Thermal runaway of Lithium-ion batteries employing LiN(SO$_2$F)$_2$-based concentrated electrolytes

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Concentrated electrolytes usually demonstrate good electrochemical performance and thermal stability, and are also supposed to be promising when it comes to improving the safety of lithium-ion batteries due to their low flammability. Here, we show that LiN(SO$_2$F)$_2$-based concentrated electrolytes are incapable of solving the safety issues of lithium-ion batteries. To illustrate, a mechanism based on battery material and characterizations reveals that the tremendous heat in lithium-ion batteries is released due to the reaction between the lithiated graphite and LiN(SO$_2$F)$_2$ triggered thermal runaway of batteries, even if the concentrated electrolyte is non-flammable or low-flammable. Generally, the flammability of an electrolyte represents its behaviors when oxidized by oxygen, while it is the electrolyte reduction that triggers the chain of exothermic reactions in a battery. Thus, this study lights the way to a deeper understanding of the thermal runaway mechanism in batteries as well as the design philosophy of electrolytes for safer lithium-ion batteries.
The application of large-format lithium-ion batteries (LIBs) with high energy density in electric vehicles requires a high level of battery safety, as the fires and explosions of batteries that take place during thermal runaway pose serious threats to the lives of passengers and property. Thermal runaway is driven by a chain of exothermic reactions that spontaneously increase the temperature of lithium-ion batteries. As a result, severe redox exothermic reactions are likely triggered at relatively high temperatures, generating a tremendous amount of heat and leading to uncontrollable rise in temperature. Thus, the removal or reduction of the main exothermic reactions during the evolution of thermal runaway is essential to guarantee the safety of lithium-ion batteries.

Flammable carbonate-based electrolytes have been widely used in commercial LIBs, and they are considered to be responsible for thermal runaway, which is reasonable, as they are among the main fuels that increase vigorous combustion. Thus, non-flammable electrolytes are considered and thought to improve battery safety. Concentrated electrolytes and localized or aqueous concentrated electrolytes have been receiving considerable attention in recent years, due to their high electrochemical performance, low volatility, and lowflammability. Many fundamental studies revealed that the unique solution structure, where all the solvent molecules and even anions are involved in the solvation sheath, endow concentrated electrolytes with different properties in comparison with the same compositional but traditional 1 M electrolytes. For example, in a battery, when the molar ratio of lithium bisamide (fluorosulfonyl) [(LiN(SO2F)2, LiFSI) to the solvent is 1:1.9, the predominant contact ion pairs (CIPs, an anion coordinating to one Li+) and aggregates (AGGs, an anion coordinating to two or more Li+) in the solvated structures guarantee the stable cycling and improve thermal properties of LIBs. Meanwhile, fire retardant solvents, such as trimethyl phosphate (TMP), triethyl phosphate (TEP), and triphosphate (trifluoroethyl) (TFEP), can be employed to develop non-flammable electrolytes due to the changes in the interface reactions that are led by the unique solvation structure of concentrated electrolytes. However, although the high thermal stability of concentrated electrolytes, which enhances battery safety as implied by the researchers, has been widely proven by ignition tests and thermogravimetric analyses (TGA), no battery safety as implied by the researchers, has been widely proven by ignition tests and thermogravimetric analyses (TGA), and mass spectrometry (DSC-TG-MS) experiments to evaluate the safety performance of graphite-LiNi0.5Co0.2Mn0.3O2 (Gr|NMC811) and Gr|NMC532 batteries with concentrated electrolytes. The thermal runaway mechanism at both the cell and material level is systematically investigated. The most promising concentrated electrolytes, LiFSI/DMC (1:1.9 by molar) and LiFSI/TMP (1:1.9 by molar), are selected in this study. The results show that the non-flammable concentrated electrolytes could not prevent LIBs from thermal runaway. Specifically, the charged anode reacts with LiFSI and releases a considerable amount of heat that triggered thermal runaway.

Results

Cycling and thermal properties of the electrolytes. Figure 1a shows that the 0.93Ah Gr|NMC811 pouch cell with the concentrated LiFSI/DMC electrolyte delivered stable charge-discharge capacities for over 300 cycles at C/3. The average coulombic efficiency was 96.6% and capacity retention was 94.5% (Fig. 1b), indicated the suppressed Al dissolution and stable SEI on the graphite during cycling. The cell with conventional 1 M LiFePO4/EC:EMC (3.7 by volume) electrolyte shows comparable electrochemical performances, the capacity retention after 300 cycles was 93.9%. The LiFSI/TMP concentrated electrolyte in the pouch cell was also investigated for two cycles before safety evaluation, and it delivered a coulombic efficiency of 99.5% (Fig. 1a). For the Gr|NMC532 pouch cells, the charge and discharge curves also demonstrated stable electrochemical performance with the concentrated electrolytes (Supplementary Fig. 1 and Supplementary Note 1).

