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(54) **THERMAL-ENERGY PRODUCING SYSTEM AND METHOD**

(52) **U.S. Cl.**
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(57) **ABSTRACT**

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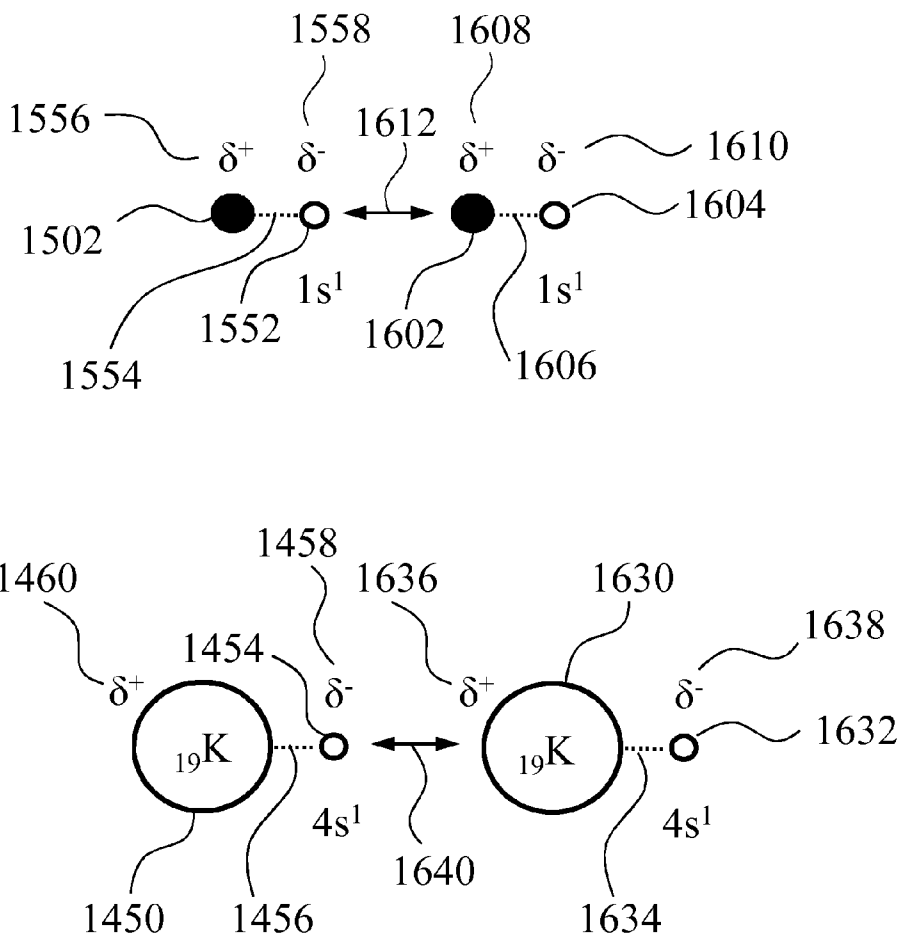
System and method for producing thermal energy is based on a very large number of nanoscale particle accelerators in a volume accelerating electrons and hydrogen ions at very high local electric fields. Nanoscale particle accelerators comprise a dielectric material possessing electric polarizability and a metallic material capable of forming an interstitial and/or electrically conductive metal hydride and capable of enhancing the local electric field by the geometry and/or by the sufficiently small dimensions of the said metallic material. Low to medium strength local electric fields are utilized for the generation of Rydberg matter and inverted Rydberg matter in the presence of a material capable of forming and storing Rydberg atoms. Destabilization of Rydberg matter and inverted Rydberg matter leads to solid state physical reactions that release energy.

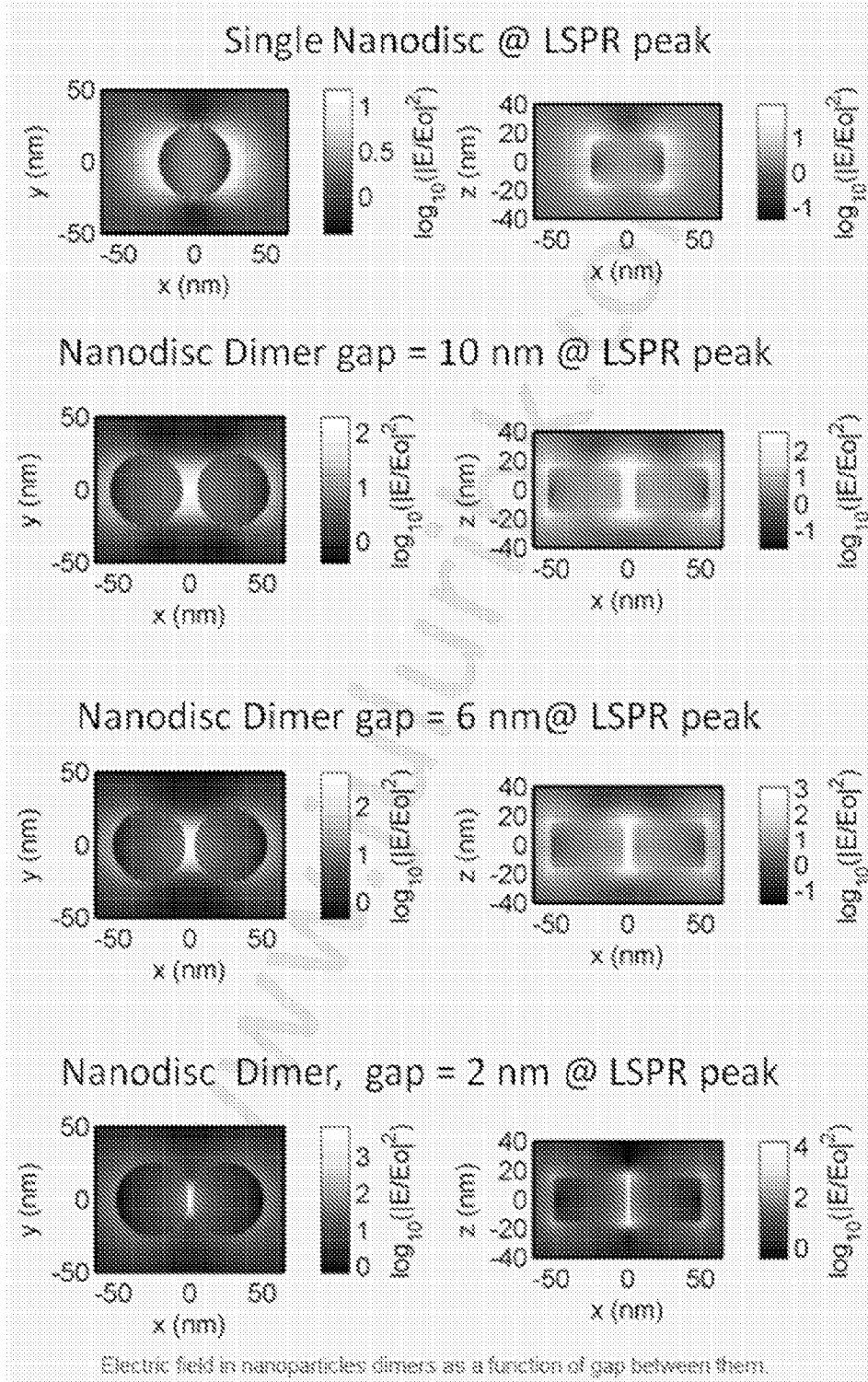
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Publication Classification

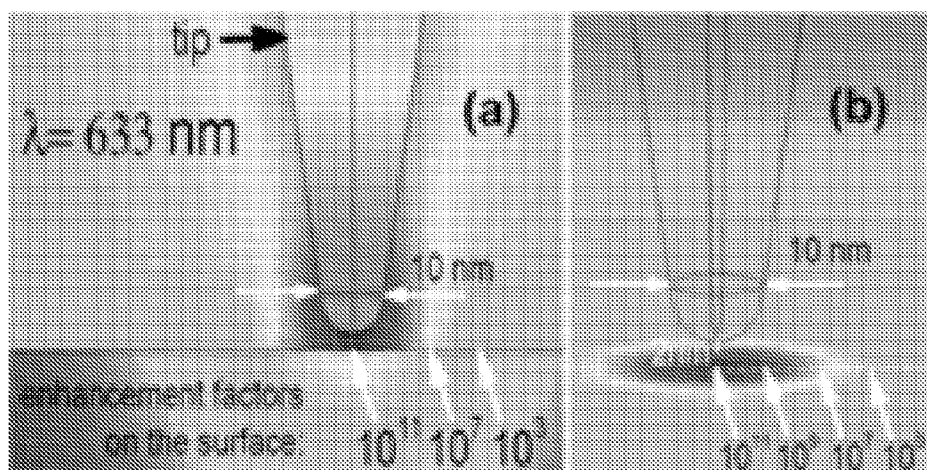
(51) **Int. Cl.**
G21B 3/00 (2006.01)





