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Computational study of novel 2H chromium ditelluride as an anode material for Li/K-ion batteries

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The goal of metal-ion battery research is to develop anode materials with high storage capacity. This study explored the potential of 2H phase $CrTe_2$, composed of two hexagonally stacked layers, as an optimal anode material for Li/K-ion batteries using Density Functional Theory (DFT). Preliminary analyses revealed that the material possesses thermodynamic, structural, and mechanical stability. A key finding was the significantly negative adsorption energy, which enhances battery stability by preventing clustering and stabilizing Li/K adsorption on the material's surface. The adsorption energy values for Li/K were calculated to be -3.7 eV and -4.63 eV, respectively. These results suggest stable lithiation and potassiation processes, with maximum storage capacities of 1395 mA h g^{-1} for Li and 1134 mA h g^{-1} for Li. Additionally, the calculated open-circuit voltages (OCVs) for Li were 0.13 V for Li in and 0.20 V for Li income. We calculated the adsorption energy, structural and electronic properties, theoretical capacity, diffusion energy, and thermal stability. The electrical conductivity of the material increased, and its metallic properties were maintained with increasing metal-ion concentration. This study highlights the potential of Li and Li in Li i

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1 Introduction

Modern hybrid vehicles and portable electronic devices have made the development of durable and efficient energy sources important. The outstanding qualities of the lithium-ion battery (LIB), including its high energy density, strong cycle performance, and lightweight design have attracted attention. LIBs are essential to the transition from oil-powered to electric-powered vehicles. They continue to serve as energy storage devices, aiding in the shift to renewable energy sources. Furthermore, the manufacturing of lithium-ion batteries has expanded dramatically in response to the growing demand for portable electronic devices, like laptops, mobiles, tablets, digital cameras, wearable devices, and drones.

The electrochemical and physical properties including the overall performance of batteries, result from the complex interaction between components such as the anode, cathode, binder, separator, and electrolyte solution. These components undergo multiple manufacturing processes that shape the structure of the battery electrodes.⁵ It is crucial to fully

understand how the various parts and production procedures interact and influence the unique qualities of the battery to achieve the best performance, quality, and cost-efficiency in LIB production.

Since Sony introduced LIBs in 1991, the technology has seen significant advancements. However, the growing demand has raised concerns about lithium scarcity. This has spurred efforts to explore alternatives, including sodium-ion batteries (NIBs), which have gained attention due to sodium's abundance and lower cost than lithium.⁶ Going beyond NIBs, there is also ongoing research into batteries utilizing other alkali metals like multivalent metal ions, including magnesium (Mg), calcium (Ca), and aluminum (Al).

As potassium is widely affordable and environmentally friendly, potassium-ion batteries (KIBs) are emerging as a competitive alternative for large-scale electrical energy storage compared to LIBs. KIBs might be a more cost-effective and environmentally responsible option for energy storage. Researchers are actively investigating different electrode materials to understand their qualities and performance to increase their efficacy and viability for future applications. The choice of electrode material greatly impacts the stability, conductivity, cost, weight, and environmental impact of batteries. The key challenge in developing these diverse secondary batteries lies in the search for suitable electrode materials that can be tailored to optimize battery performance. This quest for electrode materials is crucial to enable them to reach their greatest

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potential alternative battery technologies and address the limitations of traditional LIBs. Owing to its high particular capacity and the layered structure of electrode materials, which usually exhibit an inverse relationship with their mass density, graphite is frequently chosen as the primary electrode material. The specific capacity of these battery materials can also be enhanced due to lightweight carbon compositions.

The field of two-dimensional (2D) materials has recently seen significant research activity as a promising new class of anode materials. Their unique structure and diverse applications pique our curiosity.⁷⁻⁹ Consider the 2D material pioneer graphene, which has a remarkable theoretical Li-ion storage capacity (784 mA h g⁻¹)10 a significant surface-to-mass ratio (2965 m² g⁻¹), increased thermal conductivity, 2 and remarkable mechanical strength.¹³ Its extraordinary characteristics distinguish it from its bulk form. Besides graphene, other 2D materials like bismuthene, borophene, and their interfaces have already been extensively studied for use in lithium and non-lithium ion batteries. Various 2D materials have been extensively investigated for their application in alkali metal ion, magnesium, and zinc ion batteries, demonstrating promising characteristics such as high capacity, optimal OCV, favorable thermodynamic properties, and significant adsorption energy. These attributes highlight the potential of 2D materials as effective components in advancing battery technology. 14-18

