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Electrospinning based on benign solvents: current definitions, implications and strategies

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Electrospinning evolved as a powerful technology to produce membranes consisting of nano- to micron-scaled fibers offering unique membrane properties due to the extremely high surface-to-volume ratio and high membrane porosity. However, this technology is often far removed from a green and environmentally-friendly process, as it employs solvents with issues such as high flammability, toxicity, difficult disposal, or energy-intensive synthesis. Indeed, the most commonly used solvents in the electrospinning field are halogenated (e.g. chloroform, trifluoroethanol) and toxic solvents (e.g. dimethylformamide), the use of which is now restricted by the Chemical Control Regulation in the European Union (REACH). This is especially important when considering the commercialization of electrospun products due to the more limited solvent choice. In order to render electrospinning more attractive as a commercial technique and further improve scalability, solvent alternatives and other green routes have to be established, which adhere to societal and legal restrictions especially in regard to the environmental and health impact. Therefore, this review provides the current ecological and economical context of this technology, and summarizes recent approaches towards green electrospinning. Since electrospinning from solutions and dispersions constitutes the predominantly used form of this technique both in research and industry, the focus of this work is set on the use of solvents or solvent mixtures classified exclusively by the term "green". The approaches presented comprehensively cover the production of polymeric fibers *via* solution, emulsion, suspension, and *in situ* cross-linking electrospinning.

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1. Introduction

Electrospinning (ES) has developed into a well-known, versatile technique for the preparation of polymeric non-woven fibers. The range of applications for electrospun fibers is broad: particle filtration from liquid or gas,¹ drug-delivery,² tissue engineering,³ functional wound dressings,⁴ light-weight battery membranes,⁵ customizable sensors,⁶ sound absorption,^{7,8} etc. The success of the ES technique originates in the possibility to generate fibers from the micro- down to the lower nano-meter range by tailoring fiber diameter, surface area, porosity and other fiber surface properties, offering a very high surface-to-volume ratio superior to common textile fiber production techniques.⁹ Furthermore, the obtained nanofiber membranes possess very high air and liquid permeability, while commonly enabling diffusion of solutes and cells, which are requirements for the majority of their current applications. This versatility is further extended by the range of different materials

that can be electrospun, including biomaterials and tailored synthetic polymers. In contrast, ES of polymer solutions is limited by the use of a solvent or solvent mixtures. Standard ES compositions often include halogenated compounds due to their high solvation power for hydrophobic polymers and low boiling points. These types of solvents have caused more and more concern from a scientific, public, and industrial point of view mainly due to their high toxicity and high environmental impact. Additionally, halogenated compounds recently became strongly regulated, for example by the Chemical Control Regulation in the European Union (REACH).¹⁰ Regarding the commercialization of products using the ES technique, many companies hesitate to implement this method into their production line since it implies large know-how on solvent handling and their proper disposal, which is also extremely cost-intensive. For industrial-scale production, ES involves the use of large volumes of solvents requiring recycling or waste management facilities. This is mainly due to relatively low polymer concentrations in the spinning solutions needed to obtain a defined viscosity of the commonly used high molecular weight polymers.

Generally, solvent-free fiber manufacturing techniques, such as melt-ES,^{11,12} as well as the industrially well-established

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melt blowing^{13,14} and melt spinning,¹⁵ offer the greenest options to pursue. However, such techniques require high temperatures to melt the polymers, that can lead to chemical degradation and may not be suitable for the processing of bio-active molecules.^{16,17} Furthermore, solvent-free techniques result in fibers with usually larger diameters than their solution electrospun counterparts, but under certain circumstances, also nanofibers can be obtained.¹⁸ Melt ES in particular combines fiber diameters in the nanometer range and a solvent-free methodology, while also offering spatial precision for nano-patterning; unfortunately it is significantly less accessible.¹⁹ As this review focusses on recent accomplishments in improving the greenness of the classical solvent-based ES technique, the vast field of solvent-free ES falls beyond its scope.

Regarding classical solvent-based ES, many studies in literature have shown that it is possible to spin almost all commonly employed polymers with green solvents rather than the generally used toxic solvents. However, the significantly lower number of published articles using greener alternatives decreases their visibility dramatically. In order to emphasize greener alternatives of the ES technique with the goal to reduce the environmental impact, we highlight the implications and strategies arising from using more environmentally friendly solvents for the production of electrospun fibers.

First, we briefly highlight the latest methods and concepts behind the definition of “greenness” regarding solvent choice, followed by a comprehensive overview over recent reports of electrospun fibers generated by the most commonly used water-insoluble polymers, based entirely on green solvents. In the subsequent chapter, water-based ES techniques as the greenest options are presented, based on versatile dispersion ES methodologies to further extend the available range of polymers and combinations with active compounds in green ES. Lastly, reactive ES is introduced, which aims at enhancing the

chemical inertness and mechanical stability of weak, water-soluble polymers by performing cross-linking steps during (*in situ*) or after (post) fiber generation for support structures such as used in tissue engineering. The final conclusion discusses limitations of green ES and offers a future perspective.

2. Electrospinning with green solvents

The solvent choice probably represents the most important parameter in ES, since only a limited range of solvents and compositions can dissolve the respective polymers and subsequently achieve fiber formation *via* ES. However, many polymers are generally difficult to dissolve and, therefore, require solvents with high solvation properties. These solvents are usually toxic (*e.g.* halogenated solvents), difficult to recycle and with high environmental impact (*e.g.* ionic liquids²⁰). Consequently, the solvent choice has a tremendous effect on the environmental impact, safety and overall sustainability of the ES process.

2.1 Definition of green solvents

The definition of green solvents expresses the goal to minimize the environmental impact resulting from the chemical production until the disposal of the solvents. The simple question “what is a green solvent?”²¹ cannot be answered so easily, as it requires many criteria to be considered. Fischer *et al.* addressed this question by assessing the substance-specific hazards by quantifying the emissions and resources used over the life cycle of the solvent. For each solvent, the EHS (environmental, health, and safety) indicator was calculated by adding the scores of water hazard, persistency, chronic toxicity, irritation, acute toxicity, reactivity, fire-, and explosion-hazard and



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filtration applications. His research focus lies on functional nanofiber membranes produced by pilot-scale needleless electrospinning, using biocompatible materials, sustainable solvents and functional additives.



release potential that characterize each compound. On the other hand, the life-cycle assessment (LCA) method quantifies the cumulative energy demand (CED) taking into account the production, usage, potential recycling, and disposal of the solvent. Taken together, CED and EHS indicators provide an estimation of how green a solvent is, where the lower the score, the greener or less problematic the solvent is (Fig. 1).

Nevertheless, both the EHS indicator and CED are only estimations and do not fully reflect objective values. Indeed, a solvent can be synthesized from different commodities or raw materials and can be disposed of or recycled in different ways. Therefore, the energy demand through these different steps can vary significantly, resulting in different CED values. Also, the EHS indicator can change if new insights into the EHS impact of the solvent are revealed. Indeed more recently, new regulations and restrictions entered into force: the Global Harmonized System (GHS) as applied by the European Classification, and the Labelling and Packaging (CLP) regulation.^{22,23} Therefore, this method requires the adjustment of EHS indicators in light of the introduction of the new regulations and restrictions.

Slater and Savelski from Rowan University developed a similar method for evaluating the greenness of a solvent.²⁴ Compared to the EHS method, it takes into account occupational health and environmental hazards (threshold limit value, biodegradation, ingestion toxicity, carcinogenicity, half-time, ozone depletion, global warming potential, smog formation, acidification, soil adsorption coefficient, bio concentration factor), as well as the amount of solvent used in a process. However, peroxide formation, flash point, and CED of the solvent were not used as selection parameters. Since the definition of “green” depends on the method and its solvent score, the threshold value defining the solvent “greenness” is not always defined. For instance, ethanol (EtOH) is considered green for both EHS and LCA methods, with both having low

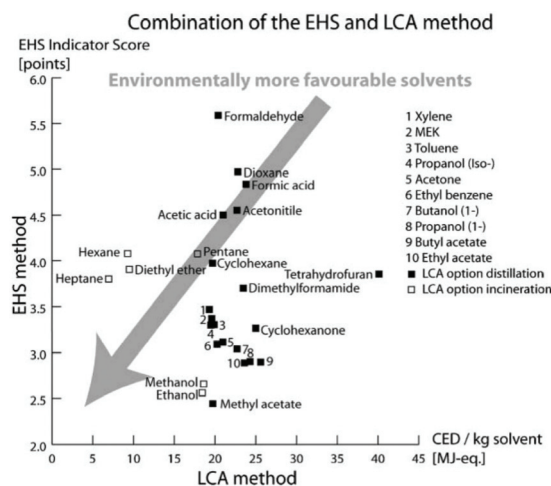


Fig. 1 Classification of 26 solvents by EHS and LCA methods representing the greenest solvents with the lowest scores. Reproduced from ref. 21 with permission from the Royal Society of Chemistry.

scores. Whereas the approach developed by Slater and Savelski can make a distinction between the “greenness” of the hydrocarbons, the EHS method considers the hydrocarbons nearly all as green.²³

In 2015, a public-private partnership European consortium comprised of pharmaceutical companies, universities, and small-medium enterprises, called the Innovative Medicines Initiative (IMI)-CHEM21, developed a ranking of the most common solvents to compare the “greenness” of processes or syntheses.^{25,26} Each solvent was classified according to benchmarks of existing guides, which were combined to limit their number to three safety, health and environmental criteria, each represented with scores from 1 to 10 and a color code (green for 1–3, yellow for 4–6, and red for 7–10). The final ranking of the solvent (recommended, problematic and hazar-



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dous) is usually derived from its least green characteristic, *i.e.* not from an average or summation of all properties, as performed in the EHS/LCA or Slater and Savelski methods. Moreover, bio-derived solvents (derived from sustainable and renewable resources) were evaluated accordingly, obtaining a final ranking. However, the environmental scoring of this method lacks the CED from LCA assessment or the carbon footprint that depends on the production process and usage of the solvent determined case by case by experts.

In the pharmaceutical sector, companies have focused on the safety and the health- and environmental-impact related to the practical use of the solvent. For instance, Pfizer developed a color-coded list of solvents using green, yellow, and red as flags for “Preferred”, “Usable” and “Undesirable”, respectively.²⁷ GlaxoSmithKline (GSK)^{28,29} and Sanofi³⁰ also used a visual “traffic light” code with tables displaying a list of solvents (Fig. 2). According to the solvent tables provided and in agreement with the EHS and Slater/Savelsky methods, the greenest solvents are water; C1–C4 alcohols, acids, ketones and esters; as well as 2-methyl tetrahydrofuran (2-MeTHF),

dimethyl sulfoxide (DMSO), acetonitrile and dimethylpropylene urea. Conclusions can also be drawn regarding the least green solvents, considered as undesirable if not already banned by REACH: chloroform, 1,2-dichloroethane, carbon tetrachloride, *N*-methyl pyrrolidone, dimethylformamide (DMF), dimethylacetamide (DMAc), benzene, hexane, 1,4-dioxane, 1,2-dimethoxyethane, diethyl ether, and 2-methoxyethanol.³¹ In 2016, GSK developed an updated solvent guide version²⁹ that retains the overall approach (color-code), but with additional 44 solvents since the last GSK guidance publication²⁵ updating the data behind all scores. All the solvents in the guide are scored assigning a value from 1 (the worst) to 10 (the best) to the following categories: incineration, recycling, biotreatment, and volatile organic compound (VOC) emissions related to the “waste” area; environmental air impact and aqueous impact; potential of exposure and health hazard; flammability & explosion potential, and reactivity & stability are related to the “safety” area; life cycle assessment. For each general area of assessment (waste, environment, health, and safety), an overall summary score (called compo-

Classification	Solvent Name	CAS Number	Composite Colour†	Boiling Point (°C)	Incineration	Recycling	Biotreatment	VOC Emissions	Aqueous Impact	Air Impact	Health Hazard	Exposure potential	Flammability & Explosion	Reactivity & Stability	Life Cycle Analysis‡
Water & Acids	Water	7732-18-5		100	4	2	4	6	10	8	10	9	8	10	10
	Acetic Acid	64-19-7		118	3	5	4	7	8	4	7	5	8	6	8
	Trifluoroacetic acid*	76-05-1		72	1	5	2	4	4	4	4	3	7	6	
Alcohols	1-Heptanol	111-70-6		178	9	8	10	9	8	4	10	7	9	10	
	Ethylene glycol	107-21-1		197	4	5	5	10	10	8	7	10	10	10	9
	1-Octanol	111-87-5		195	9	7	8	10	5	4	7	10	9	10	
	1-Butanol	71-36-3		118	6	7	5	8	9	3	7	7	8	9	5
	1-Propanol	71-23-8		97	5	3	3	6	10	4	10	7	8	10	7
	Ethanol	64-17-5		78	5	5	3	4	9	5	10	8	6	10	10
	2-Propanol	67-63-0		82	5	5	3	5	8	7	10	6	6	8	4
	t-Butanol	75-65-0		82	5	5	3	5	9	7	7	5	6	10	8
	IMS (ethanol, denatured)	64-17-5		78	5	5	3	5	9	5	4	7	6	10	
	Methanol	67-56-1		65	4	7	3	3	10	7	4	6	5	10	9
	Glycerol diacetate	111-55-7		187	5	6	6	10	6	8	10	8	10	10	
Esters	Isobutyl acetate	110-19-0		116	7	9	8	6	9	6	10	6	8	10	
	Isoamyl acetate	123-92-2		142	9	9	8	8	4	6	7	8	8	10	
	Isopropyl acetate	108-21-4		89	6	7	5	5	9	5	10	6	6	10	7
	Ethyl acetate	141-78-6		77	5	6	5	4	9	5	10	7	5	10	6
	Propylene carbonate*	108-32-7		242	4	5	6	10	10	10	10	10	10	10	
Carbonates	Diethyl carbonate*	105-58-8		126	7	9	9	7	9	8	4	5	8	10	
	Dimethyl carbonate	616-38-6		91	4	3	5	5	9	7	10	6	6	10	8
	Cyclopentanone	120-92-3		131	8	9	6	7	10	5	7	6	8	10	6
Ketones	Methylisobutyl ketone	108-10-1		117	7	8	5	7	9	3	7	6	7	9	2
	Methylethyl ketone	78-93-3		80	5	5	3	4	8	4	10	6	5	9	3
	Acetone	67-64-1		56	5	6	3	2	10	6	10	6	4	9	7
Aromatics	Anisole	100-66-3		154	8	8	8	8	7	6	7	8	7	9	5
	p-Xylene	106-42-3		138	10	9	6	7	5	2	7	7	5	10	7
	p-Cymene*	99-87-6		177	10	8	7	9	3	2	10	6	6	9	
	Toluene	108-88-3		111	10	7	6	7	7	2	7	6	5	10	7
	Trifluorotoluene	98-08-8		102	4	4	5	6	3	8	10	4	4	10	
	Pyridine	110-86-1		115	3	6	2	7	7	3	4	4	8	9	2
Hydrocarbons	Benzene	71-43-2		80	9	6	6	4	7	5	3	3	3	10	7
	Isocetane	540-84-1		99	10	4	5	6	2	5	10	7	3	10	7
	Heptane	142-82-5		98	10	4	5	6	3	5	10	6	3	10	7
	Cyclohexane	110-82-7		81	10	6	5	4	3	5	10	6	2	10	7
	Hexane	110-54-3		69	10	8	4	3	3	5	7	4	2	10	7
	Petroleum spirits	8032-32-4		55	8	9	4	2	5	5	1	6	2	10	7
Ethers	Dimethyl isosorbide*	5306-85-4		236	4	4	5	10	9	6	4	9	9	8	
	Cyclopentyl methyl ether	5614-37-9		106	8	4	5	6	4	3	4	4	6	9	4
	2-Methyltetrahydrofuran*	96-47-9		78	6	5	3	4	7	4	4	3	4	6	4
	t-Butylmethyl ether	1634-04-4		55	7	8	4	2	7	5	7	4	3	9	8
	Diisopropyl ether	108-20-3		68	9	7	6	3	5	4	10	6	4	3	9
	Tetrahydrofuran	109-99-9		65	5	5	2	3	9	3	7	5	4	6	4
	1,4-Dioxane	123-91-1		102	4	1	3	6	8	4	4	3	4	6	6
	Diethyl ether	60-29-7		35	7	7	3	1	5	3	10	4	2	6	6
	1,2-Dimethoxyethane	110-71-4		85	4	4	3	5	8	7	3	4	4	6	7
Dipolar Aprotics	Dimethyl sulphoxide	67-68-5		189	3	4	4	9	8	6	7	9	9	5	6
	Acetonitrile	75-05-8		82	3	5	1	4	10	8	7	5	6	10	4
	N-Methyl pyrrolidone	872-50-4		202	3	4	3	10	10	6	1	9	9	9	4
	N,N-Dimethylacetamide	127-19-5		165	3	6	3	9	10	6	1	7	9	9	2
	N,N-Dimethylformamide	68-12-2		153	3	6	3	8	10	4	1	6	9	9	7
Chlorinated	Dichloromethane	75-09-2		40	2	10	4	1	8	6	7	4	4	10	7
	1,2-Dichloroethane	107-06-2		84	2	7	5	5	9	7	1	2	5	10	7
	Chloroform	67-66-3		61	3	9	5	3	7	5	4	1	5	10	6
	Carbon tetrachloride	56-23-5		77	3	7	5	4	4	1	4	1	4	10	7

GSK Solvent Sustainability Guide



Column Headings Colour Key

Waste
Environment
Human Health
Safety

Composite Colour Key

Few Known Issues
Some Known Issues
Major Known Issues

*The scoring assessment for this solvent includes 4 or more data gaps, therefore there is a lower level of confidence in the solvent's placement on this guide.

