Toward High Resolution 3D Printing of Shape-Conformable Batteries via Vat Photopolymerization: Review and Perspective

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ABSTRACT High-resolution additive manufacturing offers access to the production of intricate architectures with small features that can revolutionize the fabrication of next-generation batteries. Relegated to two-dimensional sheets, commercial lithium-ion batteries consist of stacked leaflets, which are manufactured in restricted stacked or rolled geometries. By leveraging the most recent advancements of vat photopolymerization (VPP), the next-generation of shape-conformable three-dimensional batteries can be co-designed with known application requirements and provide enhanced safety and power performance based on reduced weight and dead volume. Herein, an overview of the state of the art with perspectives towards the development of electroactive photo-polymerizable resins for the direct fabrication of complete multi-material three-dimensional batteries is presented. Different approaches are described, including the formulation of composite resin through the introduction of solid electroactive particles, soluble components or metal precursors. Finally, the impact of the thermal post-processing steps on the resulting electrochemical properties of the VPP printed battery component or device is thoroughly discussed. This study paves the way towards the manufacturing of a complete high-resolution shape conformable 3D battery via VPP with enhanced power density.

INDEX TERMS Lithium-ion battery, electrodes, 3D printing, vat photopolymerization, composite.

I. INTRODUCTION

Motivated by the increasing demand of portable electronic devices and the commercialization of electric vehicles, the last decades have witnessed the development of new energy storage systems. Lithium-ion batteries (LIB) have remained the system of choice due to the promising electrochemical performances including high energy density, power density, efficiency and long cycle life. Despite the impressive commercial growth of LIB worldwide, research challenges still remain regarding the synthesis, fabrication, electrochemical performance, and safety.

In the last few years, research dedicated to LIB has been focused on the development of unconventional three-dimensional (3D) architectures. Originally initiated from the work of Long et al. [1], this focus has been motivated by the capability of 3D battery architectures to provide higher active exchange surface area and enhanced lithium-ion diffusion. Such architectures improve the electrochemical performance in terms of specific capacity, areal energy density and power density [2], [3]. Many groups have attempted to develop 3D arrangements of independent battery electrodes by inducing an electrochemical growth of nanorods/micro-tubes/post arrays onto a current collector serving as substrate followed by electrochemical plating of the active material (Fig. 1) [4]–[6]. While surface irregularities were unfortunately introduced and often resulted in short circuits (while performing the intercalation of both electrodes), these pioneering studies have established a process and geometric architecture well suited to be further improved with the design and fabrication freedom of additive manufacturing (also referred to informally as 3D printing).

Unlike conventional fabrication techniques, additive manufacturing allows the development of energy storage devices...
such as LIB based on a layer-by-layer material deposition fashion with the possibility of producing highly complex 3D geometries such as intricated gyroid design (Fig. 2a). Besides the capability of manufacturing unique 3D battery architectures for power applications, 3D printing also opens up the implementation of shape-conformable [7], [8] or structural [9] energy storage devices by reducing dead volume and weight within the final object (Fig. 2b). This represents a promising solution for producing lightweight portable devices and transportation structures including the aerospace and aeronautical sector.

Most of the early studies reported in literature relating to additively manufactured energy storage structures are focused on material extrusion (ME) due to the wide availability of inexpensive desktop printers and include research with direct ink writing [12]–[15] and fused filament fabrication [7], [8], [10], [16]–[19]. However, other additive manufacturing techniques such as vat photopolymerization (VPP), are starting to be used to construct batteries. Main advantage of VPP for battery applications is based on its improved resolution ranging from 100 μm down to 100 nm (Fig. 3). Based on the American Society for Testing and Materials, VPP includes three technologies: stereolithography using a laser to cure photoresins, digital light processing with a projected image, and two-photon polymerization (2PP) in which the intersection of two photons induces solidification. With light defining the high resolution and intricate geometric features, and a liquid feedstock providing improved surface finish, these technologies outperform in terms of resolution the less expensive thermoplastic material extrusion processes.

A summary of previous works regarding vat photopolymerization specifically employed for LIB applications is presented in this review. An overview of the different approaches employed to prepare the composite resin for the printing of electrode and electrolyte structures is discussed. Being a relatively emerging topic, an outlook of the future investigation trends and perspectives on VPP applied to LIB is comprehensively detailed.

