Designing a double-coated cathode with high entropy oxides by microwave-assisted hydrothermal synthesis for highly stable Li–S batteries

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ABSTRACT

Nowadays, Li–S batteries are considered as one of the most promising alternatives to Li-ion technology in the near future, thanks to their high specific capacity and their significantly lower environmental impact and production costs. Consequently, many efforts have been directed to tackle with the inherent issues that affect Li–S batteries. One of the main problems is the so-called shuttle effect, which basically entails the unwanted migration of lithium polysulfides (LiPSs) from the cathode to the anode side, causing the degradation of the cell. Here, we report an effective strategy to restrain the shuttle effect and increase the kinetics at the cathode of the lithium–sulfur (Li–S) battery. A functional layer including high entropy oxides (HEO) coated onto the sulfur cathode allows to exploit the HEOs capability as promoter catalysts for the conversion of LiPSs. Pure HEO powders are synthesized by fast, highly efficient microwave irradiation, followed by heat treatment at 930 °C. The formation of highly crystalline HEO is confirmed by X-ray diffraction analysis. The LiPSs adsorption capability of HEO is evaluated by UV–vis and X-ray photoelectron spectroscopy analyses. The effect of the HEO-coated sulfur cathode on the electrochemical performance of the Li–S battery is studied by cyclic voltammetry and galvanostatic charge/discharge. The cell with double-coated cathode delivers an initial discharge capacity of 1173 mAh/g at C/10 with 45% capacity retention over 500 cycles at C/5, approaching ~ 99% coulombic efficiency.

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Introduction

The lithium–sulfur (Li–S) battery is a promising technology for electrochemical energy storage, and its key contribution is the high gravimetric energy density that can be achieved with the sulfur cathode and the lithium anode. Based on the overall redox reaction $S_8 + 16\ Li^+ + 16\ e^- \rightarrow 8\ Li_2S$, the practical energy density of the Li–S battery is around 25–33% of the theoretical value (2600 Wh/kg), which is still 2–3 times higher than the best-performing Li-ion battery (LIB) [1]. The sulfur cathode is also a friendly option in terms of safety and sustainability and, due to its abundancy and low price, in fact the cost of the Li–S battery is estimated to be lower than 100 $/kWh, compared to 150 $/kWh for current LIBs [2]. Despite these advantages, implementation of the Li–S battery is required to overcome the poor power capability and short cycle life due to gradual capacity fading [3]. The drawbacks arise from the complex working mechanism at cell level, where the solid product of the discharge reaction ($Li_2S$) is accompanied by formation of various lithium polysulfides (LiPSs) intermediates that can be either soluble or insoluble in the organic electrolyte, depending on the length of the chain. This entails that soluble LiPSs may diffuse to lithium metal, where they are reduced to lower LiPSs chain, causing anode corrosion and low coulombic efficiency, and may diffuse back to the sulfur cathode in a process known as “shuttle mechanism.” [4] The diffusion of LiPSs induces the redistribution of insulating $Li_2S$ on the conductive carbon particles at the cathode, resulting in the passivation of the active surface and large overpotential for the sulfur redox conversion [5, 6].

To address these problems, nanostructured carbon hosts [7] combined with oxides/sulfides materials [8] have been employed to improve the electrochemical performance of Li–S batteries. At the same time, another approach consists of adding multifunctional interlayers in the traditional cell configuration, in order to reduce the shuttle effect creating a barrier able to retain LiPSs through physiochemical adsorption [9–11]. Although these strategies greatly improved the capacity and cycle life of Li–S cells, the complicated synthesis processes of some nanostructured materials prevent their upscaling for commercial applications [12]; moreover, in some cases, the integration of interlayers is still difficult in practical
continuous production and involves additional costs [13]. As an alternative approach to stabilize LiPSs within the cathode [14], that could be easily scaled in the continuous roll-to-toll process, is the two-step coating technique. In this cathode modification strategy, a second layer is coated onto the top of the dried sulfur-containing electrode [15]. As a result, a double-coating containing metal oxides [13] can both enable adsorption of the soluble LiPSs and enhance the kinetics of transformation processes of LiPSs to Li2S/Li2S2 [16, 17] increasing long-term performances of the cathode/electrolyte interface, significantly reducing the polarization of the Li-ion battery and increasing the rate of the capacity retention [28].