Chalcogenides and dichalcogenides are a promising class of novel electrode materials that can be used in batteries and supercapacitors. 19,20 Compounds containing one or more chalcogen anions, such as tellurium, oxygen, selenium, and sulfur are known as dichalcogenides. Various binary and ternary chalcogenide compositions are reported in the literature.²¹ Transition metal dichalcogenides (TMDs), including chromium ditelluride, are known for their semiconducting properties. The monolayer form of CrTe2 exhibits a hexagonal lattice structure and possesses intriguing electrical and optical characteristics. Its adjustable band gap is a crucial characteristic that qualifies it for use in electronic devices. Furthermore, the monolayer of chromium ditelluride exhibits strong mechanical flexibility, thermal stability, and electrical conductivity. Due to its unique properties, the chromium ditelluride monolayer has the potential to enhance battery performance. We have previously investigated magnesium diboride (MgB₂) and tungsten ditelluride (WTe₂) for their applications in batteries and the detection of volatile organic compounds (VOCs). Our findings indicate that these materials possess significant potential for exploration in various fields, including gas sensing, spintronics, battery technology, and hydrogen storage applications.²²⁻²⁴

These attributes of the 2H phase CrTe₂ have prompted us to investigate its suitability for metal ion batteries. We concentrated on investigating the possible electrochemical characteristics of the 2H phase CrTe₂ monolayer. Adsorption of Li/K changes the semiconducting behavior of the CrTe₂ to metallic. Different characteristics such as adsorption energy, binding energy, charge analysis, density of states (DOS), band structures, OCV, storage capacity, and diffusion barrier were explored. The results showed that CrTe₂ is a hopeful anode material for Li/K ion batteries. Our research concludes that

CrTe₂ monolayers outperform other studied anode materials for Li/K-ion batteries. Their high ion storage capacities and stable metallic characteristics position them as superior candidates for advancing battery technology.

2 Computational detail

The Amsterdam Density Functional (ADF) package's DFTB and BAND components were adopted using the Slater-type Orbital (STO) approach for all of the calculations.²⁵ The Generalized Gradient Approximation (GGA) in conjunction with the Perdew-Burke-Ernzerhof (PBE) technique was utilized to determine the electrons' exchange of correlation interaction.²⁶ To depict the atomic and electronic wave functions, a thorough basis set utilizing the double zeta double polarization was used.²⁷ The convergence criteria for energy (10⁻⁵ eV), gradient- and step-convergence was set to, 0.02 eV Å and 10⁻³ Å for all structures relaxation. The structures underwent full relaxation and Self-Consistent Field (SCF) cycles.²⁸

The following formula was used to calculate the adsorption energy of the Li/K atoms on CrTe₂.²⁹

$$E_{\rm ad} = \frac{E_{\rm CrTe_2(Li/K)_n} - E_{\rm CrTe_2} - nE_{\rm Li/K}}{n} \tag{1}$$

where n is the number of Li/K atoms being adsorbed, $E_{\text{Li/K}}$ is the energy of a single Li/K atom in bulk lattice, and $E_{\text{CrTe}_2(\text{Li/K})_n}$ and E_{CrTe_2} are the overall energies of the structure with and without Li/K. Using the formula in eqn (2), the value of OCV was computed to assess the material's performance as anode.³⁰

$$OCV = \frac{E_{CrTe_2} - xE_M - E_{CrTe_2M}}{xye}$$
 (2)

where E_{CrTe_2} and E_{CrTe_2M} represent the total energies of chromium ditelluride in pure and with Li/K adsorption. E_M is the cohesive energy of the metal Li/K. whereas, y is the electronic charge (y = 1) of the Li/K ion in the electrolyte.

The following formula can be used to determine the maximum theoretical capacity (CM).³¹

$$CM = \frac{x \times Z \times F}{M_{CrTe}} \tag{3}$$

where Z is the valency of fully ionized Li/K and x is the maximal concentration of metal ions. The molecular mass of Li/K ions adsorbed onto the CrTe₂ supercell is represented by $M_{\rm CrTe_2}$, and F denotes Faraday's constant (26.801 mA h mol⁻¹).

