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Buckypaper made with carbon nanotubes derived from CO₂†

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Unusual buckypapers, sheets of graphene nanocarbons (GNCs) such as carbon nanotubes, were formed with GNCs directly derived from CO₂ via molten carbonate electrolysis. Examples are presented for buckypapers made from CO₂ using either crushed or chemically washed GNCs, epoxy-infused CNTs, or GNCs pressed directly using a hot electrolyte to remove excess electrolyte.

Introduction

Buckypapers are graphene carbon nanocarbon (GNC) sheets. The majority of studies report on buckypapers are formed from the GNC consisting of carbon nanotubes. Additionally, buckypapers of graphene, graphene oxide, and carbon nano-onions have been studied. Buckypapers have demonstrated enhanced physical and chemical properties, including but not limited to high tensile strength; high electrical conductivity; high thermal conductivity; electronic shielding; magnetic shielding; electrical charge storage for use in batteries, fuel cells, and capacitors; piezo resistivity; catalytic activity; reduced friction; and targeted therapeutic activity.^{1–15} Many of these properties originate from planar sp² bonded carbons of graphene in different geometric arrangements within a buckypaper GNC sheet. Sheet properties can be modified through surface or covalent functionalization, morphology modulation, and doping.^{16–21}

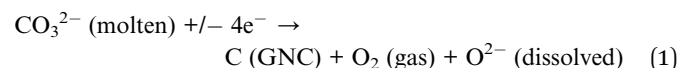
Buckypapers are often integrated within a composite to enhance the physical, chemical, or electrical properties of the buckypaper-free material.^{22–26} Buckypapers are attractive as they are lower in weight, stronger, more conductive, and more resilient for applications such as strain, fracture, chemical and motion sensors,^{4,37,38} electrochemical reactors³⁷ for water treatment;^{4,37,38} EMF shielding/absorbent coatings;^{7–9} batteries and supercapacitors;^{15,17,18,23,24} stronger plastic composites such as aeronautics^{11,12,25} and medicine.^{39,40}

Despite the extensive academic interest and wide range of applications, the acceptance and widespread use of buckypaper has been hampered by the high cost and carbon-footprint of graphene carbon nanocarbon synthesis. The manufacturing process generally includes a chemical vapor deposition (CVD) process used in the commercial production of carbon

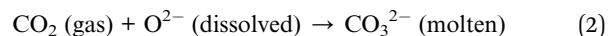
nanotubes, graphene, and carbon nano-onions. GNCs have shown promise, but their CVD production cost has remained prohibitively high in commercial settings, with the resulting products carrying both a significant cost and a significant carbon footprint.⁴¹ Currently, the price of GNCs such as CNTs, graphene, and carbon nano-onions are in the range of USD \$100,000–\$10 million per tonne. Comparatively, steel is priced at \$400 to \$700 per tonne.

To form buckypapers, CVD-formed GNCs are often first added to a liquid, then sonicated to provide a homogenous dispersion, and the liquid is filtered and/or dried off, leaving the buckypaper formed as a solid sheet of dispersed GNCs.

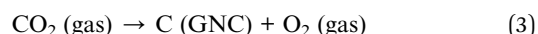
The years 2009 and 2010 marked significant milestones in the exploration of splitting CO₂ into carbon (C) and oxygen (O₂) through molten carbonate electrolysis, offering a decarbonization path to combat climate change. Building upon this progress, research in 2015 revealed that transition metal nucleus growth during this electrolysis process facilitates the direct conversion of CO₂ into pure CNTs and other GNCs.^{42,60}



CO₂ undergoes a chemical reaction with electrolytic oxide, as depicted in eqn (1), to regenerate CO₃²⁻ according to eqn (2).