II. VAT PHOTOPOLYMERIZATION

VPP is a process that selectively cures in a layer-by-layer fashion a vat of liquid photosensitive resin composed of a mixture of monomers, oligomers, photoinitiator, light absorbers and photosensitizers [21]. The chemistry behind the photo-polymerization process (free-radical) is rather simple: photons of a specific wavelength in the UV-range provide the required energy to the photoinitiator (Fig. 4a) so that the photocuring process propagates and builds a cross-linked polymeric network. Acrylate-based resins (containing for example poly(ethylene glycol) diacrylate (PEGDA)) are the most commonly used due to their high reactivity [20]. The process begins when the photoinitiator, for instance diphenyl (2,4,6-trimethylbenzoyl)-phosphine oxide, absorbs the photon energy and forms two free radicals. Then a radical opens a carbon-carbon bond in the PEGDA monomer and initiates a chain-reaction with available vinyl bonds until only dead-end polymer chains exist (Fig. 4b) [22], [23]. Tuning the resin properties can be achieved through the addition of a photo-absorber or a photosensitizer that respectively limit or promote the photoexcitation and curing depth.

III. VPP OF BATTERIES FROM RESINS LOADED WITH SOLID PARTICLES

Currently, no studies have reported the preparation and printability of highly loaded composite resins containing solid particles such as active materials and conductive additives to obtain composite LIB electrodes or ceramic electrolyte via VPP. Nonetheless, the formulation of composite resins containing active materials/fillers had been widely investigated (Table 1) for other applications including dentistry and electronics to confer unique properties and functionalities (including thermal stability, piezoelectricity, mechanical strength, magnetism or biocompatibility) as well as to decrease shrinkage upon polymerization. [24], [25]

As clearly stated by Tan et al. [24], filler loading, concentration and particle size must be thoroughly controlled to limit the viscosity of the resin, since it could obstruct the recoating process, and have a detrimental effect on the whole printability. This issue can nonetheless be overcome by slightly increasing the resin temperature during the printing [26], using smaller filler particles [27] or through the introduction of a dispersant and diluent [28]. Furthermore, when dealing with composite resin loaded with solid particles, their density must be considered, and viscosity of the resin must be thoroughly tuned to avoid filler sedimentation.

In this context, printing of a highly loaded composite resin with up to 53 vol% of alumina powder was reported by Hinczewski et al. [28]. In order to increase the ceramic fraction while maintaining a low viscosity and homogeneous suspension, a dispersant (azeotropic mixture of methyltetrahydrofuran and ethanol 60:40 vol. ratio) and a diluent (n-butyl acryloyloxy ethyl carbamate) were introduced, while the vat temperature was tuned. The dispersant was reported to act both by electrostatic and steric repulsion by dispersing the alumina particles in a low polar organic media. Similarly, other groups have reported printing of composite resins loaded with alumina (Fig. 5a) [29], [30], but also other ceramics such as silica [31]–[33], silicon nitride [31], boron nitride–silica [34] and zirconia [35].
It is worth mentioning that the main difficulty when considering printing a composite resin loaded with solid particles is the resulting optical properties. Filler particles must negligibly contribute to light scattering and be adequately clear to allow satisfactory curing depth ($C_d$). Derived from the Beer-Lambert law, $C_d$ can be expressed as (1):

$$C_d = D_P \times \ln\left(\frac{E_{max}}{E_c}\right)$$  

where $D_P$ is the penetration depth of light, $E_{max}$ (J/m$^2$) the exposure, and $E_c$ (J/m$^2$) the critical minimal exposure to initiate polymerization of the photocuring resin. For a loaded resin, $D_P$ is a function of the volume concentration of the powder, particle diameter and the refractive index difference between the UV-curable solution and the powder [28], [36].

In this framework, while the introduction of solid particles seems the most logical route to follow in order to produce LIB components, experiments are significantly limited due to the high viscosity of the composite resin as well as the considerably lower curing depth compared to traditional UV-curable resin containing only polymeric matrix and photoinitiator [37], [38]. Nevertheless, loading of resin with solid particles of electroactive materials is possible and has lately been considered for the preparation of other energy storage devices such as supercapacitors and capacitors. Park et al. [39] recently reported the preparation of a UV-curable composite resin with silver nanowires. Using DLP, the authors were able to print 3D hierarchical octet truss structure (Fig. 5b). After performing pyrolysis on the structure, a silver 3D array was obtained and used as a supercapacitor (Fig. 5c). Likewise, Yang et al. [40] reported the elaboration of 3D-printed capacitors via DLP based on the incorporation of Ag/Pb(Zr,Ti)O$_3$ filler particles (up to 18%vol) into the UV-curable polymer resin. In this study however, the green part (name given to the item immediately after the printing stage) was directly subjected to electrochemical testing without performing any additional thermal post-processes.