*A blank value for Life Cycle Analysis (LCA) indicates that this data is currently not available.

*The composite colour represents an overall categorization of the holistic sustainability of a solvent, taking all category scores into consideration.

Fig. 2 The GSK solvent sustainability selection guide showing the 55 most commonly used solvents scored and color-coded based on their greenness. In the ESI of ref. 29 is reported a chart classifying 154 solvents. Reproduced from ref. 29 with permission from the Royal Society of Chemistry.



site score) is defined as the geometric mean of each of the relevant category scores:

$$\text{Composite} = \sqrt[4]{\text{waste} \times \text{environment} \times \text{health} \times \text{safety}}$$

The color assignment of each category is based on the score: green ($x > 7.5$), amber ($3.5 \leq x < 7.5$), and red ($x < 3.5$). Instead, the composite color assignment is made by taking into account possible data gaps in the classification and giving more importance to 4 out of 10 categories (*i.e.* VOC emissions, health hazard, flammability and explosion potential, and reactivity and stability) following a decision tree assignment diagram.²⁹

According to the GSK classification (Fig. 2), anisole is the greenest aromatic solvent; DMSO and dimethylpropylene urea are the greenest among the dipolar aprotic solvents (with only some issues related to the “waste” category); formic acid (FA) and acetic acid (AcOH) can be considered green with some issues related to the fact that they are acids; ketones and alcohols can be more or less considered green; and all the halogenated compounds are considered not green.²⁹

In 2018, Clarke *et al.* published a review focusing on ionic liquids, deep eutectic solvents, supercritical fluids, switchable solvents, liquid polymers, and renewable solvents.³² Plant biomass is considered a source of renewable solvents: crops such as corn and sugar cane, agricultural residues, and agroforestry products are all considered potential raw materials for the generation of fuels and solvents.³³ However, the food *versus* fuel debate limits or hinders the use of potential food as raw material for chemical production.³⁴ The reason why biofuel/chemical production from plants remains controversial lies on the competition between food and fuel production on limited resources (land, water, labor, and capital),³⁵ since both the food and biofuel originate from crops.³⁶ Instead, chemical production using agricultural residues, grasses, and agroforestry products (lignocellulosic biomass) or waste in general, provides a renewable resource without competing directly with food. However, processing of biomass waste requires usually more energy and cost. Glucose, for example, can be obtained from the hydrolysis of starch and cellulose as renewable sources that could be extracted from wood pulp (non-food source). Subsequently, transformations of glucose can provide bioderived solvents such as limonene, γ -valerolactone, and 2-methyl tetrahydrofuran (2-MeTHF). The latter is now being used as a renewable alternative to THF showing lower toxicity.³⁷ Moreover, 2-MeTHF can be obtained from furfural, which may be derived from agricultural waste.³⁸ Nevertheless, the CHEM21 methodology categorized this solvent as “problematic” because of its high flammability.^{25,26} Limonene (both enantiomers) is considered green with some issues (color code amber) according to the GSK chart²⁹ due to the low score in the “environmental” category. Meanwhile, γ -valerolactone has better scores compared to both limonene and 2-MeTHF. Nevertheless, there is a lack of data, especially in the “environmental” and “health” categories, thus score values have been estimated *via* calculation or comparison to structurally similar solvents.

2.2 Polymers electrospun with green solvents

Many reports on the ES of polymers from single polymer solutions using solely green solvents are already available in literature, but are still a minority in the ES field. In the following section, the most common polymers used for ES with green solvents are summarized (see Table 1). ES of water-soluble polymers is not reported here, since such fibers are not stable over time because of air humidity. Therefore, a cross-linking process is needed, which is elaborated on in Section 4 “Reactive Electrospinning”.

2.2.1 Cellulose derivatives. This class of polymers refers to chemically-modified cellulose. Since cellulose is an abundant biopolymer, it is widely used in fiber manufacturing. However, as cellulose is very stable and does not melt, cellulose has to be chemically modified *via* their hydroxyl groups on the cellulose backbone. The most common functional groups are acetate, nitrate, xanthate, carboxymethyl, methyl, ethyl and hydroxyethyl to yield their cellulose derivative forms.³⁹ Due to the chemical processing under either acidic or alkaline aqueous conditions, practically all cellulose derivatives are insoluble in harsh organic standard solvents and instead are soluble in green solvents, such as water, methanol, EtOH or acetone.³⁹

Cellulose acetate (CA, acetyl content 39.8%, MW = 30 000) was successfully electrospun using a mixture of AcOH and water by Han *et al.*⁴⁰ Bead-free uniform nanofibers were obtained at concentrations higher than 17 wt% in an AcOH–water mixture 75 : 25 (w/w). Recently, Majumder *et al.* reported a formation of uniform fibers with some beads using the same type of CA and polymer concentration used in the work of Han.⁴¹ Majumder *et al.* used different types of solvent mixtures, the green ones being pure acetone, and AcOH and water (3 : 1 w/w) producing cylindrical/ribbon-shaped fibers and uniform fibers with some beads, respectively. The presence of beads using acetic acid and water can be ascribed by the fact that the RH during ES was likely higher than Han's work (Majumder and Han did not report RH); indeed high humidity can cause the formation of beads due to lower evaporation rates.⁴²

Haas *et al.* dissolved cellulose acetate in a mixture of acetone : DMSO (2 : 1 v/v) and EtOH : DMSO (1 : 1 v/v). They showed that acetone allowed it to spin fibers without beads (18 wt% CA solution) thanks to the higher volatility compared to EtOH that caused the formation of beads. Their results on DMSO-based binary solvent mixtures revealed that the transition from spraying to spinning, starting from 12 wt%, can be shifted to lower concentrations (8–10 wt%) if EtOH is replaced by acetone.⁴³

Reports on the electrospinning of other cellulose derivatives are beyond the scope of this review and the reader is referred to recent detailed review articles.^{44–47}

2.2.2 Aliphatic polyesters. This class represents polymers that contain esters as functional groups in every repeating unit in their backbone. Most frequently used polyesters for ES are polycaprolactone (PCL), poly-lactic acid (PLA), poly(lactic-co-glycolic acid) (PLGA) and poly(glycolic acid) (PGA). Aside from



Table 1 List of the most common polymers that were electrospun recently using solely green solvents

Polymer	Solvent/solvent mixture	Electric field [kV cm ⁻¹]	Fiber morphology	Ref.
CA	AcOH : H ₂ O 75 : 25 (w/w)	1.2	Fibers	40
	AcOH : H ₂ O 75 : 25 (w/w)	1.7–2.0–2.5	Beads on fibers	41
	Acetone	1.7–2.0–2.5	Cylindrical + ribbon-shaped	41
	DMSO : acetone 2 : 1 (v/v)	0.73	Fibers (650 ± 130 nm)	43
PCL	AcOH	0.75–1.0	Microfibers	48
	AcOH : H ₂ O 90 : 10 (w/w)	0.75–1.0	Beads on fibers	48
	FA	0.75–1.0	Nanofibers (88 nm)	48
	AcOH : FA 3 : 1 (w/w)	0.96	Fibers (266 nm)	49
	Acetone	0.36–1.0	Fibers	50 and 51
	Acetone	1.33	Fibers (757 ± 275 nm)	53
PVDF	DMSO : acetone = 3 : 2 (v/v)	0.64	Fibers (625 ± 113 nm)	54
PA6	FA : water 85 : 15 (v/v)	2.1	Nanofibers	55
PA11	FA : water 98 : 2 (v/v)	1.5 up to 5.0	Fibers and nanofibers	56
	AcOH : FA 3 : 1 (v/v)	3.3	Fibers (124 nm)	57
	FA : Anisole 3 : 2 (v/v)	4.2	Nanofibers (53 ± 10 nm)	58
Zein	EtOH : water 8 : 2 (v/v)	1.7	Fibers (146 ± 33 nm) with some beads	60
	EtOH : water 8 : 2 (w/w)	1.4	Fibers (126 ± 31 nm)	61
CS	AcOH : water 9 : 1 (w/w)	1.1–4.0	Fibers (140 ± 51; 130 nm)	64 and 67
PS	D-Limonene	1	Fibers (700 nm)	68
	EtOAc	1.0 to 3.0	C-shaped and ribbon-like microfibers	69 and 70
	MEK	1.0 to 3.0	C-shaped and ribbon-like microfibers	69 and 70
PMMA	EtOAc	1.0–1.4–1.8	Ribbon-like microfibers (~6.5 μm)	71
	2-propanol : water 78 : 20 (w/w)	0.59	Fibers (770 ± 45)	72
PAN	DMSO (at 80 °C)	1.3	Fibers (~550 nm)	73
	DMSO (needleless set-up)	1.4–3.1	Fibers	74 and 75

DMF and chloroform as standard solvents, these materials can usually be dissolved in AcOH, FA, and acetone. Bahrami *et al.* tested different solvent mixtures comprising AcOH, AcOH : water, FA and FA : acetone (4 : 1) using different concentrations of PCL ($M_w = 80$ kDa).⁴⁸ PCL microfibers without beads are formed using AcOH with a concentration of 15 wt%, and nanofibers with 88 ± 25 nm resulted when 20 wt% PCL solution was electrospun at 15 kV with a tip-to-collector distance of 20 cm. Using FA:acetone and 90% AcOH mixtures only beads-on-fibers were obtained.⁴⁸ Schueren *et al.* successfully spun a 10 wt% PCL in AcOH : FA = 3 : 1 at a RH of 30% obtaining fibers with an average diameter of 266 ± 39 nm.⁴⁹ Reneker *et al.* spun PCL ($M_w = 80$ kDa) solution of 15 wt% in acetone.⁵⁰ Subsequently, Royen *et al.* produced fibers without beads from a solution of 15 wt% PCL ($M_w = 40$ kDa) in acetone with CTAB (15 mol%).⁵¹

Mosher *et al.* showed the production of a blend of PLGA/PCL with AcOH and non-green solvents (DMF/DCM) and compared the mechanical properties of the produced fibers between the green solvents and the traditional solvents for ES PLGA and PCL. The fibers were produced by applying a voltage of 0.67 kV cm^{-1} for an average fiber diameter of $1.27 \pm 0.23 \text{ μm}$. Fibers produced with AcOH resulted in a 47% increase in ductility accompanied by a 29% reduction in elastic modulus compared with the traditional ES mixture (Fig. 3).⁵² Recently, PCL was also blended at different ratios together with a citric acid-based biodegradable elastomer, poly (1,8-octanediol-*co*-citrate) (POC), and electrospun from mixtures of AcOH and FA. They obtained defect-free nanofibers with diameters of 0.61–1.11 μm with wettability- and mechanical properties that were dependent on the PCL/POC ratio, post-crosslinking treatment, and fiber orientation.

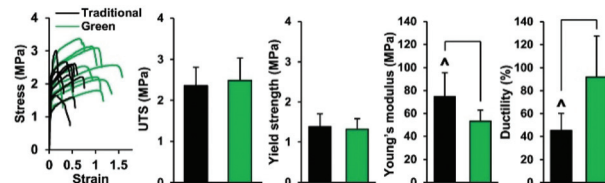


Fig. 3 Comparison of the mechanical properties of membranes electrospun using green and traditional solvents. They performed similarly in ultimate tensile strength (UTS) and yield strength, while the elastic modulus was lower and the ductility higher for the green solvents. Reproduced from ref. 274 with permission from IOP Publishing.

Finally, Casasola *et al.* were able to electrospin PLA in acetone (10% w/v). The PLA used was an amorphous polymer with an L-lactide content of around 88 wt%.⁵³

2.2.3 Fluorinated polymers. This class of polymers contains fluorine atoms on polymer backbones, which render the materials highly hydrophobic and chemically resistant. The most prominent types of fluorinated polymers that can be electrospun are poly(vinylidene fluoride) (PVDF) or its close analogue PVDF-hexafluoropropylene (PVDF-HFP). Mostly, PVDF is electrospun using pure DMF or DMF:acetone mixtures. However, DMF can be replaced by DMSO, as demonstrated recently with PVDF at 10 wt% using DMSO and acetone (3 : 2 v/v) as a greener solvent mixture.⁵⁴

2.2.4 Polyamides. Polyamides, or Nylons, can be electrospun using different green solvents, such as AcOH, FA and anisole, instead of the commonly employed chloroform or DMF. Supaphol *et al.* studied the effect of molecular weight and concentration of PA6, electrostatic field strength, and polarity on the average fiber diameter using FA : water (85 : 15 v/v) as solvent.⁵⁵



A parametric study was also performed using 85 and 98 vol% FA as solvent. The investigated parameters were polymer grade, viscosity of the solution, salt content and solvent grade, voltage, distance to the collector, nozzle size, and feeding pressure of the solution.⁵⁶

De Vrieze *et al.* performed a further parametric study by changing the ratio of AcOH and FA of different concentrations of PA6 of 10 kDa (from 12 up to 21 wt%). The study showed that a steady-state condition can be achieved using a solvent mixture from 33 to 50 vol% of AcOH. They also showed a reduction in the fiber diameter when the RH increased speculating that water acted as a plasticizer proven by the stable γ -crystals and a drop in the glass transition temperature.⁵⁷

PA11 was successfully electrospun using a mixture of FA:anisole 60:40 (v/v) by Meireman *et al.* obtaining nanofibers with an average diameter of 53 ± 10 nm.⁵⁸

2.2.5 Zein. Zein is a class of prolamine protein found in maize, thus being renewable and biodegradable. This material is clear, odorless, tasteless, hard, water-insoluble, edible, and it has a variety of industrial and food applications.