The integration of eqn (1) and (2) results in the following net reaction:



Various GNCs have been synthesized, including helical, thin-walled, magnetic, and doped CNTs with carbon nano-bamboo, nano-pearl, and nano-tree morphologies as well as single-layered or multilayered graphene (nano-platelets), hollow or concentric buckyball spheres (nano-onions), and three-dimensional structures such as graphene nano-scaffolds.^{44–55} Manipulating the electrolysis parameters, such as temperature

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and current density, allows for the tailored production of specific GNCs. For instance, lower temperatures of about 725 °C are typical for forming carbon nano-onions,⁴⁴ while a higher range from 750 to 770 °C is utilized for CNT syntheses.^{45–47,49,52–58}

This communication presents a new low-cost process for the synthesis of graphene nanocarbons sheets (GNC buckypaper). CO₂ is the sole chemical reactant in eqn (3), and 4 tonnes of CO₂ are consumed for each tonne of C_(GNC) formed. Carbanogels are GNC lattices formed by carbon capture directly through the electrolytic splitting of carbon dioxide (CO₂). The electrolysis occurs in molten carbonate. The reduced CO₂ product builds up as a carbanogel containing the GNC and excess electrolyte at the cathode. The carbanogel product is recovered hot (as a red-hot slush) or cold (as a solid) by scraping and/or pressing from the cathode. Pressing returns excess electrolyte to the electrolysis chamber. Residual electrolyte and impurities may be removed with thermal, mechanical, or electrochemical treatment. The carbanogel is crushed and compressed within a mold to form the buckypaper product.

Herein, buckypaper was composed of GNCs derived from CO₂. This buckypaper presents an opportunity to utilize the greenhouse gas CO₂ for producing stable GNCs, contributing as a reliable decarbonization methodology for the long-term removal of CO₂. Graphite, a macroscopic form of layered graphene, serves as a geological stability benchmark with a life-span spanning hundreds of millions of years, providing a stability reference for GNC materials.

Results and discussion

Design of the experiment and carbanogel formation

Lithium carbonate was purchased at a battery grade >99.5% and was used as received. As analyzed, the lithium carbonate had a composition of 99.8% (Li₂CO₃, Shanghai Seasongreen Chemical Co.). Muntz brass is a high-zinc brass alloy composed of 60% copper and 40% zinc; this material is also referred to as 280 brass. This material serves as the cathode and was purchased from <https://onlinemetals.com/> and in larger quantities from Marmetal Industries. Electrolysis was conducted in 304 stainless steel “carbon pots”. The pot acts as both the cell case and its inner walls serve as the anode.

In a 770 °C molten Li₂CO₃ environment, CO₂ underwent splitting according to eqn (1)–(3), utilizing a Muntz brass cathode and a 304 stainless steel anode at a constant electrolysis current density J of 0.4 A cm⁻². Throughout the electrolysis process, CO₂ was divided into O₂ and GNCs. The GNCs developed as a network of interconnected graphene nanocarbons and electrolyte on the cathode. An expanded description of this electrolysis procedure, including product separation from excess electrolyte and product washing, has been recently delineated.⁵⁶ The mixture of GNCs and carbonate electrolyte is referred to as a carbanogel and further refined through the electrolyte's separation process, as detailed in the ESI.†,^{56,57} Thermogravimetric analysis (TGA) of the product, conducted using a PerkinElmer STA 6000 TGA/DSC, revealed a purity exceeding 97% (with less than 3% residual oxidation impurities at 800 °C) and exhibited an inflection temperature of 608 °C,

indicating remarkable resistance to oxidative combustion, characteristic of graphene-like materials. Scanning electron microscopy was studied using a PHENOM Pro-X scanning electron microscope to examine the products of the electrolysis process at various magnifications.

Fig. 1 presents a carbanogel with reduced electrolyte content achieved through a concentrated HCl wash. In a later example, the carbanogel was instead directly pressed to remove residual electrolyte. In Fig. 1, the 770 °C molten Li₂CO₃ electrolyzed sample was observed at two different SEM magnifications (x720 and x8600). This specific example illustrates the CNT carbanogel synthesized through CO₂ electrolysis, synthesized in 770 °C molten Li₂CO₃ detached from the cooled cathode, fragment, acid washed, and utilized in the first several buckypaper GNCs. Due to its lower combustion temperature compared to GNCs, amorphous carbon is more susceptible to oxidation, and both