Based on these pioneering studies, no doubt that future works will demonstrate the energy storage capability of VPP printed LIB electrodes/ceramic electrolytes components containing solid particles. As thoroughly detailed hereafter, the electrochemical performances of the printed item can be evaluated at two different stages: 1) on the green part (before debinding and sintering steps); 2) on the sintered part (after these two thermal post-processing steps).

A. GREEN STATE ITEMS

The main feature of the production of functional battery components directly at the green state is that no thermal post-processing is needed as part of the manufacturing stages. The electrochemical properties of the printed green item can
| Solid particles | Photocurable resin | 3D-printing process and parameters | Reference |
|----------------|-------------------|-----------------------------------|-----------|
| α-Al₂O₃ (particle size: 0.5 μm; loading up to 53 vol%) | Monomer: di-ethoxylated bisphenol A dimethacrylate Dispersant: methylmethyketone and ethanol Diluent: n-bu-tylacryloxyethyl carbamate Photoinitiator: 2,2 dimethoxy-2-phenylaceto-phe-none | SLA Layer thickness: 200 μm | [28] |
| Al₂O₃ | Commercial resin | DLP Layer thickness: 25 μm Radiant exposure: 130 mJ/cm² | [29] |
| Al₂O₃ (particle size: 0.46 μm), SiO₂ (2.3 μm), Si₃N₄ (0.44 μm); loading between 40 and 55 vol% | Monomer: hexane diol diacrylate (HDDA) Dispersant: unknown Photoinitiator: unknown | SLA Layer thickness: 150 μm Radiant exposure: 100 to 5200 mJ/cm² | [31] |
| h-BN-SiO₂ (particle size: 1 μm; loading up to 50 vol%) | Monomers: Ethoxyalted (5) Pentaerythritol tetraacrylate; polyurethane acrylate; HDDA; Dispersant: capryl alcohol Photoinitiator: PEG300 | DLP Layer thickness: 50 μm | [34] |
| ZrO₂ (particle size: 0.2 μm; loading up to 55 vol%) | Monomer: HDDA; 1,1,1-trimethylolpropane triacrylate Dispersants: KOS110; KOS163; Solspere 17000 Photoinitiator: TPO | — | [35] |
| Silver nanowires (50 nm in diameter and 10 μm in length; loading: 13 wt%; for supercapacitors application) | Monomers: methacrylic acid; urethane tri-acrylate Photoinitiator: TPO | DLP Layer thickness: 30 μm Radiant exposure: 1350 μW/cm² Printed electrode performances: 3.01 mF/g (0.301 mF/cm²) at scan rate of 200 mV/s | [39] |
| Ag/Pb(Zr,Ti)O₃ (particle size: 3 μm; up to 18 vol%) | Commercial resin | DLP Layer thickness: 20-40 μm Printed electrode performance: 65 F/g at scan rate of 20 mV/s | [40] |
be directly characterized after printing. Nevertheless, poor battery performances (specific capacity, energy and power density) are expected at this stage due to the presence of a high quantity of non-electroactive polymer matrix, acting as dead material, and the low porosity of the printed item. Printed electrodes are thus expected to exhibit low electrical conductivities as well as poor electrolyte impregnation, both of which will result in poor specific capacity during cycling, even at low current densities. Printed current collectors at the green state would also exhibit low electronic conductivity for the same reasons stated before. It has been previously demonstrated that the printing of infill patterns, or of lattice electrodes with open porosity, could enhance the electrodes performance at the green state by increasing the available electroactive surface area and the liquid electrolyte impregnation [41]. Another possibility would be to add a water-soluble salt into the resin that would be removed after printing simply by soaking the printed green item in a solution, thus creating an intentional porosity. It is worth mentioning that maintaining the mechanical strength of the printed electrodes and collector is a critical feature to provide robust battery parts. The investigation of conductive parts in the green state could provide sturdy structures capable of yielding superior mechanical properties than their sintered counterparts (which tend to be brittle as the strength is only conferred by the consolidation of the electroactive material).

