



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Lithiation-Driven LiCrSe_2 Shell Growth on Metallic CrSe_2 Core Governs the Plateau–Slope Behavior

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ABSTRACT

The growing demand for high-performance lithium-ion batteries necessitates the development of cathode materials that combine high capacity, structural stability, and rapid charge–discharge capability. First-principles calculations predict that layered CrSe_2 possesses a robust framework capable of accommodating one Li^+ per formula unit while intrinsically supporting fast Li -ion diffusion. Muon spin rotation ($\mu^+\text{SR}$) measurements validate this prediction, revealing fast Li^+ diffusion in pre-lithiated CrSe_2 . Consistent with these findings, electrochemical testing demonstrates a reversible capacity of 125.3 mAh g^{-1} at 0.1 C , approaching the theoretical value of 127.7 mAh g^{-1} , with stable cycling and good rate capability. In operando X-ray diffraction and electrochemical impedance spectroscopy further reveal a reversible topotactic transition and a lithiation-driven core-shell evolution during cycling. These results show that lithiation-induced conductivity changes govern the electrochemical behavior of CrSe_2 , highlighting its potential as a high-performance cathode for LIBs. This study provides new insight into intercalation processes in layered transition-metal chalcogenides and informs the design of fast-charging electrodes.

1 | Introduction

Fast charging remains a major challenge for lithium-ion batteries (LIBs) and a key barrier to widespread adoption of electric vehicles [1–4]. While battery capacity and energy density have improved significantly [5–8], slow charging rates still limit user acceptance and practical deployment. Achieving fast-charging capability requires not only optimized electrolytes [5, 9–11] and advanced manufacturing strategies [1, 12, 13] but, more critically, electrode materials that can intrinsically support rapid

and reversible Li^+ insertion without structural degradation or polarization losses. To meet these requirements, electrode hosts must combine high Li^+ diffusivity, sufficient electronic conductivity, and robust frameworks capable of tolerating repeated ion migration.

Layered transition-metal dichalcogenides (TMDs) have emerged as promising candidates for high-rate electrode materials owing to their open van der Waals gaps, strong in-plane covalency, and intrinsic metallic conductivity [14–17]. In particular, Ti-

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[18, 19], V [16, 20, 21], and Zr-based [22, 23] TMDs are well known for their ability to undergo topotactic Li⁺ (de)intercalation while maintaining structural integrity during cycling. In contrast, chromium-based chalcogenides have received comparatively little attention despite their rich electronic structures and higher Cr³⁺/Cr⁴⁺ redox potentials [24–27], which could enable both strong electronic coupling and reversible redox activity. The mixed-anion compound CrSeS has recently been shown to accommodate one Li⁺ per formula unit and deliver exceptionally high charge–discharge rate performance in LIBs [28]. This outstanding kinetic behavior has been attributed to the expanded interlayer spacing and enhanced polarizability introduced by selenium, which together lower the Li⁺ migration barrier and facilitate ultrafast ion transport [28]. These results suggest that selenium incorporation plays a decisive role in accelerating Li⁺ diffusion within layered Cr-based hosts, suggesting that fully selenide analogues could exhibit even faster Li⁺ dynamics and superior rate performance.

Building on this insight, it is logical to examine binary CrSe₂, which contains only selenium and therefore maximizes anion size and polarizability effects [29–32]. The larger and more deformable Se²⁻ anions can effectively screen electrostatic interactions, facilitating Li⁺ migration through the layered framework and potentially enhancing rate capability beyond that of CrSeS. Using the predictions of a recently suggested descriptor for ion mobility [33, 34] this high mobility is also consistent with the relatively low electronegativity of Se compared to, e.g., oxygen. Previous μ^+ SR studies on pre-lithiated CrSe₂ have already revealed fast Li⁺ diffusion kinetics at room temperature [35], underscoring its potential as a high-rate host. Furthermore, CrSe₂ has also previously demonstrated promising performance in potassium-ion batteries by us, achieving its theoretical capacity at 0.1 C and retaining 85% capacity at 1 C [36], highlighting its structural robustness and favorable diffusion kinetics across multiple alkali systems. However, despite decades of research on intercalation chemistry in layered transition-metal dichalcogenides, the electrochemical lithiation behavior of CrSe₂ itself has never been systematically investigated. Its electronic and structural response to Li insertion, and the extent to which Se polarizability influences conductivity and diffusion, remain completely unknown. Uncovering these mechanisms is essential for understanding Cr-based TMDs and for guiding the design of next-generation fast-charging electrode materials that combine rapid ion transport with robust structural reversibility.

In this work, we investigate the lithiation dynamics of layered CrSe₂ by integrating periodic density functional theory (DFT) calculations within in operando and ex situ experiments. DFT predicts that Li⁺ occupies octahedral sites within a stable *P-3m1* lattice, driving a metallic-to-semiconducting transition that terminates at LiCrSe₂. These predictions are experimentally verified through in operando X-ray diffraction, X-ray absorption spectroscopy, electron microscopy, and μ^+ SR analysis. Electrochemical measurements reveal a distinct plateau–slope profile within 1.0–3.0 V, governed by conductivity evolution rather than multiple phase transitions. Combined structural and impedance analyses identify a conductivity-driven core-shell evolution, in which a metallic CrSe₂ core becomes progressively encapsulated by a semiconducting LiCrSe₂ shell during lithiation. These findings demonstrate that the plateau–slope behavior of CrSe₂

originates from dynamic electronic reconfiguration, establishing it as a representative model for conductivity-controlled intercalation in layered chalcogenides.