PRIOR ART

Fig. 1



PRIOR ART

Fig. 2

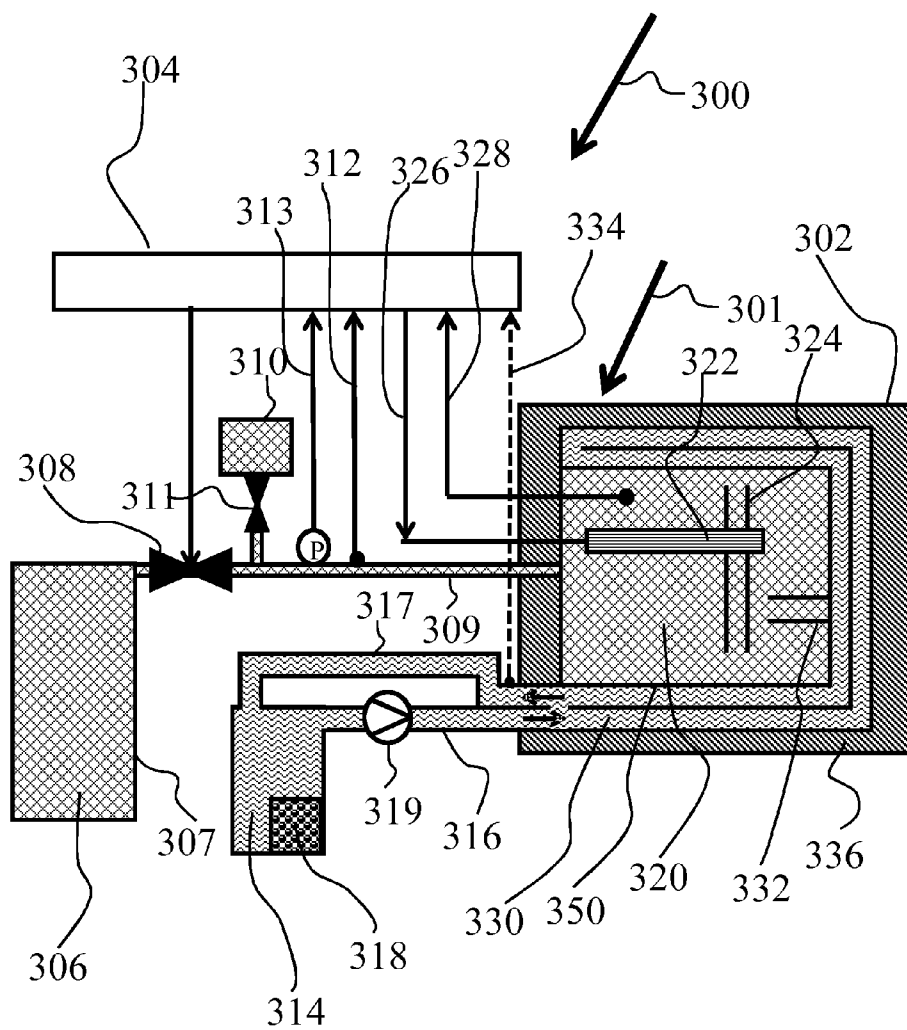


Fig. 3

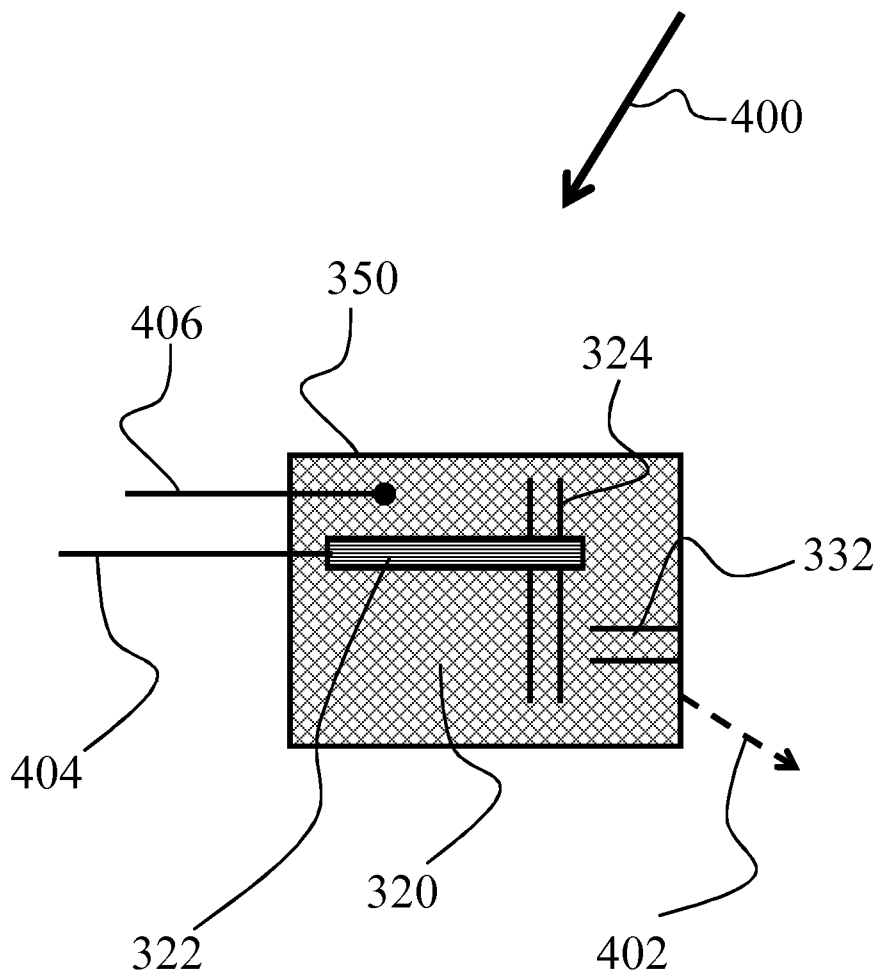


Fig. 4

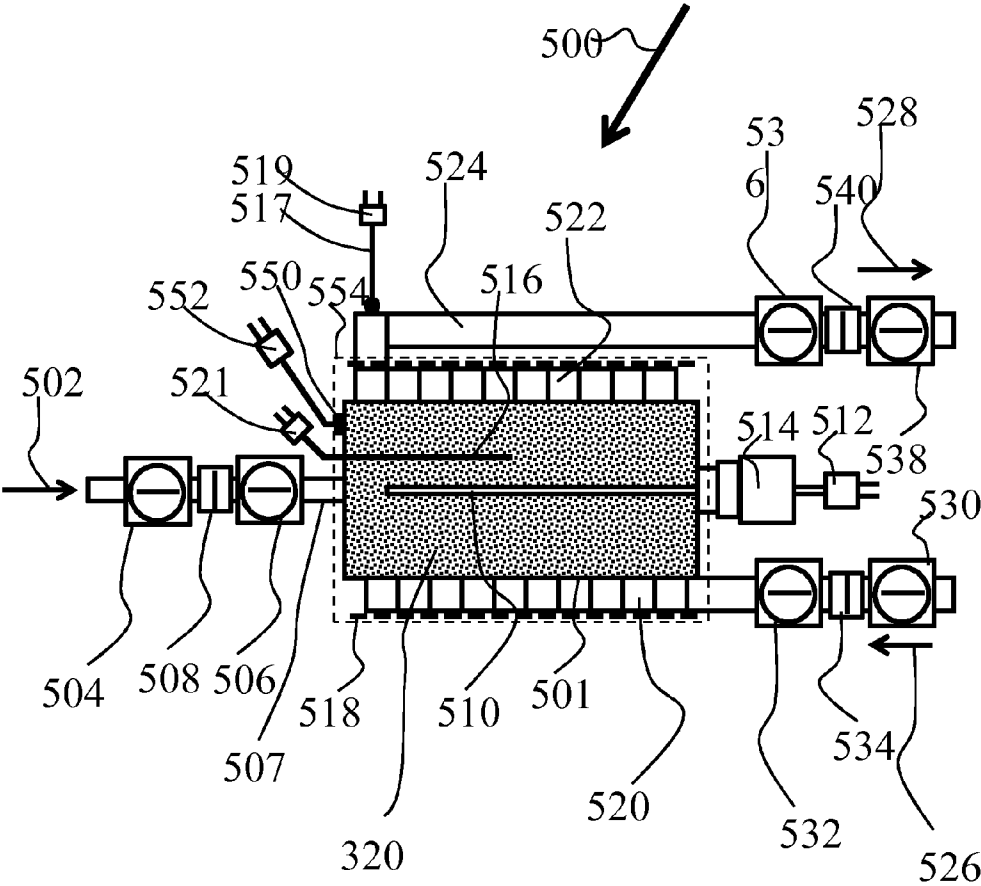


Fig. 5

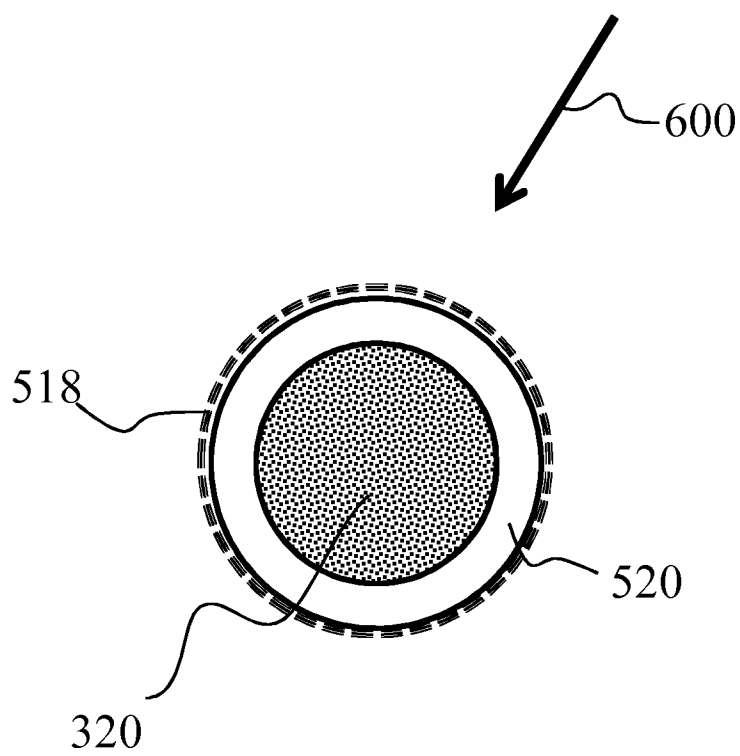


Fig. 6

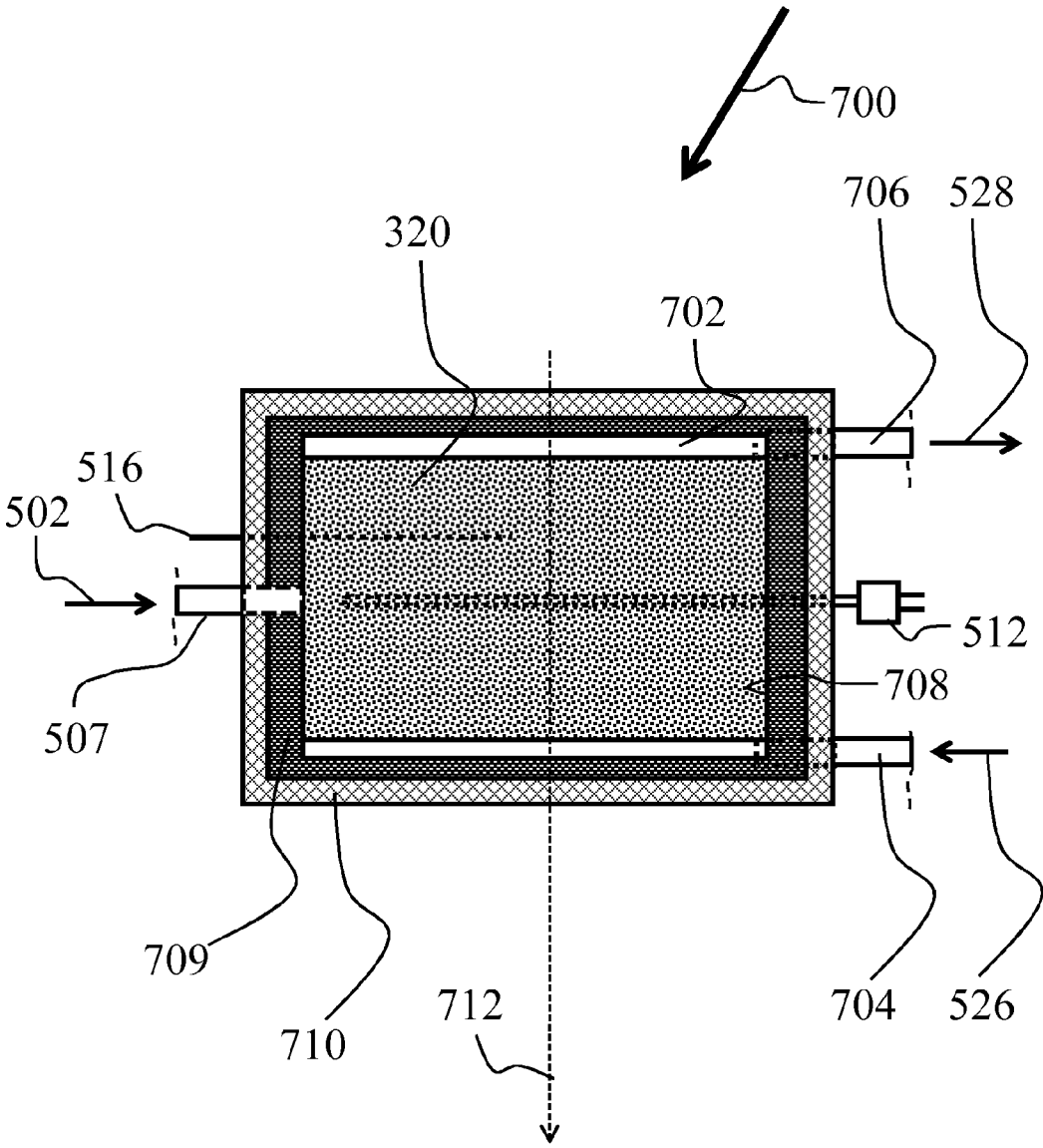


Fig. 7

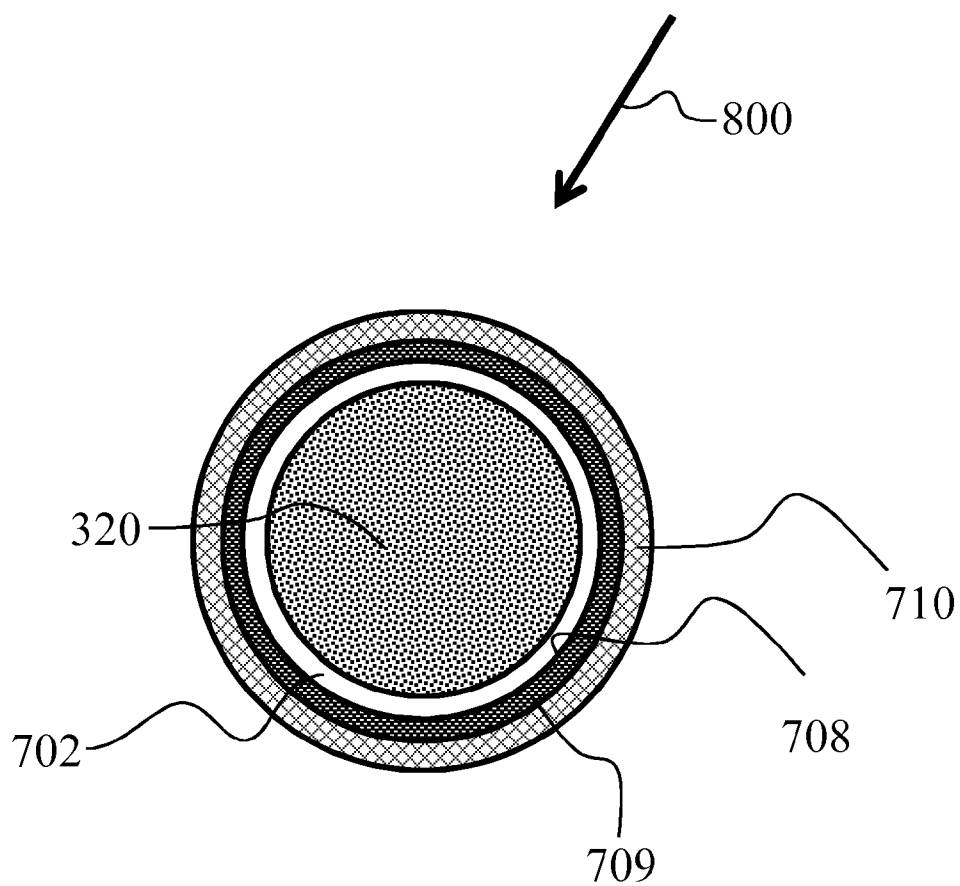


Fig. 8

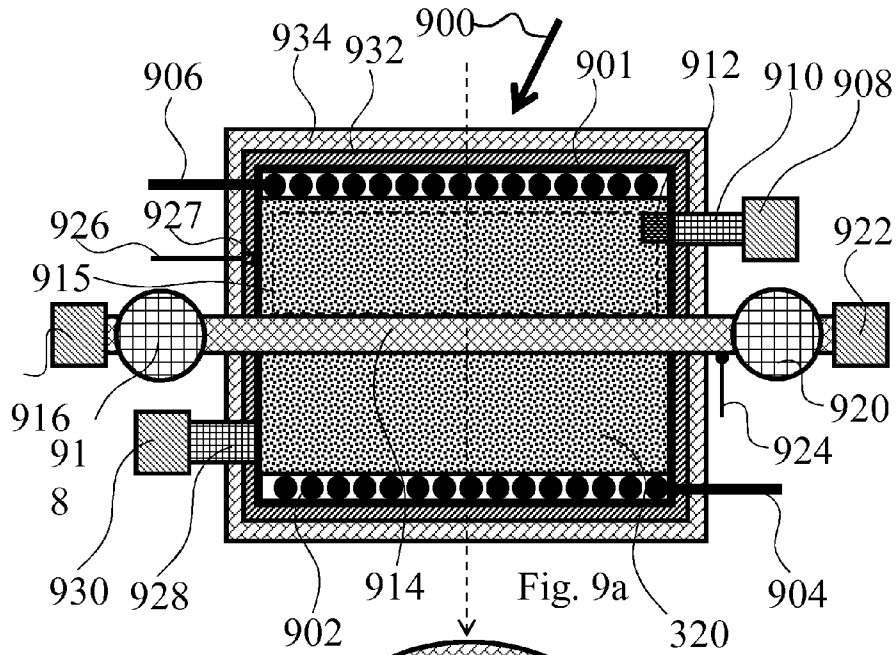


Fig. 9a

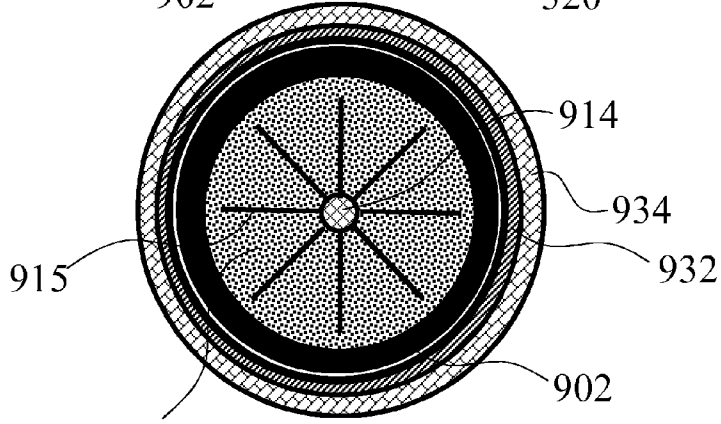


Fig. 9b

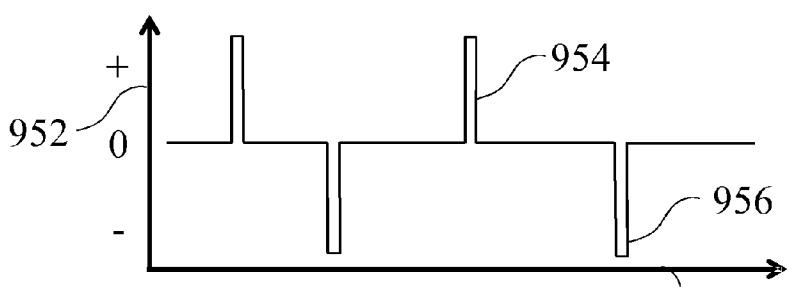


Fig. 9c

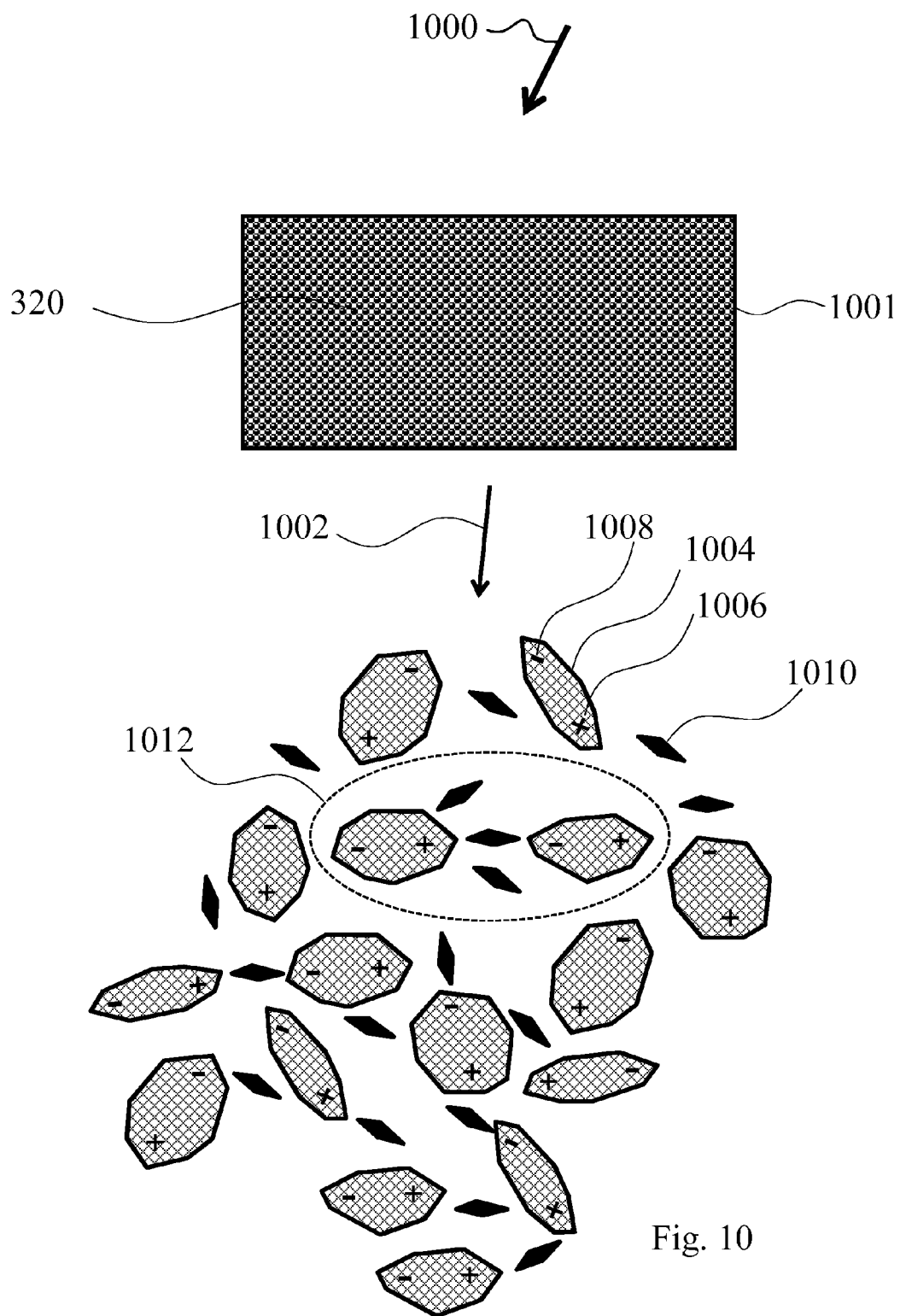


Fig. 10

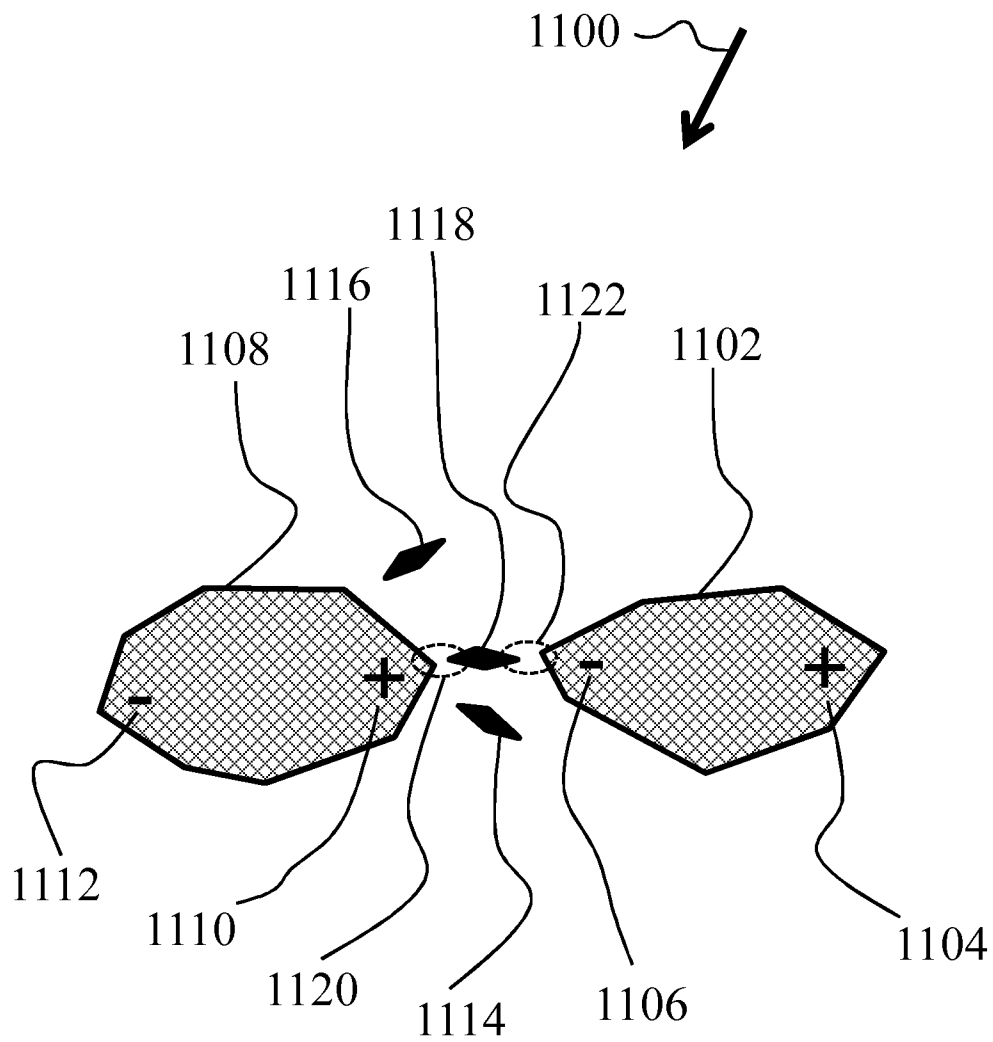


Fig. 11

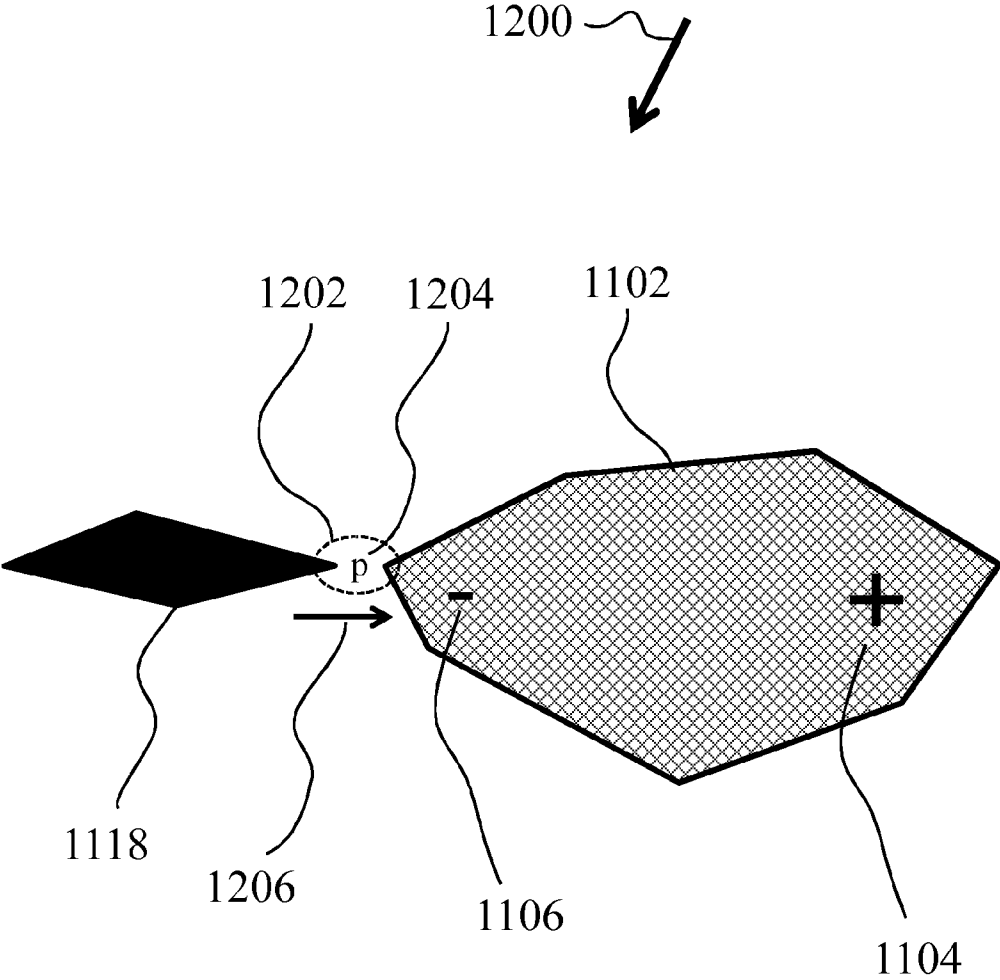


Fig. 12

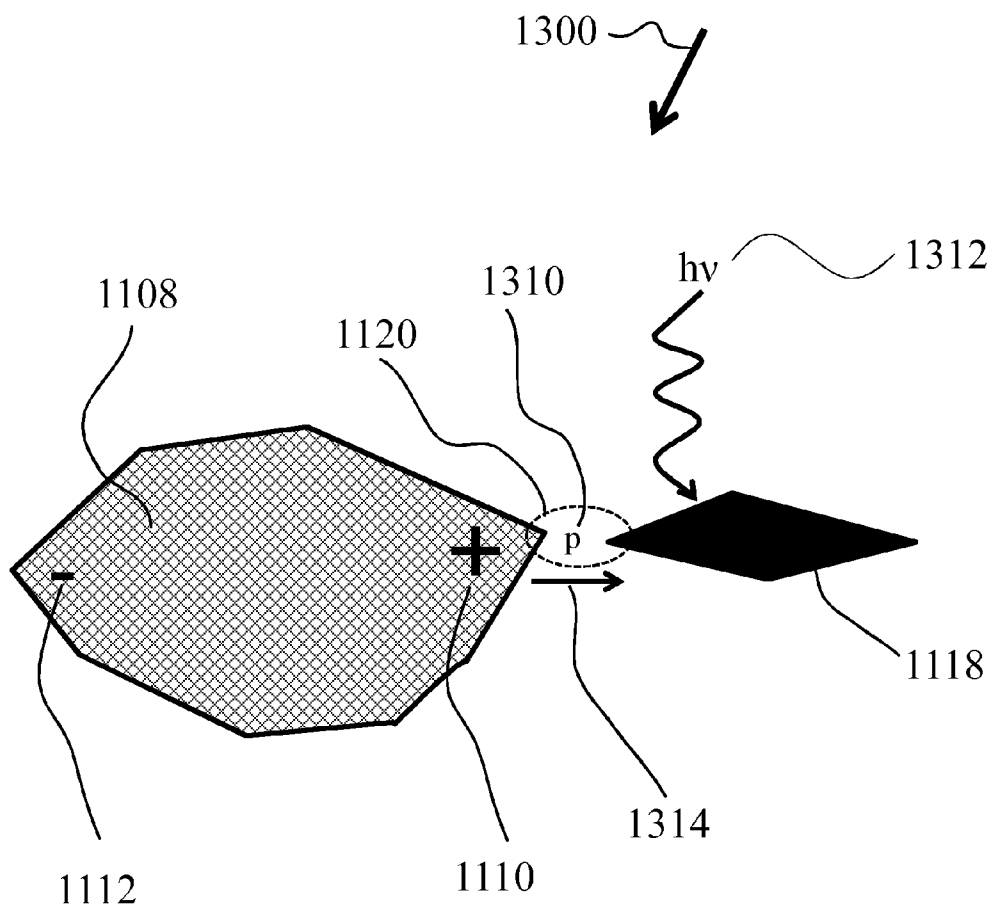


Fig. 13

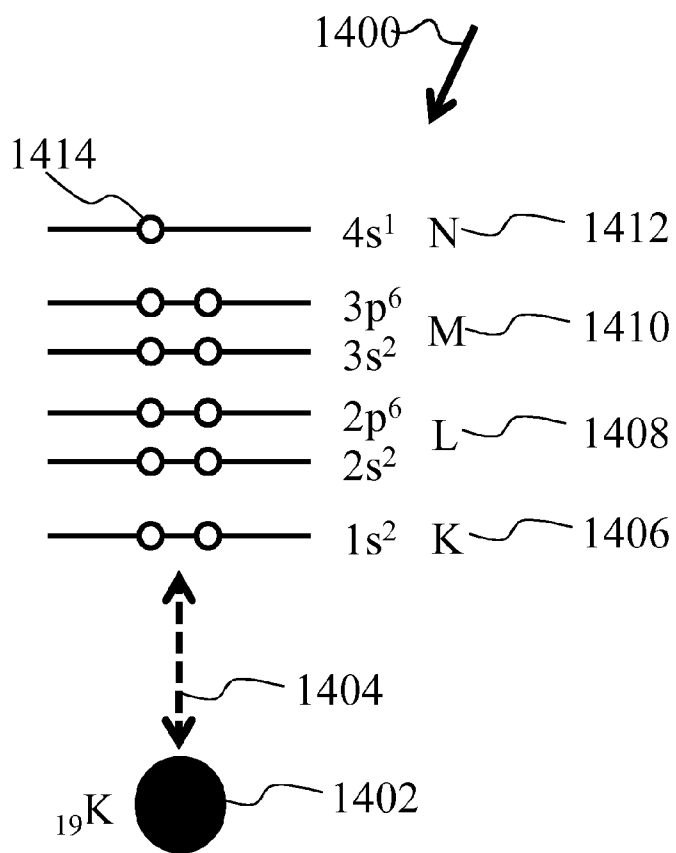


Fig. 14a

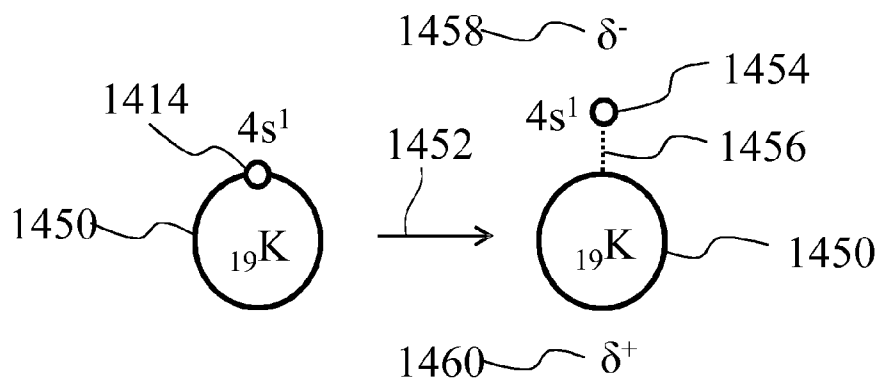


Fig. 14b

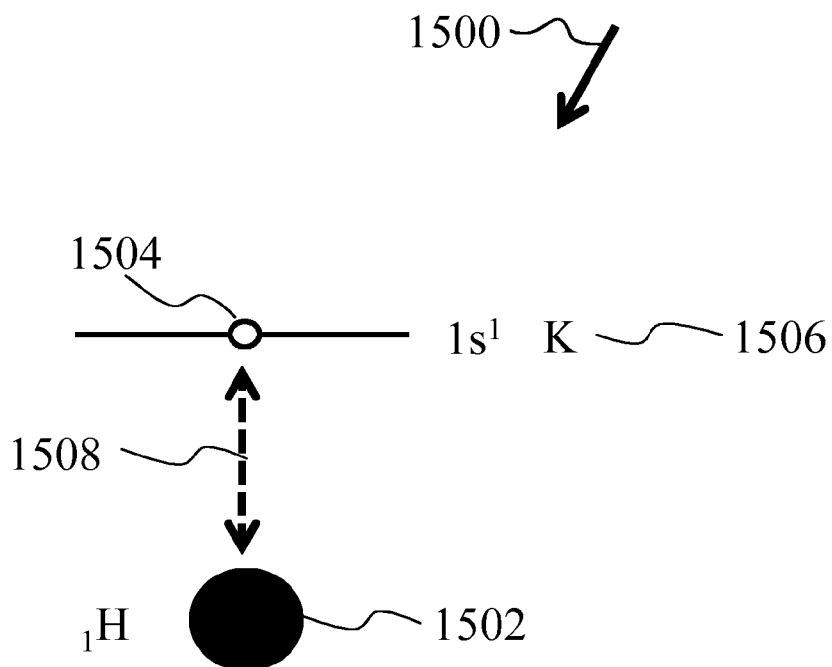


Fig. 15a

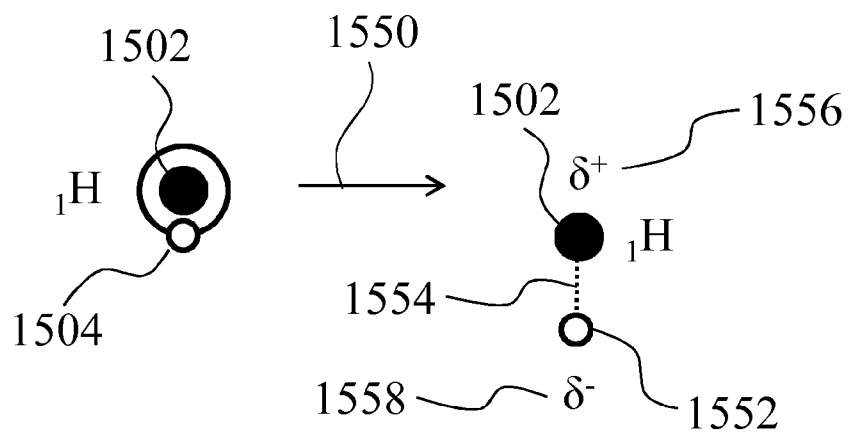


Fig. 15b

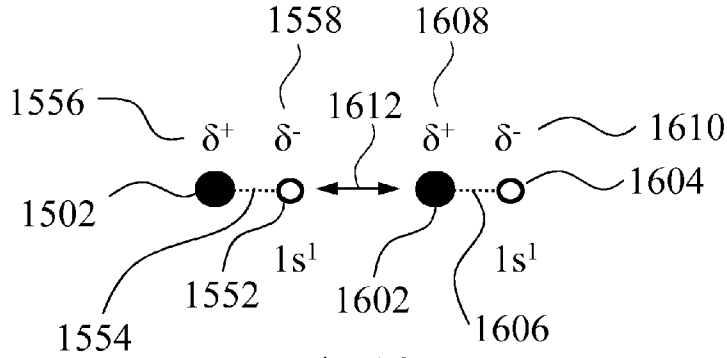


Fig. 16a

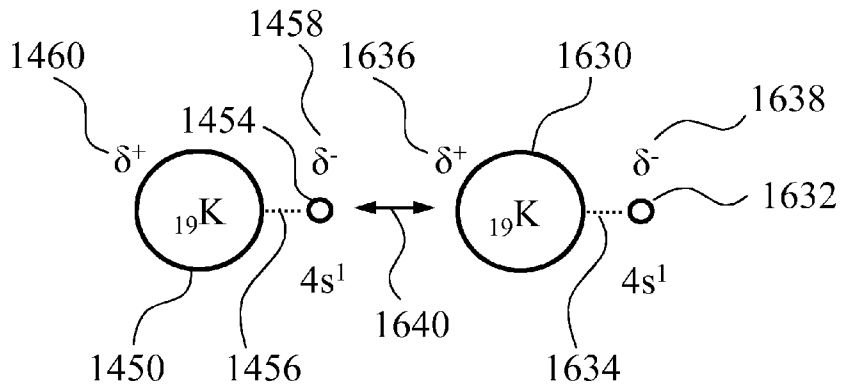


Fig. 16b

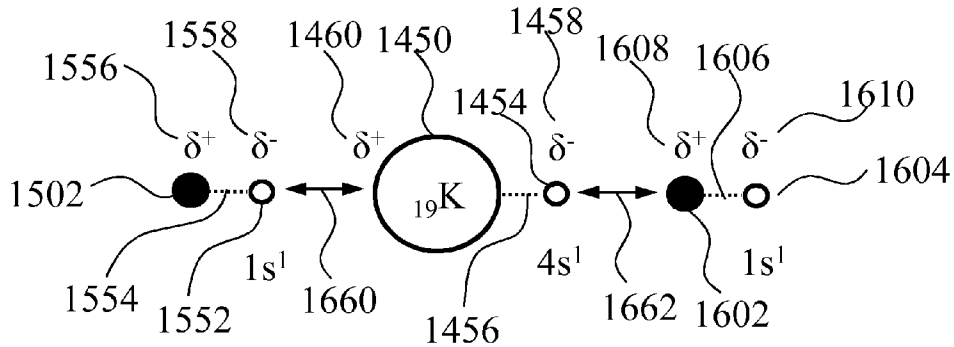


Fig. 16c

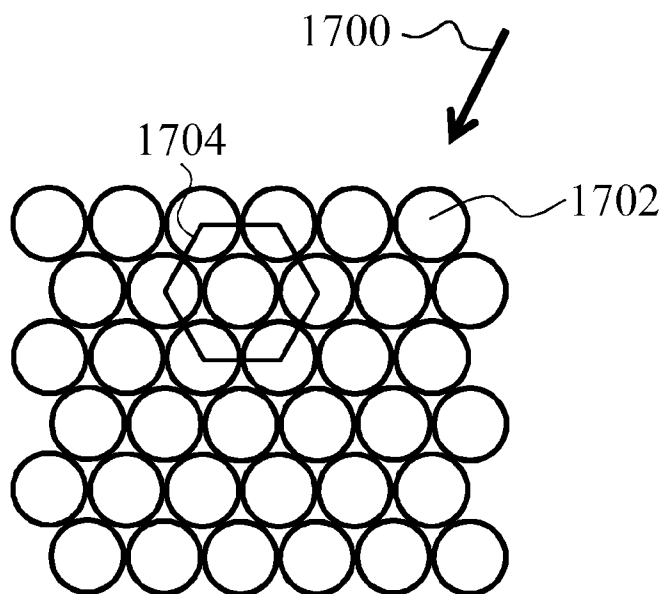


Fig. 17a

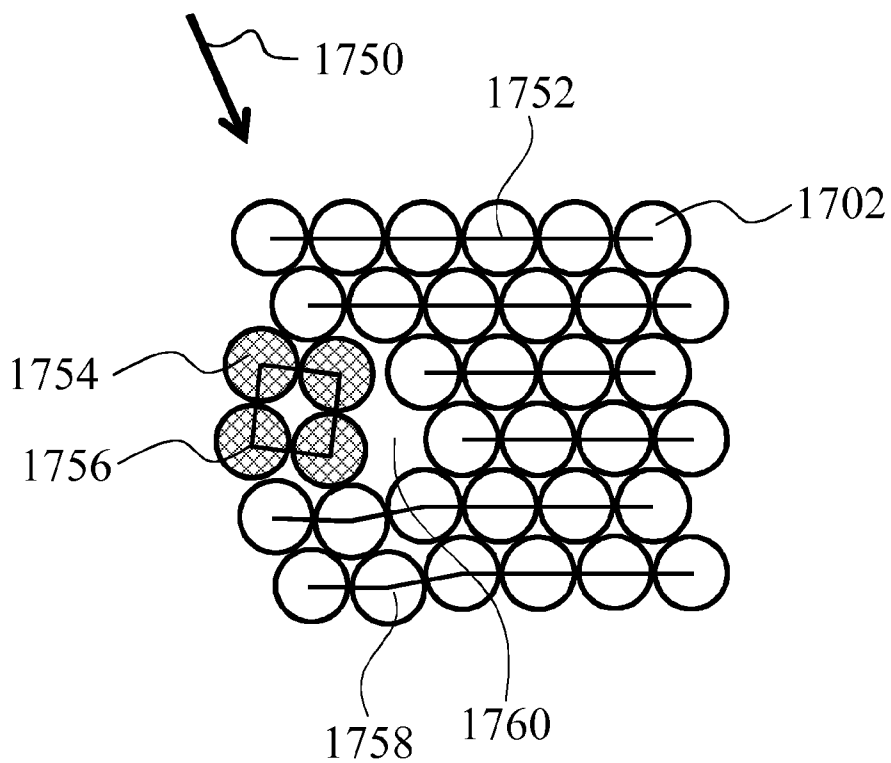


Fig. 17b

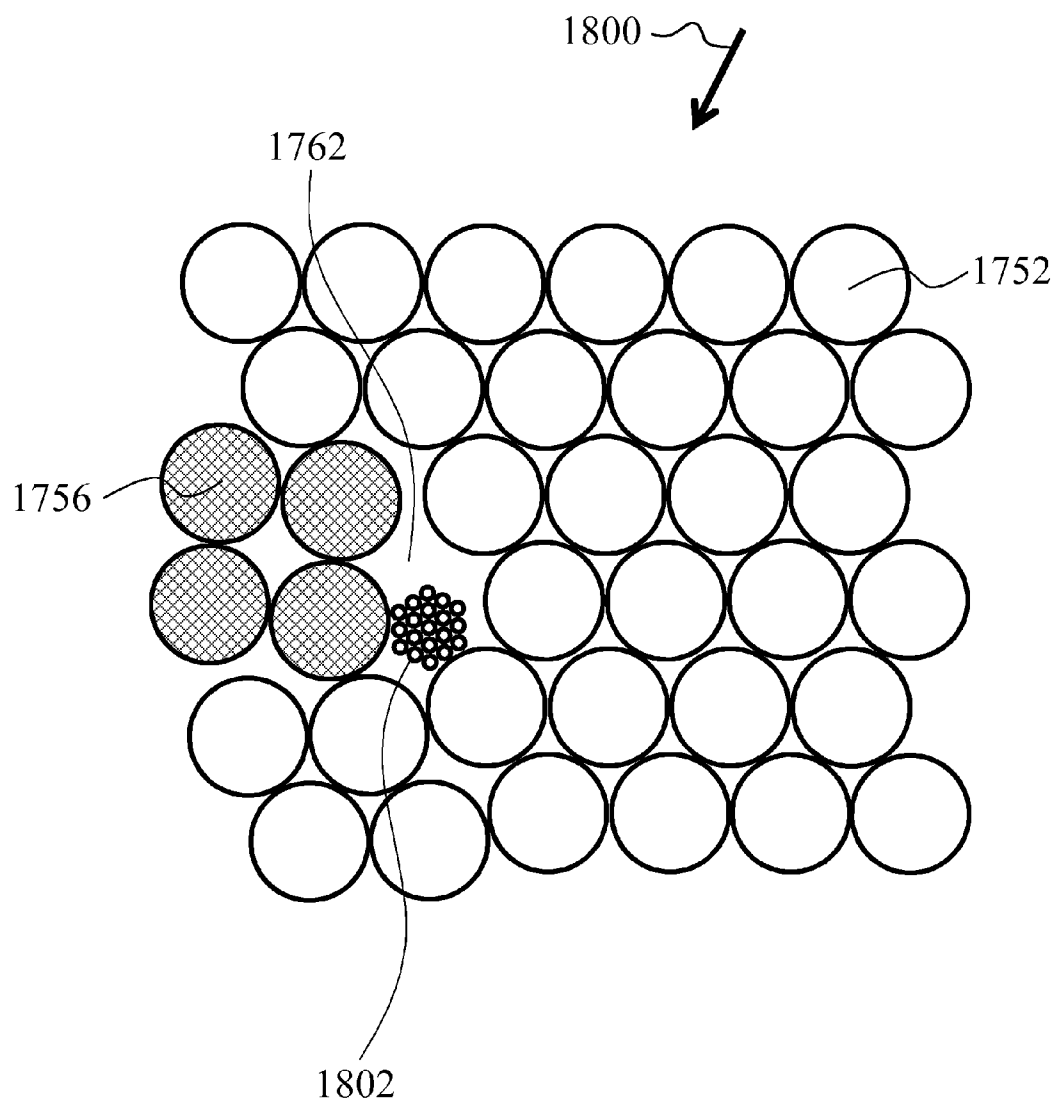


Fig. 18

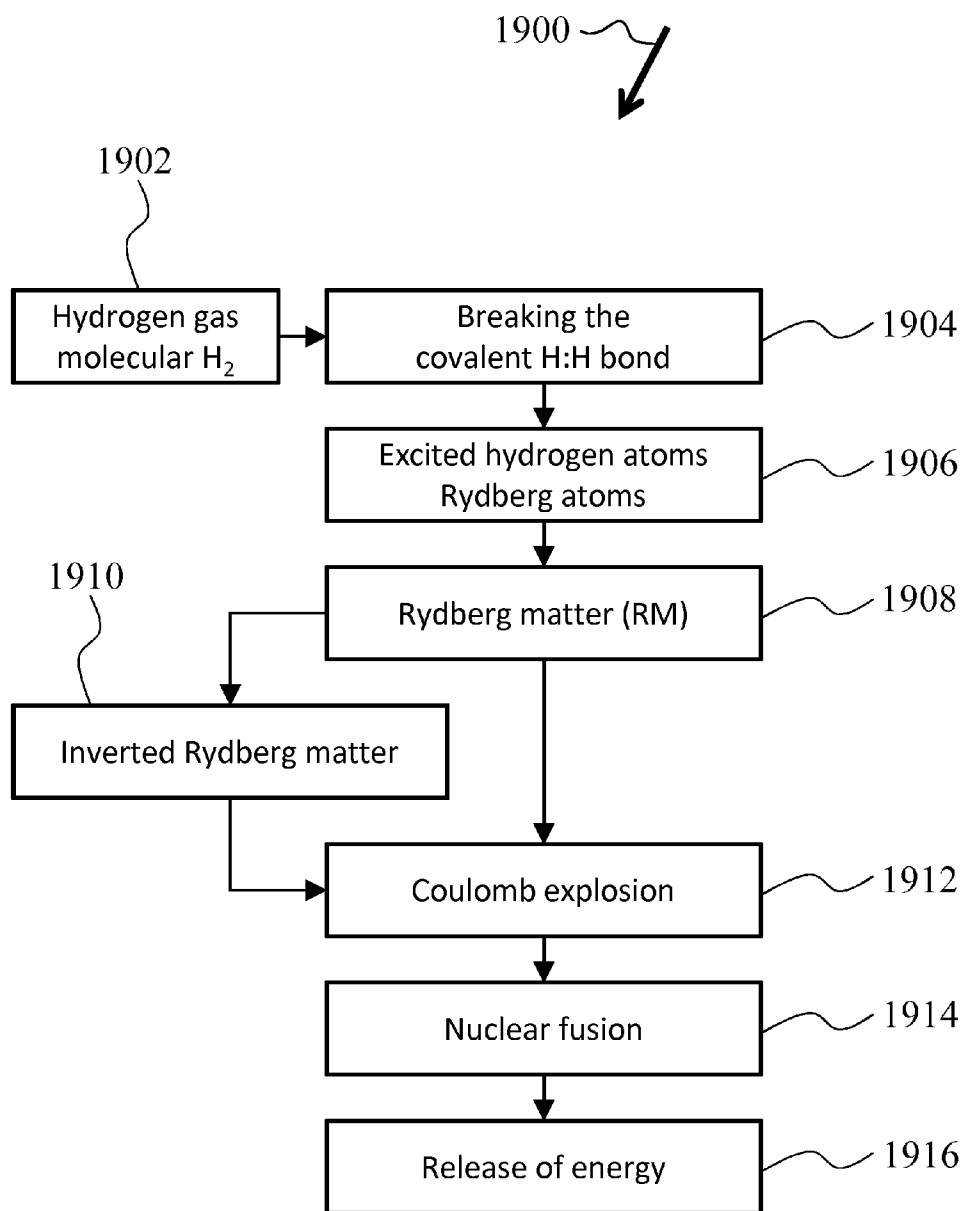


Fig. 19

THERMAL-ENERGY PRODUCING SYSTEM AND METHOD

TECHNICAL FIELD

[0001] The present invention relates generally to the production of thermal energy based on fusion reactions induced by strong electric fields.

BACKGROUND ART

[0002] According to the theory of special relativity energy has an equivalent mass and mass has an equivalent energy. The law of conservation of mass-energy in an isolated system means that the total amount of energy (energy+mass converted into equivalent energy) must be constant. On the other hand, the law of conservation of mass-energy in an isolated system means that the total amount of mass (mass+energy converted into equivalent mass) must be constant. Thus, loss of mass in the system means that energy must be released in the system. As a consequence, energy is released in the fusion reaction if the sum of masses of initial nuclei and possible elementary particles (e.g. neutrons) is larger than the mass of the final nucleus and possible elementary particles (e.g. neutrons).

[0003] According to Jeffrey A. Geuthera and Yaron Danon in a publication titled "Electron and positive ion acceleration with pyroelectric crystals", published in *Journal of Applied Physics* 97, 074109 s2005d, electric field strength of 1.35×10^7 V/cm was obtained in a lithium niobate crystal with $\Delta T = 75^\circ$ C.

[0004] W. Hu et al. present piezoelectric materials made from ternary solid solutions of $\text{BiFeO}_3\text{—PbZrO}_3\text{—PbTiO}_3$, in *Journal of the European Ceramic Society*, Volume 31, Issue 5, May 2011, Pages 801-807, which is incorporated herein as a reference. As an example of ternary solid solutions, $0.648\text{BiFeO}_3\text{—}0.053\text{PbZrO}_3\text{—}0.299\text{PbTiO}_3$ has a Curie temperature of 560° C.

[0005] P. Shiv Halasyamani and Kenneth R. Poeppelmeier have compiled and categorized over 500 noncentrosymmetric oxides by symmetry-dependent property and crystal class in *Chem. Mater.* 1998, 10, pp. 2753-2769, which is incorporated herein as a reference. Noncentrosymmetric (NCS) compounds possess symmetry-dependent properties comprising piezoelectricity and ferroelectricity.

[0006] Commercial nanopowders of metals and metal compounds are available from various companies. American Elements, 23 Rue Des Ardennes, 75019 Paris, France, sells various nanopowders, e.g. nickel oxide nanopowder (typical particle diameter 10-30 nm, specific surface area 130-150 m^2/g).

[0007] It is known that metallic hydrides are formed by most of the d-block elements (i.e., transition elements), on reacting with hydrogen. Hydrogen exists in the atomic rather than ionic form. Due to small size of hydrogen atoms when compared to the metal atoms, hydrogen atoms occupy interstitial positions in the metal lattices. Thus these are interstitial compounds and some workers regard them nearly as solid solutions.

[0008] Transition metal hydrides have been described by A. Dedieu (ed.) in a book, "Transition metal hydrides", John Wiley & Sons, 1991, ISBN: 978-0-471-18768-4, which is incorporated herein by reference.

[0009] S-block elements that have at least one stable isotope consist of hydrogen (H), lithium (Li), sodium (Na),

potassium (K), rubidium (Rb), caesium (Cs), helium (He), beryllium (Be), magnesium (Mg), calcium (Ca), strontium (Sr) and barium (Ba). In addition to transition metals, it is known that beryllium and magnesium of s-block elements also form metallic hydrides, i.e. they have low electric resistivity.

[0010] When hydrogen is absorbed into the interstices of transition metal lattices, metallic hydrides are formed. For example, palladium metal absorbs hydrogen to form palladium hydride. In some cases, the metals (e.g. palladium Pd) are used as cathodes in the electrolysis of their aqueous solutions so that metals absorb hydrogen during electrolysis and form metal hydrides such as PdH_2 . Metallic or interstitial hydrides compounds are non-stoichiometric, and their composition varies with temperature and pressure. As an example, the compositions of titanium and zirconium hydrides are often represented as $\text{TiH}_{1.7}$, and $\text{ZrH}_{1.9}$, respectively. They release hydrogen easily and are strong reducing agents, suggesting that the presence of hydrogen in its atomic state. These compounds are used as industrial reducing agents.

[0011] Complex transition metal hydrides have been described in a publication by Klaus Yvon & Guillaume Renaudin, "Hydrides: Solid State Transition Metal Complexes", *Encyclopedia of Inorganic Chemistry*, Second Edition (ISBN 0-470-86078-2), Volume III, pp. 1814-1846, which is incorporated herein by reference.

[0012] Crystal structures can be divided into 32 classes, or point groups. Ten point groups of the 32 point groups are polar. All polar crystals are pyroelectric, so the 10 polar crystal classes are sometimes referred to as the pyroelectric classes. Pyroelectric crystal classes are 1, 2, m, mm2, 3, 3 m, 4, 4 mm, 6 and 6 mm.

[0013] Piezoelectric crystal classes are 1, 2, m, 222, mm2, 4, -4, 422, 4 mm, -42 m, 3, 32, 3 m, 6, -6, 622, 6 mm, -62 m, 23 and -43 m.

[0014] Decreasing the grain size (crystallite size) of a dielectric material does not destroy the desired properties of a dielectric material. Actually, desired properties, such as the dielectric constant, are greatly improved by decreasing the crystallite size. Decreasing crystallite size of a dielectric material increased clearly the dielectric constant of the said dielectric material, as published by S. S. Batsanov, V. I. Galko and K. V. Papugin, "Dielectric permittivity and electrical conductivity of polycrystalline materials", *Inorganic Materials* 2010, vol. 46, no. 12, pp. 1365-1368, which is incorporated herein by reference.

[0015] It is generally known that hydrogen is a dielectric gas that does not conduct electricity in normal conditions. In very strong electric field (very steep voltage gradient) electron can be ripped off the hydrogen atom and plasma consisting of electrons and protons is formed. The very strong electric field has typically been created at the macroscopic level where the dimensions of the system are so large that the presence of hydrogen plasma can be observed visually by eye.

[0016] B. Naranjo, J. K. Gimzewski and S. Putterman have observed nuclear fusion driven by a pyroelectric crystal. The results were published in *Nature* 434, 1115-1117 (28 Apr. 2005), digital object identifier doi:10.1038/nature03575. However, the energy required to produce the fusion reactions in their experimental setup exceeded the energy produced by the fusion reactions. Thus, the coefficient of performance COP was below 1.

[0017] Metallic (interstitial) hydrides have been described by K. Yvon, "Hydrogen in novel solid-state metal hydrides",

Z. Kristallogr. 218 (2003) 108-116, which is incorporated herein by reference. Metallic hydrides in K. Yvon's publications comprise interstitial hydrides such as quaternary metal hydrides (the first metal, the second metal, the third metal and hydrogen) $\text{CeMn}_{1.8}\text{Al}_{0.2}\text{H}_{4.4}$, $\text{NdNi}_4\text{MgH}_4$ and $\text{LaMg}_2\text{NiH}_7$.

[0018] The decomposition temperature of the metal hydride depends on the metal hydride compound. For example, nickel hydride exists at temperatures up to hundreds of degrees centigrade ($^{\circ}\text{C}$.) as disclosed by B. Baranowski and S. M. Filipek in Polish J. Chem., 79, 789-806 (2005), which is incorporated herein by reference. On the other hand, copper hydride may decompose near room temperature in basic environment, although in acidic environment the decomposition temperature of copper hydride is higher, as disclosed by Nuala P. Fitzsimons in Catalysis Letters 15 (1992) 83-94, which is incorporated herein by reference.

[0019] Nickel compounds and manufacturing of nickel oxides, nickel hydroxides and nickel carbonate from nickel compounds have been described in Kirk-Othmer Encyclopedia of Chemical Technology (4th Edition), Vol 17, Nickel compounds, which is incorporated herein by reference.

[0020] Catalysts used for activating hydrogen by breaking chemical bonds between hydrogen atoms and other atoms and forming reactive hydrogen have been described by James H. Clark, Duncan J. Macquarrie and Mario De bruyn, in Kirk-Othmer Encyclopedia of Chemical Technology, "Catalyst, Supported", Wiley online edition 2011, <http://dx.doi.org/10.1002/0471238961.1921161614090512.a01.pub3>, and by Olaf Deutschmann, Helmut Knozinger, Karl Kochloeff and Thomas Turek, in Ullmann's Encyclopedia of Industrial Chemistry, Wiley-VCH Verlag GmbH & Co. KGaA online edition 2009, article title "Heterogeneous Catalysis and Solid Catalysts", http://dx.doi.org/10.1002/14356007.a05_313.pub2, and by Carlo Giavarini and Ferruccio Trifirò, in Encyclopaedia of Hydrocarbons, Volume II, Refining and Petrochemicals, Istituto Della Enciclopedia Italiana, Fondata da Giovanni Treccani S.p.A., Italy 2005, which are incorporated herein by reference.

[0021] Common industrial catalysts activating hydrogen comprise styrene synthesis catalysts that are examples of dehydrogenation catalysts (also known as hydrogen abstraction catalysts), ammonia (NH_3) synthesis catalysts, Fischer-Tropsch synthesis catalysts, high temperature water gas shift (HT-WGS) catalysts and hydrogenation catalysts (such as oil and fat hydrogenation catalysts).

[0022] Precursors for styrene catalysts comprise iron oxide Fe_2O_3 (red iron oxide, hematite) mixed with e.g. at least 10 wt % potassium oxide (K_2O) acting as an electron source and activation promoter, and small amounts of alumina (Al_2O_3) and chromia (Cr_2O_3) acting as structural promoters. Fe_2O_3 has rhombohedral corundum ($\alpha\text{-Al}_2\text{O}_3$) crystal structure and when it is reduced with hydrogen during the manufacturing of the styrene catalyst, black iron oxide, magnetite Fe_3O_4 with inverse spinel structure is formed. Fe_3O_4 consists of $\text{FeO}^*\text{Fe}_2\text{O}_3$ and it has iron in two oxidation states, namely Fe(II) and Fe(III). Residual Fe_2O_3 unit cells with the corundum structure induce lattice defects in the Fe_3O_4 catalyst. Addition of stable metal oxides, e.g. the said alumina and chromia and sometimes V_2O_3 with the corundum structure, preserves lattice defects in the catalyst and keeps the catalyst in active condition. Thus, a common composition of the industrial styrene synthesis catalyst is $\text{Fe}_3\text{O}_4\cdot\text{K}$, Al_2O_3 or $\text{Fe}_3\text{O}_4\cdot\text{K}$, Cr_2O_3 . These catalysts are typically used at temperatures up to about 640°C .