While Zein can be spun both from EtOH or AcOH,⁵⁹ 8:2 EtOH:water (w/w) solutions are most commonly employed,⁶⁰ as investigated comprehensively by Neo *et al.*⁶¹ The parameters studied were solution feed rate, applied voltage, tip-to-collector distance, and concentration. The authors reported that bead-less fibers were formed at 25, 30, and 35 wt% Zein solutions which are 2.1, 2.5, and 2.9 times the entanglement concentration of Zein, respectively. The results are in agreement with the observation of McKee *et al.*,^{62,63} who proposed that the entanglements between the polymers chains are enough to stabilize the jet and suppress the Raleigh instability, thus, promoting the formation of fibers rather than droplets during ES.

2.2.6 Chitosan (CS). CS is a polysaccharide, deacetylated derivative of Chitin, and is a biocompatible and biodegradable polymer mainly used in biomedical applications for its bacteriostatic properties. CS contains free amino groups that yield a positively charged polyelectrolyte and contribute to its solubility compared to chitin. It is therefore soluble in most acids, such as AcOH and FA, and also their blends with water. However, the resulting positive charges on the chitosan backbone results in repulsive forces between the polymer chains, leading to weak entanglement yet high viscosity.⁶⁴ Therefore, high acid concentrations are needed (70–90%) with low to medium molecular weights to generate pure chitosan fibers or, more commonly, polymeric blends with either hydrophilic polymers (polyethylene oxide (PEO), PVA) or hydrophobic polymers (PCL, Nylon) are used to decrease repulsion by hydrogen bonding.^{65,66}

Geng *et al.* electrospun CS fibers with an average diameter of 130 nm from aqueous 90% AcOH solution containing 7 wt% of polymer. The authors reported that the acid content should be higher than 30%, a requirement for CS fiber formation, because more concentrated acetic acid decreased surface tension of the CS solution and increased charge density. However, an AcOH solution at $\geq 90\%$ did not dissolve enough CS to reach a spinnable concentration. Instead, bead-free fibers were obtained from a molecular weight of 106 kDa,

while lower or higher molecular weights (30 and 398 kDa) showed the formation of beads. Average fiber diameters and size distributions decreased with increasing electric field and more bead defects appeared at 5 kV cm^{-1} or more.⁶⁷

Homayoni *et al.* used medium molecular weight CS of 1095 kDa and hydrolyzed it using NaOH with different treating times (from 0.75 to 48 h). The best result was obtained using 5wt% of the 48 h hydrolyzed CS (corresponding to a molecular weight of 294 kDa) in 90% AcOH. The hydrolyzed CS can also be spun in 80 and 70% AcOH producing bigger fibers.⁶⁴ In comparison with the work of Geng,⁶⁷ the applied electric field is much weaker (1.1 kV cm^{-1} instead of 4.0 kV cm^{-1}) and this can explain why the fibers are bead-less even though the molecular weight is higher.

2.2.7 Polystyrene (PS). Polystyrene, also known as the widely established Styrofoam, is a transparent, stiff material regularly employed in ES. It is soluble in a wide variety of solvents, where green solvents include methyl ethyl ketone (MEK), ethyl acetate (EtOAc), DMSO, acetone, and even natural oils, such as limonene. Shin *et al.* electrospun PS from recycled expanded PS and used the solvent α -limonene, which is a natural solvent much greener than the commonly used solvents for PS (such as THF, DMF, DMAc). The electrospun nanofiber diameter varied from 300 to 900 nm, with an average diameter of 700 nm at 30 wt% polymer concentration. α -limonene showed to be a good solvent for PS. However, the ES jet stability of PS was poor at low concentrations (below 10 wt%) compared to the other potential solvents.⁶⁸

Jarusuwannapoom *et al.* used PS solutions ($M_w = 299$ kDa, $M_n = 119$ kDa) in MEK and EtOAc.⁶⁹ EtOAc and MEK were both able to dissolve PS pellets within three and one day, respectively. Fibers and beads were observed for a 10% w/v solution of the polymer. SEM observations revealed that the fibers were c-shaped and ribbon-like. The authors stated that the occurrence of such appearance was a result of the low boiling point of the solvents (77.1°C and 79.6°C for EtOAc and MEK, respectively). Later, a similar paper was published by Manee-in *et al.*⁷⁰ In both papers, the same PS was used with the same M_w and polydispersity. However, in this work, the average fiber diameter and standard deviation of the fibers obtained from 10, 20, and 30% w/v in EtOAc and MEK are reported in the function of the change of the applied voltage and tip-to-collector distance with only EtOAc and MEK. The diameters of the bead-less-fibers ranged from 3.5 to $24 \mu\text{m}$.^{69,70}

2.2.8 Polymethylmethacrylate (PMMA). PMMA is a transparent thermoplastic frequently used as alternative to glass and due to its better environmental stability compared to *e.g.* polystyrene. While common ES solvents for PMMA include THF, DMF and chlorinated solvents, green alternatives are acetone, EtOAc or DMSO. Li *et al.* electrospun polymethylmethacrylate (PMMA, $M_w = 500$ kDa) from a polymer solution of 20 wt% in EtOAc.⁷¹ The surface of the fibers was porous due to the presence of water in the air (relative humidity (RH) $38 \pm 2\%$) that condensed on the “cold” surface of the polymer jet during the ES process, resulting in vapour-induced phase separation.



Chang *et al.* used 2-propanol:water 7.8:2 (w/w) mixture to spin PMMA with an average molecular weight of 400 kDa at a concentration of 1.5 wt%.⁷² Since RH was relatively high (45%), the formation of pores on the surface of the fibers was expected, but not documented due to the low SEM image resolution.

2.2.9 Polyacrylonitrile (PAN). PAN is a thermoplastic polymer also used as fibers in textile industry. However, the choice of green solvents is very limited to only DMSO instead of the more common solvents DMF and DMAc. Chen *et al.* e-spun PAN (80 kDa) from DMSO with acyclovir for drug release application using a syringe-needle set-up.⁷³ Due to its high boiling point (189 °C), the polymer solution in the syringe was heated to 80 °C to increase the vapor pressure of DMSO. The electrospun fibers (Fig. 4) agglutinated together when the spinning process was conducted at ambient temperature (24 °C). Conversely, at 80 °C the spinning solution became less viscous, DMSO evaporates faster, and the presence of acyclovir increased the solution conductivity. All these factors promoted the formation of bead-free fibers with a homogeneous structure, with a diameter ranging from ~400 to ~700 nm. Grothe *et al.* e-spun PAN in DMSO at 23 °C and 32% RH using a needleless ES set-up (Elmarco “Nanospider (NS) Lab”, Czech Republic).⁷⁴ The obtained fibers are not agglutinated even though DMSO was not heated during the e-spinning process. This is likely due to the higher electrode–substrate distance (24 cm instead of 12 cm used in Chen’s work⁷³), thus allowing the solvent to further evaporate enabling the formation of solid fibers once reaching the substrate. In another paper from the same research group, different concentrations of PAN in DMSO were spun.⁷⁵ At 12, 14, 16, and 18 wt%, the e-spun fibers were relatively regular in shape while increasing the solid content further to 20 wt% resulted in thicker fibers with agglutinations, which resembled a connected net. The behavior of forming net only at a higher concentration of the PAN in DMSO could be

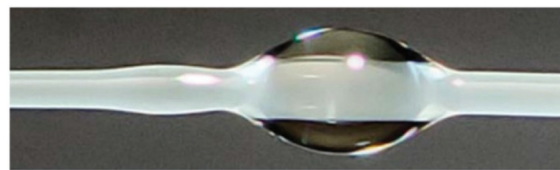


Fig. 5 Droplets of solvent on fiber spun from PAN in DMSO solution showing phase separation of the stretched polymer (in white) and the solvent (transparent droplet). Reproduced from ref. 76 with permission from Wiley-VCH Verlag GmbH & Co. KGaA.

explained by phase separation of the polymer jet under high mechanical stretching.⁷⁶ A concentrated polymer solution, under elongation, undergoes phase separation and the solvent remains on the surface of the fiber without completely evaporating (Fig. 5). This phenomenon is called “mechanotropic spinning”.⁷⁷ Kotomin *et al.* showed that the mechanotropic process has to be taken into account even for ES from polymer solutions.⁷⁶ Above all, mechanotropic spinning may be more reasonable for polymer solutions in solvents having low vapor pressures and high boiling points, like DMSO. Depending on the phase state of solution, the type of solvent, and stretching rate, both mechanisms (mechanotropic- and electro-spinning) may take place at the same time, thus causing agglutination/merging of the fibers once they are deposited onto the substrate.

3. Dispersion electrospinning

As biomolecules are most stable in aqueous environments, unfortunately, a variety of desirable active compounds is intolerant to even the majority of green solvents. Hence, a far safer and environmentally friendly approach can be found in choosing a predominantly aqueous ES medium; however, the selection of both water-soluble polymers and active compounds is limited. In order to bridge this gap, dispersion ES methodologies have been implemented in a compromise, which consists of either liquid-in-liquid (emulsion) or solid-in-liquid (suspension) approaches. While a comprehensive overview of the then-recent research into green ES was reported by Greiner and Agarwal in 2011,⁷⁸ significant progress has been made in the last decade.

3.1 Emulsion electrospinning

Emulsion ES is considered a scalable alternative to the classical blend and co-axial ES techniques, which aims at encapsulating hydrophilic and/or hydrophobic compounds within nanofibers to steer release rates (Fig. 6). Co-axial electrospinning with its channel-in-channel approach based on special needle architecture offers a stable homogeneous core-shell structure on a small scale, which is important for a range of applications as summarized elsewhere.^{79–81} In contrast, emulsion ES is in more demand as it provides scalability on both needle and needleless ES setups, and can be regarded as

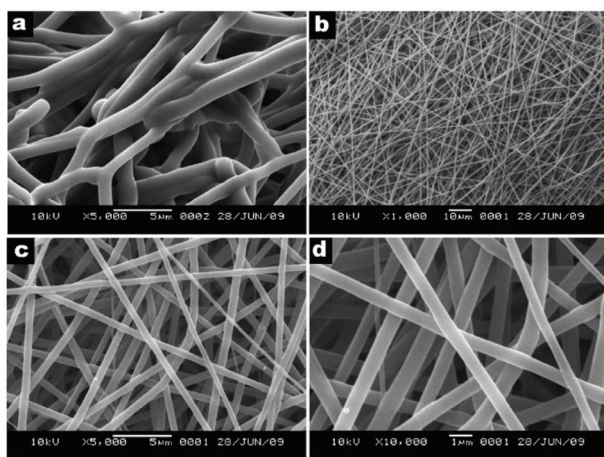


Fig. 4 SEM images of (a) PAN fibers electrospun at 24 °C, and (b–d) Acyclovir-loaded PAN fibers electrospun at 80 °C with different magnifications. Reproduced from ref. 73 with permission from Elsevier.



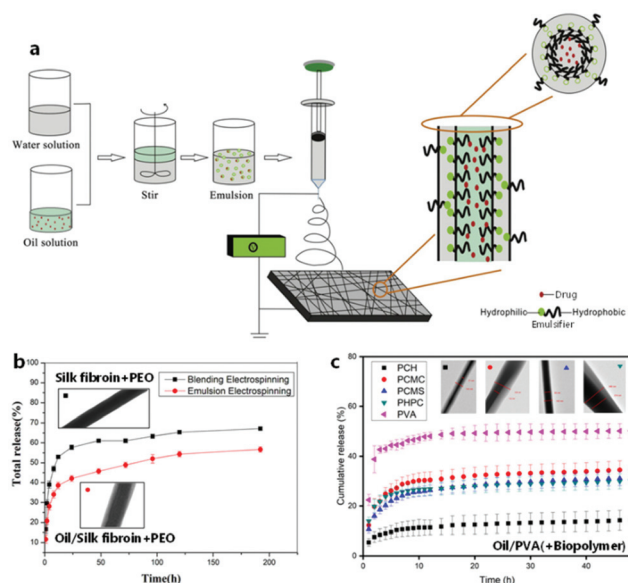


Fig. 6 Principle and effect of electrospinning from emulsion for encapsulation purposes. (a) Schematic process of emulsion electrospinning. (b) Typical release profiles from emulsion-spun and blend-spun fibers including respective TEM images. (a and b) Adapted from ref. 85 with permission from Elsevier B.V. (c) Release profiles depending on different sheath materials including respective TEM images. Adapted from ref. 109 with permission from Informa UK Limited.

a greener alternative to encapsulate hydrophobic moieties since it allows to use cheaper hydrophilic biopolymers in water as a continuous phase while avoiding or reducing organic solvents.⁸²

An emulsion constitutes a dispersion of two immiscible liquids of which one forms the continuous phase and the other a dispersed phase. By definition, emulsions are described as water-in-oil (W/O) or oil-in-water (O/W) if the water phase is the dispersed or the continuous phase, respectively. The insoluble hydrophobic phase is commonly described as the oil phase, which includes solutions based on solvents that do not necessarily constitute a viscous “oil”. Specifically, “green” oil phases commonly consist predominantly of natural oily extracts, whereas occasionally green solvents are used to facilitate phase separation. Nevertheless, “green” O/W emulsions are by far the most commonly electrospun green compositions compared to W/O emulsions. One reason for this is the fact that only few polymeric scaffold materials are soluble in natural oils alone.

In order to prevent phase separation in W/O and O/W emulsions, emulsifiers can be used, which are typically found in surfactants (e.g. Tweens, Spans), particles (e.g. calcium carbonate, silica), biopolymers (e.g. gelatin, alginate), and globular biopolymers (e.g. soy proteins, whey protein).⁸³ Therefore, the majority of modern approaches employs these naturally derived “green” biopolymers and minerals as stabilizers, which is additionally motivated by the desire to avoid potential mechanical weaknesses caused by the leaching of small-molecule surfactants.⁸⁴ While finding the right type and amount of

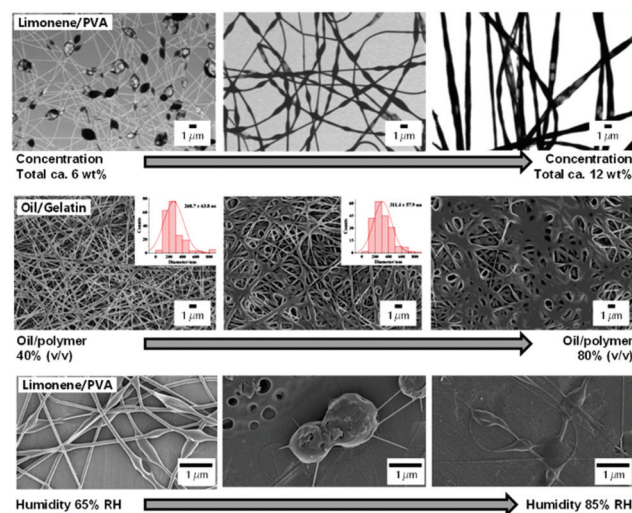


Fig. 7 Morphology of emulsion-electrospun fibers depending on emulsion composition and spinning parameters. Top: TEM images of fibers spun from emulsions containing different total matrix and oil concentrations. Adapted from ref. 97 with permission from Elsevier B. V. Middle: SEM images of fibers spun from emulsions containing different ratios of matrix to oil. Adapted from ref. 86 with permission from the American Chemical Society. Bottom: SEM images of fibers spun from an emulsion at different humidity levels. Adapted from ref. 98 with permission from Springer Science + Business Media New York.

emulsifiers to stabilize the emulsion is often a challenge in the ES of emulsions, the 2-phase composition gives rise to advantageous fibrous encapsulation structures. During fiber generation, a rapid viscosity increase occurs at the outer layer of the jet due to the higher evaporation rate of the solvent from the continuous phase gradually towards the core, displacing the dispersed second phase and effectively pushing it towards the center. The concentrated dispersed phase can then condense in the core continuously or in the shape of local reservoirs, which results in a core-sheath or beaded structure, respectively (Fig. 7).^{80,82}

3.1.2 O/W emulsion systems based on water-soluble biopolymers. A common goal of the O/W strategy is the effective encapsulation of oil-based active compounds for delayed-release within a water-soluble polymer template matrix. Biopolymers such as proteins and carbohydrates represent a particularly sustainable template material for green ES. While some oily active components can be used to form emulsions in water directly, carrier oils containing the dissolved ingredient can serve as a supporting mediator for improved phase separation and encapsulation efficiency. Making use of this methodology, Chen *et al.* used plant oil to dissolve their active anti-inflammatory compound, dexamethasone, and incorporate this into a silk fibroin and PEO sheath.⁸⁵ Stabilized by Tween 80, the electrospun emulsion yielded fibers with a well-defined core-sheath structure, which exhibited delayed-release of the active compound in comparison to fibers formed by a blend-approach without oil phase. Similarly, Zhang and co-workers used corn oil within a gelatin matrix from aqueous



acetic acid as a model to explore emulsifier-free spinning methods for food packaging and bioactive encapsulation.⁸⁶ Increasingly beaded and fused fibers were observed depending on the gelatin-oil ratio from 20% up to 80% (v/v); nevertheless, the oil load was maintained by the samples at 80% of the initial load and more during storage over several days. In a subsequent study, the addition of gum arabic into the initial emulsion formed bilayer emulsions and drastically enhanced the stability of the spun fibers by avoiding the formation of beads, owing to the intermolecular interactions between gelatin and gum arabic.⁸⁷ Also based on gelatin and a cross-linked derivative, Tavassoli-Kafrani *et al.* dispersed orange essential oil without any emulsifier in aqueous acetic acid solutions and spun to obtain smooth fibers, which encapsulated up to 70% of the oil content compared to the initial emulsion. The 80–200 nm thick fibers retained the volatile compound for up to 20 days during storage.

