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Electrospinning: designed architectures for energy conversion and storage devices

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Electrospinning is attracting close interest as a versatile fabrication method for one dimensional mesostructured organic, inorganic and hybrid nanomaterials of controlled dimensions prepared as randomly oriented or oriented continuous nanofibres that can present internal compositional organisation such as core-sheath, hollow or porous fibre, or even multichannel microtube arrangements. The dimensionality, directionality and compositional flexibility of electrospun nanofibres and mats are increasing being investigated for the targeted development of electrode and electrolyte materials. Specific properties associated with the nano-scale features such high surface to volume and aspect ratios, low density and high pore volume allow performance improvements in energy conversion and storage devices. We review here the application of electrospinning for designing architectured nanofibre materials for dye sensitised solar cells, fuel cells, lithium ion batteries and supercapacitors, with particular emphasis on improved energy and power density imparted by performance improvement to, inter alia, ionic conductivity, cyclability, reversibility, interfacial resistance and electrochemical stability, as well as mechanical strength, of electrospun electrode and electrolyte components.

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1. Introduction

Energy conversion and energy storage are ever more important in the context of escalating energy requirements and increasing world population. Future generations of efficient energy conversion and storage devices are required to meet this challenge, whereby

the conversion and storage of energy from renewable sources is bound to play an increasingly relevant role. Rechargeable, reversible and highly durable conversion and storage systems are required, consistent with a responsible utilisation of global reserves of strategic elements, and avoiding their rarefaction. The current challenges associated with the development of energy conversion and storage devices are well-documented in review articles dedicated to electrode and electrolyte developments for lithium ion batteries^{1, 2, 3} supercapacitors ^{4, 5, 6, 7, 8}, proton exchange membrane fuel cells^{9, 10, 11, 12, 13} and dye sensitised solar cells ^{14, 15, 16, 17}. To achieve high energy and power density and system efficiency, designed functional materials are needed that associate the required electrochemical properties for improved performance, with chemical and electrochemical stability, while leaving a minimally low environmental footprint. Clearly the selection of appropriate electrolyte and electrode structure and architecture is fundamental in determining device performance.

Materials with nanometer-scale dimensions have attracted a great deal of attention over the last 20 years or so. Nanomaterials and nanostructured materials, including nano-coatings and films and nanocomposites, are at the epicentre of energy materials since this corresponds to the length scale over which many of the elementary steps such as

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charge transfer or molecular rearrangement occur, and because their reduced dimensions and the effect of surface properties on their behaviour provide a high volume fraction of interfaces and favour increased reaction rates. Exploring materials chemistry down to the nanoscale to develop novel solutions for energy applications is possible by a systematic approach involving new methods for manipulation, assembly and characterisation.

Targeted development of new one-dimensional (1D) nanostructures, such as continuous nanofibres (NFs), large aspect ratio nanowires (NWs), and nanorods (NRs)^{18, 19, 20, 21}, is attracting much interest because of the dependence of their physical properties on directionality. Conventionally these materials are produced by the bottom-up approach, i.e. by selfassembly of the target material from vapour, liquid or solid phases through nucleation and growth under optimised conditions (hydrothermal synthesis,...)²². Most 1D nanomaterial synthetic methods involve multiple steps: for instance, the fabrication/removal of templates and the catalyst elaboration. The different process conditions in each step often limit the practical device quality. Among the methods for generating fibrous nanostructure, electrospinning is a top-down simple, versatile and cost-effective approach allowing the fabrication of nanofibres in a continuous process and at long-length scales.

2. Principle and applications of electrospinning

Electrospinning is a process leading to the formation of ultra-fine fibres with diameters in the micrometre to nanometre range, starting from molten polymers or polymers in solution. Some excellent reviews describing the technique are available^{23 24, 25,} ^{26, 27, 28, 29, 30, 31, 32, 33}. Electrospinning is based on the application of an electric field to a drop of fluid polymer on the tip of a spinneret. As the intensity of the electric field increases, the hemispherical surface of the solution at the tip of the capillary tube elongates to form a conical shape known as the Taylor cone (see Fig. 1). When the applied electric field reaches a critical value, the repulsive electrical forces overcome the surface tension of the drop. A charged jet of the solution is then ejected from the tip of the Taylor cone and an unstable and rapid whipping of the jet occurs between the tip and the collector leading to the evaporation of the solvent and the formation of solidified continuous, ultra-thin fibres on the collector³⁴.

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The conventional electrospinning set-up consists of three major components: a high voltage (kV) power supply, a spinneret (a syringe or pipette tip) and a grounded collector (typically a metal plate or a rotating mandrel) (see Fig. 1).

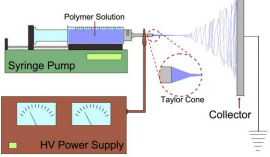


Fig. 1 : Scheme of a conventional electrospinning set-up.

Electrospun nanofibres present specific properties, such as high surface/volume and aspect ratios, leading to low density and high pore volume, and exceptional mechanical strength²³. Electrospinning, first studied by Zeleny³⁵ in 1914 and patented by Formhals³⁶ in 1934, was initially exclusively applied to polymers³⁰. During the last two decades, electrospinning has attracted renewed attention due to the huge potential that the technique presents for nanomaterials and nanotechnologies³⁷. Indeed, the new properties appearing at the nanoscale (quantum effect, superparamagnetism, electronic effects)^{38, 39} have led to intense research on nanomaterials⁴⁰. The applicability of electrospinning has been then extended to the production of hybrids (e.g. polymers loaded with chromophores, magnetic nanoparticles, biomolecules) $^{41,\ 42,\ 43,\ 44}$ and inorganic 31 32 onedimensional nanomaterials, using a range of precursor types. Electrospinning has the advantages of simplicity, efficiency, low cost, high yield and high degree of reproducibility of the obtained materials. The versatility of electrospinning is not only related to the nature of the precursors and electrospun materials, but also extends to the fibre assemblies and architectures. Modifying solution and processing parameters^{45, 46} and/or set-up geometries³³ allows the preparation of complex nanostructures with controllable hierarchical features such as nonwoven, aligned or patterned fibres, nanoribbons, nanorods, random three dimensional structures, sub-micron spring and convoluted fibres with controlled diameters^{47, 48}. The elaboration of such designed assemblies in situ (and not by a subsequent physical manipulation) allows their customisation to adapt them to specific applications. Furthermore, the possibility of co-electrospinning different solutions

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using specially designed spinnerets leads to the elaboration of core-sheath⁴⁹, hollow fibres^{50, 51, 52}, porous fibres⁵³, and multichannel microtubes^{54, 55}. The properties of the obtained designed fibres can also be tuned by the addition in situ, or by further of molecular functionalisation, species or nanoparticles.⁵⁶. It is important to note that although electrospun nanofibres are almost one-dimensional, their mats are generally disordered, unless special measures are taken, and effective techniques for producing aligned nanofibres of sufficient fibre length and at large scale are required. Bridging and alignment of electrospun fibres across slotted electrodes has been described^{33, 57} although the process is complex and the distance between the electrodes is limited. A needleimprinting method, using a movable needle system in combination with a thin conveyor belt comprised of an insulation material that slides on top of the grounded collector plate, also allows fibre alignment, and here the continuous process allows larger scale production of electrospun mats58. Continuous aligned polymer fibres have also been produced by simultaneous ejection from two needles with opposite voltages, and collecting the neutral yarn on a rotating shaft⁵⁹ while continuous aligned ceramic fibres have been produced from a multi-jet/multi-collector system whereby the the spatial orientation of the nanofibres is controlled by the inducing effect of the converging electric field generated by thin metal wire collectors⁶⁰.

This tremendous variety of electrospun nanofibres in term of materials, structures, architectures and functionalities explains the increased use of the electrospinning technique for the production of nanofibres and fabrics⁶¹ and membranes^{62, 63, 64} elaborated from them. The extensive applications⁶⁵ range from catalysis, structural applications requiring high mechanical strength, biology for tissue engineering, biomolecular machines, biosensors, nanoelectronics, optical, resonator, and opto- and magnetoelectronic devices, sensors, photocatalysis, biomedical fuel cells, solar cells, membranes (see Fig. 2)^{66, 67, 68, 69, 70}.

Due to its inherent adaptability and applicability, and as indicated recently^{71, 72}, the electrospinning technique can be applied to the preparation of different components of energy materials, from polymer electrolytes to ceramic and metallic electrodes. The aim of this review is then to give a detailed overview of recent advances in the application of materials issued from electrospinning in energy devices including fuel cells (FCs), dyesensitised solar cells (DSCs), supercapacitors and Liion batteries (LIBs). The number of published research articles on this topic has significantly increased

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especially in recent years (see Fig. 3), showing the need for monitoring the present results and identifying the future challenges.

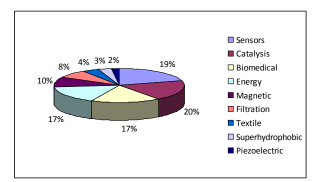


Fig. 2. Relative proportion of publications in main areas of applications of electrospinning in the period of 2000-2010 (Scifinder data: Keywords: electrospinning + type of application in the legend).

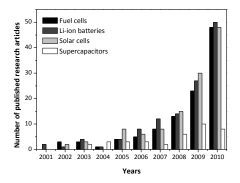


Fig. 3. Number of research articles concerning electrospun nanofibre applications in energy devices published in the period of 2001-2010 (Scifinder data: keywords electrospinning + fuel cells, electrospinning + Li-ion batteries, electrospinning + solar cells, electrospinning + supercapacitors).

3. Electrospun materials in energy conversion and storage

3.a. Electrospun nanofibres for fuel cells

Fuel cells convert the chemical energy of a fuel into electricity for power generation with high efficiency. Using pure hydrogen and air, fuel cells produce only water, thus eliminating any pollution or emission at the point of use. Hydrocarbon fuels can also be used and, although their use is associated with local carbon dioxide emissions, there is great interest in direct methanol fuel cells for portable electronics applications, and in (bio)ethanol as a renewable fuel source for fuel cells. While portable applications represent early fuel cell markets, the main drivers for development of fuel cells are stationary and transportation usage. Renewed interest over the past

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two decades triggered by the need to integrate alternative energy carriers and thus energy conversion systems to improve energy sustainability, intense research and development endeavours have been pursued to advance materials performance and robustness, to increase lifetime and reduce costs. These efforts are currently dedicated to all components of the membrane electrode assembly: electrolyte membrane^{73, 74, 75, 76} (increase conductivity and high temperature and low relative humidity, improve mechanical strength), electrocatalyst (reduce platinum loading, increase mass and specific activity), ^{77,78}and catalyst support material ^{79, 80} (stabilisation of carbon support to oxidation, alternative support materials). Elaboration of fuel cell nanomaterials of targeted architecture is increasingly considered as a powerful tool in reaching these goals, through the spatial organisation of catalytic metals to achieve efficient presentation of platinum to the reaction medium,^{81, 82, 83, 84} optimised microstructural arrangements of polymer membranes,^{85, 86} and designed nanoporosity within carbon^{87, 88}, oxide^{89, 90}, or carbide⁹¹ supports. The potential of electrospinning for fuel cell materials has been quickly recognised, and over a short period of time the literature in the field has burgeoned. Recent advances and new directions are described below and summarised in Tables 1-4.

3.a.1. Electrospun catalysts for fuel cells

Thus far there have been comparatively few studies on the use of electrospinning to synthesise electrocatalyts^{92, 93, 94, 95, 96, 97}, in contrast with its use to fabricate catalyst supports. This may be due to the difficulties in achieving an extremely fine fibre diameter (<10 nm) by electrospinning compared to other methods such as chemical vapour deposition. Nevertheless, the earliest publication was by Kim et al. regarding the synthesis of catalytic electrospun nanowires (NWs) of Pt/Rh and Pt/Ru NW with 50 nm diameter⁹⁴ (Fig. 4). The mass activity of PtRu NWs was better than the conventional catalyst of highly dispersed PtRu NPs on carbon. A NP/NW combination system was also studied, where Pt NPs on carbon were blended with electrospun Pt NWs. The mass activity of this combined electrocatalyst was higher per unit mass of Pt than using solely the conventional cathode catalysts of Pt on carbon. Shui et al.97 described the fabrication of centimetre-long nanowires of Pt by electrospinning, with the ultimate goal of removing the need for a catalyst support in the fuel cell electrode. These Pt NWs show a diameter of 5-17 nm, with the fibre formation (rather than other morphologies such as beads) dependant more on the

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type of carrier polymer (PVP) than the nature of the platinum precursor. In other recent work, Pt-Fe alloy nanowires prepared by electrospinning and dealloying were shown to present high specific activity for Another reduction⁹⁸. oxygen approach was demonstrated by Su et al.99 who fabricated polyamide-6 (PA6) nanofibres coated with a Pd layer of ~85 nm thickness for the electrooxidation of alcohol in aqueous medium. The MEA incorporating the Pd/PA6 nanofibres provided high current density and enhanced performance which was attributed to the large surface area, reduced diffusion resistance, and excellent tolerance to CO poisoning.

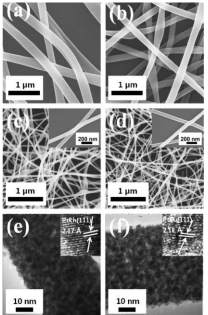


Fig. 4. FESEM images of as-spun (a) Pt-Rh precursor/PVP, (b) Pt-Ru precursor/PVP and bimetallic nanowires of (c) PtRh and (d) PtRu; insets of (c) and (d) show the diameter of bimetallic NWs. HRTEM images of (e) Pt1Rh1 NWs and (f) Pt1Ru1 NWs; insets of (e) and (f) are inverse fast fourier transform image of PtRh NWs and PtRu NWs. Reprinted from ref. 94 Copyright (2008), with permission from Elsevier.