2 | Results and Discussion

2.1 | DFT Predictions of Thermodynamic Lithiation

Comprehensive DFT investigations were performed on the entire Li_xCrSe₂ system including calculations on the electronic structure, structural stability and barriers for Li diffusion. DFT has been shown to provide reliable results for properties of batteries [37]. A thorough discussion on the underlying methodology including a structural benchmark for six different DFT flavors can be found in the [Supporting Information](#) and Table S1. Since both CrSe₂ and LiCrSe₂ can be synthesized as layered compounds, the corresponding phases were taken as starting point for the computational investigations. The calculated electronic density of states (Figure 1a) shows that pristine CrSe₂ is metallic, with the Fermi level intersecting strongly hybridized Cr 3d and Se 4p bands consistent with previous reports [36, 38]. Upon lithiation to LiCrSe₂, electrons donated by Li occupy antibonding Cr–Se orbitals, which lower the density of states near the Fermi level, open a small band gap, and transform the material from metallic to semiconducting. The substantial orbital overlap implies possible participation of both Cr and Se in charge compensation during lithiation.

Once the delithiated and lithiated phases were established, convex-hull calculations were conducted to evaluate the thermodynamic stability of possible intermediate compositions in the Li_xCrSe₂ system (Figure 1b) Additional to the *P-3m1* phase for LiCrSe₂, the *R-3m* and *C2/m* structures, known to be stable for NaCrSe₂ [39] and KCrSe₂ [40], were examined as possible Li-intercalation frameworks. Several intermediate configurations for 0.0 ≤ x < 0.08 and for 0.5 < x ≤ 1.0 are predicted to be stable or within the stability range (given the methodological inaccuracy) all of which consistently retain *P-3m1* symmetry. The lattice spacing jumps from c = 5.9 Å at x = 0.08 to c = 6.3 Å at x = 0.5 where it remains roughly constant until full lithiation indicating only a single significant structural transition at x = 0.5 for the system. The phase stability of the configurations within 0.5 < x ≤ 1.0 is likely to be driven by charge ordering effects as this region is characterized by only subtle structural changes. No additional low-energy configurations were identified outside the *P-3m1* framework, confirming that the layered lattice remains topotactically stable during Li insertion.

Within the *P-3m1* framework, the preferred Li-occupation site was further examined to clarify the intercalation mechanism. High-symmetry octahedral and tetrahedral sites between Se layers were considered, as prior studies indicate that their energy difference can be small [41]. The DFT results (Figure S1) show that octahedral coordination is consistently more stable than tetrahedral occupation across all Li contents. Configurations with Li in tetrahedral sites lie significantly above the convex-hull line, particularly at higher Li contents, indicating that tetrahedral occupation is energetically unfavorable. Although Li_xCrSe₂ (x > 1) has not been previously reported, analogous layered systems

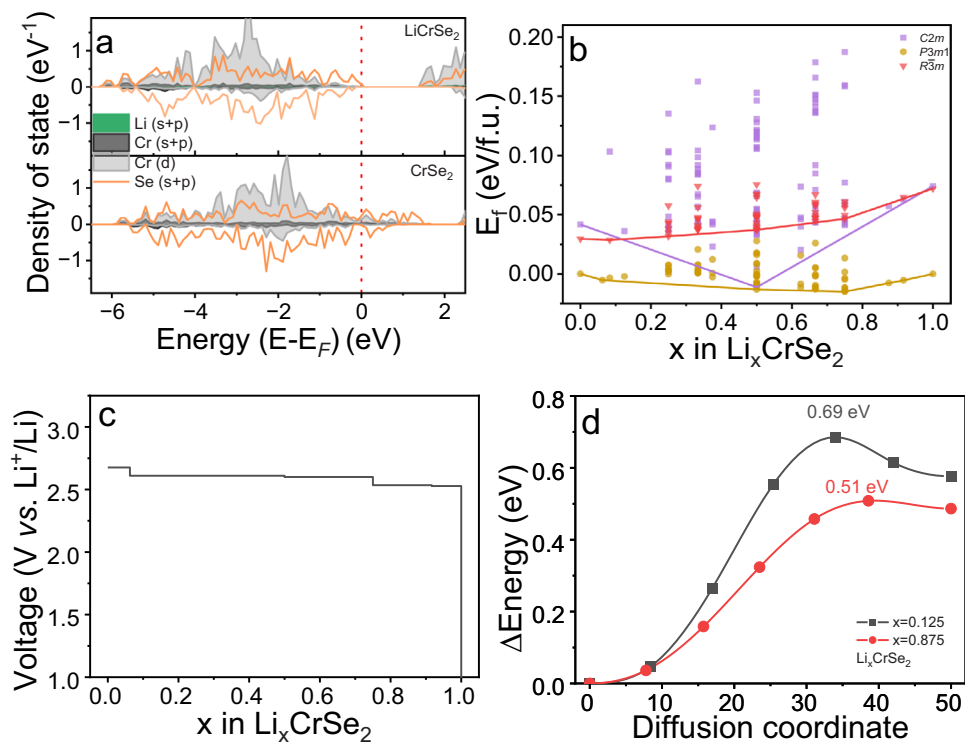


FIGURE 1 | (a) Electronic density of states of LiCrSe₂ and CrSe₂. (b) Calculated formation energies of Li_xCrSe₂ for different site occupations as explained in more detail in the [Supporting Information](#). (c) DFT-predicted intercalation voltage profile of CrSe₂ during lithiation. (d) Li-ion energy barriers of Li_xCrSe₂ for low ($x = 0.125$) and high ($x = 0.875$) lithium-vacancy phases corresponding to the diffusion path from octahedral to tetrahedral site.

such as VS₂ and TiS₂ can accommodate a second Li ion once the octahedral sites are fully occupied [42]. However, for CrSe₂ DFT calculations (Figure S2) reveal that lithiation beyond LiCrSe₂ does not yield a stable Li_xCrSe₂ ($x > 1$) phase. Instead, additional Li insertion destabilizes the structure and drives decomposition into Li₂Se and a Cr-rich residue. These findings indicate that Li intercalation in CrSe₂ is confined to octahedral sites and terminates at LiCrSe₂, beyond which further Li uptake triggers a conversion-type reaction.