[0023] In the point of catalyst activity, useful structural defects form in the boundaries between the different crystal phases (e.g. hematite/corundum). The activity of the catalyst correlates with the number of lattice defects. Good catalysts have a large number of stable lattice defects. When heated, poor catalysts rearrange their crystal lattice structures so that lattice defects are eliminated and the activity of the catalyst drops.

[0024] Paracrystalline matter has short and medium range crystal order with a lot of structural defects. Materials with paracrystalline structure have distorted lattice cells and local microstrains because of the random point defects and are utilized as very active industrial catalysts. Known paracrystalline catalysts comprise nickel-alumina, iron-alumina and copper-zinc-alumina catalysts. Preparation methods for making highly active paracrystalline catalysts have been disclosed by D. C. Puxley, I. J. Kitchener, C. Komodromos and N. D. Parkyn in "The Effect Of Preparation Method Upon The Structures, Stability And Metal/Support Interactions In Nickel/Alumina Catalysts", Studies in Surface Science and Catalysis, volume 16, 1983, pages 237-271, [http://dx.doi.org/10.1016/S0167-2991\(09\)60025-2](http://dx.doi.org/10.1016/S0167-2991(09)60025-2), which is incorporated herein by reference.

[0025] Ammonia synthesis catalysts, utilized e.g. in the Haber-Bosch method, comprise iron promoted with potassium hydroxide KOH or potassium oxide (K_2O) to increase the local electron density and with textural promoters that are stable metal oxides in process conditions, often Al_2O_3 and/or CaO, to prevent sintering of iron metal particles. For example, a typical ammonia synthesis catalyst contains about 93 wt % Fe_3O_4 , about 1 wt % K_2O , about 3 wt % Al_2O_3 and about 3 wt % CaO. These catalysts are typically used at temperatures up to about 450°C .

[0026] Fischer-Tropsch method converts carbon monoxide (CO) and hydrogen into various hydrocarbons. Fischer-Tropsch catalysts comprise cobalt (Co), iron (Fe), ruthenium (Ru) or nickel (Ni) promoted with copper (Cu) or group 1 alkali metals, e.g. potassium (K). Often Fe_2O_3 precursor doped with potassium (in the form of KOH or K_2O) is reduced with hydrogen into $\text{Fe}_3\text{O}_4\cdot\text{K}$ and utilized as the Fischer-Tropsch catalyst. These catalysts are typically used at temperatures up to about 300°C .

[0027] High temperature water gas shift method converts carbon monoxide (CO) and water (H_2O) into carbon dioxide (CO_2) and hydrogen (H_2). High temperature water gas shift catalysts comprise iron oxides doped with potassium or lanthanum niobate LaNiO_3 promoted with potassium. These catalysts are typically used at temperatures up to about 350°C .

[0028] Hydrogenation catalysts comprise platinum (Pt), palladium (Pd), rhodium (Rh), ruthenium (Ru), alloys of Pt, Pd, Rh and Ru, Raney nickel, Urushibara nickel and nickel oxide. Nickel forms stoichiometric green nickel oxide NiO with a NaCl crystal structure, non-stoichiometric black Ni_{1-x}O , wherein x is about 0.02, and black Ni(III) oxide Ni_2O_3 . Potassium is often added to the nickel catalysts for promoting catalytic activity. These catalysts are typically used at temperatures up to several hundred $^{\circ}\text{C}$.

[0029] An alkali metal, usually potassium, is essential for the activity of the styrene catalyst. In the styrene catalyst potassium forms potassium ferrites (potassium-iron-oxides), mainly KFeO_2 surface phase and $\text{K}_2\text{Fe}_{22}\text{O}_{34}$ phase with a cubic crystal structure similar to the inverse spinel structure of Fe_3O_4 (magnetite).

[0030] Crystalline materials have various types of defects. Point defects are cation and anion vacancies and interstitial atoms in the crystal structure. Vacancies can form clusters of vacancies for example in iron oxide crystals. Vacancy clusters are voids, i.e. small regions in the crystal without atoms. Line defects in crystals include edge dislocations and screw dislocations. Planar defects are stacking faults in crystals and grain boundary interfaces. Defects are essential for the high activity of catalysts. Structural promoters added to catalyst materials preserve the nickel defects in the catalyst materials lengthening the lifetime of catalysts.

[0031] Stoichiometric nickel oxide NiO has green color and it is an insulator. Nickel-deficient non-stoichiometric nickel oxide Ni_{1-x}O, wherein x is typically about 0.02, has black color and is a semiconductor. There are nickel vacancies in Ni_{1-x}O. Adding alkali metal, e.g. lithium Li, to Ni_{1-x}O makes the nickel oxide a much better conductor of electricity, a metallic metal oxide. Nickel oxide with defective crystal structure has very high catalytic activity and it is used as a hydrogenation catalyst.

[0032] Regarding the storage of hydrogen, several metal ammine salts capable of storing hydrogen in the form of ammonia have been disclosed by Rasmus Z. Sorensen, Jens S. Hummelshøj, Asbjørn Klerke, Jacob Birke Reves, Tejs Vegge, Jens K. Nørskov and Claus H. Christensen in "Indirect, Reversible High-Density Hydrogen Storage in Compact Metal Ammine Salts", J. Am. Chem. Soc. 2008, 130, pp. 8660-8668, <http://dx.doi.org/10.1021/ja076762c>, which is incorporated herein by reference. Specifically, MgCl₂ molecule is capable of binding up to 6 NH₃ molecules and forming Mg(NH₃)₆Cl₂ salt that stores over 9 wt % hydrogen and has only 2.2 mbar NH₃ vapor pressure at +27° C. Ammonia is released from the metal ammine salt by heating the salt. Ammonia gas is cracked into hydrogen gas and nitrogen gas at elevated temperatures preferably with a catalyst, e.g. Ni, Pt catalyst or a catalyst based on carbon nanotubes doped with ruthenium and potassium hydroxide.

[0033] Regarding the penetration of the Coulomb barrier around the atom nucleus, resonance of a wave function of a particle in a quantum well system has been described by David Bohm, "Quantum theory", Prentice-Hall, New York 1951, which is incorporated herein by reference. Specifically, a wave is reflecting back and forth across the potential in a quantum well, a wave coming in the quantum well from outside enhances the wave inside the quantum well and a strong standing wave is built up inside the quantum well when the system is in resonance. Further, the waveform of a proton tunnels through the Coulomb barrier to the nucleus of an atom with certain probability. Near a resonance the waveform intensity of the proton is considerable in the quantum well and the probability of fusing proton with the nucleus is increased. The metastable state of the fused nucleus may have such a long lifetime in solid state structures that it can decay in other ways than by re-emission of the incident proton or by emission of gamma-ray photons, and energy is released over relatively long time also as lower energy photons (e.g. X-ray photons) or as phonons (lattice vibrations) to the surrounding solid lattice.

[0034] When one or more electrons of an atom are excited to high principal quantum number, the excited electron is in the Rydberg state and the atom becomes a Rydberg atom. It is an electrical dipole with a positive core and a negative excited electron orbiting relatively far from the core. As a result, external electric and magnetic fields have a big effect on

Rydberg atoms. Rydberg atoms interact with each other because of the electrical dipole properties and are capable of binding together. Rydberg atoms are produced e.g. by electron impact excitation, charge exchange excitation and optical excitation. Excitation energy below the ionization energy produces Rydberg states in atoms. These Rydberg atoms are electrically polarized, which pulls Rydberg atoms together forming clusters of Rydberg atoms.

[0035] Until now elements that have been found to possess Rydberg states comprise H, Li, Na, K, Rb, Cs, N, Ni, Ag, Cu, Pd, Ti and Y.

[0036] Rydberg formula describes wavelengths of spectral lines observable in atoms that have an electron in Rydberg states. Rydberg formula for any hydrogen-like element is $1/\lambda_{vac} = RZ^2(1/n_1^2 - 1/n_2^2)$, wherein λ_{vac} is the wavelength of the light emitted in vacuum; R is the Rydberg constant for this element; Z is the number of protons in the atomic nucleus of this element (atomic number); n₁ and n₂ are principal quantum number integers such that n₁ < n₂.

[0037] Rydberg states are closely spaced and they form Rydberg series (n₂=n₁+1, n₁+2, . . . , n₁+∞). In case of hydrogen atom (Z=1) Rydberg series comprise Lyman series (n₁=1, n₂=2, 3, . . . , ∞) from 121.6 nm (10.2 eV) to 91.18 nm (13.6 eV, Lyman limit, ionization energy), Balmer series (n₁=2, n₂=3, 4, . . . , ∞) from 656.3 nm (1.89 eV) to 364.6 nm (3.4 eV, Balmer limit, ionization energy), Paschen series (n₁=3, n₂=4, 5, . . . , ∞) from 1870 nm (0.66 eV) to 820 nm (1.51 eV, Paschen limit, ionization energy), Brackett series (n₁=4, n₂=5, 6, . . . , ∞) from 4050 nm (0.31 eV) to 1460 nm (0.85 eV, Brackett limit, ionization energy), Pfund series (n₁=5, n₂=6, 7, . . . , ∞) from 7460 nm (0.17 eV) to 2280 nm (0.54 eV, Pfund limit, ionization energy) and Humphreys series (n₁=6, n₂=7, 8, . . . , ∞) from 12400 nm (0.10 eV) to 3280 nm (0.38 eV, Humphreys limit, ionization energy). Wavelengths (nm) can easily be converted to electron volts (eV) and vice versa with the equation $E(\text{eV}) \approx 1240 \text{ eVnm}/\lambda(\text{nm})$.

[0038] The electron gains energy when its principal quantum number is increased and it emits a photon when its principal quantum number decreases. When the Rydberg atom is ionized, i.e. an electron is excited with an ionization energy, the quantum mechanical unity of the Rydberg atom is lost and separate particles (positive ion and negative electron) are formed. It is possible to excite more than one electron in an atom (excluding hydrogen), but it is easier to form a Rydberg atom with a single excited electron.

[0039] Examples of first ionization energies of atoms, where an outer (valence) electron is lifted from n=1 to n=∞ and removed from the atom, are: H 13.598 eV, Li 5.392 eV, Na 5.139 eV, K 4.341 eV, Rb 4.177 eV, Cs 3.894 eV, Ni 7.640 eV, Pd 8.337 eV, Ag 7.576 eV and Ti 6.828 eV. Energies of Rydberg states of electrons are smaller than the said ionization energies.

[0040] A thorough textbook on Rydberg atoms has been written by Thomas F. Gallagher "Rydberg Atoms" (Cambridge Monographs on Atomic, Molecular and Chemical Physics), Cambridge University Press 1994, ISBN 0-521-38531-8.

[0041] Rydberg matter is a phase of matter formed by Rydberg atoms. Rydberg matter is held together by the delocalized excited electrons in Rydberg states to form an overall lower energy state of the cluster of Rydberg atoms. Lifetime of the cluster is in the order of seconds, minutes or even longer depending on the primary quantum number (n) of the Rydberg states. Rydberg matter seeks the lowest total energy of

the local system by the dipole-dipole interactions of the Rydberg atoms. Rydberg matter forms a condensed phase when the condensation energy is removed from the Rydberg matter.

[0042] Heavy Rydberg systems consisting of anion cation pairs bound together via electrostatic forces are also known, e.g. H^+H^- (proton, hydride ion). Quantum mechanics also allows an inverted system where, instead of an excited electron orbiting the core, the wavefunction corresponds to the setup where the core is in orbit around the excited electron. This kind of system is capable of forming extremely dense clusters of atoms.

[0043] Rydberg states of hydrogen molecules H_2^* have been observed at 900-1000 K on iron oxide surfaces that have been doped with potassium by Jiayi Wang and Leif Holmlid in "Formation of long-lived Rydberg states of H_2 at K impregnated surfaces", *Chemical Physics* 261 (2000) 481-488, [http://dx.doi.org/10.1016/S0301-0104\(00\)00288-3](http://dx.doi.org/10.1016/S0301-0104(00)00288-3).

