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(57) **ABSTRACT**

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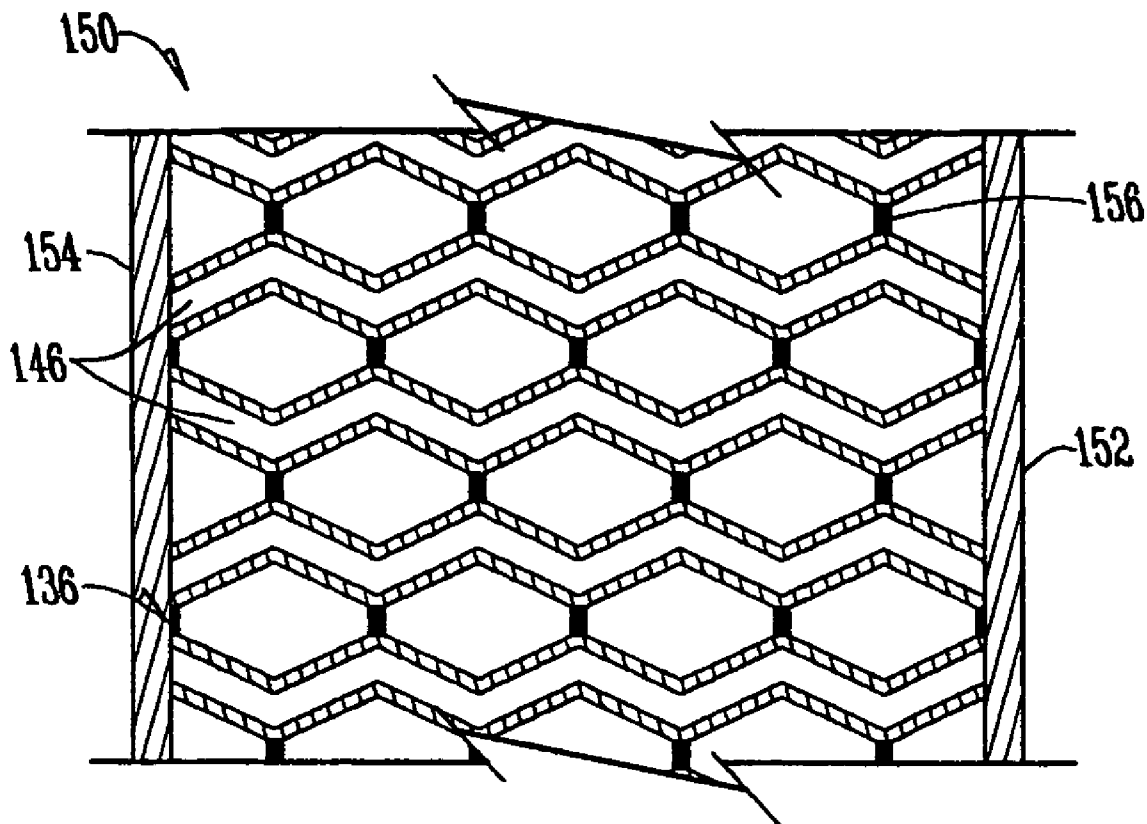
A decomposition unit that comprises a plurality of decomposition cells that comprise two metal plates in parallel spaced relation wherein each plate has a layer of molten dielectric material sprayed thereon by using a plasma spray process. Each decomposition cell also has a high voltage high frequency source attached to a metal plate to cause a discharge in the space created between the layers of dielectric material called a discharge area. The plurality of decomposition cells are then placed adjacent to one another and are connected with an insulating material to form a honeycomb configuration of cells.

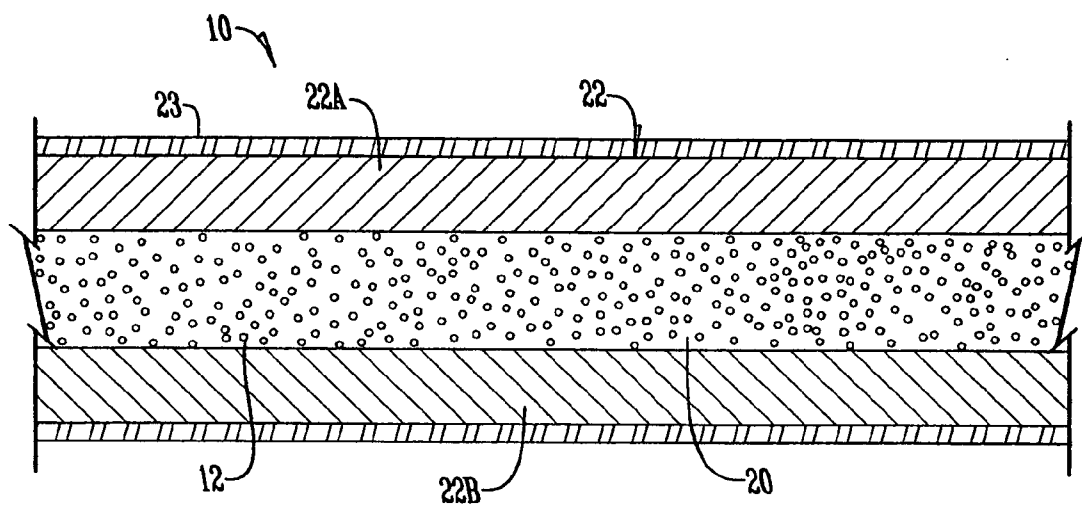
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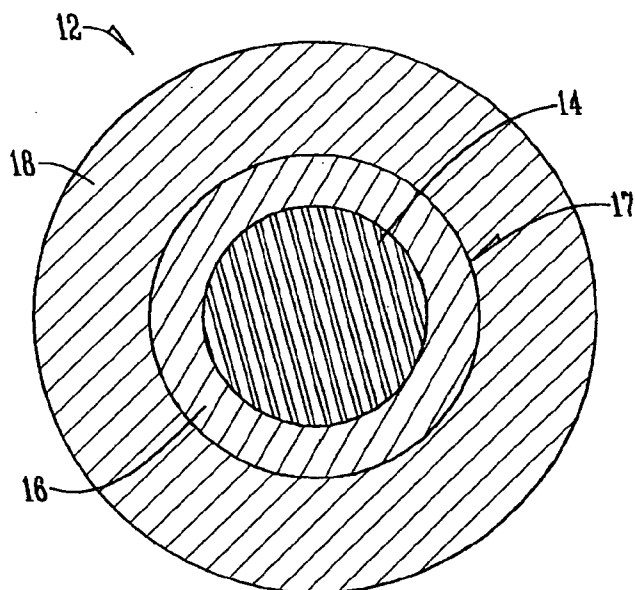
**Related U.S. Application Data**