As the printability of the complete battery in one single print (or “one-shot”) via VPP still remains one of the main targets to achieve, it is important to note that having composite resins with the same polymeric matrix would improve the adherence between the various layers and components (electrodes, current collectors, electrolyte) in the green state. While multi-material printing options [42]–[44] had been widely commercialized for material extrusion [2], [45], or inkjet additive manufacturing processes, their counterparts for VPP are still scarce as only few attempts at the laboratory scale have been carried out over the last years. These attempts have generally resulted in an exponential decrease in the production rate. Inamdar et al. [46] modified a commercial SLA system and implemented the use of a rotating vat carousel system (including 4 different tanks – Fig. 6a) with a resolution of 20 µm on the z-axis. The same group later introduced more details on the construction and programming steps during operation [47]. Using commercial resins, several multi-material complex “chess rook” parts were produced via SLA (Fig. 6b). The development of multi-material micro-SLA was further investigated by Choi et al. [48], by creating a system based on a syringe pump that controlled the dispensing of the material into a single build vat. Multi-material items were successfully produced using three different resin systems (Fig. 6c and 6d). Although the process included manual intervention for changing the materials and rinsing the parts, the combination of different materials was efficiently demonstrated both within and across layers. More recently, Khattri et al. [49] reported the development of a versatile multi-material DLP system allowing printing of UV-curable resins from three different vats. In this study, one vat was filled with isopropanol solvent to ensure computer-controlled rinsing of the printed item before switching between materials.

These pioneering results on multi-material VPP had been achieved by employing classical resins formulation (only polymer matrix and photoinitiator). No doubt that future years will witness the emergence of multi-material VPP of composite materials. Such a milestone would definitely revolutionize the additive manufacturing field and pave the way towards printability of complete functional devices for diverse applications including complete energy storage devices and other electronics. Electrochemical characterization of a complete battery that would be obtained through a VPP multi-material option is certainly envisioned.

**B. SINTERED ITEMS**

While the introduction of additional thermal post-processing steps is challenging, the main motivation for sintering printed
battery components is the improved electrochemical properties in comparison with the green state counterparts. In this case the polymer matrix is removed, leaving the resulting item free of “dead material”. As a consequence, a high degree of porosity is left and the item is constituted mainly from electroactive material. This porosity can be filled with a liquid electrolyte, which would result in an accessible route for the lithium ions. Upon cycling, sintered electrodes will exhibit greater specific capacity, capacity retention over time, energy and power density, in comparison with green state items. Since battery performances are also associated with the electronic conductivity, the presence of pure poorly conductive electroactive materials such as LiFePO$_4$, LiCoO$_2$ or Li$_4$Ti$_5$O$_12$ must be coupled with the introduction of conductive carbonaceous materials. The sintering of 3D printed ceramic electrolytes and current collectors is expected to considerably increase the ionic and electronic conductivities, respectively, but at the expense of the mechanical performance. The shrinkage occurring during the sintering of a printed item with a simple geometry can be estimated with a scaling law, but is difficult to predict for non-uniform geometries where anisotropic shrinkage can occur. In this context, the resulting shape-conformable battery, originally designed to fit perfectly in the available space, might be compromised. Other issues such as delamination between layers and eventual collapse of the printed piece can be avoided by thoroughly investigating temperature rates and dwell times during both debinding and sintering steps, for example with the use of thermogravimetric analysis. The debinding is a delicate process, the item can crack, blister, or damage, leading most of the times to structural collapse when following an improperly designed thermal cycle. To circumvent these issues, long heating cycles with controlled heating ramps are often used.

Prior to performing any electrochemical characterization, the production of a complete battery in one single print (green part) using a multi-material VPP process could also be subjected to thermal treatment to enhance the resulting performance. An important stage resides in the ability of the printed multi-material structure to maintain structural integrity from the mechanical standpoint. Here, the debinding and sintering temperatures must be tuned depending on the polymeric matrix and/or electroactive materials. Indeed, performing the thermal treatment of a printed battery composed of several different components (electrodes, electrolyte, current collectors) in a single cycle is clearly a challenging task that is yet to be overcome.

### IV. VPP OF BATTERIES VIA ALTERNATIVE APPROACHES

While 3D printing LIB components from resins loaded with solid electroactive particles has not yet been demonstrated, the initial studies of 3D printed LIB components via laser and projection VPP started to emerge from 2017 using alternative approaches: 1) 3D printing of a pure polymer scaffold structure followed by the deposition or introduction of the electroactive materials, 2) soluble components added to the resin (no thermal treatment needed) for separator, gel or solid polymer electrolyte printing, and 3) battery precursors added to the resin and electroactive materials synthesized in situ during the sintering step. A summary is displayed in Table 2.