In the Li–S battery, the homogeneous distribution of multiple metal active sites of (Co0.2Cu0.2Mg0.2Ni0.2Zn0.2)O accelerates the redox reaction kinetics of LiPSs. First-principle calculations based on density functional theory (DFT) applied to the multicomponent HEO-Li2S6 model enlightened a decrease in both Li–O and S–Ni bond distances compared to those of monometallic NiO@Li2S6 model, suggesting a stronger affinity between the LiPSs and the multiple metal species in the HEO structure [23]. Similarly, Gao et al. [29] used ((Mg0.2Mn0.2Ni0.2Co0.2Zn0.2)Fe2O4) HEO nanofibers produced by electrospinning as catalytic host of sulfur and demonstrated fast kinetics of nucleation/growth of Li2S during cell charging. The chemical adsorption of LiPSs on HEO nanofibers was demonstrated by XPS, UV analyses and electrochemical tests confirmed their catalytic activity in the conversion reaction. In this case, the S/HEO composite was prepared by combining sulfur with the host HEO via melt infusion at 155 °C for 12 h in Ar-filled autoclave, which is still the most general synthesis technique for the sulfur composite so far. Once melted, sulfur soaks in the host pore by capillarity and then creates nanocrystals bonded to the host material during cooling. However, agglomeration of S on the external surface cannot be avoided, and this sulfur is still easily dissolved in electrolytes. This could be an issue, especially during long cycling at low C rates.

In the present work, the benefits of HEO-double-coated cathode are demonstrated. Sulfur is simply mixed with carbon black (KJB) and the binder, avoiding its confinement inside the porosity of the host matrix, which means an unfavorable condition for preventing LiPSs shuttling. By building the double-layer cathode, in which the sulfur/carbon active material is sandwiched between the aluminum current collector and the (Co0.2Cu0.2Mg0.2Ni0.2Zn0.2)O HEO layer, the capacity is greatly improved and the Li–S cell can retain almost 46% of the initial capacity after 500 cycles at moderate C/5 cycling rate.
Finally, as far as high entropy oxides are concerned, there are several processing routes to obtain them [30]. As a matter of fact, the typical solid-state synthesis [24, 31, 32] of HEO requires very long ball milling time to achieve homogeneous distribution of the different oxides precursors prior to calcination. Although the calcination step is necessary, a new method for HEO preparation has been explored here, which is efficient in saving time and energy [33, 34]. In this work, the hydrothermal synthesis route is adapted with the microwave one, since the latter requires a shorter reaction time (1 h vs. 1 day) [35]. Indeed, the traditional hydrothermal method [36, 37] offers a similar approach since it consists of two steps: a first one in which the precursors solution is heated for a long time (ranging from 5 h to 2 days) inside a steel vessel and a second step which is essentially the calcination at higher temperature (above 900 °C). Compared to this method, microwave synthesis can drastically reduce the time and the energy required to obtain the intermediate powders that are then calcined in the second step, in addition to allowing a better control over the precursor synthesis and its homogeneity thanks to a more homogenous heating process.

Microwave-assisted hydrothermal synthesis is also safer than flame pyrolysis [38] and allows higher products reproducibility [34, 34]. Therefore, it is believed that these cost-effective and easy techniques for both HEO and sulfur cathode preparations can enable to build better and cheaper Li–S batteries and open up new avenues to the use of high-entropy materials in energy conversion and storage devices.

Experimental section

\((\text{Co}_{0.2}\text{Cu}_{0.2}\text{Mg}_{0.2}\text{Ni}_{0.2}\text{Zn}_{0.2})\text{O}\) HEO synthesis

