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Article

Comparative Study of Binder Stability for Aqueous Lithium-Ion and Solid-Boosted Flow Batteries

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Abstract

The replacement of polyvinylidene fluoride (PVDF) with environmentally friendly binders offers potential advantages in the development of aqueous lithium-ion batteries (ALIBs) and flow batteries (FBs) incorporating solid charge carriers (so-called solid boosters). This study investigates the electrochemical stability of ethyl cellulose and cross-linked gluten as substitutes for PVDF in LiMn₂O₄ (LMO) cathodes for aqueous Li-ion battery electrodes and solid boosters for FBs. The millimetre-scaled solid booster beads must be easily produced on a large scale, and at the same time, their charging and discharging must be reversible over long durations under electrolyte tank conditions. The binders were tested under standardized conditions for discharge capacity and cycling stability. Our results demonstrate that ethyl cellulose and cross-linked gluten can rival the electrochemical stability of PVDF, maintaining initial discharge capacities near 100 mAh g⁻¹ at 0.2C for LMO cathodes and exhibiting reasonable capacity retention over hundreds of cycles. This work supports the feasibility of sustainable electrode processing, provides promising directions for scalable, eco-friendly electrode fabrication methods, and highlights promising binder candidates for use in aqueous energy storage systems.

Keywords: lithium-ion battery; eco-friendly binder; flow battery

1. Introduction

With the rapid advancement of renewable energy systems, the need for safe, scalable, and sustainable energy storage technologies has become more pressing. Aqueous lithium-ion batteries (ALIBs) and flow batteries (FBs) with solid boosters [1] (Figure 1) have emerged as promising candidates due to their inherent safety, environmental compatibility, and potential for low-cost manufacturing. In these systems, LiMn₂O₄ (LMO) is a widely employed cathode material thanks to its thermal stability, low cost, and high voltage plateau, although its practical capacity (~100-120 mAh g⁻¹) is somewhat lower than its theoretical value (~148 mAh g⁻¹) [2,3]. One critical yet often underappreciated component in battery electrodes is the binder, which plays a vital role in ensuring mechanical cohesion and adhesion of active materials to the current collector. In traditional lithiumion batteries, PVDF is the standard binder due to its excellent chemical stability, electrochemical inertness, and mechanical properties. However, its dependence on N-methyl-2-pyrrolidone (NMP), a toxic, high-boiling solvent, and its fluorinated polymer nature pose significant environmental and regulatory challenges. This limitation becomes especially critical for large-scale applications such as flow batteries with solid booster particles, where binder processing and environmental impact are crucial factors. With global trends moving towards the elimination of per- and polyfluoroalkyl substances (PFAS), there is urgent interest in high-performance, biodegradable binder alternatives [4-6]. In fact, some battery manufacturers have already transitioned to PFAS-free, water processable binders, eliminating the use of PVDF/NMP and improving safety [4,7,8]. In response, a wide range of environmentally friendly, water-processable, or bio-based binders have been investigated. These include cellulose derivatives, proteins, starches, and other polysaccharides. Ethyl cellulose (EC), has shown promise as an alternative to PVDF due to its processability in ethanol and favourable binding properties [7]. Similarly, plant-derived proteins like gluten, particularly in a cross-linked form, offer good mechanical flexibility and water resistance after thermal curing [9]. Both materials allow for water- or ethanol-based slurry processing, eliminating the need for NMP and enabling more sustainable manufacturing practices [10]. Recent comprehensive reviews have highlighted the key factors influencing natural polymer binder performance, including adhesion, swelling behaviour, electrochemical stability, and interactions with active materials [11-13]. These characteristics for PVDF and the two bio-binders are summarized in Table 1 [14,15]. Previous studies have shown that binder degradation can severely impact long-term cycling stability, capacity retention, and coulombic efficiency of batteries [5,11,16–18]. Despite the growing interest, comparative studies evaluating multiple binders under the same conditions remain limited. This work aims to address that gap by comparing the electrochemical stability and performance of ethyl cellulose and cross-linked gluten against PVDF using LiMn₂O₄ as the active cathode material. The goal is to identify viable alternatives that combine environmental sustainability with mechanical and electrochemical reliability for both ALIBs and solid-boosted RFBs.

