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Oxidative stability and reaction mechanism of lithium bis(oxalate)borate as cathode film-forming additive for lithium ion battery

Yating Wang\textsuperscript{1}, Lidan Xing\textsuperscript{1,*}, Xianwen Tang\textsuperscript{2}, Xiangfeng Li\textsuperscript{2}, Weishan Li\textsuperscript{1,*}, Bin Li\textsuperscript{1}, Wenna Huang\textsuperscript{1}, Hebing Zhou\textsuperscript{1}, Xiaoping Li\textsuperscript{1}

1. School of Chemistry and Environment, Key Laboratory of Electrochemical Technology on Energy Storage and Power Generation of Guangdong Higher Education Institutes, Engineering Research Center of Materials and Technology for Electrochemical Energy Storage (Ministry of Education), South China Normal University, Guangzhou 510006, China
2. Guangzhou Institute of Energy of Testing, Guangzhou 510170, China
Abstract:

The oxidative decomposition mechanism of lithium bis(oxalate)borate (LiBOB) as a cathode film-forming additive has been investigated using the density functional theory calculation at the B3LYP/6-311++G(d) level, with polarized continuum model. The calculated oxidation potentials of the investigated structures decreased as the following order: carbonate (including isolate EC, PC and DMC) ≈ carbonate-BOB’ clusters. Charge distribution results show that the electron of the oxidized carbonate-BOB’ cluster was taken from BOB’, indicating the higher oxidation activity of BOB’. Decomposition mechanism analyses of EC-BOB’-e cluster indicate that breakage of the BOB’ structure is energetically favorable than EC. The most likely reaction path of this cluster is the ring opening reaction of BOB’ via two transition states, generating CO, CO₂ and radical R1, which may further terminates generating borate-containing oligomer. This oligomer is believed to play a crucial role in suppressing further oxidative decomposition of carbonate solvents.

Key words: lithium ion batteries; lithium bis(oxalate)borate; oxidative decomposition; film-forming; density functional theory

1. Introduction

Lithium ion battery has been widely applied in power portable electronic devices due to its high energy density and high power density. Developing affordable battery offering high reliability, low price and high capacity is becoming the biggest
challenge to expanding the application range of lithium ion battery in electric vehicles [1,2,3,4]. One of the most effective ways to improve the cyclic stability of the battery is to improve the stability of the electrode/electrolyte interface, which can be achieved by forming a protective solid electrolyte interface (SEI) film on the electrode surface [5,6,7,8].

Traditional electrolyte consists of carbonates and lithium hexafluorophosphate (LiPF₆). However, LiPF₆ has the defects of poor thermal stability generating HF and LiF during the charging-discharging process, which corrode the current collect and increase the interfacial impedance [9,10,11]. Recently, lithium bis(oxalate)borate (LiBOB) aroused people’s attention due to its excellent thermal and electrochemical stability[12,13,14,15]. Furthermore, LiBOB has the ability to improve the cyclic performance of the cell by forming an effective protective film on both anode and cathode surface [16,17,18,19]. Specifically, using LiBOB as lithium salt, not only prevents the decomposition of propylene carbonate (PC) on the graphite electrode, but also increases the rate capability of the cell [9]. The reductive decomposition mechanism of LiBOB has been well understood [20]. Importantly, oxidative decomposition of LiBOB also forms an protective film on the cathode surface, improving the cyclic stability of the cell at high voltage [21,22,23,24,25,26]. Ha et al. found that adding 1 wt.% LiBOB into the ethylene carbonate (EC)-based electrolyte generates a stable cathode SEI film, improving the electrochemical performance of Li/LiNi₀.₅Mn₁.₅O₄ cell [26]. There is one small fly in the ointment though, CO and CO₂ gases release during the decomposition reaction of BOB’, increasing the internal
pressure of the cells [27].