Nickel sulfate hexahydrate (NiSO\(_4\cdot6\)H\(_2\)O; Merck), copper sulfate pentahydrate (CuSO\(_4\cdot5\)H\(_2\)O; Merck), magnesium sulfate heptahydrate (MgSO\(_4\cdot7\)H\(_2\)O; Merck), cobalt sulfate heptahydrate (CoSO\(_4\cdot7\)H\(_2\)O; Merck) and zinc sulfate heptahydrate (ZnSO\(_4\cdot7\)H\(_2\)O; Merck) were used as nickel, copper, magnesium, cobalt and zinc precursors, respectively. Equimolar amounts of the metal salts were separately dissolved in deionized water to obtain 0.02 M solutions. Then, 40 mL of the aqueous salt solution (0.1 M) was added to 10 mL sodium hydroxide NaOH solution (1.2 M) and was magnetically stirred for 15 min. Subsequently, the solution was placed in the microwave oven (Milestone flexiWAVE) and exposed to microwave irradiation at 130 °C for 1 h, with a heating time of 5 min (21 °C/min). The solution was then centrifuged at 7000 rpm (by Thermo Scientific SL16 centrifuge, Thermo Fisher) to separate the precipitated powder from the solution. The powder was washed several times with deionized water and ethanol (Aldrich). After drying, the powder was treated at 930 °C, in air, for 5 h.

Structural-morphological characterization of HEO material

X-ray diffraction (XRD) analysis was carried out by a PANalytical X’Pert (Cu K\(_\alpha\) radiation) diffractometer. Data were collected with a 2D solid-state detector (PIXcel) from 10 to 90° (2θ) with a step size of 0.026° (2θ) and a wavelength of 1.54187 Å.

Field emission scanning electron microscopy (FESEM) analysis was carried out by Zeiss SUPRA™ 40 with Gemini column and Schottky field emission tip (tungsten at 1800 K). Acquisitions were made at acceleration voltage of 3 kV and working distance between 2.1 and 8.5 mm, with magnification up to 1000 kX. Transmission electron microscopy (TEM) analysis was performed by High-Resolution JEOL 300 kV.

The Brunauer–Emmett–Teller (BET) specific surface area (SSA) was determined by nitrogen physisorption at 77 K using a Micrometrics ASAP 2020 instrument. The specific surface area was calculated with the BET model in the relative pressure range of 0.07–0.30 by assuming 0.162 nm\(^2\)/molecule as the molecular area of nitrogen.

X-ray photoelectron spectroscopy (XPS) measurements were carried out using a PHI Model 5000 electron spectrometer equipped with an aluminum anode (1486 eV) monochromatic source, with a power of 25.0 W and high-resolution scan with 11.75 eV pass energy. The instrument typically operates at pressures below 5 × 10\(^{-8}\) mbar.

For LiPSs adsorption tests, a Li\(_2\)S\(_6\) solution was prepared by mixing sulfur and Li\(_2\)S at a molar ratio of 5:1 in 1,2-dimethoxyethane (DME) and 1,3-dioxolane (DOL) (1:1 by volume) for 72 h at 70 °C, under continuous stirring in argon atmosphere; all chemicals were purchased from Sigma-Aldrich. Then, the resulting brownish-red Li\(_2\)S\(_6\) solution was diluted to
1.0 mM for LiPSs adsorption test, and then, 50 mg of HEO was dispersed into such solution. The mixture was left to interact for 12 h. Then, the solid product (HEO + Li₂S₆) was filtered and dried in glove box prior to XPS analysis.

The Li₂S₆ solutions were loaded into quartz cuvette sample holders inside the argon-filled glove box, then sealed and placed in closed vials. The vials were brought out of the glove box to the Jenway 6850 double beam spectrophotometer for UV–vis analysis.

**Double-coated cathode preparation, cell assembly and electrochemical tests**

All the working electrodes were prepared by solvent tape casting method. The slurry for the “standard” S cathode (STD) was prepared using sulfur, Ketjenblack® carbon (KB, EC-300 J, AkzoNobel) and poly(vinylidene difluoride) (PVdF, Arkema) in N-methyl-2-pyrrolidinone (Sigma-Aldrich). The STD electrode composition was set at 70:20:10 by weight. Prior to casting, the slurry was ball milled at 30 Hz for 15 min, using two zirconia spheres with a radius of 0.25 cm, by a Retsch® MM40 ball miller. The slurry was then deposited on the aluminum current collector by doctor blade technique. The blade was adjusted at 200 l/m deposition by the automatic film applicator (Sheen 1133 N) with a speed of 50 mm s⁻¹. After coating, the electrode was dried at 50/150°C for 90 min. For the second layer coating, the selected ratio adopted between HEO, KB and PVDF was: 90:0:10 and 80:10:10, respectively, named as STD-HEO90 and STD-HEO80. The blade was adjusted at 200 l/m deposition. After solvent evaporation at 50°C for 90 min, disks of 1.76 cm² were punched out, vacuum dried at 40°C for 4 h (in Büchi Glass Oven B-585), and then transferred into an argon-filled dry glove box (MBrau® Labstar, H₂O and O₂ content <1 ppm) for cell assembly. In the double-coated cathode, sulfur loading was 1.0 mg/cm².