(60) **Provisional application No. 60/655,972, filed on Feb. 22, 2005.**

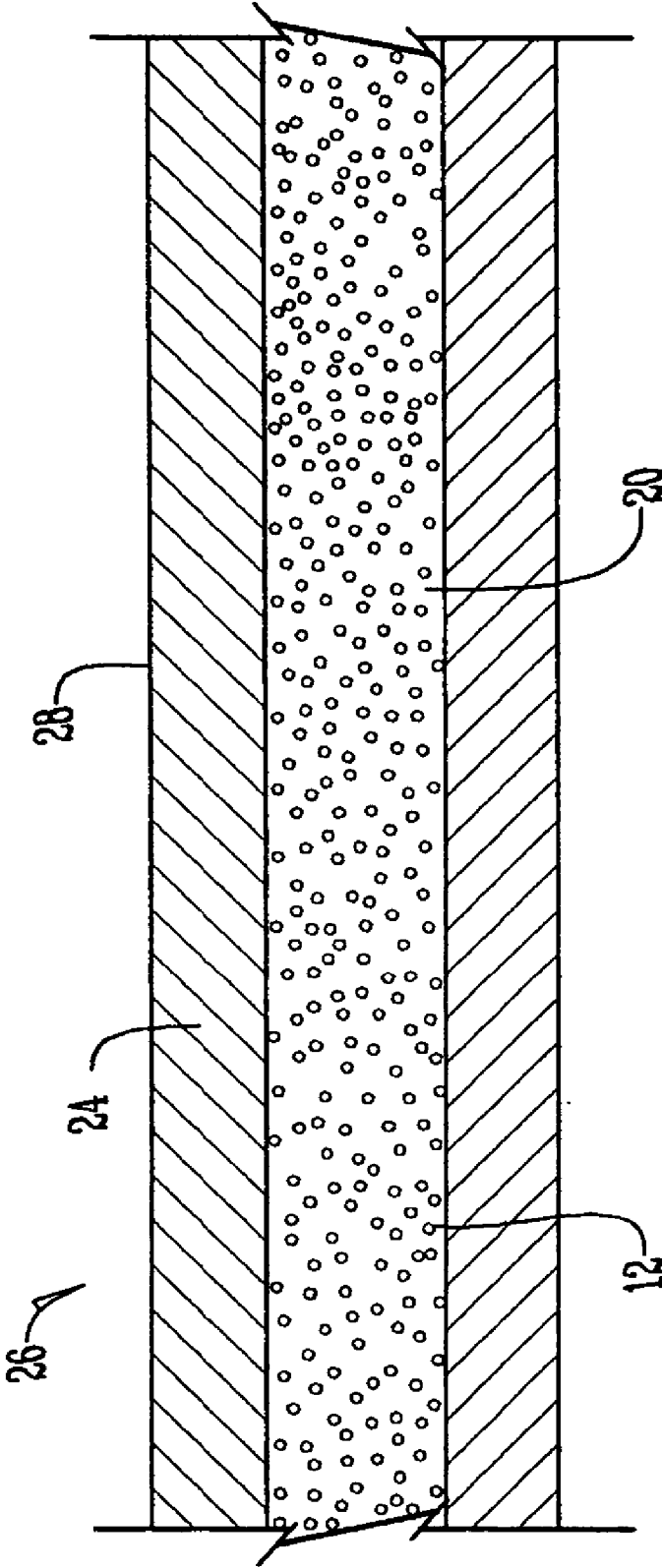




*Fig. 1*



*Fig. 2*



*Fig. 3*

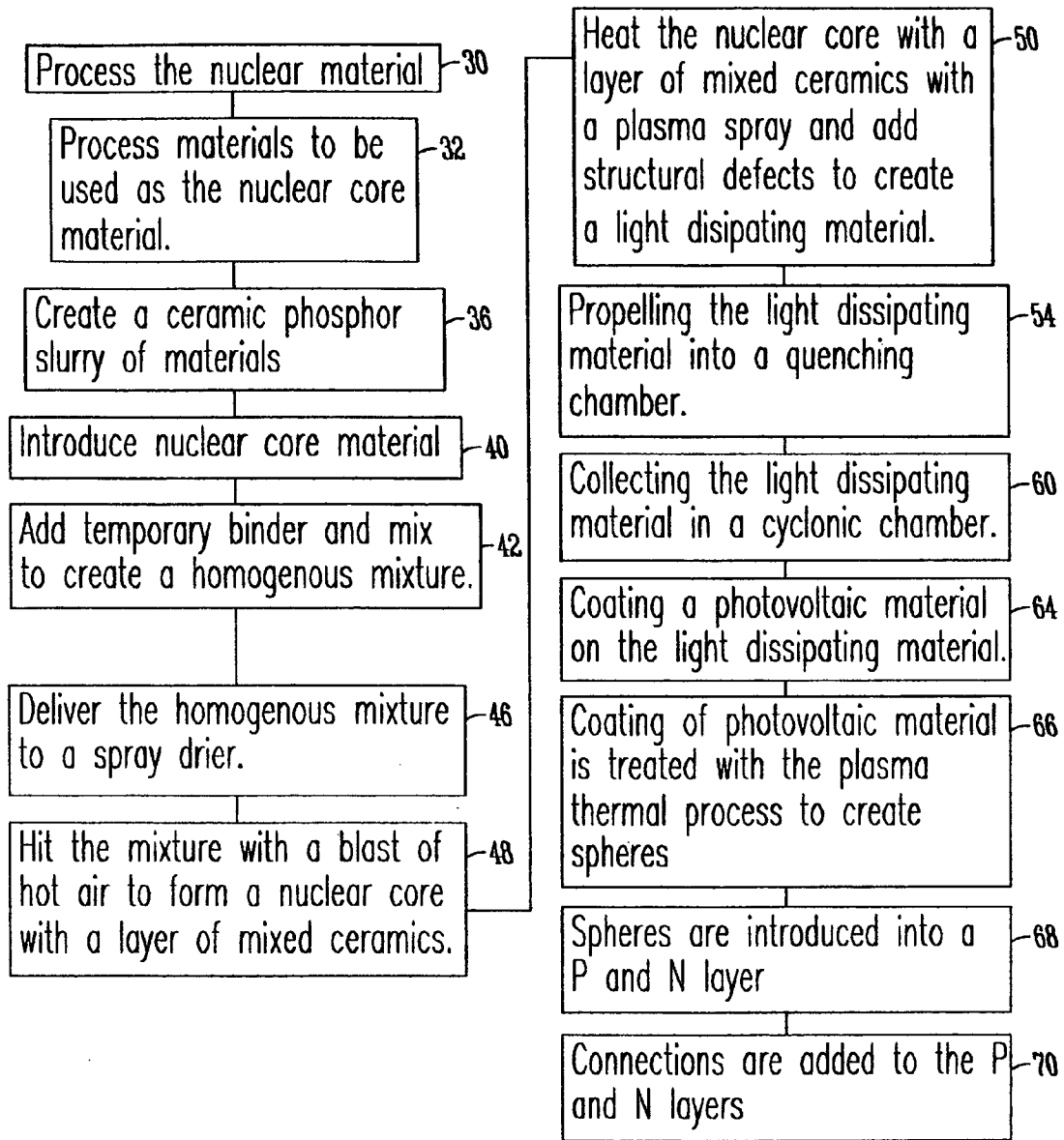
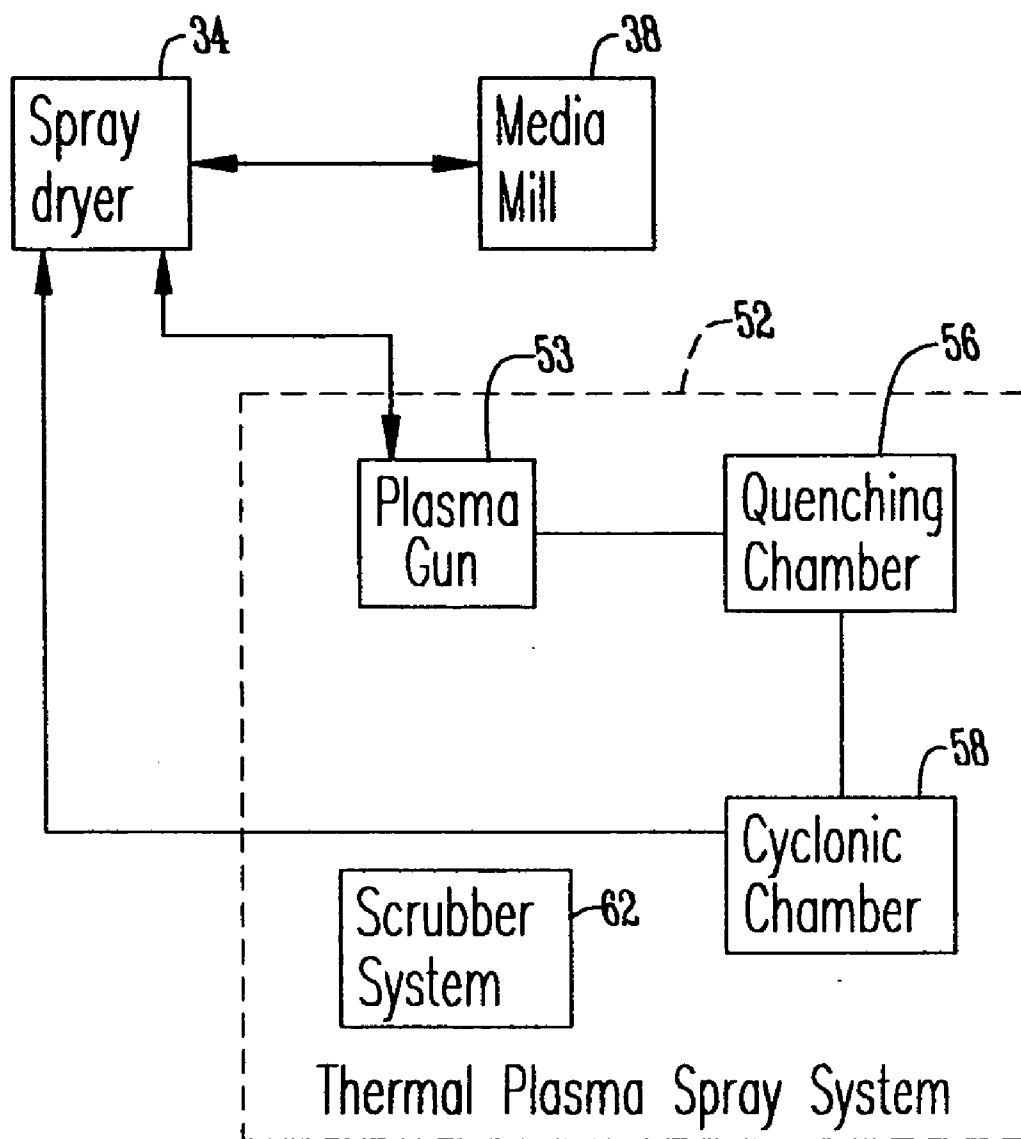
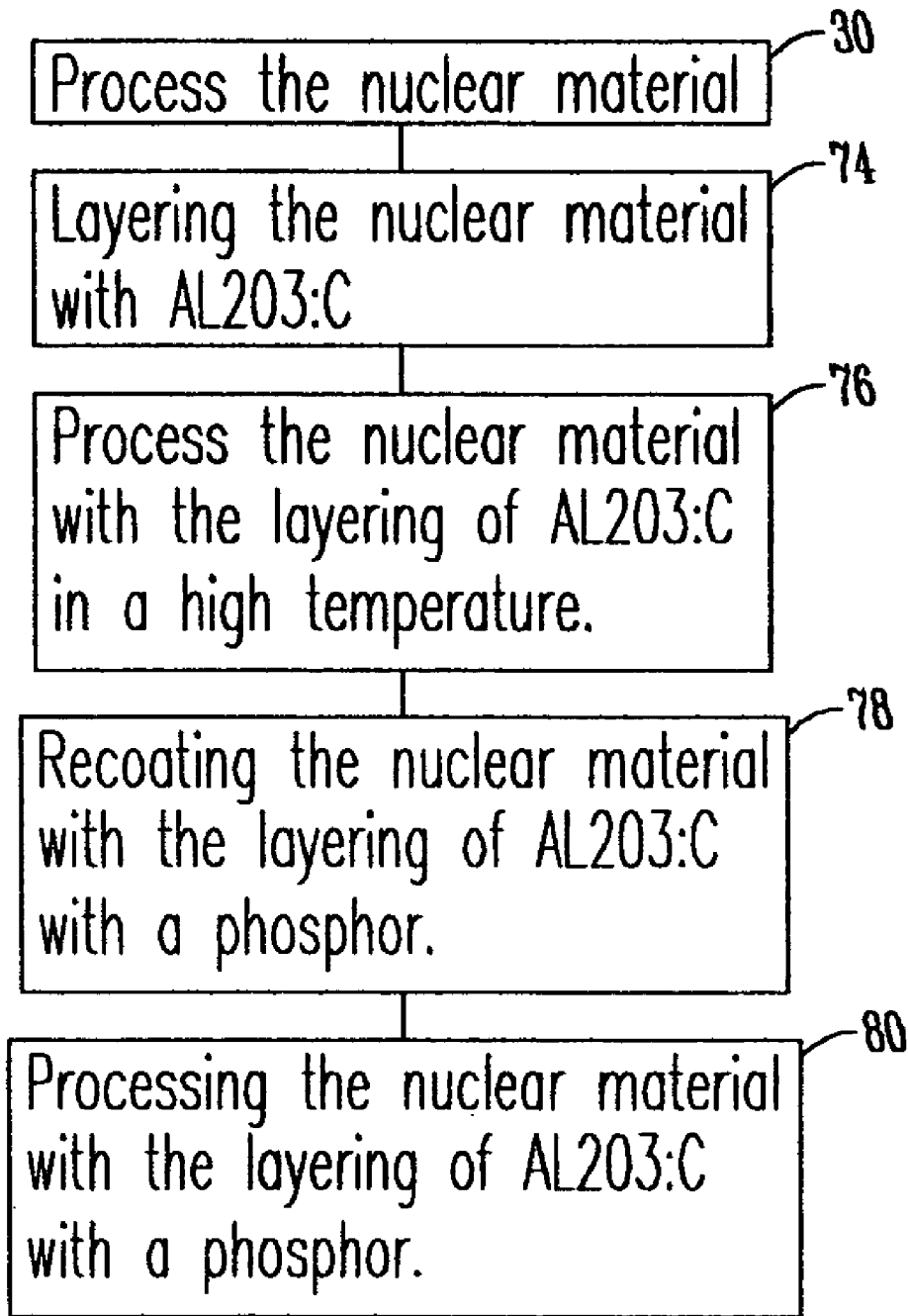