Focusing on carbohydrates instead, Tampau *et al.* used casein to stabilize and spin an acidic aqueous emulsion of carvacrol in starch.⁸⁸ Despite yielding only a small amount of well-defined fibers, which was attributed to insufficient chain entanglement, the authors achieved encapsulation efficiencies of up to 90%. In contrast, Krokida and coworkers observed smooth fibers from their emulsion-spinning of ulvan/pullulan with algae-extracts in coconut oil and Tween 20, while also encapsulating around 90% of the extract.⁸⁹ CS was employed in combination with PEO in acidic aqueous solution by Schiffman and coworkers in the surfactant-free spinning to encapsulate cinnamaldehyde.⁹⁰ The authors demonstrated the delayed release of the compound over several hours and improved antibacterial activity of the electrospun mats. In a follow-up study, the effect of molecular weight of CS, its degree of acetylation, and composition of the emulsion-components on the resulting fiber morphology was detailed.⁹¹ As expected, beaded or smooth fibers were obtained depending on the CS concentration and molecular weight, whereas increasing acetylation improved the incorporation of cinnamaldehyde while impairing the encapsulation efficiency of hydrocinnamyl alcohol. Similarly, Allafchian and coworkers conducted a comprehensive study on the composition and electro-spraying/spinning parameters of limonene and Tween 20 in *Alyssum homolocarpum* seed gum water solution.⁹² Originally focusing on spraying, their green approach also produced fibers as well as capsules with promising retention properties of limonene during storage over 90 days.

In contrast, Spano *et al.* demonstrated that the active ingredient does not always have to be incorporated into the core.⁹³ In their study, a solution of polypropylene carbonate in EtOAc was emulsified without surfactant in an aqueous blend of PEO and sericin protein. The aqueous continuous phase and hence resulting outer hydrophilic layer incorporated an antigen, inorganic fluorescent nanoparticles or magnetic nanorods *via* dispersion, whereas the subsequent release of the dispersed species over 72 h was demonstrated to depend on the exact ratio of polymeric compounds used.

3.1.2 O/W emulsion systems based on synthetic water-soluble polymers. While biopolymers are often preferred, particularly when moving towards functional fibers, the synthetic nature of water-soluble polymers such as polyvinylalcohol (PVA) and polyvinylpyrrolidone (PVP) is not considered a disadvantage. On the contrary, their well-known chemical and mechanical properties as well as spinning parameters facilitate the focus of the research on the added components. Targeting thermo-regulating textiles, Lin *et al.* achieved uniform core-sheath fibers from emulsions containing PVA and different amounts of plant oil.⁹⁴ With 7 wt% PVA, the fiber diameters ranged from 300 to 900 nm, whereas formulations with 9 wt% PVA showed a patchy structure and signs of oil leakage. Similarly, Zhou *et al.* obtained smooth core-sheath nanofibers from an emulsion containing dodecanol dodecanoate and roughly 10 wt% PVA crosslinked with glutaraldehyde.⁹⁵ As expected, the cross-linked material persisted longer for up to 10 min when exposed to water, which significantly improves the applicability compared to rapidly disintegrating PVA alone. Focused purely on the principle of reversible preservation of substances, the Crespy group electrospun peppermint oil in PVA or PVA/carboxymethyl-cellulose (CMC).⁹⁶ Subsequently, dissolution of the 300 to 1300 nm thick fibers confirmed 85–94% efficient recovery of the oil.

Despite their synthetic origin, many studies considered PVA for delayed release of oils for food and other storage applications, as a consequence of their controllable well-described properties and biocompatibility. Camerlo *et al.* demonstrated the potential of encapsulating limonene in PVA without surfactant.^{97,98} Fiber morphology shifted from beaded to uniform fibers from 5–12 wt% PVA concentration, whereas oil concentration only affected the number of beads. During storage, a slight loss of 20–35% of the volatile compound was recorded, unless conditions harsher than 25 °C and 60% RH were chosen. The effect of the humidity during the spinning was further investigated and revealed nanostructured beads at 7 wt% PVA concentration and a shift from smooth fibers to increasingly bigger beaded fibers with increasing moisture at 9 wt% PVA concentration. Similarly, Ciera *et al.* observed beaded structures as they incorporated different mosquito-repellant oils into an aqueous PVA solution and subsequently proved the repellent effect of the electrospun mat on insects.⁹⁹ Cinnamon oil as the antifungal and acaricidal active component of choice was incorporated into an aqueous PVA solution with the help of Tween 80 by Lee and coworkers.¹⁰⁰ In a comprehensive biological study, the authors established the inhibiting effect of the 250–500 nm thick and up to 30 wt% oil-containing fibers on microbes, bacteria, and fungi. Moreover, they used the analog system with phytoncide and palmarosa oil coupled with heat treatment at 170 °C for PVA layer stabilization in order to achieve both antimicrobial and water absorption capabilities, which promises potential wound treatment applications.¹⁰¹ Tampau *et al.* demonstrated the incorporation of carvacrol, both with and without Tween 85 as surfactant, into a beaded fiber structure with 75% encapsulation efficiency and higher.¹⁰² In fact, the authors suggested



that the surfactant diminished the bonding between carvacrol and the PVA matrix by competitive micelle formation, which led to a fractional release of the compound before and after pyrolysis of the PVA occurred. Biological surfactants were used by García-Moreno *et al.* to stabilize an emulsion of fish oil in acidified aqueous PVA solution to obtain smooth or beaded fibers, depending on the polymer concentration.¹⁰³

Considering the cost and efficiency of certain pharmaceuticals, carrier oils can also serve as a solvent in biological applications. Aiming for the oral delivery of the antimycotic clotrimazole, Opanasopit and coworkers employed a complex system of oleic acid as primary oil phase, as well as Tween 80 and co-surfactants, in an aqueous template of CS-EDTA (chitosan-ethylenediamine tetraacetic acid) and PVA.¹⁰⁴ Sunflower oil and Phospholipon 90 H were used by Daniels and coworkers to dissolve birch extract, which was shown to achieve prolonged release and wound healing capabilities.¹⁰⁵ Targeting antioxidant coatings instead, López-Rubio and coworkers dissolved carotene in soybean oil, which was emulsified with soy protein isolate in an aqueous PVA solution.¹⁰⁶ Expectedly, the emulsifier facilitated the formation of more stable emulsions as well as beaded fiber structures after spinning, which were annealed to remove residual incorporated water and achieve delayed gradual release. The annealing methodology was also employed by Meera Moydeen *et al.* in the preparation of core-shell PVA/dextran nanofibers, which incorporated a commonly used antibiotic.¹⁰⁷ Plant oil served as a carrier for this active substance and resisted the subsequent heat treatment at 120 °C, which was carried out to evaporate traces of water for increased intermolecular hydrogen-bonding between PVA/dextran molecules and hence improve mechanical stability of these fibers. Comparative release studies from coaxially and blend-spun nanofibers revealed a rapid release of a water-soluble ciprofloxacin salt, while the emulsion-spun antibiotic in oil exhibited efficient encapsulation with only minor release after 50 h. Subsequently, the authors further investigated the effect of the dextran content on the release kinetics using PVP as the main matrix polymer¹⁰⁸ and other polysaccharides¹⁰⁹ within an analogous system. Dextran content promoted intermolecular hydrogen bonding, thus affecting the viscosity and erosion of the polymer matrix.

Sesli Cetin and coworkers chose a system based on PVP for the encapsulation of antibiotic essential oils. Emulsions of 1–4 wt% cinnamon oil were stabilized with Cremophor RH 40 as a surfactant and successfully electrospun, although encapsulation efficiency decreased with increasing oil content.¹¹⁰ As expected, biological studies confirmed the antibacterial effect of the electrospun mat starting from concentrations of 2 wt% of essential oil after 24 h. In a follow-up study using thyme essential oil as active component and gelatin/PVP for improved durability, a similar spinning behavior was observed.¹¹¹ Fiber morphology changed from beaded to smooth fibers with increasing oil content; however, a concentration of 5 wt% and more led to issues in spinnability and stability of the membranes. The optimal membrane of this study contained 3 wt%

of the active component, showing antibacterial properties for up to 8 days.

Despite these promising properties, water-soluble sheath materials spun from an aqueous continuous phase can achieve only limited stability in hydrophilic media, which are used in biomedicine and the food industry. Furthermore, the application of O/W systems containing carrier oils requires that the application is able to tolerate the excess oily medium, which can be a considerable disadvantage.

3.1.3 W/O and other emulsion systems. Generally, a more promising green route to enhancing sheath stability for further delaying the release in aqueous environments leads *via* the use of water-insoluble degradable polymer matrices in W/O and related systems. This method has shown significant potential for targeted and delayed drug release although only applicable to water-soluble loads. Yue and coworkers recently demonstrated the encapsulation of aqueous bovine serum albumin (water phase) within a polystyrene (PS) solution in limonene (oil phase) as a green W/O example of emulsion ES.¹¹² Their work showcased the ability to tailor the fiber thickness proportionally to the chain length of the employed PS template, from around 0.6 μm up to 1.4 μm for molecular weights of 75 kDa to 350 kDa, respectively. Furthermore, an inverse yet almost linear release behavior in PBS solution was observed in relation to molecular weight, with the highest chain length achieving almost complete release after 50 days, whereas only 50% was released from the short-chain matrix after the same duration. The authors attributed this finding to the decreased entanglement derived from the more viscous solutions and hence higher porosity of the resulting fibers (with increasing molecular weight). An alternative approach to achieve a two-phase system with water can be found in the dissolution of a polymeric component in an otherwise water-miscible medium, which subsequently leads to separation from an added aqueous phase. Ferrari and coworkers utilized this method by dissolving PCL in a 3:1 mixture of acetic and formic acid and adding an aqueous PVA solution, which incorporated strontium ranelate as an osteogenic agent.¹¹³ The spun fibers were subsequently coated with gelatin for enhanced hydrophilicity before characterization, which revealed that a higher content of strontium ranelate increased elasticity, water uptake, and stimulation of osteogenic growth.

Yet another alternative, a “water in water” emulsion, can be produced by dissolving two incompatible but both water-soluble components simultaneously, which subsequently separate into two or more phases. This approach was used by Fuenmayor and coworkers to spin an emulsion consisting of both limonene-loaded cyclodextrane and pullulan in water to achieve prolonged storability of the volatile compound.¹¹⁴

In contrast, a mostly water-free emulsion approach was chosen by De Clerck and coworkers to form gellable nanofibers, in which PCL and gelatin were dissolved in a mixture of acetic and formic acid.¹¹⁵ Interestingly, this composition resulted in an emulsion forming separate PCL- and gelatin-rich phases. Different fiber diameters of 140–550 nm were achieved *via* manipulation of the polymeric content, while



optimizing the solvent ratio or addition of small quantities of water led to the formation of homogenous blend solutions instead. With an entirely different application in mind, Yu and coworkers employed a template system consisting of PVP in ethanol and acetic acid to emulsify a metalorganic titanium precursor.¹¹⁶ After spinning the sol emulsion and annealing at 500 °C, smooth TiO₂ nanofibers with a diameter of roughly 300 nm were obtained, which were ground up to nanorods for incorporation into a solar cell assembly.

Despite such achievements using emulsion systems however, clearly the more direct and convenient route for the incorporation of solid active compounds into nanofibers leads *via* the direct spinning from suspension.

3.2 Suspension electrospinning

Solid-in-liquid dispersions, termed either “suspension” or “sol” depending on particle size and visual appearance, can be electrospun similarly to emulsions in a wide range of compositions. In most cases, water-soluble matrix polymers and surfactants are dissolved in an aqueous continuous phase, to which the powdered solid particles or pre-made dispersions are added. This technique is compatible with a range of different particle types from biology to catalysis and can simply be upscaled using wire ES (Fig. 8); in fact, unstable dispersions can often lead to clogging in needle-based setups. An exception to this general procedure is only found in rather specific techniques, such as the direct near-field electrospinning (NFES) of aqueous-dispersed bacteriophages without any template polymer by Sugimoto *et al.*¹¹⁷

3.2.1 Aqueous suspensions involving a biopolymer matrix. Biopolymers present with a particularly green potential and have hence been increasingly employed as matrix polymers in suspension spinning. Gelatin fibers incorporating graphene

oxide were spun by Nirmala and coworkers, showing that graphene oxide significantly improved the mechanical strength of the still cytocompatible gelatin fibers mats for wound dressing applications.¹¹⁸ Deng *et al.* chose gelatin, which was glycation-crosslinked and contained hydroxyapatite nano-particles, to reduce water-permeability for food packaging applications.¹¹⁹ A comparison between whey protein and pullulan as encapsulation for bacteria was published by López-Rubio and coworkers, which could be applied for edible probiotics.¹⁰⁶ While the carbohydrate yielded a more fibrous structure, capsules obtained from the protein provided better cell viability. Combining both classes of biopolymers, Weiss and coworkers encapsulated nutritious pea protein isolate in maltodextrin and achieved partial crosslinking *via* heat-induced glycation.¹²⁰ Crespy and coworkers employed silica nanocapsules dispersed in aqueous dextran solutions to showcase the possibility of efficient reversible encapsulation.⁹⁶ Furthermore, cellulose-derivatives were reinforced with wood pulp¹²¹ and cellulose for aligned fibers.¹²² For the very specific purpose as a propellant, Li *et al.* dissolved nitrocellulose in aqueous acetone with dispersed boron particles followed by ES.¹²³ This necessity for co-solvents highlights the solubility limitations that many “water-soluble” biopolymers still face, which explains the predominant use of synthetic polymers in the spinning of aqueous suspensions.

