Water-responsive materials for sustainable energy applications

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Water-responsive (WR) materials that mechanically change volume in response to changes in relative humidity or water/humidity gradient can generate significantly higher energy actuation over natural muscles and conventional actuators. Recent proof-of-concept demonstrations have shown the great potential of using WR materials as high performance actuator components for various energy-related applications. For example, evaporation-driven engines which are enabled by spore based WR materials directly harvest natural evaporation of water and convert it into mechanical work and electricity, highlighting the possibility of using this untapped energy source of evaporation as an additional option for clean and low-cost energy generation and storage. Despite the growing interests of these examples, research on WR materials and their applications are still in an early stage and face multiple challenges. Namely, the fundamental mechanisms that engender material’s water-responsiveness are unclear. Additionally, current systems remain difficult to scale up, and the integration of WR materials into modern engineering systems is a critical design challenge. Here, we review the current development in this emerging category of WR materials. We discuss up-to-date studies on both natural and synthetic WR materials, which include their processing, characterization methods, as well as scientific and technical challenges that can be possibly advanced by future research endeavours.

1. Introduction

Typically, mechanical actuators or muscles convert stimuli, such as electricity, heat, light, magnetism, or chemical reaction into mechanical motion. This is different from a Fortune Teller Miracle Fish that automatically bends and moves when it sits on a wet surface, for example a sweating hand (Figure 1a). Materials that drive these motions are usually called water-responsive (WR) materials (also called humidity-responsive materials or water/humidity gradient responsive materials) as they can reversibly swell and shrink in response to changes in relative humidity (RH) or water/humidity gradients and convert the chemical potential of water into mechanical energy (Figure 1b). These materials’ WR actuation can be extremely efficient and powerful. For example, wooden wedges can function as actuators as they swell when soaked with water, and they were widely used as means of splitting rocks before the modern industrialization. While WR materials have a long history of human use as actuators, modern engineering systems have not employed this mechanism for high efficient actuation.
Over the last decade, natural WR materials that exert significant actuation energy inspired the growing studies of using materials’ water-responsiveness as a potentially efficient and extremely powerful actuators.\(^1\)\(^,\)\(^8\)-\(^10\) Many examples of natural celluloses\(^11\)-\(^13\) and protein based structures,\(^8\)\(^,\)\(^14\) synthetic polymers,\(^10\)\(^,\)\(^15\) and composites of nanomaterials\(^16\)\(^,\)\(^17\) have shown remarkable WR behaviours and programmable movements, and could offer a number advantages, including high strain/stress, energy density, and efficiency, over conventional actuators driven by other stimuli (see **Table 1**). Notably, bacterial spores have demonstrated an extremely high WR actuation energy density, reaching \(\sim 20\) MJ m\(^{-3}\), which is higher than the typical values reported for materials frequently used as actuators and artificial muscles (**Figure 1c**).\(^1\) While WR materials’ actuation could be difficult to be precisely controlled by using traditional feedback control systems (**Table 1**), their high-energy and high-efficiency actuation has recently enabled a wide range of energy-related applications, including smart structures,\(^18\)\(^,\)\(^19\) low power consuming actuation,\(^20\)-\(^22\) and electricity generation.\(^1\)\(^,\)\(^9\)\(^,\)\(^23\)\(^,\)\(^24\) For instance, using cellulose structures, architects have built weather-responsive architectural systems that can autonomously adjust their openings upon changes of local RH without using any electricity (**Figure 2a,b**).\(^18\)\(^,\)\(^19\) Smart textiles that open and close in response to human body’s sweating facilitate comfort without adjusting the temperature of the environment (**Figure 2c,d**).\(^25\)-\(^27\) WR materials based soft robots can automatically and remotely operate and conduct various tasks powered by environmental RH fluctuations (**Figure 2e,f**).\(^28\)-\(^30\) Notably, recent pioneering researches have demonstrated the potential of using WR materials to directly and efficiently harvest energy from natural evaporation and RH fluctuations under ambient conditions.\(^5\)\(^,\)\(^9\)\(^,\)\(^10\)\(^,\)\(^15\) For example, a bioinspired polymer film of polypyrrole and polyl-borate composites flips continuously when placed on a wet surface. Taking advantage of this spontaneous WR locomotion powered by the water gradient, piezoelectric elements were attached to the film to convert its mechanical energy into electricity.\(^10\) A \(\pi\)-stacked carbon nitride polymer film has also been used to demonstrate direct energy harvesting from ambient RH gradients and fluctuations. When a water droplet was placed close to the film, the minute amount of RH fluctuations caused by the water droplet is enough to drive the film actuating and walking unidirectionally.\(^15\) Furthermore, we have developed WR materials based devices, which automatically generate piston-like linear (**Figure 2g**) and rotary motion (**Figure 2h**) when placed at air-water interfaces. These devices can directly harness energy from naturally occurring or engineered evaporation and subsequently convert it to mechanical energy or electricity.\(^1\)\(^,\)\(^9\) opening a new path toward the efficient harnessing of this abundant evaporation energy from water. While this energy harvesting technique is still in its early stages, theoretical studies have predicted the great potential of water as energy source.\(^31\)

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**Table 1.** Advantages and disadvantages of various stimuli-responsive materials (modified from 32-34)

<table>
<thead>
<tr>
<th>Stimulus type</th>
<th>Representative examples</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical potential of water</td>
<td>Bacterial spores, Spider silk, Water responsive polymers</td>
<td>High stress, strain, energy density, and efficiency.</td>
<td>Difficult to control.</td>
</tr>
<tr>
<td>Electricity</td>
<td>Dielectric elastomers, Piezoelectric materials</td>
<td>Good strain, stress, and efficiency.</td>
<td>High voltage and fields are required.</td>
</tr>
<tr>
<td>Heat</td>
<td>Thermally activated shape memory alloys and shape memory polymers</td>
<td>High stress, strain, and energy density.</td>
<td>Difficult to control, low efficiency and short life cycle</td>
</tr>
<tr>
<td>Magnetism</td>
<td>Ferromagnetic shape memory alloys, Magnetic particle/polymer composites</td>
<td>Good stress and strain.</td>
<td>Bulky magnets are required, high cost.</td>
</tr>
<tr>
<td>Chemical reaction</td>
<td>Mammalian skeletal muscles, Chemo-responsive hydrogel</td>
<td>Good stress, strain, and energy density.</td>
<td>Complicated system.</td>
</tr>
</tbody>
</table>