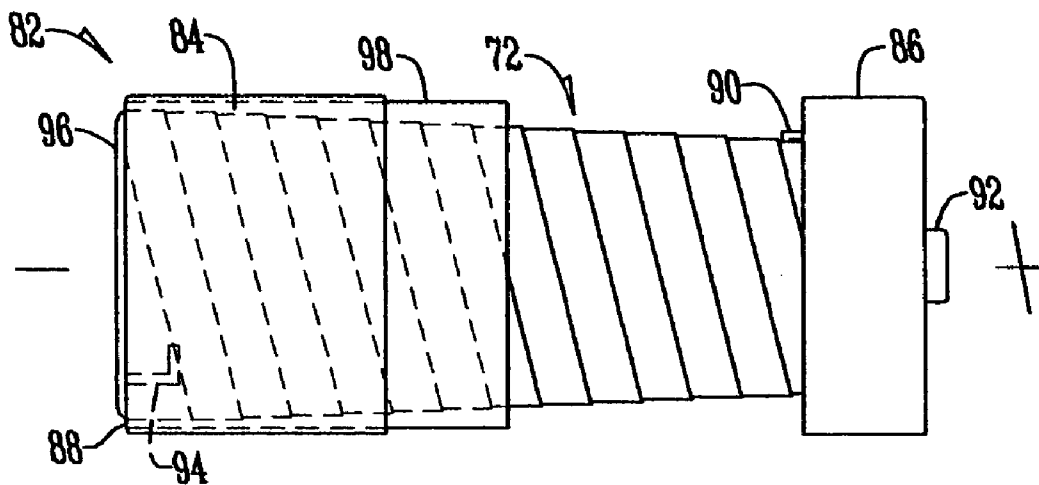
Fig. 4



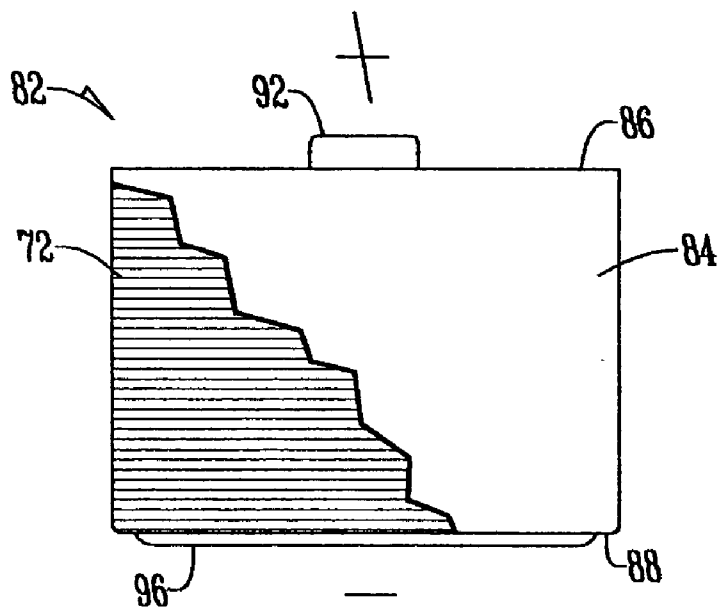
*Fig. 5*



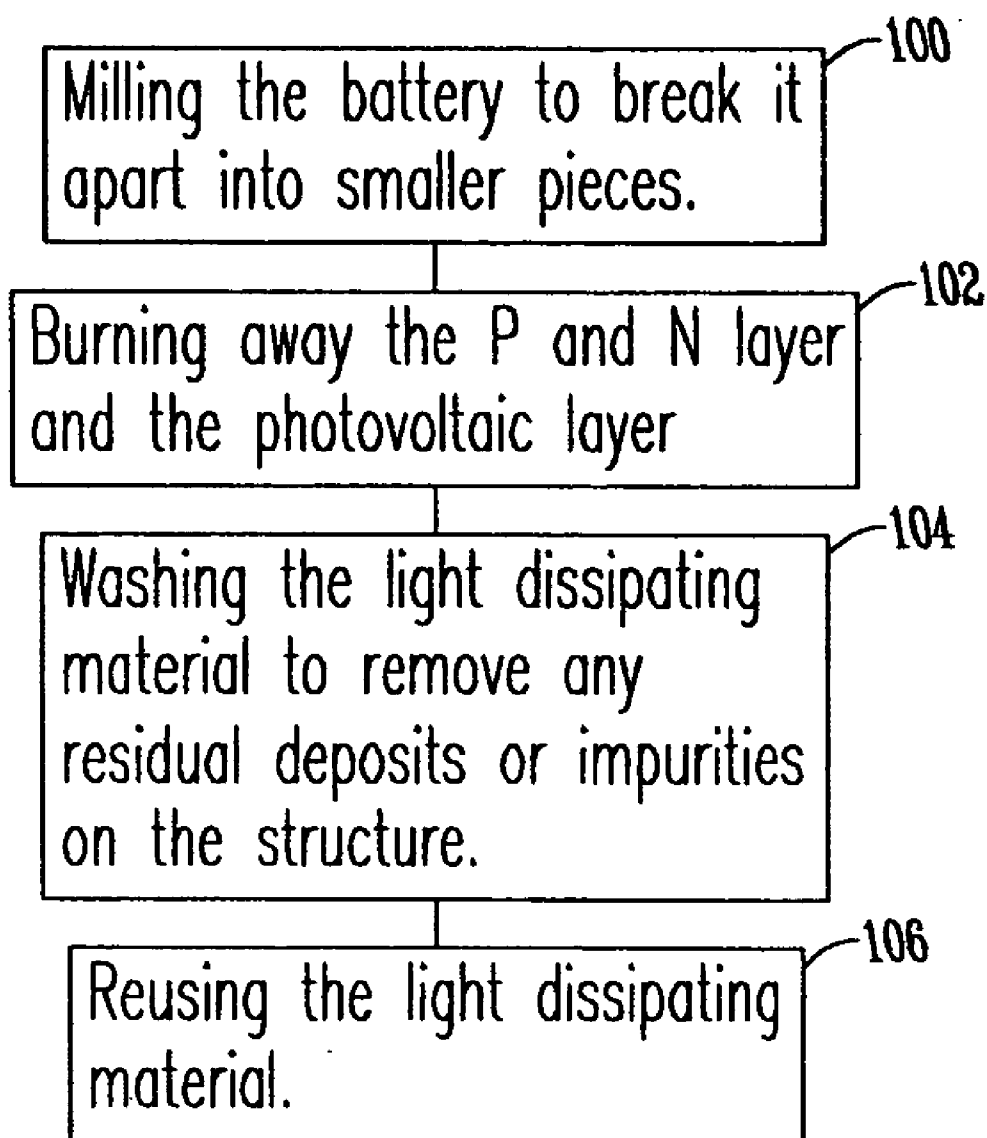
*Fig. 6*



*Fig. 7*

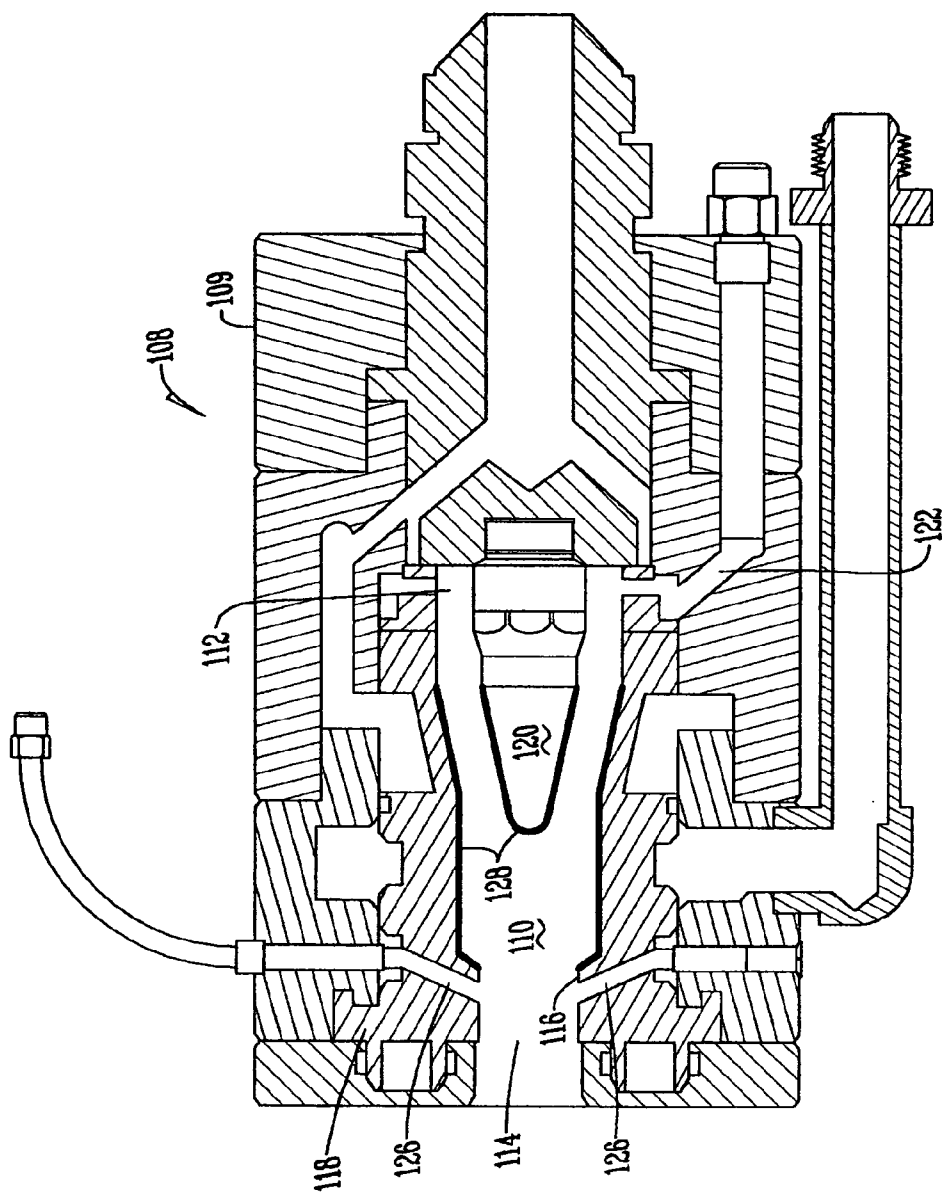