3.2.2 Aqueous suspensions involving a synthetic polymer matrix. Dispersions in water predominantly employ PVA as template polymer. Several groups published rather fundamental studies describing how the bead size of dispersed polymer particles and the PVA matrix concentration affected the morphology and porosity of resulting electrospun fibers. The incorporation of PS particles with increased particle size led to blackberry-like structures (see Fig. 9a),¹²⁴ whereas the incorporation of silicate particles and subsequent calcination gener-

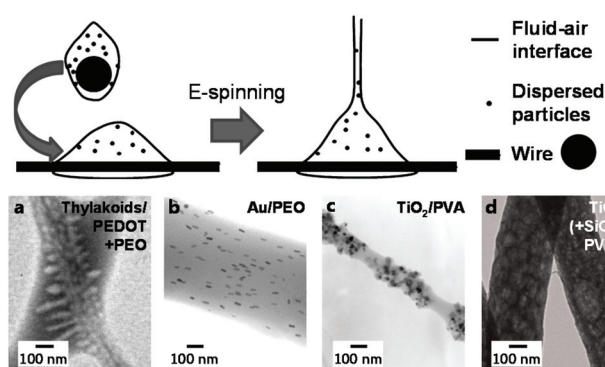


Fig. 8 Principle and morphology of suspension-electrospun fibers depending on particle type. Top: schematic process of wire electrospinning using dispersions. Adapted from ref. 164 with permission from the American Chemical Society. Bottom: selected TEM images of fibers spun from suspensions incorporating different types of particles (a–c) and after their removal (d). (a) Adapted from ref. 149 with permission from the American Chemical Society. (b) Adapted from ref. 152 with permission from the American Chemical Society. (c) Adapted from ref. 134 with permission from Springer Science + Business Media New York. (d) Adapted from ref. 167 with permission from Elsevier B. V.

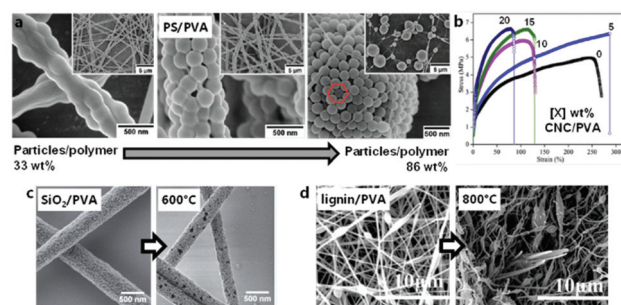


Fig. 9 Morphological and mechanical effect of particle concentration and size in suspension-electrospun fibers. (a) SEM images of electrospun fibers obtained from dispersions with increasing concentration of spherical particles. (b) Typical strain-stress curves of electrospun fibrous mats obtained from suspensions containing increasing amounts of particles. (c and d) SEM images of typical electrospun fibers obtained from dispersions containing nanoparticles (c) or microparticles (d). (a) Adapted from ref. 124 with permission from the American Chemical Society. (b) Adapted from ref. 141 with permission from Elsevier B. V. (c) Adapted from ref. 125 with permission from Wiley-VCH Verlag GmbH & Co. KGaA. (d) Adapted from ref. 144 with permission from Elsevier B. V.



ated porous structures (Fig. 9c).¹²⁵ Recently, Gonzalez *et al.* published a comprehensive study investigating compositional parameters, such as concentration and particle size of dispersed poly(methyl methacrylate-*co*-butyl acrylate) particles in the aqueous PVA solution, on the resulting fiber morphology.¹²⁶ Meanwhile, more application-focused studies have investigated the incorporation of Ag particles for their antibacterial properties for wound-healing applications. Encapsulation by ES of PVA solutions combined with subsequent crosslinking *via* glutaraldehyde (as opposed to reactive ES, see section 4.2.1) provides improved cytocompatibility and resistance to water, as demonstrated by Sun and coworkers.¹²⁷ This system was comprehensively studied by Augustine *et al.*,¹²⁸ who focused on plant-sourced Ag, and Sethuram *et al.* with an Eugenol-stabilized dispersion.¹²⁹ The higher tensile strength of the crosslinked PVA system has also been exploited in combination with multi-walled carbon nanotubes and graphene^{130,131} or with graphene oxide and a TiO₂ coating¹³² for reinforced conductive mats. Alternatively, Wei and coworkers dispersed carbon-based quantum dots for simultaneous fluorescent and electrochemical detection of peroxides and glucose.¹³³ The PVA encapsulation of metal oxides has been investigated in several studies to exploit their catalytic activities, such as the photocatalytic degradation of a dye,¹³⁴ Mn₃O₄ for the peroxide-induced oxidation of alcohols,¹³⁵ as well as calcinated fibers containing LiCoO₂ for electrode materials.¹³⁶ Focusing primarily on the stability and fluid elasticity of the suspensions, Hua and coworkers demonstrated the spinning of aluminosilicate nanotubes in a PVA matrix.¹³⁷ Other promising inorganic compounds for PVA-based suspension spinning include SiC¹³⁸ and boron nitride,¹³⁹ which produced fibrous mats with high tensile strength and thermal conductivity. Incorporation of cellulose nanofibers or cellulose nanocrystals represent greener approaches to reinforce PVA fibers for improved tensile properties (Fig. 9b).^{140,141} Furthermore, the spinning of PVA/Aloe Vera extract and dispersed CS nanoparticles can cause increasingly fused fibers and stronger films.¹⁴² In contrast, citrate-crosslinked CS was used to encapsulate cabreuva essential oil and in turn was spun from suspension by Pinotti and coworkers to manufacture beaded fibers with both antibacterial and gradual release properties.¹⁴³ An interesting alternative function was found in the dye-interaction of charged PVA fibers. Zhang *et al.* employed lignin suspended in aqueous PVA to reversibly adsorb cationic dyes onto the calcinated PVA fibers under different conditions (Fig. 9d).¹⁴⁴ Similarly, the Greiner group synthesized positively or negatively charged polyacrylate particles, which they incorporated into the PVA fibers to achieve the charge-specific dyeing of the mats depending on the dye used.¹⁴⁵

If, however, an even more water-soluble template polymer for suspension ES than PVA was required, PEO has frequently been chosen as an alternative. Colín-Orozco *et al.* incorporated water-insoluble whey protein isolate and rosemary extract into smooth PEO fibers for gradual complete release over just several hours.¹⁴⁶ A two polyester-polyether block-copolymer

was also reported on, which required no surfactants to stay dispersed and yielded slightly fused fibrous mats after spinning and re-dissolution of the PEO matrix.¹⁴⁷ Cellulose nanofibers, a polythiophene and an epoxy-crosslinker were added by Latonen *et al.* to a PEO solution to produce conductive biocompatible fibers.¹⁴⁸ A similar system was previously reported, which showcased successfully encapsulated intact thylakoid vesicles and correlated photo-induced currents and pH-changes.¹⁴⁹ This approach required EtOH as co-solvent, which was also the case in the study concerning dispersion-spinning of thermos-responsive polyacrylamide(PAAm)/CS microgels by Borges and coworkers.¹⁵⁰ Similarly, Mihindukulasuriya and Lim required this solvent system to incorporate TiO₂ particles, methylene orange, and glycerol into PEO fibers for UV-activated oxygen sensors.¹⁵¹ Other structurally interesting compositions on the other hand did not require a co-solvent. For example, internal alignment of incorporated gold nanorods was achieved simply by influencing the spinning conditions,¹⁵² while silicate platelets were incorporated into a PEO polyacrylic acid (PAA) template in a well-dispersed manner despite the stark size difference.¹⁵³ Interestingly, PAA represents another suitable matrix polymer by itself; however, it almost always requires an alcoholic co-solvent. Lei and coworkers spun a dispersion of boron nitride followed by thermal crosslinking to achieve stable thermally conductive mats for high-temperature thermoregulation.¹⁵⁴ In contrast, catalytic applications were targeted with the surfactant-aided encapsulation of Pt by Sightler *et al.*¹⁵⁵ and further improved by Kayarkatte *et al.*,¹⁵⁶ who added carbon particles to improve conductivity and prevent catalyst erosion at the same time.

Finally, water-soluble PVP has also been widely employed for suspension ES. Rare uses of aqueous spinning media include a study concerning the incorporation of poly(methyl methacrylate)-polyethyleneimine (PMMA-PEI) capsules in PVP fibers,¹⁵⁷ as well as the encapsulation of zeolites and subsequent calcination for hydrocracking catalysts.¹⁵⁸ Similarly, fibrous alumina constructs were produced for high-temperature filtration or catalysis.¹⁵⁹ Furthermore, *in situ* laser ablation enabled Uyar and coworkers to incorporate gold particles in a PVP matrix without the use of any harmful solvents whatsoever.¹⁶⁰

Only a small number of studies investigated the dispersion spinning of any other polymers from aqueous solutions, such as the incorporation of live yeast from an aqueous EtOH PAAm solution by Fan *et al.*¹⁶¹ In contrast, most other potential matrix polymers, as well as further compositions involving the above-mentioned polymers, require non-aqueous media.

3.2.3 Non-aqueous suspensions involving a synthetic polymer matrix. Virtually all PVP formulations used in suspension electrospinning are based on EtOH. This fact can be used to avoid dissolution of sensitive solid components, which are soluble in water but not EtOH, such as thermo-responsive PAAm microgels.¹⁶² Similarly, drugs like albendazole or famotidine are commonly obtained in amorphous form after spinning and crystallize over time, which can lead to physical instability of the fibers but can be prevented by dispersion



ES.¹⁶³ In contrast, surface tension can also be a deciding factor for spinning from non-aqueous suspension, which was shown in a report detailing the ES of PS and Pb particles in a PVP matrix by the same authors.¹⁶⁴ The majority of ES suspensions in non-aqueous solvents with PVP, however, is based on readily hydrolyzable organic metal-oxide precursors. For example, silicates and subsequent calcination were used to generate core-sheath-like fibers as porous ceramics.^{165,166} An interesting variation was reported by Lim and coworkers, who incorporated SiO₂ particles and a TiO₂ precursor in PVP, calcinated, and subsequently removed silicates *via* alkaline treatment to obtain porous hollow TiO₂ fibers.¹⁶⁷ Similarly, nanotubular titanate acid was suspended in a PVP solution by Gong and coworkers to ultimately obtain porous titanium nitride fibers with impressive microwave-absorption properties.¹⁶⁸ The effects of incorporated SiO₂ and hematite nanoparticles on PVP mats without calcination were separately reported by Matysiak and coworkers.^{169,170} Suspension-ES of SiO₂ particles in PVP using EtOH was also used to obtain water-resistant and flame-retardant mats (after annealing).¹⁷¹

Catalytic systems based on metal oxides and others were also obtained from green ES of entirely water-insoluble systems. Augustine *et al.* described the non-inflammatory peroxide-mediated wound-healing properties of a ZnO/PCL mat, which was obtained using acetone as dispersion medium.¹⁷² Acetic acid served as a solvent in the ES of a PCL/gelatin/graphene oxide system, which showed antibacterial behavior combined with cytocompatibility, accelerated degradation, and gradual drug release capabilities.¹⁷³

Other notable bioactive systems can be obtained using polyamides (PA, Nylon), for which FA is a green solvent. Ryu *et al.* prepared an antimicrobial and photocatalytic mat incorporating Ag-TiO₂ nanoparticles; however, they found that a PA mat with sprayed-on particles exhibited superior properties.¹⁷⁴ A similar composition was chosen by Chen and coworkers, who dispersed carbon nanotubes and generated Ag particles *in situ* to obtain conductive antibacterial mats.¹⁷⁵ The effect of the carbon nanotube concentration on the conductivity of PA fibers was further investigated by Pan and coworkers from dispersions in FA.¹⁷⁶ Meanwhile, Rahaman and coworkers combined acetic and formic acid as mixed medium and demonstrated the superior filtration and water-permeation capabilities of a PA membrane in contrast to common filter materials, whereby the PA membrane incorporated SiO₂ nanoparticles and was coated with polyvinylacetate (PVAc) for increased hydrophobicity.¹⁷⁷

Water-insoluble polymers such as PCL and PA significantly broaden the scope of potential applications for ES from dispersions. Nevertheless, even such water-resistant alternatives show degradation over time. Hence, even more durable formulations for more demanding purposes are highly desirable.

3.2.4 PTFE dispersions. Polytetrafluoroethylene (PTFE) is one of the most popular materials when exposure to aggressive chemicals or even environmental conditions is expected, thanks to its inherent inertness. Aqueous dispersions of PTFE nanospheres are broadly available and often marketed as

emulsions. However, the already solid nature of the particles following emulsion polymerization suggests that a suspension or colloidal dispersion would be a more accurate description of such products. Suitable surfactants do not necessarily contain fluorides and only constitute a fraction of such dispersions; hence, a section of this review should be dedicated to green approaches of PTFE spinning.

PTFE dispersions are commonly used in combination with a supporting template polymer for the ES of functional membranes directly, or further sintered into fibers above the PTFE melting point (approximately 330 °C). The particularly practical property of PTFE to maintain structural integrity under heat is based on its exceedingly high melt viscosity.¹⁷⁸ Despite the simplicity of the system, particle concentration and sintering conditions can significantly influence the resulting membrane morphology (Fig. 10).

PEO constitutes one of the more common matrix polymers for PTFE dispersion spinning. Feng *et al.* demonstrated that this system leads to remarkably acid- and base-resistant membranes after sintering, which was proven by unchanged tensile strength in such media.¹⁷⁹ The addition of polyamide-imide (PAI) further increased the tensile strength while diminishing elasticity.¹⁸⁰ Other important applications of PTFE fibers include triboelectric nano-generators,¹⁸¹ as well as membranes for oil/water separation.¹⁸²

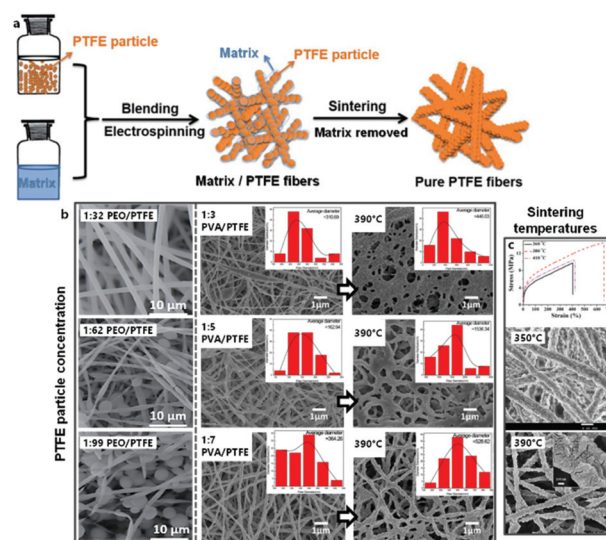


Fig. 10 Principle and morphology of the electrospinning of blends based on PTFE suspensions. (a) Schematic process of electrospinning PTFE suspensions for pure PTFE fibers. (b) Left: SEM images of typical fibers obtained from electrospinning PTFE suspensions at different ratios of matrix to PTFE. (b) Right: SEM images of typical fibers obtained from electrospinning PTFE suspensions at different ratios of matrix to PTFE and subsequent calcination. (c) Top: Typical strain-stress curves of electrospun PTFE-fiber membranes obtained after calcination at different temperatures. (c) Bottom: SEM images of typical electrospun PTFE-fiber membranes obtained after calcination at different temperatures. (a) and (b) Left, (c) top adapted from ref. 179 with permission from the Royal Society of Chemistry. (b) Right adapted from ref. 186 with permission from Springer Nature. (c) Bottom adapted from ref. 184 with permission from Elsevier B. V.