#### A. SCAFFOLD APPROACH

The initial approach for printing complex structures to be employed as scaffolds for battery components was reported by Cohen et al. [50]. The authors employed thermoplastic material extrusion and vat photopolymerization to produce complex 3D lattice structures from commercial thermoplastic filaments and UV-curable resins, respectively. On top of conductive scaffolds (polyactic acid-graphene 92:8 wt.%) printed via thermoplastic material extrusion, tri-layered arrangements of LiFePO$_4$ as cathode, LiAlO$_2$-Polyethylene oxide or Li$_{1+x}$Al$_x$Ge$_{2-x}$(PO$_4$)$_2$-Polyethylenimine membrane as separator, and Li$_4$Ti$_5$O$_12$ as anode, were built by electrophoretic deposition. The resulting micro-battery was reported to offer high reversible specific capacity and high pulse-power capability compared to commercial planar thin-film batteries. As suggested by the authors, similar work could be performed with higher resolution VPP processes.

Similarly, Zekoll et al. [51] prepared a hybrid electrolyte composed of 3D micro-channels of Li$_{1.4}$Al$_{0.4}$Ge$_{1.6}$(PO$_4$)$_3$ ceramic solid electrolyte and non-conducting polymers (epoxy polymer and polypropylene). The porous polymer framework was primarily achieved via 2PP (Fig. 7a) whereas the remaining empty space was filled with the Li$_{1.4}$Al$_{0.4}$Ge$_{1.6}$(PO$_4$)$_3$ ceramic powder. Finally, the structure was placed in a furnace to eliminate the polymer matrix and perform sintering of the Li$_{1.4}$Al$_{0.4}$Ge$_{1.6}$(PO$_4$)$_3$ 3D arrangement. Cubic, diamond-shaped and gyroidal microarchitectures were produced. Higher performances were found with the gyroid Li$_{1.4}$Al$_{0.4}$Ge$_{1.6}$(PO$_4$)$_3$ structure proving an ionic conductivity of $1.6 \times 10^{-4}$ S/cm at ambient temperature.