*Fig. 8*



*Fig. 9*





*Fig. 10*

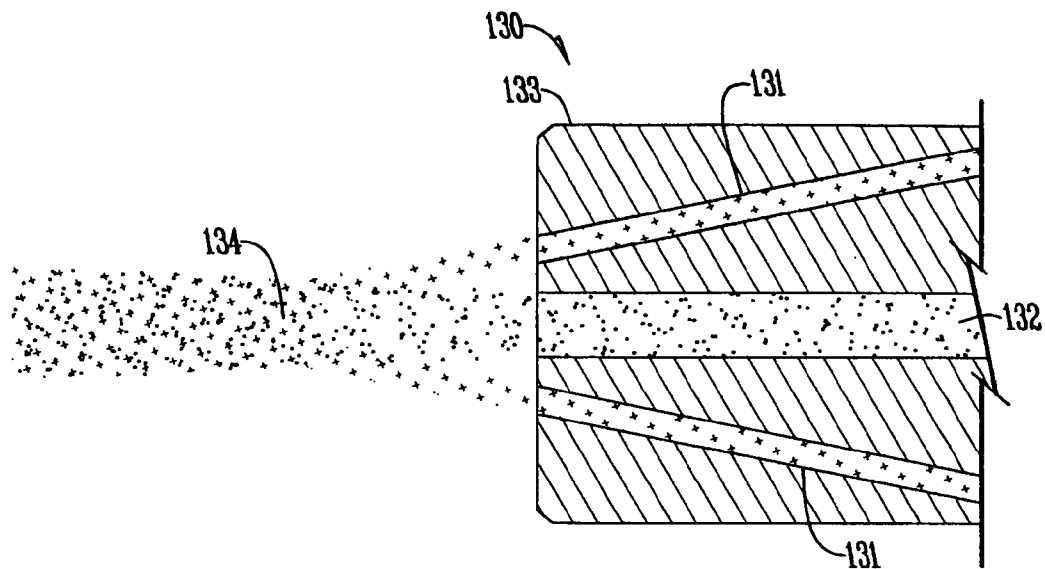


Fig. 11

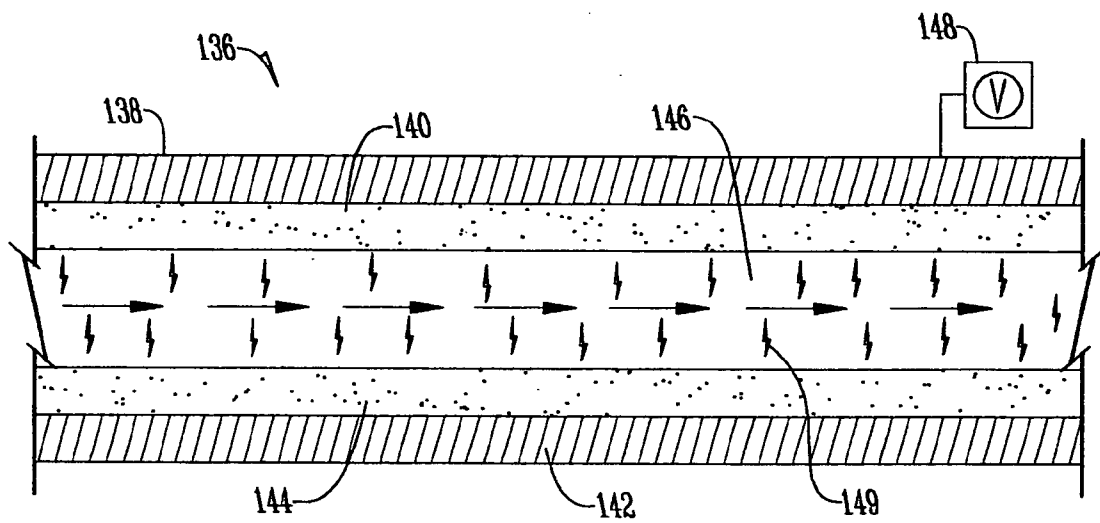
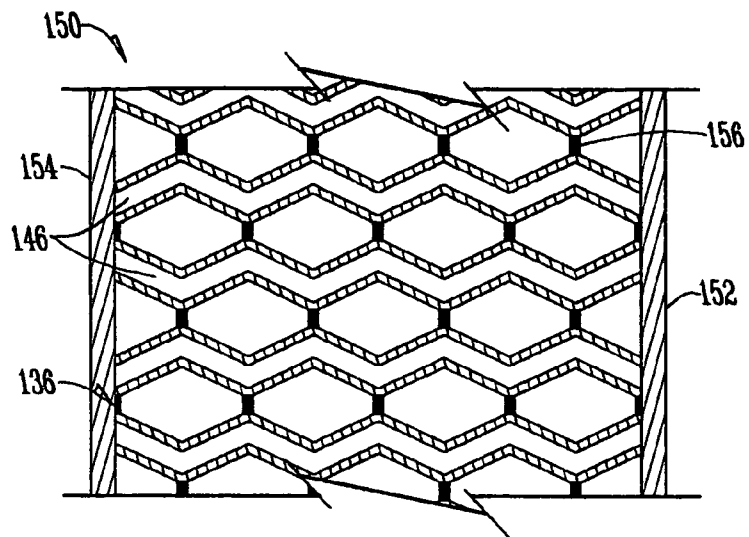
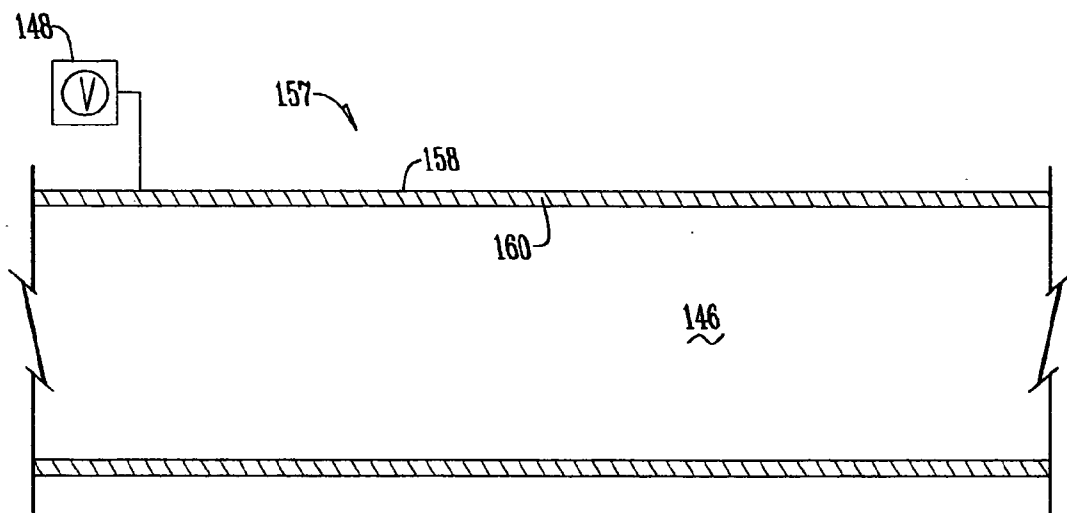