3.a.2. Electrocatalysts on electrospun support materials

The support material for electrodes is a very crucial point in fuel cell development, impacting both performance and durability. The general requirements for electrocatalyst support materials are high electronic conductivity, high specific surface area and high electrochemical and chemical stability under fuel cell operating conditions. Interaction between catalytic metals and the support should also be considered in improving the performance. Such supports allow a high dispersion and a narrow size distribution of

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catalyst nanoparticles, favouring their maximum exploitation and rapid kinetics. They can also provide a way to decrease the metal loading, and then electrode costs that are one of the major bottlenecks of PEMFC (*e.g.* in the case of noble metals such Pt)¹⁰⁰. The durability of the catalyst is also greatly dependent on its support.

Carbon (e.g. commercial Vulcan XC-72) is the most common catalyst support for fuel cell applications due to its large surface area and high electronic conductivity. However, despite these advantages and relatively low cost, the use of carbon also presents one of the main challenges for long term fuel cell operation, namely carbon corrosion. Thus, significant research efforts have been directed to develop a carbon-based or alternative material which has improved corrosion resistance. Novel nanostructured carbon materials such as nanotubes¹⁰¹ and nanofibres^{80, 102} have received much attention as catalyst supports for PEM fuel cells. Indeed, the specific 1D morphology can improve the stability of the support materials with better utilisation of the electrocatalysts and the overall better performances and lifetime of the resulting electrodes. Recently, the electrospinning technique has been used as synthesis strategy for carbon materials with controlled diameters and structures.

Carbon nanofibres (CNFs) represents a promising support material for fuel cells as CNFs have increased conductivity over carbon particles. This is due to easier electron transfer along the aligned CNF, in contrast to carbon particles where there are significant interfaces between the particles, which may add to the resistance of the system. Most studies have focused on carbon fibres derived from polyacrylonitrile (PAN) as they can be readily electrospun and their graphitisation method is well established. Phenolic resin-based carbon fibres with thin diameters and high conductivity have also been prepared^{103, 104}. Park et al.¹⁰⁵ developed electrospun PAN-based carbon nanofibres which have high surface area, shallow pores and rough surfaces leading to significantly higher Pt utilisation (69 %) in the Pt/CNF than with Pt/Vulcan XC-72R (35 %). The electrical conductivity of the electrospun CNFs carbonised at 1000 °C was also higher at 9.9 S cm⁻¹, whereas the conductivity of Vulcan XC-72R was 4.5 S cm⁻¹, which led to improved fuel cell performance of the electrospun electrode. Furthermore, it was found that the electrospun nanofibres have higher conductivity along the fibres than across them, indicating good electron transfer within the fibres.

Nataraj *et al.*¹⁰⁶ aimed to fabricate CNFs with improved morphology and conductivity by

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incorporating various amounts of heteropolyacids (HPAs) into the PAN precursor. Increasing HPA content provided increasing electrical conductivity in fibres (however HPA-containing the overall conductivity of the HPA loaded CNFs was actually lower compared to pristine CNFs carbonised at 1000 °C) and finer fibre diameters in the CNF mats, resulting in greater surface area. Another study by the same authors¹⁰⁷ investigated the influence of nickel nitrate in the place of HPA, and a similar enhancement of surface area increasing nickel nitrate concentration were observed, as well as improved thermal stability. These effects were attributed to several factors, including the inorganic particles-polymer interaction and the increased conductivity of the electrospinning solution.

Electrospun fibres have also found application in alkaline fuel cells (AFCs). For instance, iron/cobalt/carbon electrospun nanofibre (FeCo/CNF) composite electrocatalysts demonstrated comparable electrocatalytic activity and stability to commercial carbon-supported platinum (Pt/C) for ORR, a direct 4-electron reduction pathway, and better ethanol tolerance than Pt/C in an alkaline electrolyte¹⁰⁸. This approach shows the possibility to prepare advanced cathode materials for AFCs without requiring noble metal catalysts.

A series of studies by Li et al.^{109, 110, 111} investigated Pt deposited through several methods on electrospun carbon nanofibrous mats (also from PAN precursor) for DMFC applications. The carbon nanofibres improve the catalytic peak current of methanol compared to commercial Pt/C^{110} (see Fig. 5), indicating that the 1D structure of the carbon mat is beneficial to the catalytic performance of the catalyst. Another approach for Pt deposition was reported by Lin et al.¹¹² through the pre-functionalisation of the CNFs with 1-aminopyrene in order to improve Pt attachment. After chemical reduction, they obtained PtRu nanoparticles with small diameter (3.5 nm) and homogeneous distribution. The resultant PtRu/CNFs had high electrochemical surface area and good activity toward methanol oxidation.

Other polymers have also been used to produce carbon nanofibres. Xuyen *et al.*¹¹³ have prepared polyimide (PI)-based carbon nanofibres which were synthesised by the use of an electrospun poly(amic acid) (PAA) nanofibre precursor. Such a PI nanofibre mat can then be subsequently modified via hydrolysis in order to generate surface sites favourable to anchoring of Pt nanoparticles. Here the immobilisation of the Pt precursor is controlled through the degree of hydrolysis, and the size and distribution of Pt nanoparticles by the subsequent carbonisation step.

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An alternative method for the fabrication of Pt/carbon and Pt-Pd/carbon composite nanofibres has been described by Lin *et al.*^{114, 115} where Pt and Pt-Pd nanoparticles were electrodeposited directly onto electrospun carbon nanofibres. In this method, the morphology and size of nanoparticles can be controlled via the deposition time. Huang et al.¹¹⁶ have also used electrodeposition to prepare CNF-supported bimetallic Pt_xAu_{100-x} electrocatalysts, which showed good electrocatalytic performance for potential use in direct formic acid fuel cells.

As mentioned above, carbon is the most widely used catalyst support because of its high surface area and excellent electronic conductivity, however it can undergo chemical corrosion especially under stop/start

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operation conditions¹¹⁷ leading to carbon oxidation, according to^{118, 119}:

 $C + 2H_2O \rightarrow CO_2 + 4H^+ + 4e^-$

 $(E^0 = 0.207 \text{ V vs. NHE at } 25 \text{ °C})$ (1) This leads to migration and coalescence or sintering of

catalyst particles or even their detachment with a consequent loss of electrochemical surface area (ECSA) and performance^{120, 121, 122}. Furthermore, platinum is considered to be involved in accelerating this oxidative process.¹²³

Fuel cell type	Material	Mass activity (A g ⁻¹)	Fibre diameter (nm)	ECSA (m ² g ⁻¹)	Advantages	Ref.
	Pt	47	50-70	3.6-6.2	electrochemical activity toward MOR ^a	92
DMFC	PtRh and PtRu	355.8 and 449.1	50	NA ^b	stability, enhancement of electrocatalytic properties by 30% relative to PtRu/C	94
	Pd/PA6 (polyamide 6) NF	NA	Pd/PA6 322, Pd layer 85	NA	facile and cost-effective synthesis of Pd NFs	99
Direct cyclohexane fueled PEMFC	Pt and PtRh	NA	30-40	NA	performance in fuel cell tests	93
DEMEC	Pt-NP into Pt	48.3	20-40	NA	mass activity increase by 50%	95
PEMFC	Pt-Fe	91	10-20	24	increase of specific activity compared to Pt/C	98

Table 1. Electrospun nanofibres for fuel cell electrocatalysts.

Much effort is being devoted to developing novel catalyst supports or secondary supports, including conductive or semi-conductive oxides such as TiO_x^{124} , $^{125, 126}$, SiO_2^{127} , MnO_x^{128} , WO_x^{129} , SnO_2^{130} as well as carbides¹³¹ and nitrides¹³², to modify the catalyst support for PEM fuel cells and enhance the catalytic performance of the system. Metal oxides represent a very attractive alternative to conventional catalyst supports, because of their exceptional chemical stability and corrosion resistance at the potentials and pH of use in PEM fuel cells¹³³. Such metal oxide-supported catalyst systems have already given promising results in terms of catalyst performance and durability⁸⁰ and also a better tolerance to methanol (in

the case of TiO_x/C)¹³⁴ and CO ($W_{18}O_{49}$)¹³⁵ with respect to conventional Pt/C electrodes. It was shown that a modification of the carbon support surface with semiconducting oxides, such as titanium oxide, reduces the electrochemically induced corrosion of the catalyst^{136, 137, 138}. Furthermore, the electrocatalytic activity is increased in comparison with Pt/C systems due to a synergetic interaction with the metal catalyst, the so-called strong metal-support interaction (SMSI)¹³⁹. A challenge that has still not been fully resolved is the brittleness of ceramic nanofibres that can make them fragile and difficult to handle. Paths being followed include mechanical pressing, preparation of twisted yarns, as well as formation of

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composites with polymers^{140, 141}

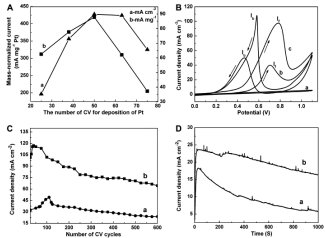


Fig. 5. (A) The plot of the peak current density of methanol oxidation on different Pt/CFM electrodes versus the cycle number of CV of Pt deposition, the potential scan rate is 100 mV s⁻¹, (B) cyclic voltammograms obtained for methanol oxidation on (a) CFM, (b) Pt/C and (c) Pt/CFM (Pt deposited by CV with 50 cycles) electrodes with the potential scan rate of 50 mV s⁻¹, (C) The plot of anodic peak current density of methanol oxidation on (a) Pt/C and (b) Pt/CFM electrode versus cycle numbers of CV with the potential scan rate of 100 mV s⁻¹, (D) Chronoamperometric curves of methanol oxidation on (a) Pt/C and (b) Pt/CFM at potential of 0.45 V. Reprinted from ref. 110 Copyright (2008), with permission from Elsevier.

Titania based materials are receiving attention as supports for PEMFC catalysts, showing excellent stability¹⁴² and promising activity for oxygen reduction¹⁴³ as well as for methanol oxidation¹⁴⁴. Nevertheless, the conductivity of stochiometric TiO_2 is poor (10^{-13} S cm⁻¹ at 298 K)¹⁴⁵, and doping with donor-type ions^{146, 147} or reducing it to nonstoichiometric compositions^{148, 149} are some of the possible methods used to enhance this property.

So far, only a few studies have been reported on the development of alternative FC catalyst supports elaborated by electrospinning and all of them deal with TiO₂. Formo et al. reported an electrocatalytic study on the methanol oxidation reaction of electrospun anatase nanofibres decorated with Pt catalysts in the form of NPs or NWs prepared by polyol reduction¹⁵⁰. The electrochemical activity and durability for the Pt nanowires and the Pt nanoparticles with a submonolayer coverage were improved compared to commercial Pt/C. Conductive titania electrospun nanofibres, obtained by Nb doping or H₂ reduction treatment, were also evaluated as Pt support materials towards oxygen reduction¹⁵¹. Pt supported on TiO₂ was more stable than Pt supported on C when subjected to 1000 voltammetric cycles in the range of 0.05-1.3V vs. RHE. Nb doped TiO₂ showed the highest stability, retaining 60 % of the ECSA after 1000 cycles while the Pt/C catalyst only retained 20 %. However, the carbon-based catalyst presented the highest ORR activity due to its larger specific area and the higher electrical conductivity of the support. Very recently, Long *et al.* developed a deposition method of Pt nanoparticles on electrospun titania fibres by microwave irradiation allowing loadings as high as 40 wt% without agglomeration¹⁵². Pt_{30wt%}/TiO₂ presented a higher activity toward methanol oxidation compared to commercial Pt_{20wt%}/C. These electrodes were very stable after electrochemical cycling and long storage. TiO₂ electrospun fibres were also used as supports for Pd nanoparticles for electrooxidation of glycerol¹⁵³. Such Pd/TiO₂ nanofibres can greatly promote glycerol electrooxidation in alkaline medium. Their application for oxidation of other alcohols such as methanol, ethylene glycol, and 1,2-propanediol was also demonstrated.

One of the greatest challenges in preparing electrospun TiO_2 -based cathodes for PEMFC is increasing their catalytic activity by improving the dispersion of Pt nanocatalysts. With that aim, new strategies are being developed such as photodeposition¹⁵⁴ or an *in situ* electrospinning-based method (see Fig. 6)⁸⁹ leading to Pt loaded conductive nanofibres with potential application as fuel cell electrodes.

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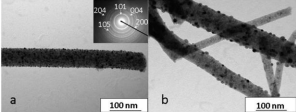


Fig. 6. Pt loaded Nb-doped TiO₂ electrospun nanofibres⁸⁹.

Concerning other non carbonaceous support materials for low temperature FC prepared by electrospinning, García-Márquez *et al.* have recently elaborated chromium nitride/carbide nanocomposite ceramic nanofibres¹⁵⁵. CrN being suggested as a suitable nonnoble electrocatalyst for cost effective fuel cells¹⁵⁶, the electrocatalytic properties of the pure fibre composites for the ORR were studied. Compared to commercial carbon-based electrodes, they showed a higher catalytic activity, but a lower current density due to lower surface area.

Few studies concerning electrospun electrodes for solid oxide fuel cells (SOFCs) have been reported as yet. The fabrication of ceramic components, such as

Table 2. Electrospun nanofibres for fuel cell electrodes

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yttria-stablized zirconia (YSZ) and gadolinia-doped ceria (GDC) in nanofibrillar structures is described^{157,} ¹⁵⁸, and recently, Ni/YSZ nanofibres, prepared by electrospinning and electroless plating, have been tested as the anodes in a commercial half cell and compared with a cell provided with conventional ballmilled powders of the same material¹⁵⁹. The peak power density for the cell with the fibre-derived anode was twice that with the powder-derived anode. This result illustrates the effect of the 1D morphology on the enhancement of ionic and electric transfer at the three-phase-boundary sites, and then of the overall performance.Electrospinning has also been recently used to prepare high porosity, surface area and conductivity Ba_{0.5}Sr_{0.5}Fe_{0.8}Cu_{0.2}O_{3-δ} fibres used as cathode material for an intermediate-temperature proton-conducting ceramic fuel cell¹⁶⁰..

