Tailor-made Functional Polymers for Energy Storage and Environmental Applications

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Abstract: CO₂ emissions into the atmosphere account for the majority of environmental challenges and its global impact in the form of climate change is well-documented. Accordingly, the development of new materials approaches to capture and convert CO₂ into value-added products is essential. Whereas the increased availability of renewable energy is curbing our reliance on fossil fuels and decreasing CO₂ emissions, the widespread adaptation of renewable energy still requires the development of high energy density batteries i.e., lithium ion batteries (LIBs). To address these energy and environmental challenges, our group has been developing porous organic polymers (POPs) with precise control over their porosity and surface chemistry for CO₂ capture, separation and conversion. To realize simultaneous CO₂ separation and conversion, we are also developing catalytically active two-dimensional membranes and POPs. In the area of LIBs, we have recognized the potential of supramolecular chemistry as a general strategy for solving the capacity-fading problem associated with high energy density electrode materials such as Li-metal, silicon and sulfur, which offer extremely high battery capacity compared to conventional LIBs. Accordingly, we have demonstrated how molecular-level design of one- and two-dimensional supramolecular polymers can be directly translated into an improved electrochemical performance in high energy density LIBs.

Keywords: Carbon capture · Energy storage · Li-ion batteries · Porous materials · Supramolecular polymers

Ali Coskun was born in Isparta, Turkey. He received his MSc and PhD degrees in chemistry at the Middle East Technical University (METU) while carrying out his research activities under the supervision of Professor Engin U. Akkaya. During his graduate studies, he developed supramolecular ion-sensing systems and investigated their application in the area of molecular logic gates. After his graduate studies, he joined the laboratory of Prof. J. Fraser Stoddart as a postdoctoral research associate at Northwestern University, where he developed dynamic metal–organic frameworks, artificial molecular machines, organic radicals and molecular electronic devices based on switchable mechanically interlocked molecules. In 2012, he started his independent career as an assistant professor at the Graduate School of EEWS and Department of Chemistry in the Korea Advanced Institute of Science and Technology (KAIST) and was promoted to the rank of Associate Professor in 2015. In 2017, he moved to University Fribourg, Switzerland as a Professor, where his research is supported by SNSF. His group develops porous organic polymers for CO₂ capture, separation and conversion, two-dimensional membranes, porous carbon materials for gas and energy storage, one- and two-dimensional supramolecular polymers for high energy density electrodes in Li-ion batteries.

1. Introduction

The consumption of fossil fuels to meet our ever-increasing energy demand is being challenged by growing concerns over anthropogenic emissions of greenhouse gases such as CO₂, which is accelerating global climate change and ocean acidification.[1] Current atmospheric levels of CO₂ average around 408 ppm, depending on the measuring location.[2] The trend, however, is a steady increase over the past century. The global carbon cycle spins off an excess of 3.3 gigatons of CO₂ per year that is unaccounted for.[3] This overflow is widely believed to be the major factor for the elevation in the atmospheric concentration of CO₂. The sources of these emissions are known to be predominantly from fossil fuel combustion. Once released into the atmosphere, it becomes much more expensive and difficult to capture CO₂ since the concentration in flue gases is about 300 times higher than it is in the air.[4] The regulation of the carbon dioxide emissions suggests the necessity to develop specific CO₂ capture technologies that can be retrofitted to existing power plants and designed into new plants with the goal to achieve 90% of CO₂ capture while limiting the increase in the cost of electricity to no more than 35%.[5] The cost analysis of carbon capture and separation (CCS) revealed that the CO₂ capture process alone accounts for 80–90% of the cost with the rest accounted for by transport and storage.[5] Therefore, the low-cost and efficient recovery of CO₂ from large emission sources is a formidable technological and scientific challenge, which has received considerable attention for several years.[6]

Presently, the most broadly implemented method is absorption using aqueous amine solutions (e.g. monoethanol amine), and this technology has been used in the natural gas industry for more than 60 years. The aqueous basic solvents selectively absorb the acidic CO₂ at ambient conditions. Once saturated, the mixture is heated up for regeneration at temperatures well above 100 °C, which itself is a quite energy intensive process and presents additional problems such as amine degradation and corrosive nature of amine solutions.[7] In this context, porous materials such as metal–organic frameworks (MOFs), zeolites, mesoporous silica, covalent organic frameworks (COFs) and porous organic polymers (POPs) have been extensively studied as possible alternatives for CO₂ capture.[4]

The separation and capture of CO₂ is only the first step in the fight
against greenhouse gas emissions. The next and the most daunting challenge is the fate of the collected gases. Safe storage in a planetary void is still heavily debated and the CO₂ market (if recycling is intended) is highly limited when compared with the massive scale of the emissions (less than 1%). Evidently, there is almost no choice but to come up with safer, economically viable products.\(^\text{[8]}\) \(^\text{[9]}\)