[0044] Density of up to 10^{29} deuterium atoms/cm³ in Rydberg matter clusters in pores in iron oxide doped with potassium and calcium was confirmed by L. Holmlid, H. Hora, G. Miley and X. Yang in "Ultra-high-density deuterium of Rydberg matter clusters for inertial confinement fusion targets", *Laser and Particle Beams* 27 (2009) pp. 529-532, <http://dx.doi.org/10.1017/S0263034609990267>. The distance between deuterons was only 2.3 pm, which indicated that the deuterium clusters consisted of inverted Rydberg matter.

[0045] Hydrogen Rydberg matter and inverted Rydberg matter on potassium doped iron oxide catalyst were exposed to 564 nm laser beam to induce Coulomb explosions and found to emit high energy particles of up to 150 eV/atomic mass unit by Shahriar Badiei, Patric U. Andersson and Leif Holmlid in "Fusion reactions in high-density hydrogen: A fast route to small-scale fusion", *International Journal of Hydrogen Energy* 34 (2009) pp. 487-495, <http://dx.doi.org/10.1016/j.ijhydene.2008.10.024>. The energy of the particles corresponds to the 109 K temperature, which indicates that favorable conditions for nuclear fusion processes can be induced in solid state matter with a relatively low power laser.

SUMMARY OF INVENTION

[0046] Utilization of novel physical phenomena at the nanoscale makes it possible to construct a compact thermal energy source with cheap common materials.

[0047] A reaction container is filled with a reaction material and is pressurized with hydrogen gas.

[0048] The reaction material comprises a dielectric material that possesses electric polarizability, a metallic material capable of forming interstitial and/or electrically conductive metal hydrides and a material promoting the formation and storage of Rydberg matter.

[0049] Hydrogen isotopes utilized in the present invention are protium H, deuterium D and tritium T (generally hydrogen).

[0050] Hydrogen molecules H_2 are activated by breaking the chemical bond between hydrogen atoms. Activation is preferably done with a material comprising a transition metal, transition metals or mixtures of transition metals capable of forming metal hydrides such as nickel and platinum group metals. Atomic hydrogen (H) is formed by activation.

[0051] Atomic hydrogen H is ionized into H^+ (proton) in strong electric field. Hydrogen ions and electrons are accelerated to high kinetic energy by the strong electric field that has steep voltage gradient.

[0052] Original source of the electric field is preferably a dielectric material that can be polarized comprising piezoelectric material (electric polarization is induced by mechanical vibration, e.g. by an ultrasonic source), pyroelectric material (electric polarization is induced by variable temperature) and/or multiferroic material (electric polarization is induced by a magnetic field). Polarization of a material creates the electric field near the material.

[0053] Fusion reactions are initiated at the nanoscale (at least one dimension smaller than about 100 nm) by the combination of three control factors: sufficiently high hydrogen gas pressure in the reaction container, sufficiently high temperature in the reaction container and the polarization of a dielectric material.

[0054] Ionized hydrogen and electrons are accelerated with the local electric field to a kinetic energy that corresponds to the strength of the local electric field. Ionized hydrogen and electrons gain kinetic energy because of the acceleration. Ionized hydrogen with sufficiently high kinetic energy tunnels with sufficiently high probability through the Coulomb barrier between the ionized hydrogen and target atomic nucleus and fuses with the target atomic nucleus. Ionized hydrogen and electrons with relatively low kinetic energy excite electrons on particle surfaces and create Rydberg atoms. Local electric field is greatly enhanced by the geometry of the metallic nanoparticles and the short distance between the nanoparticles. An electron can be ripped away from the hydrogen atom that is near the metallic tip in strong electric field or is between metallic nanoparticles in strong electric field.

[0055] Based on the above, it is an aim of the present invention to provide a method of producing energy. It is a second aim of the invention to provide a nuclear fusion system for producing thermal energy. It is a third aim of the invention to provide a fusion energy production process. It is a fourth aim of the invention to provide a fusion energy reaction material. It is a fifth aim of the invention to provide a use of hydrogen-containing Rydberg matter and/or inverted Rydberg matter.

Technical Problem

[0056] The price of energy increases continuously. The production of energy from fossil fuels is problematic due to green house gas emissions. Solar energy and wind energy suffer from variable power output. Nuclear energy is not generally favored because of catastrophic accidents in nuclear power plants. Fusion energy research has not yet produced any working solution in spite of multi-billion dollar investments.

Solution to Problem

[0057] Electric field strengths capable of accelerating hydrogen ions to kinetic energies high enough to tunnel through the Coulomb barrier of nuclei and fusing with the nuclei are generated with a novel system comprising dielectric materials possessing electric polarizability that act as electric charge sources and metallic nanopowders capable of forming interstitial and/or electrically conductive metal hydrides that act as electric field focusing and electric field strength enhancing materials and hydrogen ion sources.

[0058] Electric field strengths capable of accelerating hydrogen ions and electrons to kinetic energies high enough to excite electrons on solid surfaces to Rydberg states and

form Rydberg matter are generated with a novel system comprising dielectric materials possessing electric polarizability that act as electric charge sources, metallic nanopowders capable of forming interstitial and/or electrically conductive metal hydrides that act as electric field focusing and electric field strength enhancing materials and hydrogen ion sources and catalytic nanopowders promoting the formation and storage of Rydberg matter.

Advantageous Effects of Invention

[0059] The price of thermal energy or electrical energy produced by the present invention is less than about 1 euro-cent/3.6 MJ or kWh. The amount and cost of fuel needed for the present thermal-energy generating system is very small compared to any system utilizing fossil fuels.

BRIEF DESCRIPTION OF DRAWINGS

[0060] FIG. 1 depicts prior art describing the enhancement of electric field in very small dimensions. Reference: <http://juluribk.com/2011/04/09/electric-field-in-metal-nanoparticle-dimers/>

[0061] FIG. 2 depicts prior art describing the enhancement of electric field by the formation of surface plasmons by photons. Reference: "Modern Aspects of Electrochemistry 44, Modeling and Simulations II", Vol. 2, pp. 70-73, M. Schlesinger, ed., Springer 2009.

[0062] FIG. 3 depicts an example embodiment of the fusion system of the present invention.

[0063] FIG. 4 depicts the details of the reaction container shown in FIG. 3.

[0064] FIG. 5 depicts another example embodiment of the fusion system of the present invention.

[0065] FIG. 6 depicts the cross section of the reaction container shown in FIG. 5.

[0066] FIG. 7 depicts an example embodiment of the fusion system of the present invention.

[0067] FIG. 8 depicts the cross section of the reaction container shown in FIG. 7.

[0068] FIGS. 9a, 9b and 9c depict another example embodiment of the fusion system of the present invention.

[0069] FIG. 10 depicts a close up view of the reaction material according to an example embodiment of the present invention.

[0070] FIG. 11 depicts the enhancement of electric field strength according to an example embodiment of the present invention.

[0071] FIG. 12 depicts the acceleration of ions according to an example embodiment of the present invention.

[0072] FIG. 13 depicts the acceleration of ions according to another example embodiment of the present invention.

[0073] FIGS. 14a and 14b depict the electron shell structure of the potassium atom and the excitation of the valence electron of the potassium atom into a hydrogen Rydberg atom, respectively.

[0074] FIGS. 15a and 15b depict the electron shell structure of the hydrogen atom, and the excitation of the valence electron of the hydrogen atom into a hydrogen Rydberg atom, respectively.

[0075] FIGS. 16a, 16b and 16c depict the electrostatic attraction between two hydrogen Rydberg atoms, between two potassium Rydberg atoms, and between two hydrogen Rydberg atoms and a potassium Rydberg atom, respectively.

[0076] FIGS. 17a and 17b illustrate a structure of a defect-free crystal, and a structure of a crystal that has a structural defect, respectively.

[0077] FIG. 18 illustrates a defective crystal that has stored a cluster of Rydberg atoms in condensed phase.

[0078] FIG. 19 depicts a flow chart of the nuclear fusion process according to an embodiment of the present invention.

DESCRIPTION OF EMBODIMENTS

[0079] The reaction material of the reaction container comprises metallic material capable of forming interstitial and/or electrically conductive metal hydrides (active hydrogen material), dielectric material possessing electrical polarizability and material promoting the formation and storage of Rydberg matter. The active hydrogen material is preferably in the form of nanopowder capable of enhancing local electric field.

[0080] In an embodiment the polarizable dielectric material comprises a material or materials or a mixture of materials possessing electric polarizability, comprising pyroelectric material, multiferroic material and/or piezoelectric material. The material possessing electric polarizability is preferably in the form of powder or nanoporous material. Electric polarization of the polarizable dielectric material is induced by a controlled polarization factor comprising temperature variation, static or variable magnetic field and/or mechanical vibrations.

[0081] In an embodiment the active hydrogen material comprises generally transition metals that are capable of forming interstitial metal hydrides such as nickel, titanium, zirconium, hafnium, platinum group metals or generally metals that are capable of forming metallic metal hydrides.

[0082] In an embodiment transition metals capable of forming interstitial hydrides that have negative hydrogen ions (hydrides) and/or hydrogen with a metallic bond are utilized in the metallic nanopowder. Negative or positive hydrogen ions are pulled away by the local electric field from the surface of the metallic nanopowder. According to the definition the metallic bonding is based on the electrostatic attractive forces between the delocalized electrons (conduction electrons) and the positively charged metal ions (e.g. hydrogen ions, protons, p⁺). Thus, positively charged hydrogen exists near the surface of the transition metal hydride and that hydrogen ion can be ripped away from the transition metal hydride and accelerated with the strong local electric field until with a noticeable probability it can tunnel through the Coulomb barrier between its nucleus and another nucleus and fuse with that other nucleus releasing fusion energy.

[0083] In an embodiment the active hydrogen material comprises metallic or interstitial hydrides having partly ionic and metallic bond between metal and hydrogen. Examples of interstitial hydrides comprise transition metal hydrides. The said transition metal hydrides are preferably electrically conductive (i.e. they have low electrical resistivity). Electrical conductivity of the metal hydride is beneficial for the present invention for focusing the electric field and enhancing the local electric field strength.

[0084] Common unit for electrical resistivity (resistivity, specific electrical resistance, volume resistivity) is $\mu\Omega\text{cm}$ or Ωm . In an embodiment the resistivity of the active hydrogen material is preferably smaller than about 1000 $\mu\Omega\text{cm}$, more preferably smaller than about 500 $\mu\Omega\text{cm}$, most preferably smaller than about 100 $\mu\Omega\text{cm}$. Common unit for electrical

conductivity (specific conductance) is $S \cdot m^{-1}$. In other words, the active hydrogen material preferably has high electrical conductivity.

[0085] In an embodiment the active hydrogen material comprises hydrogen-storage alloys that are known to be used in nickel-metal hydride secondary batteries. The said hydrogen-storage alloys are optionally doped with traces of a third metal to adjust dissociation pressures and/or temperatures to ranges utilized in the present invention. Transition metal hydrides form also complexes that can be used as the active hydrogen material. Certain transition metal hydride complexes, i.e., interstitial metal hydride complexes, have metallic properties, meaning that they conduct electricity, i.e. they have sufficiently low electrical resistivity (i.e. sufficiently high electrical conductivity).

[0086] In an embodiment the active hydrogen material comprises so called AB_5 and AB_2 hydrogen storage alloys. The said AB_5 hydrogen storage alloys combine a hydride forming metal A, comprising a rare earth metal (La, Ce, Nd, Pr, Y or their mixture), with another element, comprising nickel and/or nickel doped with other metals, such as Co, Sn or Al. The said doping adjusts convenient equilibrium hydrogen pressure and convenient temperature range required for charging and discharging the AB_5 hydrogen storage alloy with hydrogen. The said AB_2 hydrogen storage alloys (Laves phases) comprise alloys containing titanium, zirconium or hafnium at the A-site and a transition metal(s) at a B-site (such as Mn, Ni, Cr and V).

[0087] In an embodiment the active hydrogen material comprises electrically conductive alloys that are known to be hydrogenation catalysts, used in the industry for example for adding hydrogen to organic compounds. Examples of hydrogen catalysts comprise cerium-magnesium alloy $CeMg_2$.

[0088] In an embodiment the active hydrogen material comprises rare earth elements having at least one stable isotope comprising Y, La, Ce, Pr, Nd, Sm, Eu, Gd, Tb, Dy, Ho, Er, Tm, Yb and Lu that are capable of forming rare earth hydrides. Examples of rare earth hydrides comprise lanthanum dihydride LaH_2 , lanthanum trihydride LaH_3 , cerium dihydride CeH_2 , non-stoichiometric cerium hydrides CeH_x , wherein x is a real number up to about 3 and rare earth metal hydrides of Y, Pr, Nd, Sm, Eu, Gd, Tb and Dy with varying metal-hydrogen composition.

[0089] In an embodiment the active hydrogen material comprises material capable of forming binary metal hydride consisting of a metal and hydrogen. Examples of formed binary metal hydrides comprise group 4B metal hydrides comprising titanium hydride TiH_2 , zirconium hydride ZrH_2 and hafnium hydride HfH_2 and group 5B metal hydrides comprising vanadium hydride VH , niobium hydride NbH , niobium dihydride NbH_2 and tantalum hydride TaH and group 8B metal hydrides comprising nickel hydrides NiH_x , wherein x is a real number bigger than 0 and smaller than about 3.

[0090] In an embodiment the active hydrogen material comprises material capable of forming ternary metal hydrides consisting of the first metal, the second metal and hydrogen. Examples of formed ternary metal hydrides comprise stoichiometric ternary hydrides, such as $FeTiH_2$, Mg_2TiH_6 , $MgTi_5H_6$, and nonstoichiometric ternary hydrides, such as $LaNi_5H_{6.7}$.

[0091] In an embodiment the active hydrogen material comprises electrically conductive materials selected from complex transition metal hydrides.

[0092] In an embodiment the hydrogen content of the active hydrogen material is altered and controlled with the hydrogen gas pressure over the metal hydride. For example, increasing the hydrogen gas pressure increases the hydrogen content of the metal hydride.

[0093] Increasing the reaction container temperature increases the number of collisions of gaseous hydrogen atoms and molecules with the solid surfaces of the nanopowders. Increasing the reaction container temperature increases the available thermal activation energy for forming atomic hydrogen. It is easier to ionize atomic hydrogen than molecular hydrogen, because molecular hydrogen is kept intact with the chemical bond between the hydrogen atoms.

[0094] In an embodiment the hydrogen content of the active hydrogen material is altered and controlled with the temperature of the said active hydrogen material. For example, decreasing the temperature increases the hydrogen content of the metal hydride.

[0095] Hydrogen gas is the primary hydrogen source. Primary hydrogen source provides hydrogen to the active hydrogen material.

[0096] In an embodiment the primary hydrogen gas source comprises hydrogen gas bottle that has pure hydrogen gas or a gas mixture having hydrogen gas mixed with other gases comprising nitrogen, helium, argon, neon, xenon, or krypton.

[0097] In an embodiment the primary hydrogen gas source comprises metal hydride that releases hydrogen gas by heating.

[0098] In an embodiment the primary hydrogen gas source comprises hydrogen gas generator based on chemical reactions comprising mixing of an acid comprising H_2SO_4 , HCl , H_3PO_4 , $HCOOH$, CH_3COOH , or a base comprising $NaOH$ or KOH with reactive metals comprising Zn, Al or Mg or reactive metal alloys comprising aluminum activated with gallium or mercury, or based on electrolysis of water or water-containing solutions.

[0099] In an embodiment the primary hydrogen gas source comprises hydrogen generator based on chemical reactions comprising mixing of water with reactive metal alloys comprising aluminum activated with gallium or mercury.

[0100] In an embodiment the primary hydrogen source is an organic compound comprising an alcohol that releases hydrogen gas when it is cracked with moist air in the presence of a catalyst comprising cerium dioxide CeO_2 doped for example with Fe, Co or Ir. Examples of the said alcohol comprise ethanol, isopropanol, n-propanol, n-butanol, sec-butanol, tert-butanol and methanol.

[0101] In an embodiment hydrogen is stored in the form of ammonia NH_3 in metal salts comprising $LiCl$, $LiBr$, LiI , $MgCl_2$, $MgBr_2$, MgI_2 , $CaCl_2$, $CaBr_2$, CaI_2 , $SrCl_2$, $SrBr_2$, SrI_2 , $BaCl_2$, $BaBr_2$, BaI_2 , $MnCl_2$, $MnBr_2$, MnI_2 , $FeCl_2$, $FeBr_2$, FeI_2 , $CoCl_2$, $CoBr_2$, CoI_2 , $NiCl_2$, $NiBr_2$, NiI_2 , $SnCl_2$, $SnBr_2$, and SnI_2 , to form metal ammine salts, preferably $MgCl_2$ to form $Mg(NH_3)_6Cl_2$, and ammonia is released by heating from the metal ammine salts, and released ammonia is cracked into hydrogen and nitrogen gases, and formed hydrogen gas is introduced to the fusion reaction container and utilized for the nuclear fusion processes.

[0102] In an embodiment the fusion process temperature is controlled by at least one means of control selected from the external heating power, mass flow rate of the cooling fluid in the cooling fluid circulation and hydrogen gas pressure of the fusion reaction container.

[0103] In an embodiment the active hydrogen material is formed in situ in the form of metallic nanopowder in the reaction container by decomposing a metal compound nanopowder into a metal oxide nanopowder and reducing the said metal oxide nanopowder into elemental metal or metallic nanopowder. As a non-limiting example, the metal compound nanopowder comprises nickel nitrate or nickel carbonate nanopowder and the decomposition into nickel oxide nanopowder is done by heating the metal compound nanopowder to the decomposition temperature. Further, the said nickel oxide nanopowder is reduced e.g. with hydrogen gas at least partially into elemental nickel metal nanopowder that serves as an example of the metallic nanopowder that is used as the active hydrogen material in the reaction container of the present invention.

[0104] Because molecular hydrogen H_2 and atomic hydrogen H is electrically neutral without an electric charge and electric field does not affect it, it is preferred to create hydrogen ions from hydrogen. Removal the electron from the hydrogen atom creates positive hydrogen ion H^+ (proton), D^+ or T^+ . Addition of electron to the hydrogen atom creates negative hydrogen ion H^- (hydride ion), D^- or T^- . Positive and negative hydrogen ions have electric charge and they can be accelerated in electric field. Positive hydrogen ions (such as H^+ , D^+ , T^+) are accelerated towards the negative pole in the electric field. Negative hydrogen ions (such as H^- , D^- , T^-) are accelerated towards the positive pole in the electric field. The proton (H^+) is the lightest hydrogen ion and it is the easiest hydrogen isotope to accelerate in electric field. Local electric field strength is greatly enhanced with nanotips and/or nanoparticles. Acceleration increases kinetic energy. Near the nanotips and between the nanoparticles the giant electric field accelerates hydrogen ions (protium ion H^+ , hydride ion H^- , deuterium ion D^+ , deuteride ion D^- , tritium T^+ ion or tritide ion T^-) to such high kinetic energies that they can enter other atom nuclei by tunneling through the Coulomb barrier and fuse with the atom nuclei releasing fusion energy.

[0105] Molecular hydrogen has a chemical bond between two hydrogen atoms. It is beneficial to break the said chemical bond by absorbing hydrogen gas into metal that can form metallic or interstitial metal hydride or by activating hydrogen by a transition metal preferably comprising nickel or a platinum group metal.

[0106] In an embodiment the activation of hydrogen is done by metal that can form metallic or interstitial metal hydride, preferably in the form of nanoparticles, inside the reaction container.

[0107] In another embodiment the activation of hydrogen is done by a transition metal or transition metal oxide preferably comprising nickel, nickel oxide, iron, iron oxide or a platinum group metal such as platinum and palladium, preferably in the form of nanoparticles, inside the reaction container.

[0108] The equilibrium of the reaction equation $MH_x \rightleftharpoons M + x/2 H_2$ depends on the surrounding temperature and pressure. At low pressure more hydrogen gas H_2 is released from the metal hydride MH_x . At high pressure more hydrogen is bound to the metal forming more metal hydride MH_x . At high temperature more hydrogen gas H_2 is released from the metal hydride MH_x . At low temperature more hydrogen is bound to the metal forming more metal hydride MH_x .

[0109] In an embodiment of the present invention a small amount of paracrystalline material doped with an element capable of forming Rydberg matter is added to the mixture of a dielectric material possessing electrical polarizability and

an element capable of forming metallic metal hydride to promote nuclear fusion in solid state matter and on surface of solid state matter.

[0110] In an embodiment of the present invention paracrystalline material utilized as a template for forming and storing Rydberg atom clusters and inverted Rydberg atom clusters in lattice defects of the said paracrystalline material comprises a metal oxide mixture made from the first metal oxide and the second metal oxide, wherein the metal of the first metal oxide is capable of changing its oxidation state in reducing atmosphere and the metal of the second metal oxide is stable and does not change its oxidation state in reducing atmosphere, the second metal oxide being a structural promoter that maximizes and stabilizes the number of lattice defects in the paracrystalline material.

[0111] In an embodiment of the present invention paracrystalline material utilized as a template for forming and storing Rydberg atom clusters and inverted Rydberg atom clusters in lattice defects of the said paracrystalline material comprise nickel mixed with alumina and/or chromia, nickel oxide mixed with alumina and/or chromia, iron mixed with alumina and/or chromia, iron oxide mixed with alumina and/or chromia and copper-zinc alloy mixed with alumina and/or chromia.

[0112] In an embodiment of the present invention paracrystalline material utilized for forming and storing Rydberg atom clusters and inverted Rydberg atom clusters is doped with an element that possesses Rydberg states due to the excitation of an electron of the element and is capable of becoming a Rydberg atom, the element comprising Li, Na, K, Rb, Cs, N, Ni, Ag, Cu, Pd, Ti and Y.

[0113] The electron of a hydrogen atom is excited from the ground energy level by the collision of accelerated electrons and protons to a higher energy level that is a Rydberg state. The energy that the colliding electron or proton must donate to the ground state electron of a hydrogen atom is only about 10.2-13.6 eV. In an embodiment electrons or protons are at least part of the operating time of the thermal-energy producing system accelerated to about 10-20 eV kinetic energy, preferably to a kinetic energy below the amount of energy required for ionizing hydrogen atom to a separate proton and a separate electron with small or moderate electric fields in small gaps between the powder particles to create hydrogen Rydberg atoms on a surface.

[0114] In an embodiment electrons or protons are at least part of the operating time of the thermal-energy producing system accelerated to about 10-100 eV kinetic energy to destabilize Rydberg matter and inverted Rydberg matter by ionization of the said matter with accelerated electrons or protons to induce Coulomb explosion in the said matter.

[0115] In an embodiment electrons of hydrogen atoms on a surface with structural defects are excited to Rydberg states with deep UV light (UVC light).

[0116] In an embodiment of the present invention styrene catalyst is utilized for enhancing nuclear fusion in a solid state system. The precursor for the styrene catalyst, hematite Fe_2O_3 , having corundum crystal structure is reduced with hydrogen gas into magnetite Fe_3O_4 . The precursor (iron oxide) is doped with alkali metal hydroxide comprising lithium hydroxide $LiOH$, sodium hydroxide $NaOH$, potassium hydroxide KOH , rubidium hydroxide $RbOH$ and/or cesium hydroxide $CsOH$ or with alkali metal oxide comprising lithium oxide Li_2O , sodium oxide Na_2O , potassium oxide K_2O , rubidium oxide Rb_2O and/or cesium oxide Cs_2O . The

alkali metal hydroxide is preferably KOH and the alkali metal oxide is preferably K_2O . Textural promoters comprising alumina Al_2O_3 and/or chromia Cr_2O_3 are added to the iron oxide. The said textural promoters are stable in process conditions in hot, highly reducing environment and they prevent the loss of lattice defects that are necessary for storing Rydberg matter and inverted Rydberg matter.

[0117] In an embodiment the typical precursor composition of the styrene catalyst applicable for the present invention comprises about 80-95 wt % Fe_2O_3 , preferably 88 wt % Fe_2O_3 , about 5-15 wt % K_2O , preferably 10 wt % K_2O , and about 1-4 wt % Al_2O_3 or Cr_2O_3 , preferably 2 wt % Al_2O_3 or Cr_2O_3 . This mixture is reduced with hydrogen gas into Fe_3O_4 :K, Al_2O_3 and Fe_3O_4 :K, Cr_2O_3 , known as styrene catalyst materials in the chemical industry. The metal oxide, in this embodiment reduced iron oxide (Fe_3O_4), adopts a new crystal structure, in this embodiment inverse spinel. Structural promoters (e.g. Al_2O_3 , Cr_2O_3) that cannot be reduced keep their original crystal structure (corundum) and induce strain and lattice defects to the inverse spinel iron oxide lattice. Styrene catalysts comprising Fe_3O_4 :K, Al_2O_3 and Fe_3O_4 :K, Cr_2O_3 are utilized in the present invention. In case of styrene catalysts it is assumed that a combination of lattice defects on catalyst particle surfaces and an element capable of forming Rydberg atoms, such as potassium, promotes the formation of condensed Rydberg matter, which enhanced solid state nuclear fusion in the present invention.

[0118] In an embodiment of the present invention ammonia synthesis catalyst is utilized for enhancing nuclear fusion in a solid state system. The precursor for the ammonia synthesis catalyst, iron oxide, typically magnetite Fe_3O_4 , has inverse spinel crystal structure that changes into body-centered or face-centered cubic crystal structure when Fe_3O_4 is reduced with hydrogen gas into elemental iron Fe. The precursor (iron oxide) is doped with alkali metal hydroxide comprising lithium hydroxide LiOH, sodium hydroxide NaOH, potassium hydroxide KOH, rubidium hydroxide RbOH and/or cesium hydroxide CsOH or with alkali metal oxide comprising lithium oxide Li_2O , sodium oxide Na_2O , potassium oxide K_2O , rubidium oxide Rb_2O and/or cesium oxide Cs_2O . The alkali metal hydroxide is preferably KOH and the alkali metal oxide is preferably K_2O . Textural promoters comprising alumina Al_2O_3 and calcium oxide CaO are added to the iron oxide. The said textural promoters are stable in process conditions in hot, highly reducing environment and they prevent the sintering of elemental iron.

[0119] In an embodiment typical composition of the ammonia synthesis catalyst suitable for the present invention comprise, before reducing the iron oxide, about 90-95 wt % Fe_3O_4 , preferably 93 wt % Fe_3O_4 , about 0.1-2 wt % K_2O , preferably 1 wt % K_2O , about 2-4 wt % Al_2O_3 , preferably 3 wt % Al_2O_3 , and about 2-4 wt % CaO, preferably 3 wt % CaO. After reducing Fe_3O_4 into elemental iron, this type of ammonia synthesis catalyst Fe: K_2O , Al_2O_3 ,CaO, preferably crushed into about 10-100 nm powder, has a lot of lattice defects and it is suggested herein that the said catalyst promotes efficiently the formation of potassium and hydrogen Rydberg atoms and, further, the formation of condensed Rydberg matter leading to enhanced rate of nuclear fusion in the reaction container of the present invention.

[0120] In an embodiment of the present invention high temperature water gas shift catalysts comprising potassium doped iron oxide Fe_xO_{3-x} :K and potassium doped lanthanum niobate $LaNiO_3$:K are utilized as Rydberg matter hatchery

(formation and storage of Rydberg atoms) for enhancing nuclear fusion in a solid state system.

[0121] In an embodiment of the present invention Fischer-Tropsch catalysts comprising metals and metal oxides of cobalt (Co, $Co_{1-x}O$), iron (Fe, $Fe_{1-x}O$), ruthenium (Ru, RuO_2 , RuO_{2-x}) and nickel (Ni, $Ni_{1-x}O$) doped with copper or group 1 alkali metals (Li, Na, K, Rb, Cs) are utilized as Rydberg matter hatchery for enhancing nuclear fusion in a solid state system within the reaction container.

[0122] In an embodiment of the present invention hydrogenation catalysts comprising platinum, palladium, rhodium, ruthenium, alloys of Pt, Pd, Rh and Ru, Raney nickel, Urushibara nickel and alkali metal doped nickel oxide, preferably Ni_2O_3 and non-stoichiometric $Ni_{1-x}O$ doped with alkali metal, preferably potassium, wherein x is a non-integer in a range of about 0.005-0.1, preferably about 0.02, are utilized as Rydberg matter hatchery for enhancing nuclear fusion in a solid state system within the reaction container.