Based on the predicted stable phases of Li_xCrSe₂ (Figure 1b), the calculated voltage profile exhibits a nearly constant plateau, centered around 2.6 V versus Li⁺/Li (Figure 1c). Although several intermediate phases were predicted from the convex-hull analysis, their voltage differences are very small, resulting in an almost flat voltage curve in which multiple transitions are not distinguishable in battery testing.

To further examine the kinetic aspect of Li transport, nudged elastic-band (NEB) calculations were performed at low ($x = 0.125$) and high ($x = 0.875$) Li contents, representing dilute and nearly filled states of Li_xCrSe₂. The calculated diffusion barriers are 0.69 eV at $x = 0.125$ and 0.51 eV at $x = 0.875$, indicating that Li migration becomes slightly easier at higher Li concentration. These barriers are only moderately higher than those reported for Li_xCrSSe (Figure S3), suggesting that although Li diffusion in CrSe₂ is somewhat slower, the kinetics remain comparable. Given that CrSSe has been reported to exhibit exceptional rate performance [28], CrSe₂ is expected to display similarly favorable kinetics, motivating its synthesis and electrochemical charac-

terization to verify the predicted lithiation behavior. Although a lower barrier was initially anticipated due to the larger and more polarizable Se²⁻ anions, the slightly higher value obtained here suggests that stronger Cr-Se covalency counterbalances this effect, leading to comparable rather than enhanced Li⁺ mobility. Furthermore, the electronegativity of Se (2.60) and S (2.69) is very similar which according to a recently suggested descriptor for ion mobility in crystalline materials based on the so-called migration number [33, 34] should also yield very similar migration barriers.

2.2 | Structural, Electronic, and Morphological Characterization CrSe₂ and LiCrSe₂

To experimentally validate the above DFT predictions, CrSe₂ was synthesized and comprehensively characterized before electrochemical testing. CrSe₂ mixed with 10 wt.% graphite (hereafter referred to as CrSe₂) was synthesized, following our previous reports [36, 38]. The resulting powder product is air-stable and was used instead of LiCrSe₂ to ensure compatibility with standard electrode fabrication procedures. Powder X-ray Diffraction (PXRD) data (Figure 2a,b) collected on CrSe₂ and LiCrSe₂ match the reported crystal structure (space group *P-3m1*), as confirmed by the agreement between experimental and modelled profiles. The refined lattice parameters of CrSe₂: $a = 3.3956(6)$ Å and $c = 5.9168(10)$ Å, and LiCrSe₂: $a = 3.65127(5)$ Å and $c = 6.29616(10)$ Å agree well with literature values (Tables S2 and S4), verifying that both phases crystallize in the expected structure without impurity peaks.

