

Optimized Temperature Effect of Li-Ion Diffusion with Layer Distance in $\text{Li}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$ Cathode Materials for High Performance Li-Ion Battery

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energy density, high voltage, and long cycle life.^[2] As one of the most widely used cathode materials, $\text{LiNi}_x\text{Mn}_y\text{Co}_z\text{O}_2$ (labeled as NMC) has been investigated extensively, due to their high reversible capacity, good environmental compatibility, and relatively high Li-ion diffusivity. In the previous works, different kinds of NMC materials with different content ratio of Ni, Co, and Mn have been developed, and their electrochemical properties have also been studied, such as $\text{Li}(\text{Ni}_{1/3}\text{Mn}_{1/3}\text{Co}_{1/3})\text{O}_2$ (111),^[3,4] $\text{Li}(\text{Ni}_{0.4}\text{Mn}_{0.4}\text{Co}_{0.2})\text{O}_2$ (442),^[5] $\text{Li}(\text{Ni}_{0.42}\text{Mn}_{0.42}\text{Co}_{0.16})\text{O}_2$ (552),^[6] $\text{Li}(\text{Ni}_{0.5}\text{Mn}_{0.3}\text{Co}_{0.2})\text{O}_2$ (532),^[7] $\text{Li}(\text{Ni}_{0.6}\text{Mn}_{0.2}\text{Co}_{0.2})\text{O}_2$ (622),^[8] and $\text{Li}(\text{Ni}_{0.7}\text{Mn}_{0.15}\text{Co}_{0.15})\text{O}_2$ (71515).^[9] For example, Noh et al. compared the electrochemical properties including the Li-ion diffusion coefficient, capacity retention, and electrochemical stabilities (25 to 55 °C) of layered NMC

cathode materials ((111), (532), (622), (71515), (811) and $\text{Li}(\text{Ni}_{0.85}\text{Mn}_{0.075}\text{Co}_{0.075})\text{O}_2$) at room temperature and found that the Ni content had a great influence on the electrochemical properties.^[10]

Solid phase diffusion coefficient (D_s) is one of the most important parameters for the active materials of the LIBs, as it determines the charge and discharge rate capability directly. In particular, for high power density applications, fast Li-ion transport in cathode materials is a key factor and must be needed. As a result, many experimental and theoretical works have been devoted to investigating the Li-ion diffusion properties in layered cathode materials.^[11,12] However, to the best of our knowledge, there is little work reported to study the relationship between the layer distance and kinetics of Li-ion diffusion in different temperatures of layered NMC cathode materials

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the distribution of the NMC particle size, the jitter of the cell voltage, and so on, may cause the measurement errors, however, no better methods can replace these methods mentioned above. Despite the existence of the errors and the disadvantage of long-time waiting, the results measured by GITT are still well accepted by scientists up to now. Besides, it is easier to implement compared with the other methods. Meanwhile, as different previous researches used different experiment conditions, different methods and different instruments, the GITT results always had a few orders of magnitude differences, and it is also difficult to compare the diffusion properties of different NMC materials from previous reported works. Therefore, it is hard to distinguish that which material has the best Li-ion diffusivity.

Here, we combined with experiments and ab initio calculations to investigate the Li-ion diffusion of NMC materials in different temperatures at the different states of Li-ion de-intercalation systematically for the first time. The D_s values of a series of different NMC materials (111, 442, 552, 532, 622, and 71515) at different temperatures from $-25\text{ }^{\circ}\text{C}$ (248 K) to $50\text{ }^{\circ}\text{C}$ (323 K) are measured by using the improved GITT method, which is developed to employ an improved model to reduce the system error of GITT method with a correction of NMC particle size distribution. After comparing the kinetics properties of different NMC materials, it is found that 622 shows the highest Li-ion diffusivity with the minimum temperature dependence. Furthermore, both ab initio calculations and X-ray diffraction (XRD) measurements suggest that the Li slab space of the NMC layered material plays a great role on the Li-ion diffusion property versus temperature.

2. Results and Discussion

Figure 1a shows the Scanning electron microscope (SEM) of 622. As the form of different NMC materials is very similar, we take 622 as an example. The spherical particles are the secondary particles which consist of a lot of primary particles with a small grain for each of them. Besides, the SEM indicates that the secondary particles size is heterogeneous and most secondary particles have a diameter about $5\text{--}20\text{ }\mu\text{m}$, as shown in Figure 1b.