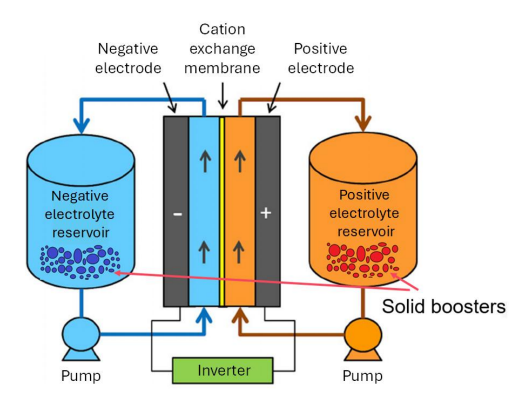


Figure 1. Schematic of a flow battery with solid boosters. Beads (e.g., LMO-based) are inserted into the electrolyte tanks to increase energy density and remain stationary during operation. Binder choice is critical - boosters must retain structural integrity during repeated cycling in aqueous media. Swelling or dissolution can lead to disintegration or system fouling. Therefore, water-stable, robust binders like ethyl cellulose and gluten are essential

Table 1. Key properties of PVDF, ethyl cellulose, and cross-linked gluten binders in aqueous battery electrodes (ALIBs/RFBs).

Property	PVDF (baseline)	Ethyl Cellulose (EC)	Cross-Linked Gluten
	Synthetic	Modified natural polymer	Natural protoin
Binder Type	fluoropolymer	(cellulose ether	Natural protein polymer (wheat
	(non-renewable)	derivative)	

			gluten), thermally cross-
			linked
Solvent Requirement	NMP (toxic, high boiling point)	Ethanol or isopropanol (benign, low toxicity)	Water-based slurry (green process); heat curing required
Mechanical Properties	Highly flexible and tough; excellent adhesion to particles Hydrophobic and	Good adhesion; moderate flexibility (sufficient for cohesion)	Flexible, rubbery network after crosslinking; strong cohesion Water-insoluble after
Water Stability	chemically inert in water (no swelling or dissolution)	Water-insoluble; minimal swelling (maintains structure)	crosslink; slight swelling possible but no dissolution
Environmental Impact	Requires toxic solvent and energy-intensive drying; not biodegradable; fluorinated (hazardous disposal)	Renewable source; solvent is low-toxic and recyclable; biodegradable over time; no fluorine	biodegradable; no
Processing Complexity	Established slurry casting with NMP, but needs strict safety controls and drying infrastructure	Simple slurry mixing in alcohol; fast drying; flammable solvent handling needed (standard lab safety)	Easiest: aqueous slurry in ambient conditions; requires thermal cross- link step (can be integrated into drying)
Electrochemical Stability	Electrochemically inert; stable in wide potential window (up to ~4.2 V vs Li ⁺ in organics)	window, no redox-active	Largely inert after curing; no significant degradation observed within water stability window (mild conditions)

2. Materials and Methods

2.1. Materials

LiMn₂O₄ (LMO, high-purity, commercial grade) was used as the active cathode material; this spinel-type lithium manganese oxide is widely employed in aqueous battery systems due to its good thermal stability, abundant raw materials, and relatively high potential plateau (~4 V vs Li/Li⁺) [3]. Carbon black (Super P, >99% C, Timcal) served as the conductive additive. The tested binders included polyvinylidene fluoride (PVDF, Solef 5130, Arkema), ethyl cellulose (EC, Sigma-Aldrich, viscosity ~10 mPa·s, 48% ethoxyl substitution), and wheat gluten (food-grade, Sigma-Aldrich) which was cross-linked in situ by heat treatment. Water (deionized, Milli-Q), ethanol (99.5%, Sigma-Aldrich) and isopropanol (IPA, 99.7%, Sigma-Aldrich) were used as solvents. In all electrode formulations, the active material fraction was fixed at 80 wt%. Binder and conductive additive contents were varied between 5–10 wt% binder and 10–15 wt% Super P (making a total of 20% binder+carbon). For example, a "5% binder" electrode contained 5% binder + 15% Super P, whereas a "10% binder" electrode contained 10% binder + 10% Super P (with LMO 80% in both cases). The electrolyte for all tests was an aqueous 1 M LiNO₃ solution (99.999%, Thermo Scientific).

2.2. Electrode Preparation and Characterization

Binder-containing slurries were prepared using appropriate solvents: NMP was used for PVDF (as per standard practice for PVDF binders), ethanol for ethyl cellulose, and deionized water for gluten. The slurry preparation for the bio-based binders followed protocols adapted from prior work on water processable electrodes [19,20]. The LMO active powder and Super P conductive carbon were mixed with the binder solution (PVDF in NMP, EC in ethanol, or gluten in water) to form a uniform

slurry. For PVDF and ethyl cellulose, the slurries were drop-cast onto the current collector (glassy carbon disk or carbon-polymer composite substrate) and dried at ambient temperature to evaporate the solvent. Gluten-based electrodes required an additional curing step: after drying the aqueous gluten slurry coating, the electrode was heated at 150 °C for 1 hour to induce thermal cross-linking of the gluten. This thermal treatment causes gluten proteins to form intermolecular bonds (e.g., disulfide bridges), yielding a water-insoluble, rubbery network that improves the binder's mechanical integrity and water resistance [9]. No additional chemical cross-linkers were used. The mass loading of active material on each electrode was kept low (on the order of 0.3–1.0 mg of LMO per cm²) to minimize diffusion limitations in aqueous testing.