Fig. 12



*Fig. 13*



*Fig. 14*

**DECOMPOSITION UNIT**

**CROSS REFERENCE TO RELATED APPLICATION**

[0001] This application is a non-provisional application gaining priority from provisional patent application Ser. No. 60/655,972 filed Feb. 22, 2005. That provisional is incorporated herein.

**BACKGROUND OF THE INVENTION**

[0002] This application relates to a device that will decompose pollutants. More specifically this invention relates to a decomposition unit for use in factory smoke stacks.

[0003] Currently, in the art of batteries, such as car batteries, a battery has a cell with one plate made of lead and another plate made of leaded dioxide and has a strong sulfuric acid electrolyte in which the plates are immersed. From this chemical reaction within the lead acid battery, electrons flow powering whatever device is connected to the battery. Though current lead acid batteries effectively power devices such as automobiles, many problems in the art remain. First, the life expectancy of an average battery in an automobile can be as little as three to four years. Additionally, current car batteries cause inefficiencies within the car motor thus lowering the miles per gallon of gasoline that a car may travel.

[0004] Batteries having a nuclear core have been developed to attempt to harness the energy from a long lasting source. The radioactive materials of these batteries have been used with chemicals known as phosphors to create light that can be converted into electricity. Though electricity has been created, because of the radioactive nature of the core material, these batteries are unsafe for everyday use.

[0005] Attempts to solve the problem of creating an nuclear-cored battery that is safe for everyday use have been made, however scientist have been unable to find a material that will effectively shield the radioactive radiation of the nuclear core material and yet still produce sufficient light that can be efficiently converted into electricity. Thus, there is a need in the art for an improved nuclear battery.

[0006] High temperature ceramics such as Al<sub>2</sub>O<sub>3</sub>, alumina and zirconium oxide in the past have been used to contain radioactive wastes such that these ceramic containers or sarcophaguses have radioactive waste material placed therein and are buried in the ground. A high temperature ceramic is defined as any ceramic material that has a melting point above 2,000 degrees Centigrade. The ceramic structure is stable and dense enough that this structure is not altered by the radioactive radiation. Nonetheless, high temperature ceramics have never been used in the field of nuclear-cored batteries because the dense structure of the ceramics is not conducive to the production of photons using a radioactive source.

[0007] Additionally, in the current art of manufacturing processes that have been developed to produce similar crystals to those that will be created in manufacturing the nuclear-cored battery are not conducive to the mass production needed to make a profit in the business community. Specifically, during the production of photoluminescent crystals the manufacturing process requires multiple steps of mixing, milling, and heating material continually. These

processes not only take a lot of time and effort, but also produce inferior crystals. Thus, there is a need for a new method of manufacturing crystals that reduces the cost to produce the crystals while increasing the quality of the crystal.

[0008] Furthermore, to assist in the manufacturing process of the nuclear-cored battery the current manufacturing equipment that would be used to manufacture the battery cause inefficiencies during the manufacturing process. Specifically, a problem exists with the nano-material production equipment, such as a plasma spray gun that will be used to manufacture the nuclear-cored battery of this disclosure. A problem with current plasma spray guns exists in that these guns use a tungsten anode and electrode that deplete into the plasma stream as the equipment is used, thus limiting the life of the anode and electrode such that current anode and electrodes within a plasma spray gun only last approximately 250 hours. Thus there is a need in the art to improve upon the life of the anode and electrode with a plasma spray gun.

[0009] Another technology that may be improved uses a similar solution as will be disclosed regarding the nano-material production equipment and this technology is known as a fuel saver. A fuel saver converts O<sub>2</sub> into O<sub>3</sub>. Currently, alumina plates are placed on top of copper plates thus creating the fuel saver and the combination of these plates are used as discharge plates within the fuel saver. Nonetheless, these fuel saver units known in the art do not yield an optimum output potential. Thus, there is a need for an improved manufacturing process to create a fuel saver, and a need for a more efficient fuel saver.

[0010] A fuel saver is one example of a decomposition cell. Decomposition cells are used in manufacturing facilities to be placed in smoke stacks to decompose pollutants into chemicals that are not harmful to the environment. Specifically, there is a need for an improved decomposition cell that will more efficiently decompose pollutants to create waste materials that are safe for the environment.

[0011] Thus, the principal object of the present invention is to use a plurality of decomposition cells to create a decomposition unit that will efficiently decompose pollutants.

[0012] Another object of the present invention is to increase the surface area of a decomposition unit to more efficiently decompose pollutants.

[0013] Another object of the present invention is to use a thermal plasma spray process to improve upon a decomposition cell.

[0014] Yet another object of the present invention is to improve the dielectric quality of a decomposition cell.

[0015] These and other objects, features, or advantages will become apparent from the specification and the claims.

**BRIEF SUMMARY OF THE INVENTION**

[0016] A decomposition unit that comprises a plurality of decomposition cells that comprise two metal plates in parallel spaced relation wherein each plate has a layer of molten dielectric material sprayed thereon by using a plasma spray process. Each decomposition cell also has a high voltage high frequency source attached to a metal plate to cause a

discharge in the space created between the layers of dielectric material called a discharge area. The plurality of decomposition cells are then placed adjacent to one another and are connected with an insulating material to form a honeycomb configuration of cells.

#### BRIEF DESCRIPTION OF THE DRAWINGS

[0017] **FIG. 1** is a sectional view of a nuclear-cored battery;

[0018] **FIG. 2** is a cut away plan view of a sphere of an nuclear-cored battery;

[0019] **FIG. 3** is a sectional view of a super magnet;

[0020] **FIG. 4** is a flow diagram of a manufacturing process of a nuclear-cored battery;

[0021] **FIG. 5** is a schematic diagram of the equipment used during the manufacturing process of a nuclear cored battery;

[0022] **FIG. 6** is a flow diagram of a manufacturing process of a nuclear-cored battery;

[0023] **FIG. 7** is a plan side view of one embodiment of a disposable battery using a layered nuclear-cored battery;

[0024] **FIG. 8** is a cut away plan side view of one embodiment of a disposable battery using a layered nuclear-cored battery;

[0025] **FIG. 9** is a sectional view of a plasma spray gun;

[0026] **FIG. 10** is a flow diagram of a recycling process of an nuclear-cored battery;

[0027] **FIG. 11** is a side plan cut away view of a plasma spray system;

[0028] **FIG. 12** is a sectional view of a decomposition cell;

[0029] **FIG. 13** is a sectional view of a decomposition unit; and

[0030] **FIG. 14** is a sectional view of a fuel saver.

#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

[0031] **FIG. 1** shows an atomic battery or nuclear-cored battery **10**. Nuclear-cored battery **10** is created by producing a plurality of energy sources in the form of spheres **12** (**FIG. 2**) that each have a nuclear core **14** that emits alpha, beta, or gamma radiation. Nuclear core **14** is comprised of any radioactive material including, uranium, uranium carbonate, uranium oxide, strontium, and strontium oxide.