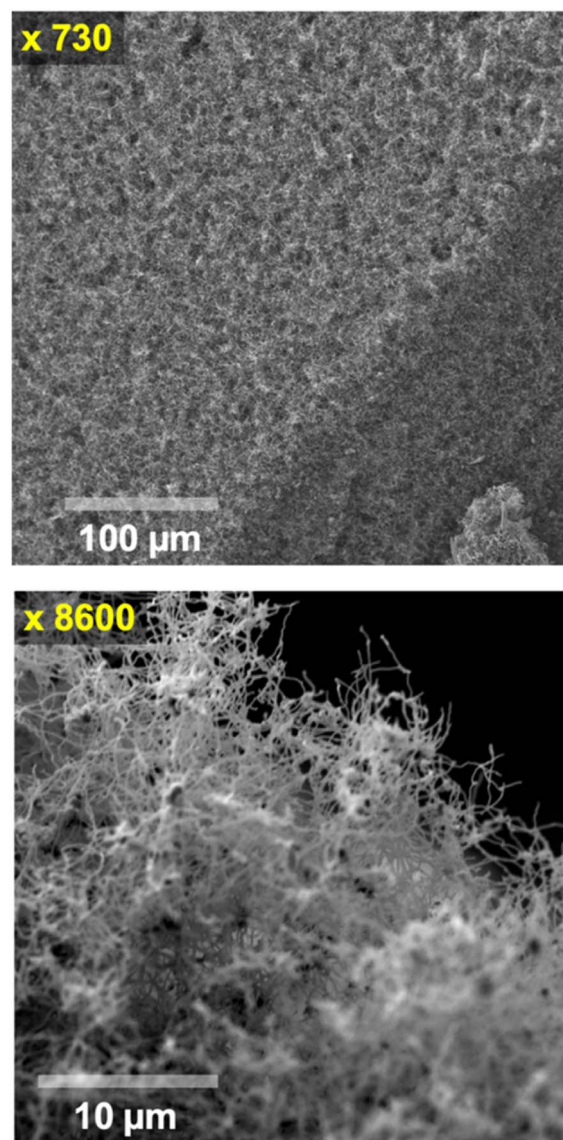


Fig. 1 Medium (top) and high (bottom) scanning electron microscopy images of the carbanogel particles comprised of carbon nanotubes formed at 770 °C Li₂CO₃ at $J = 0.4$ A cm⁻².



amorphous carbon, residual electrolyte, and metal impurities can be removed by oxidation and/or washing, as confirmed by electron dispersive spectroscopy (EDS) and TGA. Useful alternative post-electrolysis product washes include formic acid, copious water, or ammonium sulfate, although the latter two primarily remove excess electrolyte without affecting the amorphous carbon or metal impurities. Another alternative wash, combining hydrochloric acid and hydrogen peroxide, eliminates excess electrolyte, metal impurities, and amorphous carbon impurities, particularly coupled with sonication during the product wash.

The top SEM image in Fig. 1 presents a large collective size of the intertwined CNTs forming the carbanogel. One advantage of grouping nanoparticles within an intertwined macroscopic matrix is the mitigation of respiratory hazards typically associated with nanoscale particles during transportation.

Higher-resolution SEM image (Fig. 1 bottom) reveals that the GNC product comprises highly pure carbon nanotubes. The SEM shows the diverse lattice intertwined orientation of the carbanogel product. This product is utilized in subsequent buckypaper formation steps. Additionally, this structure offers electrical and thermal conductivity pathways, along with a highly porous framework suitable for accommodating polymers, catalysts, or battery intercalation. We have recently reported extensive SEM, TEM, X-ray, Raman, high angle annular dark-field (HAADF) elemental analysis/TEM of the CNT and various GNC products, as detailed in the ESI.†^{51,58}

For the electrolytic CO₂ splitting and transformation electrolyses of this study, the kilns utilize a direct (untreated) feed of 5% CO₂ from the emissions of the adjacent 860 MW (Shepard, Calgary Canada) natural gas electric power plant or direct air. The large kiln modules designed and utilized are pictured in Fig. 2. These kilns simultaneously sustain electrolysis in several carbon pots. Cathodes show an increase in the surface area to over 10 000 cm². In each case, the cathode is mounted vertically in the electrolyte, across from which are the 304 stainless steel anodes; the anodes simultaneously function as the electrolysis

anodes and as the chamber walls of the 304 stainless steel “carbon pot”. The electrolyte has a strong affinity for CO₂ from the open air. The kilns shown in Fig. 2 can also be configured for direct air use. Finally, an amine concentrator is also in place at the site that can concentrate the 5% CO₂ flue gas. From the amine concentrator as an alternative source, 98% CO₂ (containing 2% H₂O) was used as the kiln/carbon pot input, producing equivalent GNCs.