[0123] Industrial catalysts have been optimized for specific chemical processes. For example, formation of coke (solid carbonaceous material) on the catalyst surface is avoided if the process temperature is kept in a specified temperature range. The present invention does not utilize compounds that form coke and temperatures above the normal temperature range for catalytic processes can be used in the present thermal-energy producing reactor.

[0124] The probability for obtaining nuclear fusion near a single structural defect of a material is very small. Arranging a very large number of particles with surface and lattice defects to the reaction container increases the probability for nuclear fusion events per time unit within the reaction container to a noticeable and useful level. For example, if a 50 g piece of nickel is converted into 5 nm Ni nanoparticles with about 6000 atoms, about $8.55 \cdot 10^{19}$ Ni nanoparticles is obtained. Each Ni nanoparticle may be in contact with a catalyst nanoparticle that promotes the formation of Rydberg atoms and clusters. Even a very small probability for obtaining nuclear fusion near a single Ni nanoparticle becomes considerable and useful when all the $8.55 \cdot 10^{19}$ probabilities are added together.

[0125] In an embodiment of the present invention the reaction material used for the solid state nuclear fusion reactions is made by mixing dielectric material possessing electrical polarizability (the first process material) with material capable of forming interstitial metal hydrides and/or electrically conductive metal hydrides (the second process material) and with material capable of forming and storing Rydberg matter and inverted Rydberg matter (the third process material). The mixing ratio of the first, the second and the third process material may be varied in a wide range selected from 0-100 wt %.

[0126] In an embodiment the reaction material used for the solid state nuclear fusion reactions may contain about 5-80 wt % of the first process material, about 15-90 wt % of the second process material and about 1-10 wt % of the third process material.

[0127] The present invention includes the surprising finding that increasing the pressure of the reaction container with hydrogen gas increases the heat production rate to such a high value that chemical reactions (for example burning hydrogen gas with oxygen into water) are not capable of producing as much thermal energy. Increasing the pressure increases the rate of hydrogen molecule collisions with the surfaces. Nano-

particles have very high surface area and the number of hydrogen molecule collisions with the surface is very high.

[0128] The amount of thermal energy released from the reaction container is so far above the amount of energy released by any known chemical reaction that non-binding afterwards interpretation of the possible reactions in the reaction container and suggestions for a theory explain the possible reactions in the reaction container are presented herein, not to be negatively affecting to the novelty of the present invention and not to make the present invention obvious.

[0129] Without restricting to a specific theory to explain the production of thermal energy, it is herein suggested that the benefit of increasing the pressure is somehow related to the formation of high electric field strength. For example, hydrogen ions form plasma and plasma formed at high pressure possesses smaller Debye length than plasma formed at low pressure. Localized space charge regions may build up large potential drops (electric double layers) over distances of about ten Debye lengths. It is also herein suggested that the presence of very high dielectric constant material possessing electric polarizability makes it possible to decrease the Debye length down to the nanometer range in the system of the present invention. Very large potential drop over a very short distance (in other words very steep voltage gradient) leads to extremely high electric field strength that is capable of accelerating ions to high kinetic energies.

[0130] The present invention also includes the finding that increasing the temperature of the reaction container increases the fusion rate. Increasing the temperature increases the rate of hydrogen molecule collisions to the surfaces. Although the increasing temperature decreases the amount of metal hydride, higher temperature provides more high energy photons because of the strengthened thermal radiation within the pores of the porous material and between the particles in the powder. Without restricting to a specific theory to explain the production of thermal energy, it is herein suggested that photons interact with surface plasmons on the surface of the metallic nanopowder forming polaritons that proceed along the surface of the metallic nanopowder and further enhance the electric field strength especially near tips and sharp edges of the metallic nanopowder particles. Thus, increased pressure is utilized for keeping some of the hydrogen in the form of metal hydride and increased temperature is utilized for providing thermal activation energy for breaking molecular H_2 into atomic H and for forming enhanced thermal radiation for creating polaritons. As already disclosed hereinbefore, materials especially good for activating molecular hydrogen into atomic hydrogen on the surface of said materials comprise nickel, nickel oxide, and platinum group metals such as platinum and palladium.

[0131] Increasing the temperature of the reaction container shortens the wavelength of the heat radiation emitting from all hot surfaces. These photons are mostly in the infrared wavelength range when the temperature is below $500^\circ C.$, but more photons are emitted in the visible light wavelength range when the temperature goes above $500^\circ C.$ These photons can be utilized for exciting surface plasmons.

[0132] In an embodiment the reaction container with the reaction material operates under external control and the temperature of the reaction container with the reaction material is kept in a temperature range of about $100-1200^\circ C.$ during the generation of heat energy, preferably at about $300-900^\circ C.$ and more preferably at about $400-700^\circ C.$

[0133] The thermal-energy generating system of the present invention is provided with novel safety features. In an embodiment means of heating the reaction container with external power is provided for bringing the reaction container (or reaction cartridge) to the operating temperature and for providing external control of the reaction container temperature based on the feedback from the temperature measurements of the reaction container. External power is used for heating, for example, a heater cartridge placed inside the reaction container or placed to the wall of the reaction container.

[0134] In another embodiment the dielectric material possessing electrical polarizability provides an intrinsic safety feature, meaning that when the temperature of the said dielectric material increases above the Curie temperature of the said dielectric material, the said dielectric material loses polarization and, as a consequence, hydrogen ion acceleration stops, fusion of hydrogen nucleus with other nuclei stops, the generation of thermal energy (or heat energy) stops and the temperature of the said dielectric material cannot any longer increase to a higher value.

[0135] In still another embodiment the sinterability of the nanoparticles provides an intrinsic safety feature, meaning that at temperatures above the normal operation temperature of the reaction container the nanoparticles of the active hydrogen material start to sinter together forming so few large particles that the probability of fusion reactions decreases to a negligible value and heat generation based on fusion reactions stops permanently.

[0136] In still another embodiment the melting point of the nanoparticles provides an intrinsic safety feature, meaning that above the normal operation temperature of the reaction container the nanoparticles of the active hydrogen material reach the melting point and the said nanoparticles form large droplets of material that cannot sustain fusion reactions. As a result, surface area collapses to a low value, enhancement of the local electric field is lost, the probability of the fusion of hydrogen nucleus with other nuclei drops to a negligible value and generation of thermal energy based on fusion reactions stops permanently.

[0137] The surroundings of the reaction container is shielded with a heavy metal mantle (e.g. lead) for converting gamma and X-ray radiation into heat and with an optional neutron grabber mantle for stopping free neutrons. Free protons, electrons and alpha particles have short absorption depth in normal construction materials, such as steel. Walls of the reaction container made of a durable material preferably comprising steel are utilized for stopping free protons, electrons and alpha particles.

[0138] In an embodiment thermal energy generators of the present invention are clustered to increase the amount of produced thermal energy. Clusters of thermal energy generators comprise thermal energy generator units that produce more than about 1 kW/unit or more than about 5 kW/unit or more than about 10 kW/unit or more than about 25 kW/unit or more than about 50 kW/unit to produce up to multi MW or more of thermal energy power in clusters.

[0139] Referring now to the prior art published by Dr. B. K. Juluriat at the <http://juluribk.com/2011/04/09/electric-field-in-metal-nanoparticle-dimers/web> site, in FIG. 1 there are shown simulated electric field strengths obtainable between very small pieces of material. The smaller the distance between said pieces of material is the stronger is the electric field. The bottom right-hand side drawing depicts a 2-nm gap

between two rectangles. The electric field strength is enhanced by a factor of 10^4 (i.e. 10000) in the said gap. Comparing the left-hand side drawings of spheres and right-hand side drawings of rectangles of FIG. 1 it can be seen that the geometry of the pieces of material affects the electric field strength. Pieces of material with sharp edges enhance the electric field strength more than pieces of material with round shape.

[0140] Referring now to the prior art in FIG. 2, there is shown a metallic tip having an apex diameter of 10 nm and separated by a 2-nm gap from a surface. Surface plasmons coupled with the electromagnetic radiation (e.g. visible light or infrared light) enhance the electric field strength by up to a factor of 10^{11} (i.e. 100000000000) in the gap.

[0141] To clarify the term "surface plasmon", according to Wikipedia from the web site <http://en.wikipedia.org/wiki/Plasmon> "Surface plasmons are those plasmons that are confined to surfaces and that interact strongly with light resulting in a polariton. They occur at the interface of a vacuum or material with a positive dielectric constant, and a negative dielectric constant (usually a metal or doped dielectric)."

[0142] Referring now to the invention in more detail, in FIG. 3 there is shown a system 300 for producing and utilizing thermal energy. The system 300 comprises a reaction container system 301, a control system 304, a hydrogen source 306 and a secondary heat exchange unit 314.

[0143] In more detail, still referring to the embodiment of FIG. 3, the reaction container system 301 comprises reaction material 320 in a specified form, such as powder material or porous material, a heater cartridge 322 with optional heat conducting extensions 324 for distributing heat from the heater cartridge 322 to the reaction material 320, an external power line 326 connected to the heater cartridge 322, a first temperature measurement system 328 for measuring the temperature of the solid reaction material 320, a cooling fluid circulation in a mantle 330 with optional heat conducting extensions 332 for collecting and removing thermal energy from the reaction container system 301, and a second temperature measurement system 334 for measuring the temperature of the cooling fluid. Optionally, a third temperature measurement system 312 is used for measuring the external temperature for verifying that the reaction container system 301 is operating normally and thermal energy does not excessively leak to the surroundings.

[0144] Still referring to the invention of FIG. 3, the hydrogen gas source 306 comprises a housing 307 for storing and/or generating hydrogen gas, a gas control valve 308 in fluid communication with the hydrogen gas source 306 for dosing hydrogen gas from the hydrogen gas source 306 via a gas conduit 309 to the reaction container 350, a pressure measurement system 313 for measuring the gas pressure of the gas conduit 309, a surplus fluid reservoir 310 equipped with a flow control valve 311 for draining surplus hydrogen gas from the gas conduit 309 and the reaction container 350. Still referring to the embodiment of FIG. 3, the secondary heat exchange unit 314 comprises a cooling fluid circulation pump 319 in the cooling fluid conduit 316 in fluid communication with the secondary heat exchange unit 314 and the reaction container system 301.

[0145] Still referring to the embodiment of FIG. 3, the control system 304 receives input from the temperature measurement systems 328, 334 and the pressure measurement system 313 and produces control output to the valves 308, 311, and to the cooling fluid circulation pump 319.

[0146] In further detail, still referring to the embodiment of FIG. 3, the reaction container 350 is filled with the reactive material 320. The reaction container system 301 is attached to cooling fluid circulation that transfers cooling fluid between the cooling fluid mantle 330 and the secondary heat exchange unit 314 along at least two cooling fluid conduits. The inlet cooling fluid conduit 316 transports cooled cooling fluid from the secondary heat exchange unit 314 to the cooling fluid mantle 330, and the outlet cooling fluid conduit 317 returns heated cooling fluid to the secondary heat exchange unit 314. The amount of cooling fluid flowing in the cooling fluid circulation is affected by the cooling fluid circulation pump 319 and controlled by the control system 304. The reaction container 350 is attached to the gas conduit 309 that is in controlled fluid communication with the hydrogen gas source 306. The control of the hydrogen gas flow and pressurization of the reaction container (pressure vessel) 350 is performed with the gas control valve (hydrogen valve) 308 that is preferably a normally-closed valve that closes automatically in case of power failure. The hydrogen valve 308 is controlled with the control system 304. When it is necessary to increase the pressure of the pressure vessel, the control system opens the hydrogen valve 308, monitors the pressure of the gas conduit and closes the hydrogen valve 308 when the pressure reading obtained by the control system 304 from the pressure measurement system 313 shows that the target pressure range, e.g. 20-21 bar (gauge), has been reached.

[0147] The system is controlled by the heater cartridge 322 power, e.g. at maximum of 50 W, 100 W, 500 W or 1 kW heating power level, or with hydrogen gas pressure preferably above room pressure, more preferably 1 barg, 5 barg, 10 barg or 20 barg and/or by cooling fluid circulation with a flow rate of e.g. 1 liters per minute (lpm), 2 lpm, 5 lpm, 10 lpm or 50 lpm.

[0148] In an embodiment heat energy transfer is implemented with a closed loop primary coolant system arranged to receive heat energy from the reaction container system (primary heat exchanger) and a secondary coolant system arranged to receive heat energy from the closed loop primary coolant system (secondary heat exchanger). Fluid circulation in the primary heat exchange unit removes heat from the reaction container. Primary fluid is directed to the secondary heat exchange unit that has secondary fluid circulation. Heat energy transferred to the secondary fluid is utilized for heating or for generating electricity e.g. with a generator based on Rankine cycle. In an embodiment the secondary heat exchanger 314 has an electric generator 318 based on closed-loop Rankine cycle.

[0149] In an embodiment the primary fluid is directly utilized in the generator based on Rankine cycle. Rankine cycle is a hermetically sealed closed loop system meaning that the primary fluid never leaves the primary fluid circulation and can be safely used for generating electricity, which simplifies the construction of the electric generator. Hot liquid primary fluid vaporizes, the vapor powers a turbine, the turbine with a mechanical coupling with the electricity generator produces electricity, vapor is condensed to liquid after the turbine and pumped back to the primary heat exchanger inside the reaction container system.

[0150] Electricity is used for pre-heating the reaction container and controlling the system and thermal energy based on nuclear fusion is taken out of the reaction container, so that coefficient of performance COP is more than 1, preferably more than 5, still more preferably more than 10, most prefer-

ably more than 20. Part of the generated electricity is used for operating and controlling the heat generator.

[0151] Replacement reaction container system comprises valves, fittings, a heating cartridge and thermocouples. Depleted reaction container system is removed from the thermal energy generator and a new reaction container system is attached to the thermal energy generator in the field.

[0152] The construction details of the embodiment as shown in FIG. 3 are that the pressure vessel 350 may be made

radiation (helium nuclei) is converted into thermal energy by collisions between alpha particles and the surrounding material. Kinetic energy of the beta radiation (high speed electrons) is converted into thermal energy by collisions between beta radiation and the surrounding material. Generated thermal energy is utilized for heating gases or liquids. Removal of heat from the structures near the cartridge is arranged for example by flowing gas or liquid. Electricity is generated from said heated gases or liquids.

TABLE 1

Selected isotopes, isotope concentrations (%) and absorption cross sections for 2200 m/s neutrons (barn)									
³ He	0.00014	5333	In	100	193.8	¹⁷⁴ Yb	31.8	69.4	
Li	100	70.5	¹¹⁵ In	95.7	202	Lu	100	74	
⁶ Li	7.5	940	Xe	100	23.9	¹⁷⁵ Lu	97.39	21	
⁷ Li	92.5	0.0454	¹²⁹ Xe	26.4	21	¹⁷⁶ Lu	2.61	2065	
B	100	767	¹³⁰ Xe	4.1	26	Hf	100	104.1	
¹⁰ B	20	3835	¹³¹ Xe	21.2	85	¹⁷⁶ Hf	5.2	23.5	
¹¹ B	80	0.0055	¹³³ Cs	100	29.0	¹⁷⁷ Hf	18.6	373	
Cl	100	33.5	¹⁴¹ Pr	100	11.5	¹⁷⁸ Hf	27.1	84	
³⁵ Cl	75.77	44.1	Nd	100	50.5	¹⁷⁹ Hf	13.7	41	
⁴³ Sc	100	27.5	¹⁴² Nd	27.16	18.7	¹⁸⁰ Hf	35.2	13.04	
Ti	100	6.09	¹⁴³ Nd	12.18	337	Ta	100	20.6	
⁴⁸ Ti	73.8	7.84	¹⁴⁵ Nd	8.29	42	¹⁸¹ Ta	99.988	20.5	
Cr	100	3.05	Sm	100	5922	W	100	18.3	
⁵⁰ Cr	4.35	15.8	¹⁴⁷ Sm	15.1	57	¹⁸² W	26.3	20.7	
⁵³ Cr	9.5	18.1	¹⁴⁹ Sm	13.9	42080	¹⁸³ W	14.3	10.1	
⁵⁵ Mn	100	13.3	¹⁵⁰ Sm	7.4	104	¹⁸⁶ W	28.6	37.9	
⁵⁹ Co	100	37.18	¹⁵² Sm	26.6	206	Re	100	89.7	
Ni	100	4.49	Eu	100	4530	¹⁸⁵ Re	37.4	112	
⁵⁸ Ni	68.27	4.6	¹⁵¹ Eu	47.8	9100	¹⁸⁷ Re	62.6	76.4	
⁶⁰ Ni	26.1	2.9	¹⁵³ Eu	52.2	312	Os	100	16	
⁶¹ Ni	1.13	2.5	Gd	100	49700	¹⁸⁶ Os	1.58	80	
⁶² Ni	3.59	14.5	¹⁵⁴ Gd	2.1	85	¹⁸⁷ Os	1.6	320	
⁶⁴ Ni	0.91	1.52	¹⁵⁵ Gd	14.8	61100	¹⁸⁹ Os	16.1	25	
Se	100	11.7	¹⁵⁷ Gd	15.7	259000	¹⁹⁰ Os	26.4	13.1	
⁷⁴ Se	0.9	51.8	¹⁵⁹ Tb	100	23.4	Ir	100	425	
⁷⁶ Se	9	85	Dy	100	994	¹⁹¹ Ir	37.3	954	
¹⁰³ Rh	100	144.8	¹⁶⁰ Dy	2.34	56	¹⁹³ Ir	62.7	111	
Pd	100	6.9	¹⁶¹ Dy	19	600	Pt	100	10.3	
¹⁰² Pd	1.02	3.4	¹⁶² Dy	25.5	194	¹⁹⁰ Pt	0.01	152	
¹⁰⁴ Pd	11.14	0.6	¹⁶³ Dy	24.9	124	¹⁹² Pt	0.79	10.0	
¹⁰⁵ Pd	22.33	20	¹⁶⁴ Dy	28.1	2840	¹⁹⁴ Pt	32.9	1.44	
¹⁰⁶ Pd	27.33	0.304	¹⁶⁵ Ho	100	64.7	¹⁹⁵ Pt	33.8	27.5	
¹⁰⁸ Pd	26.46	8.55	Er	100	159	¹⁹⁶ Pt	25.3	0.72	
¹¹⁰ Pd	11.72	0.226	¹⁶⁴ Er	1.56	13	¹⁹⁸ Pt	7.2	3.66	
Ag	100	63.3	¹⁶⁶ Er	33.4	19.6	¹⁹⁷ Au	100	98.65	
¹⁰⁷ Ag	51.83	37.6	¹⁶⁷ Er	22.9	659	Hg	100	372.3	
¹⁰⁹ Ag	48.17	91.0	¹⁶⁹ Tm	100	100	¹⁹⁹ Hg	17	2150	
Cd	100	2520	Yb	100	34.8	²⁰⁰ Hg	23.1	60	
¹¹⁰ Cd	12.51	11	¹⁷⁰ Yb	3.06	11.4	U	100	7.57	
¹¹¹ Cd	12.81	24	¹⁷¹ Yb	14.3	48.6	²³⁵ U	0.72	680.9	
¹¹³ Cd	12.22	20600	¹⁷³ Yb	16.1	17.1				

of any sufficiently strong material such as metal, and the like. The radiation shield 302 may be made of any appropriate material 336 that stops specified radiation, such as lead in case gamma radiation must be stopped and high neutron absorption cross section material selected from Table 1 in case neutron radiation must be stopped.

[0153] Gamma and X-ray radiation is stopped and its energy is converted into thermal energy by a heavy metal shield (e.g. lead). Neutron radiation is absorbed and its released energy is converted into thermal energy by a material that has high neutron capture cross section e.g. elements or chemical compounds of lithium or boron. Kinetic energy of any released proton radiation is converted into thermal energy with the surrounding material. Kinetic energy of the alpha

[0154] Best neutron absorption is obtained with extremely high absorption cross section (more than 5000 barns) element comprising gadolinium Gd, samarium Sm.

[0155] Very good neutron absorption is obtained with very high absorption cross section (500-5000 barns) element comprising boron B, cadmium Cd, europium Eu and dysprosium Dy. Boron compounds are cheap and they are often preferred as neutron absorbing materials.

[0156] Good neutron absorption is obtained with high absorption cross section (50-500 barns) element comprising lithium Li, rhodium Rh, silver Ag, indium In, neodymium Nd, erbium Er, thulium Tm, lutetium Lu, hafnium Hf, rhenium Re, iridium Ir, gold Au and mercury Hg.

[0157] Referring now to the embodiment in FIG. 4, there is shown a reaction space 400 for generating heat 402. The

reaction system 400 comprises a reaction container 350 with optional heat collecting protrusions or heat conducting extensions 332, the said reaction container 350 being filled with reaction material 320 and pressurized with hydrogen gas, a heater cartridge 322 with optional heat distributing protrusions 324 and a power cable 404 and a thermocouple 406. Regarding commercial products, the customer receives the replacement reaction container 350 pre-filled with the required reaction material, and the pressurization of the replacement reaction container 350 with hydrogen-containing gas is preferably done after attaching the filled replacement reaction container 350 to the heat generating system.

[0158] In further detail, still referring to the embodiment of FIG. 4, the heater cartridge 322 receives electrical power along the power cable 404 and converts the electrical power into thermal energy that heats the reaction material 320 to a temperature that promotes reactions within the reaction material 320. The temperature of the reaction container 350 during the fusion reactions is more than about 0° C., preferably more than 150° C., more preferably more than 250° C. and most preferably about 350-600° C. In an embodiment, the temperature of the reaction container is preferably below the Curie temperature of the pyroelectric, piezoelectric or multiferroic material, because above the Curie temperature the polarization of the material may be lost. Depending on the material the Curie temperature can be up to 1000° C. or even higher.

[0159] Heating the reaction container 350 of FIG. 4 to high enough temperature produces certain beneficial effects. All matter emits electromagnetic radiation, the power of the radiation and the wavelength of the radiation at the maximum intensity depending on the temperature of the matter. The hotter the matter is, the higher the emitted power P is, as stated by the Stefan-Boltzmann law modified with the grey body emissivity $P = \epsilon \cdot \sigma \cdot A \cdot T^4$, wherein ϵ is the emissivity factor of the emitting surface, σ is Stefan-Boltzmann constant $5.670400 \times 10^{-8} \text{ W} \cdot \text{m}^{-2} \cdot \text{K}^{-4}$, A is the surface area of the emitting surface and T is the temperature of the emitting surface in kelvins. Also, the hotter the matter is the shorter the wavelength of the maximum intensity λ_{max} is, as stated by the Wien's law $\lambda_{max} = b/T$, wherein b is Wien's displacement constant $2.8977685 \times 10^{-3} \text{ m} \cdot \text{K}$ and T is the temperature of the emitting surface in kelvins. The wavelength of the emission maximum is in the infrared region near room temperature and moves towards visible light when the temperature of the emitting surface increases.

[0160] Still referring to the embodiment of FIG. 4, the reaction container 350 is pressurized with gas that comprises hydrogen gas and optional diluting gases such as nitrogen, helium, neon, argon, krypton or xenon or their mixtures, that decrease the concentration of hydrogen gas. Pure hydrogen gas is preferred for the pressurization. Hydrogen gas comprises at least one hydrogen isotope selected from protium H that has a proton, deuterium D that has a proton and a neutron, and tritium T that has a proton and two neutrons. After pressurization the pressure of the reaction container 350 is more than about 1 bar (gauge), preferably more than 5 bar (gauge), more preferably more than 10 bar (gauge) and most preferably about 15-30 bar (gauge), although even higher pressures are applicable. Dilution of hydrogen gas is beneficial in case the reactive materials selected for the fusion reactions tend to form an unstable heat generating system with pure hydrogen gas at high temperatures.

[0161] Various methods exist to arrange hydrogen gas to the thermal-energy producing system 300. A non-exhaustive

list of hydrogen gas sources comprise pressurized hydrogen gas bottle, metal hydrides heated to release hydrogen gas, chemical reactions releasing hydrogen gas, comprising chemical reactions between an acid and a metal (e.g. zinc, aluminum, magnesium or iron) forming a metal salt and hydrogen gas, comprising chemical reactions between a base and a metal (e.g. zinc or aluminum) forming a metal salt and releasing hydrogen gas and generating hydrogen from metal ammine salts.

[0162] In an embodiment the acid comprises sulfuric acid H_2SO_4 , hydrochloric acid HCl , acetic acid CH_3COOH or formic acid HCOOH , and the base comprises sodium hydroxide NaOH or potassium hydroxide KOH .

[0163] In an embodiment aluminum metal activated with mercury or gallium reacts with water and releases hydrogen gas.

[0164] Still referring to the embodiment of FIG. 4, the reaction material 320 comprises preferably coated porous material and/or powder material. The coated porous material comprises preferably porous crystalline material and metallic nanoparticles grown on the pore surfaces of the porous crystalline material. The porous crystalline material preferably comprises a compound or a mixture of compounds that possess electrical polarizability, i.e. are capable of having areas with a positive electrical charge (+) and areas with a negative electrical charge (-). Metallic nanoparticles comprise preferably metallic matter that is capable of forming electrically conductive metal hydrides, more preferably electrically conductive transition metal hydrides. In an embodiment metal nanoparticles are grown to the inner surfaces of pores of the porous dielectric material by thin film deposition methods comprising physical vapor deposition PVD, chemical vapor deposition CVD, atomic layer chemical vapor deposition ALCVD, atomic layer deposition ALD or molecular layer deposition MLD, preferably by the ALCVD, ALD or MLD method that are based on sequential self-saturating surface reactions capable of coating inner surfaces of pores of porous materials with uniform layer of nanoparticles, such as nickel nanoparticles. Catalytic material promoting the formation and storage of Rydberg matter and inverted Rydberg matter is added to the pore surfaces of the porous material by the said thin film deposition methods or by electrochemical methods.

[0165] The powder material in the reaction material 320 preferably comprises a mixture of a first reaction material being a compound or a mixture of compounds that possess electrical polarizability, i.e. are capable of having areas with a positive electrical charge (+) and areas with a negative electrical charge (-), a second reaction material that is metallic nanoparticles comprising preferably metallic matter that is capable of forming electrically conductive metal hydrides, more preferably electrically conductive transition metal hydrides, and a third material that is nanoparticles capable of promoting the formation and storage of Rydberg matter and inverted Rydberg matter. Method for inducing electric polarization comprise applying an electric field to multiferroic materials, applying mechanical stress or mechanical vibration to piezoelectric materials and applying temperature changes to pyroelectric materials.

[0166] Multiferroic materials preferred for the said porous crystalline material in the reaction material 320 and for the said first reaction material in the reaction material 320 utilized in the present embodiment comprise perovskite transition metal oxides, such as rare earth manganites YMnO_3 , HoMnO_3 , TbMnO_3 , HoMn_2O_5 , rare earth ferrites such as

LuFe₂O₄, bismuth ferrite BiFeO₃, bismuth manganite BiMnO₃, geometric ferroelectrics, such as BaNiF₄, BaCoF₄, BaFeF₄, BaMnF₄, spinel chalcogenides, such as ZnCr₂Se₄, boracites, such as Ni₃B₇O₁₃I, Ni₃B₇O₁₃Cl, Co₃B₇O₁₃I, doped multiferroics, e.g. Pb(Fe_{2/3}W_{1/3})O₃, Pb(Fe_{0.5}Nb_{0.5})O₃, terbium manganites TbMnO₃, TbMn₂O₅, nickel vanadate Ni₃V₂O₈, copper ferrite CuFeO₂, cobalt chromite CoCr₂O₄ and lutetium ferrite LuFe₂O₄.

[0167] Piezoelectric materials suitable for the said porous crystalline material in the reaction material 320 and for the said first reaction material in the reaction material 320 utilized in the present embodiment comprise materials selected from piezoelectric crystal classes 1, 2, m, 222, mm2, 4, -4, 422, 4 mm, -42 m, 3, 32, 3 m, 6, -6, 622, 6 mm, -62 m, 23 and -43 m. Examples of materials belonging to said piezoelectric crystal classes and suitable for the utilization in the present embodiment comprise quartz SiO₂.