The TGA curves (see Fig. 1c) show that the weight loss of the LiFSI/TMP concentrated electrolyte was only 0.7 wt% below 180 °C, which is considerably lower than that of the LiFSI/DMC electrolyte (18.2 wt%) and the diluted carbonate electrolyte (26.5 wt%). These results also indicate that the flammability of the LiFSI/DMC concentrated electrolyte is lower than that of the dilute conventional electrolyte, as less solvent was used and the vitality of the DMC was significantly changed by the solvation structure. Then, Fig. 1d shows the photographs of the separators saturated with the electrolytes during the ignition test. In comparison with the conventional electrolyte, the concentrated electrolyte with the DMC was still flammable but with a mild flame, whereas the concentrated electrolyte with the self-extinguishing solvent of the TMP did not burn completely, which proves that the concentrated LiFSI/TMP is non-flammable (see details in Supplementary Table 1 and Supplementary Note 2). According to the above thermal evaluation, concentrated electrolytes display better thermal stability and possibly lower flammability than dilute electrolytes, which is in agreement with the previous reports.

However, it was reported that the direct redox reactions between the charged cathode and anode are severe for battery chemistries of...
high energy density, which can cause thermal runaway even without electrolytes or ISC\textsuperscript{12,13}. Thus, the evaluation of battery safety based on the used electrolytes only is not sufficient, and the interaction between the electrolytes and the charged electrodes should be systematically taken into consideration.

**Safety characterization of LiFSI/DMC in Gr|NMC batteries.**

The thermal runaway features of Gr|NMC batteries with concentrated LiFSI/DMC and conventional 1 M LiPF\textsubscript{6}/EC:EMC electrolytes are compared in Fig. 2. It is observed that all the batteries were brought to the point of thermal runaway although the thermal stability of the concentrated electrolyte was obviously higher. The three characteristic temperatures of \{\(T_1, T_2, T_3\)\} were defined to describe the thermal behavior of the batteries with different electrolytes\textsuperscript{2,4,11,13}. In this study, in case of the concentrated and conventional electrolytes, \(T_1\) was located at 130 °C, and \(T_3\) was between 650 °C and 730 °C, whereas \(T_2\) exhibited a totally different value (Fig. 2). \(T_2\) was defined as the trigger temperature of the thermal runaway. At and after this critical point, the battery temperature exponentially increased and could not be shut down by any heat dissipation measures. Both the severe exothermic reactions and the ISC might be the underlying reasons. If \(T_2\) is caused by a single chemical reaction or a group of chemical reactions, this or these reactions can be defined as the trigger reaction of thermal runaway. The tremendous heat generated at \(T_2\) by the trigger reaction would immediately trigger a single exothermic reaction or a group of new exothermic reactions between the battery components, causing a dramatic rise (hundreds of degrees per second) of battery temperature. Understanding the mechanisms behind \(T_2\) is crucial for design of safer lithium-ion batteries.

For the Gr|NMC811 battery with the LiFSI/DMC concentrated electrolyte (Fig. 2a), \(T_2\) was located at 200.5 °C. A fall in the OCV took place at \(T_2\), which coincides with the sharp temperature rise. However, \(T_2\) of the cell with the conventional electrolyte reached...
213.1 °C and simultaneously with OCV falling (see Fig. 2b), which is 12.6 °C higher than that of the case with the concentrated electrolyte. $T_2$ and OCV displayed the repeatable characters (Supplementary Fig. 2), and OCV did not fall until ~213.1/214.8 °C, suggesting that the separators in the cells had the potential to stand 213.1 °C/214.8 °C or even higher temperature without ISC. Additionally, the heat generated by ISC was evaluated based on the internal resistance of battery around $T_2$.

ISC just contributed a $(dT/dt)_{ISC}$ of 0.06 °C s$^{-1}$, much lower than 1 °C s$^{-1}$ at $T_2$ (See details in Supplementary Note 3). Thus, it concludes that, for Gr|NMC811 batteries with the concentrated electrolyte, the exothermic process that leads to $T_2$ is caused by internal reactions and not by ISC. A large amount of heat was released in the battery. As a consequence, the loss of the integrity of the separator or the battery swell accompanied with the vigorous exothermic reactions, would lead to a sharp drop in the voltage$^{1, 12}$. After $T_2$, the battery temperature sharply increased to the maximum temperature ($T_3 = 652.2$ °C) in 15.4 s. The maximum $dT/dt$ during the thermal runaway was 401.2 °C s$^{-1}$. Meanwhile, the violent flame was observed in the lateral heating test (the inset in Fig. 2a), and the overall process is shown in Supplementary Movie 1, indicating that the battery with the concentrated LiFSI/DMC electrolyte was combustible during the thermal runaway even though the electrolyte demonstrated low flammability.

