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(54) LARGE STRAIN TRANSPARENT MAGNETO-ACTIVE POLYMER **NANOCOMPOSITES**

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(57) ABSTRACT

A large strain polymer nanocomposite actuator is provided that upon subjected to an external stimulus, such as a magnetic field (static or electromagnetic field), an electric field, thermal energy, light, etc., will deform to thereby enable mechanical manipulations of structural components in a remote and wireless manner.

13 Claims, 19 Drawing Sheets

FIG. 1

FIG. 2

FIG.3

FIG. 5

FIG. 8

FIG. 12B

FIG. 13

1400C-

FIG, 14C

FIG. 15A

FIG. 15B

FIG. 17

LARGE STRAIN TRANSPARENT MAGNETO-ACTIVE POLYMER NANOCOMPOSITES

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of U.S. Provisional Patent application Ser. No. 61/563,962 entitled "LARGE STRAIN TRANSPARENT MAGNETO-ACTIVE POLY-¹⁰ MER NANOCOMPOSITES" filed on Nov. 28, 2011. The entirety of the above-noted application is incorporated by reference herein.

ORIGIN OF THE INVENTION

The invention described herein was made in the performance of work under a NASA contract and is subject to the provisions of Section 305 of the National Aeronautics and Space Act of 1958, Public Law 85-568 (72 Stat. 435; 42 ²⁰ U.S.C. 2457).

BACKGROUND

Actuators and smart materials are materials that exhibit mechanical deformation in response to an external stimulus such as an electric field, thermal energy, light, and electrochemical media. Actuators are of great interest due to their current and potential applications in aerospace structural components. Specifically, these materials, when actuated, perform a number of different functions, such as deploying solar arrays, antennas, flexible packaging, etc. Actuating these materials, however, via electro-resistive heating requires electrodes and wiring to the structural components. In addition, thermal shape memory polymers necessitates ³⁵ applying stress at a temperature above switching temperature to fix the polymer shape after recovery.

SUMMARY

The following presents a simplified summary in order to provide a basic understanding of some aspects of the innovation. This summary is not an extensive overview of the innovation. It is not intended to identify key/critical elements or to delineate the scope of the innovation. Its sole purpose is to 45 present some concepts of the innovation in a simplified form as a prelude to the more detailed description that is presented later.

In an aspect of the innovation, a remote actuation of a magnetic polymer nanocomposite by a magneto-static or electromagnetic field is disclosed, which will enable mechanical manipulations of the structural components in a remote and wireles s manner which is of high value in extreme environment.

In another aspect of the innovation, a method of producing 55 a large strain nanocomposite film is provided that includes mixing predetermined amounts of iron (III) acetylactonate, manganese acetyl acetonate, dodecanoic acid, 1, 2 dodecanediol, and 6 mmol of dodecylamine to form a mixture of nanoparticles, mixing the mixture with a predetermined amount of benzyl ether under a nitrogen blanket for a first predetermined amount of time, increasing a reaction temperature to approximately 150° C. for a second predetermined time, increasing the reaction temperature to approximately 300° C. for a second predetermined time, precipitating the mixture in methanol, and centrifuging and washing mixture with excess methanol.

In accordance with yet another aspect of the innovation, the method further includes dispersing the mixture of nanoparricles in tetrahydrofuran (THE), sonicating the mixture of nanoparticles for a fourth predetermined time, dissolving the 5 dispersion in THE, and mixing the dispersions with a surfacemodified manganese ferrite suspension MnFe₂O₄.

In still yet another aspect of the innovation, a method of actuating a large strain actuator is provided and includes providing a thermoplastic polyurethane (TPU) polymer nanocomposite film having manganese ferrite $(MnFe₂O₄)$ nanoparticles, applying an external stimulus to the nanocomposite film, deforming a shape of the nanocomposite film, and actuating an object caused by the deformation of the nano- $_{15}$ composite film.

In still yet another aspect of the innovation, the method further includes removing the external stimulus, recovering the shape of the nanocomposite film, and reproducing the deformation of the nanocomposite film upon application of the external stimulus.

In still yet another aspect of the innovation the innovation, a large strain actuator is provided that includes a nanocomposite film including manganese ferrite ($MnFe₂O₄$) nanoparricles added to a thermoplastic polyurethane (TPU) polymer film, wherein the nanocomposite film experiences a deformation is greater than 10 mm when exposed to an external stimulus, whereby the external stimulus includes one of a magnetic field, an electric field, thermal energy, and light.

To accomplish the foregoing and related ends, certain illustrative aspects of the innovation are described herein in connection with the following description and the annexed drawings. These aspects are indicative, however, of but a few of the various ways in which the principles of the innovation can be employed and the subject innovation is intended to include all such aspects and their equivalents. Other advantages and novel features of the innovation will become apparent from the following detailed description of the innovation when considered in conjunction with the drawings.

40 BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is an example flow-chart of a procedure of synthesizing hydrocarbon-coated iron manganese oxide nanoparticles in accordance with an aspect of the innovation.

FIG. 2 is an example flow-chart of a procedure of preparing nanocomposite films from the synthesized nanoparticles in accordance with an aspect of the innovation.

FIG. 3 is an illustration of the polymer nanocomposite films in accordance with an aspect of the innovation.