[0032] The nuclear core **14** is surrounded by a ceramic phosphor material **16** that is in one embodiment a crystalline having a carbon defect such that the ceramic phosphor material **16** in combination with the nuclear core forms a light dissipating material **17**. In one embodiment, the ceramic phosphor material comprises a high temperature ceramic. In another embodiment this high temperature ceramic comprises a matrix having  $\text{Al}_2\text{O}_3:\text{C}$ . In yet another embodiment zinc sulfide, or another high temperature ceramic having a carbon defect is used. The ceramic material within the ceramic phosphor material **16** is used to shield and absorb the radiation emitted by the nuclear core **14** while the phosphors are excited by the radioactive radiation of the

nuclear core **14** causing the phosphors to produce energy in the form of photons. In another embodiment lanthides are used as a defect for the phosphors. The carbon defect increases the bandwidth of the ceramic material, and the lanthides are used to increase the bandwidth of the phosphors. Thus, the ceramic material prevents radiation from being emitted past the ceramic phosphor material **16**, yet this material **16** is still able to produce photons.

[0033] In one embodiment the ceramic phosphor material **16** is made into a crystalline (crystal) that is an amorphous crystalline or a structured crystalline and that is manipulated during the manufacturing process so that the photons being emitted by the material **16** are at an optimum wavelength (and thus color) to maximize the efficiency of the nuclear-cored battery **10**. One example of how the crystalline is manipulated is by adding  $\text{MO}_m(\text{Al}_2\text{O}_3):\text{Eu},\text{R}$  to the ceramic phosphor material, wherein M is chosen from one of the alkaline metals such as strontium, calcium, and barium; R is any of the lanthanides; Eu is present at a level from about 0.05% to about 10% by weight and preferably 0.1-5% by weight; and R is present at a level from about 0.05% to about 10% by weight and preferably 0.1-5% by weight. Thus the final formula of the ceramic phosphor material will comprise the matrix  $\text{MO}_m(\text{Al}_2\text{O}_3):\text{C}:\text{Eu},\text{R}$ .

[0034] Another example of a material that is added to the ceramic phosphor material **16** to manipulate the output frequency of the photons being emitted is yttrium oxysulfide doped with titanium and magnesium material that forms a crystal that emits red to orange wavelengths of light. Thus for red and orange wavelengths the ceramic phosphor material comprises the matrix  $\text{MOS}:\text{Mg},\text{Ti},\text{Eu}$  wherein M is chosen from a group consisting of  $\text{MgO}$ ,  $\text{ZnO}$ ,  $\text{ZrO}$ ,  $\text{CuO}$ , Yttrium Oxide, or Gallium Oxide.

[0035] The excitation of the base light emitter, such as  $\text{Al}_2\text{O}_3:\text{C}$ , causes the stimulation of the crystals and the combined frequency gives the final output color. Thus the output frequency of the ceramic phosphor material **16** is manipulated to any color in the visible spectrum. Below is a list of examples of different ceramic phosphors and the color wavelengths of the photons that are emitted by each depending on the amount of each element provided:

[0036] 1. Green—A polycrystalline structure of Alumina incepted with carbon and mixed with a matrix of Europium Oxide, Strontium Carbonate, also Dysprosium Oxide.

[0037] 2. Blue—A polycrystalline structure of Alumina incepted with carbon and mixed with a matrix of Europium Oxide, Strontium Carbonate, also Dysprosium Oxide.

[0038] 3. Yellow—A polycrystalline structure of Alumina incepted with carbon and mixed with a matrix of Europium Oxide, Strontium Carbonate, Barium Carbonate, also Dysprosium Oxide.

[0039] 4. Orange—A polycrystalline structure of Alumina incepted with carbon and mixed with a matrix of Europium Oxide, Strontium Carbonate, also Dysprosium Oxide. Mixed with a polycrystalline structure of Yttrium Oxysulfide and mixed with a matrix of Europium Oxide.

[0040] 5. Red—A polycrystalline structure of Yttrium Oxysulfide and mixed with a matrix of Europium Oxide, also Magnesium Titanium.

[0041] 6. White—A polycrystalline structure of Alumina incepted with carbon and mixed with a matrix of Europium Oxide, Strontium Carbonate, Neodymium Oxide, also Dysprosium Oxide.

[0042] 7. Violet—A polycrystalline structure mixed with a matrix of Europium Oxide, Calcium carbonate, also Neodymium Oxide.

[0043] Thus, each combination listed creates a separate crystalline structure depending upon the content of each element present. Each crystalline separately is unique in its interaction with different radiations produced by the nuclear core 14, and each will produce a different wavelength of visible light emitted from the crystalline.

[0044] Surrounding the ceramic phosphor material 16 is a photovoltaic layer 18 that transforms the photons into a flow of electrons to create an energy source, or sphere 12. One will appreciate that in one embodiment the photovoltaic layer 18 is made of an amorphous silicon that also is altered with defects by, for example, doping the material with magnesium in order to manipulate a stimulating frequency of the photovoltaic layer 18. Other examples of defects include titanium and chromium. Thus the output frequency of the photons generated by the ceramic phosphor material 16 is manipulated or tuned while manipulating or tuning the stimulating frequency of the photovoltaic layer 18 so that the most efficient amount of light created by the ceramic phosphor material 16 is converted into an electron flow by the photovoltaic layer 18.

[0045] After a plurality of spheres 12 are created the battery is formed by surrounding a plurality of spheres 12 with a conductive material 20 that is an intermediate layer that carries the spheres 12. This conductive material 20 comes into direct contact with the spheres 12 and in one embodiment is a conductive polymer, one example of which is a sulfidized polymer. One such conductive polymer is poly(3,4-ethylenedioxythiophene) polystyrenesulfonate. A P and N layer 22 comprising a P layer 22a and an N layer 22b sandwiches the spheres therebetween to harness the electron flow created by the photovoltaic layer 18 to create the nuclear-cored battery 10. Additionally, a layer of insulating material 23 can be used to surround the P and N layer 22.

[0046] Finally, spheres 12 in one embodiment are in powder form and will range in size from 50 microns to sub micron in size depending on the application and output. Nonetheless, in another embodiment a metal is added to the nuclear core 14 of the battery 10 in order to increase the size of the spheres 12 for macro-sized applications.

[0047] As shown in FIG. 3, in an alternative embodiment a magnetic material 24 is placed around the plurality of spheres 12 to create a super magnet 26. Specifically, the flow of electrons created by the photovoltaic layer 18 interacts with the magnetic material 24 to magnetize the outer surface 28 of the magnetic material 24.

[0048] In operation, the nuclear core 14 emits radiation, for example, beta radiation that is an electron. When the electron comes in contact with the ceramic phosphor material 16 the radioactive radiation is stopped by the ceramic, yet the electron excites the phosphors causing an electron to “jump” from a 4d valence energy level to a higher valence energy level within a phosphor. When that electron “settles” back to its original 4d state, energy in the form of a photon

is emitted. When the ceramic phosphor material 16 includes a carbon defect in its matrix the carbon defect increases the bandwidth of the phosphor allowing more photons to be generated. Furthermore, the matrix of the ceramic phosphor material 16 will determine the frequency of the photon that is being emitted from the ceramic phosphor material 16. These photons are then absorbed by the photovoltaic layer 18 to create an electron flow that is harnessed by the P and N layer 22 to cause the battery 10 to function.

[0049] When other radioactive radiations are present such as gamma and alpha radiation, the phosphors still become “excited” and produce photons, but not in the same way as beta radiation. Thus all types of radioactive material may be used as the nuclear core 14.

[0050] Manufacturing the nuclear-cored battery involves a multi-step process. FIG. 4 shows a flow chart of the multi step process used during the manufacturing of the battery and FIG. 5 shows a schematic diagram of the equipment used during this process. Processing the nuclear material is the first step 30. This is preformed in a multitude of ways depending on the initial source. If the nuclear material is of a mixed matrix of different radiation sources division is made by dissolving the materials with the mixed matrix and separating these materials via gravimetric. The weight and density of the different materials in the mixed matrix causes these materials to separate into layers making it possible to divide materials as needed.

[0051] The next step 32 is to process the nuclear material via a spray dryer 34 into a spherical metal or compound to be used as the nuclear core 14. The core 14 in one embodiment is a compound that is an oxide or carbonate that creates a stronger structure, with a higher melting point than the metal that the oxide or carbonate are derived from. Other methods and equipment such as a precipitation method or some other form of sprayer is also used to create the nuclear core 14.

[0052] At step 36 a ceramic phosphor slurry material that in one embodiment is made of a base material of Al<sub>2</sub>O<sub>3</sub>:C and phosphors is created. The ceramic phosphor slurry is a frequency alternating mixture that in one embodiment comprises strontium carbonate, europium oxide, dysprosium oxide, or the like (depending on the output frequency desired) that is mixed into a water and alumina powder. The ceramic phosphor slurry, water and alumina powder are milled together to a nano mean size to form a ceramic phosphor slurry material. This is preformed in a media mill or mixing chamber 38 or other systems. In one embodiment a carbon defect is added to the ceramic phosphor slurry material by using graphite, or another carbon additive while making the ceramic phosphor slurry.

