Exothermic behaviors of mechanically abused lithium-ion batteries with dibenzylamine

Yang Shi a, Daniel J. Noelle a, Meng Wang b, Anh V. Le b, Hyojung Yoon c, Minghao Zhang c, Ying Shirley Meng a, c, Yu Qiao a, b, *

a Program of Materials Science and Engineering, University of California — San Diego, La Jolla, CA 92093, USA
b Department of Structural Engineering, University of California — San Diego, La Jolla, CA 92093-0085, USA
c Department of NanoEngineering, University of California — San Diego, La Jolla, CA 92093-0448, USA

Highlights
- Thermal runway of lithium-ion batteries may be prevented by using DBA.
- DBA is sealed in an inert package and placed inside the battery, next to electrodes.
- Under normal condition, the DBA package does not affect the battery performance.
- As the cell is damaged, DBA is released and increases the electrolyte resistivity.

Article info
Article history:
Received 2 May 2016
Received in revised form 8 July 2016
Accepted 10 July 2016
Available online 16 July 2016

Keywords:
Lithium-ion battery
Thermal runaway
Safety
Conductivity
Electrolyte

Abstract
A thermal-runaway retardant (TRR) of lithium-ion batteries (LIBs), dibenzylamine (DBA), is investigated. In a TRR-modified LIB, DBA can be encapsulated in packages made of inert materials. When the LIB is subjected to mechanical abuse, the packages would be broken apart and the TRR is released. In nail penetration and impact tests, addition of 4 wt% DBA reduces the temperature increase of fully charged LIR-2450 cells by nearly 50%. The influence of TRR packages on the cycling performance of LIBs is negligible. The working mechanism of DBA is associated with the decrease in electrolyte conductivity, the increase in charge transfer resistance, and the reduction in lithium ion (Li\(^+\)) transference numbers.

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1. Introduction
Thermal runaway imposes significant challenges to large-scale lithium-ion batteries (LIBs). The working temperature of a LIB is usually around room temperature [1]. Upon mechanical abuse such as an impact or nail penetration, the LIB cell components may fail and internal short circuits could be formed [2]. As a result, a series of exothermic reactions would take place and the local temperature can rapidly increase by 100–200 °C in less than one minute [3].

Great efforts have been made to enhance the safety of LIBs. A common approach to suppress thermal runaway is to embed thermally triggered safety components into LIBs, such as trilayer...
separators that can close their pores when the internal temperature reaches the melting point of the middle layer, preventing further ion transport between electrodes [4,5]; low-melting-point microspheres that can cover the anode surface when they melt, preventing further ion disintercalation at the electrode-electrolyte interface [6]; positive-temperature-coefficient (PTC) additives that drastically increase the internal impedance at 110–150 °C [7–10]; microcapsules that release fire extinguishing agents at 100–130 °C [11]; and thermally polymerizable monomers that thickens electrolyte [12]. One major issue of these thermal-runaway mitigation mechanisms is that they are activated only after the local temperature rises to above 90 °C, which is dangerously close to the acceleration point of the exothermic reactions [13]. Researchers have also investigated flame retardant additives [14–18] that can reduce the electrolyte flammability, yet the battery performance is usually negatively affected [19]. Other approaches include electrode coating and doping that improve the thermal stability of the electrodes [20–26], but extensive fundamental study is still needed to enhance their efficacy and efficiency.

Mechanically triggered thermal-runaway mitigation mechanisms recently received increasing attention [27–29]. For instance, if a thermal-runaway retardant (TRR) is encapsulated in mechanically responsive packages made of materials inert to the battery environment, upon external mechanical loadings the packages can be broken apart and release the TRR. This mechanism allows for the use of aggressive chemicals to suppress the short circuit discharge and the subsequent exothermic phenomena, immediately after the battery is damaged, even before temperature increase begins. The main challenge is that the TRR must be highly efficient, such that it does not result in much increase in cell mass or volume. In this work, we investigate dibenzylamine (DBA) as a TRR candidate. The influence of TRR-containing packages on the cycling performance of LIBs is also evaluated.