Alternatively, Kolesnik *et al.* reported a procedure for ES aqueous dispersions of PTFE using PVA as water-soluble sacrificial matrix polymer.¹⁸³ The optimal sintering temperature and tensile strength of the resulting nanofibers from this system were detailed by Zhou *et al.*,¹⁸⁴ while Zhu *et al.* determined that the aerosol filtration efficiencies of these membranes matched those of commercial PTFE filters.¹⁸⁵ It was further demonstrated that the oil/water separation properties of an analogous PTFE mat could not be further improved by the incorporation of ZnO particles, however in contrast this study found that the addition of small amounts of boric acid lowered the concentration of PVA needed to form smooth fibers.¹⁸⁶ Huang and coworkers subsequently employed the same system in near-field-ES (NFES) from dispersion to 3D-print micro-meshes for effective aqueous SiO₂ particle filtration.¹⁸⁷

While PTFE systems can never be considered truly “green” due to the environmental concerns of the persistent nature of fluoropolymers and their toxic pyrolysis products,¹⁸⁸ the above-mentioned approaches can contribute to a greener manufacturing process of this still unrivalled material. Even further optimization in “greenness” is certainly under constant development, such as the focus on biopolymers as template materials. An example was recently demonstrated by Pang *et al.* with the ES and subsequent fabrication of hollow PTFE microfibers based on a pullulan template.¹⁸⁹

4. Reactive electrospinning

One of the fields with the highest need for biocompatible materials concerns biomedical applications. Specifically, the formation of tissue produced by implanted cells is influenced greatly by the scaffold onto which they are seeded. It is often preferable to use a biodegradable material scaffold in order to naturally degrade the implanted materials and leaving only the generated tissue. Usually, the common materials used in this field are synthetic and natural biodegradable polymers. Among them, hydrogels are the most promising due to their ability to retain a great amount of water, good biocompatibility, low interfacial tension, and minimal mechanical and frictional irritation upon mechanical strains. Therefore, hydrogels and other soft composites more effectively promote cell expansion and tissue formation.¹⁹⁰ The most logical option is the use of polymers that are already present in animal or human tissues, such as collagen¹⁹¹ and its hydrolyzed form gelatin,¹⁹² natural and modified polysaccharides,¹⁹³ proteins,¹⁹⁴ and glycosaminoglycans,¹⁹⁵ as well as derived polyaminoacids. Other synthetic polymers often used in this field are polyvinylalcohol (PVA), PEO and different types of acrylates. However, the mechanical properties of these hydrophilic polymers often quickly deteriorate once in contact with biological buffers. This creates a challenging obstacle for cell seeding, transfer from the cell culture, and *in vivo* implantation. Additionally, numerous findings in mechanobiology have strongly suggested that the closer the match between the elasticity modulus of the scaffold and the

intended host tissue, the lower is the immunogenic and local inflammatory response during implantation.^{196,197}

In this context, reactive ES aims at enhancing the mechanical stability of weak, water-soluble polymers by performing an *in situ* cross-linking step during or after fiber generation. This procedure provides the balance between obtaining a soft hydrogel and enhancing the mechanical stability and water stability over time.

Different methods and chemistries to cross-link electrospun fibers involving water or other green solvents are presented in this section. However, only covalent cross-linking methods will be considered herein, whereas other non-covalent approaches, such as the use of inclusion complexes, hydrophobic interactions, or the formation of electrolyte complexes, can be found elsewhere.

4.1 Reactive electrospinning designs

Different designs of ES methodologies (co-axial, direct mixing, photo-curing, and co-spinning) with a focus on how this affects fiber morphology and cross-sectional composition are presented in Table 2.

During the reactive ES process, if the fiber drawing and cross-linking are occurring at the same time, the performance of the two processes has to be especially coordinated to properly match with the targeted ES setup (*e.g.* single, multi-needle or needleless) and the available ES time window dictated by the kinetics of the cross-linking chemistry. Indeed, above a critical cross-linking degree depending on the type of polymer and the linker, the change in viscosity of the spinning solution alters the formation of a stable polymer jet and thus fiber formation. When a cross-linking reaction undergoes considerable changes in viscosity within the ES time window, as with using glyoxal and gelatin, a multi-layer scaffold with a gradient in fiber diameters can be generated.¹⁹⁸ In order to overcome the problem of change in viscosity, cross-linking chemistries with extremely fast kinetic rates (*i.e.* aldehyde-hydrazine¹⁹⁹ or blocked isocyanate-amine²⁰⁰) can be spun by separating the fluids before spinning with a double-barrel syringe (Fig. 11a). Alternatively, gelatin and genipin have also been mixed *in situ* with a customized co-axial ES setup, where the inner needle wall had 20 circular holes ($\varnothing = 0.5$ mm) at its tip.²⁰¹

UV-induced ES covers a considerable number of studies regarding reactive ES, since its kinetic is generally fast and within the time window for ES (Fig. 11b).^{202,203} Since the cross-linking is triggered by UV exposure, the cross-linking degree during spinning has to be adjusted by optimizing the power amplitude of the UV lamp as well as the distance from the needle tip. Furthermore, a recent study has shown that the UV-spinning onto a plate collector submerged in a solvent for the un-cross-linked materials can lead to strikingly different mechanical properties compared to the more conventional “dry” UV ES.²⁰⁴ This aspect can also further contribute to expanding the range of mechanical properties achievable with the desired polymer network.

In 2020, an interesting combination of reactive chemistries has been proposed by integrating reactive units within func-



Table 2 Overview of the cross-linking strategies utilized for reactive ES

Route	Reactive ES formulation	Polymer substrate		Polymer concentration [wt%]	Cross-linker (catalyst)/polymer [mol/mol]	Ref.
1	Mixing	Glutaraldehyde	CS	1.7–2.0	1 : 1	207
	Mixing		Carboxyethyl CS	3.4	1 : 1	207
	Mixing		Gelatin	10	0.07 : 1	209
	Mixing		PVA	04/07/22	90–181 : 1	213
	Mixing		Pullulan	22	0.056 (wt/wt) + 1% H2SO4	217
	Mixing	Glyoxal	Gelatin	14	1.4–6.3 : 1	198
	Mixing		Gelatin	12/16/22	5.3–6 : 1	210
	Mixing		CS/PVA	0.6 CS/6 PVA	5 vol% Glyoxal	218
	Mixing	Oxidized sugars	Carboxymethyl CS	3	1.5 (wt/wt)	208
	Mixing		Gelatin	20	5/30 (wt/wt)	211
	Mixing	PO-CMC	Gelatin	6.25–25 blend (80 : 20 wt%)	1 : 0.25	212
2	Mixing	Collagen		16	1.5–2.0 : 1 (NHS : EDC)	222
	Mixing	Recomb. Collagen Peptides/CS		8	20 mM (only EDC)	225
3	Mixing	HA /PVA/CDs		6 PVA/6 HA/0–40 CDs	1 : 2 (wt/wt NHS : EDC)	226
	Mixing	CS/PVA/MCA		0.06–0.24 CS/6–9 PVA	20–80 mM MCA	227
	Mixing	Alginate/PVA/Citric acid		1 Alginate/5 PVA	5 wt% Citric acid	228
	Mixing	PAA/PEO		5 to 12.5 PAA/5–10 PEO 25	-	230
	Mixing	Gelatin/Glucose		2.5 CMC/2.5 PEO	up to 30 wt% Glucose	231
	Mixing	CMC/PEO/MCA		4.7 CS/0.25 PEO	3–5–7–10 wt% MCA	232
	Mixing	CS/PEO/MCA			1 wt% MCA	233
4	Mixing	Di-epoxide	Gelatin	15	2–6 wt% Di-epoxide	235
5	Co-axial	Gelatin		30	0.18 wt% Genipin /polymer	201
	Mixing	Silk-fibroin		12	0.12 wt% Genipin /polymer	238
	Mixing	Gelatin		20	0.01–0.05 wt% Genipin /polymer	239
6	Mixing	PVA/gelatin core/shell		8	0.5 to 2 wt% enzyme/gelatin	240
7	Mixing	TEOS		—	1 : 0.01 : 2 : 2	241
	Mixing	TEOS		—	TEOS : HCl : EtOH : H ₂ O	242 and
	Mixing	PVA/TEOS		1.4–5.6	TEOS : HCl : EtOH : H ₂ O	243
	Mixing	M-PVA		10	0.04 wt% HCl/TEOS	246
8	Mixing + UV	PHEMA		N/A	5–10–20 wt% MPTMS/2 wt% PI	247
					318 : 1 : 1.33 (PHEMAc: EGDMA : PI)	248
9	Dual-syringe + UV	THA/PEGDA/PEO		2.5% w/v PEO/2.5% w/v THA	9.0% w/v PEGDA	250
	Mixing + UV	PVP		10	5 wt% PMC/5 wt% TTT/3 wt% PI	251
10	Mixing + UV	Gelatin/PAA-g-Az		23% w/v – 10 Gelatin	2.3% w/v - 1 wt% PAA-g-Az	192 and 252
11	No mixing (2 syringes)	Aldehyde- & hydrazide-HA/PEO		3.5 Aldehyde-HA/2.5 PEO	3.5 wt% hydrazide-HA/2.5 wt% PEO	205
	Double-barrel	Aldehyde- & hydrazide-POEGMA		7.5	N/A	253
12	Mixing under Ar	Selenol-PETox-EI		30	2–4–5–6 mol% Selenol (-SeH)	256

The first column of the table is related to the routes of reactive ES, followed by the formulation, and which polymers are used at which concentration. The 5th column, “Cross-linker/polymer”, mentions the ratio between the cross-linker and the polymer. AA=acrylic acid, BAPO = bis-acylphosphin oxide, CQ=camphorquinone, EGDMA = ethylene glycol– dimethacrylate cross-linker, HA = Hyaluronic acid, MA = methacrylic acid, MCA = multi-carboxylic acids, MMA=methyl methacrylate, MPTMS = (3-mercaptopropyl)-trimethoxysilane, M-PVA = methacrylated polyvinyl alcohol, PAA = Polyamic Acid, PEGDA = poly(ethylene glycol) diacrylate, PETox-EI = partially hydrolyzed PETox (poly(2-ethyl-2-oxazoline-)-co-ethylenimine, PI = Photoinitiator, PMC = Pentaerythritoltetrakis(2-mercaptoacetate), PO = phenyl-bis(2,4,6-trimethylbenzoyl)-phosphine oxide, POEGMA=poly(ethylene glycol) methacrylates, PAA-g-Az = Poly(acrylic acid-g-azidoaniline), PO-CMC = partially oxidized carboxymethylcellulose (bearing aldehydes groups), THA = Thiolated hyaluronic acid, TTT = 2,4,6,8-tetramethyl-2,4,6,8-tetravinylcyclotetrasiloxane.

tional block copolymers for *in situ* UV ES as well as complementarily reactive groups for covalent cross-linking.²⁰⁵ Two separate solutions of HA modified with norbornenes exclusively functionalized with hydrazides and aldehydes were oppositely spun at the same time on a rotating mandrel (Fig. 11c). Thanks to the presence of these reactive groups on the fiber surface, fiber-to-fiber cross-linking was promoted by

mechanical loading, therewith endowing the scaffold with mechano-sensitive properties.

4.2 Cross-linking of amine and hydroxyl groups

This section and onwards will be focused on a description of already adopted cross-linking chemistries, including the reac-



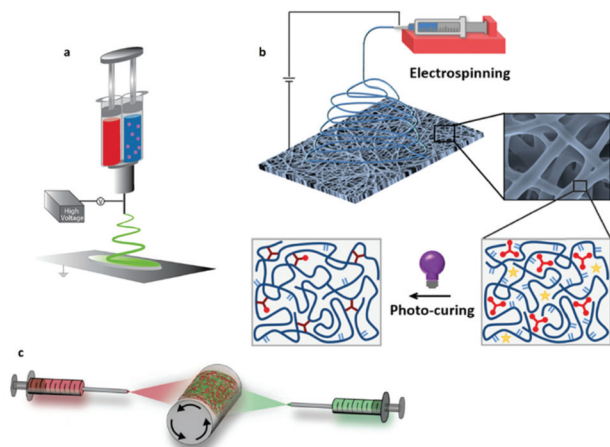


Fig. 11 Different designs for *in situ* cross-linking strategies: (a) co-axial/double-barrel electrospinning; (b) photo-induced cross-linking; and (c) the development of multi-fiber hydrogel network. (a) Adapted from ref. 199 and (b) adapted from ref. 203 with permission from the American Chemical Society. (c) Adapted from ref. 205 with permission from Wiley-VCH Verlag GmbH & Co. KGaA.

tants and eventual pre-functionalization of the polymer, experimental conditions, and residual reactive units.

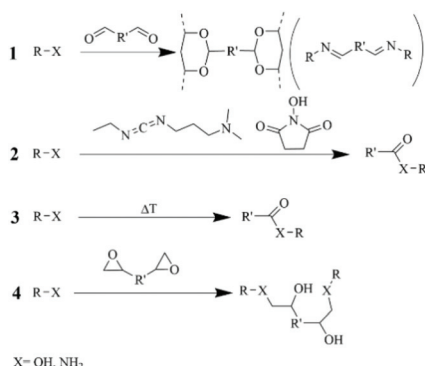
4.2.1 Di-aldehydes. Aldehydes undergo condensation reactions with 1,3-diols and primary amino groups. Therefore, model polymers for these reactions are PVA and CS, respectively (Route 1 in Scheme 1). While in the first case, the aldehyde undergoes cyclization forming acetal bridges, amine condensation with an aldehyde yielding an imine.

Glutaraldehyde has been one of the most investigated cross-linkers, probably because it is commercially available as a disinfectant and utilized for niche medical procedures, as with wart treatment.²⁰⁶ Imine cross-linking with CS or polyethyleneimine proceeds too fast and the solution cannot be spinnable unless its kinetic is quenched by using organic

acids as solvents,²⁰⁷ mostly AcOH and formic acid. Another way to slow down the reaction kinetic is to use PEO. This polymer interpenetrates the network and ensures sufficient dilution of reactive groups to enable stable ES up to 5 h. The use of an ethyl carboxylated homolog of CS (CECS) further delays the cross-linking kinetic reaching up to 6 h of stable ES with the same experimental conditions. For both polymers, reactive cross-linking did not significantly alter the fiber diameter compared to the glutaraldehyde-free membrane homolog (~60–140 nm for CS vs. ~270 nm for CECS). More recently, partially oxidized saccharides displaying two aldehyde functionalities were synthesized with sodium periodate as oxidant.²⁰⁸ Alginate di-aldehyde delays the gelation point of carboxymethyl CS up to 12 min if the two solutions are mixed on-line prior to ES and they are diluted with PEO. The resulting scaffolds have shown better biocompatibility compared to glutaraldehyde-based ones. As well as improved retention of the fiber morphology.²⁰⁸

While CS possesses a high concentration of primary amino groups, as it is composed only of aminoglycosides, gelatin and collagen are proteins with only lysine as the primary amine containing amino acid unit (e.g. concentration of amino groups of 0.286 mmol g⁻¹ in porcine skin type A vs. ~4.0–5.0 mmol g⁻¹ in CS). This feature, combined with the reactive ES with glutaraldehyde at relatively low cross-linker concentration (0.01:1 glutaraldehyde/gelatin by weight),²⁰⁹ delays the cross-linking process to yield relatively uniform fiber diameters of ~600 nm. On the other hand, although the tensile strength increases up to the lower MPa range, the fibers are extremely fragile, with elongation at break of 3%.