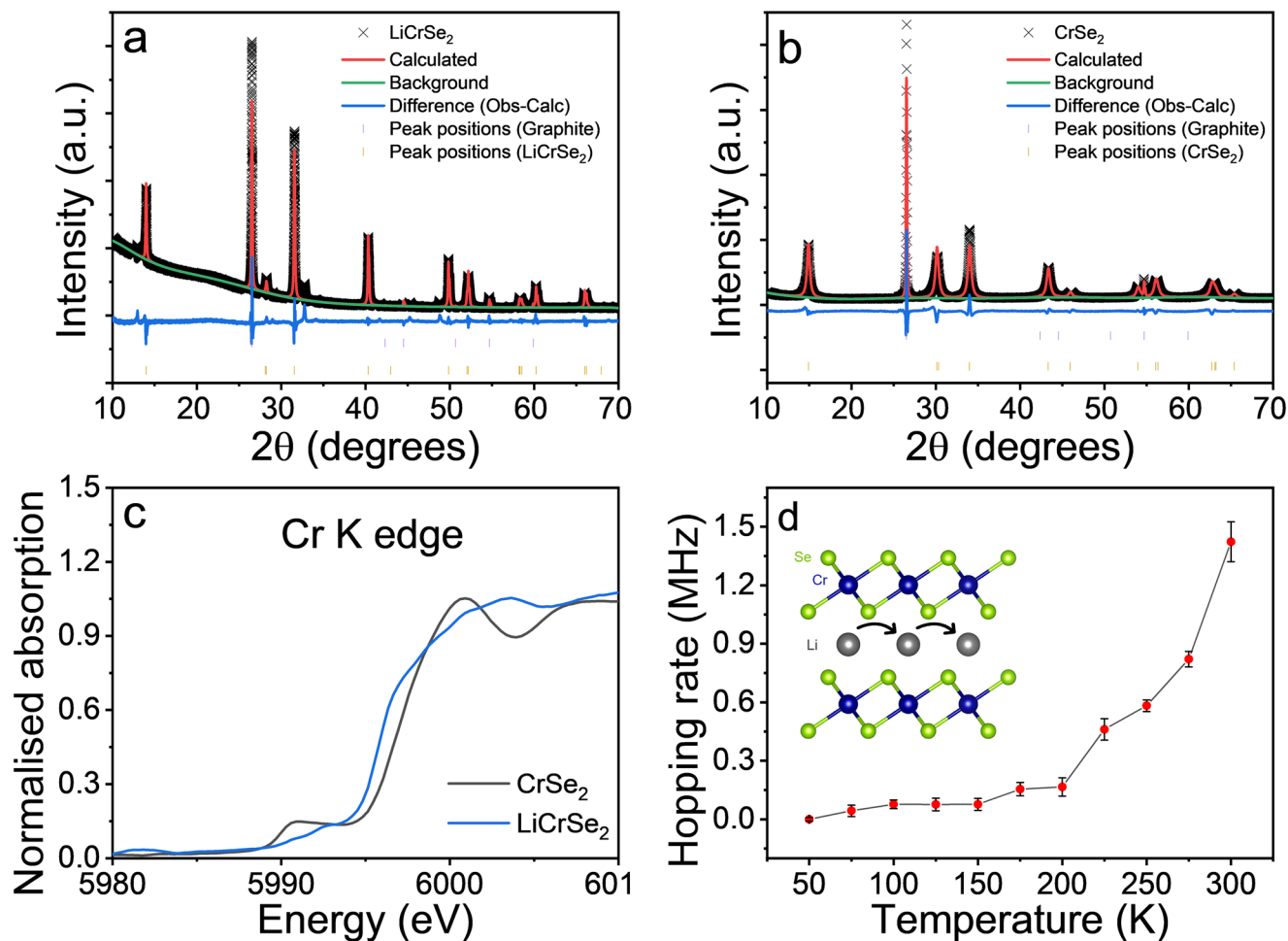


FIGURE 2 | Le Bail refinements of experimental PXR D patterns (Cu $K\alpha$) for (a) LiCrSe₂ and (b) CrSe₂. Measured data are shown as black crosses, calculated profiles as red lines, and different curves as blue lines. Magenta vertical bars indicate the expected Bragg reflection positions of the phase, (c) Cr K-edge XANES profile of CrSe₂ and LiCrSe₂, and (d) Temperature dependence of the Li-ion hopping rate (ν_{Li}) from μ^+ SR measurements.

Having confirmed the structural integrity of both compounds, their local electronic structures and oxidation state evolution were probed using X-ray absorption near-edge structure (XANES) spectroscopy (Figure 2c; Figure S4). Both CrSe₂ and LiCrSe₂ adopt edge-sharing [CrSe₆] octahedra, but the Cr–Se bond length expands from 2.47 Å in CrSe₂ to 2.54 Å in LiCrSe₂ (Table S4), consistent with Cr reduction upon lithiation. The Cr K-edge of CrSe₂ appears at higher energy than that of LiCrSe₂, consistent with a higher Cr valence in the CrSe₂ phase. However, both edge positions lie below those of Cr⁴⁺ and Cr³⁺ reference oxides, evidencing strong Cr–Se covalency arising from hybridized Cr 3d–Se 4p states [43, 44]. At the Se K edge, CrSe₂ exhibits a pre-edge feature at 12660.2 eV and a sharp absorption edge at 12669.4 eV. Upon lithiation, the pre-edge intensity decreases, and the edge shifts to 12667.4 eV, reflecting partial Se reduction. These spectral changes corroborate the DOS (Figure 1a), confirming strong Cr–Se hybridization and a transition from metallic CrSe₂ to semiconducting LiCrSe₂. The combined Cr- and Se-edge shifts indicate the participation of both elements in the (de)lithiation process.

The microstructure and morphology of CrSe₂ were investigated using scanning electron microscopy (SEM) and high-resolution transmission electron microscopy (HRTEM). SEM images (Figure S5) reveal the characteristic stacked-platelet morphology typical of layered transition-metal dichalcogenides. Energy-dispersive X-ray (EDX) mapping confirms uniform Cr and Se distributions across large areas without elemental segregation, yielding a Cr:Se ratio of $1:2.02 \pm 0.02$. HRTEM images (Figure S6) show thin, plate-like crystallites (~500 nm in lateral dimension) with well-defined lattice fringes near the platelet edges, confirming high crystallinity. Selected-area electron diffraction (SAED) patterns obtained along the [001] zone axis display sharp reflections for the (100) and (001) planes, corresponding to d-spacings of 3.3957 and 5.9182 Å, respectively, in excellent agreement with PXR D results. Fast Fourier transforms (FFT) of HRTEM images further confirm the [001] lattice orientation, while slight in-plane variations suggest weak stacking disorder between layers. Local EDX mapping of individual platelets reveals strong spatial overlap between Cr and Se signals, with a Se:Cr ratio of 2.01 ± 0.01 , consistent with bulk measurements.

While XANES and structural analyses verify the hybrid redox behavior and crystallographic stability predicted by DFT, they do not directly probe Li^+ mobility, which critically governs the kinetic performance of CrSe_2 electrodes. To evaluate intrinsic Li^+ diffusion independent of electrode architecture or electrolyte effects, muon-spin-relaxation ($\mu^+\text{SR}$) spectroscopy was employed. This technique quantifies Li^+ hopping rates in pre-intercalated compounds [35, 45, 46]. $\mu^+\text{SR}$ measurements were performed on as-synthesized LiCrSe_2 using the EMU instrument at the ISIS Neutron and Muon Source (more details in Supporting Information and Figure S7). The temperature-dependent hopping rate increases steadily up to 300 K (Figure 2d), with the onset of mobility between 150 and 175 K, similar to previous reports [35]. The jump frequency was calculated with an Arrhenius behaviour and an activation energy of 74(8) meV (Figure S8). At room temperature, the hopping rate exceeds those of benchmark layered oxides such as Li_xCoO_2 [47], LiFePO_4 [48], and $\text{LiNi}_{0.33}\text{Co}_{0.33}\text{Mn}_{0.33}\text{O}_2$ [49], confirming that LiCrSe_2 possesses intrinsically fast Li^+ diffusion kinetics. Given this superior intrinsic ion-transport behavior, LiCrSe_2 is a promising candidate for further electrochemical testing in practical battery configurations.