[0053] At step 40 the nuclear core material is introduced to the ceramic phosphor slurry material and then at step 42 a temporary binder is added (ammonia nitrate or another gas is used to create a porous structure as necessary) and the ceramic phosphor slurry material with a nuclear core material having a temporary binder is then mixed to create a homogenous mixture. Examples of the temporary binding material are methal cellulose, poly vinyl alcohol, or the like.

[0054] This homogenous mixture is processed again through the spray drier 34 to dry the homogenous mixture to form an outer shell at step 46. At step 46 the homogenous

mixture is delivered to the spray drier 34 and into the cavity of the spray drier 34 by double annulus spray nozzle or discharge wheel. The homogenous mixture is then hit with a blast of hot air that evaporates the water within the homogenous mixture and dries the temporary binding material at step 48 to form a temporarily bound layer of mixed ceramics. Atmospheric gas of nitrogen, argon, and/or carbon dioxide is used to assist in the process. In the embodiment wherein the base ceramic is Al<sub>2</sub>O<sub>3</sub>:C step 48 forms a semi ridged spherical particle with the nuclear core 14 surrounded by a temporarily bound layer of mixed ceramics having an alumina structure.

[0055] At step 50 the particle created at step 48 is subjected to a high temperature portion of the processing, or plasma thermal process using a thermal plasma spray system 52 having at least one plasma gun 53. With temperatures that are adjusted from 2,000 to over 15,000 centigrade the mixed ceramics from step 48 are brought to a molten state for a short amount of time, preferably under a minute, thus creating a layer of mixed ceramics in a molten state. While in this molten state the temporary binding material will burn out and the nuclear core with the layer of mixed ceramics becomes an amorphous structure as a result. The plasma stream sinters the layer of mixed ceramics to densify and calcinate or purify the layer. In one embodiment the layer of mixed ceramics has an alumina structure and this alumina structure is brought to a molten state for a short amount of time creating the amorphous structure.

[0056] While the particle created at step 48 is subjected to the high temperature portion of the processing structural defects are introduced. In one embodiment these defects include carbon defects and/or lanthide defects. Once a carbon defect is added the layer of mixed ceramics in combination with the nuclear core 14 becomes a light dissipating material 17. Thus, the nuclear core 14 after this high temperature processing will no longer be radioactive in nature past the layer of mixed ceramics. The radioactive decay will be transformed into light that is emitted out from the light dissipating material 17.

[0057] During step 54 the light dissipating material 17 is propelled into a quenching chamber 56 and a pair of cooling nozzles that in one embodiment emit a crosscurrent of quenching gas that is an air and gas mixture cools and incepts further amounts of carbon into the light dissipating material 17 to form a crystalline. One will also appreciate that the temporary binder provided some carbon content as it burned out but the use of carbon dioxide in the quenching gas will allow for total coverage of carbon within the light dissipating material 17. Also the use of nitrogen, or other inert gas, as a quenching gas will encourage the clarity of the crystalline allowing for a higher transfer of light from the light dissipating material 17. Rather than add the carbon defect in step 52, alternatively the carbon defect is added just in step 54. The heating of the nuclear core with a layer of mixed ceramics allows the introduction of the carbon defect at step 52, step 54, or both. The light dissipating material 17 now quenched and treated with the chamber gasses is collected by a cyclonic chamber 58 that is separate from the quenching chamber 56 at step 60. The light dissipating material 17 is then removed when collected.

[0058] Construction material in the quenching chamber 56 will be similar to that of the spray drier 34. Additionally, a

scrubber system 62 is utilized to prevent the discharge of uncoated nuclear core particles in both the spray dry process and thermal plasma spray stages.

[0059] Once the light dissipating material 17 is created the material is spray dried with a coating of photovoltaic material such as silicon by the spray dryer 34 at step 64. At step 66 this layer is treated again with the thermal plasma process to densify the silicon on the light dissipating material 17 to create the photovoltaic layer 18, thus creating the sphere 12. By using the thermal plasma process the photovoltaic layer in one embodiment has an amorphous structure. This layering technique will allow for a high strength and small particle size with each layer interacting with the next. The spray dryer 34 gives the spheres 12 their shape and one will understand that these small spherical particles are in one embodiment the form of a powder.

[0060] At step 68 the finished powder is sandwiched between organic P and N layers 22 to draw away the electrons being discharged from the photovoltaic layer 18 of the spheres 12. Leads are connected to the P and N layers 22 to transport energy to a source consumer of the electricity at step 70. The P and N layers 22 in one embodiment are applied as a spray and conform to any shape desired or as a sheet 72 (FIGS. 7 and 8) that is later inserted into a commercial product.

[0061] One will appreciate that though this method of manufacturing places a ceramic phosphor layer 16 over a nuclear core 14 to form a light dissipating material 17, that in another embodiment only the ceramic phosphor slurry undergoes the manufacturing process described to create a ceramic phosphor crystalline 16. This crystalline 16 is then used in association with the nuclear core 14 to create a light dissipating material 17.

[0062] In another embodiment seen in FIG. 6 the nuclear material is layered with the Al<sub>2</sub>O<sub>3</sub>:C first in step 74, then processed in high temperature in step 76, then recoated with the phosphor in step 78, and processed with higher temperatures to alter the output frequency at step 80. The mixing of a matrix of materials is used on low to mid output radioactive materials but high output materials will require the shielding first then the altering of the frequency. This also offers an opportunity to manipulate a carbon inception of an alumina layer. The use of pre-manufactured materials exists to create these layers. Again, a binder is used to hold the layers together temporarily until high temperature processing is implicated.

[0063] The use of a discharge circuit in one embodiment is utilized to remove unused excess electricity created by the nuclear-cored battery 10. This electricity is converted to heat or other forms of energy to dissipate excess capacity. This energy could also be redirected to a capacitor to store the electricity during sporadic and inconsistent use of the source. The reason for the use of this circuit is that the battery 10 is going to give electricity continuously without delay for the duration of the core materials half-lives.

[0064] FIGS. 7 and 8 show embodiments wherein a layered battery 82 is formed. Specifically, in this embodiment the P and N layers are applied as a spray to form a nuclear core energy source sheet 72. In this embodiment the sheet 72 is rolled or coiled into a cylinder and inserted into a plastic or metal housing or case 84 having a first and

second ends **86** and **88**. **FIG. 7** shows the coiled sheet **72** outside the case **84** and tapered; however, in use the sheet is coiled and within the case **84**. A first conductive lead **90** is electrically connected to the P layer **22a** and is attached to the first end **86** to create an anode **92** and similarly a second conductive lead **94** is electrically connected to the N layer **22b** and attached to the second end **88** to create an electrode **96**. One will understand that a layer of insulating material **98** may be attached to the case **84** to insulate the case **84** from the sheet **72**.

[0065] As shown in **FIG. 8**, in another embodiment a plurality of sheets **72** are stacked upon or are adjacent to each other within the case **84**. In this embodiment the first end **86** of the case will come into contact with a P layer **22a** of a sheet **80** to form the anode **92** and the second end **88** will come into contact with a N layer **22b** of another sheet **72** to form the electrode **96**. In this embodiment, if an insulating layer is desired, conductive leads are used to connect the P layer **22a** to the first end **86** of case **84** and to connect the N layer **22b** to the second end **88** of case **84**.

[0066] Other products that can be produced from this source of energy are: room temperature super conductors, super conducting cables/wires, resistance free polymers, infinitely formable power supplies, energy sources for: electronics, houses, cities, countries, automobiles and other forms of transportation.

[0067] When in use the product life, whether a battery, or another product using the energy source disclosed above, is determined by the material(s) of the nuclear core **14**. Therefore, a manufacturer by selecting the nuclear core material has the ability to pre-select a time limit that a product will function. This is accomplished by first testing nuclear materials by carbon dating or the like to determine a half life for the materials to provide nuclear materials having known half lives. Then a nuclear material having a known half life is selected and used as a nuclear core **14** of a nuclear-cored battery **10**. Thus, once this nuclear core ceases to produce effective radioactive radiation the product will shut down.

[0068] Another way of pre-selecting the time limit of a product that is produced from the above energy source is to attach a timing mechanism such as a timing circuit to the product that will terminate the operation of the product after a pre-selected occurrence. In one embodiment the timing mechanism is programmed to disable a product after a pre-selected period of time such as for example 10 years. In an alternative embodiment the timing mechanism disables a vehicle after a pre-selected amount of distance traveled by the vehicle. For example the timing mechanism could sense when a vehicle has driven 50,000 miles and disable the vehicle at that time.

[0069] The reason for pre-selecting the life of a product using the energy source **12** is because when a radioactive core material is used, this energy source can have the potential of lasting for trillions of years. Thus, without pre-selecting the time of the life of a product, consumers will have no need to repurchase a product. Furthermore, many devices such as DVDs, personal electronics, and others that could use the energy source **12** involve technologies that are continually being improved. Thus, products having a pre-selected life will allow for the miniaturization of many electronics and the development of new technologies to ensure products remain up to date. Thus to ensure technol-

ogy will continue to move forward, the products using the nuclear-cored battery energy source **12** will need to have a pre-selected product life.