The electrodes were assembled in 2032 coin-type cells with lithium disk (Chemetall Foote Corporation, Ø 16 mm) as counter electrode and Celgard 2500 separator, (25 μm thickness, Ø 19 mm). The electrolyte was a solution of DME and DOL 1:1 (v/v) with 1.0 M lithium bis(trifluoromethanesulfonyl)imide (CF₃SO₂NLiSO₂CF₃, LiTFSI, Sigma-Aldrich) and 0.25 M lithium nitrate (LiNO₃, Sigma-Aldrich). The solution was stored in the glove box 12 h prior to use. The electrolyte amount never exceeded 10 μL per mg of sulfur. The cell cycling performances were investigated by galvanostatic discharge/charge cycling (GC) using an Arbin LBT-21084 battery tester at room temperature. GC tests were carried out in the potential range of 1.8–2.6 V versus Li⁺/Li at different current regimes. The C-rate was calculated with respect to the theoretical capacity of sulfur (1672 mAh/g). For cyclic voltammetry (CV), the electrode potential was reversibly scanned from 1.7 to 2.8 V versus Li⁺/Li at different scan rates using a Biologic 092-11/2e potentiostat/galvanostat. Electrochemical impedance spectroscopy (EIS) was performed in coin cells to analyze the time-dependent processes that occur on the standard (STD) and modified cathodes (HEO80 and HEO90), the responses were recorded before and after eight CV cycles. The impedance was measured at OCV from 10⁵ to 5×10⁻² Hz, with a 10 mV amplitude perturbation, and the CV was performed at 0.05 mV/s from 1.7 to 2.8 V.

**Results and discussion**

**Morphological characterization of HEO material**

HEO was prepared by hydrothermal-assisted microwave method, followed by thermal treatment in air; the XRD patterns of the synthesized powders before and after calcination are detailed in Fig. 1a. The presence of clear peaks indicated that at least a fraction of the powder obtained by microwave irradiation is crystalline. Two specific crystalline phases can be recognized in the sample, which are Co(OH)₂ (card no. 98-002-6763, hexagonal, marked as )) and ZnO (card no. 98–002-6170, marked as •); nevertheless, the presence of brucite Mg(OH)₂ and Ni(OH)₂ phases cannot be excluded since their XRD reflexes are partially overlapped with those associated with the identified phases. After calcination at 900°C, the XRD characteristic peaks of the hydroxide disappeared, but the XRD pattern was still multiphasic. The diffraction peaks at 20 values of 36.8, 42.7, 62.0, 74.1, 78.2 are indicative of (111), (200), (220), (311) and (222) planes of the rocksalt crystal structure (CoCuMgNiZn)O HEO (peaks marked as §), but small peaks due to hexagonal ZnO are also observed. At the calcination temperature of 930 °C, the formation of the single rock salt
structure was complete. No trace of ZnO was detected in the diffraction pattern at 930 °C.

Additionally, it is interesting to note that the relative intensity of the main peaks, (111) and (200), of the experimental spectra reported in Fig. 1a is reversed with respect to the theoretical one of the rocksalt structure: Commonly, the peak representative of the (200) plane is higher than the one associated with the (111) plane. Moreover, the XRD pattern shows an evident peak broadening of the (200), (220) and (311) reflections, but this does not happen for (111) and (222) reflections. This could be due to anisotropy in the crystallite size or to anisotropic lattice disorder, which is a recurring event in HEO materials as a result of the displacements of the cation/oxygen in the lattice with respect to their ideal position. As observed by Berardan et al. [40], the deviation from the ideal rocksalt structure mostly depends on the thermal history of the sample, and the broadening along the (200), (220) and (311) reflections can be ascribed to higher cationic site density along these crystallographic planes [35].