FIG. 4 is a thermo-gravimetric analysis (TGA) of surfacemodified $MnFe₂O₄$ nanoparticles in accordance with an aspect of the innovation.

FIG. 5 is graphical representation of an FT-IR spectrum of the organic surface modifier in accordance with an aspect of the innovation.

FIG. 6A is an illustration of a transmission electron microscopy (TEM) of organically-modified iron manganese oxide $(MnFe₂O₄)$ nanoparticles in accordance with an aspect of the innovation.

FIG. 6B is an illustration of a high-resolution image of the organically-modified $MnFe₂O₄$ nanoparticles in accordance with an aspect of the innovation.

FIG. 6C is an illustration showing a diffraction pattern of the organically-modified MnFe₂O₄ nanoparticles in accor-65 dance with an aspect of the innovation.

FIG. 7 is a graphical representation of a wide angle x-ray scattering (WAXS) spectrum illustrating lattice spacing and crystalline structure of the organically-modified $MnFe₂O₄$ in accordance with an aspect of the innovation.

FIG. 8 is a graphical representation illustrating magnetic properties of the surface-modified MnFe₂O₄ nanoparticles in accordance with an aspect of the innovation.

FIG. 9A illustrates a chemical structure and composition of thermoplastic polyurethane (TPU) in accordance with an aspect of the innovation.

FIG. 9B is a schematic of the organically-modified $MnFe₂O₄$ nanoparticles in accordance with an aspect of the innovation.

FIG. 9C is an illustration of the organically-modified $MnFe₂O₄$ nanoparticles dispersed in tetrahydrofuran (THF) in accordance with an aspect of the innovation.

FIG. 10 is a TEM scan of the MnFe₂O₄ nanoparticles in the TPU nanocomposite film in accordance with an aspect of the innovation.

FIGS. 11A and 11B are illustrations of scanning electron microscope (SEM) back-scatter micrographs of 0.5 and 6 wt $_{20}$ % surface-modified MnFe₂O₄/TPU nanocomposite films respectively after treatment with oxygen plasma in accordance with an aspect of the innovation.

FIG. 12A is a graphical representation illustrating a transparency of the surface-modified MnFe₂O₄/TPU nanocom- 25 posite films measured over a given wavelength in accordance with an aspect of the innovation.

FIG. 12B is an illustration showing a transparency of the 0.1 wt % surface modified $MnFe₂O₄/TPU$ nanocomposite film in accordance with an aspect of the innovation.

FIG. 13 is a graph illustrating a saturation magnetization, M_s , of the nanocomposite films versus concentration of the $MnFe₂O₄$ nanoparticles in accordance with an aspect of the innovation.

FIGS. 14A and 14B are illustrations of a schematic of a film position with respect to the magneto-static field in accordance with an aspect of the innovation.

FIG. 14C is an illustration of a color-coded displacement in the y-direction (out of the plane of the figure) for a 8 wt $\%$ ₄₀ surface-modified MnFe₂O₄ film in the magnetic field in accordance with an aspect of the innovation.

FIG. 15A is a graph depicting the maximum displacement, $\delta_{\nu max}$, versus loading of the magnetic nanoparticles in accordance with an aspect of the innovation.

FIG. 15B is a graph illustrating the displacement of the surface modified $MnFe₂O₄/TPU$ nanocomposites versus the applied magnetic field in accordance with an aspect of the innovation.

FIG. 16A is a graph illustrating a film displacement versus 50 time while cycling in the magnetic field in accordance with an aspect of the innovation

FIG. 16B is a graph illustrating a film displacement in accordance with an aspect of the innovation.

FIG. 17 illustrates an example flow chart of a procedure of 55 actuating a large strain nanocomposite film in accordance with an aspect of the innovation.

DETAILED DESCRIPTION

The innovation is now described with reference to the drawings, wherein like reference numerals are used to refer to like elements throughout. In the following description, for purposes of explanation, numerous specific details are set forth in order to provide a thorough understanding of the subject innovation. It may be evident, however, that the innovation can be practiced without these specific details. In other

instances, well-known structures and devices are shown in block diagram form in order to facilitate describing the innovation.

While specific characteristics are described herein (e.g., 5 thickness), it is to be understood that the features, functions and benefits of the innovation can employ characteristics that vary from those described herein. These alternatives are to be included within the scope of the innovation and claims appended hereto.

10 While, forpurposes of simplicity of explanation, the one or more methodologies shown herein, e.g., in the form of a flow chart, are shown and described as a series of acts, it is to be understood and appreciated that the subject innovation is not limited by the order of acts, as some acts may, in accordance with the innovation, occur in a different order and/or concurrently with other acts from that shown and described herein. For example, those skilled in the art will understand and appreciate that a methodology could alternatively be represented as a series of interrelated states or events, such as in a state diagram. Moreover, not all illustrated acts may be required to implement a methodology in accordance with the innovation.

Polymer nanocomposite actuators are of great interest due to their potential applications in aerospace structural components, micro-robotics, artificial muscles, temperature-sensitive switches and valves, magneto-driven biocompatible devices, "morphing" airframe or aircraft engine structures or self-deployable structures, e.g., large area solar arrays or antennae and habitats, etc. Polymer nanocomposite actuators so are materials that undergo mechanical deformation by application of an external stimulus such as a magnetic field, electric field, light, and thermal energy.