2. Experimental

2.1. Nail penetration test

Xiamen Tob 140 mAh/g LIR-2450 rechargeable LIB cells, with the diameter of 24 mm and the thickness of 5 mm, were charged by constant current–constant voltage algorithm using a MTI BST8-WA Battery Analyzer to 4.3 V at C/5, with the cut-off current of C/50. The charged cells were disassembled and the electrode stacks together with the electrolyte were harvested with the assistance of an Argon-filled glovebox (H2O < 0.5 ppm). Empty type-2450 cell cases were modified, with two holes drilled in the cathode-side shell and covered with Kapton tapes. The harvested electrodes and electrolyte were re-assembled in the modified cell cases in the glovebox. The Kapton tapes were removed and two Tygon tubes were inserted into the holes and sealed with vacuum grease. Two thermocouples (Omega TT-K-40-25 type-K gauge 40) were attached to the upper and bottom surfaces of the cell case, respectively, ~7 mm away from the centers, and connected to a temperature logger (Omega OM-EL-USB-TC).

The cell was affixed to a polymethyl methacrylate (PMMA) holder and a steel nail was driven through the cell by a type-5582 Instron machine at the speed of 5 mm/s. After the nail penetrated the cell, the compression plate of the Instron machine was rapidly moved back, leaving the nail inside the cell. The nail diameter was 3.4 mm. Immediately prior to the nail penetration, 100 μL DBA (97%, Sigma Aldrich) had been injected by a syringe into the modified cell through the Tygon tubing. In the reference test, the same amount of pristine electrolyte was injected into the cell.

To better understand the efficiency of DBA, nail penetration tests were also carried out on unmodified LIR-2450 cells at various states of charge (SOC). The unmodified LIR-2450 cells at various SOC were prepared by discharging LIB cells that had been fully charged to 4.3 V to various cut-off discharge capacities of 35 mAh, 70 mAh, 105 mAh and 140 mAh, respectively.

2.2. Impact test

The LIR-2450 cells were charged to 4.3 V and the electrodes together with electrolyte were harvested through the same procedure as described in Section 2.1. Polyethylene/Aluminum/Polyester (PAP) trilayer (ULINE S-16893) was folded and heat sealed to form packages using an impulse sealer (Mcmaster-Carr), with 115 μL DBA or pristine electrolyte sealed inside. The packages were first at rest in the glovebox vacuum antechamber overnight, and then transferred into the glovebox for the subsequent cell assembly. The harvested electrodes and electrolyte were re-assembled into an empty type-2450 cell case, with a DBA- or electrolyte-containing package sandwiched between the top shell and the electrode. The cells containing electrolyte packages were used as reference samples.

Impact tests on the re-assembled LIB cells were performed in a table-top drop tower consisting of a stainless steel base and a polycarbonate track [27]. A stainless steel ball with the diameter of 6.35 mm was affixed at the center of the upper surface of the LIB cell. A polyurethane cover guided a stainless steel rod, the diameter of which was 12.7 mm and the length was 50.8 mm [28,29]. The rod was placed on top of the steel ball. A cylindrical stainless steel hammer with the mass of 7.7 kg was dropped onto the rod. The drop distance was 18 cm. A type-K thermocouple was attached to the outer surface of the anode-side shell and the temperature was recorded in the same way as described in Section 2.1.