Ideally, the systems capable of simultaneous separation and conversion of CO₂, i.e., catalytic membranes, could reduce the overall costs of the process. Nevertheless, considering the massive scale of CO₂ emissions, the widespread utilization of renewable energy is crucial to reduce this gap. The intermittent nature of renewable energy sources, however, requires high energy density energy storage systems such as Li-ion batteries. In this direction, Li metal and silicon as anode and sulfur as cathode materials have emerged as promising candidates as they offer significantly higher capacities compared to the conventional graphite-based LIBs.\(^\text{[11]}\)

In order to tackle these energy and environmental challenges, Laboratory of Functional Organic Materials (LFOM) focuses on (1) the development of POPs and two-dimensional (2D) membranes with precise control over porosity, surface chemistry and catalytic activity to achieve CO₂ capture, separation and conversion and (2) the development of supramolecular polymers to address the capacity fading problems associated with high energy density electrode materials in Li-ion batteries (LIBs).

### 2. Porous Materials for CO₂ Capture, Separation and Conversion

POPs have emerged as promising candidates for CO₂ capture, separation and conversion applications due to their high surface areas, low-cost, physicochemical stability, and tunable surface chemistry. Notably, synthesis of POPs is highly modular and the nature of monomeric units can dictate the function of the resulting POP. The presence of micropores (<2 nm) along with heteroatoms enables selective targeting of CO₂ via dipole–quadrupole interactions. Likewise, POPs can also be engineered to have catalytic sites for CO₂ conversion. In this direction, our group has developed series of POPs, MOFs and two-dimensional membranes for CO₂ capture, separation and conversion applications with a primary focus on sustainable synthetic and materials approaches.

#### 2.1 POPs for CO₂ Capture and Separation

Our research in this area focuses on creating sustainable bottom-up synthetic strategies without using any precious metal catalysts to achieve precise control over porosity and surface chemistry of POPs to tune their affinity towards CO₂. In this direction, we developed (Fig. 2) a new class of porous materials called azo-bridged porous organic polymers (azo-COPs) for efficient CO₂ capture and separation.

Unlike any previously reported porous material, azo-COPs showed an increase in CO₂/N₂ selectivity with rising tempera-
tures, reaching their highest selectivity value of 288 at 50 °C, thus challenging state-of-the-art CO₂ separation technologies. We explained this phenomenon by a new concept we have called ‘N₂-philicity’. Recently, we have also shown that CO₂/N₂ selectivity in these POPs can be further enhanced by incorporating CO₂-philic functionalities. In addition, considering the rich chemistry of azo-benzenes, we have synthesized chemically similar azo-polymers using different polymerization routes to clearly identify the origin of N₂-philicity, which was shown to be highly dependent on particle morphology and textural properties. We note that the initial efforts in the area of POPs mostly focused on functional groups effects, thus these findings are rather important to show the impact of textural properties on the gas affinity even for the same chemical functionality.

Recently, we advanced our research efforts in this area by synthesizing (Fig. 3) ultramicroporous POPs incorporating benzo-thiazole moieties using elemental sulfur (BTAPs) in the absence of any solvent or catalyst. Considering that elemental sulfur is a by-product of natural gas purification, this line of research demonstrates the recycling of S₈ into materials for the separation of CO₂ from natural and/or landfill gas, thus opening up new directions for the utilization of elemental sulfur, which is a low-value and abundant chemical commodity, for the development of high-value polymeric materials. BTAPs showed Brunauer-Emmett-Teller (BET) surface areas up to 750 m²/g and were found to be thermally stable up to 550 °C under N₂ and 500 °C under air conditions. We have observed sulfur loadings up to 24 wt%, which naturally increases CO₂ affinity while limiting CH₄ diffusion, thus offering a very unique porous material for the purification of CH₄ from natural gas and landfill gas streams. We have also developed POPs incorporating molecular cages/macrocycles as monomeric units and showed that their intrinsic properties such as their ability to bind guest molecules in solution can be transferred into the solid-state for the affinity-based separation of complex gas mixtures. For example, the incorporation of triazine-based cages into the POPs allowed us to introduce a so-called ‘cage effect’ to control the affinity of CO₂ towards the sorbent, leading to 15 kJ mol⁻¹ increase in CO₂ binding enthalpy compared to the control polymer without cage units.

More recently, in an effort to improve mass transport kinetics for the catalytic conversion, we have also developed porous cationic polymers with hierarchical porosity – that is the presence of interconnected micro-, meso- and macropores – which showed improved conversion yields for bulky epoxide substrates. Our research efforts in this area are currently focused towards the integration of catalytically active POPs or metal nanoparticles onto two-dimensional membranes to realize simultaneous CO₂ separation and conversion. We also explored MOFs with hierarchical porosity by using porous carbon templates or porous metal oxides, in which we controlled surface functionalities to tune the strength of interfacial interactions. We primarily used ZIF-8 due to its ease of synthesis and catalytic activity for the conversion of CO₂ into cyclic carbonates. Importantly, we observed much improved catalytic activity for these composite structures.