The innovation disclosed herein focuses on the remote actuation of a magnetic polymer nanocomposite by a mag-35 neto-static or electromagnetic field. Magneto-active materials are materials that exhibit magnetic properties coupled with mechanical deformation in a static or electromagnetic field. This type of actuation results in deformation which is recoverable upon removal of the field and is reproducible. This technology can be used for space deployable structures where a small compact, lightweight volume needs to undergo sudden large shape changes. It can also be extended to the actuation of structural components in aircrafts, e.g., wings or fan blades where a magnetic field can induce deformation of 45 components. Some examples of these technologies are given below.

Actuation and morphing of light weight structural materials have great impact in outperforming current aerospace components to the new generation of aerospace vehicles. Adaptive structures (soft and hard materials) have applications ranging from unmanned aerial vehicles (UAV), micro air vehicles (MAV), deployable antenna, satellite structures, remote light weight unlocking mechanisms, deployable structures on the Moon and Mars, morphing and adaptive wing skin. Mechanical manipulation of the structures in extreme outer space environments by wireless remote method is of great significance to space missions. Unlocking a compact volume to a large structure is essential for transportation of structures to the orbit or outer space. Adaptive materials 60 will enhance air vehicle maneuverability such as bio-inspired moving wings, where airplane wings could change depending on the altitude and mission. Shape change could result in reduced fuel consumption by change of structural components during takeoff, cruising and landing.

The innovation discloses a superparamagnetic polymer nanocomposite actuator films prepared by addition of superparamagnetic nanoparticle into the polymer films of both

thermoplastic polyurethane (TPU) and high stiffness polyimide resin. The TPU magnetic nanocomposites are called soft magneto-active materials and polyimide magnetic nanocomposites are called hard magneto-active nanocomposites.

While other approaches to magnetically responsive materials have been developed and demonstrated, the advantage of the innovation is that a much smaller amount of nanoparticles (less than 1%) is required to obtain large displacements (>10 mm) of the polymer film under an applied external stimulus. The magneto-actuation deformation increases with an io increasing magnetic nanoparticle content exponentially. As a result, these materials are much lighter in weight than other magnetically responsive materials and have other desirable attributes such as optical transparency.

In addition, nanocomposites prepared with other magnetic 15 nanoparticles, core-shell nanoparticles of a different chemical composition, exhibit resistive heating when placed in an alternating magnetic field. Nanocomposite films prepared from these nanoparticles experienced temperature rises as high as approximately 300° C. under these conditions. Such 20 temperature increases might be sufficient to initiate self-healing in nanocomposites films and fiber reinforced nanocomposites.

The TPU magneto-active polymer nanocomposites disclosed herein are both transparent and magnetically active 25 with low loading levels $(\leq 2 \le 2)$ of superparamagnetic nanoparticles. In addition, the TPU magneto-active polymer nanocomposites disclosed herein have been prepared by solvent casting as a thermoplastic elastomer. They can, however, be melt processed by injection molding, extrusion, which is 30 significantly important for high throughput industrial processes.

Magnetic actuation can be induced by applying a magnetic field (static or electromagnetic) to a magnet-active polymer composite. Magneto-active polymer composites are hybrid 35 materials composed of a polymer and magnetic material which exhibit overall magnetic properties. Magnetic nanoparticle polymer nanocomposites have great potential for large strain actuators due to their large particle number density, the large interfacial area between the magnetic nanopar- 40 ticles and the polymer matrix. Low loading levels of magnetic nanoparticles is important for aerospace applications since reduced weight is a critical driver for materials. Magnetic nanoparticles can be incorporated into soft polymer matrices to generate polymer nanocomposite actuators. This method 45 can be extended to structural components with higher glass transition temperatures to allow deformation above the glassy state.

Some known actuators include, lightweight aerogel magnetic actuators prepared by freeze-dried cellulose nanofibril 50 aerogels as templates for non-agglomerated growth of cobaltferrite, have shown actuation responses even in low magnetic fields. Coiling mechanisms and large deformation of spherical micron-sized iron particle polysiloxane have been disclosed for composites with particle loads of 20 to 77 wt %. ⁵⁵ Disclosed magnetic actuation of iron oxide (γ -Fe₂O₃) nanorods in poly (lactide-co-glycolide) biocompatible nanocomposites (10-30 wt %) could potentially stimulate cells to promote nutrient supply.

Further, epoxy/micron-sized strontium ferrite powder 60 composite (95.3 wt %) micro-actuators exhibited small deflections when tested both statically and dynamically. Electromagnetic actuation of nickel (Ni) nanowire cellulose nanocomposites (approximately 34 wt %) with both DC and AC currents generating constant and alternating magnetic field have also been disclosed. Magnetic-sensitive gels of chemically-crosslinked polymer networks with approximately 10

nm mono-domain magnetic nanoparticles undergo shape distortion when a magnetic field is applied.

Still further, the free energy of the swollen network containing both elastic and magnetic components has been studied as the basis for the shape change.