Figure 2a–c shows the Li-ion diffusion coefficient (D_s) versus different state of Li-ion de-intercalation (δ) of $\text{Li}_{1-\delta}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$ ($0 \leq \delta \leq 1$) at 50, 25, and $0\text{ }^{\circ}\text{C}$, respectively. The Li-ion diffusion coefficients of all the NMC materials are in the range 10^{-11} to $10^{-13}\text{ m}^2\text{ s}^{-1}$.

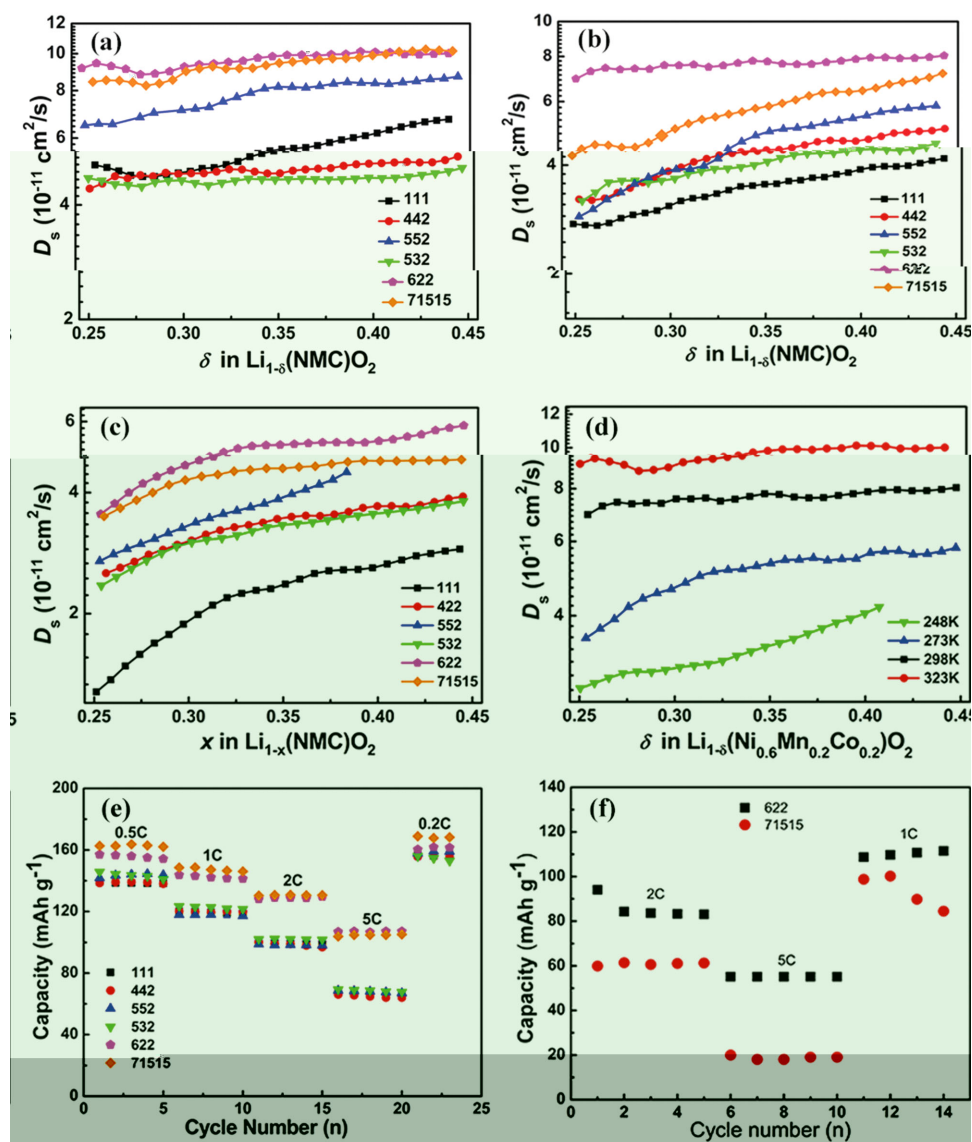


Figure 2. Diffusion coefficient of different materials versus state of charge in a) 50 °C (323 K), b) 25 °C (298 K), c) 0 °C (273 K), and d) shows D_s of 622 in different temperature. Rate capability of e) all NMC materials at 25 °C and f) 622 and 71515 at 0 °C.

$$\log(D_s) = \log(D_0) - \frac{E_a}{k} \frac{1}{T} \quad (1)$$

E_a (eV) is the activation energy for diffusion, k (eV K⁻¹) is the Boltzmann constant, and D_0 is the diffusion coefficient when the temperature goes to infinite.