Analyses and imaging were carried out using a Zeiss EVO MA15 scanning electron microscope equipped with an Oxford Instruments X-MAX energy-dispersive spectroscopy (EDS) detector. The acquired spectra were processed and quantified using Aztec software (Oxford Instruments). All measurements were conducted in variable-pressure mode to minimize surface charging of uncoated or poorly conductive samples. The accelerating voltage was set to 20 keV.

2.3. Electrochemical Measurements

Electrochemical stability was evaluated using cyclic voltammetry (CV) and galvanostatic charge-discharge cycling. A three-electrode cell (Teflon body) was assembled for testing, with Ag/AgCl (3 M KCl) as the reference electrode, a platinum wire as the counter electrode, and the prepared LMO-based electrode as the working electrode and 1 M LiNO3 as the electrolyte. All measurements were conducted at room temperature (~22 °C). CV measurements were carried out using a Gamry Reference 600+ potentiostat, typically scanning between 0.8-1.2 V vs Ag/AgCl (spanning the Mn³⁺/Mn⁴⁺ redox window of LMO in aqueous media) at 2 mV s⁻¹. These scans were used to assess the electrochemical inactivity of the binder itself and to verify that the characteristic LMO redox peaks (~0.9-1.0 V vs Ag/AgCl) appeared with each binder, indicating similar electrochemical reactions in all cases. For galvanostatic cycling, electrodes were tested in two modes: (i) a rate capability test where the discharge current was sequentially increased to evaluate performance under higher C-rates, and (ii) a long-term cycling test at a fixed current (1C rate) to assess capacity retention over hundreds of cycles. In the rate test, a LANHE battery tester (Model G340A) was used to cycle the electrodes for 6 cycles at each of several rates (0.2C, 0.5C, 1C, 2C, 5C), and then finally return to 0.2C, each segment lasting 6 cycles (total 36 cycles per test). The charge/discharge cutoff was set to a fixed capacity limit or voltage range (equivalent to ~1.0–1.2 V vs Ag/AgCl for full charge). For long-term durability, separate electrodes were continuously cycled at 1C (charge and discharge each 1 hour) for 500 cycles or until failure, using the same three-electrode configuration (with excess lithium nitrate electrolyte to avoid exhaustion of Li+). Specific capacities (mAh g-1) were calculated based on the mass of LMO active material and are reported for the discharge (lithium extraction from LMO) half-cycle.

3. Results

3.1. Cyclic Voltammetry of Electrodes

All three binder types yielded qualitatively similar cyclic voltammograms for the LMO cathodes in the aqueous electrolyte, indicating that the choice of binder did not significantly alter the fundamental electrochemical reactions of the active material. Characteristic redox peaks for the Mn³+/Mn⁴+ couple of LMO were observed around 0.9–1.0 V vs Ag/AgCl in all cases (Figure 2). There were no extra peaks or currents in the CVs that would indicate binder decomposition or parasitic side reactions within the water stability window (~0–1.23 V vs Ag/AgCl). This suggests that PVDF, ethyl cellulose, and cross-linked gluten are all sufficiently inert (after curing) under the mild potentials experienced by the LMO cathode in aqueous media. One subtle difference noted was a slightly higher internal resistance (larger peak separation) for the electrodes with 10% PVDF, consistent with its higher content of non-conductive binder. Overall, the CV results confirmed that

the alternative binders do not introduce new redox activity and that any performance differences in cycling are likely due to physical/mechanical effects rather than electrochemical side reactions.

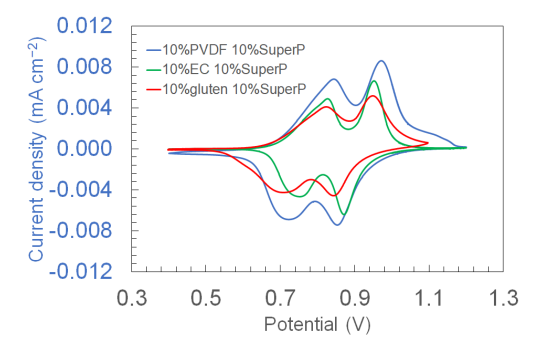


Figure 2. Cyclic voltammograms of LMO cathodes with PVDF, ethyl cellulose (EC), and cross-linked gluten binders (10% content) in 1 M LiNO₃ at scan rate of 2 mV/s All show Mn^{3+}/Mn^{4+} redox peaks, confirming electrochemical activity. The gluten-based electrode was measured in a slightly narrower potential window (~0.8–1.1 V vs Ag/AgCl) compared to PVDF and EC (~0.8–1.2 V).