#### B. SOLUBLE COMPONENTS APPROACH FOR SEPARATOR, GEL, OR SOLID POLYMER ELECTROLYTE

The synthesis of resins to serve as separator, gel polymer electrolyte, ionogel, solid polymer electrolyte after printing has also been investigated [52]–[55]. This trend can be simply justified by the easier resin formulation of these components as the incorporation of solid particles is not required (in comparison to the resin formulation for electrodes or ceramic electrolytes, where solid fillers are often incorporated). Good printability of the resin is therefore facilitated as no particle sedimentation occurs. Furthermore, the viscosity of the resin can be tuned and delamination between printed layers is not prone to occur. Moreover, it is worth emphasizing that the resulting separator, gel or solid polymer electrolyte green parts (obtained just after printing) do not require any additional thermal post-process before undergoing traditional battery assembly and cycling.
| Approach          | Battery component (chemistry) | Photocurable resin composition                                                                 | 3D-printing process and parameters | Post-treatment (if any) and resulting electrochemical performances                                                                 | Reference |
|-------------------|-------------------------------|-------------------------------------------------------------------------------------------------|-----------------------------------|----------------------------------------------------------------------------------------------------------------------------------|-----------|
| Scaffold          | Solid ceramic electrolyte (Li-ion) | Commercial resin                                                                               | 2PP Layer thickness: 1 µm         | Filling of the printed lattice structure with Li$_{1.5}$Al$_3$Ge$_{1.5}$(PO$_4$)$_3$ powder followed by sintering at 900 °C; Ionic conductivity: 1.6 × 10$^{-4}$ S/cm at ambient temperature | [51]      |
|                   | Gel polymer electrolyte (Li-ion) | Monomer: PEGDA (M$_r$=575) Photoabsorber: Sudan I Photoinitiator: TPO Liquid electrolyte: 1M LiClO$_4$ in ethylene carbonate / propylene carbonate | µSLA Layer thickness: 10 µm      | GPE ionic conductivity: 4.8 × 10$^{-3}$ S/cm at room temperature; Capacity: 1.4 µAh/cm$^2$ at 2 µA during 5 cycles in a LiFePO$_4$3D printed GPE|Li$_2$Ti$_2$O$_7$ configuration | [52]      |
|                   | Soluble components             | Monomers: N,N-dimethyl acrylamide; poly(vinylidene fluoride) Crosslinker: divinyl benzene Photoabsorber: Kemsorp 11S Photoinitiator: α-ketoglutaric acid Liquid electrolyte: LiCl in ethylene glycol | SLA                               | GPE ionic conductivity: 6.5 × 10$^{-3}$ S/cm at room temperature                                                                 | [53]      |
| Gel polymer       | Ionogel (Li-ion)               | Commercial resin I onic liquids: 1-methyl-3-(4-sulfonyl)trimidazolium para-toluensulfonate; 1-methyl-1-butyldipiperidinium sulfonate para-toluensulfonate; 1-methyl-1-3-(4-sulfoisobuty1) imidazolium methylsulfonate | SLA                               | Ionogel ionic conductivity: up to 0.7 × 10$^{-4}$ S/cm at room temperature and 3.4 × 10$^{-3}$ S/cm at 90 °C | [55]      |
| electrolyte       | Solid polymer electrolyte (Li-ion) | Monomer: PEGDA (M$_r$=1000) Photoabsorber: TPO Lithium salt: lithium bis(trifluoromethanesulfonyl)imide | SLA                               | SPE ionic conductivity: 3.7 × 10$^{-3}$ S/cm at 25 °C; Capacity: 166 mAh/g at 0.1C at 50 °C and 77% capacity retention after 250 cycles in a LiFePO$_4$3D printed SPE|Li metal configuration | [54]      |
| Precursor         | Positive electrode (Li-ion)    | Monomer: PEGDA (M$_r$=757) Photoabsorber: Tartrazine Photoinitiator: lithium phenyl-2,4,6-trimethylbenzoylphosphinate Metal precursors: lithium nitrate; cobalt nitrate hexahydrate | DLP Layer thickness: 150 µm      | LiCoO$_2$ electroactive material is synthesized upon sintering at 700 °C; Capacity: 121 mAh/g at C/40 and 76% capacity retention after 100 cycles when tested vs Li metal | [65]      |
|                   | Positive electrode (Li-S)      | Commercial resin Surfactants: polyvinylpyrrolidone; polysorbat-20 Metal precursor: Li$_2$SO$_4$, H$_2$O | DLP Layer thickness: 50 µm       | Li$_2$S-C electroactive material is synthesized upon sintering at 800 °C; First cycle capacity: 310 mAh/g of active material at C/20 and 80% capacity retention after 100 cycles when tested vs Li metal | [66]      |
For the first time in 2017, Chen et al. reported the elaboration of a gel polymer electrolyte by printing a mixture (20/80 in vol%) of UV-curable resin with a 1 M LiClO₄ in an ethylene carbonate/propylene carbonate (v/v 1/1) liquid electrolyte using the µSLA process. The authors were able to print a zig-zag membrane (Fig. 7b and 7c) exhibiting an ionic conductivity of 4.8 × 10⁻³ S/cm at room temperature. Then, the complete battery was assembled by filling both sides with electrode slurries (gel polymer electrolyte-LiFePO₄ and gel polymer electrolyte-Li₄Ti₅O₁₂) and adding an aluminum foil as a current collector. The resulting micro-battery was galvanostatically tested and exhibited a capacity of 1.4 µAh/cm² at a discharging current of 2 µA for two cycles according to the authors.

Additional work was reported by Rahman et al. to tune the mechanical properties and favor the thermal stability of the resulting printed gel polymer electrolyte membrane. They reported the preparation and printability of a UV-curable poly(vinylidene fluoride) (PVDF)/N,N-dimethyl acrylamide (DMAA)-based resin comprising LiCl as a lithium salt and ethylene glycol (EG) as solvent (Fig. 7d). An ionic conductivity as high as 6.5 × 10⁻⁴ S/cm was achieved at room temperature for the optimized membrane composition (PVDF/DMAA/LiCl/EG wt.% 5/45/5/45).