The calculation and GITT results of 532 and 622 at $\delta = 0.33$ are shown in **Figure 3a,b**, respectively, in which all curves show almost linear and the slope of 532 is slightly larger than that of 622, suggesting that the value of E_a of 532 is larger so as to be easier to be affected by the temperature. Note that the Li-ion diffusion coefficient of calculation is 1–2 orders of magnitude larger than that of GITT, which is mainly attributed to the systematic error which exists between the experiment and the calculation (see Section 4.1). Although the value of the slope from calculation is much bigger than that of measurement, the

variation trends of the Li-ion diffusion coefficients of 532 and 622 by measurement and calculation are same.

The question is why the Li-ion diffusion coefficient of 622 is larger than that of 532 and other NMCs to be the largest with the minimum temperature effect in all the NMC materials. The mechanism is investigated according to relationship of layer distance structure versus Li-ion diffusion property by combination with the experimental measurements and ab initio calculations as below.

As shown in Equation (1), the Li-ion diffusion coefficient is related to the effective activation energy E_a . As indicated in our previous work,^[12] there are two kinds of Li-ion diffusion pathways in such layered structures, namely, Li-ion diffuses from one octahedral site to next site through the oxygen dumbbell or an intermediate tetrahedral site, in which Li-ions tend to choose oxygen dumbbell hopping at the early stage of

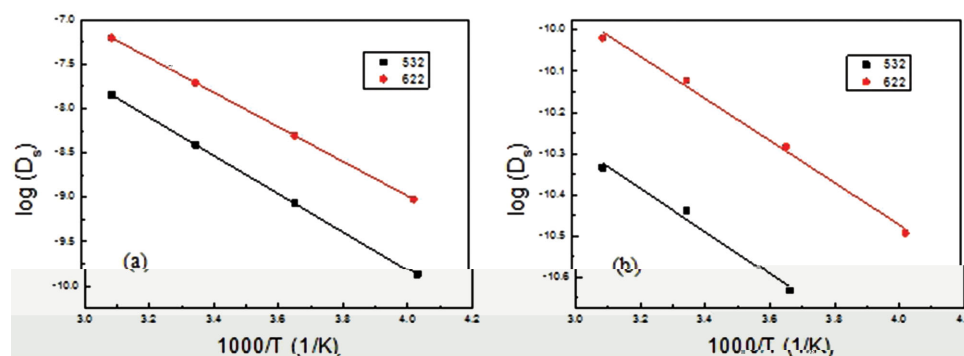


Figure 3. Diffusion coefficient comparison of 532 and 622 at $\delta = 0.33$ in a) ab initio calculation, b) GITT test.

charging (delithiation), and tetrahedral site hopping (TSH) begins to dominate when more than 1/3 Li-ions ($\delta > 0.33$) are extracted (Figure 4). The size of the oxygen dumbbell or the tetrahedral site (strain effect) as well as the electrostatic interaction between Li-ion in the activated state and the transition metal cation would contribute to the activation energy. As a result, the activation energy of Li-ions along this pathway is closely associated with the size of Li–O tetrahedrons and octahedrons. The oxygen atoms from upper and under layer would affect the diffusion barrier of Li. Li-ions that diffuse in narrow pathway are under larger forces, which leads to higher activation energy. In Figure 4b (TSH route), Li-ion diffuses from position 1 to position 3. When Li-ion is in position 2, the distance of Li and O atoms affects the work that the system has to do to transfer the Li-ion across this O atoms tetrahedron. Smaller Li slab space would decrease the size of the O atoms tetrahedron, while the energy barrier of this diffusion pathway would increase. The distance between these two O atoms is closely associated with the activation energy of this pathway. As a result, the activation barrier for Li-ion diffusion depends on the Li slab space (a type of the strain effect) and the kind of transition metals and their valence states. Previous studies^[12] and Kang^[20] reported that more Ni content with low valence state and large Li slab space are beneficial for Li-ion diffusion. Especially, the activation barrier is sensitive to the Li slab space. When Li slab space increases by 0.1 Å, activation energy along this pathway rises by 100 meV.^[20] Here, the Ni^{3+} content is the same for 532 and 622 at $\delta = 0.33$ and 0.5. Therefore, the difference of Li-ion diffusion coefficients between the two kinds of materials must come from the difference of Li slab spaces.