2.3. Characterization of cycling performance

Type-2016 coin cells were assembled to evaluate the influence of DBA-containing packages on the cycling performance. Cathode films were prepared by thoroughly mixing LiNi0.5Co0.2Mn0.3O2 (TODA America), polyvinylidene fluoride (Kynar 710), and carbon black (Timcal Super C65) powders with the mass ratio of 93:4:3. Slurries were produced in 1-Methyl-2-pyrrolidinone (Sigma Aldrich), cast on aluminum foil using a doctor blade, and dried under vacuum at 80 °C for 24 h. The electrode films had an active mass loading of about 6 mg/cm². Disk cathodes were cut, compressed using a rolling mill, dried again under vacuum for 4 h, and transferred to an Argon-filled glovebox (H2O < 0.1 ppm) for cell assembly. Small PAP packages containing ~5 μL DBA were sealed, vacuum-dried overnight, and attached to the inner surfaces of type-2016 cell cases using Kapton tapes. To assemble the cells, semicircular pieces of cathodes and lithium metal disk anodes were used. The electrolyte was 1 M LiPF6 in ethylene carbonate and ethyl methyl carbonate (EC: EMC 1:1 wt%) and the separators were tri-layer PP/PE/PP membranes (Celgard 2320). The coin cells were then allowed to rest for 2 h before the electrochemical tests were performed. Galvanostatic charge-discharge was carried out using a BST8-WA Battery Analyzer. The cells were charged and discharged at C/10 (1C = 200 mA g⁻¹) for the first cycle, followed by charging and discharging at 1C in the potential range of 3–4.3 V for all the subsequent cycles. Reference cells with the same semicircular cathodes were also assembled and cycled without DBA-containing packages.

2.4. Interaction of DBA with electrodes

To understand the interaction between DBA and the electrodes, fully charged LIR 2450 cells were disassembled and the cathode and
anode films were separated. About 100 µL DBA was dropped onto the harvested cathode or anode and the local temperature was recorded using the same thermocouple as in Section 2.1.

In addition, type-2016 coin cells were assembled in a similar way as in Section 2.3, without the PAP packages. The coin cells were allowed to rest for 2 h and charged to 4.3 V at C/10. The charged cells were disassembled in the glovebox and the cathode disks were fully washed with dimethyl carbonate (DMC). The charged cathode disks were divided into two groups. The first group were soaked in 15 mL DBA for 20 min, followed by repeated rinsing with electrolyte and DMC. The washed cathode disks were re-assembled into coin cells with Li disks; the cells were discharged to 3 V at C/10 after resting for 2 h. The second group were soaked in 100 µL DBA for 20 min. The 200 MHz NMR spectrometer in CDCl3. Mass spectra of DBA before and after the interaction with charged cathodes were recorded by a Varian Mercury Plus 400 MHz NMR spectrometer in CDCl3. Mass spectra of DBA before and after the interaction with charged cathodes were obtained by a Thermo LCQdeca molecular mass spectrometer.

2.5. Interaction of DBA with electrolyte

The ionic conductivities of electrolytes containing various amounts of DBA (0–4 wt%) were measured by a conductivity meter (DDS 307A, Shanghai Yole Instrument). Type-2016 coin cells were assembled through a similar procedure as in Section 2.4; the electrolyte was either pristine or modified with DBA. The coin cells were cycled at 1C after resting for 2 h. Electrochemical impedance spectroscopy (EIS) measurements were carried out before and after cycling in the frequency range of 106 Hz to 10−2 Hz with the signal amplitude of 10 mV, using a Solartron Impedance/Gain-Phase Analyzer.

To measure the Li⁺ transference numbers of pristine and DBA-modified electrolytes, Li metal symmetric cells were assembled. Type-2032 cell cases were used due to the increase in the thickness of cell components. Two Li metal disks were separated by a PP/PE/PP membrane, with a stainless steel spacer and a spring to ensure the tight contact between the components. The electrolytes under investigation were pristine reference (1 M LiPF6 in EC: EMC 1:1 wt %) and 5 wt% DBA modified electrolyte. Potentiostatic polarization experiments were performed on the assembled coin cells with a constant voltage of 10 mV by an Arbin BT2000 battery testing system, until a steady-state current was reached. EIS measurements were carried out before and after the polarization.