Nanocomposites of (3.5-6.5 nm) maghemite polystyrene exhibited structural supra-aggregate organization with a size of approximately 200 nm at volume fractions, $\phi \ge 5 \times 10^4$. Primary aggregates were formed at lower volume fractions (<5x $10⁴$) as shown by small angle x-ray scattering and transmission electron microscopy (TEM). The mechanical response of 1-10 wt % micron-sized $Fe₃O₄/polyvinyl$ alcohol magnetic hydrogels in low magnetic field (40mT) has also been disclosed.

Magnetic nanoparticles can be synthesized to generate different chemical compositions, shapes, sizes, and aspect ratios. These characteristics determine the magnetic strength of the nanoparticles. Magnetic nanoparticles below a critical diameter are super-paramagnetic, where the spin rotation is random, and the material can be magnetized and demagnetized upon application or removal of the magnetic field with no relaxation time. These superparamagnetic nanoparticles have single domains and respond quickly to a magnetic field above the blocking temperature. They also tend to agglomerate due to magnetic and van der Waals forces, which lower the nanoparticles surface area. The high coercivity of superparamagnetic particles is attributed to single domain effects. The increase in the aspect ratio also results in a significant increase in coercivity, i.e. the coercivity of Fe nanoparticles increased from 82 mT to 1 T when the aspect ratio was increased from 1.1 to 10. Magnetic nanoparticles have been synthesized by co-precipitation, thermal decomposition, microemulsion, and hydrothermal synthesis. Monodisperse metallic nanoparticles can be synthesizedby athermal decomposition method. This method involves reduction of organo-metallic compounds in high boiling point solvents containing surfactants as a stabilizing agent and polyol as the reducing agent.

As will be subsequently described, the innovation discloses the preparation and characterization of surface-modified manganese ferrite (MnFe₂O₄) thermoplastic polyurethane elastomer nanocomposites (0.1 wt %-8 wt %), which are capable of large deformations under applied magnetic fields. Due to the small particle size of the super-paramagnetic nanoparticles, the low particle loading (0.1 and 0.5 wt %) nanocomposites were transparent and exhibited large deformations in a static magnetic field.

Referring now to the drawings and specifically to FIG. 1, a method of synthesizing hydrocarbon-coated iron manganese oxide nanoparticles will be described. Specifically, at 102, 2 mmol of iron (III) acetylactonate, 1 mmol of manganese acetyl acetonate, 6 mmol dodecanoic acid, 10 mmol of 1, 2 dodecanediol, and 6 mmol of dodecylamine were mixed with 20-30 cc of benzyl ether under a nitrogen blanket for approximately 15 min. Then, at 104, the reaction temperature was then increased to approximately 150° C. for 30 minutes. At 106, the reaction temperature is increased to approximately 300° C. for approximately 30 minutes. At 108, the iron manganese oxide nanoparticles were precipitated in methanol after cooling and at 110, the nanoparticles are centrifuged and washed several times with excess methanol.

Referring to FIGS. 2 and 3, a method of preparing the nanocomposite film from the synthesized nanoparticles above is described. At 202, magnetic nanoparticles are dispersed in tetrahydrofuran (THE) and sonicated for approximately 5 minutes to generate visibly aggregate free dispersions. At 204, the TPU is dissolved in THE and mixed with surface-modified MnFe₂O₄/THF suspensions. At 206, the TPU/surface-modified $MnFe₂O₄/THF$ dispersions are sonicated for 30 minutes. At 208, the dispersions are solvent cast to generate nanocomposite films 300 (0.1-8 wt %) approximately 75-100 micron thick. At 210, the films are dried in a vacuum oven at a predetermined temperature and time to remove excess solvent. At 212, weight percentages of the nanocomposites are calculated based on $MnFe₂O₄$ content.

A thermogravimetric analyzer, using a controlled atmosphere of nitrogen, a temperature range of 25-800° C., and a scan rate of 10° C./minute, determines a change in weight in 10 relation to a change in temperature of the nanocomposite film. A Fourier transform infrared spectrometer was used to obtain an infrared spectrum of the nanocomposite film. In addition, a high resolution transmission electron microscopy (TEM) image of the nanocomposite film was obtained. Cryo-frac-15 tured surfaces of the nanocomposite film were exposed to air plasma for 3 minutes and another image of the nanocomposite film was obtained using a scanning electron microscope (SEM). A wide angle x-ray scattering was performed on a diffractometer configured in the Bragg-Brentano geometry 20 with Cu K- α (λ =1.5418 Å) radiation source and a linear position sensitive detector.

Magneto-mechanical testing was performed on a microload-pneumatic test rig. Tests were performed in stroke control at a rate of 0.5 mm/s. Full-field optical displacement imaging was used with a frame capture rate of 0.125 seconds. A film sample was placed vertically at a starting distance of approximately 50 mm from the magnet. A static magnet with a strength of 0.43 T (B_y (y=0)) was used. All three components of the magnetic field were measured by a triple-axis magnetometer. Only B_{ν} is acting on the film surface perpendicular to the x-z plane (B_x and B_z , were negligible and verified by the measurements). The z-variation of B_{ν} was negligible along the film z-axis within the experimental geometry constraints. The sample was moved toward the magnet using the test rig stroke, which resulted in the increasing magnetic field. The magnetic field, B_{ν} , variations with the position along the y-direction was measured in 0.5 mm increments and fit to a $6th$ order polynomial. Deflection of the film, δ_v , was monitored using the optical displacement system.