2.3 | Electrochemical Behavior of CrSe_2

To evaluate the Li-storage behavior of CrSe_2 and determine an appropriate operating voltage, Swagelok-type half-cells were assembled using Li metal as the counter electrode and 1 M LiPF_6 in EC/DMC (1:1 v/v) as the electrolyte. The galvanostatic charge–discharge (GCD) curves in Figure 3a show how the initial capacity change depending on the potential cut off during the first cycle. However, below about 0.9 V the material undergoes a conversion reaction, which leads to a sharp increase in capacity and a distinctive voltage plateau. The overall discharge capacity reaches 482 mAh g^{-1} , far exceeding the theoretical value of 127.7 mAh g^{-1} corresponding to one Li^+ per CrSe_2 . Such an unusually high capacity cannot arise from further Li intercalation, since DFT predicts that lithiation beyond LiCrSe_2 is thermodynamically unstable and leads to decomposition into Li_2Se and a Cr-rich residue (Figure S2). The low-voltage plateau is therefore attributed to a Li–Se conversion reaction, analogous to that observed in CrS_2 and CrSSe below $\sim 1.0 \text{ V}$ [28, 42]. The inset photograph in Figure S9 shows the reddish discoloration of the separator after discharge, indicative of soluble Se-containing conversion products. Consistent with this irreversible process, the first discharge delivers 482 mAh g^{-1} whereas the first charge capacity is limited to 318 mAh g^{-1} . The discharge capacity decays to 447 mAh g^{-1} at the second cycles (Figure S9).

To confirm that the sloping region observed above 1.0 V originates from intercalation rather than a conversion process, additional GCD measurements were performed with various higher cutoff voltages of 1.0, 1.25, 1.5, and 2.0 V. As shown in Figure 3b and Figures S10 and S11, the discharge–charge profiles obtained within these voltage windows are nearly identical, and more than 99% of the capacity was retained over the first five cycles. The discharge capacity increases as the cutoff is lowered from 2.0 to 1.0 V, from 64.87 to 125.3 mAh g^{-1} . Within the 1.0–3.0 V range, CrSe_2 delivers 125.3 mAh g^{-1} , very close to the theoretical value of 127.7 mAh g^{-1} corresponding to one Li^+ per CrSe_2 . These results

demonstrate that the plateau at $\sim 2.6 \text{ V}$ originates from a fully reversible intercalation process rather than from any conversion-type reaction. Cycling tests at 0.5 C (Figure S10) further show that discharging to 1.0 and 1.25 V retains approximately 85% of the initial capacity after 50 cycles, whereas cells with higher cutoffs maintain about 94%. Balancing capacity and reversibility, the 1.0–3.0 V window was therefore adopted for all subsequent electrochemical and structural analyses of the plateau–slope behavior, as it enables full theoretical capacity while preventing the irreversible Li–Se conversion reaction observed below 1.0 V.

Starting from the 2nd cycle within the optimized voltage range of 1.0–3.0 V, the discharge curve of CrSe_2 shows two distinct regions, where a plateau at around 2.6 V and a gradual slope extending to 1.0 V contribute equally to the total capacity (Figure 3b). The plateau region is in excellent agreement with the DFT results (Figure 1c) whereas the predicted voltages for $0.5 < x \leq 1.0$ contrast the experimental findings. We speculate that the applied meta-GGA functional does not describe the effects of the Cr–Se hybridization very accurately. The corresponding cyclic voltammetry (CV) curves (Figure 3c; Figure S12) display only one pair of well-defined redox peaks at 2.36 V (cathodic) and 2.62 V (anodic), representing a single, reversible intercalation–deintercalation process between CrSe_2 and LiCrSe_2 . No additional peaks are observed within the 1.0–3.0 V window. The peaks remain sharp and nearly symmetric even at higher scan rates, shifting slightly to 2.32 and 2.67 V at 1.0 mV s^{-1} due to mild polarization. The linear relationship between peak current (I_p) and the square root of the scan rate ($v^{0.5}$) (Figure S13) confirms that Li^+ transport is diffusion controlled. The calculated diffusion coefficients are $9.71 \times 10^{-9} \text{ cm}^2 \text{ s}^{-1}$ for lithiation and $7.69 \times 10^{-9} \text{ cm}^2 \text{ s}^{-1}$ for delithiation, where CrSe_2 exhibits slightly larger diffusion coefficients than CrSSe of $4.56 \times 10^{-9} \text{ cm}^2 \text{ s}^{-1}$ and $3.24 \times 10^{-9} \text{ cm}^2 \text{ s}^{-1}$ [28]. Despite this difference, the experimental diffusion coefficients of CrSe_2 is comparable to other transition metal dichalcogenides (Table S5). Rate performance tests (Figure S14) further validate the favorable Li^+ kinetics in CrSe_2 , which retaining 63% of its capacity at 1 C relative to 0.1 C. Compared with CrSSe , which maintains about 50% of its capacity even at 200 C [28], CrSe_2 exhibits slower rate performance, consistent with its slightly higher Li^+ migration barrier and lower diffusion coefficients.

Electrochemical impedance spectroscopy (EIS) was carried out during the first discharge to monitor the evolution of charge-transfer resistance (Figure 3d; Figure S15). At open-circuit voltage, the Nyquist plot exhibits a large semicircle that significantly contracts as the potential enters the plateau region (2.36–2.34 V), indicating stable and relatively low interfacial resistance. The low-frequency response remains nearly unchanged, suggesting that Li^+ diffusion is not rate-limiting in this regime. As lithiation proceeds into the sloping region (1.95–1.0 V), the high-frequency semicircle gradually enlarges, revealing a progressive increase in charge-transfer resistance and a decline in electronic conductivity. At deep discharge (1.0 V), the resistance reaches its maximum and an additional low-frequency feature emerges, evidencing sluggish interfacial kinetics in fully lithiated LiCrSe_2 [50]. In contrast, the DFT-calculated density of states for Li_xCrSe_2 (Figure S16) shows that the system remains metallic up to $x = 0.92$. However, upon lithiation the number of states at E_F decreases significantly (especially above $x = 0.75$), leaving only