N₂ adsorption/desorption analysis (Fig. 1b) of HEO shows a type IV isotherm [41], with pore size distribution that highlights the existence of mesopores (Fig. 1c), with a cumulative pore volume being 0.0847 cm³/g. The BET SSA is 32.29 m²/g, which is consistent to BET values of HEO obtained via mechanical ball milling [23]. The SSA might be an additional factor in stabilizing the cell capacity through surface adsorption of LiPSs, although here it plays a minor role than the chemical interaction [42] because the value is rather low.

FESEM analysis was performed to assess the structural morphology of HEO, and the results are shown in Fig. 2. HEO is composed of aggregated grains of particles with irregular shape exhibiting a range of size from nano- to microparticles, with well-distributed small pores located at the grain boundaries (Fig. 2a and similar in morphology and size to those obtained by hydrothermal synthesis) [35]. From Fig. 2b, the average particle size distribution is within the range of 0.5–1.5 μm, although some abnormal elongated grains of larger size can be also detected.

Energy-dispersive X-ray spectroscopy (EDX) mapping of HEO showed the presence of the five elements throughout the whole structure. The EDX signals for the Kα emission energies (Fig. 3b–f) of Mg, Ni, Zn, Co and Cu appear as homogeneous distribution throughout the entire sample at the micrometer level, confirming the chemical and microstructural homogeneity. Moreover, the average atomic composition of O, Mg, Co, Ni, Cu and Zn in HEO is 49.43%, 9.43%, 9.80%, 9.19%, 9.45% and 12.7%, respectively, which is nearly equiatomic among the metals (Fig. 3h), with no particular element segregation on the surface of the sample. TEM micrographs (Fig. 3g, i) show that the as-synthesized powder presents lattice fringes, and the average distance spacing between adjacent lattice planes is

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**Figure 1**  a XRD analysis of the sample obtained by microwave irradiation at 130 °C for 1 h (black pattern), after calcination in air at 900 °C (blue) and at 930 °C (red); b N₂ adsorption/desorption isotherms of HEO and c the pore size distribution graph.
0.241 nm (Fig. 3i), which corresponds to the typical interplanar spacing of the (111) plane in (CoCuMgNiZn)O and is consistent with the XRD data.

Prior to investigating the electrochemical behavior of double-coated cathodes, some control experiments were carried out to qualitatively reveal the interaction between HEO and LiPSs. Adsorption experiments were performed by directly adding HEO particles into the Li2S6 solution; the results are depicted in Fig. S1 of the Supporting Information. The Li2S6 solution was initially dark yellow and became almost transparent by adding HEO powder after 12 h. The UV–vis adsorption spectra of the pristine Li2S6 solution show three UV absorption bands in the 310–350 nm, 400–450 nm and 550–650 nm regions. The strong absorption of $S_6^{2-}$ is detected at around 310 nm [43], while weaker bands are noticed around 420 nm ($S_4^{2-}$) and 610 nm ($S_3^{2-}$) [44]. These characteristic bands of Li2S6 disappear when the solution is exposed to HEO powder, only a weak peak in the region from 300 to 400 nm results from the interaction between HEO and Li2S6, enlightening HEO ability to adsorb LiPSs [45].

Furthermore, additional evidence of the chemical interaction between HEO and Li2S6 is provided by XPS analysis of HEO powder before and after the Li2S6 adsorption test, and these results are shown in Fig. S2 of the Supporting Information. The most relevant interactions are observed in the high-resolution spectra of Ni2p, Mg1s and Zn2p [23]. The core peak...
of Ni2p of HEO can be fitted into two spin–orbit doublets and two shake-up satellites. The symmetrical shape of the main peaks in Ni2p and the intense satellite peak at high binding energy indicate that nickel exists both as Ni2+ and Ni3+ in the HEO material [34]. After the Li2S6 adsorption test, all peaks shift of about $\pm 0.40$ eV toward higher binding energies (Fig. S2a). The shift shows that the environment around the surface of nickel is altered, indicating a chemical interaction between nickel and Li2S6 [23]. Moreover, the high-resolution spectra of Mg1s and Zn2p show similar shifts toward higher binding energy of 0.4 eV and 0.5 eV, respectively (Fig. S2c,d). These results can be attributed to the electropositive nature of magnesium and zinc, which forces electrons away from the metal core in the LiPSs environment, resulting in the increase in the binding energy [46], Figure S2b reports the high-resolution O1s spectra, which can be de-convoluted into three peaks [47]. The peak at 529.3 eV refers to lattice oxygen, arising from the ionic metal–oxygen bond. The peak at 531.3 eV is due to adsorbed oxygen species, and the peak at 532.6 eV is due to –OH from adsorbed moisture in HEO. All peaks shift 0.3 eV toward higher binding energies after the HEO is put in contact with the Li2S6. In this regard, by DFT calculations, Zheng et al. [23] postulated an interaction between oxygen species of HEO and lithium of LiPSs, due to the interfacial affinity between HEO and LiPSs. In Fig. S2b, the 0.3 eV shift in binding energy is small and the peak intensities do not change much from the O1s of pristine HEO; accordingly, a chemical bond between lithium polysulfides and oxygen is unlikely to form. To sum up, XPS analysis suggests that the major contribution to the absorption of Li2S6 comes from the multi-cation system in HEO material.