As shown in Figure 4a, Li slabs and transition metal oxide slabs are distributed alternatively, in which d_1 is the thickness of transition metal oxide slab, and d_2 is the Li slab space, d_{001} is the distance of (001) plane. The relationship between d_1 and d_2 is given as

$$d_1 + d_2 = d_{001} \quad (2)$$

The XRD patterns of the six NMC materials we tested are shown in Figure 5a. The materials have a highly crystalline layered structure, with diffraction peaks corresponding to the R-3m space group. The splitting of the (108), (110) and (006), (102) peaks indicated well-ordered $\alpha\text{-NaFeO}_2$ structure. The (003) peaks of different NMCs are almost at the same 2θ . Based the (003) peaks of XRD, we can find that the distances of (001) plane (d_{001} , the layer distance of NMC shown in Figure 4a) of all the NMCs are almost same around 4.74 (± 0.01) Å at $\delta = 0$ as shown in Table 1, in which chemical lithium de-intercalation is used to control δ . Figure 5b,c shows the XRD of 532 and 622 at different δ , respectively. As δ increases, the (003) peak shifts as shown in the inset graphs. A summary of distances of (001) planes of all the NMCs with different δ by XRD is shown in

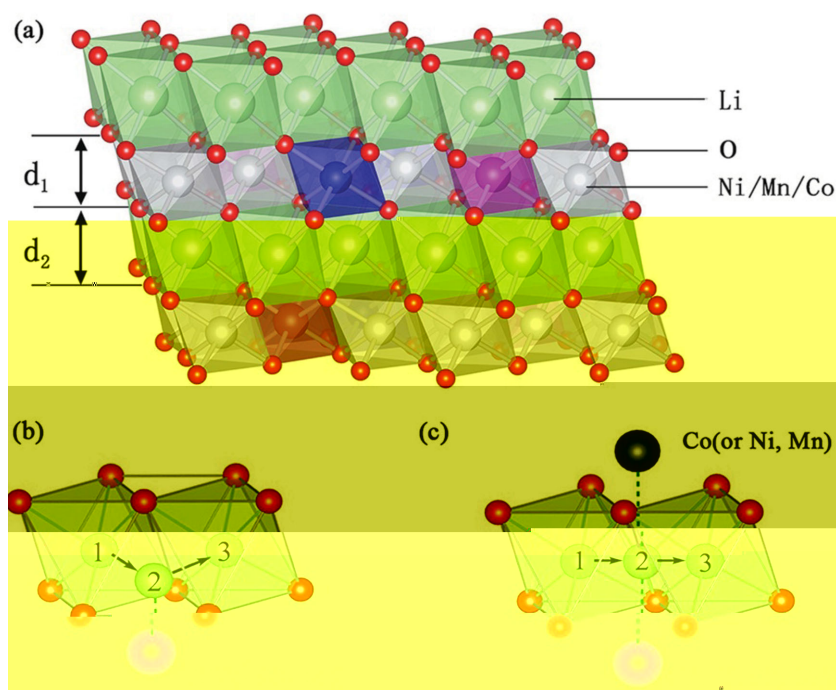


Figure 4. a) Lattice of NMC layered structure. Green atoms, Li; red atoms, O; silver/purple/blue atoms, Ni/Mn/Co transition metals. b) Tetrahedral site pathway and c) oxygen dumbbell pathway for Li-ion diffusion in NMC layered structure.