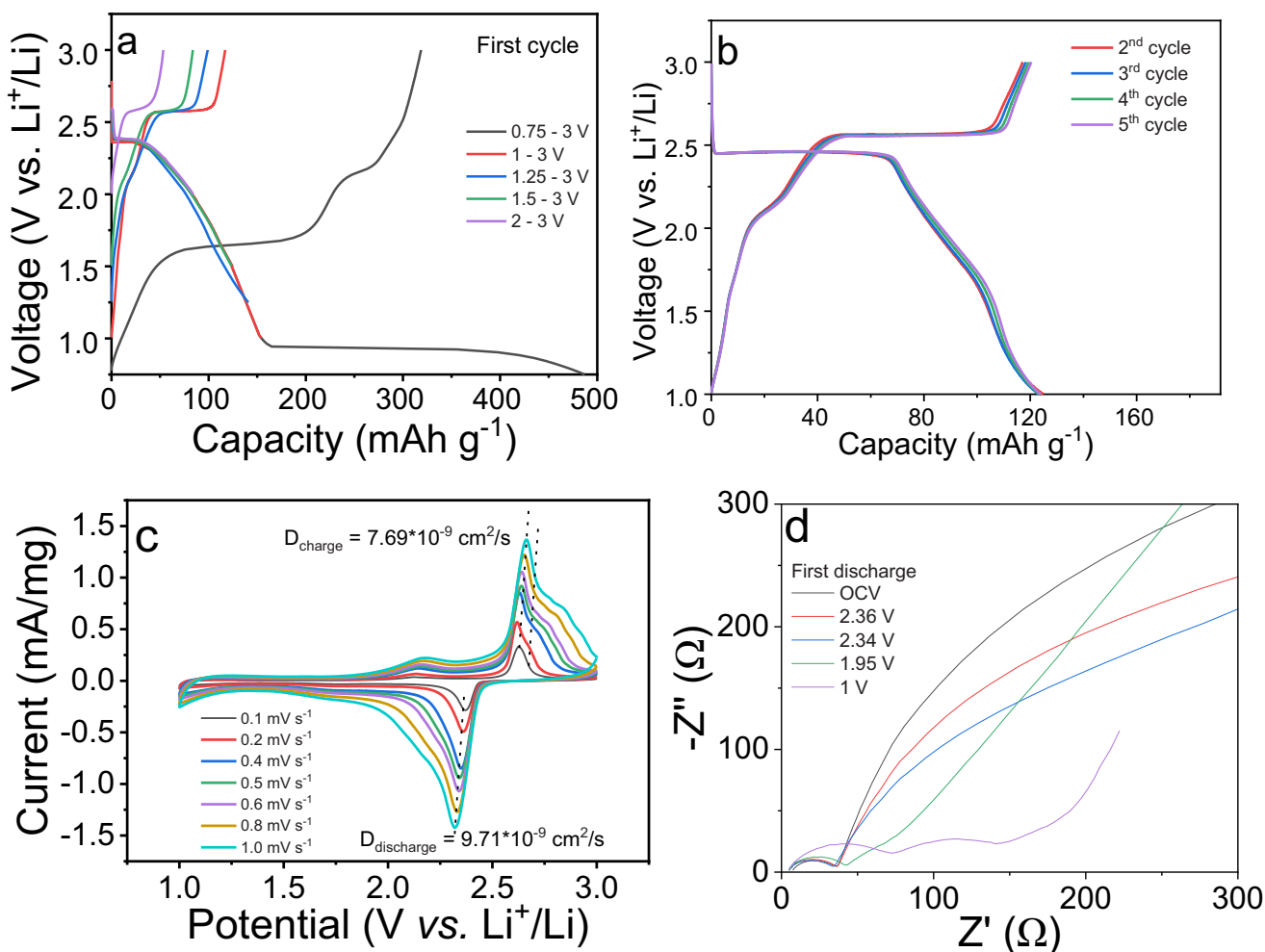


FIGURE 3 | (a) GCD profiles for the 1st cycle of CrSe₂ recorded at different lower cut off voltages (0.75, 1, 1.25, 1.5, and 2 V), (b) GCD curves (from 2nd cycle) of CrSe₂ within the voltage window of 1–3 V at 0.1C, (c) CV curves of CrSe₂ at various scan rates (0.1–1 mV s⁻¹) in the voltage range of 1–3 V, with calculated diffusion coefficients for charge and discharge processes. And (d) Electrochemical impedance spectrum measured at different potentials.

few accessible electronic states in the conduction band which may imply reduce electronic conductivity, until the band gap fully opens at $x = 1.0$. In addition, temperature-dependent resistivity measurements were performed on pressed pellets of CrSe₂ and LiCrSe₂. Pristine CrSe₂ exhibits negligibly low resistance, while LiCrSe₂ shows a temperature-dependent decrease in resistivity (Figure S17), consistent with semiconducting behavior as predicted by DFT calculations. In conclusion, the reduction in conductivity already begins within the sloping stage, rather than only at complete lithiation as could have been predicted by theory. The discrepancy between the experimental GCD profile and the DFT-predicted voltage plateau suggests that the transition in CrSe₂ occurs earlier, around $x \approx 0.5$ rather than $x = 0.92$ as predicted, prompting further investigation through *in operando* and spectroscopic analyses discussed in Section 2.4.