Electrochemical characterization

The accelerated LiPSs redox reaction is firstly studied by cyclic voltammetry (CV) measurements using both STD and STD + HEO90 cathodes. This electrochemical test was performed only on these two electrode compositions since the main purpose of this characterization was to investigate the effect of the double layer of HEO on the reactions and kinetics of Li–S cells. Consequently, it is clear that maximizing the percentage of HEO in the double layer allows highlighting the role of HEO in this sense.

As shown in Fig. 4, the CV profiles display the typical two pairs of redox peaks, due to the cathodic reduction of S to long-chain LiPSs (at 2.27 V) and further transformation of the long-chain LiPSs into lower-order Li2S2 and Li2S (at 2.05 V). The two anodic oxidation peaks account for the oxidation of Li2S to LiPSs and sulfur (at 2.34 and 2.38 V), respectively. As a matter of fact, the STD cathode (Fig. 4a, b) shows broader cathodic peaks with lower intensities than those of STD + HEO90 cathode, which is consistent with slower redox kinetics of LiPSs for both liquid/liquid and liquid/solid transformations. The CV curves highlight that HEOs mostly exert their influence on the conversion of short chain lithium polysulfides (Li2Sx, $1 < x < 4$) to final Li2S, since the reduction peak at 2.06 V and the oxidation peak at 2.35 V are more intense than those of the STD cathode. The onset potentials are shown in Table S1 in the Supporting Information, together with the peak potentials. The onset potentials were determined following the method proposed by Yuan et al. [48], and the differential CV curves are reported in Fig. S3 c,d in the Supporting Information. As depicted in Table S1, incorporation of HEO layer slightly increases the onset potentials of the reduction of both S and LiPSs, due to faster kinetics promoted by HEO. In particular, for STD cathode, a shift to lower potentials of the reduction peaks is observed from the 2nd cycle onward (Fig. 4b, S3a) These differences between the first and the following cycles reflect some redistribution of active sulfur in the STD to a less stable state, whereas the overlapping peak positions in the CV of STD + HEO90 indicate highly reversible electrochemical reaction (Fig. S3b).

To further probe the effect of HEO double-coated cathode on the redox kinetics of soluble intermediate LiPSs, CV measurements were performed at different scan rates. As shown in Fig. S4 in the Supporting Information, the linear relationship between the redox peak currents with the square root of scan rate involves that the diffusion process of LiPSs is the rate-determining step. The slopes of curves in Fig. S4 c,d,e are positively correlated with the corresponding Li+ diffusion [49] and the larger slope observed for the STD + HEO electrode, compared to that of the STD one, implies faster diffusion processes in the double-coated cathode. In particular, HEO enhances the transformation of the soluble Li2S4 to the insoluble Li2S (peak II). The poor capability of the STD to
capture LiPSs could be the reason for the lower diffusivity [50].

