Title: METHOD OF FORMING IMPRESSIONS IN A FLEXIBLE GRAPHITE MATERIAL

(57) Abstract: A method of forming impressions in a flexible graphite sheet (15) is provided. More particularly, the inventive method involves contacting a flexible graphite sheet (15) with an forming element (20) capable of oscillating against the flexible graphite sheet (15), whereby the oscillations in the element (20) are such that a plurality of impressions (14) are formed in the flexible graphite sheet (12).
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DESCRIPTION

METHOD OF FORMING IMPRESSIONS IN A FLEXIBLE GRAPHITE MATERIAL

TECHNICAL FIELD

[0001] Methods are provided for forming an impression in a flexible graphite material, especially for embossing or providing a plurality of transverse channels through a flexible graphite material. The inventive methods are particularly useful for the manufacture of materials suitable for the formation of components for flow-through capacitors, fuel cells, such as flow field plates, gas diffusion layers, and/or electrodes and such.

BACKGROUND ART

[0002] An ion exchange membrane fuel cell, more specifically a proton exchange membrane (PEM) fuel cell, produces electricity through the chemical reaction of hydrogen and oxygen in the air. Within the fuel cell, electrodes, denoted as anode and cathode, surround a polymer electrolyte to form what is generally referred to as a membrane electrode assembly, or MEA. Oftentimes, the electrodes also function as the gas diffusion layer (or GDL) of the fuel cell. A catalyst material stimulates hydrogen molecules to split into hydrogen atoms and then, at the membrane, the atoms each split into a proton and an electron. The electrons are utilized as electrical energy. The protons migrate through the electrolyte and combine with oxygen and electrons to form water.

[0003] A PEM fuel cell includes a membrane electrode assembly sandwiched between two flow field plates. Conventionally, the membrane electrode assembly consists of random-oriented carbon fiber paper electrodes (anode and cathode) with a thin layer of a catalyst material, particularly platinum or a platinum group metal coated on isotropic carbon particles, such as lamp black, bonded to either side of a proton exchange membrane disposed between the electrodes. In operation, hydrogen flows through channels in one of the flow field plates to the anode, where the catalyst promotes its separation into hydrogen atoms and thereafter into protons that pass through the membrane and electrons that flow through an external
load. Air flows through the channels in the other flow field plate to the cathode, where the oxygen in the air is separated into oxygen atoms, which joins with the protons through the proton exchange membrane and the electrons through the circuit, and combine to form water. Since the membrane is an insulator, the electrons travel through an external circuit in which the electricity is utilized, and join with protons at the cathode. An air stream on the cathode side is one mechanism by which the water formed by combination of the hydrogen and oxygen is removed. Combinations of such fuel cells are used in a fuel cell stack to provide the desired voltage.

[0004] It has been disclosed that a graphite sheet that has been provided with channels, which are preferably smooth-sided, and which pass between the parallel, opposed surfaces of the flexible graphite sheet and are separated by walls of compressed expandable graphite, can be used to form gas diffusion layers for PEM fuel cells. As taught by Mercuri, Weber and Wardrip in U.S. Patent No. 6,418,671, the disclosure of which is incorporated herein by reference, the channels can be formed in the flexible graphite sheet at a plurality of locations by a compressive mechanical impact, such as by use of rollers having truncated protrusions extending therefrom. The channel pattern can be devised in order to control, optimize or maximize fluid flow through the channels, as desired. For instance, the pattern formed in the flexible graphite sheet can comprise selective placement of the channels, or it can comprise variations in channel density or channel shape in order to, for instance, reduce or minimize flooding, control gas flow, restrict water flow, equalize fluid pressure along the surface of the electrode when in use, or for other purposes. See, for instance, Mercuri and Krassowski in International Publication No. WO 02/41421 A1.

[0005] In the method described in the Mercuri et al. '671 patent, the impact force is preferably delivered using a patterned roller, suitably controlled to provide well-formed perforations in the graphite sheet. In the course of impacting the flexible graphite sheet to form channels, graphite is displaced within the sheet to disrupt and deform the parallel orientation of the expanded graphite particles. In effect the displaced graphite is being "die-molded" by the sides of adjacent protrusions and the smooth surface of
the roller. This can reduce the anisotropy in the flexible graphite sheet and thus increase the electrical and thermal conductivity of the sheet in the direction transverse to the opposed surfaces. A similar effect is achieved with frusto-conical and parallel-sided peg-shaped flat-ended protrusions.

[0006] Compressive force may also be used to form the continuous reactant flow channel in the material used to form a flow field plate (hereinafter "FFP"). Typically an embossing tool is used to compress the graphite sheet and emboss the channels in the sheet. Unlike, the GDL, the channel(s) in the FFP do not extend through the FFP from one opposed surface to a second surface. Typically, the channel(s) is on one surface of the FFP.