2.4 | Structural Evolution of CrSe₂ During Electrochemical Testing

The structural evolution of CrSe₂ during lithiation was monitored using an *in operando* XRD setup similar to that previously employed for CrSe₂ in potassium-ion battery studies [36]. Mea-

surements were carried out over the first two discharge–charge cycles, and the resulting contour maps are shown in Figure 4a and Figure S18. The Bragg reflections observed at 14.9° and 34.0° were assigned to the (001) and (101) planes of CrSe₂, respectively. The (002) and (102) reflections overlap at approximately 30.07°, serving as reference peaks. These features agree well with the simulated XRD pattern of standard CrSe₂ (Figure 4a; Figure S19). Upon lithiation, a complete disappearance of the original CrSe₂ reflections and the emergence of new diffraction peaks of LiCrSe₂, particularly at $x \approx 0.5$ in Li_xCrSe₂ indicate a transition mechanism. The newly formed peaks at 14.1°, 28.4°, and 31.6° were assigned to the (001), mixed (002)/(102), and (101) planes of the LiCrSe₂ phase, matching the simulated pattern in Figure 4a and Figure S20. During the recharge process, the peak evolution occurred in the opposite direction to the discharge one, representing a reversible transition between CrSe₂ and LiCrSe₂ within 1.0–3.0 V.

Benefiting from the high symmetry of the *P-3m1* space group in both CrSe₂ and LiCrSe₂, structural changes during cycling can be effectively extracted through the evolution of key diffraction reflections. Specifically, the position of the (101) peak is determined by both *a*- and *c*-axis lattice parameters, while the *c*-axis

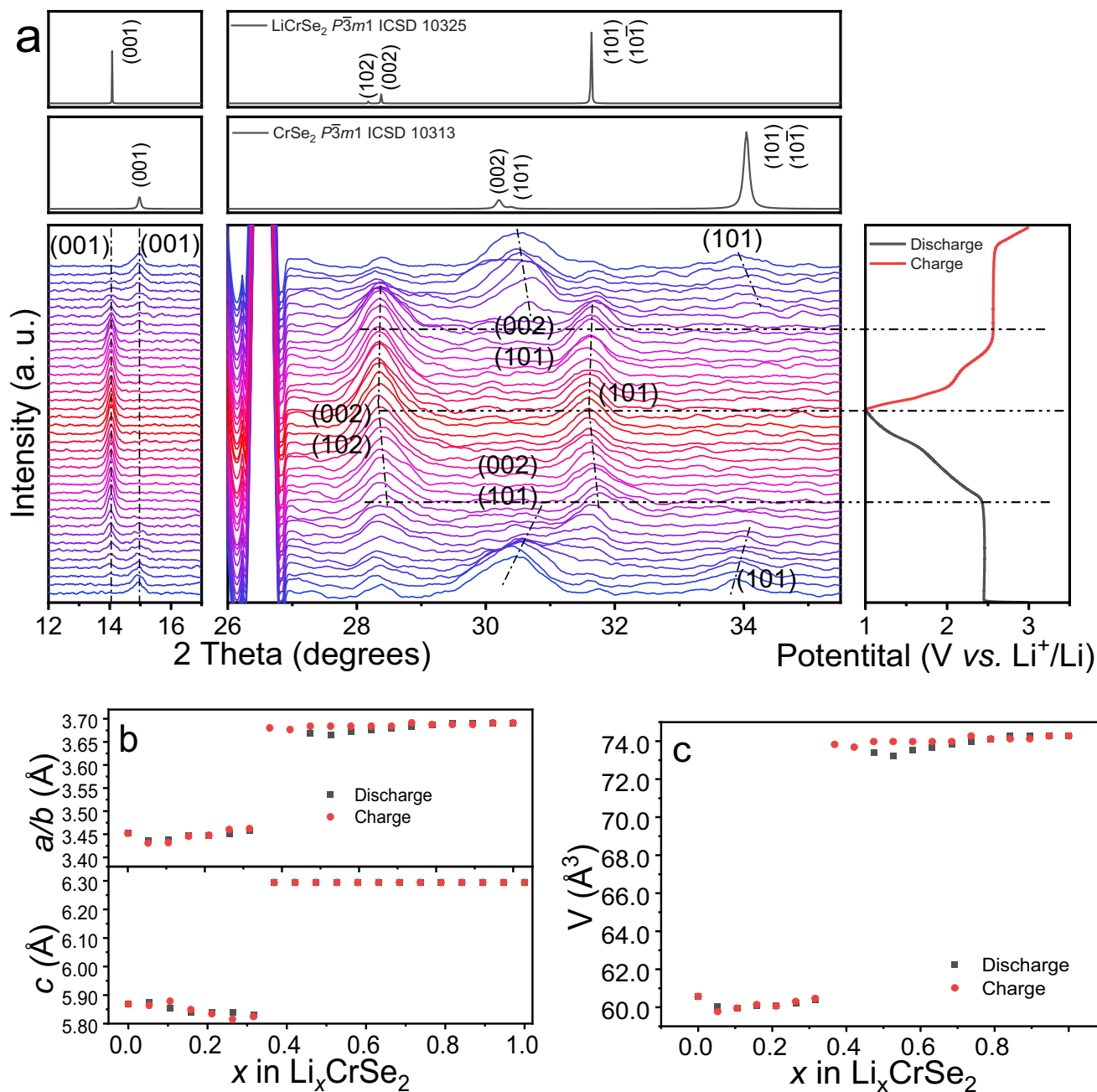


FIGURE 4 | (a) Diffraction patterns from the *in operando* XRD experiment. The right panel shows cell voltage as a function of time at second cycle, which is aligned with the diffraction patterns. Every other pattern was shown here while the intensity was sequentially offset. The cells were charged and discharged between 1.0 and 3.0 V at a rate of $C/0.1$ for 1 cycle. The simulated CrSe_2 and LiCrSe_2 patterns are on the top of in-operando XRD. (b) Evolution of lattice parameters a/b and c calculated from the *in operando* XRD during the first charge/discharge process, and (c) Unit cell volume calculated from the lattice parameters a/b and c .