3 Results and discussion

3.1 Structural and electronic properties of CrTe₂

Fig. 1(a–c) presents three different views of the optimized unit cell structures of the 2H phase $CrTe_2$. The unit cell of $CrTe_2$ contains one atom of Cr and 2 atoms of Te. Next, we created and optimized the $5 \times 5 \times 1$ supercell of $CrTe_2$. We optimized the 2H phase of the $CrTe_2$ monolayer with the P3m1 (156) space group presented in Fig. 1(d). In the 2H structure, two layers stack on top of each other, forming a hexagonal lattice. The atoms in each layer are arranged in a hexagonal configuration,

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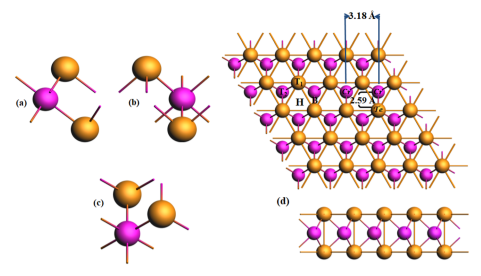


Fig. 1 The unit cell of CrTe₂ with different views (a) x (b) y (c) z and (d) optimized structure of CrTe₂ top and side view. Cr is denoted with pink and Te with golden.

and the stacking sequence repeats every two layers. Here two layers of Te atoms encircle a single layer of Cr atoms. The lattice parameter (a = b = 3.215 Å) values of MoS₂ are lower than the lattice parameter values (a = b = 3.488 Å) of CrTe₂.³² However, these are smaller than those found in the monolayers of 1T- $CrTe_2 (a = b = 3.790 \text{ Å}) \text{ and } MoTe_2 (a = b = 3.550 \text{ Å}).$ The ideal structural bond lengths for Cr-Te, Cr-Cr, and Te-Te are approximately 2.59 Å, 3.18 Å, and 3.18 Å, respectively.

Hirshfeld charge analysis revealed a notable charge transfer from Cr (1.275e) to Te atoms in the 2H-CrTe₂, increasing electrons around the Te atoms. This occurs because the Cr atom has a lower electronegativity than the Te atom.

Chromium, with its electronic configuration of [Cr] 4s¹ 3d⁵, typically forms multiple bonds influenced by its oxidation states, which range from +2 to +6, and the ligands involved. In situations where it forms more than six bonds, this involves hybridizations such as dsp² or d²sp³, utilizing its 3d, 4s, and 4p orbitals. Similarly, the electronic configuration of Te (1s² 2s² 2p⁶

3s² 3p⁶ 4s² 4p⁶ 4d¹⁰ 3d⁵ 5s²) suggests its capability to form two covalent bonds due to sp hybridization. We have applied spinpolarized calculations and found the non-magnetic nature of the CrTe₂. Our findings demonstrate that chromium ditelluride acts as the semiconductor with an indirect band gap of approximately 0.44 eV aligning with recent research. Bai et al. reported band gaps of approximately 0.52 eV using the PBE method, 0.49 eV with the PBE + SOC method, and 0.93 eV using the HSE06 functional. As mentioned in our computational details, we employed the PBE method in our calculations, so our band gap is nearly the same as the values calculated using the PBE and PBE + SOC functional. 33 Fig. 2(a) shows the DOS and (b) band structure of the pure CrTe2. Analysis of the DOS reveals that the 3d orbitals of the Cr atom significantly contribute to both the valence and conduction bands, while the 4s state contributes to the valence band. The band structure results are consistent with the DOS findings.

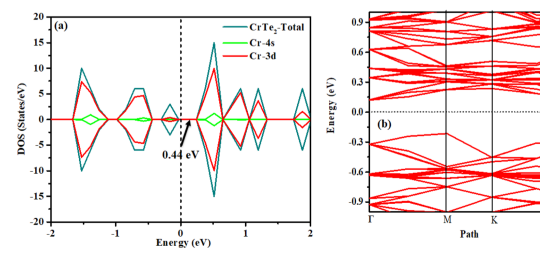


Fig. 2 Calculated (a)TDOS (b) band structure of CrTe₂.

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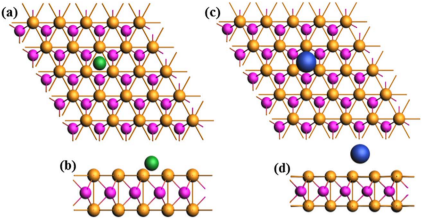


Fig. 3 Optimized structure views of Li-adsorbed CrTe₂ (a) z, (b) x, (c) Z-view of K adsorbed CrTe₂ and (d) x. The Li atom is depicted in green while K is in blue.