3.2. Rate Performance and Initial Capacity Comparison

The initial specific capacities and rate performance of LMO cathodes were strongly influenced by both the binder type and binder content. Figure 3 presents the discharge capacity versus cycle number for each binder formulation during the multi-rate test. In this test, all electrodes were initially cycled at 0.2C (cycles 1–6), then the rate was progressively increased to 0.5C, 1C, 2C, 5C in steps (each rate for 6 cycles), and finally returned to 0.2C (last 6 cycles). This protocol allows evaluation of how capacity fades with increasing C-rate and how well it recovers when returning to a low rate. Several trends are evident in Figure 3.

PVDF is widely regarded for its chemical stability and strong binding ability; however, this study shows that excessive PVDF content (10%) severely compromises electrochemical performance. The 10% PVDF sample exhibits markedly low initial specific capacity (~38 mAh g⁻¹), which remains suppressed throughout the cycling test. In contrast, the 5% PVDF sample delivers a much higher initial capacity (~98 mAh g⁻¹) and maintains it with minimal degradation. The poor performance at 10% can be attributed to over-insulation of active material particles, leading to poor electronic conductivity and reduced ionic accessibility. Excess PVDF may also form dense films that block electrolyte infiltration and increase electrode resistance [21,22]. By contrast, the 5% PVDF electrode, with more conductive carbon available, maintained good electronic conductivity and thus much better capacity at high rates. These results highlight that while PVDF is chemically stable, its non-conductive nature means the loading must be optimized to avoid degrading the electrode's power capability.

Ethyl cellulose demonstrates promising results, particularly at 10% content, where it achieves the highest initial capacity among all samples (>100 mAh g^{-1}). This suggests that ethyl cellulose forms a favourable network structure that facilitates ion transport and maintains mechanical cohesion. Unlike PVDF, a higher content of ethyl cellulose does not seem to impede performance significantly.

However, both 5% and 10% ethyl cellulose samples exhibit a gradual capacity decline, possibly due to partial solubility or mechanical fatigue of the organic binder over extended cycles. This somewhat counterintuitive result (the higher binder content yielding better high-rate retention) could be due to the 5% EC electrode having insufficient binder to maintain conductive network integrity under stress. With only 5% EC, the electrode may be more prone to micro-cracking or particle detachment as the current increases and slight volume changes or mechanical stresses accumulate, whereas 10% EC provides a more robust network that holds the composite together at high rates. Another factor is that the 5% EC electrode had 15% carbon which boosts electronic conductivity, but if the binder is too low, the carbon and active particles might not be held in close contact during rapid cycling.

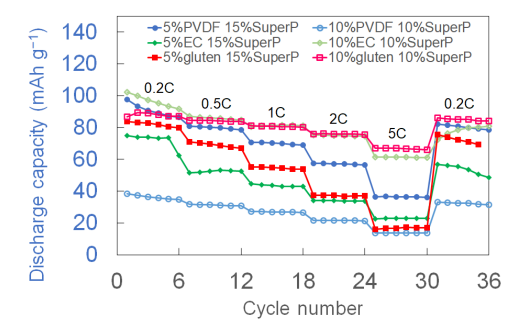


Figure 3. Discharge capacity vs. cycle number for LMO cathodes with PVDF, ethyl cellulose (EC), or gluten binders at 5% and 10% content (with Super P adjusted to 20% total). Discharge rate was increased stepwise from 0.2C to 5C, then returned to 0.2C. Lower binder content (5%) generally provided higher initial capacity and better rate performance.

Gluten-based electrodes show clear performance advantages at 10% binder content. The 10% gluten sample delivers higher initial capacity (~89 mAh g⁻¹) and significantly greater capacity retention under cycling and increasing C-rates compared to the 5% sample (~84 mAh g⁻¹). These results indicate that gluten requires a sufficiently high content to form a robust, cross-linked matrix that maintains mechanical cohesion and ensures uniform current distribution. The 5% gluten formulation, while still functional, suffers from faster capacity fading and weaker retention, likely due to insufficient binder coverage or less effective particle binding. These findings underscore that 10% gluten content is optimal within the tested range, and that binder-specific tuning is critical for bio-derived systems.

A critical component of this study involved assessing capacity retention as C-rates increased, mimicking high-power demand conditions. Among the binder types, the performance trends varied depending on the formulation. The reduced performance of 10% PVDF can be attributed to several mechanisms: (1) increased binder content dilutes active material per unit mass, (2) dense fluoropolymer films impede lithium-ion transport and reduce wettability, and (3) PVDF's non-conductive nature exacerbates internal resistance buildup. In contrast, the performance of gluten and ethyl cellulose at 10% indicates better ionic and mechanical integration without excessive impedance. For ethyl cellulose and gluten, the 10% binder content generally resulted in more stable capacities,

especially at moderate and high C-rates. These results suggest that bio-based binders may require higher content to ensure cohesion and continuous conductive networks under fast cycling conditions.

