Methods for making a neutron converter layer are provided. The various embodiment methods enable the formation of a single layer neutron converter material. The single layer neutron converter material formed according to the various embodiments may have a high neutron absorption cross section, tailored resistivity providing a good electric field penetration with submicron particles, and a high secondary electron emission coefficient. In an embodiment method a neutron converter layer may be formed by sequential supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In another embodiment method a neutron converter layer may be formed by simultaneous supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In a further embodiment method a neutron converter layer may be formed by in-situ metalized aerogel nanostructure development.
OTHER PUBLICATIONS


FIG. 1
Machine Aerogel Or Polymer Matrix To Selected Converter Layer Size

Dissolve Neutron Hardening Precursor In Supercritical CO\textsubscript{2} Fluid Above 31.1°C and 7.29 MPa

Infuse Supercritical CO\textsubscript{2} Fluid Into Aerogel Or Polymer Matrix

Lower Pressure

Reduce At Elevated Temperature

Infuse Conductive Precursor

Infuse SEE Element Precursor

FIG. 4
Machine Aerogel Or Polymer Matrix To Selected Converter Layer Size

Dissolve Neutron Hardening, Conductive, SEE Element Precursors In Supercritical CO₂ Fluid Above 31.1°C and 7.29 MPa

Infuse Supercritical CO₂ Fluid Into Aerogel Or Polymer Matrix

Lower Pressure

Reduce At Elevated Temperature

FIG. 5
Form Solution Of Alkoxide, Water, Alcohol, And Catalyst In Presence Of Metal Precursors

Add Carbon Nanotubes or Other Metal Precursors To Adjust Resistivity

Adjust Composition Of Alkoxide Solution, Water, Alcohol, And Catalyst To Control Rate Of Hydrolysis And Condensation To Form Metalized Aerogel

Dry Metalized Aerogel Using Supercritical CO₂ Fluid At 31.1°C and 7.29 MPa

FIG. 6
NANOSTRUCTURE NEUTRON CONVERTER LAYER DEVELOPMENT

CROSS-REFERENCE TO RELATED PATENT APPLICATION(S)

This patent application claims the benefit of and priority to U.S. Provisional Patent Application No. 61/777,480, filed on Mar. 12, 2013, the contents of which are hereby incorporated by reference in their entirety.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT

The invention described herein was made in the performance of work under a NASA contract and by employees of the United States Government and is subject to the provisions of Public Law 96-517 (35 U.S.C. §202) and may be manufactured and used by or for the Government for governmental purposes without the payment of any royalties thereon or therefore. In accordance with 35 U.S.C. §202, the contractor elected not to retain title.

BACKGROUND OF THE INVENTION

Ionizing radiation, and in particular neutrons, pose a hazard to crew, passengers, and equipment in the aerospace and other industries. For example, research indicates that for flights within the commercial height range, aircrew and frequent flying passengers may be subject to radiation dose levels significantly above that permitted for members of the public under statutory recommendations. Equipment and crews on spacecraft that for part or all of their flight profile enter into low earth orbit, or travel beyond low earth orbit, are subjected to even higher radiation risks than aircraft at commercial height ranges.

One hazard of neutron radiation is neutron activation, i.e., the ability of neutron radiation to induce radioactivity in most substances it encounters, including a person’s body tissues. The risk posed by radiation has long been recognized as one of the major challenges to frequent and long duration spaceflight.

To help address the risks posed by neutron radiation, effective neutron radiation absorbers and detectors are needed. However, materials for neutron radiation detection have rarely been studied extensively.

BRIEF SUMMARY OF THE INVENTION

The present invention provides methods for making a neutron converter layer. The various embodiment methods enable the formation of a single layer neutron converter material. The single layer neutron converter material formed according to the various embodiments may have a high neutron absorption cross-section, tailored resistivity providing a good electric field penetration with submicron particles, and a high secondary electron emission coefficient.

In an embodiment method a neutron converter layer may be formed by sequential supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In another embodiment method a neutron converter layer may be formed by simultaneous supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In a further embodiment method a neutron converter layer may be formed by in-situ metalized aerogel nanostructure development.

These and other features, advantages, and objects of the present invention will be further understood and appreciated by those skilled in the art by reference to the following specification, claims, and appended drawings.