[0007] In addition, and as taught by Mercuri et al. in U.S. Patent No. 6,528,199, the disclosure of which is incorporated herein by reference, a combination GDL/FFP can be provided, wherein a reactant flow channel is formed in a graphite sheet that has been provided with transverse channels. Therefore, both the fluid flow function of an FFP and the fluid diffusion function of a GDL can be combined in a single component.

[0008] The formation of embossments by compression, and especially the formation of through-channels in a flexible graphite sheet by compression, can have drawbacks, however. For example, such processes may require use of a backing material to both support the flexible graphite material during compression and to allow for complete penetration of the forming element through the flexible graphite sheet to form the transverse channels from one opposed surface to the other. Depending on the nature of the backing material employed, it may have a tendency to "crinkle" or deform during compression, which can result in a complimentary shaped flaw in the component. Also, use of a backing sheet can add additional cost to the manufacturing process. Thus there is the need to develop methods of making a flexible graphite material having embossments or through-channels that do not require the use of a backing material. Such processes are provided by the present invention.

[0009] Graphites are made up of layered planes of hexagonal arrays or networks of carbon atoms. These layered planes of hexagonally arranged
carbon atoms are substantially flat and are oriented or ordered so as to be substantially parallel and equidistant to one another. The substantially flat, parallel equidistant sheets or layers of carbon atoms, usually referred to as graphene layers or basal planes, are linked or bonded together and groups thereof are arranged in crystallites. Highly ordered graphites consist of crystallites of considerable size: the crystallites being highly aligned or oriented with respect to each other and having well ordered carbon layers. In other words, highly ordered graphites have a high degree of preferred crystallite orientation. It should be noted that graphites possess anisotropic structures and thus exhibit or possess many properties that are highly directional, e.g., thermal and electrical conductivity and fluid diffusion.

[0010] Graphites may be characterized as laminated structures of carbon, that is, structures consisting of superposed layers or laminae of carbon atoms joined together by weak van der Waals forces. In considering the graphite structure, two axes or directions are usually noted, to wit, the “c” axis or direction and the “a” axes or directions. For simplicity, the “c” axis or direction may be considered as the direction perpendicular to the carbon layers. The “a” axes or directions may be considered as the directions parallel to the carbon layers or the directions perpendicular to the “c” direction. The graphites suitable for manufacturing flexible graphite sheets possess a very high degree of orientation.

[0011] As noted above, the bonding forces holding the parallel layers of carbon atoms together are only weak van der Waals forces. Natural graphites can be treated so that the spacing between the superposed carbon layers or laminae can be appreciably opened up so as to provide a marked expansion in the direction perpendicular to the layers, that is, in the “c” direction, and thus form an expanded or intumesced graphite structure in which the laminar character of the carbon layers is substantially retained.

[0012] Graphite flake which has been greatly expanded and more particularly expanded so as to have a final thickness or “c” direction dimension which is as much as about 80 or more times the original “c” direction dimension can be formed without the use of a binder into cohesive or integrated sheets of expanded graphite, e.g. webs, papers, strips, tapes,
foils, mats or the like (typically referred to as "flexible graphite"). The formation of graphite particles which have been expanded to have a final thickness or "c" dimension which is as much as about 80 times or more the original "c" direction dimension into integrated flexible sheets by compression, without the use of any binding material, is believed to be possible due to the mechanical interlocking, or cohesion, which is achieved between the voluminously expanded graphite particles.

[0013] In addition to flexibility, the sheet material, as noted above, has also been found to possess a high degree of anisotropy with respect to thermal and electrical conductivity and fluid diffusion, comparable to the natural graphite starting material due to orientation of the expanded graphite particles and graphite layers substantially parallel to the opposed faces of the sheet resulting from very high compression, e.g. roll pressing. Sheet material thus produced has excellent flexibility, good strength and a very high degree of orientation.

[0014] Briefly, the process of producing flexible, binderless anisotropic graphite sheet material, e.g. web, paper, strip, tape, foil, mat, or the like, comprises compressing or compacting under a predetermined load and in the absence of a binder, expanded graphite particles which have a "c" direction dimension which is as much as about 80 or more times that of the original particles so as to form a substantially flat, flexible, integrated graphite sheet. The expanded graphite particles that generally are worm-like or vermiform in appearance, once compressed, will maintain the compression set and alignment with the opposed major surfaces of the sheet. The density and thickness of the sheet material can be varied by controlling the degree of compression. The density of the sheet material can be within the range of from about 0.04 g/cc to about 2.0 g/cc. The flexible graphite sheet material exhibits an appreciable degree of anisotropy due to the alignment of graphite particles parallel to the major opposed, parallel surfaces of the sheet, with the degree of anisotropy increasing upon roll pressing of the sheet material to increased density. In roll pressed anisotropic sheet material, the thickness, i.e. the direction perpendicular to the opposed, parallel sheet surfaces comprises the "c" direction and the directions ranging along the
length and width, i.e. along or parallel to the opposed, major surfaces comprises the “a” directions and the thermal and electrical properties of the sheet are very different, by orders of magnitude, for the “c” and “a” directions.