Up to date, investigations have been focused on the effects of LiBOB on cell performance, while it’s oxidative stability, film-forming and gas releasing mechanisms are still unclear. The oxidative decomposition of electrolyte on the cathode surface is closely related to the safety issues of lithium ion battery. In our previous works, we have investigated the oxidative stability and decomposition mechanisms of PC and EC by using density functional theory (DFT), and demonstrated that the presence of lithium salt anions (PF$_6^-$, ClO$_4^-$, BF$_4^-$ and B(CN)$_4^-$) and neighboring solvents significantly lower the oxidative stability and alter the reaction mechanism of PC and EC [28,29,30,31,32,33,34,35]. In this work, DFT calculations were carried out to understand the oxidative stability, film forming and gas releasing reaction mechanisms of LiBOB.

2. Computational details

All calculations were performed using the Gaussian 09 package [36]. The geometry of the investigated structures was optimized using the B3LYP level in conjunction with 6-311++G (d) basis set [37]. Frequency analyses were done with the same basis set to confirm the obtained optimized stationary point. The Gibbs free energy was obtained at 298 K. Vibration frequency and intrinsic reaction coordinate (IRC) analyses were employed to confirm all the transition states of the reaction pathways at the same level. The atomic charge distributions were computed from natural population analysis (NPA) by using natural bond orbital (NBO) theory. To
investigate the role of the electrolyte environment, the bulk solvent effect was estimated using the polarized continuum models (PCM). The acetone dielectric constant (20.5) was used to represent the solvent for PCM calculation. The oxidation potential ($E_{ox}$) was converted from the absolute oxidation potential of Li$^+$/Li by subtracting 1.4 V from the former, as shown in eq. (1).

$$E_{ox} (\text{Li}^+/\text{Li}) = \frac{G(M^+)-G(M)}{F} - 1.4 \text{ V} \quad (1)$$

where $G(M)$ and $G(M^+)$ are the free energy of the solvated complex M and its solvated oxidized form $M^+$ at 298.15 K, respectively, and F is the Faraday constant[38].

3. Results and Discussion

3.1 Oxidative stability and charge distributions

According to our previous molecular dynamic simulation [39], Li$^+$ cations are excluded out of the surface layer electrolyte, while a large amount of solvents and salt anions exist right on the positively charged cathode surface. Hence, oxidative reactions of the isolate solvent, anion, and clusters generating from the solvent and anion were considered in this work. In the LiBOB-containing LiPF$_6$/carbonate electrolyte, the carbonate solvent may interact with BOB$^-$ and PF$_6^-$, generating carbonate-BOB$^-$ and carbonate-PF$_6^-$ clusters, respectively. The optimized structures and calculated oxidation potentials of the investigated isolate carbonates (including EC, PC and dimethyl carbonate (DMC)), carbonate-BOB$^-$ and carbonate-PF$_6^-$ clusters are presented in Figure 1. The sums of NPA charge distributions of the clusters before
and after oxidation are also given in this figure. It can be seen that the oxidation potentials of the investigated carbonates and clusters decrease as the following order: isolate carbonate > carbonate-PF₆⁻ > carbonate-BOB⁻, indicating the lowest oxidation stability of the carbonate-BOB⁻ cluster in the investigated electrolytes. This is in agreement with the linear sweep voltammetric results of Ha et al. [26]. As shown in Figure 1, differently with PF₆⁻, there is no H transfer after the oxidation of carbonate-BOB⁻ cluster. The atomic charge distributions of BOB⁻ of isolate BOB⁻ and carbonate-BOB⁻ clusters before and after oxidation are summarized in Table 1. It can be seen that the electron is mainly taken from one of the five-ring of BOB⁻ after the oxidation reaction of carbonate-BOB⁻, leading to the breakage of the corresponding C-C bond (from 1.55 to 1.91 Å), while the charge on the carbonate solvent stays constant, confirming the preferential oxidation of BOB⁻. Next, we focus on the decomposition mechanism of EC-BOB⁻ cluster, since EC is the indispensable solvent in the electrolyte [40,41].