The thermal features of the Gr|NMC532 battery with the concentrated LiFSI/DMC electrolyte were also examined (see Fig. 2c). The chemical reaction was also proved to be the trigger of the separator or the battery swell accompanied with the vigorous exothermic reactions, would lead to a sharp drop in the voltage$^{1, 12}$. After $T_2$, the battery temperature sharply increased to the maximum temperature ($T_3 = 652.2$ °C) in 15.4 s. The maximum $dT/dt$ during the thermal runaway was 401.2 °C s$^{-1}$. Meanwhile, the violent flame was observed in the lateral heating test (the inset in Fig. 2a), and the overall process is shown in Supplementary Movie 1, indicating that the battery with the concentrated LiFSI/DMC electrolyte was combustible during the thermal runaway even though the electrolyte demonstrated low flammability.

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for thermal runaway (see details in Supplementary Note 3). \( T_2 \) was found to be 202.3 °C, which is close to that of the Gr|NMC811 battery with the concentrated LiFSI/DMC electrolyte. However, the Gr|NMC532 battery with the conventional electrolyte still showed \( T_2 \) at a higher temperature (241.1 °C, see Fig. 2d). It was reported that the crosstalk between the NMC532 cathode and anode was when the reactions happened at \( T_2 \). At or after \( T_2 \), the most exothermic reactions, where the cathode acted as the main reactant, were initiated. So, it was understandable that Gr|NMC811 batteries always exhibited a lower \( T_2 \) than Gr|NMC532 batteries when with the same electrolyte.

It was interesting that the value of \( T_2 \) in the Gr|NMC811 battery was very close to that of \( T_2 \) in the Gr|NMC532 battery when the same concentrated LiFSI/DMC electrolyte was used in both of them, and both values were lower than that of the \( T_2 \) in the battery with the conventional electrolyte. These phenomena indicated that the concentrated LiFSI/DMC electrolyte could not enhance the intrinsic battery safety, even though the concentrated electrolyte was more thermally stable than the conventional electrolyte. Then, the quite similar \( T_2 \) of the NMC811 and NMC532 batteries with the LiFSI/DMC concentrated electrolyte indicate that similar chemical reactions occurred at ~200 °C and that these reactions brought both batteries to the point of thermal runaway. To probe the trigger reactions in the batteries, partial cells were employed to simulate all the possible exothermic reactions in the battery.

**Contribution of the exothermic reactions to thermal runaway.**
To detect the exothermic reactions of the concentrated LiFSI/DMC electrolyte in Gr|NMC811 battery, a comparison of the temperature dependence of \( \Delta T/\Delta t \) between the full cell and the partial cells is shown in Fig. 3. In contrast to the AnEly and CaAn partial cells, the CaEly partial cell did not enter thermal runaway. No sharp temperature rise was observed from the ARC curve, and its \( T_2 \) was 290 °C. Meanwhile, the maximum \( \Delta T/\Delta t \) was even lower than 0.1 °C s\(^{-1}\), which is far lower than 1 °C s\(^{-1}\). These results illustrate that the heat generated by the reactions inside the CaEly partial cell, including the cathode material decomposition and electrolyte oxidation by the cathode material, was relatively small before 290 °C and incapable of triggering the battery thermal runaway. This result coincides with the low flammability of the concentrated electrolyte, since the combustion reflected the intensity when the electrolyte was oxidized by the oxygen in the air, and the ARC result of the CaEly partial cell reflected the maximum intensity when the electrolyte was oxidized by the charged cathode or the oxygen released by the charged cathode. The AnEly and CaAn partial cells, on the contrary, could be brought to thermal runaway (see Fig. 3). \( T_2 \) reached 200.5 °C, 202.5 °C, and 225.1 °C in the case of the full cell, AnEly, and CaAn partial cells, respectively. In addition, the maximum (\( \Delta T/\Delta t \))\(_{\text{max}} \) for the full cell, AnEly, and CaAn partial cells was 401.2 °C s\(^{-1}\), 882.9 °C s\(^{-1}\), and 164.0 °C s\(^{-1}\), respectively. Thus, it can be roughly concluded that both the AnEly and CaAn partial cells may contribute to thermal runaway in the full cell, but further analysis is needed to exactly find out which of them provided the trigger reaction and which of them provided the main reaction during thermal runaway.