Surface modification of iron manganese oxide nanoparticles is essential to provide compatibility between the nanoparticles and the thermoplastic polyurethane elastomer matrix. The synthesis method resulted in iron manganese oxide nanoparticles with an organic modifier corona on the surface. Referring to FIG. 4, a thermo-gravimetric analysis (TGA) 400 of the surface-modified MnFe₂O₄ nanoparticles shows approximately 29 wt % hydrocarbon on the surface of the $MnFe₂O₄$ nanoparticles with a degradation temperature onset of 190° C. and a maximum degradation temperature of 50 291.3° C. Referring to FIG. 5, an FT-IR spectrum 500 of the organic surface modifier is illustrated and exhibits a band at 3337 cm⁻¹ corresponding to $-$ OH stretch possibly due to 1, 2 dodecanediol, or the presence of a hydroxyl group on the $MnFe₂O₄$ surface. The $-CH$ stretch of saturated aliphatic 55 hydrocarbons generally appears in the range of 3000 to 2800 cm^{-1} , whereas the bending appears at 1500 and 1300 cm^{-1} . The stretches observed at 2922.5 and 2852.6 cm^{-1} are due to the —CH stretch in C—CH₃, and to the —CH₂ presence in the aliphatic hydrocarbon chain of the organic modifiers. The 60 absorption peaks observed at 1430.8 and 1556.6 cm^{-1} are characteristic of the --CH bending stretches.

Referring to FIG. 6A, a TEM micrograph 600A of the organically-modified $MnFe₂O₄$ nanoparticles is illustrated. The observed separation between the nanoparticles is attributed to the organic surface modifier of the $MnFe₂O₄$ nanoparticles. High-resolution imaging 600B illustrates the pres-

ence of nearly uniform spherical nanoparticles with an average diameter of 6.11 ± 0.69 nm, see FIG. 6B, measured among 250 nanoparticles. FIG. 6C is an illustration showing the electron diffraction pattern 600C of the organically-modified MnFe₂O₄ nanoparticles where the diffraction pattern corresponding to hkl indices of 220, 311, 400, 422, 511 are identified.

Lattice spacing and the crystalline structure of the organically-modified iron manganese oxide nanoparticles were studied using wide angle x-ray scattering (WAXS), see FIG. 7. The diffraction peaks from the WAXS spectrum 700 shows an excellent match to the relative hkl indices of $MnFe₂O₄$ in the PDF database. Table 1, below, lists calculated spacing $d(A)$, based on the relative diffraction peaks of the bulk WAXS spectra of synthesized MnFe₂O₄ nanoparticles and their matched hid indices.

TABLE 1

20	WAXS Diffraction Peaks							
				\mathcal{R}	4		6	
25	d MnFe ₂ O ₄ hkl	2.96 2.97 220	2.54 2.54 311	2.11 2.10 400	1.72 1.72 422	1.62 1.62 511	1.49 1.49 440	1.27 1.27 622

A material's magnetic characteristic depends on its chemical composition, size, and aspect ratio. $MnFe₂O₄$ has a Curie temperature, T, of 300° C. and is super-paramagnetic at diameters at least up to 9.9 mu. FIG. 8 is an illustration of a graph 800 illustrating the magnetic properties of the surface-modified MnFe₂O₄ nanoparticles, which were measured using an alternating-field gradient magnetometer and, owing to their small size, exhibited closed-loop, super-paramagnetic behavior. The magnetization of a permanent magnet after removal of the external magnetic field is referred to as remanence. The saturation magnetization, M, is the magnetic moment of elementary atoms per unit weight where all of the dipoles are 40 aligned parallel. The reverse magnetic field required to reduce a materials magnetization to zero while the sample is in the magnetic field is called coercivity, H. The surface-modified $MnFe₂O₄$ nanoparticles have a saturation magnetization, M, of 33.73 Am2/kg, a remanent magnetization of 125.1 mAm2/ kg, a coercivity, H_c of 0.593 mT and a coercivity of remanence, H_{av} of 4.6 mT.

TPU elastomers have been widely used as stimuli-responsive polymers due to their segregated two-phase structure. TPU is comprised of hard and soft segments, a chain extender, has a tunable glass transition temperature, and mechanical properties. Soft segments could crystallize and act as physical crosslinks enabling shape recovery effects. TPU used in this study was synthesized by polycondensation reaction of 4-4 methylenediphenylene isocyanate (MDI) and polyol using butanediol as chain extender. Its microstructure is reported to consist of 9.9% hard segments, 58.2% butanediol chain extenders, and 31.8% adipate soft segments. It has shown thermal shape memory effects when used as a host matrix for zinc nanorods and multiwall nanotubes.

The surface-modified $MnFe₂O₄$ nanoparticles were dispersed in TPU containing soft segments of aliphatic alkyl chain to generate nanocomposites films. The chemical structure and composition 900A of the TPU, and a schematic 900B of the organically-modified $MnFe₂O₄$ nanoparticles are shown in FIGS. 9A and 9B respectively. A stable dispersion of organically-modified $MnFe₂O₄$ nanoparticles in THF 900

C was obtained, see FIG. 9C, which was then mixed with a solution of TPU in THE to generate the nanocomposites films.