The aforementioned activity of the HEO materials in the Li–S cell is further demonstrated by galvanostatic charge/discharge at C/10 (first 3 cycles) followed by long-term galvanostatic cycling at C/5, which are used to evaluate the double-layer contribution and the effect of the layer composition. It was decided to further investigate this aspect by testing cathodes with two different HEO contents in the double layer (80% wt and 90% wt), to better understand which composition could actually maximize the benefits of HEO materials, previously confirmed by the results of the CVs, and at the same time to balance their low electronic conductivity ($10^{-8}$ S cm$^{-1}$) [27]. All potentials are intended versus Li$^+$/Li. As shown in Fig. 5, the STD cell displays an initial capacity of 778 mAh/g at C/10, while those of STD + HEO90 and STD + HEO80 cells are 1173 mAh/g and 1175 mAh/g, respectively. As seen, HEO double-layer allows to achieve higher specific capacity than the STD as a result of more efficient sulfur utilization. Addition of 10 wt % carbon in the double layer (STD + HEO80) does not affect the capacity values at C/10, which is the same in both STD + HEO80 and STD + HEO90 cells. Thus, the capacity at low current regime is entirely due to the presence of the HEO in the double-layer. After 250 cycles at C/5, the specific capacity of STD + HEO90 and STD + HEO80 is 528 mAh/g and 650 mAh/g, respectively. The capacity retention is 61% for STD + HEO90 and 69% for STD + HEO80, pertaining the value at the initial cycle at C/5 and after 250 cycles at the same current regime. The differences in the capacity retention between these cells are mainly due to the higher conductivity of the double layer in STD + HEO80 with additional 10 wt % carbon. Indeed, 16% capacity loss is observed by increasing the C rate from C/10 to C/5 in STD + HEO90 (Fig. 5c), whereas only 7% is observed in STD + HEO80 cell (Fig. 5d), which means a better distribution of the insulating sulfur on the conductive carbon in STD + HEO80 cathode that improves the performance at higher current rates. Therefore, addition of 10 wt % C in the double layer has the only effect of improving the capacity retention at C/5, which translates into higher capacity after 250 cycles. However, it is worth noting that STD + HEO90 retains 45% of its initial capacity after 500 cycles at C/5 (Fig. S5 of the Supporting Information), whereas the STD cell loses 61% of its initial capacity after only 250 cycles (Fig. 5a,b).

These results show that the STD + HEO90 maintains stable long-term cycling; moreover, the coulombic efficiency is approximately constant at 98.7% within 500 cycles at C/5, which means limited side reactions in STD + HEO90 cell.

The derivative-voltage profiles (dQ/dV vs V) at different cycle numbers at C/5 (Fig. 6) highlight that STD + HEO90 and STD + HEO80 cathodes induce higher contribution to the lower voltage reduction reaction due to the conversion of short-chain LiPSs to Li2S final product. In fact, the peaks at 2.05 V (in reduction) and 2.25 V (in oxidation) have higher intensity than those of the STD cathode, consistently to the trend observed in the CV measurements. In general, the derivative-voltage profiles comparison confirms higher process reversibility for the double-layer cathodes, explained by the active role of HEO in the LiPSs conversion. To further analyze the capacity characteristics of the double layer with respect to STD cathode, the capacity contribution from the upper-plateau discharge capacity ($Q_1$) and the lower-plateau discharge capacity ($Q_2$) were obtained and the
Q₂/Q₁ ratio versus cycle number is reported in Fig. 6d for both STD and STD + HEO90 cells. In this respect, Manthiram et al. [51] reported that the theoretical Q₂/Q₁ ratio should be equal to 3, since the theoretical capacity attributed to Q₁ is 419 mAh/g and that of Q₂ is 1256 mAh/g. In actual Li–S cells, however, shuttling effects and/or inappropriate reactions lower this ratio far below the theoretical. As can be seen in Fig. 6d the STD + HEO90 electrode shows higher Q₂/Q₁ values of ~2.3 at C/10 and ~2.1 at C/5 over 100 cycles than the STD (~2.2 at C/10 and ~1.8 at C/5), which means that the double layer with HEO exerts more efficient conversion of LiPSs and better sulfur utilization over 100 cycles at C/5.

Figure 5 a Long-term cycling performance of STD, STD + HEO90 and STD + HEO80 cells: first three cycles at C/10 followed by cycling at C/5. Capacity versus voltage plots for:

Figure S6 a-c reports the EIS responses for the pristine cathodes, and after eight CV cycles, the EIS results were fitted with the equivalent circuit shown in Fig. S6d.