3.2 Electronic, and adsorption properties of single Li/K atom adsorbed CrTe2

The process of adsorption and removal of Li/K can lead to structural deformation, which aids in understanding its suitability as an anode material. The materials having high adsorption energy values are favorable for anode materials. This suggests that the materials with the highest adsorption energy are famous as excellent candidates for anode applications. The optimized structure of CrTe2 after the adsorption of one Li/K atom is shown in Fig. 3. The adsorption of Li/K onto the 2H-CrTe2 monolayer is contingent upon the charge transfers and structural environment between them to identify the most constant adsorption site. Fig. 1(d) illustrates four highly symmetric adsorption sites of CrTe2 namely (i) hollow (H site) (ii) Li/K atom adsorbed at the top of Te atom (T₁ site), (iii) Cr atom (T₂ site), (iv) between the bond of Te- and Cr (B site).

A greater negative energy value indicates a more favorable location for potassium/lithium adsorption due to a stronger host contact. Table 1 shows the Li/K adsorption energy, height of the Li/K from the host material, and Hirshfeld charges at various sites. In the analysis of stable adsorption energy sites, the H site emerges as the most stable, exhibiting the lowest (most negative) adsorption energy values for both Li and K. The Li/K adsorbed CrTe₂ comes with adsorption energies of -3.7 eV/ -4.63 eV. For adsorption site H, the heights of lithium and

Table 1 Adsorption energy, Hirshfeld charge, and height of Li/K at various adsorption sites

Materials	Sites	$E_{\rm d}$ (eV)	$\Delta Q_{ m Li/K}$	Height (Å)
Li	Н	-3.70	0.346	0.02
	T_1	-3.63	0.309	0.19
	T_2	-3.62	0.323	0.15
	В	-3.65	0.321	0.15
K	Н	-4.63	0.517	0.07
	T_1	-4.59	0.515	0.11
	T_2	-4.58	0.506	0.18
	В	-4.61	0.513	0.17

potassium are 0.02 Å, and 0.07 Å respectively. At the hexagonal ring site with the Cr atom (T₂ site), the heights of lithium and potassium are 0.15 Å, and 0.18 Å respectively. At the T₁ site on top of Te, the adsorption energy values for K and Li interactions are -3.75 eV and -4.62 eV, respectively, with corresponding heights of 0.19 Å for lithium and 0.1 Å for potassium.

Masood et al. employed first-principles calculations to explore the potential of WB4 as an anode material for rechargeable alkalimetal-ion batteries. Their results indicate that Li, Na, and K adsorption on the WB4 surface leads to remarkably high conductivity. The WB4 monolayer efficiently adsorbs these ions, with notable adsorption energies of -2.516 eV, -2.356 eV, and -2.941 eV for Li, Na, and K ions, respectively. They also explored CrB₄ and MoB₄ by adsorbing Li/K as anode material. The adsorption energies for Li and K were -1.31/-1.29 eV and -1.37/-1.31 eV for CrB₄/MoB₄, respectively.^{29,34}

The MoSe₂ monolayer has been experimentally synthesized and evaluated through first-principles calculations for its potential as an anode material in different phases (1T, 1T', and 1H). The 1T'-MoSe₂ monolayer shows excellent thermal, dynamical, and mechanical stability. With an adsorption energy of -1.254 eV, it prevents the formation of metallic Li clusters and becomes metallic under Li adsorption, ensuring efficient electron transport.35 Zhao et al. found that TiB2 monolayers exhibit excellent mechanical, dynamic, and thermal stability. The negative adsorption energies for Li, Na, and Mg atoms indicate favorable adsorption and prevent metallic dendrite or cluster formation. Notably, the adsorption energy values are -1.50, -1.38, and -1.48 eV for Li, Na, and Mg, respectively.³⁶

In conclusion, CrTe₂ possesses a hexagonal layered structure with weak van der Waals forces between the layers, allowing Li/ K atoms to intercalate more easily due to the increased space and reduced resistance compared to denser materials. Furthermore, upon Li/K adsorption, CrTe2 transitions from a semiconductor to a metallic state, which enhances its electronic mobility and conductivity. This improved electron transfer between the material and the adsorbed Li/K atoms reduces the adsorption energy, as the Li/K atoms can efficiently exchange electrons with the host material.