The presence of long-chain aliphatic hydrocarbons promotes the compatibility between the inorganic MnFe₂O₄ 5 nanoparticle and the polyurethane due to the presence of aliphatic hydrocarbon moieties in the polyurethane polymer chains. This will improve the dispersion of the $MnFe₂O₄$ nanoparticles within the TPU polymer matrix. The surfacemodified $MnFe₂O₄/TPU$ nanocomposite films were prepared with particle loadings of 0.1, 0.5, 1, 2, 4, 6 and 8 wt % (0.025, 0.126, 0.252, 0.51, 1.03, 1.57, 2.13 vol. %), based upon the weight/volume of the metallic core ($\rho_{MmFe2O4}$ =4.76 g/cc and ρ_{TPI} =1.19 g/cc). The low weight/volume particle loadings of the nanocomposites were critical to achieving overall light-15 weight nanocomposites.

Referring to FIG. 10, a TEM scan 1000 examined the dispersion of 2 wt % surface-modified MnFe₂O₄ nanoparticles in the TPU nanocomposite film after cryo-microtoming of the film. A variety of nanoparticle clusters, ranging from a few nanoparticles, to larger nanoscale clusters, and micronsized aggregates were observed. Magnetic and van der Waals attractive forces result in aggregations of the nanoparticles within the film. Dispersion of 4 nm maghemite (γ -Fe₂O₃) nanoparticles in polystyrene occurred only at loading levels below 0.01 vol. % whereas, 200 mn supra-aggregates occurred at loading levels above 0.05 vol. %.

FIGS. 11A and 11B illustrate the SEM back-scatter micrographs 1100A, 1100B of the 0.5 and 6 wt % surface-modified $MnFe₂O₄/TPU$ nanocomposite films respectively after treat- 30 ment with an oxygen plasma. The more dense $MnFe₂O₄$ nanoparticle aggregates appear as bright areas on the SEM micrographs. The aggregate size ranges between 1-3 microns (average of 1. 7 microns) for 0.5 wt % and 1.1-2.9 microns (average of 2 microns) for 6 wt % MnFe₂O₄-loaded films. The 35 nanometer-size magnetic nanoparticles and clusters are not resolved at this SEM magnification. The film containing 6 wt % surface-modified $MnFe₂O₄$ nanoparticles exhibited increased nanoparticle density on one side indicating settling of the heavier $MnFe₂O₄$ nanoparticles during solvent evapo- 40 ration, see FIG. 11B. This settling effect was not observed for 0.5, 1, 2, or 4 wt % surface modified MnFe₂O₄. However, settling was more significant for 8 wt % surface-modified $MnFe₂O₄/TPU$ nanocomposite film.

A graph 1200A illustrating a transparency of the surface- 45 modified $MnFe₂O₄/TPU$ nanocomposite films measured over a wavelength range of 400-700 mn is shown in FIG. 12A. The neat TPU film showed a transmission of 97-90% in the range of 700-550 mu, while dropping from 90% to 74.4%, between the 550-400 nm range. The transmission of the 0.1 wt $\%$ 50
the 550-400 nm range. The transmission of the 0.1 wt $\%$ 50 nanocomposite films was comparable with the neat TPU film where a slight decrease in transmission was observed from 400-460 mu. The decrease in transmission of the 0.5 wt % surface modified $MnFe₂O₄/TPU$ nanocomposite in the 700-550 mn range was 91 to 75%, and 75% to 49.5% for the 55 wavelength range of 550-400 nm. Further increase in the loadings of the surface modified MnFe₂O₄ to 1 wt % resulted in a decrease in transmission from 73 to 42% for the wave length range of 700-550 nm, and a further decrease of 42-20% for the wavelength range of 550-400 nm. The TPU nanocom- 60 posite containing 2 wt % surface-modified MnFe₂O₄ didn't show a significant transmission decrease in the range of 700- 550 mn range and had a transmission of 69.35 to 33%. However, the transmission in the range of 550-492 nm significantly dropped from 33% to 20% and below 10% for the wavelengths below 470 nm. The 4 wt % surface-modified $MnFe₂O₄/TPU$ exhibited a transmission of 53.6 to 16.7% in

the range of 700-61 0 mm with a sharp drop to below 10% of wavelengths shorter than 590 nm. The TPU/surface-modified $MnFe₂O₄$ nanocomposite with particle loadings of 0.1 and 0.5 wt % were transparent, 1 and 2 wt % were semi-transparent, and 4 wt % was opaque. FIG. 12B illustrates the transparency of the 0.1 wt % surface modified MnFe₂O₄/TPU nanocomposite film 1200B, which is displayed in front of a NASA logo.

Magnetization of the surface-modified MnFe₂O₄/TPU io nanocomposites was measured to provide information about saturation magnetization, and coercivity. The coercivity of all surface-modified MnFe₂O₄/TPU nanocomposite films was in the range of 0.8 ± 0.1 mT. FIG. 13 is a graph 1300 illustrating the saturation magnetization, M_s , of the nanocomposite films versus concentration of the MnFe₂O₄ nanoparticles. The normalization of the magnetic moment versus magnetic field was performed based on the total weight of the nanocomposite film (TPU+surface modified MnFe₂O₄). The plot of saturation magnetization of the nanocomposite films versus magnetic nanoparticle concentration shows an exponential trend as follows:

$$
M_s = A\omega^B \tag{1}
$$

where A is 380.2 ± 0.033 and B is 1.02 ± 0.038 with $r^2 = 0.99$.

