Enhancing cycle life and usable energy density of fast charging LiFePO$_4$-graphite cell by regulating electrodes’ lithium level

Vallabha Rao Rikka, Sumit Ranjan Sahu, Abhijit Chatterjee, Raju Prakash, G. Sundararajan, R. Gopalan
gopy@arci.res.in

Highlights

- Cycle life of 4320 at 4C rate with LFP/graphite cell for 80% SOC-100% DOD test
- Lithium levels on the anode and cathode are pivotal factor in the cell cycle life
- Cell failure caused by nonuniform charge gradient, gas formation, and electrode deformation
- End-of-life cell impedance is not related to cell temperature but SOC-DOD window

Rikka et al., iScience 25, 104831 September 16, 2022 © 2022 The Author(s)
https://doi.org/10.1016/j.isci.2022.104831
Enhancing cycle life and usable energy density of fast charging LiFePO$_4$-graphite cell by regulating electrodes’ lithium level

Vallabha Rao Rikka, Sumit Ranjan Sahu, Abhijit Chatterjee, Raju Prakash, G. Sundararajan, and R. Gopalan

SUMMARY
Range anxiety is a primary concern among present-day electric vehicle (EV) owners, which could be curtailed by maximizing the driving range per charge or reducing the charging time of the lithium-ion battery (LIB) pack. Maximizing the driving range is a multifaceted task as charging-discharging the LIB up to 100% of its nominal capacity is limited by the cell chemistry (voltage window) and cell operating conditions. Our studies on commercial LiFePO$_4$/graphite cells show that a cycle life of 4320 is achieved at 4C rate with 80% SOC-100% DOD combination (12 min charging time), which is the highest among the works reported with this cell chemistry. Complete utilization of electrodes’ lithium during cycling resulted in the lowest cycle life of 956. This study demonstrates LIB charging-discharging protocol enabling longer driving range with quicker charging times. Besides, it might endow promising possibilities of future EV LIB packs with reduced size/weight and high safety.

INTRODUCTION
Lithium-ion batteries (LIB) are extensively employed as the power source in a wide range of applications, from portable electronic devices to electric vehicles (EV), owing to their high energy density, power density, lightweight, low self-discharge features, wide operating temperatures, and long cycle life (Tarascon and Armand, 2001; Choi and Aurbach, 2016; Bourzac, 2015). The current EV industry demands charging the battery up to 80% state of charge (SOC) within 15 min with a battery life of up to 15 years (Liu et al., 2019; Yang et al., 2018, Yang et al., 2019; Tomaszewska et al., 2019). To date, the cycle life of the LIBs is maximized by limiting the operating range of SOC, depth of discharge (DOD), temperature, and C-rate (Woody et al., 2020; Lunz et al., 2012; Kostopoulos et al., 2020; Groot et al., 2015; Sarasketa-Zabala et al., 2015; Wang et al., 2011; de Vries et al., 2015). Besides, achieving long cycle life and higher cell energy utilization simultaneously during fast charging of LIBs are the major challenges for the widespread adoption of LIBs for EVs (Yang and Wang, 2018). Several strategies have been undertaken to overcome the aforementioned challenges, but more focus has been given to either choosing a range of high energy density battery materials or structuring the design of the electrodes (Shen et al., 2021; Rohan et al., 2014). Despite this, significant capacity fade is being observed in LIBs depending on the charge-discharge parameters, i.e., SOC, DOD, operating temperature, and C-rate during cycling (Preger et al., 2020; Ecker et al., 2017).