DISCLOSURE OF THE INVENTION

[0015] One aspect of the invention is a method of making impressions in a flexible graphite material. The method includes the step of oscillating a forming element against a flexible graphite sheet, thereby forming a plurality of impressions, such as transverse fluid channels in the sheet. The plurality of transverse fluid channels pass through the sheet between first and second parallel, opposed surfaces of the sheet.

[0016] A further aspect of the invention comprises a method of embossing a flexible graphite sheet, which includes the step of oscillating a forming element against a sheet of flexible graphite, thereby forming a plurality of substantially adjacent indentations on a first surface of the sheet. In an advantageous aspect of this embodiment of the invention, at least a portion of the adjacent indentations combine to form a passage for fluid flow across the surface of the sheet.

[0017] In another aspect of the invention, the embossed sheets can be employed in the formation of components for a PEM fuel cell, including a gas diffusion layer and a flow field plate. In addition, the embossed sheets can also be used to form components for a flow-through capacitor, as described by Calarco et al. in U.S. Patent No. 6,410,128.

[0018] Practice of the present invention provides a method to achieve a high degree of variation in the density of channels in a resulting flexible graphite sheet (i.e., the number of channels per square inch of sheet), which can be of benefit in fuel cell applications; additionally, the inventive method may also be used to vary the location of the channels through, or fluid flow passages across, the material.

[0019] Furthermore, another advantage is that the material may be formed without the use of a backing material. An additional process advantage is that the inventive methods may be used in a continuous process, rather than the batch process often employed. Also, the frequency of
the oscillations in the forming element may be used to control the density of the channel pattern and the amplitude of the oscillations in the forming element may be used to control the depth of the penetration into a flexible graphite sheet. Thus, the range of channel patterns/geometries available to the user is thereby greatly expanded.

[0020] Therefore, an apparatus used to practice the inventive method may be used to configure flexible graphite materials, such as substrates used to form fuel cell components, with a wide variety of patterns of channels or indentations. The various patterns may be used to determine the most efficient pattern of the particular component or application to maximize the operating parameters.

[0021] It is to be understood that both the foregoing general description and the following detailed description present embodiments of the invention, and are intended to provide an overview or framework for understanding the nature and character of the invention as it is claimed. The accompanying drawings are included to provide a further understanding of the invention, and are incorporated in and constitute a part of this specification. The drawings illustrate various embodiments of the invention, and together with the description serve to explain the principles and operations of the invention.

[0022] Fig. 1 is one embodiment of a tool for forming a plurality of channels in a sheet of flexible graphite.

[0023] Fig. 2 is one embodiment of a contact element useful in the tool of Figure 1.

[0024] Fig. 3 is another embodiment of a tool for forming a plurality of channels in a sheet of flexible graphite.

[0025] Fig. 4 is another embodiment of a contact element useful in the tool of Figure 3.

[0026] Fig. 5 is a partial top view of a sheet of flexible graphite having a plurality of channels formed in accordance with the invention.

[0027] Fig. 5A is a partial side cross-section view of the sheet of Figure 1.
[0028] Fig. 6 is a partial side cross-section view of a sheet of flexible graphite having a plurality of indentations formed in accordance with another embodiment of the invention.

DISCLOSURE OF THE INVENTION

[00209] Embodiments of the present invention will now be described in greater detail with respect to the aforementioned drawings. Like or similar reference numerals will be used whenever possible.

[0030] The methods of the present invention relate to making impressions in flexible graphite sheets. Graphite is a crystalline form of carbon comprising atoms covalently bonded in flat layered planes with weaker bonds between the planes. By treating particles of graphite, such as natural graphite flake, with an intercalant of, e.g. a solution of sulfuric and nitric acid, the crystal structure of the graphite reacts to form a compound of graphite and the intercalant. The treated particles of graphite are hereafter referred to as “particles of intercalated graphite.” Upon exposure to high temperature, the intercalant within the graphite volatilizes, causing the particles of intercalated graphite to expand in dimension as much as about 80 or more times its original volume in an accordion-like fashion in the “c” direction, i.e. in the direction perpendicular to the crystalline planes of the graphite. The exfoliated graphite particles are vermiciform in appearance, and are therefore commonly referred to as worms. The worms may be compressed together into flexible sheets that, unlike the original graphite flakes, can be formed and cut into various shapes and provided with small transverse openings by deforming mechanical impact.