It should be noted that, the magnetic moment versus magnetic field was also normalized with respect to the weight of the magnetic nanoparticles contained in each nanocomposite film. This normalization yielded constant values for coercivity, H_c, 0.8 ± 0.1 mT and magnetization saturation, M_s, 0.04 ± 0.01 mAm²/kg.

As illustrated in Equations (2)-(6), the nanocomposite films have magnetic characteristics that result from the embedded super-paramagnetic $MnFe₂O₄$ nanoparticles. These films were placed in a static magnetic field, \vec{H} , where a magnetic force, \vec{F} , is applied that is proportional to the magnetic potential, \vec{U} . The magnetic moment, \vec{M} , is related to the magnetic field, \vec{H} , with a susceptibility, γ . The force acting on the volume of a magnetic material depends on the magnetic field moment and the rate of the magnetic field change in that direction.

$$
\frac{d\overrightarrow{M}}{d\overrightarrow{H}} = \chi \tag{2}
$$

$$
\overrightarrow{U} = \int_0^{H_0} \overrightarrow{M} d\overrightarrow{H}
$$
 (3)

$$
\overrightarrow{U} = (1/2)\chi H_O \tag{4}
$$

$$
\vec{F} = -\overline{\nabla U} \tag{5}
$$

$$
F_y = M_y \int_0^V \frac{dHy}{dy} \, dV \tag{6}
$$

Further, referring to Equations (7) and (8), the displacement of the magnetic film (8) is determined using the static deflection of a cantilever beam, where I is the moment of inertia, L, H, and b are length, width and thickness, respectively.

$$
\delta = F_y L^3 / 3EI \tag{7}
$$

$$
I = bH^3/12\tag{8}
$$

The magnetic field was induced by a static magnet with a magnetic field of B_y (y=0)=430 mT corresponding to the onset of saturation magnetization for the nanoparticles. FIGS. 14A and 14B illustrate a schematic 1400A, 1400B of the film position with respect to the magneto-static field. The test begins with the film positioned 50 mm from the magnet. The film is then moved toward the magnet at a rate of 0.5 mm/s using the test frame. Once the film is in close proximity of the magnet, the magnetic field causes the film to move gradually in the y direction. This deflection is measured using the optical displacement equipment and is given as displacement, δ_{ν} , for various points along the length of the film. FIG. 14C illustrates the color-coded displacement 1400C in the y-direction (out of the plane of the figure) for the 8 wt % surfacemodified $MnFe₂O₄$ film in the magnetic field.

Upon approaching the magnet, the film moves gradually in the y-direction where one end is fixed. However, the film eventually reaches a point where the magnetic force applied on the film is equal to the weight and the force required for the maximum deformation resulting in complete pulling of the film to the magnet. The separation distance at this point was the maximum displacement, δ_{ymax} .

FIG. $15A$ is a graph 1500\AA depicting the maximum displacement, δ_{max} , versus loading of the magnetic nanoparticles. The maximum displacement exhibits an exponential decay with the following fitting parameters:

$$
\delta_{\text{ymax}} = A \omega^B \tag{9}
$$

where $A=19.28\pm0.01$, $B=0.21\pm0.015$ with $r^2=0.99$.

Combining Equation (9) with Equation (1), an empirical equation correlating the maximum displacement, δ_{ymax} , to the film saturation magnetization, M_s , and magnetic nanoparticle weight percent can be proposed:

$$
\delta_{\text{ymax}} = 0.5 M_s \omega^{0.8l} \tag{10}
$$

This correlation suggests that the maximum displacement has a stronger dependence on the saturation magnetization than on the weight percent of the magnetic nanoparticles.

The maximum displacement increased significantly with increasing magnetic nanoparticle concentration particularly at low particle loadings and up to 2 wt %, see FIG. 15A. As expected, the nanocomposites containing more than 2 wt % reached their maximum deformation at even greater distances. For example, the maximum displacement for the 0.1 wt % (0.025 vol. %) nanocomposite is 11.1 mm whereas the maximum displacement for the 8 wt % (2.13 vol. %) nanocomposite is 30.42 mm. It is evident that surface-modified $MnFe₂O₄/TPU$ nanocomposites exhibit large displacements 55 even with a low particle load of 0.1 wt % (0.025 vol. %).

FIG. 15B is a graph 1500B illustrating the displacement of the surface modified $MnFe₂O₄/TPU$ nanocomposites versus the applied magnetic field. The displacement rate is lower for films containing low particle loads, and increases as the par- 60 ticle loading increases.

Because accurate control over actuation is critical to actuator performance, the recovery and response time of the films were examined. Cyclic deformation of the nanocomposite film containing 6 wt % $MnFe₂O₄$ nanoparticle loading was performed five times in a low magnetic field of $15.1 \text{ (mT)}₅B$ (y) _{<2}30.3 (mT).