Finally, carbon nanofibre mats produced by (gasassisted) electrospinning and solution blowing¹⁶¹ were used as electrodes in microbial fuel cells leading to high anode current densities¹⁶².

FC type	Material	Fibre diameter (nm)	Catalyst particle size (nm)	BET (m ² g ⁻¹)	ECSA (m ² g ⁻¹)	Ref
AFC	FeCo/CNFs	100-200	^a NA	NA	NA	108
	Pt/C-CFM ^b	130-170	commercial	23	NA	109
	Pt/CFM	130-170	50-200	7	NA	110
	PtRu/1-aminopyrene-CNFs	100-300	3.1-3.6	NA	32.7	112
	electrodeposited Pt/CNFs	200-300	20	NA	0.37-0.89 cm ^{2 c}	114
DMFC	Pt/Au-PAN and Pt/C/Au- PAN	740	2-6	NA	39.8 and 30.6	111
	Pt/polyimide-based CFM	≈ 200	< 3	NA	NA	113
	Pt/CFM	150	5	NA	NA	163
	Pt (NP and NW)/ TiO ₂	150-400	2-5/ 5-7×50/125	NA	56.2-16/ 17.6-16	150
	Pt/ TiO ₂	30-60	5	NA	NA	152
DAFC	Pd layer/ TiO ₂	115	61 (thickness)	NA	NA	153
DFAFC ^d	Electrodeposited Pt _x Au ₁₀₀₋ _x /CNF	≈700	118	NA	NA	116
	Pt/CNF acid	250	2.6-2.9	302	74.5	105
PEMFC	electrodeposited PtPd/CNFs	≈ 250-400	29-64	NA	NA	115
FENIFU	Pt/Nb doped TiO ₂	50-100	5	NA	5.0	151
	CrN-Cr ₃ C ₂	200-500	NA	25-110	NA	155
SOFC	Ni/YSZ	300	NA	NA	NA	159
/licrobial FC	CNF based biofilms	≈ 1000	NA	NA	NA	162

^aNA=data not available. ^bCFM=carbon fibre mat. ^cMass not reported. ^dDFAFC= direct formic acid fuel cell.

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3.a.3. Electrolyte membrane

Since electrospinning is such a versatile technique for producing nanofibre mats, it is being utilised for the development of composite fuel cell membranes with the aim of modifying the morphology and improving the mechanical properties. In the literature there have been two approaches in this particular aspect of electrospinning. The first involves electrospinning a non-conductive or less conductive polymer into a

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porous matrix, which acts as mechanical reinforcement when the pores are filled with a highly proton conducting component. An overview of the studies utilising this approach is listed in Table 3, with perfluorosulfonic acid (PFSA) ionomers such as Nafion[®] being the most commonly used polymer. In general, the matrix polymer is chosen so as to improve the mechanical or barrier properties (*e.g.* to reduce methanol crossover) of the composite membrane.

Table 3. Overview of reinforced	composite proto	n exchange membra	nes using an electro	ospun matrix.

Matrix Polymer	Filler Polymer	Details	Proton Conductivity (mS cm ⁻¹)	Ref.
Poly(vinyl alcohol)	Nafion [®]	CTAB used to facilitate electrospinning; PVA matrix crosslinked by treatment with glutaraldehyde and sulfonated	22 (70°C, 100 % RH [*])	164, 165
Poly(vinyl alcohol)	Nafion [®]	PVA matrix crosslinked by treating with glutaraldehyde	11 (70°C, 95 % RH)	166, 167
Polyvinylidene fluoride	Nafion®	PVdF electrospun from 1/1 Acetone/ DMAc	2 (65°C, hydrated)	168
Sulfonated polyethersulfone	Nafion [®]	excess Nafion [®] used to form a dense layer on one or both sides of the pore-filled membrane	~85 (RT, 100 % RH)	169, 170
Sulfonated random copolyimide	Sulfonated polyimide	proton conductivity higher compared to blend membrane from conventional (solution) methods	Up to 370 (90/10 blend at 80°C and 98 % RH)	171
Crosslinked bromomethylated sulfonated polyphenylene oxide (BPPO)	Sulfonated Poly(2,6- dimethyl-1,4- phenylene oxide) (SPPO)	BPPO electrospun from NMP and crosslinked by treating with NH ₃	30-80 (RT, 100 % RH)	172
* Dalativa humidity (0/)				

* Relative humidity (%)

Alternatively, a highly proton conducting matrix is electrospun into a porous fibre mat and this is then reinforced with a secondary polymer to provide mechanical stability. Most studies have focused on the electrospinning of PFSAs, but other sulfonated polymers such as poly(ether ether ketone) and poly(arylene ether sulfone) have also been electrospun into fibres. An overview of the publications in this area is given in Table 4.

In all cases, conductivity largely depends on the volume fraction of the proton conductive component, with a large percentage ratio of ionomer needed for high proton conductivity. In some studies, proton conductivity is reduced significantly due to the nonconductive component, however other properties of the membrane, such as mechanical and hydrolytic stability¹⁷² or reduced methanol permeability^{164, 168, 169} (for DMFC) are significantly improved. In the case of DMFC, despite their lower proton conductivity, electrospun composite membranes show comparable fuel cell performance to Nafion® due to the reduced methanol crossover and lower membrane thickness made possible by improved mechanical properties.

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Polymer Solvent		Additives/Carrier Polymer	Details / Post Treatment	Proton Conductivity (mS cm ⁻¹)	Ref.	
Nafion [®]	IPA/H ₂ O	12 wt% PAA	PAA $M_w 4.5 \times 10^5$	NA	173	
Nafion®	IPA/H ₂ O	5 wt% PVA or PEO	PEO $M_w 2 \times 10^5$; PVA $M_w 1.1 \times 10^5$	8.7-16	174	
Nafion®	MeOH	0.1 wt% high molecular weight PEO	PEO $M_w 8 \times 10^6$	1500 (single fibre of 400 nm diameter, 30°C; 90% RH)	175	
Nafion®	EtOH	PVP (1:2 PFSA/PVP wt ratio)	none	c.a. 8.5x10 ⁻³	176	
Nafion®	2/1 Alcohols/H ₂ O	PEO	PEO $M_w 2 \ge 10^5$, $3 \ge 10^5$, and $6 \ge 10^5$	NA	177	
Nafion [®]	H_2O	PEO, PVA, or PVP (very high ratio to PFSA)	large fibres (0.5-2 μ m)	NA	178	
3M perfluorosulfonic acid polymer	2/1 Propanol/H ₂ O	PEO (M _w 2 x 10 ⁵)	annealing, densification by pressing, and pore-filling with a resin	55 (80°C, 50%RH)	179	
3M perfluorosulfonic acid polymer	2/1 Propanol/H ₂ O	PAA, sulfonated polyhedral oligomeric silsesquioxane (sPOSS)	mat densified, annealed, and impregnated with a resin to form dense membrane	498 (35% sPOSS, 120°C 90%RH)	180	
Polymerised Ionic Liquid	3/1 MeCN/DMF	poly(MEBIm-BF4) , PAA	solutions of poly(MEBIm- BF ₄) was mixed with Nafion/PAA for electrospinning	7.1 x 10 ⁻⁴ (30°C, 10% RH)	181	
Sulfonated poly(ether ether ketone ketone)	DMF	none	membranes dried at 80°C for 24h	37 (spherical bead morphology at 25°C, 100% RH)	182	
Sulfonated poly(ether ether ketone)	DMAc	none	densification by pressing and welding by exposure to solvent vapours, and pore filling with an inert resin	41 (25°C 100% RH)	183	
Sulfonated poly(arylene ether sulfone)	DMAc	none	annealing, pressing, and welding of fibres by exposure to solvent vapours	86 (70% fibre volume fraction, 25°C, 100% RH)	184	
Sulfonated poly(arylene ether sulfone)	DMAc	sulfonated polyhedral oligomeric silsesquioxane (sPOSS)	mat densified, annealed, and impregnated with a resin to form dense membrane	94 (30°C, 80%RH)	185 186	
Sulfonated co- polyimide	DMF	none	pore filling with more highly sulfonated PI	~100 (80°C, 95%RH)	187	
Sulfonated polystyrene	DMF	none	fibres obtained at high concentration and PS molecular weight ($M_w 5 \times 10^5$)	NA	188	
Polyvinylidene fluoride	DMAc	Phosphotungstic acid (PWA, up to 12.8 wt%)	$PVdF M_w 2.2 \times 10^5$	~0.4 (6.7% PWA, 60°C, 100% RH)	189	
Aquivion TM	DMAc	PEO (M _w 1 x 10 ⁶)	annealing and densification by pressing	66 (120°C, 95%RH)	190	
Sulfated ZrO ₂	IPA	PVP , poly(2-acrylamido-2- methylpropanesulfonic acid) (pAMPS)	mat densified, heat treated, sulfonated and calcined to form S-ZrO ₂ membranes and pore-filled with	240 (100°C, 80% RH)	191	

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Electrospinning also introduces changes in the membrane morphology compared to conventional processing methods such as solution casting. Li et al.¹⁸² reported the electrospinning and electrospraying of sulfonated poly(ether ether ketone ketone) (SPEEKK), and found that the electrospun/sprayed membrane shows a more distinct SAXS ionomer peak, shifted to lower angles, compared to a cast membrane, indicating better phase separation and larger proton transport channels. In their study, the highest conductivity was obtained with a spherical rather than fibrous morphology, suggesting that proton transport occurs on the interface between particles rather than within the polymer itself. Since this first study, however, several others^{175, 176, 187} of single proton conducting nanofibres show greater apparent conductivity within the nanofibre than of a cast film. This is attributed to the alignment of the ionic aggregates that is believed to occur during electrospinning. Dong et al.¹⁷⁵ found that the proton conductivity increases with decreasing fibre diameter which suggests that the confinement of the nanofibre size may assist in aligning the ionic domains in the longitudinal direction. Furthermore, X-ray scattering at high relative humidity indicates that the ionic aggregates in the fibres are anisotropic and are oriented in the direction of the axis. Similarly, a study by Tamura et al.¹⁸⁷ on sulfonated polyimide nanofibres reported a much higher apparent conductivity of single nanofibres compared to cast films, with greater conductivity along the fibres than in the perpendicular direction of their membrane. The study by Pan et al.¹⁷⁶ also examined the possibility of using such single fibres in a micro-fuel cell, and a performance significantly higher than that of a conventional fuel cells was described.

In the case of PFSA, a carrier polymer is generally required to enable electrospinning due to aggregation of the PFSA chains. Electrospinning of Nafion® was first reported by Laforgue et al.¹⁷⁴ using PVA or PEO as the carrier polymer. Further studies have used PAA, PVP, or PEO, with comparable results. The presence of carrier polymer is detrimental to proton conductivity, especially when significant quantities are present, however more recent studies have minimised the carrier polymer content (or removed it through washing¹⁷⁹) and show comparable proton conductivities with cast films despite lower PFSA volume fraction in the electrospun membranes. Choi et al.¹⁸⁶ have incorporated sulfonated polyhedral oligomeric silsesquioxane (sPOSS) in order to increase the concentration of sulfonic acid groups, and

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p(AMPS)

thus were able to increase the proton conductivity of their composite membrane significantly beyond that of Nafion[®]. Some studies have also used PFSAs with shorter side chain than Nafion[®], which allows for lower EW ionomers to be used. When fibres of equal PFSA/carrier polymer ratios are compared, electrospinning with shorter side chain PFSAs was found to give smaller average fibre diameter in the case of 3M PFSA^{177, 179}, and also narrower fibre size distribution in the case of Aquivion^{TM 190} (see Fig.7).

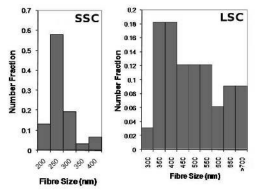


Fig. 7. Fibre size distribution of electrospun long (LSC Nafion[®])-side chain and short-side chain (SSC AquivionTM) PFSA.¹⁹⁰

Some investigations have produced a composite proton exchange membrane by filling the pores of the electrospun matrix with a curable, inert polymer for mechanical reinforcement. Studies published by Pintauro et al.^{179, 180, 184, 185, 186} used UV-curable polyurethane resin in an electrospun Nafion® or 3M PFSA matrix, and found that the proton conductivity of the composite membrane compares favourably to cast membranes, while the composite allows for lower swelling and improved mechanical properties (see Fig. 8). A different approach was reported by Takemori et al.¹⁷¹ where an electrospun sulfonated random copolyimide was pore-filled with a more highly sulfonated polyimide. The nanofibre composite membrane shows better proton conductivity and stability compared to a blend membrane of the same materials, indicating the benefits of the interpenetrating, nanofibre network morphology.

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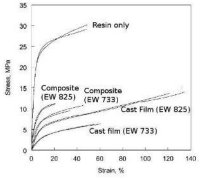


Fig. 8. Stress strain curves comparison of cast films and composite nanofibre membranes of 3M PFSA and a commercial UV-curable resin. Reprinted from ref. 179, Copyright (2010), with permission from the Royal Society of Chemistry.