Based on a similar approach, Zehbe et al. demonstrated that laser VPP can produce complex ionogels structures. In this study, the authors prepared resins composed of sulfonate-based ionic liquids (80 wt.%) (1-methyl-3-[(4-sulfobuty)]imidazolium para-toluene sulfonate, 1-methyl-1-butylpiperidiniumsulfonate para-toluensulfonate and 1-methyl-3-[(4-sulfobuty)]imidazolium methylsulfonate) mixed with a commercial UV-curable resin (20 wt.%). Upon printing, the ionogels exhibited ionic conductivities up to 0.7 × 10⁻⁴ S/cm and 3.4 × 10⁻³ S/cm at room temperature and at 90 °C, respectively. The resulting membranes were shown to be mechanically robust and thermally stable up to 200 °C.

Focused on the development of a solid polymer electrolyte via VPP, He et al. prepared a PEGDA-based resin containing phenylbis(2,4,6-trimethylbenzoyl)-phosphine oxide as photoinitiator and lithium bis(trifluoromethanesulfonyl) imide (LiTFSI) as lithium salt. A 3D-printed archimedean spiral-shaped solid polymer electrolyte exhibiting an ionic conductivity of 3.7 × 10⁻⁴ S/cm at 25 °C was obtained, and was later galvanostatically tested in an all-solid-state lithium metal battery configuration (Li|3D-solid polymer electrolyte |LiFePO₄). A capacity value of 166 mAh/g at a current density of 0.1C at 50°C and a capacity retention of 77% after 250 cycles was reported. The spiral-shaped solid polymer electrolyte was shown to greatly reinforce interfacial adhesion. A similar VPP approach could be undoubtedly transposed to other methacrylate-based solid polymer electrolytes, such as the methyl acrylate functionalized poly(D,L-Lactide)/poly(ethylene glycol) methyl ether methacrylate with LiTFSI as lithium salt and Irgacure 1173 as photoinitiator, proposed by Zaheer et al. In this study, the authors obtained a 2D film after the UV illumination step, but 3D complex geometries could theoretically be produced via VPP.

**C. PRECURSOR APPROACH: IN SITU SYNTHESIS OF THE ELECTROACTIVE MATERIAL UPON THERMAL TREATMENT**

With a view to circumvent the increased resin viscosity associated with the introduction of solid particles, alternatively, 3D printing of electrodes can be achieved by introducing soluble electroactive precursors into the resin and obtaining the electroactive material upon the sintering step. The clear advantage of this approach is that the precursor salts are mixed at the molecular level thanks to their solubility in the resin, and therefore do not promote UV light-scattering during 3D printing. The photopolymerization process is therefore more efficient and accurate compared to the photocuring of conventional powder-loaded resins. After printing, the green part must endure the thermal post-processing steps where the in situ synthesis of the electroactive material or ceramic particles occurs, to form the final self-standing item.

This approach has been optimized over the years to obtain ceramics for all kinds of applications. An example is the use of preceramic polymers such as polysiloxanes or polycarboxiloxanes, that upon thermal treatment above or at 1100°C produces SiC, SiCN, or SiOC ceramics. Similarly, Moore et al. recently demonstrated the in situ synthesis of 3D structures based on boro-phosphosilicate glasses (SiO₂–B₂O₃–P₂O₅) from a resin composed of poly(diethoxysiloxane), triethyl phosphate and trimethyl borate as soluble precursors for silicon, phosphorus and boron oxides, respectively (Fig. 8). Another example is the synthesis of...
of ultralight and ultra-strong ZrOC structures upon thermal treatment at 1200 °C of a DLP-printed structure (Fig. 9a). The resin consisted of a mixture of zirconium n-propoxide and an acrylate-based resin [62]. These examples, among others [63], [64], are a testament of the potential of electroactive oxides to be obtained following this approach.

Frequently encountered problems after sintering classical ceramic materials are non-homogeneous shrinkage, heavy mass loss, poor densification, cracking, weak mechanical performance, and undesirable surface roughness [57]. For battery materials, challenges are expected in every step of the manufacturing process. Among the expected challenges are: determining suitable precursor compounds and stoichiometries, obtaining rheological properties of the precursor resin that allow 3D printing of self-standing structures, and finding appropriate thermal post-processing conditions for the synthesis of battery materials and the elimination of non-electroactive components.