To analyze the trigger reactions, the \( \Delta T/\Delta t \) values of all the cells at 200.5 °C (\( T_2 \) of the full cell) were compared. It was observed that the \( \Delta T/\Delta t \) of the CaAn partial cell was 0.1 °C s\(^{-1}\) (see the triangle in Fig. 3) when the \( \Delta T/\Delta t \) of the full cell reached 1 °C s\(^{-1}\) (see the circle in Fig. 3), meaning that the heat released by the CaAn partial cell at this temperature was no more than 1/10 of the total heat of the full cell. In fact, the \( \Delta T/\Delta t \) of the CaAn partial cell was always ~1/10 of that of the full cell. Since the decomposition of the cathode material released little heat, the reactions inside the CaAn partial cell contributed little to \( T_2 \) and the heat accumulation before \( T_2 \) for the full cell. As a result, the CaAn partial cell cannot provide the trigger reaction. In contrast, the \( \Delta T/\Delta t \) of the AnEly partial cell was close to 1 °C s\(^{-1}\) at \( T_2 \), and the \( \Delta T/\Delta t \) curves of the AnEly and full cell almost overlapped near \( T_2 \). This demonstrates that the heat release dynamics of the AnEly and full cell were almost the same. Thus, the chemical reactions between the anode-electrolyte were exactly the same reactions that took place in the full cell. Since the battery was uniformly heated and the cell was so small that its thermal conductivity was quite good, the reactions would uniformly take place inside the cell. In addition, the exponential growth of the \( \Delta T/\Delta t \) with the temperature would be attributed to exothermic chemical reactions. That is, multiple reactions were involved between the anode and the electrolyte, and they were also initiated one after the other with a rise in temperature. In turn, the released heat led to a continuous increase in the temperature of the full cell. When the cell temperature became close to \( T_2 \), a group of vigorous reactions was initiated, enabling the full cell temperature surge at \( T_2 \). This chain reaction is very complex, and its analysis would be a great chance for future research. Overall, the rise in the temperature curves of the AnEly and full cells was very similar below \( T_2 \), which proves that the AnEly cell was responsible for the trigger reaction that led to the battery’s thermal runaway.
The surge after \( T_2 \) could be obviously observed in the full, AnEly, and CaAn cells. Then, the \( dT/dt \) gradually decreased until the cell temperature reached the maximum value. Although the maximum temperature (\( T_3 \)) of the three kinds of cells is different, the maximum \( dT/dt \) of the AnEly and CaAn cells was both brought to hundreds of orders of magnitude, which indicates that the reactions in both the AnEly and CaAn cells were mainly responsible for the exothermic reactions during thermal runaway. Also, \( T_3 \) of each of the AnEly and CaAn cells was higher than that of the full cell because some of the battery components did not exist in the AnEly and CaAn partial cells.

\( Q_{TR} \) was used to denote the intensive heat release during thermal runaway, and it can be calculated from Eq. (1), where \( M \) denotes the mass of the cell (g), and \( C_p \) signifies the specific heat capacity (J·g\(^{-1}\)·K\(^{-1}\); Supplementary Table 2 and Supplementary Note 4). The temperature range for the heat calculation is \( \Delta T (\Delta T = T_3 - T_2) \). Table 1 shows the characteristic temperatures and \( Q_{TR} \) of the cells, and the equivalent rise in temperature of full battery (\( \Delta T_{eq} \)) which was caused by the reactions in the partial cells. \( \Delta T_{eq} \) was calculated from Eq. (2). The AnEly partial cell released a \( Q_{TR} \) of 7.0 kJ with a \( \Delta T_{eq} \) of 312.6 °C, whereas the full cell released a \( Q_{TR} \) of 11.5 kJ with a \( \Delta T \) of 516.9 °C. Meanwhile, the CaAn partial cell generated a \( Q_{TR} \) of 11.3 kJ with a \( \Delta T_{eq} \) of 507.9 °C, implying that the redox reactions between the anode–cathode can also generate tremendous heat from \( T_2 \) to \( T_3 \). The AnEly partial cell, among the three partial cells, was first brought to a thermal runaway. If the AnEly partial cell contributed all the heat to the full cell, the reaction in the CaAn cell would at least provide heat of 4.5 kJ to the full cell. It indicates that the reactions in both the AnEly and CaAn cells were the main reactions during the thermal runaway. Besides, even if the cathode was completely inert under all the temperatures, the heat released by the reactions between the anode and the electrolyte can bring the full battery to thermal runaway state (see details in Supplementary Table 3 and Supplementary Note 5). Similarly, the full cell can be brought to thermal runaway but at a higher temperature if the electrolyte stays totally inert.

\[
Q_{TR} = \Delta T \sum (M \cdot C_p) \tag{1}
\]

\[
\Delta T_{eq} = \frac{Q_{TR}}{\sum (M \cdot C_p)} \tag{2}
\]

Based on the above analysis, three conclusions can be drawn for the Gr[NMC811] thermal runaway. First, the reactions between the cathode–anode contributed little to \( T_2 \) and the heat accumulation before \( T_2 \), which coincides with the low flammability of the concentrated electrolyte. Second, the reactions in the AnEly partial cell were responsible for the heat accumulation below \( T_2 \), as well as the trigger reaction of the thermal runaway. Third, the reactions in both the AnEly and CaAn partial cells were the main reactions during the thermal runaway. In the following section, the thermal stabilities of the individual materials and their mixtures are estimated to further probe the trigger reactions and main exothermic reactions during thermal runaway.