A variant of this approach using ZrO_2 nanofibres was reported by Yao *et al.*¹⁹¹, who synthesised nanofibre mats of ZrO_2 which were then sulfonated by immersion in dilute sulfuric acid. After calcination, These are referred to as dye-sensitised solar cells (DSCs) due to their used of a dye photosensitiser, and they provide a technically and economically attractive alternative because of their low cost, light weight, and high power conversion efficiency¹⁹³. In contrast to the conventional silicon systems, where the semiconductor plays the role of absorbing light as well as the charge carrier and transport, in DSCs the two functions are separated. In DSCs, the photosensitiser absorbs light to generate excited electron-hole pairs that then undergo interfacial charge separation.

A typical DSC consists of a photoanode and a counter electrode separated by an electrolyte containing an iodide $(I^{-})/triiodide (I_{3}^{-})$ redox couple. Usually the photoanode is a transparent conducting oxide (TCO) coated with a film of a wide bandgap semiconductor (typically TiO2 nanoparticles) and the counter electrode is a TCO coated with a thin layer of platinum. Light is absorbed by the photosensitiser (a charge-transfer dye) linked to the oxide surface. The photo-electrons are then injected from the dye into the conduction band of the semiconductor and finally transported to the counter electrode through an external circuit. Here, the electrons are transferred to the electrolyte that carries them back to the dye molecule through a redox process, completing the circuit. Unlike conventional silicon solar cells, the use of dye-sensitisers with broad absorption bands in combination with nanostructured oxides allows DSCs to harvest a large fraction of sunlight.

So far, conversion efficiencies of 11 $\%^{194, 195}$ have been obtained, but the theoretical limit of 30% for a single junction semiconductor photovoltaic device¹⁹⁶

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the S-ZrO₂ mats were impregnated with crosslinked poly(2-acrylamido-2-methylpropanesulfonic acid) (pAMPS) to fabricate a composite membrane having high IEC and in which the authors suggest the existence of a continuous conducting pathway along the organic-inorganic interface.

3.b. Electrospun materials for dye-sensitised solar cells

Solar cells convert light into electricity and are based on the charge separation at an interface of two materials of different conduction bands. They have been dominated for the most part by inorganic solidstate p-n junction devices, usually Si-based, with high performance but very high material and production costs. Since its first discovery in 1991, a third generation of photovoltaic devices has emerged¹⁹².

remains far from being reached. One of the strategies to improve these devices is the introduction of new material morphologies, which sparks the recent interest in the integration of electrospun materials in DSCs in relation to all the components: photoanodes, counter electrodes and electrolytes. The effect and benefits of the 1D morphology on the obtained performances compared to the conventionally prepared devices is discussed in the following paragraphs and summarised in the Tables 5 and 6.

3.b.1. Photoanodes

There are two main requirements for the optimisation of DSC photoanodes which would lead to optimal light-to-electricity conversion efficiencies.

Firstly, charge transport must be enhanced while minimising charge recombination¹⁹⁷, which can be achieved by controlling the morphology, porosity and crystallinity of the photoanode material as these properties strongly influence the charge transport and recombination processes. For instance, electron transport in widely used mesoporous titania nanoparticle-based photoelectrodes depends on surface states and particle morphology as well as on interparticle connectivity. Recently, the use of onedimensional (1D) nanostructured materials in DSC, such as nanotubes, nanorods, nanobelts, nanowires and nanofibres, has been investigated^{198, 199, 200, 201}. 1D morphology shows better electron transport properties compared to conventional NP-based systems with disordered geometrical structures and interfacial interference in electron transport²⁰². Indeed, in

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monodimensional nanostructures (without exciton-like trap states²⁰³) electron transport is faster as such structures present lower grain boundaries due to their interconnectivity and high surface areas. The direct electrical pathways provided by the NFs ensure the rapid collection of charge carriers generated throughout the device.

Another requirement is to maximise the light harvesting properties of the device. Thus, photoanodes should possess a large surface area and high porosity in order to anchor the maximum amount of dye sensitisers²⁰⁶ (or quantum dots²⁰⁴), in which regard the 1D geometry of NFs gives promising results.

Furthermore, nanofibres provide an open structure which allows better pore-filling with the electrolyte compared to conventional NP-based electrodes. Such infiltration of a viscous electrolyte into a fibre network improves the contact with the semiconductor and helps

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the regeneration of oxidized dyes, leading to enhanced energy conversion efficiency²⁰⁵.

In this context, several recent studies describe 1D photoanode materials (mainly TiO_2 anatase) obtained by electrospinning (see Table 5). In order to further increase the dye loading and the overall efficiency of the solar cell, the electrospun nanofibres have also been shortened into nanorods (by mechanical pressure, grinding or ultrasonic bath) and DSCs prepared by using these rods alone or in combination with nanofibres and/or nanoparticles. One of the issues to be addressed for the electrospun fibre based DSCs is the poor adhesion of nanofibrous materials to conductive substrates. In order to solve this key problem, several approaches have been used and will be discussed here.

Material	Fibre diameter (nm)	Dye	V _{oc} (V)	J _{sc} (mA cm ⁻²)	FF	η (%)	Ref.
TiO ₂ (anatase)	150-700	N3	0.738- 0.849	3.60-9.88	0.45-0.51	1.34-4.14	206
TiO ₂ (anatase)	150	N3	0.78	9.45	0.57	4.2	207
TiO ₂ (anatase + rutile crystals)	$\approx 400-800$	N3	0.77-0.78	10.52- 11.24	0.56-0.58	4.60-5.02	210
TiO ₂ (anatase)	≈ 300 NR 15×60-100	N3	0.70	14.77	0.60	6.2	Erreur ! Signet non défini.
TiO ₂ (anatase) NFs + NPs	250	N719	0.64-0.71	15.9-22.4	0.66-0.72	7.10-10.3	209
TiO ₂ (anatase) NRs + NPs	100-200	N719	0.71-0.73	16.4-19.3	0.61-0.64	7.27-8.70	208
TiO ₂ (anatase) aligned NWs	100	N719	0.78	5.71	0.64	2.87	212
TiO ₂ (anatase)	80-100	QD (CdS/CdSe)	0.33-0.69	1.93-9.74	0.31-0.48	0.31-2.69	Erreur ! Signet non défini.
Nb doped TiO ₂ (anatase)	150	D131	NA	9.74-10.0	NA	NA	213
TiO ₂ (rutile)	100-200 NRs	N3	0.748- 0.815	12.14- 12.37	0.4547	4.17-4.56	211
ZnO	200-500	N719	0.60	3.58	0.62	1.34	217
ZnO	NA	N719	0.57-0.63	4.58-9.14	0.57-0.66	1.77-3.02	218
Al doped ZnO	40-160	N719	0.33	6.24-6.34	0.26	0.54-0.55	219
SnO ₂ /ZnO	NA	D149	0.57	2.99	0.59	1.01	220
ZnO NWs/C NFs	50/1160	black dye	0.17-0.35	2.60-11.2	0.25-0.28	0.1-1.1	221
ITO	200	N719	0.64-0.65	9.97-10.20	0.57-0.59	3.65-3.97	222

Table 5. Electrospun fibres for DSC photoanodes and the resulting photovoltaic parameters^{*} obtained under air-mass (AM) 1.5 and 100 mW

^{*}Power conversion efficiency of photovoltaic cells is given by: $\eta = FF \times V_{oc} \times J_{sc} / P_{in}$ with $FF = V_{max} \times J_{max} / V_{oc} \times J_{sc}$ where P_{in} is the power of the incident light; V_{oc} the open-circuit voltage; J_{sc} the short-circuit current density; FF=fill factor; the voltage and current density at the maximum power point are V_{max} and J_{max} , respectively.

One possibility is the use of a pre-deposited titania

layer and a solvent post-treatment to soften the

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electrospun anatase nanofibre mats to improve their adhesion to the substrate²⁰⁶. These treatments improved the photocurrent and the conversion efficiency of the corresponding device, which reached a value of 4.14 %. A similar result was obtained using TiO_2 pre-deposited film by spin-coating²⁰⁷. Furthermore, it was shown that the effective electron diffusion coefficient of the NF-based DSC is similar to that of high efficiency nanocrystalline samples. The low electron lifetime and obtained efficiency were explained by the existence of surface traps in the nanofibres.

Another strategy to overcome this problem is to deposit the TiO₂ electrospun nanofibres on TiO₂ nanoparticles. Such an approach was used by Chuangchote *et al.*^{208, 209} and showed an improvement of the light harvesting properties and of the overall efficiency (up to 10.3 %) by combining these morphologies (see Fig. 9). Furthermore, the coexistence of NFs or NRs and NPs enhanced the porosity and thus the penetration of the liquid electrolyte and the amount of the adsorbed dye.

The issue of electrolyte penetration into the photoanode material was studied in particular for quasi-solid electrolyte DSCs (cf. § 3.b.3). Nanorod electrodes were used to fabricate such devices and led to efficient photocurrent generation and an overall conversion of 6.2 % Erreur ! Signet non défini. A hot pressing procedure was successfully used to increase the fibre adhesion to the conductive glass. In another study, electrospun TiO₂ NFs were used to epitaxially grow TiO₂ rutile crystals. This additional layer modified the photocurrent generation of the prepared DSCs and the energy conversion efficiency increased 30% after post-treatment, reaching 5 %²¹⁰. In recent studies, the NR morphology seems to show superior performance compared to continuous nanofibres. Rutile 1D NFs and NRs based DSCs were analysed and the higher efficiency of the latter (4.6 vs 1.8%) was attributed to a strong electronic coupling efficiency of photoexcited electrons. The short size of the rods causes a decrease in the resistance and an increase of the currents²¹¹.

A further strategy to improve light harvesting and charge collection can be the use of vertical arrays of TiO₂ nanowires. The fabrication of such vertically aligned materials by electrospinning followed by a top-down process is very recent (see Table 5)²¹².

Nb doped anatase nanofibres have been recently explored as photoanode materials for dye-sensitised solar cells²¹³. The defects derived from doping play an important role in their photovoltaic properties. The electron diffusion and mobility improved with doping, however the BET surface area decreased and the

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charge recombination increased.

ZnO has been investigated as an alternative to TiO₂ photoanode material for DSCs, as it possesses similar electron injection process and bandgap as that of TiO₂ (3.3 eV vs. 3.2 eV for anatase)²¹⁴. In particular one dimensional ZnO nanomaterials, which show increased conductivity path, have attracted great attention²¹⁵. In spite of the higher charge transport and electron mobility, 1D ZnO has poor chemical stability in acidic dye solutions and low surface area compared to titania NP films (and consequently adsorb a minor amount of dye)²¹⁵, resulting in energy conversion efficiencies which are always lower than TiO₂ based systems (6.58 % for NP based DSCs²¹⁶ and around 1% for 1D ZnO based DSCs²¹⁷). Many strategies have been applied in order to overcome these drawbacks, and again to improve the fibre adhesion on the substrate and consequently to enhance the overall performance of the systems. The use of electrospun ZnO fibres in dye-sensitised solar cells was first published in 2007²¹⁷. To enhance the electrical contact between the nanofibres and the electrode, hot pressing of the ZnO/PVAc precursor NFs was performed before their calcination. This treatment resulted in the formation of dense networks of twisted nanofibres, which further improved the direct conduction paths for efficient electron collection as well as the porosity and surface area. Nevertheless, the efficiency of these systems was low (1.34 %). A higher energy conversion efficiency of 3.02 % was achieved when a Zn(OAc)₂ treatment followed by calcination was deposition²¹⁸. performed after the fibre Electrochemical impedance spectroscopy results showed the effective suppression of the back reaction between photoelectrons (in the ZnO conduction band) and I_3^- (in the electrolyte) in the $Zn(OAc)_2$ treated systems, leading to the observed improved performance. In order to further enhance its optical and electrical properties, ZnO NFs have also been doped with Al²¹⁹, which enhances their adhesion to fluorine-doped tin oxide (FTO). This was achieved by introducing an Al-doped ZnO seed layer prepared by electrospinning before the deposition of the Al-doped ZnO electrospun nanofibres. It was shown that NFs did not crack nor peel away after the calcination step, as this layer facilitated the release of interfacial tensile stress during calcination and improved the interfacial adhesion to the conducting substrate. As a result, the efficiency improved when compared to DSC without seed layers (0.54-0.55 % vs 0.03-0.04 %).

Electrospun nanofibres were also used to form light-scattering layers on photoanode nanomaterials. Such layers could reflect the unabsorbed photons back into the NW anode and improve the DSC efficiency²²⁰

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For instance, Wu *et al.* deposited SnO_2/ZnO nanofibres by electrospinning on the surface of ZnO NWs and formed a film by methanol vapour treatment followed by high-temperature calcination. J_{sc} , V_{oc} , and FF of the nanofilm/ZnO NW DSCs were all enhanced compared to those of the ZnO NW DSCs.

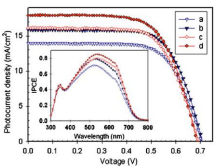


Fig. 9. Photocurrent density-voltage characteristics of NP electrode (a, c) and NP/NF electrode (b, d) at AM1.5 global illumination. **Inset:** Incident photon-to-current conversion efficiency spectra for those electrodes. The area of electrode was 0.25 cm². The thicknesses of TiO₂ particle layers were 8.4 μ m (a, b) and 15.5 μ m (c, d). Reprinted from ref. 209, Copyright (2008), with permission from American Institute of Physics.

brittleness and high price. Recently, 1D nanostructured networks such as carbon nanotubes and silver nanowires revealed good performances as potential alternatives. However, they present some disadvantages like high junction resistances for CNTs and limited aspect ratios and high price for silver NWs. In this context, electrospinning has been used to prepare copper nanofibre networks by calcination of copper acetate/PVA electrospun fibres that were subsequently reduced in a H_2 atmosphere²²³. The power efficiency of organic solar cells using copper nanofibre electrode materials was 3.0 %, a value comparable to devices made with standard ITO electrodes. Cu fibres have ultrahigh aspect ratios (up to 100000), very low junction resistances and great flexibility. Furthermore, their preparation bv electrospinning is low-cost and easily scaleable.