Specifically focused on LIB. Yee et al. [65] reported the synthesis of LiCoO2 via projection VPP. In this case, an aqueous PEGDA-based resin was loaded with Co(NO3)2.6H2O and LiNO3 in stoichiometric ratios, from which a hydrogel was printed and calcinated at 700 °C (Fig. 9b and 9c). SEM-energy dispersive X-ray analysis demonstrated that 7.5 atomic% of foreign elements (C, Na, P, Al, S and Si) were left after the calcination process, but the authors stated that this amount can be reduced by better controlling the thermal post-processing conditions. From this 3D structure an initial capacity of 121 mAh/g (theoretical capacity of 140 mAh/g) at C/40 in half-cell versus lithium was obtained. The capacity retention over 100 cycles at C/10 was shown to be 76%. Capacity loss can be explained by the large ohmic impedance due to the electrode thickness, lack of conductive additives, and progressive structural collapse upon cycling.

In regular ceramics, any kind of porosity must be avoided because its presence can lead to cracking during thermal post-processing. In contrast, 3D printed electrodes require a certain micro-porosity, so that adequate electrolyte impregnation is allowed. To address this issue, Yee et al. [65] worked on the tuning ratio of PEGDA-to-LiCoO2, and found that by increasing the lithium precursor, it was possible to increase the micro-porosity without excessively increasing the Ohmic impedance. Two other methods to promote micro-porosity are also mentioned. The first is to increase the surface area-to-volume ratio, in order to evacuate in a controlled manner the thermal decomposition gases. The second is to reduce the formation rate of decomposition gases through a combined temperature and vacuum protocol.

On a similar note, Saccone et al. [66] recently published the production of a robust and expansion-tolerant Li2S-C positive electrode for Li-sulfur batteries. Li2S-C was obtained from the pyrolysis at 800 °C of 3D printed electrodes via projection VPP (Fig. 9d) from an acrylate-based UV-curable resin containing Li2SO4.H2O and surfactants. Surprisingly, the structure presented promising first discharge capacity of 310 mAh/g at C/20 and a retention of 80% after 100 cycles when tested in half-cell configuration versus lithium. It is worth noting that these pioneering studies [65, 66], cycled independent 3D electrodes against a lithium foil (planar structure), resulting in 2.5D batteries [67], as the lithium-ion diffusion is relegated only to two dimensions. In all, these promising efforts demonstrated that VPP of precursor UV-curable resins for battery applications is achievable and highlighted the potential to manufacture 3D designs that were previously thought impossible.

Another important feature to consider when designing battery architectures is the layer resolution. Conventional VPP machines can go as low as few micrometers with optimized printing parameters (exposure time and brightness). Alternatively, 2PP, the highest-resolution additive manufacturing technology currently, could be used to reach details in the nanoscale. At the present time, there are no examples of battery 3D printing via 2PP. However, 2PP printing of preceramic resins followed by in situ synthesis has been reported for other applications [68]–[72]. Brigo et al. [68] obtained
dense SiOC structures with height of 100 μm and 450 nm features after calcination at 1000 °C of the 2PP printed structures. The precursor resin was a commercial acrylate siloxane loaded with bis(dimethylamino)benzophenone as a radical initiator [68]. Similarly, Greer’s group printed zinc oxide architectures from a PEGDA-based resin loaded with zinc nitrate hexahydrate [70] and nickel lattices from an acrylic-based resin containing nickel acrylate (Fig. 10) [71]. These works pave the way towards the fabrication of ceramic electrode and electrolyte 3D structures via 2PP.

V. CONCLUSION

3D printing techniques allow the development of energy storage devices such as LIB based on a layer-by-layer material deposition, offering the possibility to manufacture shape-conformable structures with intricate 3D geometries to enhance the power performance, while reducing electrochemically inactive weight and volume. VPP has the potential to revolutionize the fabrication of batteries based on the improved spatial resolution (main advantage) compared to other additive manufacturing subcategories such as material extrusion. Although VPP has not been extensively used to fabricate batteries for now, the demand for shape-conformable and high-power devices makes this topic particularly promising. The work on ceramic resin formulation and printing that has been previously reported for other applications such as dentistry or electronics, can certainly be transposed to batteries. An important research threshold still involves the composite resin development through the introduction of solid electroactive particles or by employing a precursor approach. The impact of the added fillers on the viscosity of the resin and the light path are the main two key-points that must be studied thoroughly. Finally, studying the impact of the printing parameters as well as the development and optimization of multi-material printers will undoubtedly pave the way towards the manufacturing of a complete high-resolution shape conformable 3D battery via VPP.

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FIGURE 10. (a) Two magnifications of an octet lattice structure printed via 2PP using a resin containing nickel precursors. (b) The same nickel lattice with nanoscale features obtained after sintering [71].
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