**Table 1** Thermal runaway features of the partial cells and full cell.

| Cells | Mass (g) | \( T_1 \) (°C) | \( T_2 \) (°C) | \( T_3 \) (°C) | \( \Delta T \) (°C) | \( (dT/dt)_{max} \) (°C S\(^{-1}\)) | \( Q_{TR} \) (kJ) | \( \Delta T_{eq} \) (°C) |
|-------|---------|----------------|----------------|---------------|----------------|-----------------|----------------|-----------------|
| CaEly | 13.9    | 220.3          | -              | 288.3         | 68             | -               | 0.8            | 37.4            |
| AnEly | 13.9    | 130.1          | 202.5          | 712.2         | 582.1          | 882.9           | 7.0            | 312.6           |
| CaAn  | 17.5    | 140.7          | 225.1          | 724           | 583.3          | 164             | 11.3           | 507.9           |
| Full cell | 24.5 | 135.3          | 200.5          | 652.2         | 516.9          | 401.2           | 11.5           | -               |

**Thermal stability of LiFSI/DMC in Gr[NMC811] battery.** A DSC-TG-MS test was used to characterize the thermal stability of the cell components. By enumerating all the thermal reactions of the individual and mixed cell components, the reactions inside the battery during the thermal runaway evolution can be screened out. As the chemical reactions in the AnEly partial cell were targeted as the trigger reaction of thermal runaway. Then, first, all the possible reactions among the lithiated anode, concentrated LiFSI/DMC electrolyte, electrolyte components, and delithiated cathode were measured (Fig. 4 and Supplementary Table 4).

Both the lithiated anode (An) and the An+DMC mixture exhibited a broad and mild exothermic peak at ~289 °C (the dotted box and the inset in Fig. 4a), where the corresponding \( \Delta H \) for both of them was ~70 J g\(^{-1}\). This exothermic peak can be attributed to the reaction between the lithiated graphite and the polyvinylidene fluoride-based binder\(^{12}\), which coincides with what was indicated by the plat TGA curve (Fig. 4d). In detail, the weight was kept constant in the temperature range between the room temperature and 550 °C, which is consistent with the fact that no gases or volatilizable liquids were produced in the reactions between the lithiated graphite and the binder. The addition of DMC did not change it, indicating that no reaction between the An and DMC occurred. For LiFSI, the endothermic peak at ~145 °C can be attributed to the melting of LiFSI, as no weight loss at this temperature occurred (Fig. 4d). As for the concentrated electrolyte, the peak at ~145 °C disappears (LiFSI/DMC and An+LiFSI/DMC curves in Fig. 4a). The exothermic peak at ~350 °C might be associated with the thermal decomposition of FSI\(^{-36,37}\), as 65% of the weight loss can be determined according to Fig. 4d, and the NO\(_2\) (\( m/z = 46 \)) and SO\(_2\) (\( m/z = 64 \)) gases were released at ~350 °C because of the S–F and S–N breakage in the FSI\(^{-36,37}\). The concentrated electrolyte and LiFSI showed quite a similar exothermic peak at ~350 °C, which indicates the high thermal stability between LiFSI and DMC. Besides, the change from the crystal to the solution state did not change the decomposition behavior of LiFSI.