CO₂ buckypaper (BP) with CNTs

Fig. 3 presents a typical example of a GNC buckypaper produced from CO₂. In this initial example, the GNC buckypaper was generated through electrolysis to convert CO₂ into carbanogel. The carbanogel product was cooled, removed from the electrode, ground, washed, dispersed, and molded to CNT buckypaper. Specifically, the process involved utilizing a 304 stainless steel case within a 770 °C Li₂CO₃ molten electrolyte, employing a Muntz brass cathode and a 304 stainless steel anode to yield the CNT carbanogel product (Fig. 1). Subsequently, this carbanogel product underwent cleaning with hydrochloric acid (HCl) and 0.2 grams of the washed product was mixed in 300 mL of isopropyl alcohol, followed by sonication for 30 minutes to ensure even dispersion.

The dispersed isopropanol CNT mixture was then transferred into a vacuum filter assembly (Whatman nylon membrane filter; 0.2 μm Pore, 47 mm diameter), pressed, the liquid was extracted under vacuum, dried, pressed, and detached by lifting from the membrane filter. The GNC buckypaper had a thickness of 96 μm, and a measured conductivity of 11 000 S m⁻¹ ($\sigma = \text{length}/(R, \text{resistance}, x \text{ cross section})$). This conductivity is consistent with our measurements of carbonate-synthesized CNTs and may be increased an order of magnitude by boron doping (addition of a boron salt during the electrolysis).^{52,53}

Composite, larger, and improved dispersion BP from CO₂

Buckypaper strengtheners can include, but are not limited to epoxies, resins and other polymers, cementitious materials, and



Fig. 2 The Genesis Device® kiln modules facilitate large-scale CO₂ molten carbonate electrolysis and was used in this study. Carbon Corp operates these modules in Calgary, Canada, for onsite decarbonization.



Fig. 3 A 4.7 cm diameter carbon nanotube buckypaper prepared via the electrolysis of CO₂ to form a carbanogel, grinding of the product, dispersion in isopropanol, pressing and filtering, and forming on a 0.2 μm pore nylon membrane.



metals. Catalysts to expedite chemical or electrochemical reactions can also be included. Dopants to affect the physico-chemical properties of the BP can be added. Effective GNC dopants can enhance the conductivity, catalytic activity, and battery storage capacity properties. Dopants include boron, nitrogen, sulfur, and phosphorous.^{52–54} A magnetic material is one or more of those with ferromagnetic properties, paramagnetic properties, diamagnetic properties, and any combination thereof.^{4,46}

Jetset-Metlab epoxy is a rapidly cured BPA epoxy. Its resin comprises 80–90 wt% propane, 2,2-bis[*p*-(2,3-epoxypropoxy)phenol] polymers, and 10–20 wt% alkyl (C12–14) glycidyl ether. The hardener consists of 60–70 wt% diethylenetriamine, 30–40 wt% bisphenol A, less than 0.8% aminoethylpiperazine, and less than 0.2% ethylenediamine. In a manner similar to the buckypaper shown in Fig. 3, buckypaper (BP) was formed from 0.25 g of the carbanogel. However, the buckypaper was not pressed, resulting in a considerably thicker (1133 μm) BP with an order of magnitude lower density (0.2 g cm^{-3}). 3.6 g of the Jetset-Metlab epoxy was mixed with 0.38 g of the Jetset-Metlab hardener. 0.38 g of the epoxy mix was spread on the BP. A silicone sheet was placed over the epoxy-BP, compressed with a heavy (0.5" thick) stainless steel plate, and cured overnight. The silicone sheet was readily removed and the epoxy was infused into the BP and hardened. The composite, again formed with CNT from CO_2 , was readily handled and highly resistant to fracturing or breakage. In an upcoming paper, we will present that compared to samples without CNTs, even low levels of added CNTs (1.5 to 2 wt% CNT) increase the tensile strength of several epoxies. Specifically, Timber Cast, Varathane and Jetset-Metlab epoxies' tensile strengths were increased by 40 to 60% in ASTM D638 "dogbone" type V mold samples tested with an ETM-10 kN Computer Controlled Electronic Universal Testing Production Machine (Shore D Durometer). The hardness was also increased. Equivalent tests on the BP resin composite, upon completion, will be reported in an expanded paper.