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*Figure 2.* (a) Cellulose based smart architectural system opens when the local environment is dry and (b) closes when the local environment is humid.\(^19\) Smart textiles with ventilation flaps (c) remain flat when the skin is dry and (d) open when the skin is sweating.\(^27\) (e-f) A soft robot built with a bilayered WR structure move forward during hydration and dehydration processes.\(^26\) (g) An engine generates oscillatory motions when placed above water. The oscillatory engine illuminates light emitting diodes (LEDs) when connected to an electromagnetic generator. (h) A rotary engine, powered by water evaporation from the wet paper, can continuously rotate and drive a miniature car (weighing 0.1 kg). Reproduced from Ref. 19, 27, 36, 9, with permission from John Wiley and Sons, The American Association for the Advancement of Science, Nature Publishing Group.*
Despite growing interests in materials’ WR actuation, development of WR materials still faces challenges in scaling up, meeting efficiency expectations, and integrating them into the existing engineering systems. To this end, we intend to provide a brief overview of recent studies on WR materials. We discuss WR behaviours of newly discovered natural materials and efforts on processing natural materials and hybrid materials with natural components. We also discuss synthetic WR materials, which include their structures, water-responsiveness, and fabrication methods. To promote development of WR materials for energy-related applications, we review quantitative characterization methods that are used to evaluate the WR performance and potential for practical applications. Finally, we share our vision of remaining scientific and engineering challenges, as well as future interdisciplinary research directions that could accelerate the growth of this emerging field of WR materials.

2. Current Water-Responsive (WR) Materials

2.1. Natural WR Materials

Biological systems have developed remarkable WR materials that power their daily tasks, such as plants’ dispersing or burying their seeds driven by daily RH cycles (Figure 3a-d). These natural WR behaviours are a consequence of their cellulose-based microstructures that mechanically deform in response to RH changes. In addition to plants’ cellulosic structures, many animal fibres and microbes were found to exhibit WR behaviours more significantly than plants. While the biological function of these animal fibres’ and microbes’ water-responsiveness is still not fully understood, they could potentially serve as low-cost and high-performance natural components for WR actuators and artificial muscles. In this section, we review current progresses on natural WR materials and hybrid/engineered WR materials composed of natural components.

![Figure 3](image-url)

**Figure 3.** Natural WR materials. (a) Wheat awns repeatedly bend and straighten fuelled by the daily RH cycles, leading to the propulsion of its seed into the soil. (b) The scales of pine cones open and release pine seeds when the environment is dry. (c) Ice plant’s seed capsule opens when the environment is wet due to the hygroscopic deformation of its unique keel structures. (d) The filaree seed’s awns bury the seed by drilling, as they wind and unwind in response to fluctuations of ambient RH. (e) Spider-dragline silk lifts 9.5 mg of plastic weight during repeated cycles of wetting and drying. (f) Bacillus subtilis spores expand and contract in response to changes in RH which can potentially generate a high amount of work.

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2.1.1. Plants

The dispersal and burial of many plants’ seeds rely on their WR structures’ predictable mechanical motions. For example, fuelled by the daily RH cycles, wheat awns repeatedly bend and straighten, leading to the insertion of its seed into the soil (Figure 3a). The scales of pine cones open and release pine seeds when the environment is dry (Figure 3b), while ice plant’s keel structures open up its seed capsule when the environment is wet (Figure 3c). The filaree seed drills into the soil as its own winds and unwinds in response to fluctuations of ambient RH (Figure 3d).
Recent studies show that many of the plants’ WR motions are controlled by bilayer structures with both active and passive layers. These bilayer structures, which usually consist of stiff crystalline cellulose microfibrils embedded in an amorphous matrix of polysaccharides, aromatic compounds, and structural proteins, can amplify the WR actuation of the active layers, but also create different types of motions, such as linear, bending, and coiling motions, to complete their vital tasks.\(^{11, 12, 37-39}\) For instance, the bending of pine cone’s scales relies on a bilayer structure that is composed of two kinds of tissues—sclerids in the outer surface and sclerenchyma fibres in the inner surface, and each surface has distinct mechanical and WR properties. When RH changes, sclerids in the outer surface deform more than sclerenchyma fibres in the inner surface, resulting in bending of the scales (Figure 3b).\(^{11, 38}\) While sclerids and sclerenchyma fibre cells consist of similar amount of cellulose (~20% volume fraction), the contrast in the alignment of cellulose microfibrils differentiates their stiffness, hygroscopic expansion,\(^{11}\) and water transport behaviours.\(^{40}\) Wheat awns’ bending motions are also controlled by a bilayer structure similar to pine cone scales, where cellulose microfibrils align in different directions per active and passive layers, leading to internal stresses to curve the entire awn structure (Figure 3a).\(^{32}\) Likewise, ice plant seed capsule opens up when exposed to humid environment as a result of swelling and bending of hygroscopic keels at the bottom of the seed capsule (Figure 3c).\(^{13}\) Dandelion seed pappi opens and closes in response to RH changes by anisotropic swelling and shrinking of pulvinus compartments.\(^{41}\) In addition to the bending motions, various plants use coiling motions to drive their seeds into the soil.\(^{42, 43}\) Most of the coiling motions depend on similar bilayer structures, but hygroscopic movements of these two distinctive layers are at a tilted angle to each other. For instance, the Geraniaceae seeds have bilayered awns where the tilted arrangement of cellulose helices within the active inner layer leads to the coiling of awns when responding to changes in RH (Figure 3d).\(^{42}\)

2.1.2. Animal Fibres
In addition to plants, many animal fibres exhibit outstanding water-responsiveness, such as spider silk that can shrink and expand up to ~50% in length when exposed to dry and wet environments (Figure 3e).\(^8\) Silk produced by some spider species can generate extremely high mechanical stress and WR energy density.\(^{45, 46, 47}\) For example, spider dragline silk’s WR stress and energy density can reach 80 MPa and ~500 kJ m\(^{-3}\), respectively, surpassing that of mammalian muscles and many artificial muscles.\(^8\) While it is still unclear why certain kinds of spider silks exhibit such a significant WR performance, researchers speculate that spider silk’s water-responsiveness plays an important role in maintaining structural integrity of its webs and catching prey.\(^{46, 47}\) Despite spider silks’ remarkable water-responsiveness, the impracticality of spider silk farming prevents their usage in engineering applications.\(^{48}\) In contrast, silkworm silk that has a similar primary protein structure as spider silk is readily available in large quantities.\(^{49}\) Several groups looked into the potential of using silkworm silk as an alternative WR material. While studies have found that silkworm silk changes length in response to continuous RH cycles, the WR stress only reaches ~20 MPa, which is much lower than that of dragline spider silk.\(^8\) Other than silk fibres, camel hair, sheep hair, and goat hair also deform and recover their shapes upon immersion in water and subsequent drying process.\(^{50, 51}\) The proposed mechanism is based on water molecules’ diffusing into the amorphous regions of these fibres and the protein’s intermolecular hydrogen bonds becoming disrupted, rearranging toward higher entropy states and leading to WR deformations.\(^{52, 53}\)

