 920°C at 25°C/minute; held for 15 minutes; and then allowed to cool to room temperature.

The resulting sample was mounted in mounting resin (obtained under the trade designation "TRANSOPTIC POWDER" from Buehler, Lake Bluff, IL) in a cylinder of resin about 2.5 cm in diameter and about 1.9 cm high. The mounted section was prepared using conventional polishing techniques using a polisher (obtained from Buehler, Lake Bluff, IL under the trade designation "ECOMET 3"). The sample was polished for about 3 minutes with a diamond wheel, followed by 5 minutes of polishing with each of 45, 30, 15, 9, 3, and 1-micrometer slurries. The mounted, polished were sputtered with a thin layer
of gold-palladium and viewed using a scanning electron microscopy (SEM) (Model JSM 840A from JOEL, Peabody, MA).

An SEM micrograph of Example 1 is shown in FIG. 10, where the sample has crystalline beads (particles) 11 and pores 13 in matrix 12.

Example 2

Example 2 was prepared as described in Example 1, except (a) the composite mixture contained 18 grams of crystallized beads of composition 1 (ALZ) and 12 grams of glass beads of composition 1 (ALZ), (b) the glass beads were jet milled to an average particle size of 10 microns size. The ratio of the average size of the crystalline beads to the average size of the glass beads was 12.5:1.

An SEM micrograph of Example 2 is shown in FIG. 11, where the sample has crystalline beads (particles) 21 and pores 23 in matrix 22.

Example 3

Example 3 was prepared as described in Example 1, except (a) the abrasive grains of composition 1 (ALZ) were screened to retain -35+45 mesh fraction, and (b) the glass beads of composition 1 (ALZ) were jet milled to an average particle size of 10 microns size. The ratio of the average size of the crystalline beads to the average size of the glass beads was 43:1.

An SEM micrograph of Example 3 is shown in FIG. 12, where the sample has crystalline beads (particles) 31 in matrix 32.

Example 4

Example 4 was prepared from a composite mixture containing 15 grams of composition 1 (ALZ) glass beads (prepared as described in the sections entitled "Glass Beads Preparation", above) and 15 grams of composition 2 (AY) abrasive grains (prepared as described in the sections entitled "Glass Beads Preparation" and "Crystallization of Glass Beads", above). The glass beads were screened to obtain the -50+70 mesh size fraction (i.e., the fraction collected between 300-micrometer opening size and 212-micrometer opening size screens). The abrasive particles were screened to obtain the -
35+45 mesh size fraction (i.e., the fraction collected between 500-micrometer opening size and 355-micrometer opening size screens). The ratio of the average size of the crystalline beads to the average size of the glass beads was 1.6:1.

Figure 13 is a SEM micrograph of Example 4 material, where the sample has crystalline beads (particles) 41 and pores 43 in matrix 42.

Example 5

Example 5 was prepared as described in Example 3, except glass beads of composition 1 (ALZ) were screened to obtain the -230+270 mesh size fraction (i.e., the fraction collected between 63-micrometer opening size and 53-micrometer opening size screens) were used instead of the glass beads in the —50+70 mesh size fraction. The ratio of the average size of the crystalline beads to the average size of the glass beads was 7.4:1.

Figure 14 is a SEM micrograph of Example 5 material, where the sample has crystalline beads (particles) 51 and pores 53 in matrix 52.

Various modifications and alterations of this invention will become apparent to those skilled in the art without departing from the scope and spirit of this invention, and it should be understood that this invention is not to be unduly limited to the illustrative embodiments set forth herein.
What is claimed is:

1. A method of making an abrasive article comprising glass-ceramic abrasive particles in a ceramic matrix each comprising a mixture of metal oxides, wherein the glass-ceramic abrasive particles and the matrix each comprise at least 35 percent by weight $\text{Al}_2\text{O}_3$, based on the total weight of the glass-ceramic abrasive particle and the matrix, respectively, the method comprising:
   - providing a mixture comprising at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, and wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 to provide a body;
   - heating the mixture so that the glass of the ceramic particles coalesces to provide a body; and
   - heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic,
   to provide the abrasive article.

2. The method according to claim 1, wherein the ratio of the first average size to the second average size is either greater than 1.5:1 or less than 1:1.5.

3. The method according to claim 1, wherein the abrasive article is an abrasive wheel.

4. The method according to claim 3, wherein the abrasive wheel has a porosity in the range from 35 to 65 percent by volume.

5. The method according to claim 1, wherein the abrasive article is a honing stick.
6. The method according to claim 1 wherein the abrasive article has a porosity in the range from 35 to 65 percent by volume.

7. The method according to claim 1 wherein the glass-ceramic abrasive particles comprise crystallites having average size no greater than 200 nanometers.