The buckypaper from CO_2 presented in Fig. 3 is scalable. However, thinner BPs, consisting of BP formation with a decrease in the grams of carbanogel per unit area of the membrane, were increasingly difficult to detach from the filter paper. Intermediate diameter BPs could be formed and removed. Specifically, when 0.3 g of ground carbanogel was washed in HCl, dispersed by sonication for 30 minutes in isopropanol, and vacuum filtered through a 270 mm Whatman 0.2 μm pore (grade 4) nylon filter, the BP was formed but could not be removed in one piece from the filter paper.

A modified dispersion method to facilitate BP detachment was sought. Polyvinylpyrrolidone (PVP) was utilized to promote the dispersion of CNTs.⁵⁹ 0.05 g of 10 000 molecular weight PVP (PVP10, from Sigma Aldrich) was added along with 0.10 g of CNTs, rather than 0.20 g, ground from the washed CNT, and dispersed in 300 mL of isopropanol with 15 minutes, rather than 30 minutes, of sonication. The CNTs were prepared in a manner similar to those presented in Fig. 1, also in Li_2CO_3 , but at a lower temperature of 750 $^\circ\text{C}$ and a lower current density of 0.09 A cm^{-2} . Further, the carbanogels were more extensively

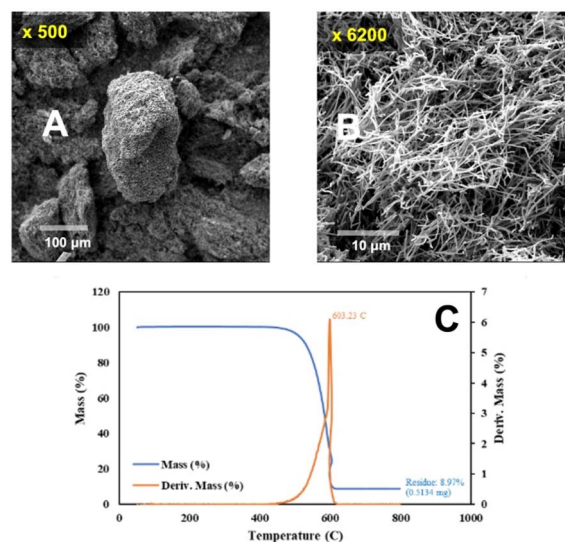


Fig. 4 (A) Medium and (B) higher magnification scanning electron microscopy images of the carbanogel particles composed of carbon nanotubes. (C) TGA of the carbanogel product, formed in 750 $^\circ\text{C}$ Li_2CO_3 at $J = 0.09 \text{ A cm}^{-2}$.

ground using a Magic Bullet Blender. The CNT's SEM and TGA are presented in Fig. 4.

Once again, in Fig. 4, the carbanogel particles are evident in the lower magnification SEM (panel A), and the TGA temperature of inflection of 603 $^\circ\text{C}$ is high, although the oxidized 9% residual (panel D) is higher than that in the previous sample. Fig. 5 presents a typical example of a GNC buckypaper produced with the PVP isopropanol-dispersed CNT from CO_2 . The unpressed BP is considerably thinner at 58 μm than the unpressed sample prepared without the PVP dispersant.