Finally, TiO_2^{224} and ZnO^{225} nanofibrous networks have found very recent use in hybrid solar cells based on poly(3-hexylthiophene) and metal oxides.

3.b.2 Counter electrodes

The conventional counter electrode in DSCs is platinum, due to its high electrocatalytic activity for I_3^- ions reduction¹⁹³. Nevertheless, Pt is expensive and the corrosive I^{-}/I_3^- redox couple can reduce its activity and affect the long-term stability of the overall device²²⁶. Recent studies deal with the replacement of Pt in order to decrease the costs and enhance the

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We can also cite the use of electrospun materials $(e.g. carbon)^{221}$ as templates for the growth of ZnO NWs. These composites can be used as an anode material for the fabrication of flexible DSCs.

So far, several studies focused on the active layer on which photosensitive dye is adsorbed have been described here. Another strategy to increase the efficiency of electron transport and improve DSC efficiency is to maximise the interface between the conductive glass electrode and the active layer. Recently, electrospinning of transparent and conductive indium tin oxide (ITO) was performed from a mixture of inorganic precursors and PVP and the resulting nanofibres deposited onto ITO-glass substrates. As an electrode of DSC²²², improvement leading to 3.97 % efficiency was achieved, due to a reduction in grain boundaries, an efficient high charge collection, and a rapid electron transport.

Electrospun nanofibres can also be used to replace the conductive glass substrate for DSCs which is usually ITO. Indeed, ITO films present several bottlenecks to the real application, including

durability of DSCs. Some of them are still based on Pt,

but in the form of nanoparticles dispersed on conductive materials. For instance, Pt impregnated on mesoporous Nb-doped TiO₂ films resulted in large active surface area and high mechanical stability²²⁷. Other studies have revealed that carbonaceous materials including graphite, carbon black, and CNTs, can exhibit electrocatalytic performance comparable to Pt^{226, 228}. Such materials, in contrast to Pt, are abundant and low-cost, which could facilitate the introduction into the market of dye-sensitised solar cells. For example, electrospun carbon nanofibres have been used recently, as an alternative to platinum for triiodide reduction in DSCs (see Fig. 10)²²⁹. Such counter electrodes exhibited low charge-transfer resistance, large capacitance, and fast I3- reduction rates. The obtained current density was comparable to those of conventional Pt based cells, although the open circuit potential was lower. The efficiency was also slightly lower because of the lower fill factor, probably due to high total series resistance, R_{Stot}. Thus, although the lower price of such carbon-based conterelectrode makes them an attractive alternative to Pt, the main challenge is in fabricating thinner and highly porous carbon NFs in order to reduce the thickness of the counter electrode and also to decrease R_{Stot} to make their performance more competitive.

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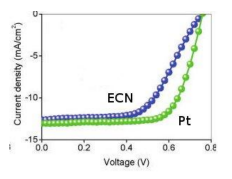


Fig. 10. Current density-voltage (*J-V*) curves of the DSCs based on electrospun carbon nanofibres (ECN) and Pt counter electrodes. Reprinted from ref. 229, Copyright (2010), with permission from American Chemical Society.

3.b.3. DSC electrolytes

The conventional electrolytes for DSCs are liquids (e.g. triiodide/iodide redox couple in organic solvents) leading to overall efficiencies around 11%¹⁹⁴. The major drawbacks to practical use of such systems is the poor long-term stability due to the volatility of organic solvents, the difficulty in robust sealing, and the leakage of electrolyte in case of breakage. In order to overcome these problems, many studies have been carried out on alternatives to replace the liquid electrolytes. For instance, solid-state electrolytes such conductors²³⁰ and polymers²³¹, that as hole unfortunately show low solar-to-electricity а conversion efficiency because of their low electron injection efficiency. A better alternative is represented by quasi-solid-state DSCs using polymer gel electrolytes²³² presenting high thermal stability, negligible vapour pressure, good contacting and filling properties with the nanostructured electrodes, and higher ionic conductivity²³³.

So far, several types of polymer gel electrolytes have been used in quasi-solid-state DSCs, such as polyacrylonitrile $(PAN)^{234}$, poly(ethylene oxide) $(PEO)^{235}$, polymethylmethacrylate $(PMMA)^{236}$, and poly(vinylidenefluoride-co-hexafluoropropylene)

(PVdF-HFP)²³⁷. In particular, the latter shows high ionic conductivity and stability at room temperature.

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However, the complex preparation technology and the poor mechanical strength of these gel polymer based DSCs represent a bottleneck to their introduction to the market. To overcome this problem, the electrospinning of such polymers has been performed with the aim of integrating the resulting fibrous, easy to obtain and low-cost materials as electrolytes (see Table 6).

In 2002, Drew et al. reported an "electrospun photovoltaic cell" including electrospun fibres of PAN with an azo-dye (Congo Red) with or without TiO₂ nanoparticles.²³⁸ This system was still based on a liquid junction and showed photocurrents higher than those obtained with a spin-coated film of the same materials. In 2008, Priya et al. prepared an electrospun PVdF-HFP membrane and used it for the first time to form quasi-solid-state DSCs²³⁹. Its solar-to-electricity conversion efficiency was similar to that obtained for a conventional liquid electrolyte (7.3 % vs. 7.8 %) while its long-term stability was higher. Indeed, the electrospun membrane electrolyte effectively encapsulated the organic solvent containing the $I_3/I^$ redox couple and promoted a strong interfacial contact between the dye-adsorbed TiO₂ electrode and the Pt counter-electrode. Other polymer electrolytes for quasi-solid state DSCs have been fabricated with electrospun PVdF-HFP^{240, 241, 242}, and DSC devices were fabricated using them²⁴¹. The photovoltaic performance of DSC devices using such nanofibre films showed better results compared to DSC devices using the same material films obtained by spin-coating (see also Table 6)²⁴⁰. In addition, it was found that the photocurrent density and efficiency of DSC using electrospun PVdF-HFP nanofibres in electrolytes were not proportional to the electrolyte conductivity. A further improvement was achieved very recently by using electrospun PVdF-HFP/polystyrene(PS) blend nanofibres (3:1) which resulted in a 5.3% conversion efficiency²⁴². Lastly, the stability of the electrospun polymer electrolyte-based DSC has been compared to that of the liquid electrolyte-based DSC: after deliberately provoking holes and removing glasses from the cell, the fibrous electrolyte maintained its integrity while the liquid one underwent leakage (see Fig. 11).

Table 6. Electrospun fibres for DSC electrolytes and the resulting photovoltaic parameters obtained under AM 1.5 and 100 mW cm⁻² of light intensity.

Material	Fibre diameter (nm)	Dye	Ionic conductivity (m S cm ⁻¹)	V _{oc} (V)	J _{sc} (mA cm ⁻²)	FF	η (%)	Ref.
PVdF-HFP	800-1000	N3	4.53	0.75	12.3	0.57	5.21	241
PVdF-HFP/ (PS)	800-1200	N719	1.89/2.65	0.71/0.76	11.4/11.8	0.64/0.66	5.37/5.75	242

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PVdF-HFP	600	N719	10-2	0.76	15.57	0.62	7.3	239

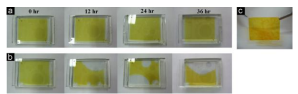


Fig. 11. Stability testing of (a) the electrospun PVdF-HFP nanofibre electrolyte cells, (b) the liquid electrolyte cells, and (c) soaked electrospun PVdF-HFP nanofibre after 36 h. Reprinted from ref. 242, Copyright (2011), with permission from Elsevier.

3.c. Electrospun nanofibres in lithium-ion batteries

Lithium-ion batteries (LIB) have attracted much research attention and industrial development due to their high energy density, low gravimetric density, long cycle life and flexible design²⁴³. Indeed, since the 1990s, non-aqueous rechargeable Li-ion batteries have established a strong market position especially for portable electronic devices. In such systems, electrical energy is generated by conversion of chemical energy via redox reactions at the electrodes²⁴⁴. In the "rocking-chair" LIB, reactions follow an insertion mechanism: lithium cations are reversibly inserted/extracted into/from a host matrix and this process is accompanied by an electron flow through an external circuit. Upon charging, Li⁺ are released by the cathode and inserted at the anode, while upon discharging, Li⁺ are extracted from the anode and inserted into the cathode. Similarly to fuel cells, the Li-ion cell consists of an anode and cathode separated by an electrolyte, which is an electronic insulator but a Li⁺ conductor. A conventional cell uses a carbon/graphite anode, a lithium-metal oxide (LiCoO₂, LiNiO₂, LiMn₂O₄) cathode, and an organic electrolyte of lithium hexafluorophosphate (LiPF₆) salt with ethylene carbonate (EC)-organic solvent mixture.

So far, LIB have enabled the development of small and portable electronic devices, but future challenges concerning larger and more demanding applications such as electric or hybrid electric vehicles necessitate significant advances in battery technology as they require fast charging and discharging at high power rates. Thus, more improvements are needed in various aspects such as energy density, power, cycling life, charge/discharge rate, safety and cost² in order for LIBs to achieve satisfactory performance in these demanding applications.

One of the strategies used to improve the efficiency and the durability of LIBs is to maintain a low weight, volume and cost by designing and fabricating nanomaterials and nanostructured materials for use as electrodes and electrolytes^{245, 1}. In this regard, the versatility of electrospinning makes them highly suitable to prepare both these components with the benefits of smaller, nanostructured geometries and architectures and reduced manufacturing costs. In this next section, recent advances in the areas of electrospun cathodes, anodes, and separator membranes for LIBs are described and summarised (see also Tables 7-9).

3.c.1 Electrode materials

In order to achieve high specific energy and energy density, electrode materials for rechargeable Li-ion batteries must possess a high specific charge and charge density, generate a high cell voltage, and have a high reversibility of electrochemical reactions at both electrodes to maintain the specific charge for thousands of charge-discharge cycles. Nanostructured materials have been extensively explored to prepare high-performance LIB electrodes, due to the multiple advantages that they possess^{246, 247}. Firstly, the nanometre size significantly increases the rate of Li⁺ insertion/extraction because of the short diffusion length for lithium-ion transport within the particles, enhancing the rate capability and power density. The same is also true for electron transport, which is also enhanced in nanomaterials. Secondly, their high surface area allows a high electrode/electrolyte contact area that can increase the rate capability of the device. Finally, the range of composition of solid solutions is often more extensive for nanoparticles, and the strain associated with insertion/extraction is often better accommodated, avoiding destruction of the material structure (electrochemical grinding) and improving the cycle life of the cell.

Despite these advantages, the use of nanomaterials in LIB also introduces new challenges. The large surface area can enhance the reaction between electrodes and electrolyte, leading to an increase in solid-electrolyte interfacial (SEI) layer area, self discharge, and decrease in cycle life. Furthermore, the low packing density of nanostructured materials can result in lower volumetric energy density compared to micrometric and bulk materials. There may also be issues regarding the complexity of the synthetic methods and the difficulty of accurately controlling the size of nanomaterials, which may result in greater processing and manufacturing costs. Taking these challenges into account, among the wide range of morphologies nanowires show great potential in this

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area because of their better percolation behaviour than particles^{1, 248}. Soft-chemistry routes and template syntheses are usually employed to synthesize nanostructured electrodes, and after sintering treatment they often result in large grain size and aggregation, losing the high surface areas and its associated merits²⁴⁹. In this regard, electrospinning presents a simple and versatile alternative approach for preparing nanostructured anode and cathode hosts for lithium-ion batteries^{Erreur ! Signet non défini.} and the main results in this area will be discussed here.

Cathode materials

Recent years have witnessed a growing development of the electrospinning of transition-metal oxide with different phases (layered, anatase, spinel, etc...) as cathodes for lithium-ion batteries. By modifying the electrospinning configuration and the post-treatments, a wide variety of morphologies and architectures were studied such as nanowires²⁵⁰, nanobelts²⁵¹, nanonuggets²⁵², biaxial²⁵³ and triaxial²⁵⁴ core-sheath, and hollow²⁵⁵ fibres.

Among the materials available, most commercial batteries use $LiCoO_2$ in the cathode due to its good performance in terms of specific energy density (theoretical capacity 140 mA h g⁻¹) and cycle life²⁵⁶. Nanostructured $LiCoO_2$ fibre electrodes have been prepared by electrospinning with the aim of achieving a fast solid state diffusion due to the short diffusion distance of Li⁺ cations^{257, 258}. Gu *et al.*²⁵⁸ prepared

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LiCoO₂ fibre electrodes with large surface area and small pores, with a high initial discharge capacity of 182 mA h g⁻¹ compared with ca. 140 mA h g⁻¹ for conventional powder and film electrodes. However, such nanostructured cathodes suffered from a large loss of capacity during the charge-discharge process resulting from the dissolution of cobalt and lithium cations forming Li₂CO₃ and CoF₂ impurities. A promising strategy to increase stability and cycling performance is by coating of the nanofibres. For instance, using coaxial electrospinning, LiCoO₂ fibres²⁵³ and $LiNi_{0.8}Co_{0.1}Mn_{0.1}O_2$ tubes²⁵⁵ covered with a MgO shell were prepared. The MgO coating avoided impedance growth by protecting the surface from passive film formation during cycling. The core-sheath fibre electrode showed improved reversibility and cyclability. After 40 cycles, 90 % of the initial discharge capacity of LiCoO2/MgO was maintained, compared to 52 % for the uncoated fibre electrode. The hollow fibres showed still better performances in terms of capacity and cycling stability (see Fig. 12 and Table 7). Similarly. electrospun core-shell Li(Ni_{1/3}Co_{1/3}Mn_{1/3})O₂/Li(Ni_{1/2}Mn_{1/2})O₂ fibres exhibited an enhanced cycle stability compared to core $Li(Ni_{1/3}Co_{1/3}Mn_{1/3})O_2$ fibres with the loss of 24.5 % of the initial capacity after 40 cycles²⁵⁹. A lithium phosphorous oxynitride (LiPON) layer deposited by radio-frequency sputtering was also effective in stabilising LiCoO₂ fibre performance, with a loss of 0.11 % per cycle after 100 cycles at the discharge rate of 0.05 mA cm^{-2 260}.