Hirshfeld charge transfer analysis was used to assess the interaction between Li/K and the CrTe₂; the results are displayed in Table 1. The CrTe₂ monolayer gained electronic charge from adsorbed Li/K. When a single Li/K atom is adsorbed on different sites of CrTe₂, it transfers a charge to the CrTe₂ and behaves as an electron donor. The electron transfer from lithium and potassium to the CrTe₂ indicates that Li and K atoms are adsorbed onto the CrTe₂ material, suggesting a redox reaction occurring in the electrode material during battery operation, consistent with previous studies and findings (Table 2).

At the H site, Li/K transfers 0.346e/0.517e to the atoms of the six-membered CrTe ring. For the T_2 site, Li donates 0.323e while K donates 0.506e to the CrTe ring. At the T_1 site, Li/K transfers a charge of 0.309e/0.515e, respectively. When a Li/K atom is adsorbed on the link between Cr and Te (B site), the transferred charge values are 0.321e and 0.513e. The findings indicate that the strongest contact between Li and Cr is towards Cr atoms in

Table 2 Comparison of single Li/K adsorption energy (eV) with the literature

Material	K	Li	Ref.	
CrTe ₂	-4.63	-3.7	This work	
WB_4	-2.94	-2.51	34	
CrB_4	-1.37	-1.31	29	
MoB_4	-1.31	-1.29	29	
TiC_3	_	-0.93	37	
AlB_4	_	2.14	38	

the monolayer, implying that the newly created Cr–Cr bond is more electronegative. By the adsorption energy findings, the H site appears to be the most stable due to Li/K's strong charge transfer value.

The absorption of a single Li/K atom transforms the semi-conducting nature of CrTe₂ into a conductor. Fig. 4(a and c) presents the DOS of Li/K adsorbed CrTe₂, while Fig. 4(b and d) shows the corresponding band structure. The DOS results in Fig. 4(a and c) are in good agreement with the band structure in Fig. 4(b and d).

Using the ELF, we assessed the structure's bonding properties as illustrated in Fig. 5. A molecule with an ELF value of less than 0.5 shows that it is ionic in nature whereas a high ELF value (>0.5) indicates primarily covalent bonding. An ELF value of 0.5 signifies a metallic bond.³⁹

From Fig. 5(a) purple regions around the Te atoms in host material CrTe₂ initially show covalent bonding characteristics between Cr and Te with an ELF value of 0.75. Nevertheless, following Li/K adsorption red color around the Li/K atom shows that the structure has an ionic bonding nature with an ELF value of 0.25. The structural stability is enhanced by the coexistence of ionic and covalent bonds, which is beneficial for the adsorption of Li and K ions.

3.3 Effect of lithium/potassium concentration and stability

The system's average adsorption energy was determined by gradually adding Li/K atoms to the host material using eqn (4).

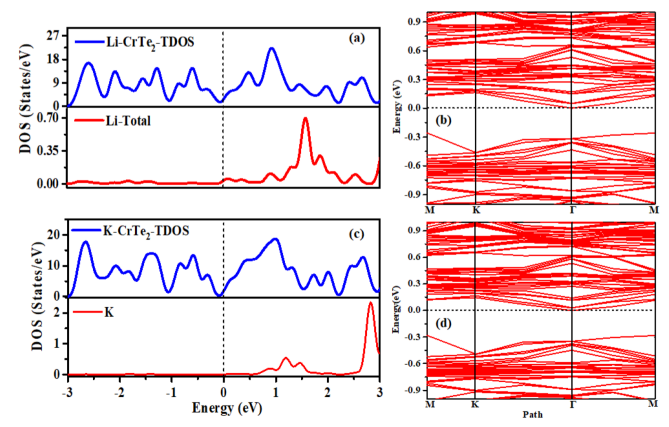


Fig. 4 (a) DOS- (b) band structure of Li-adsorbed CrTe₂ (c) DOS- (d) band structure of K-adsorbed CrTe₂.

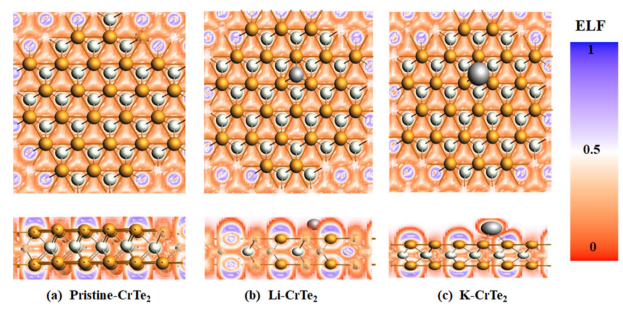


Fig. 5 The ELF maps of (a) pristine (b) Li- (c) K-adsorbed CrTe₂.