Direct press CO_2 buckypaper

The electrolyte has a strong affinity for CO_2 , directly from air ($p\text{CO}_2 = 420 \text{ ppm}$ and climbing), or directly sourced from an industrial stack. Industrial CO_2 feedstocks tested vary from $p\text{CO}_2 = 50\,000 \text{ ppm}$ (from the Shepard Energy Centre 860 MW natural gas power plant, Calgary, Canada) to $p\text{CO}_2 = 980\,000 \text{ ppm}$ (from a liquid amine concentrator onsite at Alberta Carbon Capture Technology Centre, ACCTC, at the plant). The electrolysis provides a continuous source of oxide (eqn (1)), reacting with CO_2 (eqn (2)). The capture and transformation of CO_2 was determined by ^{13}C isotopic labeling,⁶⁰ and the real-time simultaneous measurement of CO_2 and the eqn (1) co-product O_2 was done as presented previously.⁵⁵ In the first examples, buckypapers were formed with carbanogels by the transformation of CO_2 in the air. In this next example, the BP was formed from 50 000 ppm CO_2 fed from the Shepard Energy Centre natural gas power plant exhaust.

Fig. 6 presents an alternate methodology for the preparation of a GNC buckypaper derived from CO_2 . This GNC buckypaper was produced from CO_2 transformed into carbanogel, following the process outlined in the previous example. However, instead of chemical washing, the electrolyte content of the carbanogel





Fig. 5 A 4.7 cm diameter carbon nanotube buckypaper prepared via the electrolysis of CO₂ to form a carbanogel, grinding of 0.10 g of the product, dispersion with PVP in isopropanol, pressing, filtering, forming on the 0.2 μm pore nylon membrane, and lifting from the membrane.

was reduced through direct compression while the carbanogel was hot (containing the GNC product and residual, excess electrolyte). Specifically, the carbanogel, containing the solid GNC and molten electrolyte, was generated at the cathode and then compressed at 500 psi through layers of 60 mesh 304 alloy stainless steel mesh while hot, as recently delineated.⁵⁶ Notably, the 250 μm mesh pore size is significantly larger than that of GNC but smaller than that of the carbanogel particle size. The small area of the lighter brown discoloration in Fig. 6 is where the GNC buckypaper adhered to the mesh screen during separation.

During compression, the larger size of the intertwined CNTs in the carbanogel is retained above the mesh, while the electrolyte passes through. The resulting GNC buckypaper's thickness is linearly proportional to the starting mass of the carbanogel and approximately inversely proportional to the applied pressure.



Fig. 6 A 35 cm diameter carbon nanotube buckypaper prepared via the electrolysis of CO₂ to form a carbanogel, and pressing of the carbanogel product while molten to separate residual carbanogel electrolyte from the graphene nanocarbon, carbon nanotube, and product.

The GNC buckypaper shown in Fig. 6 has a diameter of 350 mm, and larger versions, approximately two-fold in size, were produced. A 3700 psi example is seen in our recent carbanogel electrolyte extraction study.⁵⁶ Unlike other buckypaper formation protocols, which call for a prerequisite dispersion step such as sonication, to provide a homogeneous distribution of the GNC components, the GNC components are already homogeneously distributed in the unground carbanogel. Hence, the BP formation process does not need a prerequisite GNC dispersion.

CO₂ BP generalization and future potential

A broad array of filter pore sizes and solvents are found to be effective for forming buckypaper from CO₂ transformed into the carbanogel using these BP formation methodologies. Varying the CO₂ electrolysis parameters, including temperature, current density, and cathode, anode, and electrolyte composition, generates alternative GNCs for buckypaper (such as CNOs,⁴⁴ to be described for buckypaper in a subsequent study).

Forces to align, rather than disperse, the CNMs can additionally be applied during the carbanogel buckypaper formation and or liquids added before the pressing to maintain more even layering of the carbanogel particles. The alignment can provide directional, anisotropic properties to the carbanogel buckypaper and provide enhanced carbanogel buckypaper properties, including but not limited to enhanced strength, conductivity, and directional interactions with visible and other electromagnetic radiation. The applied alignment forces can be linear, radial, cylindrical, or spherical to produce directional geometries of anisotropy.^{23,60–64}