[0031] Graphite starting materials for the flexible sheets suitable for use in the present invention include highly graphitic carbonaceous materials capable of intercalating organic and inorganic acids as well as halogens and then expanding when exposed to heat. These highly graphitic carbonaceous materials most preferably have a degree of graphitization of about 1.0. As used in this disclosure, the term “degree of graphitization” refers to the value g according to the formula:

$$g = 3.45 \cdot d(002)$$

0.095
where \( d(002) \) is the spacing between the graphitic layers of the carbons in the crystal structure measured in Angstrom units. The spacing \( d \) between graphite layers is measured by standard X-ray diffraction techniques. The positions of diffraction peaks corresponding to the (002), (004) and (006) Miller Indices are measured, and standard least-squares techniques are employed to derive spacing which minimizes the total error for all of these peaks. Examples of highly graphitic carbonaceous materials include natural graphites from various sources, as well as other carbonaceous materials such as carbons prepared by chemical vapor deposition and the like. Natural graphite is most preferred.

[0032] The graphite starting materials for the flexible sheets used in the present invention may contain non-carbon components so long as the crystal structure of the starting materials maintains the required degree of graphitization and they are capable of exfoliation. Generally, any carbon-containing material, the crystal structure of which possesses the required degree of graphitization and which can be exfoliated, is suitable for use with the present invention. Such graphite preferably has an ash content of less than twenty weight percent. More preferably, the graphite employed for the present invention will have a purity of at least about 94%. In the most preferred embodiment, such as for fuel cell applications, the graphite employed will have a purity of at least about 99%.

[0033] A common method for manufacturing graphite sheet is described by Shane et al. in U.S. Patent No. 3,404,061, the disclosure of which is incorporated herein by reference. In the typical practice of the Shane et al. method, natural graphite flakes are intercalated by dispersing the flakes in a solution containing \( \text{e.g., a mixture of nitric and sulfuric acid,} \) advantageously at a level of about 20 to about 300 parts by weight of intercalant solution per 100 parts by weight of graphite flakes (pph). The intercalation solution contains oxidizing and other intercalating agents known in the art. Examples include those containing oxidizing agents and oxidizing mixtures, such as solutions containing nitric acid, potassium chlorate, chromic acid, potassium permanganate, potassium chromate, potassium dichromate, perchloric acid, and the like, or mixtures, such as for
example, concentrated nitric acid and chlorate, chromic acid and phosphoric acid, sulfuric acid and nitric acid, or mixtures of a strong organic acid, e.g. trifluoroacetic acid, and a strong oxidizing agent soluble in the organic acid. Alternatively, an electric potential can be used to bring about oxidation of the graphite. Chemical species that can be introduced into the graphite crystal using electrolytic oxidation include sulfuric acid as well as other acids.

[0034] In a preferred embodiment, the intercalating agent is a solution of a mixture of sulfuric acid, or sulfuric acid and phosphoric acid, and an oxidizing agent, i.e. nitric acid, perchloric acid, chromic acid, potassium permanganate, hydrogen peroxide, iodic or periodic acids, or the like. Although less preferred, the intercalation solution may contain metal halides such as ferric chloride, and ferric chloride mixed with sulfuric acid, or a halide, such as bromine as a solution of bromine and sulfuric acid or bromine in an organic solvent.

[0035] The quantity of intercalation solution may range from about 20 to about 150 pph and more typically about 50 to about 120 pph. After the flakes are intercalated, any excess solution is drained from the flakes and the flakes are water-washed. Alternatively, the quantity of the intercalation solution may be limited to between about 10 and about 50 pph, which permits the washing step to be eliminated as taught and described in U.S. Patent No. 4,895,713, the disclosure of which is also herein incorporated by reference.

[0036] The particles of graphite flake treated with intercalation solution can optionally be contacted, e.g. by blending, with a reducing organic agent selected from alcohols, sugars, aldehydes and esters which are reactive with the surface film of oxidizing intercalating solution at temperatures in the range of 25°C and 125°C. Suitable specific organic agents include hexadecanol, octadecanol, 1-octanol, 2-octanol, decylalcohol, 1, 10 decanediol, decylaldehyde, 1-propanol, 1,3 propanediol, ethyleneglycol, polypropylene glycol, dextrose, fructose, lactose, sucrose, potato starch, ethylene glycol monostearate, diethylene glycol dibenzoate, propylene glycol monostearate, glycerol monostearate, dimethyl oxylate, diethyl oxylate, methyl formate, ethyl formate, ascorbic acid and lignin-derived compounds, such as sodium
lignosulfate. The amount of organic reducing agent is suitably from about 0.5 to 4% by weight of the particles of graphite flake.