BRIEF DESCRIPTION OF THE SEVERAL VIEWS OF THE DRAWINGS

The accompanying drawings, which are incorporated herein and constitute part of this specification, illustrate exemplary embodiments of the invention, and together with the general description given above and the detailed description given below, serve to explain the features of the invention.

FIG. 1 is a graph of neutron shielding effectiveness for various materials;
FIG. 2 is a graph of electrical conductivity over frequency for various carbon nanotube volumes;
FIG. 3 is an image of a silver nanoparticle infused single wall carbon nanotubes (“SWCNT”) polymer composite morphology;
FIG. 4 is a process flow diagram illustrating an embodiment method for forming a neutron converter layer by sequential supercritical fluid metallization of a porous nanostructure aerogel or polyimide film;
FIG. 5 is a process flow diagram illustrating an embodiment method for forming a neutron converter layer by simultaneous supercritical fluid metallization of a porous nanostructure aerogel or polyimide film; and
FIG. 6 is a process flow diagram illustrating an embodiment method for forming a neutron converter layer by in-situ metalized aerogel nanostructure development.

DETAILED DESCRIPTION OF THE INVENTION

For purposes of description herein, it is to be understood that the specific devices and processes illustrated in the attached drawings, and described in the following specification, are simply exemplary embodiments of the inventive concepts defined in the appended claims. Hence, specific dimensions and other physical characteristics relating to the embodiments disclosed herein are not to be considered as limiting, unless the claims expressly state otherwise.

The word “exemplary” is used herein to mean “serving as an example, instance, or illustration.” Any implementation described herein as “exemplary” is not necessarily to be construed as preferred or advantageous over other implementations.

The various embodiments will be described in detail with reference to the accompanying drawings. Wherever possible, the same reference numbers will be used throughout the drawings to refer to the same or like parts. References made to particular examples and implementations are for illustrative purposes, and are not intended to limit the scope of the invention or the claims.

As used herein “a high neutron absorption cross-section” may be a neutron absorption cross-section at or above 1.9 barns. For comparison purposes, the neutron absorption cross-sections of various materials are illustrated in Table 1.

<table>
<thead>
<tr>
<th>Material</th>
<th>Density (g/cm³)</th>
<th>Neutron Abs x-sections (barns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminum</td>
<td>2.7</td>
<td>0.212</td>
</tr>
</tbody>
</table>
As used herein “a high electron emission coefficient” may be a secondary electron emission coefficient (“SEE”) greater than 1.

As used herein “tailored resistivity” may be resistivity greater than or equal to 10^7 Ohms/cm and less than or equal to 10^9 Ohms/cm.

Materials for neutron radiation detection that can provide a high neutron absorption cross-section, high electron emission coefficient, and tailored resistivity have rarely been studied. Multiple layers have been used to attempt to achieve a high neutron absorption cross-section, high electron emission coefficient, and tailored resistivity, but there are a number of disadvantages to using multiple layers, in particular, the inability to achieve a material with a high neutron absorption cross-section, high electron emission coefficient, and tailored resistivity without disrupting other functions of the material. Multiple layer material has required the use of large amounts of filler material to achieve a high neutron absorption cross-section, high electron emission coefficient, and tailored resistivity. The use of filler material has resulted in increasing the weight of the multiple layer material because fillers are generally denser than the matrix of the multiple layer material, complexity in manufacture of the multiple layer material, and cost increases for the multiple layer material as larger amounts of neutron attenuating filler material are added. Additionally, processability of the multiple layer material decreases as filler volume increases and negative impacts on other desirable properties of the multiple layer material occur as filler volume increases.

The present invention provides methods for making a neutron converter layer. The various embodiment methods enable the formation of a single layer neutron converter material. The various embodiments may enable the development of a neutron converter layer formed as a one layer porous nanostructure or a one layer solid film. The single layer neutron converter material formed according to the various embodiments may have a high neutron absorption cross-section, tailored resistivity providing a good electric field penetration with submicron particles, and a high electron emission coefficient. In some embodiments, a high neutron absorption cross-section may be achieved by the use of lithium (Li), boron (B), and/or gadolinium (Gd) as precursors. In some embodiments, a high electron emission coefficient may be achieved by the use of Magnesium Oxide (MgO) and/or Cesium Iodide (CsI) as precursors.