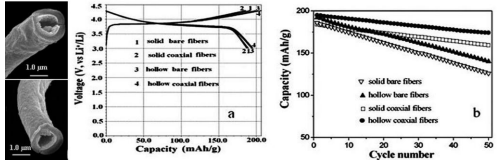


Fig. 12. SEM micrographs of hollow coaxial fibres obtained at (a) 0.4 and (b) 0.5 molar ratios of PVP/(Ni²⁺ + Co²⁺ + Mn²⁺). The initial charge-discharge curves (c) and capacity retention versus cycle number plots (d) of the fibre electrodes at a current density of 20 mA g⁻¹. Reprinted from ref.255, Copyright (2008), with permission from American Chemical Society.

Table 7. Electrospun	cathodes and the corresp	ponding electrochemica	l performance.

Material	Fibre diameter (nm)/structure	Crystallite size (nm)	BET surface (m ² g ⁻¹)	Discharge capacity 1 st cycle (mAh g ⁻¹) [rate (C)]	Discharge capacity n th cycle (mAh g ⁻¹) [rate (C)]	Ref.
LiCoO ₂	500-2000	20-35	18	182 [0.14]	n=2, 173, n=20, 123 [0.14]	258

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1000-2000 shell 10-100	<10	20	181 [0.14]	n=40, 163 [0.14]	253
60-80/ 3D architecture	NA^*	NA	129 [NA]	n=100, 120	260
1000-2000 30-60 shell 300- 500 wall/ hollow coaxial	NA	30	195 [0.14]	n=50, 174 [0.14]	255
< 5000 shell 300	20-50	NA	192 [0.1]	n=2, 171, n=5, 166 [0.1]	259
≈100-200 NFs/nanonuggets	NA	3-6	≈500-800 [1], ≈240-150 [1.5]	[1]; n=50,	252
100-300	18-28	NA	166 [0.05], 140 [0.1], 125 [0.2], 98 [0.5], 71 [1], 37 [2]	$n=50, \approx 140$ [0.1], ≈ 120 [0.2]	263
≈400-1000/ triaxial	NA	NA	160 [0.06], 130 [0.6], 80 [6]	n=20, ≈160 [0.06], ≈130 [0.6], ≈80 [6]	254
100/ shell 2-5	NA	NA	169 [0.1], 162 [0.5], 150 [1], 114 [5], 93 [10]	n=100, 146 [1]	264
40-70×10-20/ nanobelts	NA	NA	350 [NA]	n=25, 270 [NA]	251
100-200	50×100 NRs	NA	275-390 [NA]	n=50, 187-201 [NA]	250
300-800	70-80	NA	300 [0.1], 160 [1.0]	n=20, 200 [0.1], 112 [1.0]; n=50, 150 [0.1]	265
350	≈100	97	377 [NA]	n=10, 347, n=40, 344 [NA]	266
	shell 10-100 60-80/ 3D architecture 1000-2000 30-60 shell 300- 500 wall/ hollow coaxial < 5000 shell 300 ≈100-200 NFs/nanonuggets 100-300 ≈400-1000/ triaxial 100/ shell 2-5 40-70×10-20/ nanobelts 100-200 300-800	shell 10-100 $\langle 10$ $60-80/$ NA* $1000-2000$ NA $30-60$ shell 300- NA 500 wall/ NA hollow coaxial $\langle 5000$ $\langle 5000$ 20-50 $\approx 100-200$ NA $100-200$ NA $100-300$ 18-28 $\approx 400-1000/$ NA $100/$ NA $100/$ NA $40-70 \times 10-20/$ NA $100-200$ NA $100-200$ NA $300-800$ $70-80$	shell 10-100 $\langle 10$ 20 $60-80/$ NA* NA $3D$ architecture NA* NA $1000-2000$ $30-60$ shell 300- NA 30 500 wall/ NA 30 hollow coaxial < 5000 $20-50$ NA < 5000 $20-50$ NA $3-6$ $\approx 100-200$ NA $3-6$ $100-300$ $18-28$ NA $100-300$ $18-28$ NA $\approx 400-1000/$ NA NA $100'$ NA NA $40-70 \times 10-20/$ NA NA $100-200$ 50×100 NA $100-200$ 50×100 NA $300-800$ $70-80$ NA	shell 10-100 $\langle 10$ 20 $181 [0.14]$ $60-80/$ 3D architectureNA*NA $129 [NA]$ $1000-2000$ $30-60 shell 300-500 wall/NA30195 [0.14]hollow coaxial< 5000shell 30020-50NA192 [0.1]\approx 100-200NFs/nanonuggetsNA3-6\approx 500-800 [1],\approx 240-150 [1.5]100-30018-28NA1-66 [0.05],140 [0.1],125 [0.2],98 [0.5],71 [1], 37 [2]\approx 400-1000/triaxialNANA160 [0.06],80 [6]100/shell 2-5NANA162 [0.5],150 [1], 114 [5], 93 [10]40-70 \times 10-20/nanobeltsNANA350 [NA]100-20050 \times 100NRsNA300 [0.1],160 [1.0]300-80070-80NA300 [0.1],160 [1.0]$	shell 10-100 $\zeta 10$ 20 $181 [0.14]$ $[0.14]$ $n=2, 128,$ $n=100, 120$ $[NA]3D architectureNA*NA129 [NA]n=2, 128,n=100, 120[NA]1000-200030-60 \text{ shell } 300-50NA30195 [0.14]n=50, 174[0.14]hollow coaxial< 5000$

*NA=data not available

Furthermore, the particular 3D structure electrode of the latter material, presenting highly accessible surface areas and continuous networks seems very promising for the design of novel batteries that maximise power and energy density.

LiFePO₄ is a highly promising candidate for positive electrode material because of its large specific capacity (170 mA h g⁻¹), high discharge potential, good thermal stability, low toxicity, low cost and safety. However, it suffers from low ionic and electronic conductivity, leading to low rate capability and high impedance in LIBs. One of the most investigated strategies to avoid this limitation is coating this material with carbon in order to introduce an electron path²⁶¹. Furthermore, such a coating protects the surface from undesired side reactions leading to Fe³⁺ and Li₃PO₄²⁶². Several studies are devoted to composite LiFePO₄/carbon nanofibres obtained by electrospinning^{254, 263, 264}. Very recently, Zhu *et al.* reported on singlecrystalline thin (100 nm) carbon-coated LiFePO₄ nanowires²⁶⁴. The carbon coating and the threedimensional connectivity of the network ensured high conductivity that, together with the liquid electrolyte, led to fast Li⁺ transport. Very good rate performance and cycling capability were obtained. For instance at 0.1 C discharge rate, the capacity was close to theoretical (169 mA h g⁻¹) and after 100 cycles at 1 C it was 86% of the theoretical value (146 mA h g⁻¹). Furthermore, after cycling, the morphology of the carbon-coated LiFePO₄ fibres was maintained.

Vanadium pentoxide has high theoretical capacity (ca. 400 mA h g^{-1}), but the performance of conventional bulk V₂O₅ powder cathodes is limited by low electronic conductivity, slow lithium diffusion rate and structural instability upon charge/discharge cycling. To overcome these problems, the use of electrospinning to prepare nanostructured V₂O₅

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positive electrodes is growing ^{250, 251, 265}. For instance, Cheah *et al.* have prepared single phase electrospun V_2O_5 fibres that present an initial discharge capacity of 320 mA h g⁻¹ and excellent coulombic efficiency ($\approx 100\%$) throughout 50 charge-discharge cycles²⁶⁵. This study highlights once again the importance of the network morphology of electrospun fibres to increase the electroactive surface area, thus improving the accessibility of electrolyte to the cathode material and the overall efficiency of the Li-ion batteries. The use of mesoporous nanofibres further improves the performance especially in term of cycling stability²⁶⁶.

Anode materials

Cycle life and rate capability of lithium ion batteries strongly depend on the properties of the anode material, while their electrical storage capacities depends on how much lithium can be held in said anode. Most current batteries use carbon as its anode, replacing metallic lithium found in early batteries. Carbon provides several advantages such as safety, low cost and greater cell life, however it also has lower cell voltage, specific charge (theoretical capacity of 372 mA h g⁻¹), and rate capability. In order to improve these properties, researchers have investigated control of the anode nanostructures in order to increase its performance with regards to the above parameters.

Carbon nanofibre (CNF) anodes have been prepared through a combination of polymer electrospinning and thermal treatments. PAN derived fibres treated at 1000 °C exhibited the highest reversible capacity for pure carbon (ca. 450 mA h g⁻¹), and a good rate capability (350 mA h g⁻¹) (at a charge current of 100 mA h g⁻¹) due to their particular microtexture, comprising a highly disordered structure, defects, and dangling bonds²⁶⁷. CNFs prepared from a bicomponent polymer mixture (PAN/polypyrrole) also showed a relatively large reversible capacity (ca. 360 mA h g⁻¹), high rate capability, good cycle performance and structural integrity after 50 charge/discharge cycles, attributable to their large surface area and extended interface with electrolyte, leading to short pathway for charge and electron transport²⁶⁸.

Studies on anode materials also concern electrospun carbon fibres loaded with metal or oxide nanoparticles. For instance, CNF have been doped with silicon^{269, 270} and $tin^{271, 272, 273, 274}$ NPs. Indeed, Si and Sn have high theoretical capacity (4200 mA h g⁻¹

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and 992 mA h g^{-1} , respectively), but suffer from poor cycling performance owing to large volume changes by up to 400% for Si and nanoparticle aggregation upon the alloying and dealloying reaction with Li⁺ ions²⁷⁵. One of the strategies proposed to overcome these drawbacks is by dispersing nano-sized particles within carbon nanofibres that act as structural buffers, particle stabilisers, and electroactive materials, and eliminate the need for binding or conducting additives. The effect of the surrounding confinements of nanoparticles is fundamental, and was studied in the case of Si by Choi et al.²⁷⁰. Hard confinements contributed to suppressing the volume expansion of Si NPs during charge/discharge cycles. The improvements on reversible capacity and cycle stability were mainly attributable to the formation of nanofibrillar networks favouring electronic and ionic transport through the surrounding confinements as in the case of Si@CNF. It was shown that an important role is also played by the electrical conduction of the surrounding material. Carbonisation temperature is also a key factor for electrospun NPs/C anode materials since affecting their morphology and electrochemical performance. Yu et al. prepared tin nanoparticle-dispersed carbon (Sn/C) nanofibres by stabilisation of electrospun SnCl₄/PAN fibres and subsequent carbonisation at different temperatures²⁷⁴. Sn/C nanofibres at 700 and 850 °C presented the highest charge (785.8 and 811 mA h g⁻¹) and discharge $(1211.7 \text{ and } 993 \text{ mA h g}^{-1})$ capacities due to the unique feature of reticular nanofibre geometries.

Fibrous carbon anodes have also been loaded with transition-metal oxide nanoparticles such as Co₃O₄²⁷⁶, $Fe_3O_4^{277}$ and MnO_x^{278} . Such materials have been widely investigated as anode materials in LIBs, because of their high theoretical capacities, safety, non-toxicity and low cost²⁷⁹. However, they have a poor electronic conductivity and a large volume change during repeated lithium insertion/de-insertion. The formation of composite nanofibres by insertion of nanometric metal oxides in electrospun CNFs is again a promising solution. One example is provided by the preparation of MnOx/CNFs by electrospinning combined with electrodeposition; high reversible capacity (618 mA h g⁻¹), good capacity retention, and great morphological integrity were shown after cycling (see Fig. 13).

The exceptional improvement of anode performance (see Table 8) upon the incorporation of a Li-inactive metal such as Ni in CNF²⁸⁰ is particularly noteworthy.

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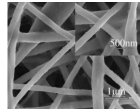


Fig. 13. SEM images of MnO_x/CNFs after 100 charge/discharge cycles at a constant current density of 50 mA g⁻¹. Deposition time: 10 h. The inset shows MnO_x/CNFs with higher magnification. Reprinted from ref. 278, Copyright (2010), with permission from Elsevier.

The last typology of electrospun anode materials for LIBs is represented by pure metal oxide fibres^{281, 282, 283, 284, 285, 286, 287}. Among them, SnO₂ and TiO₂ have been intensively investigated²⁸⁸. Due to their particular porous structure, electrospun SnO₂ fibres consisting of nanoparticles were shown to facilitate electrolyte diffusion, charge transfer and the mechanical strain due to the volume changes during charge/discharge **Erreur**! Signet non défini. The electrochemical performance

was improved, reaching high specific capacity (446 mA h g⁻¹ after 50 cycles at a 100 mA g⁻¹ rate), and high rate capability (477.7 mA h g⁻¹ at 10 C). However, it was considered that the large irreversible capacity and capacity fading that were observed probably derived from side reactions of the large surface area with the electrolyte. The performance was further improved by exploiting the versatility of electrospinning and preparing SnO₂ porous nanotubes, that delivered a high discharge

Table 8. Electrospun anodes and the corresponding electrochemical performance.