Overall, these findings underscore the importance of binder-specific optimization: while lower PVDF content enhances capacity retention, higher gluten or ethyl cellulose content may improve performance, particularly under high-rate cycling. Future work should examine binder-additive synergies, cross-linking behaviour, and long-term effects of formulation on ionic/electronic transport pathways.

3.3. Long-Term Cycling Stability

While rate performance tests the immediate capacity under stress, long-term cycling reveals how each binder sustains the electrode over many charge–discharge cycles. Figure 4 shows the discharge capacity retention over 500 galvanostatic cycles at 1C for LMO electrodes with 10% binder (PVDF, EC, or gluten; all with 10% Super P). These conditions accelerate any binder degradation or electrode deterioration, as 1C cycling for hundreds of cycles in aqueous media is a stringent test (notably, LMO itself can suffer capacity fade from Mn dissolution in aqueous electrolytes over long term).

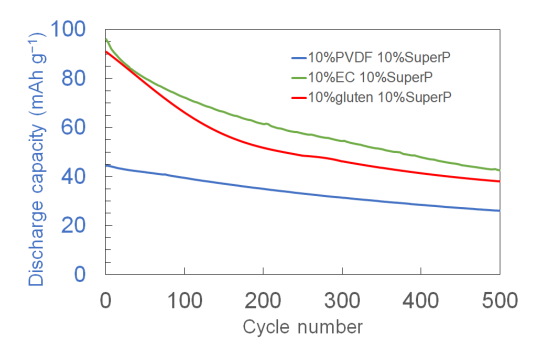


Figure 4. Long-term cycling performance of LMO electrodes at 1C in 1 M LiNO₃ with 10% binder (PVDF, ethyl cellulose (EC), or cross-linked gluten) and 10% Super P. The EC-based electrode shows the highest initial capacity (~95 mAh g⁻¹) and retains ~50 mAh g⁻¹ after 500 cycles. The gluten-based electrode starts near 90 mAh g⁻¹ but fades more rapidly, stabilizing below EC after ~300 cycles. PVDF shows the lowest capacity throughout (~45 mAh g⁻¹ initially), but with the slowest degradation rate. Results highlight the trade-off between initial performance and long-term stability across different binder systems.

As seen in Figure 4, the PVDF-based electrode displays a relatively stable but low capacity over 500 cycles. It begins around 40 mAh g⁻¹ at cycle 1 (consistent with the earlier observation of suppressed capacity for 10% PVDF at 1C) and very gradually fades to about 26 mAh g⁻¹ by cycle 500. The capacity decay is mild and linear, indicating that PVDF itself remains chemically and mechanically intact; the loss is likely due to LMO's known failure modes (e.g., Mn leaching, or electrolyte/interface changes) rather than the binder. This aligns with PVDF's reputation for excellent inertness and mechanical resilience – it imposes performance limitations (low initial capacity) but does not dramatically break down over time.

The ethyl cellulose 10% electrode starts with a much higher discharge capacity (around 100 mAh g⁻¹ at cycle 1, thanks to EC's low initial impedance). However, it exhibits a more pronounced fade over the 500 cycles. By 100 cycles, the EC electrode's capacity drops to ~80 mAh g⁻¹; by 300 cycles, it is around ~60 mAh g^{-1} ; and by the end of 500 cycles, it approaches ~50 mAh g^{-1} . Despite this decline, it still slightly surpasses the PVDF electrode's capacity at the end of test. The more significant fading in the EC-bound electrode could stem from a gradual deterioration of the binder's binding ability – for instance, slow swelling, mechanical fatigue (micro-cracks forming under repeated strain), or minor dissolution of any un-crosslinked polymer chains. Additionally, because the EC electrode started at a higher capacity, it may be more susceptible to LMO's intrinsic issues (like Mn dissolution) which are exacerbated at higher utilization levels. Nonetheless, the EC binder maintained the electrode cohesion well enough that catastrophic failure was avoided; the capacity decay flattened out in later cycles, suggesting that after an initial drop, the system stabilized. This result is promising, because even without any special additives or optimizations, the EC binder allowed for a 2-3 times higher capacity than PVDF in early cycles and still comparable capacity after 500 cycles. Minor improvements (such as cross-linking EC or blending with a co-binder) could further improve its stability.