[0037] The use of an expansion aid applied prior to, during or immediately after intercalation can also provide improvements. Among these improvements can be reduced exfoliation temperature and increased expanded volume (also referred to as "worm volume"). An expansion aid in this context will advantageously be an organic material sufficiently soluble in the intercalation solution to achieve an improvement in expansion. More narrowly, organic materials of this type that contain carbon, hydrogen and oxygen, preferably exclusively, may be employed. Carboxylic acids have been found especially effective. A suitable carboxylic acid useful as the expansion aid can be selected from aromatic, aliphatic or cycloaliphatic, straight chain or branched chain, saturated and unsaturated monocarboxylic acids, dicarboxylic acids and polycarboxylic acids which have at least 1 carbon atom, and preferably up to about 15 carbon atoms, which is soluble in the intercalation solution in amounts effective to provide a measurable improvement of one or more aspects of exfoliation. Suitable organic solvents can be employed to improve solubility of an organic expansion aid in the intercalation solution.

[0038] Representative examples of saturated aliphatic carboxylic acids are acids such as those of the formula H(CH₂)ₙCOOH wherein n is a number of from 0 to about 5, including formic, acetic, propionic, butyric, pentanoic, hexanoic, and the like. In place of the carboxylic acids, the anhydrides or reactive carboxylic acid derivatives such as alkyl esters can also be employed. Representative of alkyl esters are methyl formate and ethyl formate. Sulfuric acid, nitric acid and other known aqueous intercalants have the ability to decompose formic acid, ultimately to water and carbon dioxide. Because of this, formic acid and other sensitive expansion aids are advantageously contacted with the graphite flake prior to immersion of the flake in aqueous intercalant. Representative of dicarboxylic acids are aliphatic dicarboxylic acids having 2-12 carbon atoms, in particular oxalic acid, fumaric acid, malonic acid, maleic acid, succinic acid, glutaric acid, adipic acid, 1,5-pentanedicarboxylic acid, 1,6-hexanedicarboxylic acid, 1,10-
decanedicarboxylic acid, cyclohexane-1,4-dicarboxylic acid and aromatic dicarboxylic acids such as phthalic acid or terephthalic acid. Representative of alkyl esters are dimethyl oxylate and diethyl oxylate. Representative of cycloaliphatic acids is cyclohexane carboxylic acid and of aromatic carboxylic acids are benzoic acid, naphthoic acid, anthranilic acid, p-aminobenzoic acid, salicylic acid, o-, m- and p-tolyl acids, methoxy and ethoxybenzoic acids, acetoacetamidobenzoic acids and, acetamidobenzoic acids, phenylactic acid and naphthoic acids. Representative of hydroxy aromatic acids are hydroxybenzoic acid, 3-hydroxy-1-naphthoic acid, 3-hydroxy-2-naphthoic acid, 4-hydroxy-2-naphthoic acid, 5-hydroxy-1-naphthoic acid, 5-hydroxy-2-naphthoic acid, 6-hydroxy-2-naphthoic acid and 7-hydroxy-2-naphthoic acid. Prominent among the polycarboxylic acids is citric acid.

[0039] The intercalation solution will be aqueous and will preferably contain an amount of expansion aid of from about 1 to 10%, the amount being effective to enhance exfoliation. In the embodiment wherein the expansion aid is contacted with the graphite flake prior to or after immersing in the aqueous intercalation solution, the expansion aid can be admixed with the graphite by suitable means, such as a V-blender, typically in an amount of from about 0.2% to about 10% by weight of the graphite flake.

[0040] After intercalating the graphite flake, and following the blending of the intercalant coated intercalated graphite flake with the organic reducing agent, the blend is exposed to temperatures in the range of 25° to 125°C to promote reaction of the reducing agent and intercalant coating. The heating period is up to about 2 hours, with shorter heating periods, e.g., at least about 10 minutes, for higher temperatures in the above-noted range. Times of one-half hour or less, e.g., on the order of 10 to 25 minutes, can be employed at the higher temperatures.

[0041] The above described methods for intercalating and exfoliating graphite flake may beneficially be augmented by a pretreatment of the graphite flake at graphitization temperatures, i.e. temperatures in the range of about 3000°C and above and by the inclusion in the intercalant of a lubricious additive.
[0042] The pretreatment, or annealing, of the graphite flake results in significantly increased expansion (i.e., increase in expansion volume of up to 300% or greater) when the flake is subsequently subjected to intercalation and exfoliation. Indeed, desirably, the increase in expansion is at least about 50%, as compared to similar processing without the annealing step. The temperatures employed for the annealing step should not be significantly below 3000°C, because temperatures even 100°C lower result in substantially reduced expansion.