3. Results and discussion

Nail penetration creates a short circuit with consistent discharge dynamics. Fig. 1a shows the experimental setup. The DBA amount is 4 wt% of the total mass of electrodes, current collectors, separator, and electrolyte. The measured temperature profiles of the two thermocouples respectively attached to the top and bottom surfaces of LIB cells are similar, with the difference less than 1–2 °C. Fig. 1b shows typical temperature profiles measured from the top surface. The addition of DBA reduces the temperature increase (ΔTmax) by about 50%, from −75 °C in reference cells to −40 °C in DBA-modified cells. The temperature ramp rates (Fig. 1c) are calculated by differential analysis of the temperature profiles. Both reference and DBA-modified cells reach their maximum ramp rates (Rmax) in about 8 s; DBA reduces Rmax by −30% from −1 °C/min to −0.7 °C/min.

A simple model is used to estimate the generated heat over time, based on the following assumptions: (1) The temperature of the cell is represented by the thermocouple measurement; (2) The convective heat transfer is considered and the radiative heat transfer is ignored [30]; and (3) the heat capacity of the added electrolyte or DBA is much smaller than that of the cell components. The heat generation rate (q′) can be calculated by:

\[
q' = (m_e C_{pe} + m_s C_{ps}) \frac{dT}{dt} + Ah(T - T_{amb})
\]

where \( m \) indicates mass and \( C_p \) indicates specific heat; subscript “e” stands for the electrode stack containing electrolyte and subscript “s” stands for the stainless steel case together with the nail; \( T \) is the temperature of the cell and \( T_{amb} \) is ambient temperature; \( A \) is the surface area exposed to air, including both the cell and the nail, and \( h \) is the convective coefficient. The values of these parameters are given in Table 1. The accumulated heat (q) is calculated by:

\[
q = \int_0^T \left[(m_e C_{pe} + m_s C_{ps}) \frac{dT}{dt} + Ah(T - T_{amb})\right] dt
\]
Charged cathode disks are then soaked up in DBA. After the cathodes are thoroughly washed to remove remaining DBA, they are reassembled into coin cells with pristine electrolyte. The specific discharge capacities of the two types of reassembled cells are 164 mAh/g and 149.2 mAh/g, respectively (Fig. 3b). For the reassembled reference cell, the initial voltage drop is due to the voltage relaxation. The voltage drop of the cell containing DBA-treated cathode is larger than that of the reference cell. In a parallel test, charged cathode materials were allowed to react with DBA in order to determine the reaction products, but no change is observed in the \(^1\)H NMR spectra (Supplementary Fig. S1) and mass spectra (Supplementary Fig. S2) of DBA before and after the reaction, suggesting that the reaction products of DBA are not in liquid phase. Moreover, no gas generation is observed during the reaction of DBA.

Table 1  
The parameters used in Eqs. (1) and (2).

<table>
<thead>
<tr>
<th>(m_e (\text{kg}))</th>
<th>(m_s (\text{kg}))</th>
<th>(C_P_e (\text{J/kg K}))</th>
<th>(C_P_s (\text{J/kg K}))</th>
<th>(A (\text{m}^2))</th>
<th>(h (\text{W/m}^2 \text{K}))</th>
<th>(T_{\text{amb}} (\text{K}))</th>
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<tr>
<td>(2.2 \times 10^{-3})</td>
<td>(7.2 \times 10^{-3})</td>
<td>1100 ([31])</td>
<td>500</td>
<td>(1.8873 \times 10^{-3})</td>
<td>7.9 ([32])</td>
<td>298</td>
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</tbody>
</table>
with the charged cathode. It is possible that a solid layer of reaction products is deposited on the cathode surface. The initial voltage drop and the capacity loss might be attributed to this additional resistive layer. Note that the DBA-treated cathode could still deliver 91% of the discharge capacity of the reference cathode, and the fact that DBA-cathode reaction is exothermic (Fig. 3a) is against the observed reduction in $\Delta T_{\text{max}}$. Therefore, the chemical reactions between the cathode and DBA should not be a major factor that influences the heat generation in nail penetration and impact tests.