The GNC components in the BP can be aligned mechanically, electrically, or magnetically during the BP formation or a combination thereof to further enhance the BP properties, including enhancements of the BP strength and/or electrical and thermal conductivities and EMF absorption, and will be presented in an expanded article. The mechanical alignment can be achieved with the application of shear force, such as by pulling or spinning during the BP preparation steps or dragging a piston applying the formation pressure. Alternatively, the shear force can be directionally applied to increase rather than align the GNC entanglement. The electrical and/or magnet alignment is achieved with the application of an orienting electrical or magnetic field during the BP preparation or processing stages. Magnetic GNCs are prepared to incorporate magnetic materials, such as metals or metal carbides, during the GNCs preparation stage. The decrease in the distance of the magnetic field is greater than that for the electric field, and the magnetic alignment is more offset than the electric alignment effect by the competing random disorder of Brownian motion, which increases with temperature and modes of freedom and decreases with increasing molecular mass and viscosity. Hence, the alignment is enhanced by the decrease in temperature or molecular mass (as exemplified by multiwalled *versus* single-walled carbon nanotubes) and increases with viscosity.

The carbanogel buckypaper may be used alone, such as in liners, heat retardants, or shields, or in combination, such as laminates, with other materials to impart improved properties to other materials. CNT materials have displayed a shape-



memory property (spontaneously returning to their original shape) under thermal, mechanical, electrical, or magnetic activation conditions which can be incorporated into BP.^{65–74} This shape-memory effect is promoted by anisotropic properties in the buckypaper. The electrical and thermal conductivity of buckypaper present superior properties for their applications as heating elements. Both shape-memory and heating element behavior have been studied alone and in combination with other material layers.⁶⁸

Incentivized carbon mitigation

Innovative approaches, such as employing high-solubility molten pathways and harnessing renewable energy for electrolysis, have shown promise in reducing energy consumption and production costs. In the formation of graphene nanocarbon composites (GNCs) *via* CO₂ to carbon nanoallotrope technology (C2CNT®), the process solely requires CO₂ as a reactant. The energy required for electrolysis to convert CO₂ into GNCs falls within the range from 0.8 to 2 volts.⁶⁰ When produced in bulk, they are approximately \$1000 per tonne.⁵³ These are akin to the costs of the industrial aluminum production electrolytic process, which involves splitting aluminum oxide to yield commercial-grade aluminum metal.⁵³ C2CNT® costs will be further reduced as solar and wind energy are increasingly used as alternative power sources.^{75–84}

An estimate of cost and decarbonization benefits is exemplified in the context of one application. Typical ratios of plastic resin to fiber infusion are in the range of 60 : 40. The price index for plastic materials and resins has been \$300–\$400 per tonne over the last three years.⁸⁵ Hence, the infusion of resin in buckypaper may decrease the material costs to the order of \$500 per ton buckypaper resin composite while significantly lowering the quantities of the material to achieve the properties needed for various applications. The buckypaper-resin composite provides the capability of enhancing the strength, tearing resistance, and fracture toughness several folds while increasing the electrical and thermal conductivity by several orders of magnitude. Reducing the amount of plastic needed to achieve the desired properties through buckypaper resins made with GNCs from CO₂ provides the incentive of both cost and carbon footprint reduction.

Conclusions

Buckypapers with GNCs, such as carbon nanotubes made from CO₂, are demonstrated to form a range of conductive, liquid-dispersed and unusual direct molten pressed forms of buckypapers, all prepared from lattices of intertwined carbon nanotube carbanogels synthesized by the electrolytic splitting of CO₂ in molten carbonates. As with industrial aluminum production by the electrolytic splitting of aluminum oxide, the production of GNCs by the electrolytic splitting of carbon dioxide is inexpensive, providing a cost-effective path for the synthesis of buckypapers. The only reactant used in the production of the GNCs is CO₂ (3.7 tonnes of CO₂ are consumed for each tonne of C_(GNC) formed), providing a decarbonization path to mitigate

the greenhouse gas CO₂. The captured carbon, transformed from CO₂, provides a storage buffer to remove CO₂. Furthermore, the removed CO₂ in the form of GNCs is highly stable, with the potential to permanently sequester CO₂ from the carbon cycle. It is anticipated that with the further alignment and shape-memory effects that can be achieved with buckypapers, a range of useful lightweight, high strength, and high conductivity applications can be demonstrated with these new materials.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

There are no conflicts to declare.

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