[0168] Pyroelectric materials suitable for the said porous crystalline material in the reaction material 320 and for the said first reaction material in the reaction material 320 utilized in the present embodiment comprise materials selected from pyroelectric crystal classes 1, 2, m, mm2, 3, 3 m, 4, 4 mm, 6 and 6 mm. Examples of materials belonging to said pyroelectric crystal classes and suitable for the utilization in the present embodiment comprise quartz SiO₂.

[0169] Piezoelectric and/or pyroelectric minerals suitable for the said porous crystalline material in the reaction material 320 and for the said first reaction material in the reaction material 320 utilized in the present embodiment comprise awillite, alunite, aminoffite, analcime, bastnasite-(Ce), batisite, bavenite, bertrandite, boracites, bromellite, brucite, brushite, buergerite, burbankite, caledonite, clinohedrite, colemanite, diopside, dravite, edingtonite, elbaite, epistilbite, flagstaffite, gismondine, gmelinite-(Na), gugiaite, helvine, hemimorphite, hilgardite, hydrocalumite, innelite, jarosite, jeremejevite, junitoite, langbeinite, larsenite, leucophanite, londonite, meliphante, mesolite, mimetite, natrolite, neptunite, nitrobarite, olsacherite, pharmacosiderite, pirssonite, pyromorphite, quartz, rhodizite, schorl, scolecite, searlesite, shortite, spangolite, sphalerite, stibiocolumbite, stibiotantalite, struvite, suolunite, thomsonite, thornasite, tilasite, tugtupite, uvite, weloganite, whitlockite, wulfenite and yugawaralite.

[0170] Still referring to the embodiment of FIG. 4, metallic nanoparticles in the reaction material 320 comprise elements that are capable of forming electrically conductive and/or interstitial metal hydrides comprising transition metals that have at least one stable isotope comprising group 3B elements scandium Sc, yttrium Y, lanthanum La, lanthanides (cerium Ce, praseodymium Pr, neodymium Nd, samarium Sm, europium Eu, gadolinium Gd, terbium Tb, dysprosium Dy, holmium Ho, erbium Er, thulium Tm, ytterbium Yb and lutetium Lu), actinides (thorium Th and uranium U), group 4B elements titanium Ti, zirconium Zr, hafnium, group 5B elements vanadium V, niobium Nb, tantalum Ta, group 6B elements chromium Cr, molybdenum Mo, tungsten W, group 7B elements manganese Mn, rhenium Re, group 8B metals iron Fe, ruthenium Ru, osmium Os, cobalt Co, rhodium Rh, iridium Ir, nickel Ni, palladium Pd, platinum Pt, group 1B elements copper Cu, silver Ag, gold Au and group 2B elements zinc Zn, cadmium Cd and mercury Hg. In periodic table of elements where groups of elements are marked with numbers 1-18, transition metals are in groups 3-12.

[0171] Particle size of the metallic nanoparticles in the reaction material 320 is preferably smaller than about 1 μm, more preferably smaller than about 100 nm, even more preferably smaller than about 30 nm, most preferably smaller than about 10 nm. The smaller the metallic nanoparticles are the larger is the active surface area of the said metallic nanoparticles. Metallic nanoparticles have preferably sharp edges and/or tips to enhance the local electric field strength.

[0172] Referring now to the invention in more detail, in FIG. 5 there is shown a reaction container system 500 for generating thermal energy. The reaction container system 500 comprises active powder material 320 in a reaction container 501.

[0173] In more detail, still referring to the embodiment of FIG. 5, the reaction container 501 is pressurized with hydrogen gas from the direction indicated with an arrow 502 through gas line manual isolation valves 504 and 506 along the hydrogen gas line 507 to the reaction container 501.

[0174] There is a gas line fitting 508 between the gas line manual isolation valves 504, 506.

[0175] In certain example embodiment of the present invention the cooling fluid circulation is arranged with a cooling fluid tube coil 522 (also referred to as cooling fluid circulation 520) placed around the reaction container 501.

[0176] In certain example embodiment of the present invention the cooling fluid circulation is arranged with a cooling fluid mantle essentially surrounding the reaction container 501.

[0177] Cooled cooling fluid enters the cooling fluid tube coil through inlet manual isolation valves 530 and 532 as indicated with the inlet flow direction arrow 526.

[0178] There is a cooling fluid inlet fitting 534 between the inlet manual isolation valves 530, 532. Cooling fluid collects thermal energy from the reaction container 501 and becomes hot while flowing through the cooling fluid tube coil 522. Heated cooling fluid leaves the cooling fluid tube coil 522 through outlet manual isolation valves 536 and 538 as indicated with the outlet flow direction arrow 528.

[0179] There is a cooling fluid outlet fitting 540 between the outlet manual isolation valves 536, 538. The temperature of the powder material is measured with the thermocouple 516 that has a thermocouple connector 521. The temperature of the heated cooling fluid arriving from the cooling fluid tube coil 522 is measured with the thermocouple 517 having a thermocouple connector 519 and attached to the outlet conduit 524.

[0180] The heater cartridge 510 is placed into the powder material 320 through the refilling port fitting 514. Electric power plug 512 attached to the heater cartridge 510 is used for connecting an external electric power cable to the heater cartridge 510.

[0181] In another embodiment the reaction container system 500 is equipped with a piezoelectric transducer 550 attached to the reaction container 501. The piezoelectric transducer 550 with the power plug 552 converts electrical pulses to mechanical vibrations that make the reaction container 501 and the reaction material 320 vibrate. Electric power plug 552 attached to the piezoelectric transducer 550 is used for connecting an external electric power cable to the piezoelectric transducer 550.

[0182] In yet another example embodiment at least one piezoelectric transducer is placed in direct contact with the powder material 320 inside the reaction container 501.

[0183] In certain example embodiments the cartridge system 500 is equipped with an electrical coil 518 placed around the reaction container 501. Electric current in the electrical coil 518 induce a magnetic field inside the said electrical coil.

[0184] The reaction container 501 is surrounded by a radiation shield mantle 554 that stops any residual radioactive radiation arriving from the powder material 320. The radiation shield mantle converts radioactive radiation into thermal energy and donates the said thermal energy to the cooling fluid tube coil 522.

[0185] To avoid thermal energy losses the reaction container system 500 is thermally insulated by a thermal insulation mantle (not shown in FIG. 5) that surrounds the reaction container system 500.

[0186] The reaction container system 500 is replaced by closing the gas line manual isolation valves 504, 506 and closing the inlet manual isolation valves 530, 532 and closing the outlet manual isolation valves 536, 538, disconnecting power cables from the electric power plugs 512, 552, disconnecting thermocouple connectors 519, 521, opening the gas line fitting 508 and opening the cooling fluid inlet fitting 534 and cooling fluid outlet fitting 540. The used reaction container system 500 now isolated from the surroundings is removed from the installation point and a new reaction container system with active reaction material is attached to the gas line and cooling line fittings and cable plugs and thermocouple connectors that were opened before removing the used reaction container system. The reaction container system 500 becomes in fluid communication with the hydrogen source and the cooling water circulation by opening the outlet manual isolation valves 536, 538, opening the inlet manual isolation valves 530, 532 and opening the gas line manual isolation valves 504, 506. The reaction container 501 is pressurized with hydrogen gas.

[0187] Referring now to the invention in more detail, in FIG. 6 there is shown a cross section 600 of the parts inside the radiation shield mantle (554 in FIG. 5) used for generating thermal energy. In the cross section there are seen the reaction material 320, the cooling fluid circulation 520 and the electrical coil 518 for inducing magnetic field inside the said coil.

[0188] In more detail, still referring to the embodiment of FIG. 6, the reaction material 320 generates thermal energy by the fusion of hydrogen with nuclei. The fusion process is controlled by the magnetic field induced by the electrical coil 518. Magnetic field inside the coil in the powder material polarizes the multiferroic phase of the powder material 320. Heat energy is transported mostly by thermal conduction from the powder material 320 to the fluid in the cooling fluid circulation 520, the said fluid preferably been high thermal capacity liquid (e.g. water or molten metal), solution or gas (helium).

[0189] Referring now to the invention in more detail, in FIG. 7 there is shown a reaction container system 700 according to an embodiment comprising reaction material 320, hydrogen gas line 507 that is used for pressurizing the reaction container 708 with hydrogen gas, a heater cartridge with a power plug 512 for preheating the reaction material 320 to the optimum reaction temperature range, a thermocouple 516 for measuring the temperature of the reaction material 320, an internal cooling mantle 702 with a cooling fluid inlet 704 and a cooling fluid outlet 706 for collecting heat from the reaction material 320, a radiation shield mantle 709 surrounding the

powder material and a thermal insulation mantle 710. A cross section of the reaction container system 700 is indicated with the line 712.

[0190] In more detail, still referring to the embodiment of FIG. 7, the powder material 320 is pressurized with hydrogen gas flowing through the hydrogen gas line 507 along the direction indicated with an arrow 502. The heater cartridge is heated with electricity and the generated heat energy preheats the reaction material 320 to a suitable reaction temperature range.

[0191] In further detail, still referring to the embodiment of FIG. 7, the hydrogen gas in the reaction material 320 is in equilibrium with the metal hydrides forming a part of the reaction material 320, meaning that increasing the hydrogen gas pressure increases the amount of metal hydrides in the reaction material 320 and decreasing the hydrogen gas pressure decreases the amount of metal hydrides in the reaction material 320.

[0192] Still referring to the embodiment of FIG. 7, hydrogen is activated into atomic hydrogen and ionized into hydrogen ions such as proton in the reaction material 320. Hydrogen ions are accelerated in the high electric field strength areas in the reaction material 320 to high kinetic energy and fused by tunneling through the Coulomb barrier with the nuclei of the target isotopes in the reaction material 320 creating new isotopes from the target isotopes and releasing energy in the form of thermal energy and radioactive radiation, essentially gamma radiation or X-ray radiation, that is converted into thermal energy by radiation-absorbing materials.

[0193] Still referring to the embodiment of FIG. 7, hydrogen ions and electrons are accelerated in low or medium electric field strength areas in the reaction material 320 to relatively low kinetic energy and contacted with powder surfaces having elements capable of forming Rydberg atoms. Rydberg atoms being electrical dipoles are attracted together to form condensed phase Rydberg matter that is stable until the excited electrons return to the ground level or the matter is at least partly ionized and Coulomb explosion tears the Rydberg atom cluster into separate ions that are accelerated away from the cluster because of the repulsive forces between ions. Accelerated ions, such as protons or deuterons, hit surrounding target atoms, such as nickel atoms, and some of the accelerated ions are capable of tunneling through the Coulomb barrier of target atoms fusing with the nuclei of the target atoms creating new isotopes from the target isotopes and releasing energy in the form of thermal energy and radioactive radiation, essentially gamma radiation or X-ray radiation, that is converted into thermal energy by radiation-absorbing materials.

[0194] The construction details of the embodiment as shown in FIG. 7 are that the walls of the reaction container and the cooling fluid mantle are made of materials preferably comprising metals and metal alloys known to be suitable for nuclear reactor construction, such as fine-grained low alloy ferritic steel and pressure vessel steel, e.g. the 20MnMoNi55 alloy. The radiation shield mantle is made of materials that stop radioactive radiation, preferably comprising lead for stopping gamma radiation and materials with high nuclear cross section selected from Table 1, e.g. boron or gadolinium, for stopping neutron radiation. In an embodiment the cooling fluid contains fluid-soluble metal compounds that absorb

radioactive radiation releasing thermal energy, e.g. boron compounds such as sodium borate for absorbing neutron radiation.

[0195] Referring now to the invention in more detail, in FIG. 8 there is shown a cross section 800 of the reaction container system 700 presented in FIG. 7 comprising reaction material 320 in a reaction container 708, the cooling fluid mantle 702, the radiation shield mantle 709 and the thermal insulation mantle 710.

[0196] In more detail, still referring to the embodiment of FIG. 8, thermal energy released from the reaction material 320 or generated within the radiation shield mantle 709 from the radioactive radiation heats the cooling fluid flowing in the cooling fluid mantle 702. Radioactive radiation, mostly gamma and X-ray radiation, released from the reaction material 320, is absorbed by the radiation shield mantle 709 and converted into thermal energy that heats the cooling fluid flowing in the cooling fluid mantle 702. Thermal insulation mantle 710 limits heat losses from the fusion container to the surroundings.

[0197] Referring now to the invention in more detail, in FIG. 9a there is shown a reaction container system 900 that initiates fusion processes by electromagnetic induction comprising a reaction container 901 (a rectangle with thick black line) that contains reaction material 320, a cooling fluid tube 914, a metal coil 902, a radiation shield mantle 932, a thermal insulation mantle 934, a hydrogen gas port 910 and a refill port 928.

[0198] In more detail, still referring to the embodiment of FIG. 9a, the hydrogen gas port 910 has a fitting 908 for attaching the hydrogen gas port 910 to the external hydrogen gas source and a particle filter 912 for preventing the flow of reaction material 320 to the external hydrogen gas source. The cooling fluid tube 914 has manual valves 918, 920 for isolating the cooling fluid tube from external cooling fluid lines and fittings 916, 922 for disconnecting the cooling fluid tube 914 from the external cooling fluid lines. The cooling fluid tube has optional fins or protrusions 915 for enhancing the collection of thermal energy from the reaction material 320. A thermocouple 924 is attached to the wall of the cooling fluid tube 914 for measuring the temperature of the cooling fluid in the cooling fluid tube 914. A thermocouple 927 with a measurement cable 926 is attached to the wall of the reaction container 901 for measuring the temperature of the reaction container 901. A filling port 928 with a blind plug 930 is attached to the reaction container 901 for removing depleted reaction material 320 from the reaction container 901 and filling the reaction container 901 with new reaction material 320.

[0199] In further detail, still referring to the embodiment of FIG. 9a, the metal coil 902 with power cables 904, 906 is used for inducing variable magnetic field inside the reaction container 901. The variable magnetic field pre-heats the reaction material by electromagnetic induction to the temperature that is suitable for the fusion reactions. After the pre-heat time the metal coil 902 is used for inducing quickly variable magnetic field in the reaction material 320 for polarizing the dielectric material (multiferroic material) present in the reaction material 320 and/or inducing voltage to the metallic material (metallic nanoparticles) present in the reaction material. The frequency of the current going through the metal coil and inducing the magnetic field is preferably in the range of about 10 Hz-100 MHz. The shape of the current pulses can be sine wave, square wave, step wave, sawtooth wave, triangular

wave or an arbitrary waveform easily generated by a computer-controlled arbitrary waveform generator equipped with a power amplifier.

[0200] Still referring to the embodiment of FIG. 9a, in an embodiment the dielectric material present in the reaction material 320 is an electric insulator material (polarizable or non-polarizable) and it is used for keeping metallic nanoparticles present in the reaction material 320 separated from each other so that there are a large number of very small gas gaps in the nm-range between metallic nanoparticles, and the metal coil 902 is used for creating a variable magnetic field that induces electric potential (voltage) to the metallic nanoparticles that focus the local electric field caused by the induced electric potential to very high electric field strength suitable for accelerating hydrogen ions (e.g. protons) and causing fusion reactions that release fusion energy.

[0201] Still referring to the embodiment of FIG. 9a, the cooling fluid used in the cooling fluid tube 901 for transporting thermal energy comprises a material that is liquid or gas near room temperature, selected from liquids comprising water, solutions containing water, gallium metal alloys, such as galinstan (Ga—In—Sn alloys), and heat transfer oils, such as Shell heat transfer oil S2, and gases comprising helium.

[0202] Referring now to the invention in more detail, in FIG. 9b there is shown a cross section of the fusion container comprising the cooling fluid tube 914, reaction material 320, the metal coil 902, the radiation shield mantle 932 and the thermal insulation mantle 934.

[0203] In more detail, still referring to the embodiment of FIG. 9b, the cooling fluid tube 914 has thermally-conductive fins or protrusions 915 that enhance the transfer of thermal energy from the reaction material 320 to the cooling fluid flowing in the cooling fluid tube 914. The material, size and shape of the fins or protrusions 915 and the cooling fluid tube 914 is optimized in such a way that electromagnetic induction from the metal coil is still capable of inducing voltage to the electrically-conducting phases of the reaction material 320. The thermal insulation mantle 934 prevents thermal losses (thermal conduction, convection, heat radiation) from the fusion container to the surroundings, so that the thermal energy can efficiently be collected by the cooling fluid in the cooling fluid tube 914.

[0204] The construction details of the embodiment of the present invention as shown in FIG. 9b are that the thermally-conductive fins or protrusions 915 are made of materials that have good thermal conductivity such as aluminum, copper or silicon carbide. The cooling fluid tube 914 and the reaction container 901 are preferably made of materials suitable for constructions used in environment that have radioactive radiation, such as fine-grained low alloy ferritic steel and pressure vessel steel, e.g. the 20MnMoNi55 alloy. In an embodiment the corrosion resistance of the cooling fluid tube is enhanced with an internal and/or external cladding of a corrosion-resistant material such as zirconium, e.g. Zircaloy. In an embodiment the diffusion of hydrogen gas from the reaction material volume to the cooling fluid tube 914 is prevented by a coating that comprises preferably dense metal compounds such as amorphous tantalum pentoxide, titanium dioxide, aluminum oxide or silicon dioxide made by known coating methods.

[0205] Referring now to the invention in more detail, in FIG. 9c there is shown a curve of variable voltage that has positive voltage peaks 954 and negative voltage peaks 956.

The magnitude of the voltage is presented in the y-axis **952** and the time scale is presented in the x-axis **950**.

[0206] In more detail, still referring to the embodiment of FIG. **9c**, according to the method of preheating the reaction container by electromagnetic induction and inducing a voltage to the electrical conductors inside the cartridge, the voltage applied to the induction coil **902** is preferably alternating voltage that has a frequency and intensity depending on the heating or induction power that is needed. The said frequency can be up to the radio frequency range (RF-range, MHz range) or even higher.

[0207] In further detail, still referring to the embodiment of FIG. **9c**, the driving voltage may comprise positive and/or negative pulses that have certain duration based on pulse-width modulation.

[0208] Still referring to the embodiment of FIG. **9c**, short and strong voltage pulses create strong variable magnetic field in the reaction material **320** that induces variable voltage to electrically conductive particles in the reaction material **320**. The said variable voltage creates an electric field around the electrically conductive particles (such as titanium, zirconium, hafnium and/or nickel nanoparticles). The electric field is focused with the geometry and dimensions of the electrically conductive particles to very high electric field strength. Ions such as protons are accelerated by the electric field to high kinetic energy. Some of those ions collide with target isotopes, tunnel through the Coulomb barrier shielding nuclei of the target isotopes and fuse with the nuclei forming new isotopes heavier than the original collision target isotopes were before the fusion process and at the same time release energy in the fusion process. Although the probability for the fusion between a single selected target isotope nucleus and a single proton is extremely small, the number of target isotope nuclei and protons in the reaction cartridge (fusion cartridge) is so large and so many collisions between target isotope nuclei and protons occur per time unit that the total sum of probabilities for fusion reactions becomes favorable for the thermal energy production with the coefficient of performance COP clearly above 1.

[0209] For example, when 100 g of natural nickel consisting of stable nickel isotopes, that has a density of 8.9 g/cm^3 , is divided into 5-nm particles (in this example spheres to simplify the calculations, although more useful geometries with sharp tips and edges can also be applied in the invention), the number of Ni nanoparticles is $100 \text{ g} / 8.9 \text{ g/cm}^3 * (10^7 \text{ nm})^3 / \text{cm}^3 / (4/3 * 3.14159 * (2.5 \text{ nm})^3) = 11.236 * 10^{21} / 65.45 \text{ pieces} = \text{about } 1.717 * 10^{20} \text{ pieces}$. Because the atomic weight of nickel is 58.69 g/mol and 100 g of nickel contains $100 / 58.69 \text{ mol} = 1.704 \text{ mol}$ of nickel and 1 mol contains about $6.022 * 10^{23}$ nickel atoms, each 5-nm nickel nanoparticle contains about $1.704 * 6.022 * 10^{23} / 1.717 * 10^{20} = \text{about } 6000$ nickel atoms of various nickel isotopes.

[0210] Referring now to the invention in more detail, in FIG. **10** there is shown the reaction container core **1000** comprising a container or pressure vessel **1001** holding reaction material **320** that in more detail, as indicated with an arrow **1002**, comprises dielectric material **1004** that possesses electric polarizability having positive **1006** and negative **1008** electric poles due to the polarization, metallic material **1010** capable of forming interstitial and/or electrically conductive metal hydrides, and optionally catalytic material capable of forming and storing Rydberg matter and inverted Rydberg matter (not shown in FIG. **10**).

[0211] In more detail, still referring to the embodiment of FIG. **10**, the reaction container core **1000** has walls in the pressure vessel **1001** that keep the reaction material **320** inside the reaction container core **1000**.

[0212] In further detail, still referring to the embodiment of FIG. **10**, an area **1012** is selected for detailed description illustrated in FIG. **11**.

[0213] The construction details of the embodiment of the invention as shown in FIG. **10** are that the walls in the pressure vessel **1001** are made of a material strong and leak-tight enough to hold high pressure (e.g. over 10 bar gauge) inside the reaction container even after bombardment with radioactive radiation without any considerable leak of gases from the reaction container **1000** to the surroundings of the said reaction container. The material of the walls in the pressure vessel **1001** comprises preferably metals and metal alloys known to be suitable for nuclear reactor construction, such as fine-grained low alloy ferritic steel and pressure vessel steel, e.g. the 20MnMoNi55 alloy. The dielectric material **1004** comprises material that can be electrically polarized with magnetic field, such as multiferroic materials, or that can be electrically polarized with mechanical stress or vibrations, such as piezoelectric materials or that can be electrically polarized with variable temperature, such as pyroelectric materials.

[0214] Still referring to FIG. **10** the construction details of the embodiment of the invention are that the metallic material **1010** comprises material that is capable of forming interstitial metal hydrides and/or electrically conductive metal hydrides, such as transition metal hydrides, e.g. nickel hydrides or titanium hydrides. The dielectric material **1004** comprises porous material that provides large internal surface area in the pores and/or powder that provides large outer surface area on the surface of particles. The metallic material comprises powder that provides large outer surface area on the surface of particles of the metallic material. The reaction material **320** comprises a mixture of the dielectric material **1004** and the metallic material **1010**. In an embodiment in the mixture there is preferably a porous dielectric material coated with metallic nanoparticles so that the nanoparticles are located on the inner surface of the pores of the porous material. In an embodiment in the mixture there are preferably dielectric material **1004** particles having preferably 10-10000 nm size mixed with metallic material **1010** nanoparticles having preferably 0.5-100 nm size, more preferably 2-10 nm size. The metallic material **1010** nanoparticles comprises preferably transition metals, e.g. titanium, zirconium, hafnium or nickel. The dielectric material **1004** comprising multiferroic, pyroelectric and/or piezoelectric material is selected from multiferroic material that become electrically polarized in magnetic field and/or pyroelectric materials selected from pyroelectric crystal classes 1, 2, m, mm2, 3, 3 m, 4, 4 mm, 6 and 6 mm, and/or from piezoelectric materials selected from piezoelectric crystal classes 1, 2, m, 222, mm2, 4, -4, 422, 4 mm, -42 m, 3, 32, 3 m, 6, -6, 622, 6 mm, -62 m, 23 and -43 m.

[0215] Referring now to the invention in more detail, in FIG. **11** there is shown a schema **1100** that comprises dielectric particles **1102**, **1108** that possess electrical polarizability and metallic nanoparticles **1114**, **1116**, **1118** that comprise elements capable of forming interstitial and/or electrically conductive metal hydrides.

[0216] In more detail, still referring to the embodiment of FIG. **11**, the dielectric particle **1108** has a positive electric

pole **1110** pointing towards the metallic nanoparticle **1118** and a negative electric pole **1112** pointing away from the metallic nanoparticle **1118**. The dielectric particle **1102** has a negative electric pole **1106** pointing towards the metallic nanoparticle **1118** and a positive electric pole **1104** pointing away from the metallic nanoparticle **1118**.

[0217] In further detail, still referring to the embodiment of FIG. **11**, the positive electric pole **1110** and the negative electric pole **1106** create an electric field between the said poles **1110** and **1106**. The metallic nanoparticle **1118** between the electric poles **1110** and **1106** focuses the electric field to a small volume and thus assists the formation of the very strong electric field **1120** between the positive electric pole **1110** and the metallic nanoparticle **1118** and the formation of the very strong electric field **1122** between the metallic nanoparticle **1118** and the negative electric pole **1106**. The voltage gradient is extremely steep in the very strong electric field **1120**, **1122**.

[0218] Still referring to the embodiment of FIG. **11**, ions can be accelerated to high kinetic energy in the very strong electric fields **1120**, **1122** (in other words nanoscale particle accelerators are utilized). Positive ions (such as proton, p^+) are accelerated towards the negative electric pole **1106** and negative ions (such as hydride ion, H^-) are accelerated towards the positive electric pole **1110**. Polarity of the dielectric particles **1108**, **1102** can be quickly altered by the polarization control factors comprising variable magnetic field, mechanical vibrations and/or variable temperature, the polarization control factor being chosen according to the material of the dielectric particles **1108**, **1102**.

[0219] Still referring to the embodiment of FIG. **11**, ions accelerated to high kinetic energy can fuse with the nuclei of the metallic nanoparticle **1118** and the nuclei of the dielectric particles **1108**, **1102**, releasing fusion energy e.g. in the form of gamma radiation.

[0220] Referring now to the invention in more detail, in FIG. **12** there is shown a schema **1200** comprising a particle **1102** that possesses electrical polarizability and a nanoparticle **1118** that comprises an element or elements capable of forming interstitial and/or electrically conductive metal hydrides.

[0221] In more detail, still referring to the embodiment of FIG. **12**, the particle **1102** is polarized and it has a negative electric pole with negative electric charge **1106** and a positive electric pole with positive electric charge **1104**. There is an area **1202** with very high electric field strength between the particle **1102** and the nanoparticle **1118**. The electric field accelerates ions towards the electric pole as indicated with the arrow **1206**. In an embodiment positive hydrogen ions (p^+ , protons) are accelerated towards the negative electric pole **1106**.

[0222] In further detail, still referring to the embodiment of FIG. **12**, the protons **1204** acquire kinetic energy in the area of very strong electric field **1202**. The protons arrive to the surface of the polarized particle **1102** colliding with the atoms of the polarized particle **1102**. Some of the accelerated positive hydrogen ions (protons, p^+) **1204** tunnel through the Coulomb barrier of the atoms of the polarized particle **1102** and fuse with the nuclei of the said atoms forming new isotopes that may also comprise non-stable isotopes, and releasing fusion energy.

[0223] Still referring to the embodiment of FIG. **12**, in case the polarized particle **1102** comprises lithium tetraborate,

accelerated protons fuse with lithium and boron nuclei in the polarized particle **1102** releasing fusion energy.

[0224] Still referring to the embodiment of FIG. **12**, the polarized particle **1102** and the metallic nanoparticle **1118** may be a part of a larger agglomerate that immobilizes particles and nanoparticles in such a way that at least some of those particles can maintain small gas gaps between those particles.

[0225] Referring now to the invention in more detail, in FIG. **13** there is shown a schema **1300** comprising a particle **1108** that possesses electrical polarizability and a nanoparticle **1118** that comprises an element or elements capable of forming interstitial and/or electrically conductive metal hydrides.