Material	Fibre/NP diameter (nm)/structure	Crystallite/pore size (nm)	BET surface (m ² g ⁻¹)	Discharge capacity 1 st cycle (mAh g ⁻¹) [rate (C)]	Discharge capacity n th cycle (mAh g ⁻¹) [rate (C)]	Ref.
C NFs	200-300	0.5-5	NA	1000 [0.08]	n=2, 500 [0.08]	267
C NFs	≈ 150-300	NA	NA	621 [0.13]	n=20, 479, n=50, 454 [0.13]; n=2, 440 [0.27]; n=2, 345 [0.54]	268
C NFs + Si NPs	NA	NA	NA	1710 [0.1] 870 [0.33]	n=5, 1150, n=50, 1100 [0.1]; n=30, ≈700 [0.33]	269
C NFs + Si NPs	300-500 50-80 Si	NA	NA	1060-780 [0.01]	n=50, 410-380 [0.01]	270
C NFs + Sn NPs	$\frac{100-1000}{\text{Sn}} \approx 1$	NA	NA	816 [NA]	n=20, 400 [NA]	271
C NFs + Sn/SnO _x NPs	≈50-500 Sn 20-40	NA	352	1192 [0.03]	n=40, ≈520 [0.03]	272
C NFs + Sn NPs	150-350 Sn 30-40 150	NA	NA	1211.7 [0.025]	n=2, ≈800 [0.025] n=100, ≈780	274
C NFs + Sn@C NPs	hollow fibres wall thickness 30 Sn 100@10	NA	NA	1156 [0.1]	$[0.1]; n=100, \approx 450$ [5]	272
C NFs + Ni NPs	≈ 150-200 Ni 20	NA	NA	795 [0.13], ≈570 [0.8], ≈450 [1.34], ≈400 [2.7]	n=50, ≈500 [0.13], ≈400 [0.8], ≈350 [1.34], ≈250 [2.7]	280
C NFs + MnO _x NPs	≈ 150-200 MnO _x 21-42	NA	NA	≈650 [NA]	n=2, 558, n=50, ≈500 [NA]	278
$C NFs + Fe_3O_4 NPs$	≈ 350 Fe ₃ O ₄ 8-52	NA	NA	1551 [0.22] 623 [1.73]	n=7, 763, n=80, 1007 [0.22]	277
SnO ₂	≈50-80	10	NA	1650 [0.1]	n=50, ≈480 [0.13]	Erreur ! Signet non défini.
SnO ₂	220 nanotubes	pores 11	28	1650 [0.23]	n=2, ≈1100 n=50, 807 [0.23]; n=40,	Erreur ! Signet non

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	wall thickness 25				≈350 [2.4]; n=30,	défini.
					≈150 [10]	
TiO ₂	60	10	54	175[0.45]	n=50, ≈90 [0.45]	248
	100-200/	<100	NT A	192 [0.5],	n=30, 140 [0.5],	289
Li ₄ Ti ₅ O ₁₂	3D cross-bar NFs	<100	NA	170 [1.5]	87 [1.5]	289
						Erreur !
Co_3O_4	200	<50	NA	1336 [0.5]	n=40, 604 [0.5]	Signet non
						défini.
Co ₃ O ₄ /C	200-300	50-100	NA	1146 [0.11]	n=20, 861 [0.11]	276
					$-5 \sim (50)$	Erreur !
Mn_3O_4	100-300	20-25	NA	2200 [NA]	n=5, ≈650, n=50, ≈450 [NA]	Signet non
					II-30, ~430 [INA]	défini.
A a an Aul	50	6				Erreur !
Ag or Au/ TiO ₂	NP 5-10		25-53	180-300 [1]	n=2, 160-170[1]	Signet non
ΠO_2	NP 3-10	pores 3.5				défini.
				164 [0.2]		Erreur !
$Li_4Ti_5O_{12}$	< 1000	NA	169		n=1200, 100 [22]	Signet non
+ graphene				110 [22]		défini.

capacity of 807 mA h g⁻¹ after 50 cycles and retained a high fraction of their theoretical capacity even after cycling at high rates^{Erreur ! Signet non défini.} Lu *et al.* prepared 3D architectures stacked from a cross-bar array of aligned electrospun fibres²⁸⁹. They compared cathodic performances of 3D networks of anatase TiO₂ and spinel Li₄Ti₅O₂ and they found that the first was destroyed after Li⁺ insertion, while the latter showed a

stable architecture and a good reversibility in electrochemical cycles.

Recently LIB anodes based on titania nanofibres loaded with 10% Au or Ag nanoparticles showed improved performance compared to the bare material. The inserted metallic NPs not only decreased the charge transfer, but also enhanced the electronic conductivity of the composite TiO_2 . The capacity was improved by 20% or more compared to bare 1D TiO_2 especially for the material doped with gold.

Finally, another promising example of metal oxide anode material obtained by electrospinning is tricobalt tetraoxide, known for its high reversible capacity and electrochemical stability. Co_3O_4 electrospun nanofibres delivered an initial discharge capacity of 1336 mA h g⁻¹ that, after 40 cycles, was maintained at 604 mA h g⁻¹. This result can be ascribed to the high surface area developed by the NFs^{Erreur ! Signet non défini}.

3.c.2 Li-ion battery separators

Polymer electrolytes have received significant attention in lithium-ion battery applications since they have the advantages of being safer than liquid electrolytes. In general, the polymer electrolyte uses a polymer gel consisting of an electrolyte solution encapsulated in a polymer such as poly(vinylidene fluoride-co-hexafluoropropylene) (PVdF-co-HFP). Despite their advantages, polymer gel membranes also have drawbacks due to poor mechanical properties and sometimes unsatisfactory thermal stability. In some cases, this can be improved by, for example, using highly crystalline PVdF homopolymer, however the crystallinity of the polymer hinders the migration of lithium ions, and as such is detrimental to the ionic conductivity of the membrane and thus to the performance of the battery.

In order to overcome this problem, a nanoporous or microporous matrix presents an attractive alternative to a gel, as such a matrix provides better interconnectivity for ionic conduction within the membrane. Earlier work focused on other methods such as phase inversion or casting of a polymer gel, whereby precise control of the porosity may be difficult. Furthermore, it may be difficult to completely remove residual solvent from within the matrix, which may affect electrochemical properties or stability of the final separator. Electrospun membranes offer an alternative with high electrolyte uptake and good homogeneity with some degree of control over the morphology of the membrane. An overview of the studies on such membranes is given in Table 9.

The majority of the publications in this field have utilised PVdF and its copolymers, although several studies have also used polyacrylonitrile. In both cases it is generally found that the electrospun separator performs better than gel membranes or phase inversion membranes due to its high uptake of electrolyte and the interconnected conduction pathway within the membrane. A comparative study by on membranes made by electrospinning and phase inversion³⁰⁵ found that electrospun membranes not only have greater porosity, but also a more uniform structure and significantly higher ionic conductivity leading to better cell performance. A study on a PAN membrane separator has also shown better cycle life and thermal stability than a conventional

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polyolefin microporous separator³⁰⁰ (see Fig. 14).

Several studies have also added inorganic additives such as SiO₂ or Al₂O₃ in order to increase cell performance. Generally, addition of inorganic oxides increases the ionic conductivity of the membrane, which in some studies with PAN fibres^{302, 309} was attributed to the smaller average fibre diameters (up to a certain inorganic content) due to the electrostatic repulsion of the inorganic particles during electrospinning, leading to membranes with greater porosity and electrolyte uptake. In contrast, other studies on PVdF-HFP^{297, 307} concluded that the presence of silica produced the opposite effect on fibre diameter, but a similar increase in ionic conductivity and electrolyte uptake up to a certain SiO₂ content which was attributed to a decrease in PVdF crystallinity.

Another approach is the development of a multilayered composite membrane using one or more electrospun layers. A study using electrospun PVdF-CTFE/Al₂O₃³⁰⁶ deposited on both sides of a porous polyethylene membrane shows better stability and cycling performance as the electrospun layer helps confine the electrolytes within the membrane and also increases electrolyte

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uptake. To decouple these effects, in another study a trilayer membrane of PVdF-HFP and PAN was fabricated through continuous electrospinning with the different polymer solutions (a PVdF-HFP layer sandwiched between two PAN layers and vice versa). The multilayered composite did not perform as well as a membrane electrospun from a solution-based blend of the two polymers, indicating that there are more beneficial interactions between them when electrospun from the same solution, resulting in a greater porosity and electrolyte uptake despite larger average fibre diameter.

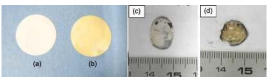


Fig. 14. Electrospun PAN nanofibre separator (a, b) and conventional polyolefin separator (c, d) before and after thermal stability tests. Reprinted from ref. 300, Copyright (2008), with permission from Elsevier.

Material	Av. fibre diameter (nm)	Anode/cathode materials	Conductivity (1M LiPF ₆ , mS cm ⁻¹)	Discharge capacity 1 st cycle (mAh g ⁻¹) [rate (C)]	Discharge capacity after (n) cycles (mAhg ⁻¹)	Ref.
PVdF	250	none	1.7	NA^{*}	NA	290
PVdF	450	MCMB ^a /LiCoO ₂	>1	≈135 [0.5]	≈135 (50)	291
PVdF-HFP	500	MCMB/LiCoO ₂	>1	≈150 [0.5]	≈150 (n=50)	292
PAN	330	MCMB/LiCoO ₂	>1	145 [0.5]	136 (n=150)	293
PVdF, PVdF- HFP	420	MCMB/LiCoO ₂	1.04	141 [1]	NA	294
PVdF-HFP	1 µm	Li/LiFePO ₄	4.7	125 [1]	117 (n=100)	295
PVdF- HFP/IL ^b	~700	Li/LiFePO ₄	2.3	149 [0.1]	152 (n=24)	296
PVdF- HFP/SiO ₂	1-2 µm	Li/LiFePO ₄	8.06	170 [0.1]	NA	297
PVdF- HFP/NP ^c	1-5 µm	Li/LiFePO ₄	7.2 ^c	164 ^d [0.1]	156 (n=30)	298
PI ^e	300-625	MCMB/LiNi _{0.8} Co _{0.2} O ₂	NA	~320 [0.1]	~305 (n=21)	299
PAN	380 / 250	graphite/LiCoO ₂	NA	124.5 ^f [0.5]	102.7 (n=250)	300

Table 9. Studies on electrospun lithium battery separators

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AR	TIC	LE 1	YPE

PVdF- HFP/SiO ₂	2 µm	Li/LiFePO ₄	4.3	170 [0.1]	160.5 (n=80)	301
PAN/SiO ₂	270	graphite/LiCoO ₂	11	139 [0.5]	127 (n=150)	302
PVdF	~125	Li/LiFePO ₄	N/A	177 [0.1]	150 (n=10)	303
PVdF- HFP/PAN ^g	320-490	Li/LiFePO ₄	3.9-6.5	145 [0.1]	136 (n=50)	304
PVdF-HFP	1 µm	Li/LiFePO ₄	4.32	140 [0.1]	128 (n=50)	305
PVdF- CTFE/Al ₂ O ₃	2-4 µm	MCMB/LiCoO ₂	0.49	143.5 [0.1]	131.8 (n=200)	306
PVdF/SiO ₂	490	none	4.7	NA	NA	307
PVdF-CTFE	230	Sn-C/ LiNi _{0.5} Mn _{1.5} O ₄	>2	120 [0.33]	114 (n=10)	308
PAN/LTO ^h	230	Li/LiFePO ₄	1.95	162 [0.2]	156 (n=50)	309

^{*}NA=data not available. ^aMCMB=mesocarbon microbeads. ^bRoom temperature ionic liquids incorporated in the membrane. ^cMetal oxide nanoparticles of SiO₂, Al₂O₃, or BaTiO₃ incorporated in the membrane. ^dQuoted value is of BaTiO₃-containing membrane. ^ePI used as a matrix to confine gel electrolyte containing PVdF or PAN. ^fValue is of PAN membrane of 380 nm fibre diameter. ^gMultilayered composite membrane. ^hLTO=Lanthanum titanate oxide

3.d. Electrospun fibres for supercapacitors

Electrochemical supercapacitors work by storing an

electrical charge in a metal/electrolyte interface by means of two mechanisms: electrostatic charge transfer for electrochemical double layer capacitors (EDLC) and faradaic charge transfer for pseudocapacitors310. Although the concept was industrialised a long time ago, it is undergoing renewed interests due to recent demands for energy storage devices with high power density and rapid charge/discharge rate in applications such as digital electronics and medical implants. Supercapacitors complement batteries and conventional capacitors, being used to deliver high power pulses, load-levelling or electricity storage. In an EDLC, the capacitance is determined by the surface area accessible to the electrolyte, and thus 1D nanomaterials such as carbon fibres have attracted significant attention due to their high surface area⁷. More recently, electrospinning has been investigated as a versatile and simple fabrication method for the highly porous, nanofibrous electrode and separator materials for supercapacitors. An

overview of the developments in this area is given in Table 10.

As many commercial supercapacitors are based on porous carbon^{311, 312}, several studies have focused on elaboration of electrospun carbons. The most part concerns poly(acrylonitrile) (PAN)-based precursor, such as the study by Kim et al.³¹⁴ on activated carbon nanofibres (ACNF), fabricated by stabilisation and carbonisation of electrospun nanofibres obtained from this polymer, followed by activation using steam to create pores and increase the surface area of the material. Other precursors, such as PBI^{316, 318}, PI³¹⁷, and isotropic pitch³²³ have also been reported. The performance of these ACNFs greatly depends on the activation temperature that can lead to an optimum surface area. Higher activation temperature increases the volume fraction of mesopores (even though the overall surface area may decrease at very high temperatures due to fusing of the pores), which improves the specific capacitance at high current densities due the faster transport of solvated ions enabled by mesopores³¹³. Such carbon nanofibres can also be bonded together by the addition of a thermoplastic such as PVP to the precursor, which would partially melt during the early stages of pyrolysis and creates inter-fibre linkages.