FIG. 16A is a graph 1600A illustrating the film displacement versus time while cycling in the magnetic field. The imposed cyclic period time was 25 seconds with the film having an approximate 3 second lag time. The maximum 5 displacement of the films from cycle-to-cycle was constant and reproducible within the experimental conditions. The films also returned to their original position as the cycle returned to the low value of the magnetic field.

To determine the loss and hysteresis of the 6 wt % nanocomposite film in the magnetic field (15.1 mT<B(y)<30.3 mT) the film displacement 1600B is plotted in FIG. 16B for all five cycles. The traces from all five cycles were identical within experimental error. Neither hysteresis nor permanent deformation could be discerned from this test.

Referring to FIG. 17, a method of actuating a large strain nanocomposite film will be described. At 1702, a thermoplastic polyurethane (TPU) polymer nanocomposite film having manganese ferrite (MnFe₂O₄) nanoparticles therein as disclosed herein is provided. At 1704, an external stimulus is 20 applied to the nanocomposite film. At 1706, a shape of the nanocomposite film is deformed, and at 1708 an object is actuated, whichis caused by the deformation of the nanocomposite film. At 1710, the external stimulus is removed and at 1712, the initial shape of the nanocomposite film is recovered. Finally, if necessary, at 1714, the deformation of the nanocomposite film is reproduced upon application of the external stimulus.

To summarize, surface modified $\text{MnFe}_2\text{O}_4/\text{TPU}$ nanocomposite films with nanoparticle loading between 0.1 and 8 wt% were prepared by solution mixing followed by solvent casting. All of the films exhibited superparamagnetic behavior and the saturation magnetization increased with increasing nanoparticle content. Nanocomposite films were transparent or semi-transparent when the surface modified $MnFe₂O₄$ 35 nanoparticle loading was less than 2 wt %. Films with nanoparticle loadings of 4 wt % and higher were opaque. Large displacements (>10 mm) of all magnetic nanocomposite films were observed when a static magnetic field was applied. The maximum displacement increased with increasing mag-4o netic nanoparticle content. The proposed empirical correlation between the maximum displacement, saturation magnetization, and magnetic nanoparticle loading suggests a linear dependence of the maximum displacement to the saturation magnetization and a correlation to the nanoparticle weight 45 percentage. TEM and SEM micrographs show variable dispersion ranging from small nanometer-sized clusters to more abundant micron-sized aggregates.

> What has been described above includes examples of the innovation. It is, of course, not possible to describe every conceivable combination of components or methodologies for purposes of describing the subject innovation, but one of ordinary skill in the art may recognize that many further combinations and permutations of the innovation are possible. Accordingly, the innovation is intended to embrace all such alterations, modifications and variations that fall within the spirit and scope of the appended claims. Furthermore, to the extent that the term "includes" is used in either the detailed description or the claims, such term is intended to be inclusive in a manner similar to the term "comprising" as "comprising" is interpreted when employed as a transitional word in a claim.

What is claimed is:

1. A method of producing superparamagnetic $MnFe₂O₄$ 65 nanoparticles comprising:

mixing iron (III) acetylactonate, manganese acetyl acetonate, dodecanoic acid, 1, 2 dodecanediol, dodecylamine, benzyl ether under a nitrogen blanket for a first amount of time;

increasing the mixture temperature to approximately 150° C. for a second amount of time;

increasing the mixture temperature to approximately 300° C. for a third amount of time;

precipitating the nanoparticles in methanol; and

centrifuging and washing nanoparticles with excess methanol.

2. The method of claim 1, wherein the nanoparticles are used to create a nanocomposite film for use as an actuator that deforms upon exposure to a magnetic field.

3. The method of claim 2, wherein the first amount of time is approximately 15 minutes.

4. The method of claim 2, wherein the second amount of time is approximately 30 minutes and the third amount of time is approximately 30 minutes.

5. The method of claim 1 further comprising:

dispersing the mixture of nanoparticles in tetrahydrofuran (THF);

sonicating the mixture of nanoparticles for a fourth amount of time;

dissolving the dispersion in THF;

mixing the dispersions with a surface-modified manganese ferrite suspension MnFe₂O₄;

generating an aggregate free dispersion;

sonicating the dispersions for a fifth amount of time; solvent casting the dispersion to generate a nanocomposite

film; drying the nanocomposite films for a sixth amount of time.

5 6. The method of claim 5, wherein the fourth amount of time is approximately 5 minutes.

7. The method of claim 5, wherein the fifth amount of time is approximately 30 minutes.

8. The method of claim 5, wherein the nanocomposite films io s approximately 0.1-8 wt % superparamagnetic nanoparticles.

9. The method of claim 5, wherein the nanocomposite film is approximately 75-100 microns thick.

 10 . The method of claim 1, further comprising redispersing 15 and reprecipitating the superparamagnetic MnFe₂O₄ nanoparticles and recentrifuging and rewashing the superparamagnetic $MnFe₂O₄$ nanoparticles to purify the nanoparticles and remove excess reactants.

11. The method of claim 10, wherein the produced super-20 paramagnetic MnFe₂O₄ nanoparticles are further mixed with a polymer to produce a magnetic polymer nanocomposite.

12. The method of claim 11, wherein the polymer is a thermoplastic polyurethane (TPU).

13. The method of claim 11, wherein the polymer is a 25 polyimide resin.