[0226] In more detail, still referring to the embodiment of FIG. **13**, the particle **1108** is polarized and it has a negative electric pole with negative electric charge **1112** and a positive electric pole with positive electric charge **1110**. The surface of the nanoparticle **1118** interacts with thermal radiation. Thermal radiation photon **1312** that has energy $h\nu$, wherein h is the Planck constant and ν is the frequency of the photon, excites surface plasmon that is a traveling wave oscillation of electrons and forms a polariton that propagates along the surface of the nanoparticle **1118**. It is not yet clear to what extent the said polariton is capable of enhancing the local electric field strength especially near the tip pointing towards the electric pole **1110**. There is an area **1120** with very high electric field strength between the particle **1108** and the nanoparticle **1118**. The electric field accelerates ions towards the matching electric pole as indicated with the arrow **1314**. In very high electric field electrons can be ripped away from atoms forming ions. In an embodiment very high electric field **1120** ionizes hydrogen and forms low-temperature hydrogen plasma consisting of electrons and protons. In an embodiment positive hydrogen ions (p^+ , protons) are accelerated towards the nanoparticle **1118**. Accelerated protons **1310** arrive to the surface of the nanoparticle **1118** colliding with the atoms of the nanoparticle **1118**. Some of the accelerated protons **1310** tunnel through the Coulomb barrier of the atoms of the nanoparticle **1118** and fuse with the nuclei of the said atoms forming new isotopes and releasing energy. In an embodiment accelerated protons **1310** fuse with the nuclei of nickel atoms in the nanoparticle **1118** and form new isotopes that are heavier than the original nickel isotopes releasing energy. The amount of released energy is estimated in Examples hereinafter.

[0227] Referring now to the invention in more detail, in FIG. **14a** there is shown the electron shell structure of the potassium atom, and in FIG. **14b** there is shown the excitation of an electron of the potassium atom to a Rydberg state.

[0228] In more detail, still referring to the embodiment of FIG. **14a**, potassium atom **1400** has nucleus **1402** containing 19 protons (${}_{19}K$) and a variable number of neutrons depending on the potassium isotope. Stable potassium isotopes are ${}^{39}K$ and ${}^{41}K$, wherein the numbers 39 and 41 denote the number of nucleons consisting of protons and neutrons in potassium nuclei. Potassium also has a long-lived radioactive isotope ${}^{40}K$. The electron shell structure of potassium is $1s^2 2s^2 2p^6 3s^2 3p^6 4s^1$. Large distance from the small nucleus **1402** to the first electron shell, K shell **1406**, filled with two 1s electrons, is indicated with the dashed arrow **1404**. L and M shells **1408**, **1410** are also completely filled with electrons. The outer shell, N shell **1412**, has one 4s electron that acts as a valence electron capable of forming a chemical bond

between atoms. This valence electron (in this case the 4s electron) is important for the formation of the Rydberg atom. The valence electron 4s can be removed from potassium atom with about 4.34 eV ionization energy and Rydberg states of the 4s electron utilized in an embodiment of the present invention are below this ionization energy level.

[0229] Still referring to the embodiment of FIG. 14b, the single 4s electron 1414 orbits the potassium atom. The other electrons and the nucleus of the potassium atom are inside the circle 1450. When the 4s electron 1414 is excited 1452 to the Rydberg state of the electron 1454, the distance between the 4s electron and the filled electron shells of the potassium atom increases and the potassium atom becomes potassium Rydberg atom. The excited electron is not removed from the potassium atom but the electron 1454 is still orbiting the potassium atom 1450 and it is part of the potassium atom as indicated with the dashed line 1456. Because the orbit of the excited electron 1454 is far away from the potassium atom 1450 and the angular velocity of the excited electron 1454 is relatively small, the side of the excited electron becomes negatively charged δ^- 1458, and the opposite side of the potassium atom becomes positively charged δ^+ 1460. The potassium Rydberg atom behaves like an electric dipole and it will be affected by external electric and magnetic fields.

[0230] Referring now to the invention in more detail, in FIG. 15a there is shown the electron shell structure of the hydrogen atom, and in FIG. 15b there is shown the excitation of the electron of the hydrogen atom to a Rydberg state.

[0231] In more detail, still referring to the embodiment of FIG. 15a, hydrogen atom 1500 has nucleus 1502 containing one proton (^1_1H) and a variable number of neutrons depending on the potassium isotope. Stable hydrogen isotopes are ^1H (protium) and ^2H (deuterium), where the nucleus has 0 and 1 neutron, respectively. Radioactive hydrogen isotope ^3H (tritium) has two neutrons. All these hydrogen isotopes ^1H , ^2H and ^3H can be used for generating Rydberg atoms and Rydberg matter. Large distance from the small nucleus 1502 to the first electron shell, K shell 1506, filled with one is electron 1504, is indicated with the dashed arrow 1508. This valence electron (in this case the 1s electron) is important for the formation of the Rydberg atom.

[0232] In more detail, still referring to the embodiment of FIG. 15b, the single 1s electron 1504 orbits the nucleus 1502 of the hydrogen atom. Neutral hydrogen atom does not have any other electrons. When the 1s electron 1504 is excited 1550 to a Rydberg state of the electron 1552, the distance between the 1s electron and the nucleus of the hydrogen atom increases and the hydrogen atom becomes hydrogen Rydberg atom. Methods for exciting the electron comprise collision with accelerated protons and electrons and exposure to coherent electromagnetic radiation. The excited electron is not removed from the hydrogen atom but the electron 1552 is still orbiting the nucleus 1502 of the hydrogen atom and it is part of the hydrogen atom as indicated with the dashed line 1554. Because the orbit of the excited electron 1552 is far away from the nucleus 1502 of the hydrogen atom and the angular velocity of the excited electron 1552 is relatively small, the side of the excited electron becomes negatively charged as indicated with the symbol δ^- 1558, and the opposite side of the hydrogen atom becomes positively charged as indicated with the symbol δ^+ 1556. The hydrogen Rydberg atom behaves like an electric dipole and it will be affected by external electric and magnetic fields.

[0233] Referring now to the invention in more detail, in FIG. 16a there is illustrated the attractive force between two hydrogen Rydberg atoms, in FIG. 16b there is illustrated the attractive force between two potassium Rydberg atoms, and in FIG. 16c there is illustrated the formation of a small Rydberg atom cluster.

[0234] In more detail, still referring to the embodiment of FIG. 16a, there is the first hydrogen Rydberg atom comprising hydrogen nucleus 1502 and an excited electron 1552 bound 1554 together forming the first electric dipole and the second hydrogen Rydberg atom comprising hydrogen nucleus 1602 and an excited electron 1604 bound 1606 together forming the second electric dipole. The first electric dipole has a side with a positive electric charge as indicated with the symbol δ^+ 1556 and a side with a negative electric charge as indicated with the symbol δ^- 1558. The second electric dipole has a side with a positive electric charge as indicated with the symbol δ^+ 1608 and a side with a negative electric charge as indicated with the symbol δ^- 1610. The first electric dipole and the second electric dipole are attracted and bound together by electrostatic forces as indicated with the arrow 1612.

[0235] In more detail, still referring to the embodiment of FIG. 16b, there is the first potassium Rydberg atom comprising potassium nucleus ^{19}K with K, L and M electron shells 1450 and an excited electron 1454 bound 1456 together forming the third electric dipole and the second potassium Rydberg atom comprising potassium nucleus ^{19}K with K, L and M electron shells 1630 and an excited electron 1632 bound 1634 together forming the fourth electric dipole. The third electric dipole has a side with a positive electric charge as indicated with the symbol δ^+ 1460 and a side with a negative electric charge as indicated with the symbol δ^- 1458. The fourth electric dipole has a side with a positive electric charge as indicated with the symbol δ^+ 1636 and a side with a negative electric charge as indicated with the symbol δ^- 1638. The third electric dipole and the fourth electric dipole are attracted and bound together by electrostatic forces as indicated with the arrow 1640.

[0236] In more detail, still referring to the invention of FIG. 16c, there is a small cluster of Rydberg atoms consisting of two hydrogen Rydberg atoms and one potassium Rydberg atom. The first hydrogen Rydberg atom with the excited electron 1552 in Rydberg state and the potassium Rydberg atom with the positive electric charge side 1460 are attracted and bound together by electrostatic forces as indicated with the arrow 1660. The second hydrogen Rydberg atom with the positive electric charge side 1608 and the potassium Rydberg atom with the excited electron 1454 in Rydberg state are attracted and bound together by electrostatic forces as indicated with the arrow 1662. Although the illustrated string comprises only three Rydberg atoms, longer strings are also feasible. Clusters of Rydberg atoms in sheet geometry are also feasible. The strings and sheets of Rydberg atoms may comprise single elements such as hydrogen Rydberg atoms. The strings and sheets of Rydberg atoms may also comprise more than one element such as hydrogen and potassium Rydberg atoms to form mixed element Rydberg matter. The formation of electrostatic bonds between Rydberg atoms releases energy that must be removed from the Rydberg atom cluster to form stable Rydberg matter. Energy released from the condensation of Rydberg atoms to Rydberg matter is

dissipated for example by phonons (lattice vibrations) to the surrounding crystal lattice or by the evaporation of nearby atoms or molecules.

[0237] Referring now to the invention in more detail, in FIG. 17a there is illustrated a crystal lattice without any defects, in FIG. 17b there is illustrated a crystal lattice with a defect induced by the change in crystal structure.

[0238] In more detail, still referring to the embodiment of FIG. 17a, the crystal lattice 1700 has atoms 1702 forming a hexagonal crystal lattice as indicated with the hexagon symbol 1704 drawn on the lattice.

[0239] In more detail, still referring to the embodiment of FIG. 17b, the mixed crystal lattice 1750 has the first crystal lattice shown with open circle atoms 1702 and the second crystal lattice shown with shaded atoms 1754. The first crystal lattice has hexagonal structure. Most of the first crystal lattice is without any mechanical strain as indicated with straight lines 1752. The second crystal lattice has cubic structure as indicated with the square symbol 1756 drawn on the lattice.

[0240] The first crystal structure and the second crystal structure differ from each other and they do not fit together. The second crystal lattice has induced mechanical strain to the first crystal lattice as indicated with the curved lines 1758. As a result, a discontinuity area (defect) is formed between the two crystal structures. In this example the formed defect is a void 1760 without any atoms. The void 1760 is utilized for the enhanced generation of Rydberg matter and storing the Rydberg matter or inverted Rydberg matter.

[0241] Referring now to the invention in more detail, in FIG. 18 there is illustrated the mixed crystal lattice structure with a structural defect housing a particle of inverted Rydberg matter.

[0242] In more detail, still referring to the embodiment of FIG. 18, the void 1762 between the first crystal lattice having atoms 1752 and the second crystal lattice having atoms 1756 is now occupied with a cluster of atoms 1802 forming a particle of inverted Rydberg matter. The particle may comprise hydrogen atoms in inverted Rydberg matter form.

[0243] Still referring to the embodiment of FIG. 18, the first crystal lattice comprises the first material that is often a metal oxide such as alumina Al_2O_3 and the second crystal lattice comprises the second material that is often a metal oxide such as iron oxide Fe_3O_4 . Although in this illustration the amount of the first material with the first crystal structure (hexagonal for the illustration purposes) is much larger than the amount of the second material (cubic for the illustration purposes), the ratio of crystal structures can vary in a wide range. For example, in case of a dehydrogenation catalyst (such as styrene catalyst) comprising inverse spinel cubic Fe_3O_4 doped with potassium generally in the form potassium oxide K_2O and also doped with structural promoters such as corundum structure alumina Al_2O_3 and/or chromia Cr_2O_3 , the amount of the second material (inverse spinel Fe_3O_4 doped with alkali metal such as potassium) 1756 is much larger than the amount of the first material (hexagonal Al_2O_3 and/or Cr_2O_3) 1752.

[0244] Still referring to the embodiment of FIG. 18, the illustrated mixed crystal structure material that is capable of enhancing the formation of Rydberg atoms and storing the said Rydberg atoms is preferably a constituent of the reaction material (320 in FIG. 4) comprising a compound or a mixture of compounds possessing electrical polarizability and a compound or a mixture of compounds capable of forming electrically conductive metal hydrides.

[0245] Referring now to the invention in more detail, in FIG. 19 there is shown a flow chart 1900 of a method for generating energy by solid state nuclear fusion.

[0246] In more detail, still referring to the embodiment of FIG. 19, hydrogen gas is originally molecular H_2 1902. Between two hydrogen atoms in the hydrogen molecule there is a chemical bond that must be broken. Breaking the covalent H:H chemical bond 1904 forms active hydrogen atoms H^* , wherein * denotes an unpaired electron, from the hydrogen molecule. Metals capable of forming metallic metal hydrides are utilized for breaking the H:H bond. Hydrogenation catalysts, such as Fischer-Tropsch catalysts, and dehydrogenation catalysts, such as styrene catalysts, and/or ammonia synthesis catalysts are utilized as promoters for enhancing the formation of active hydrogen atoms.

[0247] Still referring to the embodiment of FIG. 19, hydrogen atoms are excited into Rydberg atoms 1906, where an excited electron in hydrogen atom is in a Rydberg state that is above the ground state of the said electron and below the ionization energy level of the said electron. Excitation methods comprise excitation by accelerated electrons or protons and excitation by coherent electromagnetic radiation. Electrons or protons are accelerated to sufficient kinetic energy with the electric field generated by the dielectric materials possessing electrical polarizability. Rydberg atoms, being electrical dipoles, are attracted together into clusters to form Rydberg matter, shortened as RM 1908. Alkali metals, such as potassium K, used in catalysts activating hydrogen bond, form potassium Rydberg atoms with less excitation energy than the hydrogen atoms require for forming hydrogen Rydberg atoms. Alkali metals are examples of elements that possess Rydberg states and are capable of forming mixed element Rydberg matter, e.g. potassium-hydrogen Rydberg matter. In a nanopowder system that has large number of Rydberg atom clusters on the surface of powder particles, e.g. Fe_3O_4 :K particles, some of those clusters may enter such a quantum mechanical state that they invert their structure to form very dense inverted Rydberg matter 1910 where positively charged atom cores orbit close to negatively charged electrons.

[0248] Still referring to the embodiment of FIG. 19, the strength of the local electric field is increased to create higher energy electrons or protons that hit some of the Rydberg matter spots destabilizing Rydberg matter and inducing Coulomb explosion 1912 of the Rydberg matter and inverted Rydberg matter. Destabilization means that enough energy is added to an electron in Rydberg state so that the electron is lifted from the Rydberg state to the ionization energy level, which results in separate positive atom and negative electron and the quantum mechanical wave function of the Rydberg atom collapses.

[0249] Still referring to the embodiment of FIG. 19, in the beginning of the Coulomb explosion the atoms still have fixed position in the Rydberg matter meaning that they must have wide range of kinetic energy values. This is because of the Heisenberg uncertainty principle, which states that $\Delta x \Delta p \geq \hbar/2$, wherein Δx is the uncertainty of the position of the particle, Δp is the uncertainty of the momentum of the particle, and \hbar is reduced Planck's constant. Another way of expressing the system state is to apply the uncertainty principle of quantum mechanics, which states that $\sigma_x \sigma_p > \hbar/2$, wherein σ_x is the standard deviation of position and σ_p is the standard deviation of momentum. Fixed positions of atoms in a cluster of Rydberg atoms leads to a large range of momentum values for the

said atoms when they leave the cluster during the Coulomb explosion. Some of the atoms leaving the Rydberg matter cluster have very high kinetic energy, because the momentum wavefunction becomes spread out, and have an increased probability of tunneling through the Coulomb barrier of the surrounding atoms to induce nuclear fusion **1914**.

[0250] Still referring to the embodiment of FIG. 19, energy is released **1916** from the nuclear fusion between two nuclei, often between the nucleus of hydrogen and the nucleus of e.g. nickel.

[0251] Depending on the time how long the nucleus is in excited state due to the nuclear fusion, different kinds of de-excitation paths are feasible. Short time for the nucleus in excited state leads to the release of energy in the form of high energy gamma ray photons. Long time for the nucleus in excited state and resonance between the nucleus and the surrounding crystal lattice may lead to the release of de-excitation energy in the form of relatively low energy photons, such as X-ray photons, deep UV photons, or phonons (lattice vibrations) which transmit heat energy through the lattice.

[0252] Based on the above a method of producing energy according to the present invention, comprises the steps of

[0253] providing a reaction container (**350**) comprising reaction material (**320**), the reaction material (**320**) being formed by an electrically polarizable dielectric material and metallic material,

[0254] pressurizing the reaction container (**350**) with hydrogen gas,

[0255] activating hydrogen molecules in the hydrogen gas to provide atomic hydrogen,

[0256] polarizing the dielectric material to produce an electric field,

[0257] pulling hydrogen ions with the electric field from the metallic surface or ionizing the atomic hydrogen in the electric field to provide hydrogen ions, and

[0258] accelerating hydrogen ions in the electric field, wherein a part of the accelerated hydrogen ions tunnels through a Coulomb barrier between the hydrogen ions and atomic nuclei of the reaction material to fuse the hydrogen ions with the atomic nuclei of the reaction material to release energy.

[0259] In a preferred embodiment, the resistivity of the active hydrogen material is smaller than $1000 \mu\Omega\text{cm}$, preferably smaller than $500 \mu\Omega\text{cm}$, in particular smaller than $100 \mu\Omega\text{cm}$. The active hydrogen material comprises, for example, a hydrogen storage alloy, an electrically conductive hydrogenation catalyst, a material capable of forming binary metal hydride consisting of a metal and hydrogen, or a material capable of forming ternary metal hydride consisting of a first metal, a second metal and hydrogen.

[0260] A nuclear fusion system (**300**) of the present kind, for producing thermal energy, comprises

[0261] a reaction container (**350**),

[0262] reaction material (**320**) within the reaction container (**350**), the reaction material comprising electrically polarizable dielectric material and metallic material,

[0263] hydrogen gas source (**306**) connected to the reaction container (**350**) for pressurizing the reaction container (**350**) with hydrogen gas, and

[0264] heat exchange unit (**314**) for removing thermal energy produced in the reaction container.

[0265] The system further comprises

[0266] means for polarizing the dielectric material in order to produce an electric field within the reaction material,

[0267] means for activating hydrogen molecules into hydrogen atoms and ionizing hydrogen atoms in order to accelerate the hydrogen ions in the electric field so that they can tunnel through a Coulomb barrier between the hydrogen ions and atomic nuclei of the reaction material to fuse the hydrogen ions with the atomic nuclei of the reaction material to release energy.

[0268] In particularly preferred embodiments the system contains

[0269] a temperature measurement system (**328, 334**) for measuring the temperature of the reaction material (**320**) and from the heat exchange unit (**314**),

[0270] a pressure measurement system (**313**) for measuring hydrogen gas pressure, and

[0271] a control system (**304**) adapted to receive input from the temperature measurement system (**328, 334**) and the pressure measurement system (**313**) and to control the heat exchange unit (**314**) and/or hydrogen gas pressure, and optionally the heater (**322**).

[0272] The hydrogen gas source (**306**) preferably comprises a pressurized hydrogen gas bottle, metal hydrides heated to release hydrogen gas, or a source of chemical reactions releasing hydrogen gas, or a combination thereof.

[0273] A fusion energy production process according to the present technology comprises the steps of

[0274] providing a matrix of porous reaction material,

[0275] filling the pores of the matrix with hydrogen molecules,

[0276] breaking at least part of the covalent bonds of hydrogen molecules by activation to produce hydrogen atoms, and

[0277] exciting at least part of the hydrogen atoms into hydrogen Rydberg atoms so as to form Rydberg matter.

[0278] Further, in the process, at least part of the Rydberg matter is collided with ions or electrons accelerated in electric fields inside the reaction material so as to induce a Coulomb explosion of the Rydberg matter to produce high kinetic energy hydrogen ions, and at least part of the high kinetic energy hydrogen ions are fused with the atomic nuclei of the reaction material so as to release fusion energy.

[0279] In a preferred embodiment, the process comprises using metal capable of forming metallic metal hydride for breaking the covalent bonds of hydrogen molecules.

[0280] A fusion energy reaction material according to the present technology comprises a porous or powder mixture of electrically polarizable dielectric material, preferably in porous or powdery form, metallic material capable of forming metallic metal hydride, preferably in nanoparticle form, and a material capable of promoting the formation of Rydberg matter upon interaction with active hydrogen. There is also provided the use of hydrogen-containing Rydberg matter and/or inverted Rydberg matter as an intermediate material for providing high-energy hydrogen ions capable of fusing with other atomic nuclei in a fusion energy production process.

[0281] The following non-limiting examples illustrate the present technology.

Example 1

[0282] Nickel nanopowder having an average particle size of 10 nm was mixed with pyroelectric lithium tetraborate

$\text{Li}_2\text{B}_4\text{O}_7$ crystallite powder having particle size range of about 100 nm-1000 nm. $\text{Li}_2\text{B}_4\text{O}_7$ crystallite powder was prepared by mechanically crushing commercial $\text{Li}_2\text{B}_4\text{O}_7$ crystals to powder. The powder mixture is placed to the reaction cartridge. The reaction container was connected to a hydrogen gas line receiving hydrogen gas from a pressurized hydrogen gas bottle. The reaction container was also connected to the cooling fluid circulation. The reaction container was pressurized with hydrogen gas to 20 bar (gauge) and slowly heated to 400° C.

[0283] It is assumed that the pyroelectric crystallite powder was polarized by the temperature changes within the reaction material. The temperature of the reaction material was altered with external control (cooling fluid circulation) to keep the pyroelectric crystallite powder polarized. The system started to produce gamma radiation that had specific gamma photon energies. Generated thermal energy was removed by the cooling fluid circulation from the reaction container. The amount of collected thermal energy was much larger than the energy used for pre-heating the reaction container. After the test the reaction cartridge was de-pressurized and let to cool to room temperature for several days. The reaction material obtained from the cooled reaction container contained possibly some helium gas and traces of copper and beryllium that were not present in the original reaction material before the experiment. The construction materials used for the reaction container were originally free of copper and beryllium.

Example 2

[0284] The experimental setup was the same as used in Example 1 but nickel nanopowder was replaced with titanium nanopowder and lithium tetraborate was replaced with piezoelectric quartz SiO_2 powder. Externally controlled mechanical vibrations (ultrasonic source) provided the original electric field by polarization of the piezoelectric material. A lot of thermal energy was produced during the experiment. The COP was over 10. After the reactions the reaction material obtained from the reaction container possibly contained traces of vanadium isotopes and phosphorus that were not present in the original reaction material, although contamination from the steel used for the construction is not entirely excluded.

[0285] Secondary nuclear reactions forming stable isotopes from non-stable isotopes release more energy along time depending on the half lifes of the non-stable isotopes until the system consists only of stable isotopes. It is not yet certain how far along the titanium isotope chain it is possible to proceed. It is herein hypothesized that lighter titanium isotopes are fused with hydrogen into heavier titanium isotopes via non-stable vanadium isotopes.

[0286] It is not yet known how extensive and fast is the deterioration of the crystal structure of polarizable dielectric materials while operating the system at conditions favorable for fusion. The probability of proceeding further in the transmutation chain from the just created element to the next heavier element (a proton added) is possibly weakened locally after the first fusion reaction but the extent of deterioration that destroys locally the favorable fusion reaction conditions (high local electric field strength) for the transmutation is not yet clear.

Example 3

[0287] The experimental setup was the same as used in Example 1 but nickel nanopowder was replaced with zirconium

nanopowder and lithium tetraborate was replaced with multiferroic BiFeO_3 powder. Externally controlled magnetic field provided the local electric field by polarization of the multiferroic material. It is hypothesized that hydrogen was fused with zirconium because quite a lot of thermal energy was released accompanied by noticeable gamma radiation. After the reactions the reaction material obtained from the reaction container possibly contained traces of niobium and molybdenum isotopes that were not present in the original reaction material, although contamination from the steel used for the construction cannot be entirely excluded.

Theoretical Example 4

[0288] The experimental setup is the same as used in Example 1 but nickel nanopowder is generally replaced with transition metal nanopowder that is capable of forming a metallic or interstitial hydride having electrical conductivity.

Theoretical Example 5

[0289] The experimental setup is the same as used in Example 1 but instead of nanopowder mixtures, nanoporous pyroelectric, piezoelectric or multiferroic material is coated with transition metal nanoparticle thin film that is capable of forming a metallic or interstitial hydride having electrical conductivity. Nanopores provide sufficient surface area for colliding noticeable amount of hydrogen ions with the surface.

Example 6

[0290] A method of operating the thermal energy generator is presented herein.

[0291] Initiate the cooling media circulation around the reaction cartridge (control the mass flow rate based on the reaction cartridge temperature)

[0292] Increase the temperature of the reaction container with the heating means, e.g. with the heater cartridge e.g. to over 300° C. or to over 400° C.

[0293] Increase the pressure of the reaction container with hydrogen gas above room pressure, e.g. to over 10 bar gauge or to over 20 bar gauge.

[0294] Polarize the material possessing electric polarizability by creating mechanical vibrations in the reaction cartridge volume (used for polarizing electrically piezoelectric material) or by creating magnetic field in the reaction cartridge volume (used for polarizing electrically multiferroic material) or by changing the temperature in the reaction cartridge volume (used for polarizing electrically pyroelectric material, temperature change is induced by controlling the mass flow rate of the circulated cooling medium and by controlling the electric power going to the heater cartridge).

[0295] Collect thermal energy from the reaction cartridge with the heated circulated cooling medium.

Example 7

[0296] Instead of using mass defect and binding energy values, the amount of energy released in the fusion process is herein calculated directly from the fusion reaction equation: isotope x+hydrogen->isotope y+energy, wherein the total amount of energy (energy+energy equivalent of mass) is always constant in the isolated system.

[0297] The atomic mass (m_a) is the mass of a specific isotope, most often expressed in unified atomic mass units. The atomic mass is the total mass of protons, neutrons and electrons in a single atom.

[0298] The amount of energy released from the fusion of nickel with hydrogen is now estimated.

[0299] Natural nickel ${}_{28}\text{Ni}$ contains 0.680769 mole fraction of stable ${}^{58}\text{Ni}$ isotope. Fusion of nickel with hydrogen produces copper and releases energy as follows.

[0300] ${}^{58}_{28}\text{Ni}$ (57.9353429 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{59}_{29}\text{Cu}$ (58.939498 u) + 0.003670 u (3.41839426 MeV)

[0301] ${}^{59}\text{Cu}$ has 81.5 s half life and it emits β^+ (positron) to form ${}^{59}\text{Ni}$.

[0302] ${}^{59}_{29}\text{Cu}$ (58.939498 u) \rightarrow ${}^{59}_{28}\text{Ni}$ (58.9343467 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00460272 u (4.287249656 MeV)

[0303] Further, a positron β^+ annihilates immediately with an electron β^- .

[0304] β^+ (0.00054857990943 u) + β^- (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0305] ${}^{59}\text{Ni}$ has 76000 year half life and it emits β^+ to form ${}^{59}\text{Co}$, which is a stable isotope of 27Co.

[0306] ${}^{59}_{28}\text{Ni}$ (58.9343467 u) \rightarrow ${}^{59}_{27}\text{Co}$ (58.9331950 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00060312 u (0.56178224 MeV)

[0307] β^+ (0.00054857990943 u) + β^- (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0308] On the other hand, the life time of ${}^{59}\text{Ni}$ is so long that it can be fused with ${}^1\text{H}$ before the transmutation into ${}^{59}\text{Co}$.

[0309] ${}^{59}_{28}\text{Ni}$ (58.9343467 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{60}_{29}\text{Cu}$ (59.9373650 u) + 0.004806732 u (4.477278589 MeV)

[0310] ${}^{60}\text{Cu}$ has 23.7 min life time and it emits β^+ to form stable ${}^{60}\text{Ni}$ isotope.

[0311] ${}^{60}_{29}\text{Cu}$ (59.9373650 u) \rightarrow ${}^{60}_{28}\text{Ni}$ (59.9307864 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00603002 u (5.616722514 MeV)

[0312] β^+ (0.00054857990943 u) + β^- (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0313] Natural nickel ${}_{28}\text{Ni}$ contains 0.262231 mole fraction of stable ${}^{60}\text{Ni}$ isotope.

[0314] ${}^{60}_{28}\text{Ni}$ (59.9307864 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{61}_{29}\text{Cu}$ (60.933458 u) + 0.005154 u (4.800402128 MeV)

[0315] ${}^{61}\text{Cu}$ has 3.333 h half life and it emits β^+ to form stable ${}^{61}\text{Ni}$ isotope.

[0316] ${}^{61}_{29}\text{Cu}$ (60.933458 u) \rightarrow ${}^{61}_{28}\text{Ni}$ (60.9310560 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00185342 u (1.726386678 MeV)

[0317] β^+ (0.00054857990943 u) + 13 (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0318] Natural nickel ${}_{28}\text{Ni}$ contains 0.011399 mole fraction of stable ${}^{61}\text{Ni}$ isotope.