The cross-linked gluten binder showed a mixed outcome. Initially, the 10% gluten electrode matched the ethyl cellulose's high capacity (~90 mAh g^{-1}) in the first few cycles, underlining that it provides excellent initial performance (no significant kinetic barriers). It decreases exponentially in the 500 cycles, dropping to ~45 mAh g^{-1} by the end. While it performs well initially, its long-term durability in aqueous environment needs improvement. It is possible that optimizing the curing process (temperature/time) or adding a secondary cross-linker could substantially improve longevity. Indeed, previous studies with modified or co-crosslinked biopolymer binders have achieved hundreds of stable cycles [23,24], so there is room to refine the gluten system.

In summary, for 10% binder long-cycle tests, PVDF provides lower capacity but rock-solid stability, ethyl cellulose provides high capacity but with a steady fading that still retains useful capacity by 500 cycles, and gluten provides high capacity initially but suffered an exponential loss during 500 cycles, indicating the need for further binder optimization. The fact that EC and gluten allowed much higher capacity utilization of LMO initially is encouraging for applications where energy density is paramount, but ensuring long-term stability will be key for these alternatives to truly replace PVDF in commercial systems. It should be noted that in a full cell or practical system, cycle life might be limited by other factors (e.g., the counter electrode or electrolyte); nonetheless, improving binder longevity (perhaps through composite binders or additives) is an important direction for future work.

3.4. Electrode morphology studies

Scanning electron microscopy studies were carried out for the electrode that contained 10% gluten. The SEM pictures show how uniformly the binder covers particles and how the microstructure evolves after cycling (Figure 5).

Elemental analysis by EDS provided postmortem insights into binder and electrode degradation, complementing the morphological observations obtained by SEM. This analysis is particularly valuable when evaluating novel or bio-based binders such as gluten. Since LMO is known to undergo gradual Mn dissolution in aqueous electrolytes—especially during prolonged cycling—chemical changes at the electrode surface can serve as indirect indicators of binder performance. A binder that forms a protective coating around the active material may help limit electrolyte penetration and thereby reduce Mn leaching.

Table 2 summarizes semi-quantitative EDS results averaged over multiple spectra collected from distinct regions of the electrode surface in both fresh and post-cycled LMO electrodes containing 10% gluten binder (with Super P). Although the EDS maps and spectra showed heterogeneity, the

averaging approach allows for the identification of meaningful comparative trends. After electrochemical cycling, a reduction in Mn signal was observed (from 42 to 34 wt.%), likely indicating Mn dissolution or redistribution. Concurrently, an increase in carbon content (from 24 to 33 wt.%) may reflect binder degradation, Mn loss, and increased exposure of the conductive carbon (Super P).

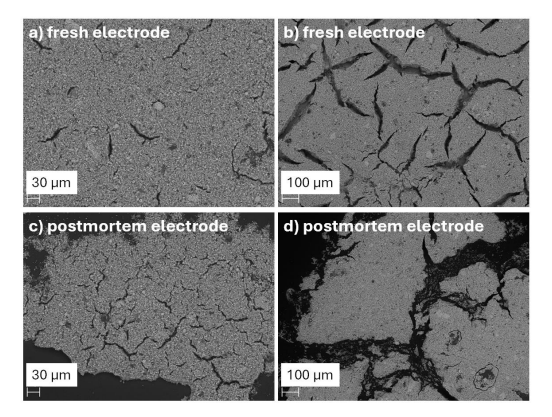


Figure 5. SEM images of LMO electrodes with 10% gluten binder: (a, b) fresh electrode at two magnifications (30 μ m and 100 μ m scale), showing a relatively uniform surface with moderate cracking; (c, d) postmortem electrode after 500 cycles, revealing increased surface fragmentation, deep cracking, and signs of binder degradation and delamination.

Table 2. Averaged EDS results for LMO electrodes with 10% gluten binder before and after 500 cycles. Values represent the mean elemental composition (wt.%) from multiple regions across the electrode surface. These data provide a comparative overview and differ from the individual EDS spectra shown in Figure 6, which illustrate local variations.

Element	Fresh electrode	Post mortem electrode
Manganese (Mn)	42	34
Carbon (C)	24	33
Sulphur (S)	0.09	0.01
Nitrogen (N)	3.1	2.4

Notably, the sulfur signal—originating from sulfur-containing amino acids in gluten—dropped significantly (from 0.09 to 0.01 wt.%), suggesting considerable binder degradation, particularly through disruption of disulfide cross-links. A decline in nitrogen content (from 3.1 to 2.4 wt.%) was also observed, which may indicate denaturation, leaching, or electrochemical decomposition of the protein matrix. A similar pattern is visible in the area-averaged SEM-EDS spectra shown in Figure 6, supporting these findings.