2.1.3. Microbes
Bacillus spores, which we employed to build the first engines that directly harvest evaporation energy, are another type of natural WR materials.\(^1\). \(^9\) Bacillus spores are dormant cells formed by Bacillus bacteria when they are under harsh conditions, such as lacking of nutrients. Despite their dormancy, Bacillus spores can rapidly and powerfully expand and shrink in response to changes in RH (Figure 3f).\(^1, 54, 55\) When local RH is alternated between 10% and 90%, their diameters can change by as much as ~15% within 0.1 s.\(^7\) More significantly, within 10 s, the energy exerted by Bacillus spores in a cycle of desorption and absorption of water is more than 20 MJ m\(^{-3}\), surpassing that of existing actuators and muscles (Figure 1c).\(^1\) Later, multiple types of microbes, including wild type Bacillus subtilis and genetically modified Escherichia coli (E. coli) were also found to exhibit WR behaviours.\(^{27}\) While these microbes’ WR actuation are not as powerful as spores, researchers have demonstrated smart garment prototypes by printing Bacillus subtilis on pre-patterned latex surfaces (Figure 2c,d).

2.1.4. Hybrid/Engineered Natural WR Materials
Inspired by WR actuation of these natural examples, many natural WR materials, such as wood,\(^{18, 19}\) papers,\(^{56}\) natural pollen,\(^{57}\) and cellulose nanocrystals,\(^{16}\) have been used to develop programmable actuators that reversibly and quickly change their structures in response to RH changes. Microbes’ outstanding WR properties illustrate the applicability as building blocks for ultra-high energy actuators. However, efficiently scaling up their water-responsiveness at sub-micrometre scale into a feasible engineered structure remains as a challenge. In our previously demonstrated evaporation energy harvesting devices (Figure 1g,h), we scaled up the spore’s nanoscale WR behaviours by mixing them with adhesive and depositing the mixture on micrometre thick plastic films.\(^9\) Similar to the bilayer structures found in plants, these spore based films change curvatures in response to RH changes. Patterning equally spaced spore layers on both sides of the plastic tape creates linearly expanding and contracting structures. We estimated the WR energy density of the entire strip to be approximately 17 kJ m\(^{-3}\), which is close to that of mammalian skeletal muscles, but these strips fail to match the possible energy density of the spore (by ~1000 times). In addition to directly using natural materials or mixing them into a matrix to form composites, some natural materials’ water-responsiveness could be dramatically improved by simple
processes that alter their microstructures. Our group recently reported that regenerated Bombyx (B.) mori silk can be processed with simple solvent treatments to change their secondary structures (Figure 4a-d), which dramatically increases B. mori silk’s WR energy density to 1.6 MJ m⁻³, surpassing dragline spider silk’s energy density (Figure 4e). These examples show the possibility of using biological matters for manufacturing WR actuators potentially on an industrial scale.

Figure 4. Example of regenerated biomaterials for WR actuation (a) Regenerated Bombyx (B.) mori silk fiber is deposited on thin polyimide substrates and subsequently treated with (b) water vapor or (c) methanol, which increases silks β-sheet crystallinity. (d) Schematics of silk’s microstructures, water molecules, silk and polypeptide layers in (a-c). (e) WR energy density was calculated by estimating the work done by silk layer to curve the entire bilayer structure when RH is changed between 90% and 10%. WA 45°C and WA 60°C represent water vapor treatments at 45 °C and 60 °C, and MeOH 1min and MeOH 1 h represent methanol treatments for 1 min and 1 h. Water and methanol treatments increase silk’s WR actuation energy densities, surpassing that of spider silk, mammalian skeletal muscle, and muscles of black blow fly and desert locust. Reproduced from Ref 58 with permission from John Wiley & Sons, Inc.

2.2. Synthetic WR Materials

Instead of using or engineering natural WR materials, a great deal of research focuses on synthetic approaches to develop WR materials. The relatively new field of synthetic WR materials has been growing rapidly. By mimicking natural structures, synthetic WR materials with programmed motions have spurred many potential applications, including soft robotics,⁶⁸ smart textile,⁷⁵, ⁷⁶ smart windows,²²,⁷², ⁷³ electricity generator,¹⁰, ⁷⁴-⁷⁶ and sensors.⁷⁷, ⁷⁸ Here, we review recent progresses on synthetic WR materials, especially polymers and materials with inorganic components. We discuss their WR properties, programmable motions, as well as fabrication methods.

2.2.1. Polymer Based WR Materials

The redistribution of secondary bonds between polymer molecules when water molecules diffuse in or out of polymers’ amorphous regions could lead to their WR behaviors.²¹, ⁸², ⁸³ This is analogous to the cyclic WR actuation of spider silk under alternating RH where disruption and reformation of the hydrogen bonding network between silk protein molecules cause entropy-driven uncoiling/recoiling of the spider silk.⁵² Since hydrogels are cross-linked polymers with rich hydrophilic groups that are capable of absorbing/desorbing a large amount of water while maintaining their structural integrity, many of them exhibit WR properties. Current examples include hydrogels of sodium polyacrylate (pNaAc),⁸⁰ agarose,⁷⁴, ⁷⁶ poly(ethylene glycol) (PEG),⁸⁴ acrylamide,⁸⁴ polyethylene glycol-diacylate (PEG-DA),¹⁰, ⁸³, ⁸⁵ methacrylic anhydride,⁸⁶ sodium alginates,⁸⁷ polyacrylamide xerogel,⁸² and poly(N-isopropylacrylamide)-co-acrylic acid).²¹ WR behaviours of these hydrogels are highly dependent on their hydrophilicity and crosslinking degrees. For instance, a higher degree of swelling occurs in the region with less crosslinking, as polymer molecules have less structural restriction to move.²⁰, ⁸⁰, ⁸⁷-⁸⁹ Thus, hydrogel’s WR behaviours can be easily changed by controlling their crosslinking using a variety of methods, such as UV light irradiation²⁰ and divalent ions diffusion.⁸⁰, ⁸⁷, ⁸⁹ Figure 5a describes an ‘ionprinting’ process where a copper electrode imprints Cu²⁺ ions into the anionic gel to locally reduce gel’s water-responsiveness.⁸⁰ With different patterns of imprinted ions, the anionic gel’s WR movement can be programmed in many ways (Figure 5b).²¹ Despite programmable water-responsiveness, hydrogels’ WR speeds are usually very slow. The pNaAc hydrogel in Figure 5b takes ~30 min to absorb water and complete its swelling.⁸⁰ In addition, hydrogels’ mechanical strength and WR actuation stress are relatively low. Integrating inorganic materials into hydrogels appears to improve their mechanical and WR properties. Functionalized silica nanoparticle embedded hydrogels had high mechanical strength of ~110 MPa and WR stress of ~10 MPa,⁸⁶ but it is still a challenge for these multi-components hydrogels’ mechanical and WR properties to reach that of other WR materials.