can be independently monitored through the position of the (001) reflection (details in Supporting Information, Formulas 1 and 2). From the fully delithiated CrSe_2 of the second cycle (3.0 V), the extracted lattice parameters are $a = 3.452$ Å and $c = 5.869$ Å, while the fully lithiated CrSe_2 at the end of the second discharge (1.0 V) shows expanded values of $a = 3.691$ Å and $c = 6.294$ Å. These values agree well with both the chemically synthesized CrSe_2 and LiCrSe_2 in this work and previously reported data (Tables S2 and S3). The evolution of the lattice parameters during cycling, extracted from Figure 4a, is summarized in Figure 4b,c. The a -axis remains nearly constant at ~ 3.452 Å during the

plateau stage of discharge ($x < 0.5$), then expands gradually and stabilizes at 3.691 Å upon further lithiation. During charging, it fully returns to its original value. In contrast, the c -axis remains unchanged at ~ 5.869 Å during the early discharge, then increases to 6.294 Å at deeper lithiation and stays constant thereafter as has been predicted by DFT. Compared with other $P-3m1$ materials, the c -axis expansion in CrSe_2 exhibits 7.67%, which is slightly higher than that of CrSeS (7.44%) [28], VS_2 (6.6%) [20], and TiS_2 (5.22%) [49]. This corresponds to a significant lattice volume expansion of approximately 22% (Figure 3c). Notably, despite this large lattice expansion, the unit cell parameters of electrochemi-

cally intercalated LiCrSe_2 and deintercalated CrSe_2 remained in excellent agreement with chemically synthesized LiCrSe_2 and CrSe_2 (Tables S2 and S3). Although lithiation involves relatively large volume change, no pronounced mechanical degradation is observed under the cycling conditions investigated. The as-synthesized CrSe_2 exhibits an intrinsically broad particle-size distribution (Figure S5), which is largely preserved after electrode preparation. SEM images collected after 20 cycles show that the particles maintain similar sizes and overall morphology, with no clear evidence of particle fracture or pulverization (Figure S21).

The *in operando* XRD (Figure 4a) revealed only a single structural transition during lithiation which is in line with the DFT investigations. This transition coincides precisely with the voltage inflection from the plateau to the sloping region in the GCD profile, indicating that the structural and electronic evolutions occur simultaneously. The cross-sectional Scanning Transmission Electron Microscopy analysis of solids extracted from electrodes with the electrochemical experiment halted at approximately $x \approx 0.5$ in the $\text{Li}_{0.5}\text{CrSe}_2$ phase (Figure S22) provides further insight into the lithiation mechanism. One region displays well-ordered lattice fringes consistent with pristine CrSe_2 , while adjacent regions exhibit expanded and locally bent lattice planes that can be attributed to the LiCrSe_2 phase. This observation suggests that lithiation proceeds heterogeneously within a single particle. Combining these observations with the DFT-calculated density of states for CrSe_2 and LiCrSe_2 , the process can be rationalized as a progressive conductivity change driven by lithiation. During lithiation, Li intercalates initially at the particle surface, forming a LiCrSe_2 shell around a metallic CrSe_2 core. In this stage, corresponding to the plateau region, the metallic core ensures high overall conductivity, consistent with the low and stable interfacial resistance observed by EIS (Figure 3d). As lithiation proceeds, the LiCrSe_2 shell thickens, and once its semiconducting layer reaches a critical thickness, the overall electronic conductivity of the core-shell structure decreases sharply. This increase in charge-transfer resistance manifests as the voltage drop and transition from the plateau to the sloping region. Therefore, we hypothesize that the observed plateau-slope behavior originates from the dynamic formation of a conductive-semiconductive core-shell configuration within CrSe_2 during lithiation, rather than from multiple discrete phase transitions.

3 | Conclusions

In summary, combined theoretical and experimental investigations reveal that CrSe_2 undergoes a single, reversible topotactic intercalation process within the voltage range of 1.0–3.0 V. DFT calculations reveal that Li insertion is limited to octahedral sites, maintaining the *P-3m1* framework and inducing a transition from metallic to semiconducting behavior upon full lithiation. Experimental XRD and XANES confirm the structural stability and hybrid Cr-Se redox character of the CrSe_2 - LiCrSe_2 pair. Electrochemical measurements identify a distinct plateau-slope discharge profile, governed by conductivity evolution rather than multiphase transitions. *in operando* XRD reveals only one structural conversion, while EIS and DOS analyses collectively indicate that the transition from plateau to slope arises from the

formation and growth of a LiCrSe_2 shell surrounding a metallic CrSe_2 core. This core-shell evolution progressively increases interfacial resistance and triggers the observed voltage slope. The $\mu^+\text{SR}$ measurements further confirm fast intrinsic Li^+ diffusion in LiCrSe_2 , supporting its excellent kinetic properties. Overall, CrSe_2 exemplifies conductivity-governed intercalation in layered electrodes, where a gradual conductivity change drives the transition from plateau to slope. The proposed core-shell evolution offers a plausible explanation for this behavior, providing a foundation for future TMDs material studies aimed at directly resolving and potentially controlling such conductivity-driven transformations.

Several challenges must still be addressed before CrSe_2 can be considered for practical applications. First, the synthesis demonstrated here is restricted to laboratory scale, therefore the development of scalable preparation method will be required. LiCrSe_2 is sensitive to ambient conditions, which indicates that controlled processing or appropriate surface-stabilization strategies will be necessary for practical electrode fabrication. Nevertheless, the mechanistic insights obtained in this work establish a useful framework for understanding how conductivity changes influence charge-discharge behavior in Li-ion cells. This creates a clear route toward rational optimization of the electronic structure in chalcogenide-based electrodes, especially since density-functional theory provides an effective way to guide such optimization in a systematic manner through the density of state calculations.

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Conflicts of Interest

The authors declare no conflicts of interest.

Data Availability Statement

The data that support the findings of this study are openly available in NOMAD repository at <https://dx.doi.org/10.17172/NOMAD/2024.07.18-1>, reference number 202407181.

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