Fig. 6 shows the optimized structures of fully lithiated and potassiated CrTe₂. The concentration of Li/K atoms in CrTe₂ was gradually increased to calculate the adsorption energy by the following eqn (4).

ave
$$E_{\text{ads}} = \frac{1}{n} (E_{\text{MCrTe}_2} - E_{\text{CrTe}_2} - nE_{\text{M}})$$
 (4)

Here, E_{MCrTe_2} is the energy of the Li/K adsorbed CrTe₂, while E_{CrTe_2} is the energy of the pure CrTe₂. E_{M} is the reference energy of Li/K calculated using a bulk lattice with a bcc unit cell, where n represents the number of metal atoms.

Li/K atoms were initially positioned on the top side of the CrTe₂ at the most favorable H site and then on the lower side. 16

Li and 13 K atoms were adsorbed one by one on the studied structure. The declining trend shown in Fig. 6(a) indicates that the lithiation/potassiation energy value approaches zero as more Li/K atoms are introduced, demonstrating the feasibility of the exothermic reaction. The potassiated structure shows positive adsorption energy when the 14th K atom is adsorbed onto $CrTe_2$, indicating that the host material will not adsorb additional atoms. The adsorption energy becomes progressively more negative upon the adsorption of 17th and 18th Li. However, with the 20th and 21st Li adsorption on $CrTe_2$, the adsorption energy starts to decrease towards zero, reaching values of -0.10 and -0.05, respectively. Therefore, we halted further calculations.

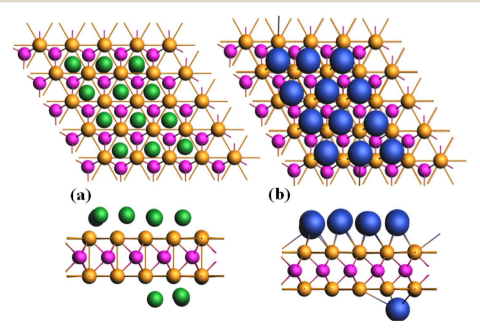


Fig. 6 The fully optimized structures (a) lithiated (b) potassiated CrTe₂.

The geometrical structures are found to be rather stable within the estimated adsorption energy ranges of -3.7 eV to -0.20 eV for Li and -4.63 eV to -0.13 eV for K, respectively, as the concentrations of Li and K grow. Its stability guarantees a structure free of dendrites, allowing for safe lithiation/potassiation.

Structural stability is determined by the change in the area after the adsorption of Li/K in $CrTe_2$. A structure is considered stable if it experiences minimal volume expansion after adsorption. To assess the effect of Li/K adsorption on $CrTe_2$, we calculate the change in area (ΔS) as a function of the concentration of Li/K, representing the volume expansion. This is calculated by eqn (5) given below.

$$\Delta S = \frac{\left(S_{\text{Li/K}}@_{\text{CrTe}_2} - S_{\text{CrTe}_2}\right)}{S_{\text{CrTe}_2}} \times 100\%$$
 (5)

where $S_{\rm Li/K@CrTe_2}$ represents the volume of fully lithiated/potassiated CrTe₂ and $S_{\rm CrTe_2}$ represents the volume of the pure CrTe₂. The adsorption of Li/K on CrTe₂ did not change the bond lengths of the structure. Consequently, no volume expansion occurs during the adsorption process. This stability improves the cycle capacity during the charging and discharging, making CrTe₂ a beneficial anode material for Li/K ion batteries.

3.4 OCV and theoretical capacity of Li/K adsorbed CrTe₂

The average OCV and theoretical capacity are critical attributes, as they reflect the higher capacity and energy density of metalion batteries in their electrochemical performance. OCV is a key feature of rechargeable ion batteries, offering insight into their functionality and storage capacity. Eqn (2) is used to compute the OCV value, and Fig. 7(b) illustrates the average OCV for Li/K at various atomic concentrations. As shown in Fig. 7(b), no negative values are observed, indicating a preference for Li/K ions to adsorb on CrTe₂ rather than forming

metallic states. This underscores the material's suitability compared to the conventional graphite electrode.⁴⁰ The concentration of Li atoms peaks near an OCV of 3.8 V. The electrode potential voltage profile ranges between 3.8 V and 0.20 V for Li and 4.6 V to 0.13 V for K, showing a decreasing trend with increasing Li content. Low OCV range anodes for high-capacity materials are essential to ensure proper operation (Fig. 8).