Besides hydrogels, other polymer materials, such as cellulose papers,⁵⁶, ⁹⁰-⁹² stretched nylon 6 films,⁹³ polypyrrole/polylborate film,¹⁰ α-stacked carbon nitride polymer,¹⁵ cellulose stearoyl ester films,⁹⁴ polyimide-based systems,⁹⁵, ⁹⁶ have demonstrated water-responsiveness upon exposure to RH changes or water/humidity gradient. Figure 5c presents a spontaneous locomotion of a polypyrrole/polylborate film when it is placed on a moist substrate. The film consists of polypyrrole that serves as a rigid backbone and polylborate that can easily alter its intermolecular hydrogen bonding structures when in contact with water. When responding to water gradients, such structure exerts a high contractile stress, reaching 27 MPa.¹⁰ Some hydrophobic polymers can be also modified to exhibit water responsiveness. For example, polylamide polymers that were modified by adding highly polar groups, such as ester-sulfone and carboxylic acid, showed a similar locomotion as polypyrrole/polylborate film when placed on moist surfaces.⁹⁶ Certain shape-memory polymers (SMPs), including polyester copolymer,⁹⁷ poly(lactic-co-glycolic acid),⁹⁸ poly(ethylene glycol),⁹⁹, ¹⁰⁰ polyurethane,¹⁰¹ poly(vinyl alcohol),¹⁰², ¹⁰³ epoxy,¹⁰⁴ agarose/polyacrylamide,¹⁰⁵ were also programmed to be WR. These SMPs usually require
“programming steps” which involves deforming the polymer at high temperature above its glass transition temperature (T_g) to program a temporary shape that could be maintained at a lower temperature. As water molecules penetrate into the SMPs, water disrupts physical crosslinks (usually hydrogen bonding), and causes the SMPs to return to their permanent shapes. Many of SMPs are biocompatible and can actuate upon immersion in water at an elevated temperature, which makes them great candidates for biomedical applications, including drug delivery and cell scaffolds. Recent research found that integrating cellulose nanofibres into polyurethane matrix dramatically decreases its responsive time to less than 1 min, which is possibly due to a faster water absorption facilitated by these cellulose nanofibres. Notably, many polymers’ WR movements can be extremely rapid. For example, the π-stacked carbon nitride polymer can curl up within ~50 ms when the irradiation of ultraviolet light locally heats samples and disrupts the equilibrium state. Hydrophobic nanoparticle coated cellulose stearoyl ester films starts bending within 1 s when put on moist surfaces.

Figure 5. Polymer based synthetic WR materials. (a) Crosslinking gradient within a hydrogel is created by the ‘ionoprinting’ process where a copper electrode imprts Cu²⁺ ions into the anionic gel with application of electric potential at the electrode. (b) Stresses induced by the lines ionoprinted with a copper wire anode fold a 3D gel. This 3D shape is conserved when the gel shrinks in ethanol. (c) Water gradient across polypyrrole/polyol-borate film leads to spontaneous WR locomotion when in contact with wet surface. (d) Bending of bilayer structures of PEDOT:PSS/PVDF film upon RH changes was induced by differential swelling between hygroscopic PEDOT:PSS layer and inert PVDF film. (e) Flying dragonfly and (f) blooming flower were assembled with WR PEDOT:PSS/PVDF. Plant’s bilayer-inspired design of polymer composite for predetermined WR actuation. Magnetic particles were aligned within the polymer matrix to mimic (g) pine cone’s and (h) wheat awn’s bending motion and (i) orchid tree seed pod’s twisting motion. Reproduced from ref 80, 10, 81, 44 with permission from Nature Publishing Group, The American Association for the Advancement of Science and Elsevier.

Polymer based WR materials are amenable to various established methods to change their chemical and mechanical structures to control their water-responsiveness, thus yielding advantages in programming motions for different applications. Hydrophilic groups (e.g. –OH, –NH, –O-) on polymer chains can affect the bulk polymer’s water absorption behaviours. It has been demonstrated that a higher degree of substitution of hydroxyl with stearoyl esters resulted in less curling of the cellulose stearoyl ester film in response to alternating RH. In addition to these simple methods that alter polymer’s WR properties, many researchers have resorted to biomimetic approaches to achieve various types of WR motions. Inspired by natural bilayer structures that facilitate bending, twisting, and coiling motions, polymeric materials are constructed with various bilayer structures where individual layers swell in different degrees upon RH changes, leading to varied motions as a result of inhomogeneous internal stresses. Using this bilayer structure, RH-driven dragonfly wings that bend during hydration (~4.6 s) and dehydration (~6.3 s) and an artificial flower which blooms in response to RH changes were demonstrated. Similarly, bilayer structures with electrospun polyethylene oxide (PEO) nanofibres on a stretched poly(dimethylsiloxane) (PDMS) substrate respond to RH changes within 1-10 s, mimicking the rapid closure of the Venus flytrap leaf. In another research, biomimetic structures with different alumina platelets aligned within gelatin polymer matrix achieved bending and twisting motions, emulating WR motions of pine cones, wheat awns, and orchid tree seed pods. In addition, liquid crystalline (LC) polymers demonstrated a similar programmability of WR actuation. Typically, LC polymer consists of a network of rigid mesogens and flexible linker regions. These WR LC polymers absorb water and swell along the direction perpendicular to the alignment of mesogens. By controlling the mesogens’ alignment, various
mechanical motions, including bending, folding, and curling, were successfully programmed.113-115