In areas exhibiting severe degradation, sulfur was no longer detectable in the EDS surface maps. The spatial variability of sulfur and nitrogen signals is consistent with the inherently inhomogeneous distribution of biopolymer binders. These findings underscore the susceptibility of cross-linked gluten to degradation under prolonged aqueous cycling and suggest that further optimization—such

as improved chemical cross-linking or protective surface coatings—may be required to enhance long-term binder stability.

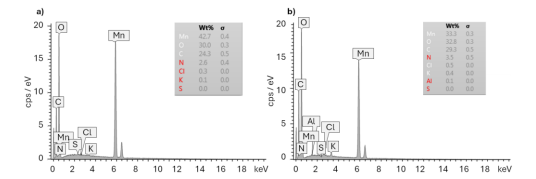


Figure 6. SEM-EDS area-averaged spectra of LMO electrodes with 10% gluten binder a) before and b) after 500 cycles in aqueous electrolyte. Larger-area scans highlight changes in surface composition, including reduced Mn and S signals post-cycling, indicating active material loss and binder degradation.

4. Discussion

Mechanical and Interfacial Considerations

The performance of electrode binders in lithium-ion batteries is governed not only by their chemical and electrochemical stability, but also by their influence on mechanical cohesion, ionic transport, and electrode morphology. This study compared three types of binders - polyvinylidene fluoride (PVDF), ethyl cellulose, and gluten—each tested at 5% and 10% weight content in the electrode. The results reveal significant differences in initial specific capacity, long-term cycling behavior, and capacity retention under increasing C-rates. The above results illustrate that binder selection impacts not only environmental sustainability but also cell performance metrics. It is therefore useful to discuss the mechanical and interfacial properties of these binders in the context of electrode function.

PVDF is well-known for its excellent mechanical strength and flexibility, which translates into strong adhesion of active material particles to each other and to the current collector. PVDF's ductile, non-reactive fluoropolymer structure imparts robust cohesion, allowing electrodes to endure volume changes and stresses without cracking or delamination [25]. This partly explains the long cycle life observed with PVDF – it simply does not break apart.

Ethyl cellulose, being a cellulose derivative, provides good adhesive strength as well, though it is somewhat more brittle than PVDF. EC forms a polymer network that adequately binds particles; while it may not be as elastomeric as PVDF, prior studies have shown it can maintain electrode cohesion under repeated cycling [5,25]. One advantage of EC is that it does not swell or dissolve in water, so after drying it remains intact in the aqueous electrolyte, avoiding the extreme swelling that water-soluble binders (like carboxymethyl cellulose or polyacrylic acid) would experience [6]. This helps preserve the electrode structure, though some gradual mechanical degradation (as seen by capacity fade) can still occur, potentially due to minor swelling or softening in the electrolyte over time.

Cross-linked gluten yields a different mechanical profile, as thermally cured gluten forms a flexible, rubbery network. The heating causes the proteins (gliadin and glutenin fractions) to cross-link via mechanisms like disulfide bond formation, which significantly improves film toughness and elasticity [26]. The resulting gluten binder can accommodate strain and maintaining contact between particles during cycling, much like PVDF.

In our short-term and rate tests, the gluten-bound electrodes did not show excessive cracking or early capacity loss, indicating decent mechanical resilience (Figure 5). However, the eventual capacity

loss by ~500 cycles suggests that the gluten binder might have slowly absorbed water or undergone structural fatigue that PVDF and EC did not. It is worth noting that, unlike PVDF and EC, gluten is a polypeptide that could be more susceptible to chemical degradation (e.g., hydrolysis) over long periods in water, especially if not fully cross-linked. Ensuring a high degree of cross-linking or blending gluten with a secondary polymer (to reinforce it) could mitigate this. The absence of any soluble components from the gluten binder is a positive – once cured, it does not dissolve – so the failure is likely mechanical rather than chemical dissolution.

Interfacial characteristics are also relevant. A good binder should facilitate close contact between the electrolyte and the active particles (for ionic transport) while also ensuring electronic connectivity via the conductive additive network. PVDF, being hydrophobic, can in some cases limit electrolyte wetting at the electrode interface, whereas more polar or hydrophilic binders (like many natural polymers) might allow better electrolyte penetration. Ethyl cellulose is modestly polar (due to – OC_2H_5 groups) and gluten is hydrophilic until cross-linked (after which it becomes water-insoluble but still slightly hydroscopic). In our experience, the EC and gluten slurries wetted the substrate and particles well, and the cured coatings did not inhibit electrolyte access. Indeed, the high capacities achieved suggest efficient ionic transport in those electrodes. PVDF electrodes, especially at 10% loading, might have had regions less accessible to the aqueous electrolyte due to PVDF's hydrophobicity and the reduced pore volume (because PVDF can dry to form a film that partly occludes pores). This difference in wetting could partly explain why the 10% PVDF sample underperformed at even moderate rates, whereas the bio-binders allowed better usage of LMO.