SOC and DOD levels are important performance parameters of the LIBs, indicating the energy utilization of a cell per cycle. Although higher SOC and DOD (△SOC) levels result in higher cell energy utilization per cycle and longer driving range per charge in EVs, the associated cycle life and safety-related issues should be critically analyzed to achieve the desired LIB performance during fast charging. The high SOC levels often lead to fast decay in the cycle life and the potential safety issues in the lithium-ion cell as the anode potential reaches close to the lithium plating potential (Gao et al., 2018; Edge et al., 2021; Wikner et al., 2021). In addition, most cathode materials, such as spinel and layered-typed cathodes, experience structural degradation in the form of oxygen evolution at low SOC levels. On the other hand, olivine-type cathodes could potentially undergo transition metal dissolution, which can result in an inventory loss of lithium, the loss of active material, and a significant decrease in capacity (Zhan et al., 2018). The high DOD levels can also result in capacity loss, impact electrode stability, and lead to anode current collector corrosion (Gu et al., 2014; Saxena et al., 2016). Lowering the SOC and DOD levels might reduce the aforementioned
issues and potentially lead to lower cell energy utilization per cycle and range anxiety in EVs. While the individual effect of SOC and DOD on the cycle life of LIB is well known, the combinatorial effect of SOC-DOD on the cycle life of LIB is still very little understood (Kupper et al., 2018). Hence, identifying the optimum operating SOC-DOD combination for LIB is pivotal to meet the desired mileage, short charging time, long battery life, and safety in current and future generation EVs. Moreover, the cell cycle life fade is primarily due to the degradation of the electrodes and cell components during cycling (Li et al., 2014; Palacín, 2018). Therefore, a thorough understanding of the cell degradation mechanisms is vital to identify the optimum SOC-DOD combination to accomplish desired cell performance and long cycle life at fast charging rates.

This work essentially brings out a comprehensive study on the cycle life of 2.5 Ah, 26650-type cylindrical LiFePO₄/graphite lithium-ion cells at a constant 4C charge-discharge rate with various SOC and DOD combinations. LiFePO₄/graphite cells are promising candidates for examining cycle life due to their excellent structural stability during cycling at high SOC-DOD levels and high C-rates (Kang and Ceder, 2009; Asenbauer et al., 2020). A test matrix was designed with combinations of SOC and DOD in the range of 80%–100%, which is on par with the current global demands for fast charging EV applications.

RESULTS AND DISCUSSION

Figure 1 shows CT cross-section images of the pristine lithium-ion cell. The cell comprises an aluminum metal casing and a jelly roll consisting of an anode, a cathode, and a separator. The pristine cell did not show any deformation in the jelly roll (Figure 1A). A hollow cylindrical nickel tube is placed at the center
of the jelly roll to stabilize the shape of the jelly roll. The hollow tube also provides a clear path for the gases to flow from the cell base to the pressure release valve located at the top of the cell. Four anode tabs and four cathode tabs are located inside the jelly roll and connected to the electrical bus bar at the top and bottom of the cylindrical cell, respectively, shown in Figure 1A. The four tabs appear as bright semi-circular arches in the CT cross-section image (taken from the dotted line portion in Figure 1A), as shown in Figure 1B. The anode bus bar is connected to the top lid, while the cathode bus bar is connected to the bottom of the cell casing.

Figure 1C shows a cross-sectional CT scan image of the electrodes and separator layers. A-line profile (Figure 1D) was acquired along the diameter of the jelly roll in Figure 1B. The jelly roll consists of 38 turns of electrodes and separators as calculated from the line signals in Figure 1D. The jelly roll includes high atomic number elements (Al, Cu, Fe, and Ni) and low atomic number elements (Li, C, P, and O). Elements with higher atomic numbers typically display a higher grayscale value and appear brighter than those with lower atomic numbers. The width and length of the cathode were measured to be 5.55 cm and 172 cm, respectively, whereas, for the anode, it was 5.75 cm and 172 cm. The higher width of the anode over the cathode could prevent lithium plating during fast charging (Son et al., 2013).

The BOL cell capacity got altered significantly by the different SOC-DOD test conditions due to the variation in the cycling lithium, Li leftover/lithiation and Li leftover/de-lithiation of the anode and cathode, as shown in Figure 2, Table S1 and Figure S2. A representative diagram is shown in Figure 3. The terms “Li leftover/lithiation” and “Li leftover/de-lithiation” refer to the incomplete delithiation and lithiation of the electrodes, respectively. Anode lithiation level (SOC%) is described as the sum of cycling lithium level and Li leftover/de-lithiation level of the anode, while the cathode lithiation level (SOC%) is the sum of cycling lithium level and Li leftover/de-lithiation level of the cathode.