Neutron shielding materials for aerospace applications are being developed under the Materials International Space Station Experiments ("MISSSE") program. Emerging materials such as boron nitride nanotubes ("BNNT") and single wall carbon nanotubes ("SWCNT") as well as B, hexagonal boron nitride (h-BN), and Gd nanoparticles have been studied using a neutron exposure lab with a 1 Curie (Ci) americium/beryllium source. The preliminary study indicates that BNNT, h-BN, and Gd exhibited excellent neutron radiation shielding effectiveness compared with polyethyl-ene. Polymers containing high nitrogen (N) composition, such as polyimides, showed good neutron shielding effectiveness compared with non-nitrogen containing polymers. All N, B, and Gd possess high neutron absorption cross-sections compared with other elements and exhibited excellent neutron shielding effectiveness (i.e., above 0.1 mm^-1) as illustrated in the graph shown in FIG. 1.

Tailoring physical properties of nanocomposites has been the main focus of research activities, such as private industry ("PT") research activities, throughout the last decade to generate multifunctionalities for specific aerospace applications of interest. Especially for sensor and actuator applications, electrical conductivity and dielectric properties were effectively controlled as a function of the degree of dispersion, concentration, and orientation of the nano-inclusions. For example, the electrical conductivity can be controlled by several orders of magnitude with less than a 0.05% volume of carbon nanotubes as seen in FIG. 2.

Supercritical fluid ("SCF") metal infusion has been studied and a novel metallized nanotube polymer composites ("MNPC") has been developed to incorporate functional metals on the nanotube surface preferentially inside of a polymer matrix. Various metals (such as silver (Ag), gold (Au), platinum (Pt), palladium (Pd), iron (Fe), cobalt (Co), and nickel (Ni)) have been successfully metalized inside of a polymer and a SWCNT polymer composite. The metal particle size, infusion depth, and distribution may be controlled as a function of the SCF infusion conditions (e.g., time, temperature, and pressure). A silver nanoparticle infused SWCNT polymer composite morphology is shown in FIG. 3. Bright round dots represent reduced silver nanoparticles deposited on the SWCNT surface predominantly.

The following section describes embodiment methods to develop a nanostructure with high neutron absorption cross-section, high electron emission coefficient, and tailored resistivity. In the various embodiment methods a porous aerogel nanostructure (e.g., silica) with nanoparticles offering three functions may be processed systematically. Different approaches are performed to achieve the proposed nanostructure.

In an embodiment method a neutron converter layer may be formed by sequential supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In another embodiment method a neutron converter layer may be formed by simultaneous supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In a further embodiment method a neutron converter layer may be formed by in-situ metalized aerogel nanostructure development.

FIG. 4 is a process flow diagram illustrating an embodiment method 400 for forming a neutron converter layer by sequential supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. In step 402a a nanostructured aerogel or polymer matrix (e.g., a commercially purchased nanostructured aerogel or polyimide film) may be machined to a selected (e.g., appropriate to the intended application) dimension for the converter layer. In step 404 neutron hardening precursors (e.g., B and/or Gd) may be dissolved in a supercritical carbon dioxide (CO2) fluid above 31.1 degrees Celsius and 7.29 MPa (72.0 bar). In step 406 the supercritical CO2 fluid with the precursors dissolved in it may be infused into the aerogel or polymer (i.e., polyimide) matrix. In step 408 the pressure may be lowered, thereby trapping the infused metal precursors into the internal pores and surfaces of the aerogel or polymer matrix uniformly while the highly diffusive CO2 escapes rapidly. In step 410 the trapped and deposited metal precursors may be reduced.

### TABLE 1-continued

<table>
<thead>
<tr>
<th>Material</th>
<th>Density (g/cm³)</th>
<th>Neutron Absorption Sections (barns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Boron BN</td>
<td>2.27, BNNT 1.37</td>
<td>710 (²⁹B: 3835)</td>
</tr>
<tr>
<td>Gadolinium</td>
<td>79</td>
<td>40000</td>
</tr>
<tr>
<td>Lead</td>
<td>11.34</td>
<td>0.28</td>
</tr>
<tr>
<td>Titanium</td>
<td>4.54</td>
<td>5.0</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>gas</td>
<td>1.9</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>gas</td>
<td>0.33</td>
</tr>
<tr>
<td>Carbon</td>
<td>1.8-3.5</td>
<td>0.0035</td>
</tr>
</tbody>
</table>

at an elevated temperature to create nanoparticles. In step 412 conductive precursors may be infused, and in step 414 high SEE element precursors (e.g., MgO and/or CsI) may be infused to provide appropriate conductivity and SEE, respectively to the neutron converter layer.