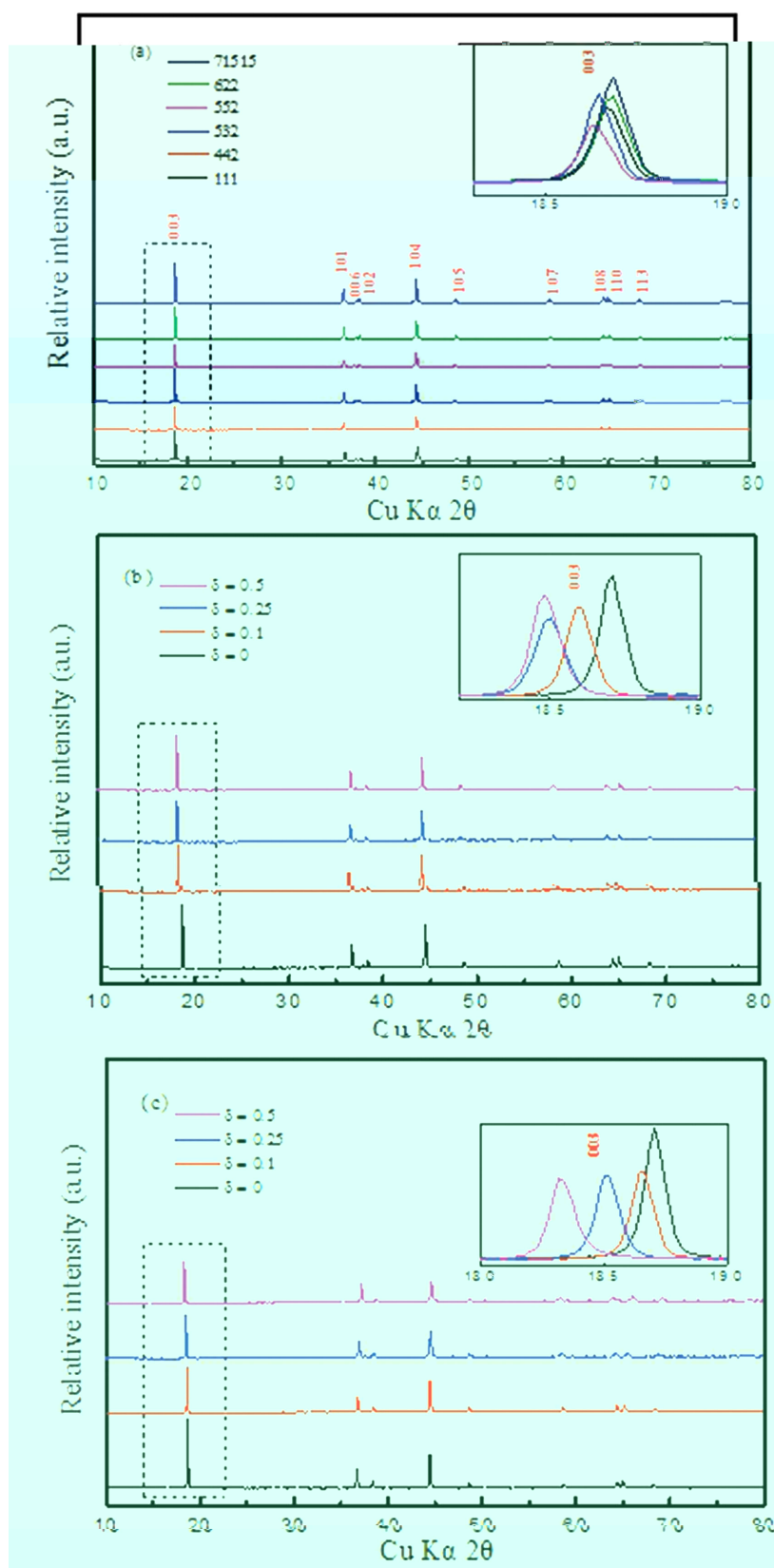


Figure 5. a) XRD of NMC materials. XRD of b) 532 and c) 622 after chemical lithium de-intercalation. The data in the dashed line box is magnified in the inset graph.

Table 1, in which displacements of the (001) peaks shift to lower diffraction angles, meaning that the (001) plane distances of all the NMCs increase with Li-ion de-intercalation process (increasing δ). Note that the (001) plane distances of 622 increase more than that of 532 with the increasing δ .

By using ab initio calculation models, the (001) plane distances and Li slab spaces of different NMCs are calculated as shown in **Tables 2 and 3**, respectively. Note that measured (001) plane distances of 333, 442, and 552 around $4.74 (\pm 0.01)$ Å are a little bit larger than those of calculation around $4.68 (\pm 0.02)$ Å, in which $x(\text{Ni}) = y(\text{Mn})$ in $\text{Li}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$. However, measured and calculated (001) plane distances of NMC 532, 622, 71515 are very close around $4.73\text{--}4.78$ Å at $\delta = 0$ (see **Tables 1 and 2**), in which $x(\text{Ni}) > y(\text{Mn})$ in $\text{Li}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$. The difference of transition metal oxide slab is caused by ionic radius of Ni, Co, and Mn. According to Equation (2), when transition metal oxide slab is smaller, Li slab would share larger space. The calculation data of transition metal oxide slabs at $\delta = 0, 0.33$, and 0.5 is shown in Table 3. (110) plane distance measured by XRD at $\delta = 0, 0.1, 0.25$, and 0.5 can indirectly indicate distances of transition metal oxidation octahedral inside their slabs as shown in Table S4 (Supporting Information).