2.3 Continuous MOF Membranes for CO₂ Separation

The synthesis of MOF membranes can be achieved by a wide variety of methods including solvent evaporation diffusion, hydro/solvothermal methods, microwave-assisted synthesis and ultrasonic methods. In general, ‘in situ’ growth or ‘seeding and secondary growth’ procedures were adopted to fabricate MOF crystals on various substrates. The presence of pinhole defects, grain boundary defects and intercrystalline cracks, however, can significantly reduce the separation performance of MOF membranes. Since the developed methods have a high methodological dependency on the types of substrates, it is hard to control the formation of cracks in large-scale production, which is one of the crucial barriers that postpone the application of MOF membranes in practical applications. In an effort to tackle this problem, we introduced (Fig. 5) the interfacial templation concept in which graphene oxide was used as a template on the anodic aluminum oxide (AAO) substrate to grow HKUST-1.

The significance of this approach originates from its broad applicability to various substrates. We were able to show the formation of a continuous and crack-free MOF membrane structure, which could also, in principle, address the brittleness of inorganic thin-film membranes based on MOFs.
frameworks (CTFs) can be used as artificial SEI layers to stabilize the surface of Li metal anodes.

### 3.1 Elemental Sulfur-mediated Polymer Synthesis for Li-S Batteries

Li-S batteries are considered as next-generation energy storage systems due to their light-weight, low-cost and high gravimetric capacity. However, the major obstacle that still remains (Fig. 6A) is the dissolution of Li-polysulfide intermediates and their shuttling during electrochemical reactions, which leads to a significant capacity decay over cycling. Moreover, the insulating nature of sulfur is also an important challenge.\[24\] Li-polysulfides can be targeted (Fig. 6B) either through Li\(^+\) (a hard acid) or sulfur anion (a soft base) by engaging various supramolecular interactions.\[25\] In this direction, we have demonstrated\[26\] (Fig. 6C) the synthesis of a CTF with chemically confined sulfur and a high sulfur content of 62 wt% under catalyst- and solvent-free reaction conditions from 1,4-dicyanobenzene and elemental sulfur. Our synthetic approach introduced a new way of preparing CTFs under environmentally benign conditions by the direct utilization of elemental sulfur. We were able to show that the homogeneous sulfur distribution due to the \textit{in situ} formation of the polymer network, and chemical sulfur impregnation within the micropores of CTF as well as the presence of hetero atoms effectively suppress the dissolution of polysulfides into the electrolyte. Furthermore, the conjugated nature of the resulting framework facilitates electron and ion transport, which leads to a high-performance lithium–sulfur battery. Notably, when compared to sulfur-doped porous carbon, we observed a significant improvement in the electrochemical performance of the Li-S cell. In order to realise sulfur contents above 80 wt%, we recently demonstrated

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**Fig. 6.** A) Typical discharge profile of Li-S batteries demonstrating the dissolution region and various redox processes involved during lithiation. B) Various supramolecular approaches to target Li-polysulfides and (C) our synthetic approach for organosulfur polymers with high sulfur contents based on elemental sulfur-mediated synthesis of covalent triazine frameworks. Figure reproduced with permission of the Publisher.\[25\]
(Fig. 6C) chemical impregnation of elemental sulfur via SN\(_{\text{Ar}}\) chemistry\(^{27}\) and C-S insertion.\(^{28}\) In particular, SN\(_{\text{Ar}}\) chemistry facilitated the dehalogenation and nucleophilic addition reactions of perfluoroaryl units with nucleophilic sulfur chains, achieving a high sulfur content of 86 wt% in the resulting CTF. The SF-CTF, exhibited a specific capacity of 1138.2 mAh g\(^{-1}\) at 0.05C, initial Coulombic efficiency of 93.1%, and capacity retention of 81.6% after 300 cycles, by utilizing homogeneously distributed sulfur within the micropores and nitrogen atoms of triazine units, thus offering high binding affinity toward lithium polysulfides.

Relatively low conductivity of CTF backbone, however, limited both sulfur mass loading and the rate performance of resulting electrodes. To address this challenge, we combined 1D charged polypyrrole with 2D CTF in the presence of sulfur to form a 3D network with high electrical and ionic conductivity as well as suitable charged docking sites on the backbone of polypyrrole for the polysulfide anions.\(^{29}\) The resulting polymer network enabled stable cycling performance for sulfur mass loadings up to 4.0 mg\(_{\text{sulfur}}\) cm\(^{-2}\).