Eqn (3) was used to determine the maximum capacity of $CrTe_2$ upon Li/K adsorption. The maximum capacity at the highest concentration of Li/K on $CrTe_2$ is 1395 mA h g^{-1} and 1134 mA h g^{-1} , respectively. These values exceed the useable capacity of the graphite anode, which is 372 mA h g^{-1} . Furthermore, the computed capacity for K is higher than the blue phosphorene monolayer's (570 mA h g^{-1}) capacity,⁴¹ and Ti_3C_2 MXene (191 mA h g^{-1}).⁴² On the other hand, Li has a greater estimated capacity than graphene (784 mA h g^{-1}),⁴² and germanium sulphide (256 mA h g^{-1}).⁴³

In this comparison, the maximum theoretical capacity for YS_2 in a lithium-ion battery is 350 mA h g^{-1} and the diffusion barrier is 0.33.44 T Zaho et al. studied lithium-ion batteries using TiC2. Their study results show a theoretical capacity of 622 mA h g^{-1} , a voltage of 0.94 V, and a diffusion barrier (0.11 eV).45 C. Eamas et al. worked on lithium-ion batteries using Ti₂C, finding a theoretical capacity of 440 mA h g⁻¹, a voltage of 0.44 V, and a diffusion barrier (0.27 eV).46 Working with Ti₂BN monolayer, Y. Y. Wu et al. discovered a theoretical capacity of 889 mA h g^{-1} , a voltage of 0.024 V, and a diffusion barrier (0.24 eV) for lithium-ion batteries.47 H. R. Jiang et al. worked on potassium-ion batteries using BP, finding an open circuit voltage of 0.27 V and a diffusion barrier of 0.155 eV.41 Working with 2D MON₂, X. Zong et al. discovered an open circuit voltage of 1.11 V and a diffusion barrier of 0.49 eV when studying potassium-ion batteries. 48 For potassium-ion batteries, K. Dou et al. found a voltage of 0.024 V and a diffusion barrier of 0.24 eV (Table 3).49

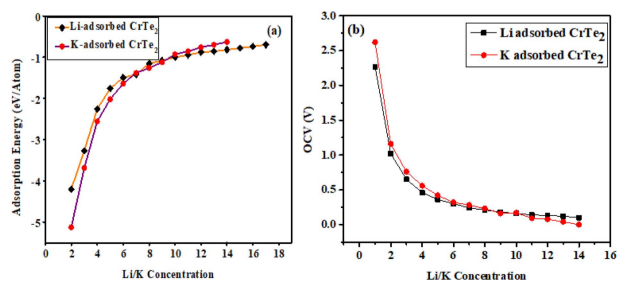


Fig. 7 (a) Average adsorption energy (b) OCV profile upon loading the Li/K on CrTe₂.

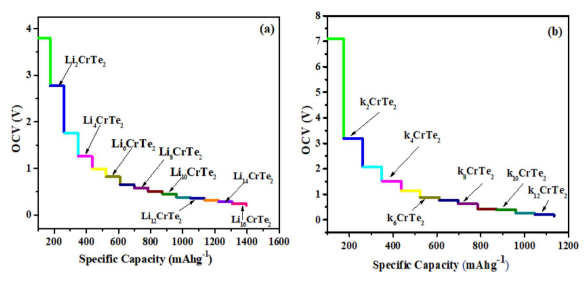


Fig. 8 Relation of OCV and specific capacity (a) Li (b) K adsorbed CrTe₂.

Table 3 An analysis of a few popular anode materials for LIBs/KIBs regarding diffusion barrier (eV), open circuit voltage (V), and specific theoretical capacity (mA h q^{-1})

Material Atom		Specific capacity (mA h g^{-1})	Diffusion barrier (eV)	OCV (V)	Reference
CrTe ₂	Li	1395	0.16	0.20	This work
YS_2	Li	350	0.33	_	44
TiC_2	Li	622	0.11	0.94	45
Ti_2C	Li	440	0.27	0.44	46
Ti ₂ BN	Li	889	0.24	0.024	47
CrTe ₂	K	1134	0.26	0.13	This work
BP	K	_	0.155	0.27	41
MoN_2	K	_	0.49	1.11	48
PC	K	_	0.26	0.69	49