[0070] In an embodiment wherein a product uses a timing circuit to pre-select the time of the life of a nuclear cored product, this product will need to be recycled. The steps for recycling a nuclear-cored battery are shown in **FIG. 9**. Recycling of the nuclear-cored battery can be accomplished by first milling the battery to break it apart into smaller pieces, as represented in step **100**. Then the pieces undergo a thermal burn, such as in a kiln to melt way the P and N layer and the photovoltaic layer as shown in step **102**. Remaining after the thermal burn is the light dissipating material **17** that is either chemically treated with an acid to etch the ceramic within the light dissipating material **17** or physically treated with a circulating wash to remove any residual deposits or impurities on the light dissipating material such as excess carbon, as shown in step **104**. Thus, the light dissipating material **17** may then be reused in another application as shown in step **106**.

[0071] **FIG. 10** shows an improved thermal plasma spray gun **108** that is one example of one embodiment of thermal plasma spray gun **53** used during the manufacturing of the nuclear-cored battery **10**. The plasma spray gun **108** has a housing **109** with a plasma stream conduit **110** that extends from an inlet end **112** to a discharge end **114** having a discharge aperture **116**. Within the plasma stream conduit **110** is a discharge dielectric anode **118** and a discharge dielectric electrode **120**. In communication with the plasma stream conduit **110** are gas feed conduits **122** that extend through the housing **109** of the plasma spray gun **108** such that a single gas, or mixture of gasses, is exposed to the anode **118** and electrode **120** within the plasma stream conduit **110** to create a stream of plasma therein. Supply conduits **126** extend through the plasma spray gun **108** and are in communication with the discharge end **114** of the plasma stream conduit **110** to supply powdered metals or ceramics to the plasma stream to create a molten material.

[0072] A voltage supply is electrically connected to the plasma stream conduit **110** to supply voltage to the conduit **110** to create an electrostatic discharge that will convert feed gases into a plasma stream. This voltage supply may be integrated as a circuit that is part of the plasma gun **108** or may be a voltage supply that is remotely located from the plasma gun **108**.

[0073] The use of hydrogen, nitrogen, helium, and/or argon is used to produce the plasma stream. A hydrogen nitrogen combination will generate sufficient heat with the ability not to interact with the structure and alter the nuclear core with a layer of mixed ceramics introduced to the plasma spray gun **108**. A high-energy electrostatic discharge through the gas causes the plasma phase of the gas to be generated. The gas is then ejected from the plasma stream conduit **110** of the plasma spray gun **108**, and metallic or ceramic powders are introduced into the stream via the supply conduit **126** where the heat is transferred to the powders.

[0074] The anode **118** and electrode **120** create an electrostatic discharge causing the formation of the plasma gas. During this electrostatic discharge high amounts of energy cause a pitting of the surfaces of the anode **118** and electrode **120**. To solve this problem the anode **118** and electrode **120** are milled to remove 2-20 mills and a dielectric material



such as alumina is deposited onto the anode **118** and electrode **120** surfaces to create a dielectric barrier **128**, preventing the pitting from the discharge of the static field, thus increasing the efficiencies of the unit and allowing for a higher purity in the end product. This dielectric material may be applied to the anode **118** and electrode surfaces using a thermal plasma process to spray molten dielectric material onto the anode **118** and electrode **120**. Furthermore, in one embodiment the dielectric material may be doped with another material, such as for example, magnesium.

[0075] FIG. 11 shows a reconfigured plasma spray system **130** that is an example of one of the embodiments of plasma spray system **60** that is used during the creation of the nuclear-cored battery. Specifically, this embodiment shows a reconfigured plasma system **130** that will more efficiently handle a liquid stream of material thus creating a wider spray area. By using this configuration the spray dry process may be eliminated from the processing.

[0076] Specifically, the plasma spray system **130** of FIG. 11 shows a plurality of smaller plasma jets **131** that are configured in a 3-12 inch diameter around a centrally located jet **132** that is a liquid generating device within a housing **133**. The jets **131** are one embodiment of the spray gun **108** shown in FIG. 10 that generate a plasma stream by utilizing an anode **118** and electrode **120** in combination with gases from gas feed conduits **122** within a plasma stream conduit **110** having an inlet end **112** and discharge end **114**. Similarly, the centrally located jet **132** is also one embodiment of the spray gun **108** wherein a material powder, such as metallic powder is fed into the discharge end **114** of the plasma stream conduit **110** via supply conduit **126** so that molten, or liquid metal is discharged by the liquid generating device **132**. In one embodiment the centrally located jet **132** has a double annulus spray head that generates a stream of atomized liquid.

[0077] The plurality of plasma spray jets **131** are positioned so that their plasma streams will intersect at a point **134** along the path of the atomized liquid and thus become part of the atomized liquid stream. With this system in place a smaller particle is produced and fewer steps are required to produce the same product.

[0078] FIG. 12 shows a decomposition cell **136** that functions to cause the decomposition and production of materials in a discharge field. The decomposition cells **136** are placed in a conduit in order to break pollution down into its simplest components to minimize pollution. For example, in one embodiment, a decomposition cell **136** is placed in a smoke stack of a manufacturing facility to convert pollutants into environmental safe oxygen or carbon.

[0079] The decomposition cell **136** of FIG. 12 is manufactured by taking a first metal plate **138** such as copper and using a plasma gun **53** or **108** to spray a molten dielectric material such as alumina onto the metal plate **138** to create a first dielectric layer **140**. One will appreciate that by using the plasma spray gun **53** or **108** to spray the molten dielectric material on the metal plate **138**, an optimum contact area between the dielectric layer **140** and the plate **138** is achieved to create a more efficient decomposition cell **136**. Furthermore, by using the plasma spray gun **53** or **108**, magnesium oxide may be doped into the molten dielectric material such as alumina to further increase the efficiency of the decomposition cell **136**. Similarly, the molten dielectric

material is then sprayed onto a second metal plate **142** to create a second dielectric layer **144**.

[0080] Next the plates **138**, **142** are placed in parallel spaced relation to create a discharge area **146** wherein air is able to flow through the cell **136**. Thus during a discharge process, when the two metal plates **138**, **142** are electrically connected to a high voltage high frequency source **148** and when voltage is supplied to the two metal plates **138**, **142** an electrostatic discharge **149** occurs in the discharge area **146**, thus decomposing pollutants flowing therethrough and filtering the air. One will appreciate that the high voltage high frequency source **148** may be supplied by a circuit that is part of the decomposition cell **136** or a by a voltage source remote to the cell **136**.

[0081] As best shown in FIG. 13, a plurality of decomposition cells **136** are placed together to form a decomposition unit **150**. The decomposition unit **150** of FIG. 13 has a honeycomb configuration of decomposition cells **136** that extend between first and second side walls **152**, **154** thus creating a plurality of discharge areas **146** that can be separated by an insulating material **156**. Because of the plurality of discharge areas **146** in combination with the honeycomb configuration of the unit **150** the surface area of the discharge area **146** within the decomposition unit **150** increases, thus causing more pollutants to be decomposed as the pollutants flow through the decomposition unit **150**.

[0082] In one embodiment shown in FIG. 14, the decomposition cell **136** is used primarily to convert  $O_2$  to  $O_3$ . In this embodiment the decomposition cell **136** is referred to as a fuel saver **157**. The fuel saver **157** specifically is created by taking a rolled copper body **158** and thermally applying an alumina matrix **160** thereto. One will understand that by thermally applying the alumina to the rolled copper the surface area between the copper and alumina is increased while minimizing the gap between the copper and alumina matrix. Additionally, the alumina is doped with magnesium oxide to change the oxygen state in the final product from an  $O_3$  thus yielding a higher output during the discharge process. Because  $O_3$  has more chemical bonds than  $O_2$ ,  $O_3$  burns much more intensely than  $O_2$ . Thus a fuel saver is used in an engine to convert  $O_2$  to  $O_3$  within the engine to provide an improved fuel system that creates optimum gas mileage for the engine.

[0083] It will be appreciated by those skilled in the art that other various modifications could be made to the device without the parting from the spirit in scope of this invention. All such modifications and changes fall within the scope of the claims and are intended to be covered thereby.

What is claimed is:

1. A decomposition unit comprising:

a plurality of decomposition cells secured between a pair of sidewalls to form a honey comb configuration; and

the decomposition cells having a first metal plate with a first layer of dielectric material in spaced relation to a second metal plate with a second layer of dielectric material.

2. The decomposition unit of claim 1 wherein insulating material is disposed between each decomposition cell.

3. The decomposition unit of claim 1 wherein the first layer of dielectric material is applied by spraying a molten dielectric material onto the first metal plate.

4. The decomposition unit of claim 1 wherein the second layer of dielectric material is applied by spraying a molten dielectric material onto the second metal plate.

5. The decomposition unit of claim 1 wherein the dielectric material is doped with magnesium.

6. The decomposition unit of claim 1 wherein the decomposition cells further have a high voltage source electrically connected to the first metal plate.

7. A decomposition unit comprising:

a plurality of decomposition cells secured between a pair of sidewalls to form a honey comb configuration;

the decomposition cells having a first metal plate with a first layer of dielectric material in spaced relation to a second metal plate with a second layer of dielectric material; and

wherein the first layer of dielectric material is applied by spraying a molten dielectric material onto the first metal plate.

8. The decomposition unit of claim 7 wherein the second layer of dielectric material is applied by spraying a molten dielectric material onto the second metal plate.

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