[0043] The annealing of the present invention is performed for a period of time sufficient to result in a flake having an enhanced degree of expansion upon intercalation and subsequent exfoliation. Typically the time required will be 1 hour or more, preferably 1 to 3 hours and will most advantageously proceed in an inert environment. For maximum beneficial results, the annealed graphite flake will also be subjected to other processes known in the art to enhance the degree expansion — namely intercalation in the presence of an organic reducing agent, an intercalation aid such as an organic acid, and a surfactant wash following intercalation. Moreover, for maximum beneficial results, the intercalation step may be repeated.

[0044] The annealing step of the instant invention may be performed in an induction furnace or other such apparatus as is known and appreciated in the art of graphitization; for the temperatures here employed, which are in the range of 3000°C, are at the high end of the range encountered in graphitization processes.

[0045] Because it has been observed that the worms produced using graphite subjected to pre-intercalation annealing can sometimes “clump” together, which can negatively impact area weight uniformity, an additive that assists in the formation of “free flowing” worms is highly desirable. The addition of a lubrious additive to the intercalation solution facilitates the more uniform distribution of the worms across the bed of a compression apparatus (such as the bed of a calender station conventionally used for compressing (or “calendering”) graphite worms into flexible graphite sheet. The resulting sheet therefore has higher area weight uniformity and greater tensile strength. The lubrious additive is preferably a long chain
hydrocarbon, more preferably a hydrocarbon having at least about 10 carbons. Other organic compounds having long chain hydrocarbon groups, even if other functional groups are present, can also be employed.

[0046] More preferably, the lubricious additive is an oil, with a mineral oil being most preferred, especially considering the fact that mineral oils are less prone to rancidity and odors, which can be an important consideration for long term storage. It will be noted that certain of the expansion aids detailed above also meet the definition of a lubricious additive. When these materials are used as the expansion aid, it may not be necessary to include a separate lubricious additive in the intercalant.

[0047] The lubricious additive is present in the intercalant in an amount of at least about 1.4 pph, more preferably at least about 1.8 pph. Although the upper limit of the inclusion of lubricious additive is not as critical as the lower limit, there does not appear to be any significant additional advantage to including the lubricious additive at a level of greater than about 4 pph.

[0048] The thus treated particles of graphite are sometimes referred to as “particles of intercalated graphite.” Upon exposure to high temperature, e.g. temperatures of at least about 160°C and especially about 700°C to 1200°C and higher, the particles of intercalated graphite expand as much as about 80 to 1000 or more times their original volume in an accordion-like fashion in the c-direction, i.e. in the direction perpendicular to the crystalline planes of the constituent graphite particles. The expanded, i.e. exfoliated, graphite particles are vermiciform in appearance, and are therefore commonly referred to as worms. The worms may be compressed together into flexible sheets that, unlike the original graphite flakes, can be formed and cut into various shapes and provided with small transverse openings by deforming mechanical impact as hereinafter described.

[0049] Flexible graphite sheet and foil are coherent, with good handling strength, and are suitably compressed, e.g. by roll-pressing, to a thickness of about 0.075 mm to 3.75 mm and a typical density of about 0.1 to 1.5 grams per cubic centimeter (g/cc). From about 1.5-30% by weight of ceramic additives can be blended with the intercalated graphite flakes as
described in U.S. Patent No. 5,902,762 (which is incorporated herein by reference) to provide enhanced resin impregnation in the final flexible graphite product. The additives include ceramic fiber particles having a length of about 0.15 to 1.5 millimeters. The width of the particles is suitably from about 0.04 to 0.004 mm. The ceramic fiber particles are non-reactive and non-adhering to graphite and are stable at temperatures up to about 1100°C, preferably about 1400°C or higher. Suitable ceramic fiber particles are formed of macerated quartz glass fibers, carbon and graphite fibers, zirconia, boron nitride, silicon carbide and magnesia fibers, naturally occurring mineral fibers such as calcium metasilicate fibers, calcium aluminum silicate fibers, aluminum oxide fibers and the like.

[0050] The flexible graphite sheet can also, at times, be advantageously treated with resin and the absorbed resin, after curing, enhances the moisture resistance and handling strength, i.e. stiffness, of the flexible graphite sheet as well as “fixing” the morphology of the sheet. Suitable resin content is preferably at least about 5% by weight, more preferably about 10 to 35% by weight, and suitably up to about 60% by weight. Resins found especially useful in the practice of the present invention include acrylic-, epoxy- and phenolic-based resin systems, fluoro-based polymers, or mixtures thereof. Suitable epoxy resin systems include those based on diglycidyl ether of bisphenol A (DGEBA) and other multifunctional resin systems; phenolic resins that can be employed include resole and novolac phenolics. Optionally, the flexible graphite may be impregnated with fibers and/or salts in addition to the resin or in place of the resin. Additionally, reactive or non-reactive additives may be employed with the resin system to modify properties (such as tack, material flow, hydrophobicity, etc.).