3.5 Diffusion analysis

Lithium diffusion plays a crucial role in the charging and discharging rates of electrode materials, making the analysis of ion mobility essential for enhancing LIB capacity. The rate capability, particularly important for high-power applications like electric vehicles, depends on the transport of lithium ions and electronic conductivity.50,51 To address these issues, the diffusion barrier must be calculated. In this work, diffusion properties were determined using the climbing-image nudged elastic band (CI-NEB) method, which finds the highest energy point along the diffusion path.52 The two selected diffusion pathways for Li/K are the hollow site on top of Cr/Te and the hollow site (H-Cr/Te-H). The diffusion barriers were found to be 0.26 and 0.53 eV for K, and 0.16 and 0.17 eV for Li. Therefore, CrTe₂ significantly enhances Li/K mobility and improves the charging/discharging capabilities when used as an anode material in Li/K ion batteries.

Regarding thermal properties, we computed the thermal energies of K and Li atoms at temperatures of 300 K and 400 K using the following relation. 53

$$E = \frac{3}{2}K_{\rm B}T\tag{6}$$

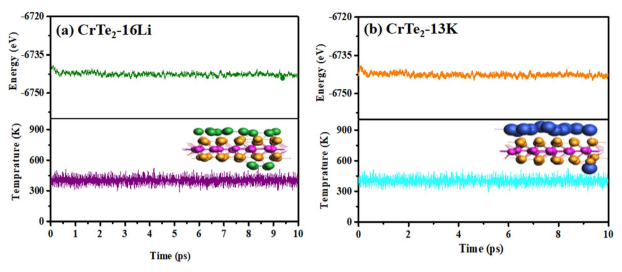
Here $K_{\rm B}$ is the Boltzmann constant and T is the temperature in Kelvin (K). The calculated thermal energies are 6.21×10^{-2} and 8.28×10^{-21} at 300 K and 400 K respectively, which are significantly lower than their corresponding diffusion barriers. This implies that the likelihood of Li and K atoms overcoming these barriers and diffusing is negligible. Based on these diffusion barriers, $CrTe_2$ -Li/K materials are energetically stable and demonstrate good rate capability, making them suitable as anodes for Li/K ion batteries.

3.6 Dynamical and thermal stability

In this research, we performed *ab initio* molecular dynamics (AIMD) simulations on a $5 \times 5 \times 1$ supercell of CrTe₂ at 400 K, to assess thermal stability. The key factors in determining the optimal Li/K concentration are the negative average adsorption energy and the material's thermal stability throughout the charging/discharging process. Structural changes with Li/K insertion, shown in Fig. 6, revealed no structural deformation during the fully lithiated/potassiated process, indicating structural stability.

The AIMD simulations over 10 ps at 400 K, illustrated in Fig. 9, evaluated the recovery dynamics of the CrTe₂ anode. The

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The energy and temperature fluctuation vs. time steps (a) Li-CrTe₂ and (b) K-CrTe₂.

minimal energy variation during this MD timescale underscores the thermal stability of Li/K-adsorbed CrTe2 at 400 K, as the structure remains intact with no indications of bond breaking, thereby confirming the monolayer's durability for Li/K. This suggests the stability and feasibility of the proposed material as an anode for Li/K ion batteries.

Conclusions

DFT calculations were employed to explore CrTe2 as an anode material for Li/K ion batteries. The binding energy values for the most stable H site are -3.7 eV for Li and -4.63 eV for K. The charge transfer from Li/K to CrTe₂ is 0.34/0.51 eV, which is essential for anode materials to prevent clustering during the adsorption process. Li/K adsorbed CrTe2 exhibits a favorable OCV of 0.20/0.13 V and a storage capacity of 1395 mA h g⁻¹ for Li and 1134 mA h g⁻¹ for K during the storage process. These values surpass those of the commercially available graphite anode material. In addition, 2H-CrTe2 has an ultra-low diffusion barrier, indicating rapid mobility of Li/K atoms. The diffusion energy barrier is 0.16 eV for Li and 0.26 eV for K. Our proposed material demonstrated excellent cycling performance due to the absence of volume shrinkage. These results suggest that 2H-CrTe₂ is a promising candidate for Li/K ion batteries.

Data availability

No datasets were generated or analysed during the current study.

Conflicts of interest

The authors declare no conflict of interest.

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