In our experiments, we found that with the increase of δ , d_{001} of all NMCs become bigger (see Table 1), and the calculations also show that d_{001} increases with δ (Table 2). However, transition metal oxide slabs (d_1) of NMCs decrease with the increase of δ because the Ni ionic radius decreases during the Ni oxidation from Ni^{2+} to Ni^{3+} , or Ni^{3+} to Ni^{4+} . d_1 and d_2 of NMC materials are calculated, as shown in Table 3, in which the transition slab space (d_1) decreases as the increasing of δ . According to Equation (2), Li slab space increases with the increasing of δ , which is consistency with the variation of d_2 in Table 3. Note that d_1 of 111 is the smallest. That is because in NMC111, the content of Mn^{4+} and Co^{3+} is the largest among all the NMC materials. Due to the smaller size of Mn^{4+} (53 pm) and Co^{3+} (54.5 pm) compared with Ni^{3+} (60 pm) and Ni^{2+} (69 pm), NMC111 has the smallest transition slab space (d_1). These calculation and measurement data indicate that transition metal oxide slab (d_1) of 622 is the smallest among all the NMC materials except 111, while the Li slab (d_2) of 622 is largest comparing to that of 532 and 71515 (see Table 3).

The question is why the layer distance (d_{001}) and the Li slab (d_2) of 622 at various Li-ion de-intercalation (δ) are larger than that of other NMCs. As Ni content rises in NMC material

Table 1. The distance of (001) plane of a series of NMC materials analyzed from XRD. The value of the error is 0.01 Å.

δ	(001) plane distance [Å]					
	333	442	552	532	622	71515
0	4.729	4.751	4.745	4.753	4.735	4.734
0.1	4.773	4.777	4.783	4.769	4.754	4.757
0.25	4.798	4.790	4.797	4.795	4.791	4.777
0.5	4.819	4.810	4.812	4.799	4.839	4.828

Table 2. The (001) plane distance in different NMC with $\delta = 0, 0.33$, and 0.5 from ab initio calculations.

δ	(001) plane distance [Å]					
	333	442	552	532	622	71515
0	4.664	4.705	4.686	4.766	4.785	4.768
0.33	4.672	4.721	4.708	4.778	4.792	4.781
0.5	4.678	4.729	4.717	4.788	4.797	4.787

Table 3. Transition slab space (d_1) and Li slab space (d_2) in ab initio calculation with $\delta = 0, 0.33$, and 0.5 .

	333	442	552	532	622	71515
d_1 [Å] (Calculated)	$\delta = 0$	2.040	2.091	2.076	2.154	2.084
	$\delta = 0.33$	2.032	2.080	2.046	2.099	2.076
	$\delta = 0.5$	2.028	2.073	2.033	2.069	2.065
d_2 [Å] (Calculated)	$\delta = 0$	2.624	2.614	2.610	2.612	2.701
	$\delta = 0.33$	2.640	2.641	2.662	2.679	2.716
	$\delta = 0.5$	2.650	2.656	2.684	2.719	2.732

Table 4. The average activation energy of Li-ions in different NMC.

	Average activation energy [meV]				
	111	442	532	622	71515
$\delta = 0.33$	514	495	503	475	484
$\delta = 0.5$			471	457	

from 532–622–71515–811, average transition ions radius increase because ionic radius of Ni^{2+} and Ni^{3+} is higher than those of Co^{3+} and Mn^{4+} as shown in Table S1 (Supporting Information). However, as the valence of Ni rises, it causes the average radius of Ni to decrease. These two mechanisms compete with each other, and there is a balance point near 622 which results in that Li slab space in 622 is larger than those in 532 and 71515.

The height of Li–O octahedrons is used to indicate the Li slab space. So the size of Li–O octahedrons and tetrahedrons along the Li-ions diffusion pathway is closely associated with

the activation energy of Li-ions. For example, if the size of Li–O octahedrons reduces, the interaction between Li atoms and O atoms would increase. Therefore, Li-ions in 622 can diffuse easily due to the larger size of Li–O octahedrons and tetrahedrons than other NMC materials. On the other hand, Li slab space changes with the δ variation. Li-ions diffusion is accompanied by electrons transfer and redox of transition metal ions, and Li slab gets larger in the early stage of charging process due to the removal of $\text{O}^{2-}\text{--Li}^+\text{--O}^{2-}$ bonds across the slab, which is consistent with the variation of d_2 in Table 3. The activation energies of Li-ions diffusion of 532, 622 are calculated when $\delta = 0.33$ and 0.5 along TSH pathway with Ni atom nearby as shown in Table 4, along which diffusion pathways with Ni atom are the advantageous pathways with low activation energy in NMC materials.^[12] Obviously, the activation energy of 622 is much lower than that of 532 due to the larger layer distances.