8. The method according to claim 1 wherein the glass-ceramic abrasive particles and the ceramic particles each contain not more than 10 percent by weight collectively As₂O₃, Bi₂O₃, B₂O₃, GeO₂, P₂O₅, SiO₂; TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides, based on the total weight of the glass-ceramic abrasive particles and the ceramic particles, respectively.

9. The method according to claim 1 wherein the glass-ceramic abrasive particles are graded to a specified nominal grade.

10. The method according to claim 1 wherein the glass-ceramic abrasive particles have a density of at least 90 percent of theoretical density.

11. The method according to claim 1 wherein the forming at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles includes consolidating the glass present in the ceramic particles is conducted at a temperature in a range from 800°C to 1200°C.

12. The method according to claim 11, wherein the forming at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles includes consolidating the glass present in the ceramic particles is conducted at a pressure 100 MPa.

13. A method of making one of a cutting tool or a cutting tool insert comprising glass-ceramic abrasive particles in a ceramic matrix each comprising a mixture of metal oxides, wherein the glass-ceramic abrasive particles and the matrix each comprise at least
35 percent by weight Al₂O₃, based on the total weight of the glass-ceramic abrasive particle and the matrix, respectively, the method comprising:

- providing a mixture comprising at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles comprising glass, wherein the plurality of glass-ceramic abrasive particles has a first average size, and wherein the plurality of ceramic particles comprising glass has a second average size, and wherein the ratio of the first average size to the second average size is either greater than 1.4:1 or less than 1:1.4 to provide a body;
- heating the mixture so that the glass of the ceramic particles coalesces to provide a body; and
- heat treating the glass to crystallize at least a portion of the glass to provide a glass-ceramic,

to provide the cutting tool or cutting tool insert.

14. The method according to claim 13, wherein the ratio of the first average size to the second average size is either greater than 1.5:1 or less than 1:1.5.

15. The method according to claim 13, wherein the glass-ceramic abrasive particles comprise crystallites having average size no greater than 200 nanometers.

16. The method according to claim 13, wherein the cutting tool or cutting tool insert has a density of at least 85 percent of theoretical density.

17. The method according to claim 13, wherein the glass-ceramic abrasive particles and the ceramic particles each contain not more than 10 percent by weight collectively As₂O₃, Bi₂O₃, B₂O₃, GeO₂, P₂O₅, SiO₂, TeO₂, V₂O₅, alkaline oxides, and alkaline earth oxides, based on the total weight of the glass-ceramic abrasive particles and the ceramic particles, respectively.

18. The method according to claim 13, wherein the glass-ceramic abrasive particles are graded to a specified nominal grade.
19. The method according to claim 13, wherein the glass-ceramic abrasive particles have a density of at least 90 percent of theoretical density.

20. The method according to claim 13, wherein the forming at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles includes consolidating the glass present in the ceramic particles is conducted at a temperature in a range from 800°C to 1200°C.

21. The method according to claim 20, wherein the forming at least a plurality of glass-ceramic abrasive particles and a plurality of ceramic particles includes consolidating the glass present in the ceramic particles is conducted at a pressure 100MPa.
Fig. 6

Fig. 7
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

B24D 3/00(2006.01)i, B24D 3/06(2006.01)i, B24D 3/14(2006.01)i

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 8 B24B 29/00, B32B 5/16, C03B 19/10, C03C 12/00, C04B 35/06, C08K 3/40, C08K 7/16

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
Korean Utility models and applications for Utility models since 1975
Japanese Utility models and applications for Utility models since 1975

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
eKIPASS (KIPO internal) & keywords glass, ceramic, metal, oxide, abrasive, particle

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>US 6,254,981 B1 (RICHARD B CASTLE) 3 July 2001 see column 7, line 12 to column 10, line 53 and fig 1</td>
<td>1-21</td>
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<td>US 6,482,758 B1 (J K RICHARD WEBER et al ) 19 November 2002 see column 4, line 17 to column 6, line 21</td>
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<td>US 6,669,749 Bl(ANATOLY Z ROSENFLANZ et al ) 30 December 2003 see column 5, line 11 to column 10, line 24 and fig 1</td>
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Date of the actual completion of the international search

08 JUNE 2007 (08.06.2007)

Date of mailing of the international search report

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<th>Patent family member(s)</th>
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<tbody>
<tr>
<td>US 6,254,981 B1</td>
<td>03-07-2001</td>
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<td>19-11-2002</td>
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<tr>
<td>US 6,669,749 B1</td>
<td>30-12-2003</td>
<td>AU200 134695 A1</td>
<td>14-08-2001</td>
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<td></td>
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<td>KR1020030004338 A</td>
<td>14-01-2003</td>
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<td>WO0156950 A1</td>
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