The reaction velocity of the imine condensation between aldehydes and primary amines from lysine subunits in gelatin can be considerably reduced when a shorter di-aldehyde is utilized, such as glyoxal, achieving several hours of stability using standard ES. Glyoxal bears two aldehydes directly linked together and its reactivity is strikingly different from glutaraldehyde. After the reaction of glyoxal with an amine, the resulted imine further undergoes cyclization together with another mole of the imine (directly linked to aldehyde) group to yield imidazole group. Longer kinetic times for cross-linking can then be utilized for the development of gradient scaffolds with increasing fiber diameter (e.g. from 60 to 680 nm) from bottom to top.¹⁹⁸ Despite the use of a cross-linker in excess of up to 6.7 times the amount of primary amines, the fibers showed an elongation at break at 180% and a lower Young's modulus of 880 kPa. Contrary to this, nanoin-indentation measurements of electrospun fibers from gelatin and glyoxal mixtures showed much lower elastic moduli down to 2.1, 3.2, and 10.9 kPa when the needle size was 2.1, 1.4, and 0.9 mm, respectively (e.g. compared to the 0.8 mm utilized by the previous work).²¹⁰ Other cross-linking strategies for gelatin by using di-aldehydes include reactive ES with alginate²¹¹ and carboxymethylcellulose.²¹² When cross-linked with carboxymethylcellulose di-aldehyde, the fiber Young's moduli are between 10 and 15 MPa.



Scheme 1 Cross-linking reactions occurring with polyelectrolytes bearing amine and hydroxyl groups. Route 1 yields acetal and imine groups whether the functional group was a hydroxyl groups (1,2-diols or 1,3-diols) or an amine group. Route 2 and 3 depict the reactions with R-X and carboxylic groups enabled by EDC/NHS and temperature, respectively. Route 4 shows the crosslinking behavior of epoxides.

When di-aldehydes are reacting with 1,3-diols, as in the model compound PVA, a condensation reaction occurs to form an acetal bridge between the macromolecules without addition of catalyst. In contrast, the imine formation *via* amine-aldehyde reaction is too slow to proceed *via* reactive ES at room temperature without any catalyst. This issue has been tackled by the group of Khan,²¹³ who added hydrochloric acid as catalyst to match the cross-linking within the ES window. They have shown that the formation of stable electrospun fibers upon soaking in water could be produced only above a glutaraldehyde/PVA molar ratio of 90:1. Moreover, *in situ* cross-linking enables stable ES of fibers at lower concentrations down to 4 wt%, which results in the formation of smaller fibers with an average diameter of 178 nm instead of 233 nm. This setup has been utilized also in other studies for producing conductive fibers when blended with semi-conductor ionomers,²¹⁴ as a substrate for ultra-thin layer chromatography,²¹⁵ and enzyme immobilization.²¹⁶ Similarly, recent work has shown that this method is valid also for polysaccharides.²¹⁷ Specifically, sulfuric acid was added to the formulation with glutaraldehyde to obtain *in situ* cross-linking of pullulan fibers. The cross-linking kinetic is even slower when glyoxal instead of glutaraldehyde is utilized as di-aldehyde cross-linker, similar to what was observed when reacting it with amines. In this case, CS added to the blend is the primary target for the initial cross-linking of amino groups, which increases the entanglements of the polymer blend favoring its early-stage mechanical integrity.²¹⁸

4.2.2 Carbodiimide chemistry. Cross-linking can also occur by amide or ester condensation from carboxyl/amine and carboxyl/hydroxyl functionalities. However, amide and ester condensation does not occur spontaneously, thus the use of temperature (see section 4.2.3) or catalysts is required. Common strategies for amine/carboxyl bioconjugation in proteins include the use of reactant displaying the highly reactive carbodiimide moieties, as in the water-soluble molecule 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC)²¹⁹ (route 2 in Scheme 1). The carbodiimide reacts first with the carboxylic units to form an O-acylisourea intermediate, which then specifically reacts with primary amines/hydroxyl groups to form an amide/ester bond, leaving an isourea by-product. However, the O-acylisourea intermediate is prone to hydrolysis, thus being short-lived in aqueous solutions. The addition of *N*-hydroxysuccinimide (NHS) and its sulfonated homolog (Sulfo-NHS) stabilizes and maximizes the yield of ester/amide coupling.^{220,221}

The kinetics of this reaction is normally too fast to permit a stable ES process. A breakthrough was achieved by Meng *et al.*, who used an excess of NHS/EDC to delay the coupling reaction to match the duration of the ES process.²²² This concept was used to cross-link collagen in an EtOH/PBS buffer solution yielding fibers with diameters of 420 nm, which was roughly doubled when compared with un-cross-linked fibers with the same solvent system. Storage humidity of the membrane after ES was found to play a key role in the preservation of the fiber architecture. Unfortunately, fibers were stable in water only

after production at relatively low humidity conditions (*e.g.* 43 and 53% after 3 and 1 day, respectively), with only the scaffold produced at 33% RH showing substantial retention of the fibrous structure. Indeed, water droplets condensing on the fiber, on one hand, increase the cross-linking conversion, but on the other hand, they provoke the fusion and merging of the fibers, therewith corrupting the fiber architecture and its porosity.

A similar strategy in terms of NHS/EDC amount was pursued to cross-link gelatin in combination with a thermo-sensitive polymer, poly(*N*-isopropylacrylamide) (PNIPAM), although utilizing a toxic solvent (*e.g.* 2,2,2-trifluoroethanol) for controlled drug release.²²³ Despite PNIPAM is not expected to contribute to cross-linking, several fiber-to-fiber coagulation areas are observed, which results in a loss of fibrous morphology. However, the gelatin-PNIPAM electrospun fibers could be potentially produced using water as a solvent since waterborne ES of PNIPAM has already been performed.²²⁴

A recent work, which combined CS and recombinant gelatin peptides, has shown complete retention of nanofibrous morphology when cross-linked with EDC in the absence of NHS as *in situ* cross-linking strategy.²²⁵ Contrary to conventional post-cross-linking by EDC, where the porosity decreases by one-third, the fibers retain a high surface-to-volume ratio with porosities above 60%, and consequently, their water uptake is much higher (1314% instead of 927%).

On the other hand, ester condensation does not occur fast enough to provoke cross-linked fibers during the ES process. Indeed, a polysaccharide bearing carboxylic functionalities, hyaluronic acid (HA), completes the cross-linking only after annealing it at 60 °C for 24 h, obtaining fibers with an average diameter around 200–300 nm and stable at all RH conditions.²²⁶

4.2.3 Thermal esterification and amidation. Herewith, procedures involving temperature-triggered condensation of carboxylic acids with hydroxyl groups/amines functionalities are depicted (route 3 in Scheme 1). Carboxylic acids and hydroxyl/amines may not be present on the polymer chain of blends. In that case, a low molecular weight molecule acting as a cross-linking agent can be added into the polymer solution.

Different polymer blends bearing carboxylic and amine/hydroxyl units have been thermally cross-linked with this approach (120–170 °C from few minutes up to hours), namely PVA/CS,²²⁷ PVA/alginate,²²⁸ PVA/poly(acrylic acid) (PAA),²²⁹ polyamic acid/PEO,²³⁰ gelatin,²³¹ carboxymethyl-cellulose,²³² and CS.²³³ For most of these approaches, multi-carboxylic acids, namely citric acid,^{227,228,234} succinic acid,²²⁷ itaconic acid,²³³ 1,2,3,4-butanetetracarboxylic acid,^{227,232} were added to the ES formulation to facilitate the diffusion of the reactive functional groups within a suitable distance for the thermal crosslinking to occur. A similar strategy was employed for this scope in other works, where the authors have explored the OH-rich cross-linkers glucose²³¹ and cellulose nanocrystals.²²⁹ A systematic study of the respective morphologies of PVA/CS electrospun blends has clarified that the higher the carboxyl functionality of the multi-acid cross-linker, the higher its stability



in water and retention of fibrous morphology without any substantial difference in the water uptake.²²⁷ This observation is in agreement with the observed decrease of its elongation at break from 43% with succinic acid compared to 20% with 1,2,3,4-butanetetracarboxylic acid.

4.2.4 Epoxy coupling. Epoxides are highly reactive towards nucleophiles, such as hydroxyls, amines, and thiols, due to the strain on their 3-membered ring structure. The products are ethers and substituted amines, respectively (route 4 in Scheme 1). However, analysis of the gelatin cross-linking degree in the presence of 1,4-butanediol diglycidyl ether showed that the reaction yield is only 10% when 2 wt% of the cross-linker is used for a reaction time of 24 h whereas it increases up to 70% at 6 wt% and 72 hours.²³⁵ Cross-linking provokes a shrinking of the fiber diameter, down to 280 nm at 72 hours, although the fibers are still relatively brittle with elongation at break between 15 and 35%.

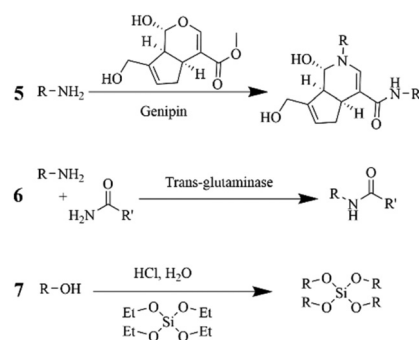
4.2.5 Genipin. Genipin is a hydrolytic product from geniposide extracted from gardenia fruits with cytotoxicities 5000 to 10 000 times lower than conventional glutaraldehyde-based cross-linking methods. Genipin can be considered as a bi-functional cross-linker in the presence of polymers with pendant amino groups to form stable amine and amide linkages (route 5 in Scheme 2). Although the formation of a cross-linking network reduces its crystallinity,²³⁶ the scaffold is still relatively brittle with respective elongation at break of 4.5–5.0% similar to the un-cross-linked CS.²³⁷ Arginine and lysine residues in fibroin undergo cross-linking with genipin in FA after mixing for 2, 15, and 24 h before ES.²³⁸ The increase in conductivity at higher mixing times causes a reduction of the fiber diameter and, as expected the higher the cross-linking degree, the lower is the extent of fiber dilatation upon swelling in water. The same amino acids are also responsible for the cross-linking with genipin at 1–5 wt% with gelatin.²³⁹ ES of the precursors immediately after mixing leads to fibers that still require storage in water vapor saturated conditions to complete the cross-linking reaction.

An innovative design for co-axial spinning of gelatin and genipin is described by Gualandi *et al.*²⁰¹ In this case, through the inner needle that contained several holes, a genipin solution could diffuse out for thorough mixing with the sheath

gelatin flow. Although the resulting fibers in a range of 320–350 nm were initially stable with minimal fused fiber-to-fiber intersections, the fibers still required a thermal and EtOH treatment to preserve their stability in water. Overall, despite its low toxicity and promising cell compatibility in combination with different polymeric substrates, the costs of genipin and the fiber generation process are still intolerable for most applications.

4.2.6 Enzymatic cross-linking. Transglutaminase is an enzyme able to catalyze the cross-linking reaction between the carboxamide groups of glutamine residuals and primary amino groups from lysine residuals (route 6 in Scheme 2). Only one example has been reported recently for the cross-linking of gelatin by utilizing enzymatic reactive ES.²⁴⁰ In this study, a co-axial needle for spinning PVA as core flow and gelatin pre-mixed with transglutaminase at different mixing times were utilized. Although relatively smooth fibers with an average diameter of 270 nm were presented, no mechanical characterization and nor its biodegradation profile in water was reported.

4.2.7 Silane. Hydroxyl groups can react in the presence of multi-functional organosilane (*i.e.* tetraethylorthosilicate (TEOS)) to yield siloxane bridges between the polymer substrate and the cross-linker in the so-called sol-gel reaction (route 7 in Scheme 2). This synthetic pathway requires a catalytic amount of acids and/or bases to initiate the reaction. Few studies have reported the direct spinning of TEOS to form silica fibers. The solution is *per se* not spinnable unless the sol-gel reaction of TEOS is initiated beforehand.²⁴¹ An optimum window of cross-linking degree, achieved when the viscosity is 100–200 mPa s, represented a good compromise between the smaller size of the agglomerates (less than 3.5 nm) and the concentration of EtOH, which resulted in the spinning of smooth fiber constructs and the mechanical integrity of the scaffold.²⁴² Interestingly, the surface of these fibers was almost superhydrophobic once deposited on the collector with a water contact angle approaching 150°, followed by a sudden switch to superhydrophilic surfaces after exposing to 100 days of highly humid environment due to the substitution reaction of the hydrophobic ethoxyl groups (Si-OCH₂CH₃) to the hydroxyl groups (Si-OH) over time.²⁴³ This transition could be accelerated by heating the substrate above 400 °C for a few hours. Sol-gel cross-linking occurs when TEOS is mixed with di-functional siloxane prepolymers. However, bi-functional pre-polymers in combination with TEOS do not contain sufficient reactive units per volume to yield a sufficient cross-linking degree forming merged fibers.^{244,245} PVA, especially at high de-acetylation degrees, yields plenty of hydroxyl functionalities and enables the use of green solvents like water and EtOH. A first investigation of PVA-TEOS reactive ES has shown that PVA can be spun at relatively low concentrations (down to 1.4 wt%), thanks to the increase in viscosity during the reactive cross-linking.²⁴⁶ To reach the optimal viscosity, the two components still needed to be pre-cross-linked during thermal treatment at 60 °C for 1 h. The resulting fibers showed diameters down to 50 nm at 10 wt% of TEOS. Unfortunately, at such concen-



Scheme 2 Cross-linking reactions specific for amino and hydroxyl groups using (5) Genipin, (6) Transglutaminase, and (7) TEOS.



tration, the water stability of the scaffold was relatively poor, since the minimum ratio at which stable cross-linked scaffolds were produced was 27 : 7 TEOS/PVA. A twist to this strategy was combining the thiol-ene induced coupling of thiol-derived organosilane with methacrylated PVA during UV-assisted ES in DMSO.²⁴⁷ The product of this reaction was not a membrane scaffold, but rather an assembly of nanowires with an average length of 800 nm and diameters of 35–55 nm, which is rather peculiar among the different reactive ES processes.

4.3. *In situ* photo-induced cross-linking

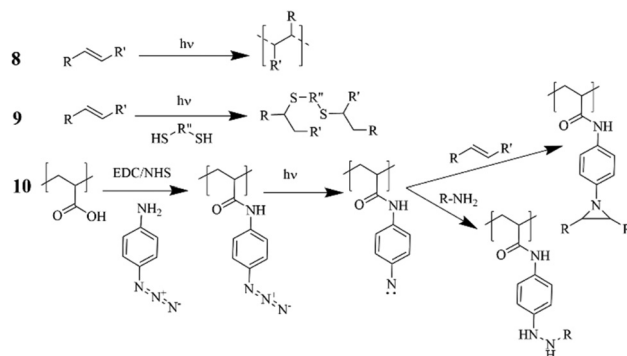
The simultaneous UV-induced cross-linking of double bonds during the ES process is an appealing alternative to cross-link on demand the electrospun scaffolds and retain the high porosity features of this technique. The mechanism is mainly a radical-initiating polymerization of the terminal and substituted alkene or methacrylate groups.

While substituted alkenes are generally formed by inserting functional monomers during the polymerization process, terminal alkenes and (meth)acrylates are formed by post-functionalization of the end groups or the pendent units of the macromolecule. Alternatively, even the direct polymerization of mono and di-functional (meth)acrylates, after thermal pre-conditioning can offer appealing versatility in terms of the nature of the functional groups incorporated into the polymer.

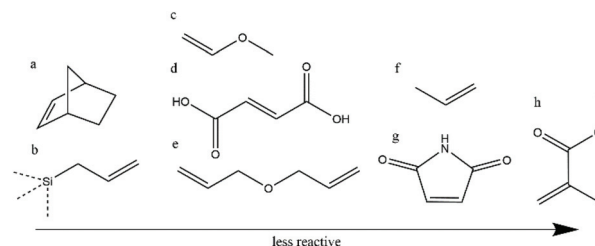
A twist to this approach has included in the past few years the use of thiol as co-reactant in these radical reactions, thereby remarkably expanding the window of polymer architectures achievable with this technology. Finally, other photo-induced cross-linking mechanisms, such as the recently explored nitrene chemistry will also be debated in this section.