The effects of DBA addition on the electrolyte are investigated. As shown in Fig. 4a, the ionic conductivity of pristine electrolyte is 9.23 mS/cm. The 5 wt% and 10 wt% DBA-modified electrolytes have reduced ionic conductivity of 7.59 mS/cm and 6.38 mS/cm, respectively. Coin cells assembled with pristine electrolyte as well as 5 wt% or 10 wt% DBA containing electrolyte are cycled at 1C and the cell potentials dynamics over time are shown in Fig. 4b. The cell with 5 wt% DBA has a larger polarization and a greatly reduced capacity compared to the reference cell; the cell with 10 wt% DBA has the lowest capacity and cannot be normally charged and discharged. Clearly, DBA significantly increases the cell impedance. EIS measurements are performed on the cells before and after 5 charge-discharge cycles. Fig. 4c and d shows the Nyquist plots,
equivalent circuits, and fitted curves, with \( R_s \) being the electrolyte resistance, \( R_{sei} \) the solid electrolyte interface (SEI) resistance, and \( R_{ct} \) the charge-transfer resistance [33]. Table 2 shows the resistance values according to the fitting results of the equivalent circuit. It is evident that the DBA-modified cell has not only a larger electrolyte resistance, but also much higher SEI and charge-transfer resistances.

The effect of DBA addition on the \( \text{Li}^+ \) transport property is further investigated through the measurement of \( \text{Li}^+ \) transference numbers by a potentiostatic polarization method [34,35], in which a constant voltage of 10 mV (\( \Delta V \)) is applied on a \( \text{Li} \)-metal symmetric cell. During the polarization, the current drops from the initial value (\( I_0 \)) to a steady-state value (\( I_{ss} \)). Fig. 5a and b display the current drop with time, and the insets show the Nyquist plots from EIS measurements before and after polarization. In the corresponding equivalent circuit, \( R_1 \) and \( R_2 \) are the \( \text{Li} \)-electrolyte interfacial resistances, and \( R_e \) is the electrolyte resistance [35,36]. The \( \text{Li}^+ \) transference numbers \( t_{\text{Li}^+} \) is given by Ref. [37]:

\[
t_{\text{Li}^+} = \frac{I_{ss}}{I_0} \left( \frac{\Delta V - I_0 R_0}{\Delta V - I_{ss} R_{ss}} \right)
\]

where \( R_0 \) and \( R_s \) equal to the summations of \( R_1 \) and \( R_2 \) before and after polarization, respectively. Table 3 shows the values of \( I_0 \) and \( I_{ss} \) during polarization, the resistance values from the fitting results, and the calculated \( \text{Li}^+ \) transference numbers of reference and DBA-modified electrolyte. The \( \text{Li}^+ \) transference numbers of reference electrolyte is 0.48, which is consistent with the data in open literature [38]. The addition of 5 wt% DBA decreases the \( \text{Li}^+ \) transference number to 0.23.

The testing data show that the addition of DBA increases the electrolyte resistance and the charge transfer resistance, and decreases the \( \text{Li}^+ \) transference number, all of which would suppress \( \text{Li}^+ \) transportation, suggesting that the modification of electrolyte is a major reason of the hindered heat generation and the reduced \( DT_{\text{max}} \) in nail penetration and impact tests. The increased resistances and reduced \( \text{Li}^+ \) transference number may result from the Lewis base characteristic of DBA. Previous studies reported that adding Lewis acid to an electrolyte could improve the conductivity by forming complex with salt anions [39]. The Lewis acid serves as an anion receptor, enhances the dissociation of ion pairs, and increases the population of free cations [40]. The dissociation of ion pairs contributes to a high \( \text{Li}^+ \) transference number. DBA is a Lewis base and an electron donor, which may repel the \( \text{PF}_6^- \) anions and decrease the dissociation of \( \text{Li}^+ \) solvation shell and \( \text{PF}_6^- \) pairs, resulting in the decrease in \( \text{Li}^+ \) transference number and conductivity.