[0319] ${}^{61}_{28}\text{Ni}$ (60.9310560 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{62}_{29}\text{Cu}$ (61.932584 u) + 0.006297 u (5.865433492 MeV)

[0320] ${}^{62}\text{Cu}$ has 9.673 min half life and it emits β^+ to form stable ${}^{62}\text{Ni}$ isotope.

[0321] ${}^{62}_{29}\text{Cu}$ (61.932584 u) \rightarrow ${}^{62}_{28}\text{Ni}$ (61.9283451 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00369032 u (3.437385552 MeV)

[0322] β^+ (0.00054857990943 u) + β^- (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0323] Natural nickel ${}_{28}\text{Ni}$ contains 0.036345 mole fraction of stable ${}^{62}\text{Ni}$ isotope.

[0324] ${}^{62}_{28}\text{Ni}$ (61.9283451 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{63}_{29}\text{Cu}$ (62.929598 u) + 0.006573 u (6.122143868 MeV)

[0325] ${}^{63}\text{Cu}$ is a stable isotope of ${}_{29}\text{Cu}$.

[0326] Natural nickel ${}_{28}\text{Ni}$ contains 0.009256 mole fraction of stable ${}^{64}\text{Ni}$ isotope.

[0327] ${}^{64}_{28}\text{Ni}$ (63.9279660 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{65}_{29}\text{Cu}$ (64.927790 u) + 0.008002 u (7.453107062 MeV)

[0328] ${}^{65}\text{Cu}$ is a stable isotope of ${}_{29}\text{Cu}$.

[0329] Because copper hydrides are unstable at high temperatures and fusion reactions are somehow enhanced by the presence of electrically conductive metal hydride (in this example hydrides of nickel), it is possible that in this exemplar case the chain of nuclear transmutions stops to copper and copper does not transmutate to heavier elements.

Example 8

[0330] The electron is removed from hydrogen atom by ionization and a free proton is formed. The proton is accelerated by the very steep voltage gradient (very strong electric field between the metallic nanoparticle and the lithium tetraborate crystallite) towards the negative electric pole in lithium tetraborate. The amount of energy released from the fusion of hydrogen with lithium and boron in lithium tetraborate is now estimated. Although the fusion process involves free protons, the electron belonging to hydrogen is present in the fusion reaction equation to keep zero electric charge on both sides of the equation.

[0331] Natural lithium ${}_{3}\text{Li}$ contains 0.0759 mole fraction of stable ${}^6\text{Li}$ isotope.

[0332] ${}^6_3\text{Li}$ (6.015122795 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^7_4\text{Be}$ (7.01692983 u) + 0.006017997 u (5.605523551 MeV)

[0333] The half life of ${}^7\text{Be}$ is 53.22 days and it transmutes by electron capture (EC) into stable ${}^7\text{Li}$ isotope.

[0334] ${}^7_4\text{Be}$ (7.01692983 u) \rightarrow ${}^7_3\text{Li}$ (7.01600455 u) + 0.00092528 u (0.861861319 MeV)

[0335] Natural lithium ${}_{3}\text{Li}$ contains 0.9241 mole fraction of stable ${}^7\text{Li}$ isotope that can be fused with hydrogen. Natural boron ${}_{5}\text{B}$ contains about 19.9 at % of ${}^{10}\text{B}$ isotope and about 80.1 at % of ${}^{11}\text{B}$ isotope. The molecular weight of lithium tetraborate $\text{Li}_2\text{B}_4\text{O}_7$ is 169.1218 g/mol. One mole of $\text{Li}_2\text{B}_4\text{O}_7$ contains 2 mol Li (13.88 g) and 4 mol B (43.24 g). Further, 1 mol $\text{Li}_2\text{B}_4\text{O}_7$ contains 0.152 mol ${}^6\text{Li}$, 1.848 mol ${}^7\text{Li}$ and 0.796 mol ${}^{10}\text{B}$, 3.204 mol ${}^{11}\text{B}$.

[0336] $\text{Li}_2\text{B}_4\text{O}_7 + 3\text{H}_2 \Rightarrow 2\text{Li} + 2\text{p}$, $4\text{B} + 4\text{p} + 3.50_2 \Rightarrow$ helium + oxygen gas + energy

[0337] The nuclear reactions are as follows.

[0338] ${}^7_3\text{Li}$ (7.01600455 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^8_4\text{Be}$ (8.00530510 u) + 0.01852448 u (17.25481407 MeV)

[0339] The half life of ${}^8\text{Be}$ is 6.7×10^{-17} s and it fissions into two stable ${}^4\text{He}$ isotope atoms.

[0340] ${}^8_4\text{Be}$ (8.00530510 u) \rightarrow $2 \times {}^4_2\text{He}$ (2 \times 4.00260325415 u) + 0.00009859 u (0.091834225 MeV)

[0341] Natural boron ${}_{5}\text{B}$ contains 0.199 mole fraction of stable ${}^{10}\text{B}$ isotope.

[0342] ${}^{10}_5\text{B}$ (10.0129370 u) + ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{11}_6\text{C}$ (11.0114336 u) + 0.009328432 u (8.689061271 MeV)

[0343] The half life of ${}^{11}\text{C}$ is 20.334 min and it transmutes by positron emission (β^+) into stable ${}^{11}\text{B}$ isotope.

[0344] ${}^{11}_6\text{C}$ (11.0114336 u) \rightarrow ${}^{11}_5\text{B}$ (11.0093054 u) + β^+ (0.00054857990943 u, 0.5109989 MeV) + 0.00157962 u (1.47135293 MeV)

[0345] Positron β^+ annihilates electron β^- .

[0346] β^+ (0.00054857990943 u) + β^- (0.00054857990943 u) \rightarrow 2×0.5109989 MeV = 1.0219978 MeV

[0347] One mole natural boron ${}_{5}\text{B}$ contains 0.801 mol of stable ${}^{11}\text{B}$ isotope.

[0348] ${}^{11}_{5}\text{B}$ (11.0093054 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{12}_6\text{C}$ (12.0000000 u)+0.017130432 u (15.95631219 MeV)

[0349] ${}^{12}\text{C}$ isotope is stable.

[0350] The amount of energy released in the fusion process is $0.152*4.0*10^6*96.48$ kJ+ $1.848*17.2*10^6*96.48$ kJ+ $3.204*8.7*10^6*96.48$ kJ= $5814*10^6$ kJ, which corresponds to 1615000 kWh (thermal). Burning diesel in air releases thermal energy 38.6 MJ/liter. On the other hand, fusing about 170 g of lithium tetraborate with about 6.05 g of hydrogen releases about 5814000 MJ, which is equal to burning about 150000 liters (150 m³) of diesel.

[0351] It can be understood that the present invention provides an energy source that is very compact and has far higher energy-producing capacity than any energy source based on burning fossil fuels or hydrogen gas fuel cell.

Example 9

[0352] The amount of energy released from the fusion of titanium with hydrogen is now estimated.

[0353] Natural titanium ${}_{22}\text{Ti}$ contains 0.0825 mole fraction of stable ${}^{46}\text{Ti}$ isotope.

[0354] ${}^{46}_{22}\text{Ti}$ (45.9526316 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{47}_{23}\text{V}$ (46.9549089 u)+0.005547732 u (5.167490449 MeV)

[0355] The half life of ${}^{47}\text{V}$ is 3206 min and it transmutes by positron emission into stable ${}^{47}\text{Ti}$ isotope.

[0356] ${}^{47}_{23}\text{V}$ (46.9549089 u) \rightarrow ${}^{47}_{22}\text{Ti}$ (46.9517631 u)+ β^+ (0.00054857990943 u, 0.5109989 MeV)+0.00259722 u (2.41920663 MeV)

[0357] Positron β^+ annihilates electron β^- .

[0358] β^+ (0.00054857990943 u)+13 (0.00054857990943 u) \rightarrow $2*0.5109989$ MeV= 1.0219978 MeV

[0359] One mole of natural titanium ${}_{22}\text{Ti}$ contains 0.0744 mol of stable ${}^{47}\text{Ti}$ isotope.

[0360] ${}^{47}_{22}\text{Ti}$ (46.9517631 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{48}_{23}\text{V}$ (47.9522537 u)+0.007334432 u (6.831730031 MeV)

[0361] The half life of ${}^{48}\text{V}$ is 15.9735 d and it transmutes by positron emission into stable ${}^{48}\text{Ti}$ isotope.

[0362] ${}^{48}_{23}\text{V}$ (47.9522537 u) \rightarrow ${}^{48}_{22}\text{Ti}$ (47.9479463 u)+ β^+ (0.00054857990943 u, 0.5109989 MeV)+0.00375882 u (3.50119056 MeV)

[0363] Positron β^+ annihilates electron β^- .

[0364] β^+ (0.00054857990943 u)+ β^- (0.00054857990943 u) \rightarrow $2*0.5109989$ MeV= 1.0219978 MeV

[0365] One mole of natural titanium ${}_{22}\text{Ti}$ contains 0.7372 mol of stable ${}^{48}\text{Ti}$ isotope.

[0366] ${}^{48}_{22}\text{Ti}$ (47.9479463 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{49}_{23}\text{V}$ (48.9485161 u)+0.007255232 u (6.757958399 MeV)

[0367] The half life of ${}^{49}\text{V}$ is 329 d and it transmutes by electron capture into stable ${}^{49}\text{Ti}$ isotope.

[0368] ${}^{49}_{23}\text{V}$ (48.9485161 u) \rightarrow ${}^{49}_{22}\text{Ti}$ (48.9478700 u)+0.00064610 u (0.60181631 MeV)

[0369] Natural titanium ${}_{22}\text{Ti}$ contains 0.0541 mole fraction of stable ${}^{49}\text{Ti}$ isotope.

[0370] ${}^{49}_{22}\text{Ti}$ (48.9478700 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{50}_{23}\text{V}$ (49.9471585 u)+0.008536532 u (7.951438097 MeV)

[0371] ${}^{50}\text{V}$ isotope of ${}_{23}\text{V}$ element has such a long half life ($1.4*10^{17}$ a) that it is practically stable.

[0372] Natural titanium ${}_{22}\text{Ti}$ contains 0.0518 mole fraction of stable ${}^{50}\text{Ti}$ isotope.

[0373] ${}^{50}_{22}\text{Ti}$ (49.9447912 u)+ ${}^1_1\text{H}$ (1.007825032 u) \rightarrow ${}^{51}_{23}\text{V}$ (50.9439595 u)+0.008656732 u (8.063399589 MeV)

[0374] ${}^{51}\text{V}$ is a stable isotope of ${}_{23}\text{V}$.

[0375] Primary fusion reactions between titanium isotopes and hydrogen release a lot of energy.

[0376] The fusion of 1 mol ${}_{22}\text{Ti}$ (47.867 g) with 1 mol ${}^1_1\text{H}$ (1.0078 g) releases 0.0825 mol* 96.48533 kJ/mol* $5.167490449*10^6+0.0744$ mol* 96.48533 kJ/mol* $6.831730031*10^6+0.7372$ mol* 96.48533 kJ/mol* $6.757958399*10^6+0.0541$ mol* 96.48533 kJ/mol* $7.951438097*10^6+0.0518$ mol* 96.48533 kJ/mol* $8.063399589*10^6=652667538$ kJ= 652667 MJ= 181296 kWh of energy that is converted to thermal energy within the space surrounded by the gamma radiation shield made of, e.g., lead metal.

[0377] In case the released energy, about 650 000 MJ, is utilized directly as thermal energy, the amount of thermal energy is comparable to burning about 16800 liters of diesel oil, because burning diesel releases thermal energy about 38.6 MJ/liter (10.7 kWh/liter).

[0378] In case the released energy, about 180 000 kWh, is utilized in an electric generator based on e.g. Rankine cycle that has about 30% efficiency at relatively low fluid temperatures (e.g. 400° C.), 54000 kWh of electric energy is produced. Assuming that an electric car travelling at 80 km/h consumes about 20 kWh/100 km, the amount of electric energy produced from almost 48 g of titanium and slightly over 1 g of hydrogen is enough for driving that electric car for 270000 kilometers. Estimating that up to 10% of the electricity (COP=10) is used for operating the fusion system, about 240000 km driving distance is still feasible with the single fuel cartridge.

Example 10

[0379] Reaction material for the thermal-energy producing system was prepared from the following constituents. Nickel nanopowder (40 g) having an average particle size of 10 nm was mixed with 10 g of multiferroic bismuth ferrite BiFeO_3 crystallite powder having particle size range of about 100 nm-1000 nm. BiFeO_3 crystallite powder was prepared by mechanically crushing commercial BiFeO_3 sputtering target to powder. The precursors for the catalyst enhancing the formation of Rydberg matter comprised 85 wt % iron oxide Fe_2O_3 , 12 wt. % potassium hydroxide KOH and 3 wt % aluminum oxide Al_2O_3 . The precursor mixture was heated to 400-450° C. in the presence of hydrogen gas to form Fe_3O_4 : $\text{K}_2\text{O}, \text{Al}_2\text{O}_3$. The calcined catalyst powder was then mechanically crushed to catalyst nanopowder that had particle size range of about 10-100 nm. About 2.0 g of the catalyst nanopowder was added to the Ni— BiFeO_3 mixture and the powder mixture was placed to the reaction cartridge.

[0380] The reaction container was connected to a hydrogen gas line receiving hydrogen gas from a pressurized hydrogen gas bottle. The reaction container was also connected to the cooling fluid circulation. The reaction container was pressurized with hydrogen gas to 20 bar (gauge) and slowly heated to 400° C. with an electric resistance heater. The reaction material was capable of standing temperatures up to at least about 630-650° C. without losing its ability to generate heat energy.

[0381] Variable current was applied to the metal coil surrounding the reaction material in the reaction container to polarize the multiferroic material and sustain the generation of heat energy.

[0382] The presence of some water vapor impurity in the reaction container possibly helped to keep BiFeO_3 and Fe_3O_4 ; K_2O , Al_2O_3 powders in active form by preventing the reduction of iron oxides into elemental iron and sintering of the powders.

[0383] It is assumed that the multiferroic crystallite powder was polarized by the variable magnetic field within the reaction material and the local electric field due to the polarization was capable of accelerating electrons or protons to excite electrons to Rydberg states. The variable magnetic field was generated with an alternating current fed to a metal coil around the reaction container. The frequency of the alternating current could be adjusted up to the megahertz (MHz) range to provide control of the solid state fusion reactions inside the reaction container.

[0384] The system produced mostly relatively low energy photons (X-ray photons or deep UV photons) and the gamma radiation was very weak. In spite of that the system generated at least 5 kW of thermal energy with less than 1 kW input power. It is herein hypothesized that excitation state of the metastable fused nucleus (e.g. nickel-hydrogen) was so long-lived that the excitation state of the nucleus was capable of decaying via the energy transfer to K-shell electrons and resulting in X-ray photon emission. Generated thermal energy was removed by the cooling fluid circulation from the reaction container. The amount of collected thermal energy was at least 6 times larger than the energy used for pre-heating and controlling the reaction container ($\text{COP} > 6$). After the tests the reaction cartridge was de-pressurized and let to cool to room temperature for several days while the amount of residual radiation was monitored. Highly radioactive isotopes were not observed.

INDUSTRIAL APPLICABILITY

[0385] Apparatuses and methods based on various embodiments of the present invention produce very cheap thermal and electric energy. Targets for the utilization of the thermal energy produced by the reaction container comprise real estates for heating or cooling, farms, factories, houses, blocks of flats, green houses, private persons, private companies or public companies melting ice and snow from streets, roads, bridges and air ports. Applications for the utilization of thermal energy also comprise adsorption cooling especially in tropical or subtropical climates for the cooling of buildings, production of purified water by distillation or by freezing water into ice with adsorption cooling and producing fresh water from melted ice, and unit processes in chemical industry where solutions are fractionated into separate components, solutions are evaporated until a solid product is obtained or moist products are dried.

[0386] Targets for the utilization of electricity made from the thermal energy produced by the reaction container comprise farms, houses, blocks of flats, other real estates, green houses, factories, water purification plants, automotive industry, vehicles, cars, trucks, trains, ships and air planes.

[0387] It will be appreciated by those skilled in the art that various modifications and changes can be made without departing from the scope of the invention. Similar other modifications and changes are intended to fall within the scope of the invention, as defined by the appended claims.

1. A method of producing energy, comprising providing a reaction container (350) comprising reaction material (320), the reaction material (320) comprising electrically polarizable dielectric material and metallic material, pressurizing the reaction container (350) with hydrogen gas, activating hydrogen molecules in the hydrogen gas to provide atomic hydrogen, polarizing the dielectric material to produce an electric field, pulling hydrogen ions with the electric field from the metallic surface or ionizing the atomic hydrogen in the electric field to provide hydrogen ions, and accelerating hydrogen ions in the electric field, wherein part of the accelerated hydrogen ions tunnels through a Coulomb barrier between the hydrogen ions and atomic nuclei of the reaction material to fuse the hydrogen ions with the atomic nuclei of the reaction material to release energy.

2. The method according to claim 1, wherein the metallic material is capable of forming active hydrogen material comprising interstitial and/or electrically conductive metal hydrides, such as transition metal hydrides, in particular nickel, titanium, zirconium, hafnium, platinum group metal or other metal capable of forming metallic metal hydride.

3. The method according to claim 2, wherein the resistivity of the active hydrogen material is smaller than $1000 \mu\Omega\text{cm}$, preferably smaller than $500 \mu\Omega\text{cm}$, in particular smaller than $100 \mu\Omega\text{cm}$.

4. The method according to claim 2 or 3, wherein the active hydrogen material comprises a hydrogen storage alloy, electrically conductive hydrogenation catalyst, material capable of forming binary metal hydride consisting of a metal and hydrogen, or material capable of forming ternary metal hydride consisting of a first metal, a second metal and hydrogen.

5. The method according to any of the preceding claims, wherein the metallic material comprises transition metal having hydrogen in the form of hydride and/or hydrogen with a metallic bond.

6. The method according to any of the preceding claims, wherein the metallic material is in the form of nanopowder comprising metallic nanoparticles.

7. The method according to claim 6, comprising enhancing and focusing the electric field locally by the metallic nanoparticles.

8. The method according to any of the preceding claims, wherein the dielectric material comprises piezoelectric material, pyroelectric material and/or multiferroic material, which is polarized by mechanical vibration, temperature variation, and/or magnetic field, respectively.

9. The method according to any of the preceding claims, comprising initiating the fusion reactions at the nanoscale, at least one dimension being less than 100 nm.

10. The method according to any of the preceding claims, wherein the dielectric material is in the form of a powder or nanoporous material.

11. The method according to any of the preceding claims, wherein the reaction material comprises powdery material and/or porous material.

12. The method according to claim 11, wherein the reaction material comprises coated porous material comprising porous electrically polarizable crystalline material and metal-

lic nanoparticles arranged on pore surfaces of the porous electrically polarizable crystalline material.

13. The method according to any of the preceding claims, comprising keeping the temperature of the reaction material at a range of 100-1200° C., preferably at 300-900° C., in particular at 400-700° C.

14. The method according to any of the preceding claims, wherein the reaction material further comprises material promoting the formation and storage of Rydberg matter, said material preferably being arranged near the electrically polarizable dielectric material or to the surface of the electrically polarizable dielectric material.

15. The method according to claim **14**, comprising accelerating electrons in the electric field in addition to hydrogen ions and wherein the electric field strength is capable of producing a kinetic energy for the hydrogen ions and electrons high enough to excite electrons in the reaction material to Rydberg states and to form Rydberg matter.

16. The method according to claim **14** or **15**, comprising colliding at least part of the Rydberg matter with ions or electrons accelerated in an electric field so as to induce a Coulomb explosion of the Rydberg matter to produce high-energy hydrogen ions, and

fusing at least part of the high-energy hydrogen ions with atomic nuclei of the reaction material so as to release energy.

17. The method according to any of claims **14-16**, wherein said material promoting the formation and storage of Rydberg matter is in the form of catalytic nanopowder.

18. The method according to any of claims **14-17**, wherein the reaction material comprises paracrystalline material doped with an element capable of forming Rydberg matter.

19. The method according to claim **18**, wherein the paracrystalline material comprises

a metal oxide mixture comprising a first metal oxide and a second metal oxide, the metal of the first metal oxide being capable of changing its oxidation state in reducing atmosphere and the metal of the second metal oxide is stable and does not change its oxidation state in reducing atmosphere,

nickel mixed with alumina and/or chromia,

nickel oxide mixed with alumina and/or chromia,

iron mixed with alumina and/or chromia,

iron oxide mixed with alumina and/or chromia, or

copper-zinc alloy mixed with alumina and/or chromia.

20. The method according to claim **18** or **19**, wherein the doping element capable of forming Rydberg matter possesses Rydberg states due to the excitation of an electron of the element and is capable of becoming a Rydberg atom, the element preferably comprising Li, Na, K, Rb, Cs, N, Ni, Ag, Cu, Pd, Ti or Y.

21. The method according to any of claims **14-20**, wherein in the reaction container, at least part of the electrons or protons are accelerated to 10-20 eV kinetic energy, preferably to a kinetic energy below the amount of energy required for ionizing hydrogen atom, to create hydrogen Rydberg atoms.

22. The method according to any of claims **14-21**, wherein the material promoting the formation and storage of Rydberg matter is capable of promoting the formation of potassium and/or hydrogen Rydberg atoms, in particular potassium isotope ^{39}K and/or ^{41}K Rydberg atoms and/or hydrogen isotope ^1H , ^2H and/or ^3H Rydberg atoms.

23. The method according to any of claim **22**, wherein the potassium and/or hydrogen Rydberg atoms form clusters of Rydberg atoms to form Rydberg matter.

24. The method according to any of claims **14-23**, wherein the material promoting the formation and storage of Rydberg matter comprises

styrene catalyst,

ammonia synthesis catalyst,

high temperature water gas shift catalyst comprising potassium doped iron oxide and/or potassium doped lanthanum niobate,

Fischer-Tropsch catalyst comprising metals and metal oxides of cobalt, iron, ruthenium and/or nickel doped with copper or group 1 alkali metals, or

hydrogenation catalyst comprising platinum, palladium, rhodium, ruthenium, alloys of Pt, Pd, Rh and Ru, Raney nickel, Urushibara nickel and alkali metal doped nickel oxide, preferably Ni_2O_3 and non-stoichiometric Ni_{1-x}O doped with alkali metal, preferably potassium, wherein x is a non-integer in a range of about 0.005-0.1, preferably about 0.02.

25. The method according to any of claims **14-24**, wherein in the reaction material

the amount of said dielectric material is 5-80 wt %,

the amount of said metallic material is 15-90 wt %, and

the amount of said material promoting the formation and storage of Rydberg matter is 1-10 wt %.

26. The method according to any of claims **14-25**, wherein the electric field is adapted to accelerate hydrogen ions and electrons to a first kinetic energy sufficient to form Rydberg atoms in the reaction material,

the Rydberg atoms are attracted together to form condensed Rydberg matter,

the condensed Rydberg matter is destabilized by ionization of the said condensed Rydberg matter to induce Coulomb explosion so as to accelerate the hydrogen ions separated from the condensed Rydberg matter due to repulsive force to a second kinetic energy, and

at least part of the accelerated hydrogen ions tunnels through a Coulomb barrier between the hydrogen ions and atomic nuclei of the reaction material so as to release energy.

27. The method according to any of the preceding claims, wherein the energy released is removed from the reaction container as thermal energy.

28. The method according to any of the preceding claims, wherein the reaction container is shielded with a heavy metal mantle for converting radiation released in the fusion process into thermal energy.

29. A nuclear fusion system (**300**) for producing thermal energy, the system comprising

a reaction container (**350**),

reaction material (**320**) within the reaction container (**350**), the reaction material comprising electrically polarizable dielectric material and metallic material,

hydrogen gas source (**306**) connected to the reaction container (**350**) for pressurizing the reaction container (**350**) with hydrogen gas,

heat exchange unit (**314**) for removing thermal energy produced in the reaction container,

wherein the system further comprises

means for polarizing the dielectric material in order to produce an electric field within the reaction material,

- means for activating hydrogen molecules into hydrogen atoms and ionizing hydrogen atoms in order to accelerate the hydrogen ions in the electric field so that they can tunnel through a Coulomb barrier between the hydrogen ions and atomic nuclei of the reaction material to fuse the hydrogen ions with the atomic nuclei of the reaction material to release energy.
- 30.** The system according to claim **29**, comprising a heater (**322**) for heating the reaction material (**320**).
- 31.** The system according to claim **29** or **30**, comprising temperature measurement system (**328**, **334**) for measuring the temperature of the reaction material (**320**) and from the heat exchange unit (**314**), pressure measurement system (**313**) for measuring hydrogen gas pressure, and a control system (**304**) adapted to receive input from the temperature measurement system (**328**, **334**) and the pressure measurement system (**313**) and to control the heat exchange unit (**314**) and/or hydrogen gas pressure, and optionally the heater (**322**).
- 32.** The system according to any of claims **29-31**, wherein the hydrogen gas source (**306**) comprises a pressurized hydrogen gas bottle, metal hydrides heated to release hydrogen gas, or source of chemical reactions releasing hydrogen gas.
- 33.** The system according to any of claims **29-32**, wherein the electrically polarizable dielectric material comprises piezoelectric material and said means for polarizing the dielectric material to create electric field comprise a transducer (**550**) for inducing mechanical vibrations to the piezoelectric material for creating said electric field.
- 34.** The system according to any of claims **29-33** wherein the electrically polarizable dielectric material comprises multiferroic material and said means for polarizing the dielectric material to create electric field comprise an electrical coil (**518**) for inducing a magnetic field to the multiferroic material for creating said electric field.
- 35.** The system according to any of claims **29-34**, comprising a cooling fluid mantle (**702**) around the reaction container (**708**), a radiation shield mantle (**709**) around the cooling fluid mantle (**702**), and a thermal insulation mantle (**710**) around the radiation shield mantle (**709**).
- 36.** The system according to any of claims **29-35**, wherein the reaction material comprises dielectric material in the form of particles (**1004**, **1102**, **1108**) having a size of 10-10000 nm mixed with metallic material in the form of nanoparticles (**1010**, **1114**, **1116**, **1118**) having a size of 0.5-100 nm.
- 37.** The system according to any of claims **29-36**, wherein the reaction material further comprises material promoting the formation and storage of Rydberg matter.
- 38.** A fusion energy production process, comprising providing a matrix of porous reaction material, filling the pores of the matrix with hydrogen molecules, breaking at least part of the covalent bonds of hydrogen molecules by activation to produce hydrogen atoms, exciting at least part of the hydrogen atoms into hydrogen Rydberg atoms so as to form Rydberg matter, colliding at least part of the Rydberg matter with ions or electrons accelerated in electric fields inside the reaction material so as to induce a Coulomb explosion of the Rydberg matter to produce high kinetic energy hydrogen ions, and fusing at least part of the high kinetic energy hydrogen ions with the atomic nuclei of the reaction material so as to release fusion energy.
- 39.** The process according to claim **38**, comprising using metal capable of forming metallic metal hydride for breaking the covalent bonds of hydrogen molecules.
- 40.** The process according to claim **38** or **39**, comprising using a catalyst for activating the hydrogen.
- 41.** The process according to any of claims **38-40**, comprising using electrons or hydrogen ions accelerated in an electric field or electromagnetic radiation for exciting the hydrogen atoms.
- 42.** The process according to any of claims **38-41**, wherein the Rydberg matter comprises a mixed-element Rydberg matter including hydrogen Rydberg atoms and other Rydberg atoms.
- 43.** The process according to any of claims **38-42**, wherein providing the target matter in the form of an electrically polarizable porous matrix, providing the hydrogen molecules in the form of pressurized gas conveyed to the pores of the porous matrix, and polarizing the porous matrix to induce nanoscale electric fields into the porous matrix for exciting the hydrogen atoms and/or accelerating the collision ions or electrons.
- 44.** A fusion energy reaction material comprising a porous or powder mixture of electrically polarizable dielectric material, preferably in porous or powdery form, metallic material capable of forming metallic metal hydride, preferably in nanoparticle form, and material capable of promoting the formation of Rydberg matter upon interaction with active hydrogen.
- 45.** Use of hydrogen-containing Rydberg matter and/or inverted Rydberg matter as an intermediate material for providing high-energy hydrogen ions capable of fusing with other atomic nuclei in a fusion energy production process.

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