However, the addition of the lithiated anode results in obvious changes in the thermal behavior of LiFSI, which signifies the reaction between them. As seen in Fig. 4a, for the mixture of LiFSI and An when the lithiated anode was in contact with the concentrated LiFSI/DMC electrolyte, the sample showed a sharp exothermic peak (602.9 J g\(^{-1}\)) at 209.6 °C, which was eight times greater than that of the anode alone, while no exothermic peaks took place at 350 °C. Similarly, for the An+LiFSI, NO\(_2\) and SO\(_2\) gases were released at 210.9 °C, and they were accompanied by an intensive \( \Delta H \) of 757.9 J g\(^{-1}\). In addition, the TGA curve (Fig. 4d) showed that a weight loss of almost 20% took place at ~210 °C, while the weight loss at ~350 °C was <3%, indicating that most of the LiFSI powders reacted with the lithiated anode and produced gas or volatile products. It is reasonable that the DSC curve of the An+LiFSI/DMC sample did not show a LiFSI decomposition, as the heat released by the 3% residue of the LiFSI was too little. Furthermore, for the fresh graphite (Li\(_0\)C\(_6\)), Li\(_0\)C\(_6\) + LiFSI
displayed an endothermic peak at 145 °C without the intensive exothermic behavior at ~210 °C, demonstrating the considerable heat produced in LiC₆ + LiFSI from the chemical reaction between intercalated lithium and LiFSI. It was also reported in ref. 38 that the battery employing the LiFSI-based electrolyte showed an exothermic peak of 1300 J g⁻¹ at ~200 °C, attributed to the chemical reduction of the FSI⁻ anion by the lithiated anode. For comparison, the thermal behaviors of LiPF₆ and An were also investigated, it demonstrated that LiPF₆ was not involved in the trigger or main reactions of thermal runaway, coincided with the different Tₛ values of the concentrated and conventional electrolytes (see details in Supplementary Fig. 3 and Supplementary Note 6). The exothermal peak at 350 °C can be attributed to the phase transition. The thermal behavior of Ca+LiFSI/DMC implies that the NMC811 cathode barely reacts with the concentrated LiFSI/DMC electrolyte before 320 °C (see details in Supplementary Note 6). The exothermal peak at 350 °C and 380 °C in Ca+LiFSI/DMC may not be triggered in CaEly partial cell as ARC system will enter into a cooling mode if the cell does not go to thermal runaway at a pre-set temperature of 290 °C. Thus, the exothermic reaction after 320 °C contributed less to the thermal runaway which coincides with the thermal runaway behavior of the CaEly partial cell. Additionally, Ca+Separator was also investigated and with a small heat generation (Supplementary Fig. 4 and Supplementary Note 8). The Ca+An sample exhibited two major exothermic peaks that were centered at 239.5 °C and 279.4 °C, respectively. The ΔH values of the two exothermic reactions were calculated, and they were found to be 834.0 J g⁻¹ in total, where it was believed that they emanated from the consumption of the cathode-produced oxygen by the anode. The reaction between the cathode and the anode also significantly contributed to the heat during thermal runaway, but it was not the trigger reaction, which coincides with the Qₜₚ analysis and thermal runaway behavior of the CaAn partial cell.

Two conclusions can be drawn from Fig. 4. First, the LiFSI can be reduced by the charged anode with gas evolution and the released intensive heat at ~210 °C, which was the trigger reaction that brought the battery to the point of thermal runaway. Second, the reaction between the fully charged cathode and the anode generated great heat, which significantly contributed to the released heat during thermal runaway, but it was not the trigger reaction.

Thermal stability of LiFSI/DMC in Gr|NMC532 battery. It is widely accepted that charged cathodes take part in the thermal runaway process and that the NMC532 battery is more thermally and chemically stable than the NMC811 battery when the other battery materials are the same. According to research on the NMC811 battery with the LiFSI/DMC concentrated electrolyte, the charged cathode only participated in the thermal runaway, while the trigger reaction was led by the anode and the concentrated electrolyte. Also, the Gr|NMC532 battery was further used to confirm thermal runaway evolution when employing concentrated LiFSI/DMC electrolytes (Fig. 5). Similar to the case of the Gr|NMC811 battery, the An and Ca individual samples resulted in a very weak exothermic peak, with ΔH values of 68.2 and 46.8 J g⁻¹, respectively (the insert in Fig. 5a and Supplementary Table 5), indicating that without a strong oxidizer or reductor, An and Ca cannot cause great damage by thermal decomposition. As for their mixtures, two peaks appeared at 272.1 °C and 394.3 °C with a total ΔH of 709.3 J g⁻¹. This intensive heat was considered to bring the Gr|NMC532 battery with the conventional electrolyte to thermal runaway. However, for the concentrated electrolyte, the LiFSI exhibited an energetic peak of ~350 °C (Fig. 5a). In contrast, the An+LiFSI mixture only showed one exothermic peak at 210.5 °C with a ΔH of 767.8 J g⁻¹ and with a simultaneous release of NO₂ and SO₂ gases (see Fig. 5a–c). This change indicates that reactions took place between An and LiFSI. Similar to the results in the Gr|NMC811 battery, where the An induced the damage of the S–F and S–N bonds.
their mixture during the DSC measurement. However, the TMP could not prevent the exothermic reactions of lower than that of the LiFSI/DMC concentrated electrolyte.