The bilayer structure is not always necessary to program polymers’ WR properties as the addition of stiff fillers within polymer system alone can induce inhomogeneous stresses that result in different kinds of movements. For example, 3D printed composites composed of stiff cellulose fibres embedded in soft acrylamide achieved complicated movements similar to the blooming of flowers16 and glass fibres attached to a poly(N,N-dimethylacrylamide) hydrogel base simulated the WR opening and closing of dandelion pappi.41 Using a similar concept, cellulose nanocrystals (CNCs) were integrated into liquid crystalline structures to improve their water-responsiveness and achieve designed deformation.116, 117 Moreover, integrating nanopores that promote water transport inside polymers can also be used to program various WR motions.25, 71, 118 For example, by taking advantage of inherent nanoscale molecular channels within perfluorosulfonic acid ioniomer (PFSA) films, researchers fabricated WR actuators with both bending and helical actuations.25, 71 In a similar fashion, an engineered protein, Consensus Tetratricopeptide Repeat protein (CTPR18), forms nanochannels that affect the CTPR18 film’s WR behaviours with programmed bending and twisting.118 By decreasing the length of the nanochannels by half, responsive time of the CTPR18 film was reduced by ten times.

Figure 6. 1D and 2D nanomaterials based synthetic WR materials. (a) Graphene oxide fibres with selectively reduced regions by laser irradiation generate various motions in response to RH changes.22 (b) Twisted CNT yarns with a polymer (poly(diallyldimethylammonium) chloride) sheath expand when RH is low and contract when RH is high in linear direction.119 (c) Bilayered actuators consisting of aligned CNTs on a GO substrate programs bending and twisting motions.120 (d) A film of with free standing titanium oxide nanocapillaries bends by capillary forces generated in response to RH changes.121 Reproduced from Ref 29, 119, 120, 121 with permission from Angewandte Chemie International Edition, The American Association for the Advancement of Science, American Chemical Society, and Nature Publishing Group.

2.2.2. WR Materials with Inorganic Components

1 dimensional (1D) and 2 dimensional (2D) based materials have been extensively investigated for mechanical actuation in response to thermal and electrochemical stimuli.122, 123 Recently, they have demonstrated a great potential for developing WR actuators.17, 22, 124-126 For example, graphene oxide (GO) 2D materials curve rapidly when put on a wet surface.69, 79, 127 Bilayer films comprising reduced graphene oxide (rGO)/GO bilayers generate bending motions upon alternating RH, because GO has more hydrophilic groups such as hydroxyl, epoxy and carboxyl groups than rGO, making GO layers more prone to swell.23, 70, 128 In addition, bilayer structures of GO on hydrophobic polymer films, such as polyvinylidene fluoride,130 polydimethylsiloxane,130 and polypropylene,132 have demonstrated rapid WR actuation. In another research, hydrophilic groups of selective regions of GO fibres were reduced by laser beam irradiation.29 Careful selection of laser-induced reduction of GO fibres leads to predetermined motions such as partial folding, twisting, and bending as GO absorbs more water than rGO (Figure 6a). Recent research also uses CNT to develop WR materials, which show forceful and fast WR actuation.17, 22, 119, 133, 134 Notably, when expanding and contracting in response to RH cycles, CNT hybrid twisted yarns exhibit a large tensile stroke up to 78% and an extremely high energy density of 1.8 MJ m⁻³ which is higher than actuation energy density of conventional nickel titanium (NiTi) shape memory alloy (1 MJ m⁻³) (Figure 6b).119 Since aligned CNTs tend to induce lateral swelling, the orientation of CNTs coated on the GO substrate directly programs the CNT/GO films’ movement that is analogous to plants’ WR twisting (Figure 6c).120 The WR speed for these CNT/GO films was as fast as 0.08 s which is comparable to the snapping speed of the Venus flytrap.

In addition to 1D and 2D WR based materials, materials with other inorganic components have also been engineered to exhibit water-responsiveness. For example, some hydrophilic ceramics and metals with nano-scale porous structures, including titanium oxides,121, 125 silica,136 gold,137 can actuate strongly due to water capillarity. When responding to changes in RH, a titanium oxide film with well aligned nanocapillaries exhibits extremely high energy density of 1.25 MJ m⁻³ (Figure 6d).121 A nanoporous gold film prepared by electrochemical dealloying process also shows dramatic water-responsiveness and its energy density reaches 0.15 MJ m⁻³.137 Interestingly, some metal-organic frameworks (MOFs), an important class of inorganic-organic hybrid materials with highly nanoporous structures, have shown WR properties.138-140 For example, a
specific type of MOF crystal, MIL-88B-Fe that is synthesized in dimethylformamide (DMF), changes its unit cell’s volume by almost 25% upon dispersing in water.\textsuperscript{138} Such WR behaviours of MOF crystals have been mixed with the poly(vinylidene fluoride) polymer to achieve macroscopic WR actuation, whose motion can be programmed by patterning the MOF/polymer composites using chemical etching.\textsuperscript{139, 140}

2.3. Actuation performance comparison of natural and synthetic WR materials

Despite the recent demonstrations of synthetic WR materials, to the best of our knowledge, their performance is still not as good as natural materials, and the design criteria are not well established. We summarize and compare reported values of WR strain, stress, energy density, and response time of current natural and synthetic WR materials in Table 2. While some synthetic materials, such as CNT hybrid yarn\textsuperscript{119} and titanium oxide films,\textsuperscript{121} exhibited relatively high WR energy density, most of synthetic WR materials’ performance is still far below that of natural examples such as bacterial spores.\textsuperscript{4}