Temperature effects of D_s in NMC materials are shown in Figure 2, in which the variation of D_s at 0°C is larger than those at 50°C indicating that the Li-ion diffusion is more likely to rely on the Li slab space at low temperature, because the thermal motion of Li-ions is one of the factors that facilitate the diffusion process. Ions at high temperature can diffuse spontaneously even self-destruct. It can be concluded that D_s is dominated by the increase of δ at low temperature such as 0°C . When the temperature rise exceeds a threshold value, D_s becomes not so sensitive to δ and the variation of D_s is smaller when δ increases. Interestingly, the curve of D_s versus δ variation is flatter for 622 compared to other NMC materials (Figure 2b) at each temperature. The spacing between Li slabs is one of the important factors that affects the activation energy.^[12,20] If the spacing is enlarged, the energy barrier will be lowered, so Li-ions can migrate more easily to lead to less sensitive to the Li slab space variation during the de-intercalation process. Therefore, 622 is the most promising NMC materials, for it has a larger

Li-ion diffusion coefficient, the minimum temperature dependence, a better thermal adaptability and a smaller δ dependence.

3. Conclusion

Li-ion diffusion properties versus layer distance for a series of NMC materials at different temperatures are studied systematically for the first time. Among all the NMC materials, the Li-ion diffusion coefficient of NMC622 is the largest with a smaller fluctuation with the temperature. Our experimental measurements

and ab initio calculations suggest that the large Li slab space accounts for the large Li-ion diffusivity with lower the activation energy along the diffusion pathway. This study reveals that a large Li slab space would facilitate the Li-ion diffusion with less temperature dependence and also predicts that 622 is the most promising materials for LIBs applied in multi-temperature environments.

4. Experimental Section

The series of NMC were synthesized by the co-precipitation method followed by solid-state reaction.^[21] The 3%–5% excess of lithium was added in order to make the molar ratio for Li / (Ni + Mn + Co) around unity. The sintering procedure was done at 800 °C for 8–12 h and then the sample was quickly cooled down to the room temperature. The synthesis conditions for these NMC cathode materials (111, 442, 552, 532, 622, and 71515) were the same, so the influence of the synthesis temperature to the *d*-space and diffusion properties is little. Electrochemical experiments were performed using cells consisting of cathode electrode made of NMC, anode electrode of metal Li and electrolyte of EC: DEC.

Electrochemical measurements were carried out by MACCOR battery test cabinet (Model MC-16 Battery Test System, 5 V, 5 A). The electrochemical cells are consisted of NMC and Li metal as the positive electrode and negative electrode, respectively. The electrolyte was LiPF₆ in EC: DEC mixture solvent. During the GITT experiments, the cells were kept in a thermostatic tank (RKS-80L, 220 V, 50 Hz) to guarantee that the temperature was constant during the measurements. Prior to the measurements, all the cells were activated by 3 charge/discharge cycles, and then they were charged again to 4.35 V for the GITT measurements. During the GITT measurements, the cells were repeatedly discharged for 15 min at 0.05 C and rested for 45 min, until the voltage reached 3.3 V. SEM (ZEISS SUPRA 55) and laser particle analyzer (Mastersizer3000E) were used to determine the homogeneity and distribution of the particle diameters. X-ray diffraction (D8 Advance) was used to measure the spacing between layer distances of the NMC materials at different state of Li-ion deintercalation of Li_{1- δ} (Ni _{x} Mn _{y} Co _{z})O₂ (0 ≤ δ ≤ 1). Element

Ab Initio Calculation Details: As the accuracy of the GITT model has its limitation, the accurate data provided by GITT is a finite region ($0.2 < \delta < 0.5$).^[16] Therefore, first-principle calculation is needed to serve as a necessary supplement. All ab initio calculations were performed using the Vienna ab initio simulation package (VASP).^[23] Energies used for Li mobility study are calculated with the spin-polarized generalized gradient approximation to density functional theory, using the Perdew–Burke–Ernzerhof (PBE) exchange–correlation parameterization.^[24] Compared with PBE, the newer WC can give improvements for lattice constants, crystal structures, and metal surface energies for solids.^[25] For cohesive energies, WC is nearly as accurate as PBE. As PBE can well deal with layered $\text{Li}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$ cathode materials (such as crystal structure, cohesive energy, and electronic structures),^[26,27] we also adopt it in this work. Another reason is that WC is not implemented in VASP package, which is the most widely used ab initio package in studies of lithium batteries. Not only the PW91 and PBE functional usually can give reasonable results for battery materials, but also many well established methods are also implemented in VASP package. For example, ab initio MD and climbing-image nudged elastic band (cNEB) method^[28] are two useful tools to investigate the Li-ion diffusion properties.