Non-flammable LiFSI/TMP in Gr|NMC811 battery. The release of DMC below \( T_2 \) was an interference factor when analyzing the combustion of the Gr|NMC battery with the LiFSI/DMC concentrated electrolyte, as the small amount of free DMC is flammable though the flammability is greatly reduced. Then, the non-flammable concentrated LiFSI/TMP electrolyte was also examined in the Gr|NMC811 battery to further verify the vigorous reactions between the lithiated graphite and the LiFSI. Unfortunately, the battery was brought to thermal runaway at a temperature, the sample was transferred for the XPS analysis gated (Supplementary Fig. 6 and Supplementary Note 10). The XPS analysis of An+LiFSI (Fig. 6d), the trigger reaction was also demonstrated to be LiC\(_6\) + LiFSI.

Post-test analysis. An XPS analysis was conducted on the DSC residue to further support the thermal runaway mechanism. During the DSC measurement, the reaction of the LiFSI with the lithiated anode was ceased at 230 °C, which was at the end of the exothermic peak (Supplementary Fig. 5a). After cooling to room temperature, the sample was transferred for the XPS analysis (Supplementary Fig. 5b–e). The byproducts validated the chemical reactions between the LiFSI and LiC\(_6\) during the thermal runaway, supposing that the tremendous heat was initiated by breaking the S-F and S-N bonds with the formation of Li\(_2\)CO\(_3\), Li\(_2\)SO\(_4\), LiF, and so on (see details in Supplementary Note 9). The XPS analysis of An+LiFSI/TMP was also investigated (Supplementary Fig. 6 and Supplementary Note 10).
NMC batteries to thermal runaway. The mechanism of LiFSI-based concentrated electrolytes in Gr|NMC batteries. The considerable heat generated by the reaction of LiFSI in the Gr|NMC811 battery with the concentrated LiFSI/TMP electrolyte was demonstrated. The inset shows the combustibility of battery in the lateral heating test.

Conventional fire retardant agents could not contribute to thermal runaway for the batteries with conventional flammable electrolytes, but it is not the only contributor to the heat output during thermal runaway, where the fire retardant agent could not work. The flammability of electrolytes does contribute to thermal runaway for the batteries with conventional flammable electrolytes, but it is not the only contributable reaction and it is even not the biggest contributor to the strong cathode oxidization like the NMC811. Thus, the interactions between the charged electrodes and electrolytes should be fully considered in battery safety assessments. These findings provide valuable insights into the thermal runaway mechanisms of concentrated electrolytes, including water-in-salt aqueous electrolytes in LIBs.

Discussion

This study revealed that batteries with LiFSI-based concentrated electrolytes also undergo thermal runaway. In the conducted experiments in this study, although the fire retardant agent acted as a solvent, lithium salt acted as a strong oxidant, and the fire retardant agent could not hinder the reactions between the lithium salt and the lithiated anode. Also, the reactions between the anode and the cathode were found to significantly contribute to the heat output during thermal runaway, where the fire retardant agent could not work. The flammability of electrolytes does contribute to thermal runaway for the batteries with conventional flammable electrolytes, but it is not the only contributable reaction and it is even not the biggest contributor to the strong cathode oxidization like the NMC811. Thus, the interactions between the charged electrodes and electrolytes should be fully considered in battery safety assessments. These findings provide valuable insights into the thermal runaway mechanisms of concentrated electrolytes, including water-in-salt aqueous electrolytes in LIBs.

Methods

Materials and batteries. The solvents of EC (ethylene carbonate) and EMC (ethyl methyl carbonate) and the electrolytes of LiFSI/DMC (1:1.9 by molar), LiFSI/TMP (1:1.9 by molar), and 1 M LiPF6 in EC:EMC (3:7 by volume) were purchased from Dodochem Ltd. The moisture content of the solvents and electrolytes was <20 ppm. 0.95 Ah Gr|NMC811 and 1.2 Ah Gr|NMC532 pouch cells with a ceramic coated PE separator were used in this work. The Gr|NMC811 and Gr|NMC532 dry cells were manufactured by Guangdong Canrd New Energy Technology Co., Ltd. The LiFSI/DMC and LiFSI/TMP concentrated electrolytes were injected into the dry cells with 3.6 mL per Ah, and the formation of all the batteries was performed at C/10 under 45 °C. The cells were cycled in the voltage range of 2.85–4.2 V at 1/3 C under 25 °C. The energy densities of the Gr|NMC811 and Gr|NMC532 batteries were 191 Wh kg⁻¹ and 182 Wh kg⁻¹, respectively (see details in Supplementary Table 6 and Supplementary Note 11). All the batteries were charged to 4.2 V before the disassembly and measurements.

Battery safety evaluation. Battery safety was systematically assessed at the cell and material levels in this study as shown in Fig. 7. In detail, the thermal properties of the concentrated electrolytes were characterized by an ignition test, TGA and DSC. The safety performance of the battery was evaluated by an ARC test. The combustibility of the battery was measured by lateral heating. DSC-TG-MS was used to probe the reactions during the thermal runaway evolution, revealing the thermal runaway mechanism. Partial cells were used to simulate the reactions between different battery components in the battery environment, and an ARC test was employed to record the heat flow. The reactions between the different battery components were also investigated with DSC-TG-MS, and the results provided the reaction dynamics in the material level.