<p>| Table 2. Representative examples of WR materials with promising actuation properties for energy related applications |
|--------------------------------------------------|----------------|-------------|------------------|------------------|------------------|</p>
<table>
<thead>
<tr>
<th>WR material type</th>
<th>Material description</th>
<th>WR actuation motion</th>
<th>Strain (%)</th>
<th>Stress (MPa)</th>
<th>Energy density (kJ m(^{-3}))</th>
<th>Response time (s)</th>
<th>Ref</th>
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<td>Natural</td>
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<td>1.2</td>
<td>48~54*</td>
<td>~360</td>
<td>141</td>
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<tr>
<td>Spider silk</td>
<td>Linear</td>
<td>2.5</td>
<td>~50</td>
<td>500</td>
<td>2-3</td>
<td>8</td>
<td></td>
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<tr>
<td>Bacterial spore</td>
<td>Linear</td>
<td>11.7</td>
<td>-</td>
<td>21300</td>
<td>0.1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Rat tail tendon</td>
<td>Linear</td>
<td>~5</td>
<td>~100</td>
<td>2500*</td>
<td>-</td>
<td>142</td>
<td></td>
</tr>
<tr>
<td>Engineered natural</td>
<td>Regenerated silk film</td>
<td>Bending</td>
<td>2</td>
<td>14</td>
<td>1626</td>
<td>-</td>
<td>58</td>
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<tr>
<td>Twisted degummed silk fibre</td>
<td>Linear</td>
<td>3.3</td>
<td>12.5</td>
<td>172</td>
<td>67</td>
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<td>Synthetic polymer</td>
<td>Photoactive dopant/agarose hybrid film</td>
<td>Bending</td>
<td>-</td>
<td>-</td>
<td>7.4**</td>
<td>1.2</td>
<td>75</td>
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<tr>
<td>Bioinspired polymer</td>
<td>Bending</td>
<td>2.5</td>
<td>27</td>
<td>114</td>
<td>0.1</td>
<td>10</td>
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<tr>
<td>(PEE-Ppy)</td>
<td>-</td>
<td>-</td>
<td>1250</td>
<td>625*</td>
<td>-</td>
<td>93</td>
<td></td>
</tr>
<tr>
<td>Nylon 6 film</td>
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<td>2.5</td>
<td>22</td>
<td>&lt;30</td>
<td>86</td>
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<tr>
<td>Functionalized silica embedded hydrogel fibre</td>
<td>Linear</td>
<td>12.5</td>
<td>~10</td>
<td>625*</td>
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<td></td>
</tr>
<tr>
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<td>CNT hybrid yarn</td>
<td>Linear</td>
<td>14.3</td>
<td>22.5</td>
<td>1800</td>
<td>&lt;6</td>
<td>119</td>
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<tr>
<td>Graphene fibre</td>
<td>Rotational</td>
<td>-</td>
<td>-</td>
<td>~520**</td>
<td>3.2</td>
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<tr>
<td>Titanium oxide film with nanocapillaries</td>
<td>Bending</td>
<td>~0.9</td>
<td>-</td>
<td>1250</td>
<td>&lt;10</td>
<td>121</td>
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<td>19</td>
<td>150</td>
<td>120-180</td>
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*estimated based on the reported strain and stress values ** estimated based on the reported power density and response time

2.4. Fabrication Methods

Low cost and industrial processability make synthetic WR materials, especially polymer based WR materials, promising for large-scale manufacturing. Currently, constructing a bilayer structure that contains active and passive layers is one of the most popular strategies in imparting various WR motions for applications.\textsuperscript{28, 61, 65, 67, 76, 81, 87, 106-111} In these bilayer structures, active layers mostly consisting of hydrophilic components, such as cellulose papers,\textsuperscript{65, 73} polyethylene oxide (PEO),\textsuperscript{28, 109} sodium alginate,\textsuperscript{82, 87} and chitosan,\textsuperscript{68} are adhered onto passive layers usually consisting of hydrophobic components, such as polyvinyl chloride (PVC),\textsuperscript{108} polypropylene (PP),\textsuperscript{65, 73, 146, 147} polyimide (PI),\textsuperscript{78, 148} polyvinylidene fluoride (PVDF),\textsuperscript{81, 87} and polytetrafluoroethylene (PTFE).\textsuperscript{62, 108} To fabricate these bilayer structures, one of the most widely used methods is solvent-casting, where solution with active components is poured onto passive substrates and subsequently allowed for drying to form films (Figure 7a). Using the solvent-casting method, stiff fillers can be easily incorporated into a polymer matrix to program multiple WR motions as described in the previous sections. Examples include casting agarose solution onto aligned glass fibre fillers\textsuperscript{76} and embedding magnetic field aligned metallic particles in gelatin matrix (Figure 5d-f).\textsuperscript{44} Despite the simplicity of the traditional solvent-casting method, it is usually difficult to control the uniformity of these casted films. Spin coating, another facile method to produce highly repeatable and uniform films, was used to fabricate WR bilayer structures. As shown in Figure 7b, a PEDOT:PSS/PDMS bilayer WR actuator with a well-controlled homogeneous and reproducible thickness was fabricated by spin coating.\textsuperscript{67} To fabricate this bilayer structure, a passive PDMS layer was first spin coated onto a silicon wafer, followed by air plasma treatment to enhance the adhesion between the PDMS layer and the PEDOT:PSS layer. PEDOT:PSS was then spin coated on the top of plasma-treated PDMS substrate. Finally, the PEDOT:PSS/PDMS bilayer WR actuator was laser cut into a predetermined shape and peeled off from the silicon substrate.\textsuperscript{67}
Figure 7. Examples of fabrication methods to prepare synthetic WR structures. (a) Fabrication of cellulose stearoyl esters (CSE) films by solvent-casting method where CSE solution was poured onto substrate and dried to form films. (b) Spin coated PEDOT:PSS films on a silicon wafer were laser cut into predetermined shapes. (c) Fabrication of a WR “flower” with patterned spore/adhesive composites using photolithography. (d) 3D printing of silicon beads/polydimethylsiloxane WR structures with complicated geometries. (e) Electrospinning set up consisting of a syringe pump, a high voltage supply, and a collector prepares nanofibers of polymer solution. (f) Wet spinning of alginate solution into a coagulant bath forms solidified WR fibres. Further twisting process amplifies WR actuation of the wet spun alginate fibre. Reproduced from Ref 94, 67, 143, 144, 28, 145 with permission from Nature Publishing Group, John Wiley & Sons, Inc., The American Association for the Advancement of Science, The Royal Society of Chemistry.