3.2 Oxidative decomposition mechanism of EC-BOB⁻ cluster

For comparison, the oxidative decomposition reaction of isolate BOB⁻ anion was also investigated in this work, as shown in Figure 2. The calculated oxidation potential of BOB⁻ is 5.97V vs. Li⁺/Li, which is almost the same with that of EC-BOB⁻ cluster, and 1.14 V lower than that of isolate EC. Similarly with the oxidative reaction of EC-BOB⁻ cluster, a breakage of C-C bond can be observed after oxidation of the isolate BOB⁻, as shown in Figure 2. The O-B bond (in the breakage ring of BOB⁻)
increases from 1.48 to 1.52 Å. Hence, a subsequent breakage of this bond was investigated. It can be found that the oxidized BOB requires only a small energy barrier of 15.75 kJ/mol (TS1) to release CO₂, which means that gas releasing reaction of the oxidized BOB is easy to take place.

A further decomposition of M1 via C-O bond of another five-ring (TS2, $\Delta G = 108.23$ kJ/mol) was carried out next. With the breakage of the C-O bond, the five-ring structure was completely destructed generating CO, CO₂ and radical R1, as shown in M2. Gas products of both CO and CO₂ have been detected during the oxidative decomposition of LiBOB-containing electrolyte [26,42]. According to the reaction energy, it is obvious that generating CO₂ is easier than CO.

For EC-BOB-e cluster, there are 7 possible decomposition pathways, as shown in Figure 3, while the corresponding reaction energy was summarized in Figure 4. Decomposition reactions starting from BOB and EC were represented as Path 1 and Path 2 (see Figure 3), respectively. It can be easily found that, except there is a neighboring EC in the cluster, the decompositions of Paths 1 to 3 are similar to those of the isolate BOB (see Figure 2). The existence of the EC solvent does not alter the reaction energy and decomposition products of BOB, as shown in Figure 4. The reaction energy profile of BOB is almost overlapping with that of Path 1 to 3. Decompositions of Path 4 and Path 5 are corresponding to the ring-opening reactions of EC, with the activation energy of 228.4 (TS3-M1) and 231.2 (TS4-M1) kJ/mol, respectively, which are significantly higher than TS1 and slightly higher than TS2. This indicates that further decomposition of the residue structure of BOB in M1 is
kinetically favorable to EC.

As mentioned above, initial decomposition of EC (breakage of C-O bond) in the EC-BOB\(^{-}\)-e cluster is defined as Path 2. The decomposition reaction energy via TS5 is 225.4 kJ/mol, which is higher than all the decomposition reactions of BOB\(^{-}\) (TS1=19.3kJ/mol, TS2-M1=164.7kJ/mol, TS6-M5=13.92kJ/mol and TS7-M5=207.2 kJ/mol), indicating that the decomposition of EC in EC-BOB\(^{-}\)-e cluster is unfavorable. By comparing the decomposition reactions of EC in Path 2 (TS5), Path4 (TS3-M1=228.4kJ/mol) and Path 5 (TS4-M1 = 231.2kJ/mol), we can find that the participation reaction of BOB\(^{-}\) (Path 4) has little influence on the decomposition reaction energy of EC.

According to the above discussion, Path 1 to 3 is the most possible while Path 2 to 7 is the least possible decomposition reaction. Additionally, the relative free energy of M3 is 167.4 kJ/mol, which is energetically unfavorable. Hence, the decomposition products of Path 7 (M6) and Path 4 (M3) are not considered in this work.