Table 10. Electrospun materials for supercapacitors and the corresponding properties and	l performance.
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Material	Fibre diameter (nm)	Specific surface area (m ² g ⁻¹)	specific capacitance (F g ⁻¹) / current density or scan rate	Ref.
PAN	300	850-1230	120 / 1000 mA g ⁻¹	314, 315

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PBI	250	500-1220	125-178 / 4 mA	316
PAA	1-2 µm	900-1450	175 / 1000 mA g ⁻¹	317
PAN, PBI, PI, IPP ^a	50-2000	1000-2500	120-210 / 1000 mA g ⁻¹	318
PAN/ZnCl	200-350	310-550	120-140 / NA	319
PAN/Ag	200-500	NA	190-250 / NA	320
PAN/CNT ^b	~600	810	280 / 1000 mA g ⁻¹	321
PAI ^c	350	1080-1360	150 / 10 A	322
PAN/IPP	750	722-1724	75.5-143 / 1 mA cm ⁻¹	323
PAN/Ni	210	NA	164 / 200 mA g ⁻¹	324
PVP/PAN	~390-800	~530-810	~160-220 / NA	325
PCS^d	~3-10 µm	1740-3116	NA	326
PVP/PEDOT ^e	350	NA	19 / 1000 mA g ⁻¹	327
TiO ₂ -RuO ₂	< 1000	20	687 / 10 mV s ⁻¹	331
110 ₂ -KuO ₂	N 1000	20	460 / 1000 mV s ⁻¹	551
Pt-RuO ₂	200-300	NA	409.4 / 10 mV s ⁻¹	334
Ft-KuO ₂	200-300	NA	322 / 1000 mV s ⁻¹	554
RuO_2 - Ag_2O	40-70	NA	$173 / 10 \text{ mV s}^{-1}$	332
$RuO_2 - RuO_2 \cdot nH_20$	400-500	59	886.9 / 10 mV s ⁻¹	333
KuO_2 - KuO_2 · <i>n</i> II_2 0	400-500	59	$pprox 620 / 2000 \text{ mV s}^{-1}$	555
RuO_2-MnO_x	RuO ₂ 400	NA	208.7 / 10 mV s ⁻¹	337
composite mat	MnO _x 100			
$RuO_2-Mn_3O_4$	150-600	NA	293 / 10 mV s ⁻¹	329
LaNiO ₃ -Mn O _x	< 1000	NA	$160 / 10 \text{ mV s}^{-1}$	336
V_2O_5	500-600	3-10	190-250 / 5 mV s ⁻¹	328
SnO ₂ /C	400-500	NA	$187 / 20 \text{ mV s}^{-1}$	335
coaxial TiN/VN	core 300	169	247.5 / 2 mV s ⁻¹	338
	shell 660	109	160.8 / 50 mV s ⁻¹	330
and instanting with the second	bONT	manatal a CDAL mala (amile	inite dDCC astronytheory that it as	PEDOT 1 (2.4

^aIPP= isotropic pitch precursor ^bCNT= carbon nanotubes ^cPAI=poly(amide imide) ^dPCS=polycarbomethylsilane ^cPEDOT=poly(3,4-ethylenedioxythiophene)

In order to improve the capacitance of carbon electrodes, some studies have reported the addition of components such as silver³²⁰, nickel³²⁴, or carbon nanotubes³²¹ (CNTs). It was found that the conductive fillers improve the capacitance of the electrode, although the exact mechanism for this effect remains unclear. The addition of CNT is believed to improve the specific capacitance by improving the conductivity of the material, the effect of which becomes more obvious at higher current densities. However, the study using a Ag additive did not show a similar correlation of conductivity and capacitance, as the capacitance continue to increase even when the conductivity remains unchanged. Other investigations have focussed on $ZnCl_2$ as an additive, which improved the capacitance by enhancing the stabilisation and carbonisation of PAN nanofibres³¹⁹. The ZnCl acted as a dehydrating agent, eliminating water and catalysing graphite formation. Furthermore, as it decomposed, it produced chlorine gas, which etched the carbon fibres and created micropores on their surface.

Very recently, electrospinning has also been utilised to produce a flexible supercapacitor³²⁷. The all-textile supercapacitor utilised PEDOT-coated electrospun PVP electrodes and an electrospun PAN separator. Although its performance was limited and utilised only half of the electrode capacitance, the supercapacitor assembly was fully flexible with no delamination of the layers upon bending (see Figure 15).

Pseudo-capacitors are currently attracting much attention as energy storage devices due to their high capacitances (200-1300 F g⁻¹) in comparison to EDLCs (50-150 F g⁻¹)⁷. Transition metal oxides such as RuO₂, MnO_x^{329, 330} and V₂O₅³²⁸ are attractive pseudo-supercapacitor electrodes, because of their several possible oxidation states, high electrical conductivity, and electrochemical stability³¹⁰.

RuO₂ fulfils these requirements, and due to its high cost it is studied as nanomaterial or in composite materials. For instance, capacitors based on RuO₂ electrodeposited onto electrospun TiO₂ nanorods show a high rate capability (see Table 9)³³¹. Similarly, RuO₂-Ag₂O composite nanowires produced 300% higher capacitance compared to the single RuO₂ NW electrodes³³².

 RuO_2 in its hydrated form (*i.e.* $RuO_2 \cdot nH_2O$) is a promising pseudo-capacitors material because of its high specific theoretical capacitance, high proton conductivity, and excellent electrochemical cyclability. To overcome possibly its only drawback apart from cost, the low electronic conductivity, the versatility of electrospinning in combination with other synthesis methods gives a solution, allowing to

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prepare conductive core fibres coated with the active material.

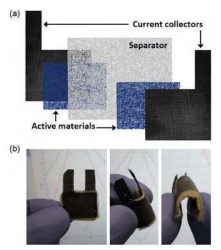


Figure 15. Schematic representation of the components in the all-textile supercapacitor (a) and photographs the flexible supercapacitor (b). Reprinted from ref. 327, Copyright (2011), with permission from Elsevier.

Electrochemical capacitors were then fabricated by depositing a thin hydrous RuO₂ layer onto a crystalline RuO₂ nanofibrous mat by cyclic voltammetry. Such electrodes showed a high specific capacitance (886.9 F g⁻¹ at a scan rate of 10 mV s⁻¹, based on the mass of the RuO₂·*n*H₂O coating) and high rate capability³³³. Highly porous and conductive electrospun Pt fibres were also used as core support for electrochemically deposited hydrous RuO₂ overlayers³³⁴. The resulting composite electrode presented good performance with a capacity loss of only 21.4% passing from 10 to 1000 mV s⁻¹. The same strategy is also used for producing other composite materials useful in electrochemical capacitors, such as SnO₂ on C nanofibres³³⁵ and MnO_x on LaNiO₃ fibre mats³³⁶.

Electrospinning is also capable of fabricating composite mats by co-depositing different materials with different properties, as in the case of layer-by-layer stacked MnO_x -RuO₂, in which the first offers a facile ion pathway and the latter a fast electron pathway. The multi-stacked morphology provided high surface area and high porosity allowing an efficient permeability of the electrolyte and a good surface activity³³⁷. Coaxial electrospinning was also exploited to prepare core-sheath titanium nitride-vanadium nitride fibres, which combined higher specific capacitance of VN and high rate capability of TiN³³⁸, exhibiting high specific capacitance and high rate capability (Table 9).

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4. Conclusions and future perspectives

Development of new functional materials for energy-related applications with targeted nanomaterial architectures is essential to meet the performance and durability targets of energy conversion and storage applications. Along with many other avenues of research under investigation, 1D nanostructured materials are likely to significantly contribute to meeting some of these challenges as the structural control of nanomaterials can be as important as the materials themselves.

Amongst the fabrication method for nanofibres and nanowires, electrospinning is undergoing renewed intense interest and current advances indicate a high potential of electrospun materials in energy related applications. It has been established as a relatively facile method which provides researchers with great control over the fibre morphology. Electrospun nanofibrous mats show good homogeneity and porosity, and the application of electric field and stretching that occurs during electrospinning may also confer beneficial effects in certain cases, such as for ionomers.

Several challenges remain to be overcome before large scale use of electrospun materials in energyrelated applications, and this is partly due to the fact process parameters affecting that the the electrospinning technique are still not fully understood. Thus, there are still limitations on the type materials that can be synthesised of bv electrospinning, and it may be difficult to fully eliminate defects such as beads and obtain completely homogeneous nanofibres. For applications such as nanofibrous catalysts and supports, supplementary developments will be needed should further reduction of fibre diameters be required. Additional challenges are related to scale-up of electrospinning and downstream processing, these being essential steps for the introduction of electrospun materials in industrial application in energy devices. Progress with multi-jet emitters/nozzles increases both productivity and cover area of large scale nanofibre production339, 340 Furthermore, although advances in aligning and bridging adjacent fibres have been made, still further solutions must be found to temper the fragility of ceramic electrospun nanofibre networks, for which elimination of the carrier polymer while maintaining fibre integrity and composition represent additional challenges that can impede transfer to the target application.

Despite the challenges, electrospinning has been shown to be capable of producing nanofibre mats of excellent uniformity and further developments have

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brought about more complex architectures such as core-shell and hollow fibres by coaxial electrospinning, as well as novel composite materials by the use of co-electrospinning. As we look for solutions to meet new technological requirements, electrospinning is emerging as one of the principal methods in the synthesis of nanofibres as it is a powerful and versatile technique which would be relatively simple to scale-up and which appears attractive from an industrial perspective.

Acknowledgements

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Abbreviations

1D	One-dimensional
NF	Nanofibre
NW	Nanowire
NR	Nanorod
NP	Nanoparticle
FC	Fuel cell
DSC	Dye-sensitised solar cell
LIB	Li-ion battery
CNF	Carbon nanofibres
CFM	Carbon fibrous mat
PEMFC	Proton exchange membrane fuel cell
AFC	Alkaline fuel cell
DMFC	Direct methanol fuel cell
DAFC	Direct alcohol fuel cell
DFAFC	Direct formic acid fuel cell
SOFC	Solid oxide fuel cell
ECSA	Electrochemical surface area
ORR	Oxygen reduction reaction
MOR	Methanol oxidation reaction

Notes and references

PFSA	Perfluorosulfonic acid
PAN	Poly(acrylonitrile)
PVA	Poly(vinyl alcohol)
PEO	Poly(ethylene oxide)
CTAB	Cetyl trimethylammonium bromide
DMAc	Dimethylacetamide
sPOSS	Polyhedral oligomeric silsesquioxane
PWA	Phosphotungstic acid
BPPO	Bromomethylated sulfonated polyphenylene
	oxide
SPPO	Sulfonated Poly(2,6-dimethyl-1,4-phenylene
	oxide)
poly(ME	EBIm-BF ₄)
	Poly(1-(2-methacryloyoxy)ethyl-3-
	butylimidazolium tetrafluoroborate
MeCN	Acetonitrile
DMAc	N,N-dimethylacetamide
DMF	N,N-dimethylformamide
IPA	Isopropyl alcohol (2-propanol)
PMMA	Polymethylmethacrylate
PVdF	Polyvinylidenefluoride
PVdF-H	FP
1,01,11	• •
1 / 01 11	Poly(vinylidenefluoride-co-
1 / 01 11	
PVdF-C	Poly(vinylidenefluoride-co- hexafluoropropylene)
	Poly(vinylidenefluoride-co- hexafluoropropylene)
	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE
	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co-
PVdF-C	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene
PVdF-C PI	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide
PVdF-C PI PS	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene
PVdF-C PI PS PVAc	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate
PVdF-C PI PS PVAc AM	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass
PVdF-C PI PS PVAc AM V _{oc}	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage
PVdF-C PI PS PVAc AM V _{oc} J _{sc}	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density
PVdF-C' PI PS PVAc AM V_{oc} J_{sc} FF	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor
PVdF-C PI PS PVAc AM V_{oc} J _{sc} FF η	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency
PVdF-C PI PS PVAc AM V_{oc} J _{sc} FF η R _{ct}	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance
PVdF-C PI PS PVAc AM V_{oc} J _{sc} FF η R _{ct} TCO	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance Transparent conducting oxide
PVdF-C PI PS PVAc AM V _{oc} J _{sc} FF η R _{ct} TCO ITO	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance Transparent conducting oxide Indium tin oxide Fluorine-doped tin oxide Solid-electrolyte interface
PVdF-C PI PS PVAc AM V _{oc} J _{sc} FF η R _{ct} TCO ITO FTO SEI LiPON	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance Transparent conducting oxide Indium tin oxide Fluorine-doped tin oxide Solid-electrolyte interface Lithium phosphorous oxynitride
PVdF-C PI PS PVAc AM V _{oc} J _{sc} FF η R _{ct} TCO ITO FTO SEI LiPON EDLC	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance Transparent conducting oxide Indium tin oxide Fluorine-doped tin oxide Solid-electrolyte interface Lithium phosphorous oxynitride Electrochemical double layer capacitors
PVdF-C PI PS PVAc AM V _{oc} J _{sc} FF η R _{ct} TCO ITO FTO SEI LiPON	Poly(vinylidenefluoride-co- hexafluoropropylene) TFE Poly(vinyledenefluoride-co- chlorotrifluoroethylene) Polyimide Polystyrene Polyvinylacetate Air-mass Open-circuit voltage Short-circuit current density Fill factor Light-to-electricity conversion efficiency Charge-transfer resistance Transparent conducting oxide Indium tin oxide Fluorine-doped tin oxide Solid-electrolyte interface Lithium phosphorous oxynitride

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