The assembly of the partial cells is shown in Supplementary Fig. 7 and Supplementary Note 12, and the detailed description is presented in ref. 11. After removing the cathode from the fully charged Gr|NMC811 battery, the cell that only
retains the lithiated anode and concentrated LiFSI/DMC was sealed and named as the AnEly partial cell. Similarly, the CaEly partial cell was prepared. As for the CaAn partial cell, the charged cathode and anode were washed with DMAC and naturally dried, and then the cathode and anode were rolled without a separator to assemble the CaAn partial cell13. All the above processing steps were performed in an argon-filled glove box in which the oxygen and water contents were controlled below 0.1 ppm. The mass of active materials in AnEly, CaEly, and CaAn were examined by a lateral heating test39. In detail, a ceramic heater, 28 mm in diameter was inserted into the internal central positions of the tested partial and full cells as shown by the red arrows (Supplementary Fig. 7a). Then, the cells were resealed with a silicone sealant that can maintain tightness and elasticity even at 343 °C. The micro-thermocouples probed the real-time temperature of the cells during the ARC measurement.

**DSC-TG-MS test protocol.** A DSC-TG-MS test was used to characterize the thermal properties and gaseous products of the cell components. The experiments were performed on NETZSCH STA449F5-QMS403D. The anode and cathode samples were acquired by disassembling a fully charged battery. The electrode was firstly washed with DMAC to remove the adsorbed electrolyte, and then the cathode and anode powders were scratched from the electrodes in an argon-filled glove box. All the samples were pressed in a aluminum crucible in the glove box and heated from 50 °C to 600 °C at a heating rate of 10 °C·min⁻¹ in a highly pure flowing argon atmosphere. The examined individual cell components or their mixtures by the DSC-TG-MS test are listed in Table 2. These solutions covered the key exothermic reactions in the battery33. In the DSC results, the heat flow was normalized by the total weight of Ca + An - LiFSI/DMC, which was calculated based on the measured weight of the test sample and the corresponding weight ratio of the test sample in Ca + An - LiFSI/DMC. The compressing proportion of Ca + An - LiFSI/DMC is the same as that of the full cell. By these means, the peak intensity implies the heating effect on the battery. The heat generations (ΔH) normalized by both the individual component weight and total weight are shown in Supplementary Tables 4 and 5.

**XPS analysis.** To probe the byproducts of the reactions between the lithiated anode and the concentrated electrolyte, the sample for XPS (X-ray photoelectron spectroscopy) analysis was prepared as follows. First, a mixture of the full charged anode and certain amounts of the electrolyte was heated to 230 °C with a heating rate of 10 °C min⁻¹. Then, the solid residues were cooled to room temperature in flowing argon, collected, and analyzed by XPS, which was collected with a K-Alpha + spectrometer using the Thermo Fisher Scientific Co. The binding energy (BE) scale was calibrated with a C 1 s peak at 284.8 eV.

**Data availability**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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### Table 2 Samples for DSC-TG-MS test.

| Sample no. | Composition | Mass (mg) | Scanning rate (°C min⁻¹) |
|------------|-------------|-----------|--------------------------|
| 1          | Cathode (Ca)| 4.0       | 10                       |
| 2          | Anode (An)  | 3.4       | 10                       |
| 3          | LiFSI/DMC   | 4.6       | 10                       |
| 4          | An-LiFSI/DMC| 3.4 + 4.6| 10                       |
| 5          | An-LiFSI    | 3.4 + 2.2| 10                       |
| 6          | An-LiFSI    | 3.4       | 10                       |
| 7          | LiFSI/DMC   | 3.4       | 10                       |
| 8          | Ca+LiFSI/DMC| 4.0 + 4.6| 10                       |
| 9          | Ca+LiFSI/DMC| 4.0 + 2.2| 10                       |
| 10         | LiFSI       | 4.0       | 10                       |
| 11         | Ca+An       | 4.0 + 3.4| 10                       |
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Author contributions
M.O. supervised this project. J.H. and X.N.F. conceived the original concept. J.H. and L.W. analyzed the data and wrote the paper. L.L., Y.N., A.O., X.M.H., X.B.H., and I.O. revised the manuscript and triggered helpful discussion. D.R. helped J.H to conduct the ARC tests and data analysis. Y.L., W.R., and Y.L.L. helped J.H. to conduct DSC-TG-MS and XPS tests. All authors contributed to discuss the results.

Competing interests
The authors declare no competing interests.

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