Other methods, including photolithography, 3D printing, and fibre spinning were also used to fabricate WR structures and program their motions. Figure 7c shows a representative example of using photolithographic method to make a pre-designed motion of a WR “flower” that consists of an active layer of spore/UV-curable adhesive composites. After mixing spores into UV-curable adhesive and depositing them on a passive polyimide substrate, UV light exposure cures spores/adhesive composites into patterns, which subsequently drive the blooming motions of the artificial flower in response to RH changes. A similar photolithographic method was used in developing WR LC polymers, where the orientation of its internal mesogens was determined by UV curing. Recent development of 3D printing with the ability to fabricate structures with complex topography from nanoscale to macroscale and to extrude various types of materials from plastics to metals offers rapid prototyping and programming of WR motions. By integrating WR components into printable polymers, several 3D printed WR architectures have demonstrated complicated and programmable WR actuation (Figure 7d). Fibre spinning is another method that has been not only used by animals, but also widely used in industry to produce fibres with precisely controlled morphology from a wide range of polymer solutions. Electrospinning is one of the spinning methods that are often used to prepare fibres at nanoscales. By using the electrospinning whose setup simply
consists of a high voltage source, a syringe pump, and a collector, researchers deposited aligned hygroscopic PEO nanofibers in different directions on passive substrates to achieve various motions (Figure 7e).\textsuperscript{28, 109} In addition to electrospinning, wet spinning process is a more industry-relevant technique to manufacture fibres. The wet spinning method was applied to fabricate WR alginate fibres by extruding sodium alginate solution through a syringe needle and solidifying the solution in a calcium chloride coagulation bath (Figure 7f).\textsuperscript{145} The wet spun alginate fibre’s diameter was well-controlled by the spinneret aperture and collecting speed of the rolling collector. Subsequently, the alginate fibres were twisted to amplify its WR actuation.\textsuperscript{145} Similarly, WR GO fibres were also produced by direct wet spinning of GO suspension (20 mg ml\textsuperscript{-1}) into methanol solution with 3 M KCl through a single-capillary spinneret.\textsuperscript{124}

### 3. Water-Responsiveness Characterization

Methods

Understanding and further improving the performance of newly developed WR materials as actuators and energy conversion components are critical for their potential energy-related applications. However, commercially available equipment has a limited capability for characterizing materials’ WR properties. By taking advantage of existing techniques used to characterize actuators and muscles, researchers have developed various methods to measure materials’ WR properties.\textsuperscript{1, 8, 9, 32, 47, 58, 143} These properties involve WR materials’ characteristics under equilibrium (steady-state) conditions, including WR strain (%), stress (MPa), energy density (KJ m\textsuperscript{-3}), and efficiency (%); and non-equilibrium (dynamic) conditions, including response time (s), specific power (W kg\textsuperscript{-1}) or power density (W m\textsuperscript{-3}), and cycle life.\textsuperscript{1, 8, 9, 32, 47, 58, 143} To promote fundamental study of materials’ WR behaviours and inspire development of new characterization methods and equipment, we briefly discuss currently available approaches to study WR materials’ both steady-state and dynamic properties.

#### 3.1. Steady-state WR Behaviours

WR actuation relies on pressure changes induced by chemical potential difference of water inside the material and its vapour phase outside. When RH changes, the system’s thermodynamic equilibrium changes from one state to another, leading to materials’ WR actuation. WR actuation stops when the whole system re-reaches its equilibrium state. The steady-state WR properties characterize WR properties under equilibrium conditions.

**Strain and stress** are general parameters to evaluate an actuator’s or muscle’s displacement normalized by its original length in the direction of actuation and the actuation force normalized by its cross-sectional area, respectively. WR strain is usually measured when the material of interest is exposed to two extreme RH conditions (~10% and ~90% RH). By varying

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Figure 8. (a) The WR strain and stress of poly(3,4-ethylenedioxythiophene)/polyvinyl alcohol/copolymer of acrylic acid and 2-acrylamido-2-methylpropanesulfonic acid was measured under dry and wet conditions with different loads.\textsuperscript{154} (b) WR stress of spider silk was measured by using a tensile testing machine with a humidity-controlled chamber. WR stress to maintain the strain of spider silk was recorded under high RH condition.\textsuperscript{47} (c) Measured radius of curvature (inset) was used to estimate the plane stresses at the spore/rubber interface and WR energy density of the spore layer.\textsuperscript{1} (d) Illustration of a thermodynamic cycle created by controlling the force and RH using an AFM.\textsuperscript{1} Reproduced from ref 154, 47, 1 with permission from American Chemical Society, Elsevier and Nature Publishing Group.
external loads applied during the strain test, WR stress can also be measured. For example, WR strain and stress of a poly(3,4-ethylenedioxythiophene)/polyvinyl alcohol/copolymer of acrylic acid and 2-acrylamido-2-methylpropanesulfonic acid film was measured under dry and wet conditions with different loads (Figure 8a). The maximum WR stress that stops the material’s WR actuation is called as blocking stress. To measure WR strain and stress, traditional tensile testing machines could be modified to be environmental-controlled. Figure 8b shows an example of how WR strain and stress of spider silk were measured using a tensile testing machine within a chamber where the RH was precisely controlled. Most of these WR strain/stress testing methods require standing alone samples with a minimum size of millimetre scales. However, many of newly developed WR materials are either fabricated into bilayer structures or too small to be tested by traditional tensile testing methods. For WR bilayer structures, WR stress can be estimated by monitoring their curvature changes at different RHs. As shown in Figure 8c, the plane stress of WR active layer at 10% RH can be estimated by equation (1):1, 58, 143

\[ \sigma_x = \frac{E_2}{\sigma(1-v^2)} \left( \frac{1}{R_x} - \frac{1}{R_y} \right) \]  

where \( \sigma_x \) is the surface stress along the direction of the observed curvature, \( E \) is Young’s modulus of the passive substrate, \( v \) is Poisson’s coefficient for the passive substrate, \( t \) is the thickness of the passive substrate and \( R_x, R_y \) are the radii of the curvature. For WR materials at nanoscale, we have developed an environmental-controlled AFM where local RH can be varied while monitoring the height change and actuation force of WR materials. By using this method, we have measured the WR strain of various nanoscale spores in response to alternating RH between ~15% to ~90%.1

**Energy density**, also known as work density, measures the amount of work done by a WR actuator in one hydration and dehydration cycle normalized by its unit volume. For the bilayer structure shown in Figure 8c, since work done by the active layer is equivalent to elastic energy stored in the curved active/passive bilayer structure, we can estimate WR materials’ actuation energy \( (U) \) by using equation (2):1, 58, 143

\[ U = \frac{(E_1 I_1 + E_2 I_2)}{R^2} \]  

where \( E_1 \) and \( E_2 \) are the Young’s modulus of the active layer and passive substrate, respectively. \( I_1 \) and \( I_2 \) are the area moment of inertia of the active layer and the passive substrate, respectively. Energy density can be subsequently calculated by dividing WR materials’ volume in the active layer.