4.3.1 Photo-polymerization of functional (meth)acrylate monomers and macromonomers. Functional (meth)acrylates can be used in reactive ES only when part of the polymerization is carried out beforehand in order to reach the optimal viscosity for the spinning process. In this case, poly(hydroxyethyl)methacrylate (PHEMA) fibers have been produced by initiating the polymerization with a thermal initiator and by continuing it during the ES process with a photo-initiator (route 8 in Scheme 3).²⁴⁸ The average fiber diameter of the swollen nanofibers in water ranged between 50 and 800 nm whereas atomic force microscopy imaging has shown the elastic recovery of the scaffold. Unfortunately, this was the only procedure involving green solvents since all the other methods reported in the literature used DMF or halogenated solvents to perform functionalization reactions and ES.

4.3.2 Thiol-ene photo-polymerization. Polymers with terminal/internal alkenes can also undergo photo-induced addition with thiols (route 9 in Scheme 3). This chemistry remarkably expands the window of polymer architectures that can be obtained with reactive ES. However, their reactivity varies depending on the type and degree of substitution of the double bond, according to the following order: norbornene, vinyl silane > allyl ether, vinyl ether, fumarate > propene, maleimide >> methacrylate (Scheme 4).²⁴⁹



Scheme 3 Overview of the different photo-induced cross-linking chemistries using (8) vinyl groups, (9) vinyl groups and thiols, and (10) polyacrylic acid and 4-azidoaniline forming nitrene intermediate under UV-light that can further react with amines or vinyl groups on polymers, thus acting as a cross-linker.



Scheme 4 Chemical structure of molecules undergo photo-addition with thiols in order of reactivity: (a) norbornene, (b) vinyl silane, (c) vinyl ether, (d) fumarate, (e) allyl ether, (f) propene, (g) maleimide, and (h) methacrylate.

Ji *et al.* used a double syringe setup containing aqueous solutions of hyaluronic acid functionalized with thiol groups and poly(ethylene oxide) in the first syringe and poly(ethylene glycol) di-acrylate (PEGDA) in the second syringe.²⁵⁰ The reactive ES process initially produced a uniform distribution of fibers, with a mean diameter of 90 ± 15 nm. Moreover, the thiol functionalized HA and the PEGDA underwent cross-linking reaction at RT in a time frame of 10 min without using UV light. The so prepared fiber network enabled fibroblasts to form 3D dendritic networks penetrating within the scaffold.

Not only macromonomers but also small multi-functional thiol-ene blocks can be easily translated into non-woven mats by reactive ES. Thiolated derivatives from pentaerythritol were utilized as tetra-functional blocks with cyclic siloxanes bearing four²⁵¹ terminal alkene per molecule. When adding PVP to the tetra-functional polysiloxane/thiolated pentaerythritol formulation, phase separation occurred and core-shell morphologies were obtained.²⁵¹ These fibers were generated in a mixture of EtOH and EtOAc.

4.3.3 Nitrene formation. Besides the alkyne-azide cyclo-addition, phenyl azide groups can also disproportionate photo-chemically leading to a highly reactive nitrene intermediate (route 10 in Scheme 3). This can either promote addition reac-



tions to an alkene or form a dehydroazepine intermediate and subsequently react with primary amines. The latter route has been utilized in the literature for the reactive ES of gelatin. Azide polymers were synthesized from polyacrylic acid and 4-azidoaniline with EDC/NHS coupling chemistry (route 2 in Scheme 1). Gelatin has been shown to undergo cross-linking when blended 9:1 with these reactive units.^{192,252} Their Young's modulus of 5.5 ± 1.0 kPa were lower than gelatin cured with glutaraldehyde, which suggests a lower cross-linking degree.²⁵² However, their cytotoxicity was lower than glutaraldehyde-based scaffolds, with better adhesion of fibroblast and good potential for cell scaffold in osteogenesis.¹⁹²

4.4 Other cross-linking mechanisms

4.4.1 Hydrazone click chemistry. One of the most recent chemistries for reactive ES is the conjugation of aldehyde or ketone groups and hydrazide moieties to yield hydrazone bond formation readily at RT (route 11 in Scheme 5). This chemistry, besides being highly reactive and compatible with the reactive ES time window, forms hydrazone bonds that are stable under neutral to alkaline conditions (pH = 7–9) and labile under slightly acidic conditions (pH = 4–5). This could turn useful for the controlled release of active agents at the targeted pH conditions. The group of T. Hoare pioneered the use of this chemistry by utilizing water-soluble polymer precursors derived from PEG, using methacrylated analogs of PEG (POEGMA) co-polymerized with acrylated monomers bearing aldehyde and hydrazide functionalities.²⁵³ In all studies, double-barrel syringes kept POEGMA functionalized with aldehydes separated from hydrazide before *in situ* mixing (Fig. 11a), since the gel time of the polymer mixture was around 45 min. Although the nanofiber average diameter increased from 0.34 ± 0.08 μm to 1.33 ± 0.20 μm , the textured nanofibrous structure was preserved in the swollen state. The structures, showing a swelling degree of 91% after few minutes, were capable of retaining an elastic modulus of 2.1 kPa over at least 40 cycles.

A follow-up study on the application of this type of cross-linking strategy for tissue engineering demonstrated the direct addition of biological cells to the ES solution in order to have cells inside or attached to the electrospun fibers, thus the cells were distributed in the whole scaffold already.¹⁹⁹ Cell viability was ensured by using water as the solvent and by the good bio-inertness of this coupling chemistry. The generated fiber membranes not only showed higher cell viability and proliferation compared to a bulk cell-loaded hydrogel of the same material, but also maintained this high cell viability and proliferative

capacity following a freeze/thaw cycle without requiring any cryoprotectant. Recently, the same group has also shown that, depending on the amount of PEO utilized as ES aid as well as on the weight ratio between hydrazide and aldehyde-based polymers, pure fibers to beaded fibers to bead network morphologies with tunable bead sizes can be fabricated.²⁵⁴ When utilizing ketone precursors, as with poly(acrylamide)-co-poly(diacetone acrylamide), the reaction proceeded slightly slower in the presence of a di-functional cross-linker, adipic dihydrazide, thus enabling the production of fibers with diameters of 200–220 nm by precursor mixing before usage.²⁵⁵

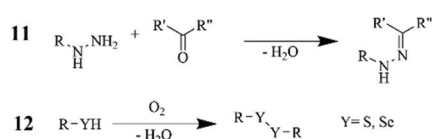
A new concept in the field of reactive ES for tissue engineering has been proposed by Davidson *et al.*, where fiber-to-fiber cross-linking could be triggered by mechanical loading, thereby mimicking the force-responsive properties of the extra-cellular matrix.²⁰⁵ To achieve this novel mechano-sensitive hydrogel, they have utilized hyaluronic acid as a functional polymer for the selective attachment of aldehyde or hydrazide units. Aldehyde functionalities in polysaccharides can be easily obtained by oxidizing the glucoside ring with sodium periodate as previously discussed (see section 4.2.1), whereas hydrazine moieties were introduced by reacting to the carboxylic groups in hyaluronic acid with adipic dihydrazide. Additionally, both polymers contain norbornene units for UV-induced reactive ES according to the previously described thiol-ene chemistry (*e.g.* dithiothreitol is also added as a bi-functional thiol linker) (see section 4.3.3). The mixed fiber scaffold with both unreacted hydrazide and aldehyde fiber network has been achieved by spinning the two different polymers with separate needles onto a common rotating mandrel. The advantage of this chemistry also offers the possibility to reverse the cross-linking in acidic pH (acetate buffer, pH = 4.3), thus restoring the fiber architecture similar to the state before its plastic deformation induced by *in situ* fiber-to-fiber cross-linking.

4.4.2 Selenide chemistry. Selenols (like thiols) undergo oxidative couplings upon exposure to oxygen, yielding diselenide bonds (route 12 in Scheme 5). Interestingly, the cleavage of this bond can be accelerated either by thermal or photo-induced stimuli to tune the overall biodegradation time of the scaffolds.²⁵⁶ Functionalization with selenide units was conveniently achieved by reacting the secondary amino groups present in the copolymer of 2-oxazoline and ethyleneimine according to the above-mentioned ring-opening reaction. Most importantly, this hydrogel was prepared by utilizing only water as solvent.

4.4.3 Further potential cross-linking mechanisms. Other chemistries for reactive ES currently using toxic solvents for the functionalization of the polymer and/or for the ES process are reported, which potentially offer further green ES strategies by substituting these solvents accordingly.

Specifically, the reaction of the trimetaphosphate cyclic anion with 1,3-diols leads to ring-opening and cycle re-arrangement directly with the polymer backbone, for example with pullulan and dextran blends.^{257–259}

In a grafting approach, (meth)acrylation of polymers bearing hydroxyl/amine moieties can be achieved with smaller building blocks such as glycidyl methacrylate,^{260–262} 2-isocya-



Scheme 5 Other cross-linking strategies involving hydrazone bond (11), and disulfide and diselenide formation (12).



noethyl-methacrylate,²⁶³ methacrylic anhydride,^{264,265} and acryloyl chloride.^{266,267} Subsequently, the (meth)acrylate units can be crosslinked following the route 8 or 9 of Scheme 3.

Isocyanate coupling is another potentially transferrable strategy to cross-link polymers bearing amino and hydroxyl groups, which yields urea and urethane bridges by reacting 1,6-hexamethylenediisocyanate (HMDI) in the presence of 1,4-diazabicyclo[2,2,2]octane (DABCO)²⁰⁰ or blocked isocyanates^{268,269} and hexamethylene-1,6-diaminocarboxysulfonate.^{270–272}

Despite their potential, certainly the latter approaches and others based on quite toxic components (*e.g.* glutaraldehyde, epoxides, isocyanate, *etc.*) risk higher cytotoxicity in tissue engineering applications due to residual crosslinkers, impairing the scaffold transplantation. Therefore, selection of chemical routes involving less toxic crosslinker/catalyst (*e.g.* carboxylic acid, genipin, enzymes, *etc.*) is generally preferred.

5. Conclusions and future perspectives

Methods to achieve green ES, including exclusively sustainable and non-hazardous solvents, are under constant development and have been practically available for most common types of polymers. Available solvent chart tables that classify the solvents according to their greenness offer the possibility to easily select solvents based on their health and safety properties, environmental impact and disposal requirements. Since classical ES requires the use of copious volumes of solvents, the choice of solvent has a large impact on the sustainability of the production process. Especially the use of halogenated solvents should further be avoided due to their environmental and safety impact. Hence, greener solvent alternatives have been listed, among which the most common ones are water, EtOH, MeOH, AcOH, acetone and DMSO. By employing water-based spinning solutions, the fields of emulsion and suspension-based spinning offer vast potential and unrivalled opportunities for encapsulation and immobilization of active agents, by either completely avoiding or at least drastically minimizing the use of organic solvents. Another sophisticated strategy to avoid organic solvent completely is the suspension electrospinning, which allows the generation of fibers from undissolved polymer particles, and can be sintered after the ES process. To this end, various compositions and conditions to prepare not only carrier membranes for drug delivery and storage of volatiles, but also functional membranes for filtration and catalysis from dispersions have been summarized. In contrast, reactive ES was shown to be yet another promising technique to achieve green manufacturing conditions, whereas it often escalates in complexity compared to the more traditional ES approaches. Nevertheless, it offers the formation of particularly stable 3D fiber networks using water-soluble polymers and green solvents. And practically, the only necessary steps to achieve cross-linking reactions are the addition of a suitable cross-linking agents and curing conditions, according to the type of functionalities of the polymer.

Despite these options, the final selection of solvents relies on the assessment by the user. Limiting the solvent choice to green solvents inherently reduces the possibilities to optimize the ES parameters of the spinning process for desired fiber and membrane production. Also, green solvents usually have higher boiling points, which renders them safer to work with, but can cause a higher energy demand, such as through higher required voltages, more time consuming and complex manufacturing processes, or higher temperature and/or lower air humidity. Furthermore, the polymer solubility sometimes depends on other polymer properties, such as conformation, molecular weight and polydispersity. For example, not all aliphatic polyesters and even just PLAs are equally well soluble in acetone, although they are synthesized from the same monomers.²⁷³ Another limitation of using green solvents in ES is the limitation in mixing different types of polymers in one spinning solution (polymer blend), particularly without the solvation capabilities of halogenated solvents.

As the environmental and ecological awareness of consumers is increasing from year to year, also the production and material selection of products that were not in focus yet, will have to adapt for a more green manufacturing practices. Members of the ES industry might be hesitant to introduce unfamiliar green solvent alternatives, therefore we hope that placing more emphasis on reactive and dispersion electrospinning could offer sufficient incentive for implementing these alternative green pathways. Apart from the obvious sustainability and customer interest benefits, this allows for an extension of the choice of materials for blends which are otherwise impossible to find a good co-solvent for, *e.g.* blends of hydrophilic and hydrophobic polymers or incorporation of additives and active compounds.

As restrictions for the use of certain solvents are increasing, solvent alternatives, especially to replace halogenated solvents, are increasingly investigated in the electrospinning field. While most reports still use classical solvents, literature has shown that the most common polymers can be electrospun with green solvents to obtain membranes exhibiting largely the same properties. In the end, it can and hopefully will be just a matter of habit to first select green solvents. After all, the drive towards safe, environmentally conscious and sustainable ES-manufacturing is only increasing.

Abbreviations

AcOH	Acetic acid
CA	Cellulose acetate
CED	Cumulative energy demand
CS	Chitosan
CMC	Carboxymethylcellulose
DMAc	Dimethylacetamide
DMF	Dimethylformamide
DMSO	Dimethylsulfoxide
EDC	1-Ethyl-3-(3-dimethylaminopropyl)carbodiimide
EHS	Environmental, health, and safety indicator



ES	Electrospinning
EtOAc	Ethyl acetate
EtOH	Ethanol
FA	Formic acid
LCA	Life-cycle assessment
MEK	Methyl ethyl ketone
2-MeTHF	2-Methyl tetrahydrofuran
NHS	N-Hydroxysuccinimide
Nylon-6	see PA6
Nylon-11	see PA11
O/W	Oil-in-water
PA6	Polyamide 6 (Nylon-6)
PA11	Polyamide 11 (Nylon-11)
PAA	Polyacrylic acid
PAAm	Polyacrylamide
PAN	Polyacrylonitrile
PCL	Polycaprolactone
PEO	Polyethylene oxide
PHEMA	Poly(hydroxyethyl) methacrylate
PLA	Poly-lactic acid
PLGA	Poly(lactic-co-glycolic acid)
PMMA	Polymethylmethacrylate
PS	Polystyrene
PTFE	Polytetrafluoroethylene
PVA	Polyvinylalcohol
PVDF	Poly(vinylidene fluoride)
PVP	Polyvinylpyrrolidone
REACH	Chemical Control Regulation in the European Union
RH	Relative humidity
SEM	Scanning electron microscopy
TEM	Transmission electron microscopy
TEOS	Tetraethylorthosilicate
THF	Tetrahydrofuran
UV	Ultraviolet light
W/O	Water-in-oil

Author contributions

J. A., G. H. and C. T. conceptualized, compiled and drafted the review. J. A., G. H. and F. I. edited the manuscript, supported by C. T. F. I. and R. M. R. supervised the work. All authors checked and modified the manuscript, and agreed to the final version.

Conflicts of interest

There are no conflicts to declare.

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