For all three cathodes, a semicircle at high frequencies is observed which intersects the Z_Rs axis in Rs and then at Rₛ + Rₑ, the semicircle diameter represents the charge transfer resistances (Rₑ). For the three pristine electrodes, this value is around 36 Ω, while after eight CV cycles, it slightly increases for the double-layer electrodes, though it remains stable for the standard (within the experimental error). This increment is more noticeable with the increase of the amount of HEO in the double layer. After eight CV cycles, the change of Rₑ is associated
with the formation of more resistive cathode electrolyte interface (CEI) on the HEO-modified electrodes, and the higher increment is observed for STD + HEO90 cathode. At lower frequencies, a linear dependence between $Z_{Re}$ and $-Z_{Im}$ is observed, this frequency range contains the long constant time processes, as the ion diffusion of reactive ions and non-reactive ions which can accumulate in the electrode pores. This is fitted with a CPE (shown in the equivalent circuit as W in Fig. S6d). By comparing with the pristine cathodes, a more capacitive behavior for STD + HEO80 and STD + HEO90 electrodes is observed, as the responses have a higher slope, while the STD cathode has less capacitive behavior since the slope is lower. After eight CV cycles, the behavior becomes less capacitive and the slope at low frequencies decreases. Furthermore, there are no significant differences between STD + HEO80 and STD + HEO90. In summary, the double-layer cathode shows a $R_{ct}$ increment after cycling, that is mostly associated with a more resistive CEI due to the presence of the second layer.

Additionally, the rate capability of cells was evaluated at various rates from C/10 to 1C, as shown in Fig. 7. In this case, the STD + HEO80 cell displayed the best performance, from the initial capacity of about 1000 mAh/g at C/10 to capacity values of 930, 840 and 520 mAh/g at C/5, C/2 and 1C, respectively. The results of the rate capability highlight that negligible benefits are achieved at high C rates with STD + HEO90 cell compared to the STD one, in which 70 wt % S is simply mixed with 20 wt % C and 10% of binder by ball-milling. In fact, the voltage hysteresis between charge and discharge curves

![Figure 6](https://example.com/figure6.png)
showed high overpotential for STD + HEO90 particularly at C/2 (Fig. S7c of the Supporting Information), mostly due to the low conductivity of S ($5 \times 10^{-30}$ S cm$^{-1}$) and of HEO in the electrode. However, and unlike the STD cell, when the C rate is reduced to C/10 the voltage hysteresis of the STD + HEO90 cell abruptly decreases (Fig. S7d) and most of the reversible capacity, $\sim 900$ mAh/g, is recovered with very slow capacity rate degradation of 0.15% per cycle from 45 to 100th cycles, which confirms the synergistic affinity of multi-element in HEO to LiPSs, resulting in stable cycling performance (Fig. 7b).

Conclusions

In summary, a smart and rapid preparation of Li–S battery electrode with a HEO layer coated on the sulfur cathode has been developed, which is suitable for large-scale production and exhibits a favorable electrochemical performance and steady cyclic efficiency. In particular, when compared with the traditional melt infusion process, the double-layer approach allows to better discriminate the role of HEO, since the sulfur is simply physically mixed with the carbon black and the oxide powders. This procedure is particularly effective for long cycling performance at low C rates (C/10), while additional 10 wt % C in the double layer is required to ensure appreciable rate capability. Furthermore, a novel and simple synthesis strategy by microwave irradiation has been proposed to produce HEO materials. This experimental procedure presents several advantages when compared with others such as flame pyrolysis or typical hydrothermal route: It is safer than the former and dramatically faster than the latter. We have demonstrated the synergic contribution of the multi-elements in HEO to absorb LiPSs, i.e., one of the main issues regarding Li–S batteries, and the improvement of the electrochemical kinetics.

Overall, the material is electrochemically stable and the compositional uniformity and consistency in charge/discharge processes ensured efficient and stable Li–S cell operation, with a coulombic efficiency approximately constant at 98.7% within 500 cycles at C/5. The cycling results clearly highlight the fact that the presence of HEO is beneficial to the performances of the Li–S cell, since the cycling stability and specific capacity are higher than those exhibited by the standard sulfur electrode.

In conclusion, our findings can significantly contribute to counter the unwanted effects caused by LiPSs, bringing Li–S batteries closer to the industrial scalability also leveraging on rapid and sustainable preparation processes, such as the microwave-based one presented in this study.

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