PBE + U approach is performed to calculate accurate structural properties and electronic properties with spin polarization. The U value of Ni, Mn, and Co is set to be 6.4, 3.5, and 3.3 eV, respectively. U values are applied based on the simulation parameters of $\text{Li}(\text{Ni}_x\text{Mn}_y\text{Co}_z)\text{O}_2$ models from the Materials Project website (www.materialsproject.org). All the layered structure in this work are fully relaxed and optimized to reach their minimum energy with space group R3m. The plane wave cutoff energy is 450 eV. Based on the reciprocal lattice size of NMC models, k -points is set from $4 \times 4 \times 2$ to $6 \times 6 \times 2$, which is used for sampling in Brillouin zone. Structures with different arrangement of transition metal atoms in each layer have been tested to find those structures with the lowest total energy in every NMC materials, which make sure that all models in our calculation are the most energetically favorable structure of NMC materials. According to previous report,^[26] we apply antiferromagnetic structure in our NMC layered structure to relax the lattice and calculate structural properties instead of other magnetic structures. Various experiments and calculations show that the transition metal compounds exhibit some kinds of ordering,^[6,29,30] such as ower or honeycomb ordering in $\text{LiMn}_{1/2}\text{Ni}_{1/2}\text{O}_2$.^[29] Taking 622 as an example, we calculated two random distribution structures. The calculated lattice parameters of them (a , b , and c) are 5.730, 12.436, and 14.021 Å and 5.740, 12.385, and 14.017 Å, respectively. The energy of the former is –659.094 eV, and the later is –659.424 eV. Comparing the lattice parameters to the ordered structure whose lattice constants, a , b , and c , are 5.781, 12.382, and 14.100 Å, respectively, we can find the difference is not significant. But the ordered structure has an obviously lower energy, –661.157 eV. Thus, in our calculations, different ordering of transition metal ions in $\text{LiMn}_x\text{Ni}_y\text{Co}_z\text{O}_2$ have been tried to determine the ions arrangement in the slab. As all the ions in NMC layered structure are fully relaxed to the minimum energy, the lattice in our models is the most stable and energetically favorable structure. Structural properties can be read from these structures, such as Li slab space and transition metal slab space.

The Li-ion diffusion coefficients for these materials mentioned above were calculated by the ab initio MD simulation, which was implemented in VASP. MSD (Mean square displacement) analysis could be used to find the MSD versus time through the data from MD simulation. Time step was set to be 3 fs in MD analysis

$$\text{MSD}_i = |r_i - r_0|^2 = (x_i^2 + y_i^2 + z_i^2) - (x_0^2 + y_0^2 + z_0^2) \quad (6)$$

$$D = \lim_{t \rightarrow \infty} \frac{1}{2d} |r_i - r_0|^2 \quad (7)$$

where x_i , y_i , z_i refer to the current position of atoms and x_0 , y_0 , z_0 stand for the original site of atoms, $\frac{1}{2d}$ is the constant related to diffusion dimension and $d = 2$ in layered materials for 2D diffusion.

As a result of diffusion between Li-ions position, the diffusion coefficient depends on the barrier energy between Li-ions, as it is shown in Equation (8)

$$D_s = D_0 e^{-\frac{E_a}{kT}} \quad (8)$$

It should be noticed that E_a is effective activation energy instead of actual activation energy to describe some energy barrier along diffusion pathways. In NMC materials, there is no observable structure displacement below 600 K. As in Equation (8), $\log(D_s)$ is linear to $1/T$, to compare the results from first-principle calculation with experimental results, linear fitting was applied to reach the diffusion coefficient at room temperature.

Also, cNEB method was performed to investigate the migration barrier of Li-ions in NMC layered materials.^[28] cNEB was a complementary to NEB method. In this work, cNEB was applied in a $3 \times 3 \times 1$ supercell with five images as intermediate states. A series of images along the migration pathway were fully relaxed with a spring constant as constraint condition. Total energy of these series of intermediate states were calculated to describe the migration barrier along this pathway.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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