Lastly, it is important to consider the broader implications for solid-boosted FBs. The binder in each booster particle must withstand mechanical abrasion (particles colliding) and continuous contact with flowing electrolyte. From our findings, a binder like PVDF would likely hold particles together but raises sustainability and processing concerns (needing NMP, etc.), whereas ethyl cellulose and gluten could enable a fully green processing (water/alcohol-based). Ethyl cellulose looks particularly promising as it combined high performance with decent stability. Gluten showed that natural materials can work, but its long-term stability in an FB tank is questionable without further improvements. Still, even if gluten required periodic re-curing or replacement, its environmental benignity and cost (being essentially a food byproduct) could be advantageous. One possible mitigation for gluten's degradation could be coatings on the booster particles – for instance, a thin protective coating (like alginate or a shellac) over the gluten-bound particle to seal it from direct water exposure, while still allowing ion transport. These strategies are beyond the scope of the present study but represent promising directions for future research.

5. Conclusions

This study demonstrates that both ethyl cellulose and cross-linked gluten are viable, eco-friendly alternatives to PVDF in aqueous lithium-ion battery electrodes. Replacing PVDF with these natural binders dramatically improves the sustainability of electrode manufacturing: it enables safer processing (no toxic NMP fumes), faster electrode drying, and eliminates the need for costly solvent recovery systems. These advantages make the bio-binders highly attractive for large-format energy storage systems, where environmental regulations and safety requirements can drive up costs. The only processing trade-off is that gluten binders require an extra curing step (heating to 150 °C) to achieve water resistance, but this can be conveniently integrated into the standard drying process and is a one-time treatment. Electrochemically, our comparative tests showed that ethyl cellulose and gluten can replace PVDF without major sacrifices in performance. In fact, at 5% binder loading, both alternatives enabled higher initial capacities (~90–100 mAh g⁻¹ at 0.2–1C) than a traditional 10% PVDF electrode (~40 mAh g⁻¹ at 1C). Ethyl cellulose excelled in initial performance, and even though it exhibited some capacity fade over long cycling, it maintained a useful capacity (>50% of initial) over 500 cycles. Cross-linked gluten delivered comparable initial results and stable medium-term cycling (tens of cycles), though its long-term durability needs improvement as currently implemented. Notably, both ethyl cellulose and gluten binders produce electrodes that are completely free of

fluoropolymers and toxic solvents, aligning with the push for greener battery technologies. This means an entire cell can be made without PFAS or hazardous components when using these binders (especially in combination with an aqueous electrolyte), greatly simplifying end-of-life disposal or recycling. Each alternative binder has its strengths. Ethyl cellulose offers a very straightforward transition for industry – it uses a volatile, low-toxicity solvent (alcohol) and requires no special treatment, while delivering reliable mechanical and electrochemical performance close to PVDF. Crosslinked gluten allows an even greener process (water-based, no organic solvents at all) and, after curing, shows surprisingly good mechanical robustness (on par with soft polymers) due to protein network formation. The weaknesses observed – namely, the slightly higher capacity fade for ethyl cellulose and the exponential capacity loss for gluten – are not seen as fundamental barriers but rather as issues that can be addressed with further optimization (for instance, improved cross-linking density for gluten, or binder mixes to stabilize ethyl cellulose). The fact that these binders achieved near-PVDF performance in our tests is a significant outcome, as it underscores that moving away from PVDF is not only feasible but can be done without compromising cell function in aqueous systems.

In aqueous lithium-ion batteries, the use of ethyl cellulose or gluten as binders allows for the elimination of toxic and environmentally harmful components, complementing the safer water-based electrolyte. In solid-boosted flow batteries, these binders can simplify manufacturing and enhance recyclability, as the absence of fluorinated polymers facilitates material recovery and reduces environmental risks from disposal. This study demonstrates that sustainable binders like ethyl cellulose and cross-linked gluten can enable environmentally friendly energy storage systems without compromising reliability. With further optimization of formulation and long-term stability, biodegradable, PFAS-free binders could be confidently adopted in large-scale battery applications.

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Abbreviations

The following abbreviations are used in this manuscript:

PVDF Polyvinylidene Fluoride ALIBs Aqueous Lithium-Ion Batteries

FBs Flow Batteries

LMO Lithium Manganese Oxide (LiMn₂O₄)

NMP N-Methyl-2-Pyrrolidone

PFAS Per- and Polyfluoroalkyl Substances

EC Ethyl Cellulose CV Cyclic Voltammetry

SEM Scanning Electron Microscopy

EDS Energy-Dispersive X-ray Spectroscopy

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