FIG. 5 is a process flow diagram illustrating an embodiment method 500 for forming a neutron converter layer by simultaneous supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. Method 500 is similar to method 400 described above with reference to FIG. 4, except that in method 500 the neutron hardening, conductive, and SEE element precursors are applied together. In step 502 neutron hardening precursors (e.g., B and/or Gd), conductive, and SEE element precursors (e.g., MgO and/or CsI) may be dissolved in a supercritical carbon dioxide (CO₂) fluid above 31.1 degrees Celsius and 7.29 MPa (72.0 bar). Dissolution of the precursors is continued at an elevated temperature to create nanoparticles. In step 608 supercritical fluid metallization of a porous nanostructure aerogel or polyimide film. Method 600 may provide uniformly distributed functional nanoparticles incorporated into an aerogel nanostructure.

What is claimed is:

1. A method for forming a neutron converter layer, comprising:
   - machining an aerogel or polymer matrix to a selected converter layer size;
   - dissolving a neutron hardening precursor in a supercritical carbon dioxide (CO₂) fluid above a temperature of 31.1 degrees Celsius and a pressure of 7.29 MPa;
   - infusing the supercritical CO₂ fluid with the dissolved neutron hardening precursor into the aerogel or polymer matrix;
   - lowering the pressure to trap the infused neutron hardening precursor in the aerogel or polymer matrix;
   - infusing a conductive precursor into the reduced aerogel or polymer matrix;
   - infusing a secondary electron emission coefficient (SEE) element precursor into the reduced aerogel or polymer matrix.

2. The method of claim 1, wherein the neutron hardening precursor is boron or gadolinium.

3. The method of claim 2, wherein the SEE element precursor is magnesium oxide or cesium iodide.

4. The method of claim 3, wherein the neutron converter layer has a high neutron absorption cross-section, a high electron emission coefficient, and a tailored resistivity.

5. A method for forming a neutron converter layer, comprising:
   - machining an aerogel or polymer matrix to a selected converter layer size;
   - dissolving neutron hardening precursor, a conductive precursor, and a secondary electron emission coefficient (SEE) element precursor in a supercritical carbon dioxide (CO₂) fluid above a temperature of 31.1 degrees Celsius and a pressure of 7.29 MPa;
   - infusing the supercritical CO₂ fluid with the dissolved neutron hardening precursor, conductive precursor, and SEE element precursor into the aerogel or polymer matrix;
lowering the pressure to trap the infused neutron hardening precursor, conductive precursor, and SEE element precursor in the aerogel or polymer matrix; and reducing the aerogel or polymer matrix including the trapped infused neutron hardening precursor, conductive precursor, and SEE element precursor at an elevated temperature.

6. The method of claim 5, wherein the neutron hardening precursor is boron or gadolinium.

7. The method of claim 6, wherein the SEE element precursor is magnesium oxide or cesium iodide.

8. The method of claim 7, wherein the neutron converter layer has a high neutron absorption cross-section, a high electron emission coefficient, and a tailored resistivity.

9. A method for forming a neutron converter layer, comprising:
   forming a solution of an alkoxide solution, water, alcohol, and a basic catalyst in the presence of metal precursors;
   adjusting a composition of the alkoxide solution, water, alcohol, and the basic catalyst to control a rate of hydrolysis and condensation and form a metalized aerogel having radiation hardened nanoparticles and secondary electron emission coefficient (SEE) nanoparticles; and
   drying the metalized aerogel having radiation hardened nanoparticles and SEE nanoparticles using a supercritical carbon dioxide (CO₂) fluid at a temperature of 31.1 degrees Celsius and a pressure of 7.29 MPa to form a single layer neutron converter material.

10. The method of claim 9, wherein the metal precursors are selected from the group consisting of Gd₂O₃, B₂O₃, MgO, CsI, and any combinations thereof;
    wherein the radiation hardened nanoparticles include boron or gadolinium, and wherein the secondary electron emission coefficient (SEE) nanoparticles include magnesium oxide or cesium iodide.

11. The method of claim 9, further comprising adding a quantity of carbon nanotubes to adjust a resistivity of the metalized aerogel having radiation hardened nanoparticles and SEE nanoparticles.

12. The method of claim 11, wherein the neutron converter layer has a high neutron absorption cross-section, a high electron emission coefficient, and a tailored resistivity.

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