[0051] Alternatively, the flexible graphite sheets of the present invention may utilize particles of reground flexible graphite sheets rather than freshly expanded worms. The sheets may be newly formed sheet material, recycled sheet material, scrap sheet material, or any other suitable source.
[0052] Also the processes of the present invention may use a blend of virgin materials and recycled materials.

[0053] The source material for recycled materials may be sheets or trimmed portions of sheets that have been compression molded as described above, or sheets that have been compressed with, for example, pre-calendering rolls, but have not yet been impregnated with resin. Furthermore, the source material may be sheets or trimmed portions of sheets that have been impregnated with resin, but not yet cured, or sheets or trimmed portions of sheets that have been impregnated with resin and cured. The source material may also be recycled flexible graphite PEM fuel cell components such as flow field plates or electrodes. Each of the various sources of graphite may be used as is or blended with natural graphite flakes.

[0054] Once the source material of flexible graphite sheets is available, it can then be comminuted by known processes or devices, such as a jet mill, air mill, blender, etc. to produce particles. Preferably, a majority of the particles have a diameter such that they will pass through 20 U.S. mesh; more preferably a major portion (greater than about 20%, most preferably greater than about 50%) will not pass through 80 U.S. mesh. Most preferably the particles have a particle size of no greater than about 20 mesh. It may be desirable to cool the flexible graphite sheet when it is resin-impregnated as it is being comminuted to avoid heat damage to the resin system during the comminution process.

[0055] The size of the comminuted particles may be chosen so as to balance machinability and formability of the graphite article with the thermal characteristics desired. Thus, smaller particles will result in a graphite article which is easier to machine and/or form, whereas larger particles will result in a graphite article having higher anisotropy, and, therefore, greater in-plane electrical and thermal conductivity.

[0056] If the source material has been resin impregnated, then preferably the resin is removed from the particles. Details of the resin removal are further described below.
[0057] Once the source material is comminuted, and any resin is removed, it is then re-expanded. The re-expansion may occur by using the intercalation and exfoliation process described above and those described in U.S. Patent No. 3,404,061 to Shane et al. and U.S. Patent No. 4,895,713 to Greinke et al.

[0058] Typically, after intercalation the particles are exfoliated by heating the intercalated particles in a furnace. During this exfoliation step, intercalated natural graphite flakes may be added to the recycled intercalated particles. Preferably, during the re-expansion step the particles are expanded to have a specific volume in the range of at least about 100 cc/g and up to about 350 cc/g or greater. Finally, after the re-expansion step, the re-expanded particles may be compressed into flexible sheets, as hereinafter described.

[0059] If the starting material has been impregnated with a resin, the resin should preferably be at least partially removed from the particles. This removal step should occur between the comminuting step and the re-expanding step.

[0060] In one embodiment, the removing step includes heating the resin containing regrind particles, such as over an open flame. More specifically, the impregnated resin may be heated to a temperature of at least about 250°C to effect resin removal. During this heating step care should be taken to avoid flashing of the resin decomposition products; this can be done by careful heating in air or by heating in an inert atmosphere. Preferably, the heating should be in the range of from about 400 °C to about 800 °C for a time in the range of from at least about 10 and up to about 150 minutes or longer.

[0061] Additionally, the resin removal step may result in increased tensile strength of the resulting article produced from the molding process as compared to a similar method in which the resin is not removed. The resin removal step may also be advantageous because during the expansion step (i.e., intercalation and exfoliation), when the resin is mixed with the intercalation chemicals, it may in certain instances create toxic byproducts.
Thus, by removing the resin before the expansion step a superior product is obtained such as the increased strength characteristics discussed above. The increased strength characteristics are a result of in part because of increased expansion. With the resin present in the particles, expansion may be restricted.

In addition to strength characteristics and environmental concerns, resin may be removed prior to intercalation in view of concerns about the resin possibly creating a run away exothermic reaction with the acid.

In view of the above, preferably a majority of the resin is removed. More preferably, greater than about 75% of the resin is removed. Most preferably, greater than 99% of the resin is removed.

Once the flexible graphite sheet is comminuted, it is formed into the desired shape and then cured (when resin impregnated) in the preferred embodiment. Alternatively, the sheet can be cured prior to being comminuted, although post-comminution cure is preferred.

In accordance with Figs.5 and 5A, a sheet of flexible graphite 12 prepared as described above and having a plurality of channels 14 is shown. In this embodiment, channels 14 extend from a first surface 16 of sheet 12 to a second surface 18 of sheet 12. Preferably, first surface 16 and second surface 18 are generally parallel opposed surfaces. In practice, sheet 12 may be resin-impregnated or not impregnated, as described herein.