### 3.3 Termination reactions of radicals

Initial decomposition products of EC-BOB\(^{-}\)-e cluster contain gaseous products, such as CO and CO\(_2\), aldehyde and radicals R1 and R3. The gaseous products have been found in the oxidative reaction of LiBOB-containing electrolyte [26,42,43,44,45]. Similarly to the decomposition reaction of M1 to M2 (See Figure 3), radical R3 may decompose via a C-O bond in the five-ring structure (See Figure 2, TS 2), generating R1 and gaseous products, CO and CO\(_2\). Termination reactions of R1 are
investigated, as presented in Figure 5. Two R1 radicals may terminate forming P1, while three R1 radicals generate borate-containing oligomer P3. This kind of borate-containing oligomer has been found on the cathode surface after cycling in the 1 wt.% LiBOB-containing electrolyte, by X-ray photoelectron spectroscopy measurement. And this borate-containing film is believed to play an important role in suppressing further oxidative decomposition of solvents and improving the stability of the electrode/electrolyte interface [26].

4. Conclusions

The reaction mechanisms of LiBOB as cathode film-forming additive have been investigated by density functional theory calculations in this work. Calculated oxidation potentials of the investigated structures decreased as the following order: carbonate (including EC, PC and DMC) > BOB\(^-\) (isolate) \(\approx\) carbonate-BOB\(^-\) clusters. Charge distribution results show that the electron of the oxidized carbonate-BOB\(^-\) cluster was taken from BOB\(^-\), indicating that BOB\(^-\) is oxidized preferentially to the investigated carbonate solvents. Decomposition reactions of the oxidized EC-BOB\(^-\) cluster show that decompositions of BOB\(^-\) are energetically favorable to EC. The most likely reaction path of this cluster is the decomposition of BOB\(^-\) via two transition states, generating CO, CO\(_2\) and radical R1, which may further terminates generating borate-containing film. All of these oxidative decomposition products are in good agreement with the experimental detections. And the borate-containing film is believed to play a crucial role in suppressing further oxidative decomposition of
carbonate solvents and improving the cyclic stability of the cell.

Additionally, we found that the existence of EC does not alter the oxidative stability and decomposition reaction energy of BOB⁻.

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1124-1129.


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Table 1 NPA Atomic charges of BOB\(^{-}\) before and after oxidation.

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<th>Isolated BOB(^{-})</th>
<th>EC-BOB(^{-})</th>
<th>PC-BOB(^{-})</th>
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<td>Before</td>
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<td>Before</td>
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Figure captions:

**Figure 1** Optimized structures of isolate carbonates, carbonate-BOB' and carbonate-PF$_6$ clusters before and after oxidation (-e), together with the calculated oxidation potentials (V vs. Li$^+$/Li). Sum of NPA charge of the oxidized clusters in red and blue circle is relative to 0 and -1 e, respectively.

**Figure 2** Optimized structures and geometric parameters (bond length in Å, dihedral angle in degree, relative free energy in kJ/mol) of the stationary points and the transition states (TSs) for the oxidative decomposition of isolate BOB'.

**Figure 3** Optimized structures of the stationary points and the transition states (TSs) for the oxidative decomposition reactions of EC-BOB'-e cluster.

**Figure 4** Relative free energy of oxidative decomposition of BOB'-e and EC-BOB'-e clusters (from Path1 to Path 7).

**Figure 5** Possible polymerization reactions and relative free energies (kJ/mol) of R1.
Figure 1 Optimized structures of isolate carbonates, carbonate-BOB' and carbonate-PF₆⁻ clusters before and after oxidation (-e), together with the calculated oxidation potentials (V vs. Li⁺/Li). Sum of NPA charge of the oxidized clusters in red and blue circle is relative to 0 and -1 e, respectively.

Figure 2 Optimized structures and geometric parameters (bond length in Å, dihedral angle in degree, relative free energy in kJ/mol) of the stationary points and the transition states (TSs) for the oxidative decomposition of isolate BOB'.
Figure 3 Optimized structures of the stationary points and the transition states (TSs) for the oxidative decomposition reactions of EC-BOB-e cluster.

Figure 4 Relative free energy of oxidative decomposition of BOB-e and EC-BOB-e clusters (from Path1 to Path 7).
Figure 5 Possible polymerization reactions and relative free energies (kJ/mol) of R1.
The most possible oxidative decomposition reaction path of EC-BOB^− cluster.