The current study on coin cells demonstrate the basic concept of including DBA in battery cells. It is envisioned that the conditions in large-sized pouch cells are different from coin cells, in terms of cell size, mass, electrolyte-electrode mass ratio, among others. Using TRR for thermal-runaway mitigation of pouch cells will be an important topic of our future study.

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**Table 2**

<table>
<thead>
<tr>
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<th>After assembly</th>
<th>After 5 cycles</th>
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<tr>
<td>( R_s ) (( \Omega ))</td>
<td>50</td>
<td>75</td>
</tr>
<tr>
<td>( R_{sei} ) (( \Omega ))</td>
<td>250</td>
<td>1050</td>
</tr>
<tr>
<td>( R_{ct} ) (( \Omega ))</td>
<td>58</td>
<td>65</td>
</tr>
<tr>
<td>( R_{sei} ) (( \Omega ))</td>
<td>254</td>
<td>1040</td>
</tr>
<tr>
<td>( R_{ct} ) (( \Omega ))</td>
<td>450</td>
<td>1700</td>
</tr>
</tbody>
</table>

**Fig. 4.** (a) Ionic conductivity of DBA-modified electrolyte. (b) Typical voltage profiles of the reference and DBA-modified cells. Nyquist plots, equivalent circuits, and fitting plots: (c) immediately after reassembly and (d) after 5 charge-discharge cycles.
4. Conclusions

Dibenzylamine (DBA) is identified as a promising thermal runaway retardant (TRR) of lithium-ion batteries (LIBs). Upon nail penetration or impact, the temperature increase of a fully charged LIR-2450 LIB cell with 4 wt% DBA could be reduced by ~50%. Embedding DBA-containing packages in coin cells does not affect the cell capacity and the cycling performance up to 100 cycles. Charged anode does not react with DBA. Exothermic reactions between DBA and charged cathode are observed, but it is not a governing factor of the reduced heat generation of damaged cells. The major working mechanism of DBA is associated with the increase in electrolyte and charge transfer resistances as well as the decrease in Li⁺ transference number.

Acknowledgement

This research is supported by Advanced Research Projects Agency – Energy (ARPA-E) under Grant No. DE-AR0000396, for which we are grateful to Dr. Ping Liu, Dr. John Lemmon, Dr. Grigorii Soloveichik, Dr. Chris Atkinson, and Dr. Dawson Cagle. Special thanks are also due to Dr. Jiang Fan and Dr. Dengguo Wu at the American Lithium Energy Corporation for the help with lithium-ion battery design and processing.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.jpowsour.2016.07.034.

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Table 3

<table>
<thead>
<tr>
<th></th>
<th>(I_0)</th>
<th>(I_a)</th>
<th>(R_{00})</th>
<th>(R_{ss})</th>
<th>(R_{0})</th>
<th>(R_{a})</th>
<th>(I_{ss})</th>
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<tr>
<td>Reference cell</td>
<td>151.102</td>
<td>74.454</td>
<td>0.271</td>
<td>0.548</td>
<td>12.345</td>
<td>22.032</td>
<td>0.48</td>
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<tr>
<td>DBA-modified cell</td>
<td>183.487</td>
<td>43.687</td>
<td>0.402</td>
<td>0.715</td>
<td>8.304</td>
<td>28.218</td>
<td>0.23</td>
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</table>

Fig. 5. Polarization curves of Li metal symmetric cells with (a) pristine electrolyte and (b) electrolyte containing 5 wt% DBA; the insets show the Nyquist plots before and after polarization.
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