3.2 Supramolecular Polymeric Binders for Si Anodes in Li-ion Batteries

We recognized the potential of supramolecular chemistry as a general strategy for solving the capacity-fading problem associated with silicon anodes, an approach which offers extremely high battery capacity compared to conventional graphite-based anodes. Compared to one lithium per six carbon atoms in the fully lithiated state of graphite, silicon can accommodate up to 4.4 Li atoms, giving rise to a theoretical capacity of 4200 mAh g\(^{-1}\).

This process, however, leads to a significant volume expansion (~300%), causing particle pulverization, morphology change and loss of electrical contact and an unstable solid electrolyte interface (SEI).\(^{11}\) Our research in this area focuses on the development of supramolecular polymeric binders to alleviate these challenges. We have systematically investigated various noncovalent interactions along with polymer structures – namely, linear, hyperbranched, graft and network polymers and correlated with the electrochemical performance. For example, our investigation\(^{30}\) of one-dimensional supramolecular polymeric binders incorporating Meldrum’s acid revealed that the most critical binder property for silicon anodes in lithium ion batteries is the self-healing effect facilitated by a series of noncovalent interactions such as ion–dipole\(^{31}\) and host–guest complexation.\(^{32}\) These investigations suggested the importance of covalent crosslinking while maintaining dynamic nature of supramolecular interactions. Thus, we turned our attention to mechanically interlocked molecules in the form of rotaxanes. In this direction, we demonstrated\(^{33}\) stable cycling performance for the full cell with an areal capacity of 2.88 mAh cm\(^{-2}\) for Si microparticle anodes by engaging polyrotaxane-based polymeric binders (Fig. 7).
We were able to show that the incorporation of polyrotaxane to conventional polyacrylic acid binder imparts extraordinary elasticity to the polymer network originating from the ring sliding motion of polyrotaxane and this unique binder combination keeps even pulverized silicon particles coalesced without disintegration, enabling stable cycle life for silicon microparticle anodes at commercial-level areal capacities.

3.3 Supramolecular Polymers as Artificial SEI Layers for Li-metal Anodes

The Li-metal anode, with the lowest electrochemical potential (−3.04 V vs SHE) and the highest theoretical specific capacity (3860 mAh g⁻¹), has aroused discernable attention. However, the uneven growth of Li dendrites during cycling has impeded the practical cell settings, causing an unstable SEI layer, electrolyte consumption, poor CE stability and safety risks by short-circuits. The uncontrolled Li plating and interface stability could be guided and enhanced by a liothiphilic polymer layer. In this direction, our group has introduced (Fig. 8) a 2D polymer (CTF) with rationally designed binding sites for Li⁺ ions to suppress Li-dendrite growth and improve the electrochemical performance.

Our strategy for the CTF synthesis involved the cyclotrimerization of 1,4-dicyanobenzene using LiTFSI as a catalyst under ionothermal conditions and the in situ generation and homogeneous distribution of LiF particles across the framework during synthesis. After loading on the airlaid-paper (AP), as a host to store Li, uniform Li plating was achieved over the entire electrode by the liothiphilicity of CTF originating from the triazine moieties and interface stabilization by LiF. Based on this synergistic effect, the resulting Li-metal anode exhibited ultra-stable CE at 10 mA cm⁻² for 220 cycles and long lifetime at 5 mA cm⁻² for 700 cycles, giving a useful design principle for Li-metal stabilization and expanding the application of POPs into this emerging battery technology. Our group is currently further expanding this strategy to various dynamic molecular and supramolecular polymers to effectively stabilize the Li-metal anode surface.

4. Future Perspectives

The future of porous organic polymers for CO₂ capture and conversion is quite promising. There are, however, still significant challenges present to improve their catalytic performance and cost effectiveness. In this direction, the use of POPs as porous organocatalysts or as supports for single-atom catalysis is rather interesting. Aside from CO₂, these systems can also be used as heterogeneous catalysts for various chemical transformations although their textural properties should be carefully tailored to realize hierarchical porosity for improved mass transport kinetics. Catalytic two-dimensional membranes featuring catalytically active POPs, MOFs or metal nanoparticles could enable the realization of simultaneous separation and conversion of CO₂. Since the membrane will allow constant removal of product(s) from the reaction mixture, it could, in principle, improve the conversion yields. The development of POPs incorporating organic cages/macrocycles for selective separation of complex mixtures as well as the applications of POPs in the area of LIBs are emerging research topics. In the area of lithium ion batteries, supramolecular chemistry is expected to play a major role in addressing the challenges associated with high energy density electrode materials such as sulfur, Li-metal and silicon. In addition, liquid, polymer and solid electrolytes are also expected to benefit significantly from this approach. Our research efforts to achieve these goals are currently in progress.

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