As noted hereinabove, the density of the channels 14 may be varied as desired. When sheet 12 is to be used in the formation of GDL, sheet 12 preferably has a channel density of at least about 215 channels per sq. centimeter. More preferably, the channel density is at least about 310 channels per sq. centimeter, most preferably at least about 390 channels per sq. centimeter.

The inventive method comprises oscillating a forming element 20 against flexible graphite sheet 12 in order to form impressions (i.e., channels, embossments such as fluid flow passages, etc.) therein. Embodiments of forming element 20 are shown in Figs. 1 and 3. As depicted in Figs. 1 and 2, forming element 20 comprises a contact element 22, shown
as a needle, and an oscillating element 24 in operative connection with contact element 22, so as to cause contact element 22 to oscillate in such a manner as to emboss or perforate sheet 12. Contact element 22 may assume any desired shape, depending on the nature of the embossment desired. As illustrated, contact element 22 assumes a cylindrical, flat-ended configuration. Alternatively, contact element (denoted 22a in Fig. 4) may have a tapered end, preferably having a degree of taper of greater than about 10° and less than about 90°. However, depending on the shape of channels 14 desired, contact element 22 may have a corresponding cross-section, such as square, oval, circular, rectangular, etc. Additionally, oscillating element 24 comprises any apparatus that is able to create oscillations in contact element 22 with respect to sheet 12.

[0069] Although virtually any apparatus that can provide a vibratory motion can be adapted for use as oscillating element 24, one apparatus found to be effective is a device intended to be used for marking, called a pin marking system, such as a Pinstamp® device available from Telesis Technologies, Inc. of Circleville, Ohio. Indeed, any tool that is capable of moving contact element 22 back and forth from sheet 12, or vibrating contact element 22 along the surface of sheet 12, may be used as oscillating element 24. In fact, it will be noted that relatively simple vibratory devices, such as the motors found in household items like electric shavers, can be employed to create the oscillations in contact element 22.

[0070] A controller may be added to the forming element 20 to control or vary the frequency or nature (such as amplitude or orientation of the oscillations with respect to flexible graphite sheet 12) of the oscillation of contact element 22. For instance, the frequency and nature of the oscillations can be employed to produce different patterns and/or geometries (such as cross-sectional shape) of channels 14 without requiring additional equipment. Additionally, manipulation of the amplitude of the oscillation includes the ability to vary the penetration of contacting element 22 into sheet 12. For example the amplitude of oscillation may be varied such that contacting element 22 may form a channel that extends from first surface 16 of sheet 12, through sheet 12, and out second surface 18 of sheet 12, as
illustrated in Fig. 5A. In another example, illustrated in fig. 6, the amplitude may be manipulated such that forming element 20 may form an indentation 14a in surface 16 of sheet 12. In comparison to channel 14, indentation 14a will not extend to the second surface of sheet 12. Combinations of channels and indentations (including indentations of varying depths) can also be created.

[0071] By use of the inventive apparatus, the forming of channels through, and/or embossments in, graphite sheets can be accomplished without the use of a backing material, and certainly not a sacrificial backing material. In addition, control of the channels and embossments, in terms of size, shape and placement, can be controlled and varied to a greater degree than in conventional perforating/embossing techniques.

[0072] All cited patents and publications referred to in this application are incorporated by reference.

[0073] The invention thus being described, it will be obvious that it may be varied in many ways. Such variations are not to be regarded as a departure from the spirit and scope of the present invention and all such modifications as would be obvious to one skilled in the art are intended to be included within the scope of the following claims.
What is claimed is

1. A method of forming impressions in a flexible graphite sheet comprising contacting a flexible graphite sheet with a forming element capable of oscillating against the flexible graphite sheet, whereby the oscillations are such that a plurality of impressions are formed in the flexible graphite sheet.

2. The method of claim 1, wherein the plurality of impressions comprise transverse channels passing through the sheet.

3. The method of claim 1, wherein the plurality of impressions comprise indentations in a surface of the flexible graphite sheet.

4. The method of claim 3, wherein the indentations combine to form a passage across a surface of the flexible graphite sheet.

5. The method of claim 1 which further comprises controlling the frequency of oscillations.

6. The method of claim 1 which further comprises controlling the amplitude of oscillations.

7. The method of claim 2 wherein the channel density in the flexible graphite sheet is at least about 210 channels/sq. centimeter.

8. The method of claim 1 which further comprises providing a forming element which comprises a contact element capable of oscillating and contacting the flexible graphite sheet to make impressions therein.

9. The method of claim 8 wherein the contact element has a tapered end.

10. The method of claim 9, wherein the degree of taper of the contact element is greater than about 10° and less than about 90°.

11. The method of claim 9, wherein the contact element has a cross-section shape selected to provide the desired shape or geometry of impressions.

12. The method of claim 8, wherein the forming element further comprises an oscillating element which is in operative contact with the contact element, and is capable of creating oscillations in the contact element.