In addition, we customized an AFM setup to measure the maximum work done by nanoscale WR materials within single hydration and dehydration cycle and developed a computer program to create the thermodynamic cycle where WR materials are subjected to periodically varying forces and RH. The thermodynamic cycle with changing forces and RH levels is composed of four stages as illustrated in Figure 8d. In stage I, the spores rest at low RH. The AFM probe is put in contact with the spores using a negligibly small force—just enough to maintain contact. In stage II, the cantilever exerts a predetermined force (loading). The spores respond by reducing their overall height. In stage III, the spores are subjected to high RH while the force is maintained constant. Spores respond to increasing RH by expanding and pushing the cantilever away from the surface. In stage IV, the force is reduced back to zero. The cycle is completed by lowering the RH to the initial level. Energy density can be determined by dividing the work done in one thermodynamic cycle to the volume contributing to the WR work.1

**Efficiency or energy conversion efficiency** is the ratio of materials’ WR actuation energy output to its energy input. Usually, energy input is the chemical potential differences of absorbed/desorbed water molecules and energy output is the mechanical work generated by the WR. Therefore, energy conversion efficiency of the WR material can be estimated from its energy density, the amount of absorbed/desorbed water, and water’s chemical potential differences. Many existing techniques can accurately measure materials’ water exchange, including dynamic vapour sorption (DVS) which was used to measure the WR silk’s mass changes at predefined RH cycles.58

**3.2. Dynamic WR Behaviours**

Materials’ WR dynamic behaviours describe their water-responsive ness when they are out of equilibrium conditions. **Response time** (or speed) of a WR material shows how fast the material re-reaches the equilibrium during WR actuation. Response time \( (r) \) of a bilayer WR material can be calculated by fitting WR bending to an exponential curve \( 1/r = a + be^{-r/t} \) using the least squares method where \( r \) is the radius of the curvature, \( a \) and \( b \) are regression coefficients, and \( t \) is the time. Figure 9a shows that, by fitting a spore/polymide bilayer structure’s curvature changes over time to the exponential curve, the response time was estimated as 4.2-14.8 s when the bilayer structure experiencing alternating RHs between 10 % and 90 %. **Power density** (specific power) that describes how fast a WR actuator can release a certain amount of energy can be estimated by dividing its energy density by its response time (Figure 9b). **Cycle life** (or lifetime) is the number of actuation that the WR material can perform until any major attenuation of its performance or failures of its structures. Figure 9c shows a negligible reduction in WR displacement of spore/polyimide bilayer structures after 1 million hydration and dehydration cycles. Another study of degummed silkworm silk fibres that were twisted to form torsional silk muscles demonstrated a reversible torsional stroke upon exposure to ultrasonically generated fog for multiple cycles in an open circulating environment (20%RH, 25 °C) (Figure 9d-e).26
4. Perspectives for Future Research

WR materials that powerfully deform during hydration and dehydration cycles have enabled various energy-related applications, including ambient RH fluctuation and natural evaporation energy harvesting.\(^{1, \ 9, \ 10, \ 15}\) artificial muscles,\(^{119}\) smart textiles,\(^{27}\) and smart building materials.\(^{18, \ 19}\) However, the use of WR materials in energy-related applications is mainly restricted by the limited knowledge about how to design high-efficiency WR actuators. Despite a great number of demonstrations, it remains difficult to understand the fundamental mechanisms of materials’ water-responsiveness and energy transfer processes, especially at molecular level. Many high-performance WR materials contain hierarchical and nanoporous structures such as bacterial spores,\(^{3}\) CNT hybrid yarns,\(^{119}\) and engineered protein films.\(^{118}\) It is very likely that, water molecules at solid-liquid interfaces in these nanopores exhibit very different physical and chemical properties compared to bulk water, which could play a crucial role in their overall energy conversion processes. Thus, to fully understand WR mechanisms, it is essential to understand water’s behaviour at the nano- and sub-micro scale, including chemical potential difference induced pressure and structural changes. Further studies on nanococonfined water within WR materials using both experimental and computational approaches could help reveal the mystery behind materials’ powerful WR behaviours.

The device design is another challenge to demonstrate the feasibility of using WR materials for practical applications. Since there is very limited commercially available engineering system that can be used to work with this new kind of WR actuators, researchers are often required to develop their own devices. For example, we have designed oscillatory and rotary engines with mechanical feedback systems to demonstrate the potential of using WR bacterial spores to directly and continuously harvest energy from natural evaporation.\(^{1,\ 9}\) However, a proper mechanical design usually requires a great amount of interdisciplinary work, which could be challenge for individual research groups. We envision that increasing collaborative efforts between scientists and engineers could dramatically speedup the development of various devices that could be used to demonstrate WR materials’ potential, and even to be commercialized.

In addition, current limitations of commercially available equipment for characterizing and fabricating WR materials also restrict their development. As discussed, some researchers have modified existing analytical techniques to test materials’ WR properties, including WR strain, stress, efficiency, responsive speed, energy density and power density, which are of great importance for energy-related applications. For examples, a tensile testing machine and an AFM that were customized to be RH-controlled can measure WR strain and stress of materials at millimetre-scale and nanoscale, respectively.\(^{1, \ 8, \ 47}\) These studies provide great examples to standardize characterization methods specifically for materials’ water-responsiveness and promote future development of commercial testing equipment. Despite excellent examples of natural and synthetic WR materials we discussed, in particular, bacterial spores,\(^{1}\) natural and regenerated silk,\(^{8, \ 58}\) and CNT based composites,\(^{119}\) efficiently scaling up materials’ water-responsiveness is also critical for practical applications. Many of current proof-of-concept demonstrations are based on millimetre-scale or centimetre-scale WR materials of film and fibre structures.\(^{10, \ 15, \ 19, \ 58, \ 119, \ 124, \ 145}\) Further studies on fabrication of WR structures by utilizing existing industrial manufacturing techniques seem promising. For instance, 3D printing that manufactures complicated geometries could be a great avenue to program and scale up materials’ water-responsiveness. We hope our review on current progresses of WR materials could inspire future studies to establish general rules for design and fabrication of high-efficiency WR materials.

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Up-to-date studies of water-responsive materials for energy-related applications are reviewed. Future research endeavours could advance scientific and technical challenges.

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