As the lithiation level of the electrodes changed based on the test conditions, the areal capacity (Ah/cm²) and the capacity ratio of the electrodes (anode to cathode and cathode to anode) are changed accordingly, which in turn change the effective C-rate experienced by the electrodes for a given current (A) (Xu et al., 2015; Wang et al., 2012). The electrode’s current density (A/cm²) and areal capacity (Ah/cm²) were used to determine the effective C-rate (Heubner et al., 2019). The electrode capacity and C-rate ratio for all the test conditions were calculated and shown in Table S1, represented in Figure 3. The BOL results indicate that the effective C-rate of the electrode becomes greater than the applied C-rate as the lithiation level of the electrode decreases. If the effective C-rate is higher than the applied C-rate for a given current (A), a significant amount of lithium-ion depletion occurs inside the electrode (Heubner et al., 2019). In this case, 9 different SOC-DOD test conditions result in different effective anode and cathode C-rates in the cells for an applied 4C rate current (10 A). Thus, the magnitude and nature of the cell aging might also vary over the course of continuous charge-discharge cycles. It has been reported that anode aging limits...
Figure 3. Representative plots for FCFD, FCPD, PCFD, and PCPD test conditions: Percentage of cycling lithium, Li\text{leftover/de-lithiation} and Li\text{leftover/lithiation} levels of the anode and cathode; electrode capacity, and C-rate ratio.
the rate capability and the cycle life of a lithium-ion cell (Lewerenz et al., 2017; Li et al., 2018). Therefore, to identify the primary aging effects on the cycle life of a lithium-ion cell, all the 9 SOC-DOD test conditions are classified into four groups based on the lithiation and delithiation level of the anode, i) fully charged and fully discharged (FCFD), ii) fully charged and partially discharged (FCPD), iii) partial charged and fully discharged (PCFD), and iv) partial charged and partial discharged (PCPD). In all the cases, the discharge capacity in Table S1 is considered as the cell capacity as that is the active capacity cycling between cathode and anode.

The cell tested at the T100-100 test condition corresponds to the FCFD group. T100-100 test condition showed the capacity of the anode and cathode equal to 2.5 Ah at 4C. Hence, the effective C-rate on the anode and cathode equals the applied 4C. The lithium-ion cells tested at T100-90 and T100-80 conditions correspond to the FCPD group. The cathode is not completely lithiated at the end of the cell discharge due to 10% (T100-90) and 20% (T100-80) un-delithiation of the anode, hence reducing the cell capacity to 2.25 Ah and 2 Ah for T100-90 and T100-80 test conditions, respectively. Thus, the effective C-rate on the cathode is 1.11 times and 1.25 times higher than that of 4C for T100-90 and T100-80, respectively. While the effective C-rate on the anode is 4C.

While the effective C-rate on the cathode and anode differs based on the test conditions. An important observation in this class was that the effective C-rate on both the cathode and anode remains above 4C. The EOL cycle number (capacity retention of 80%), BOL and EOL temperature and impedance (Ro) of the cell were recorded for all test settings (Table 1), detailing the experimental method depicted in Figure 4.

Herein, we correlated capacity fading with cell temperature rather than cell impedance over cycles. Since impedance measurements were only taken once every 100 cycles under OCV conditions, it is possible that these results do not reliably reflect the impact of cell temperature on capacity fading over subsequent charge discharge cycles. Nevertheless, the progression of the impedance over the cycles was depicted in the Figure S3.

**Table 1. EOL cycle number, cell temperature, and cell impedance (BOL and EOL) for 9 test conditions**

| Test condition | EOL (Number of cycles) | BOL temperature(°C) | Cell impedance (Ro) mΩ at 1KHz |
|----------------|-----------------------|---------------------|-------------------------------|
| T100-100       | 956                   | 36.4                | 47.6                          |
| T100-90        | 2000                  | 35                  | 46.5                          | 14.78 | 30.80 |
| T100-80        | 3200                  | 35                  | 46.5                          | 14.78 | 25.86 |
| T90-100        | 2300                  | 35                  | 44                            | 14.78 | 27.64 |
| T90-90         | 2500                  | 34.5                | 47.2                          | 14.78 | 21.15 |
| T90-80         | 1970                  | 34.5                | 46                            | 14.78 | 37.8   |
| T80-100        | 4320                  | 35                  | 43.6                          | 14.78 | 55.8   |
| T80-90         | 2650                  | 35                  | 46.3                          | 14.78 | 22.33 |
| T80-80         | 2100                  | 34                  | 46.3                          | 14.78 | 49.2   |

Figure 5 shows the life cycle data (capacity retention and cell temperature vs. full equivalent cycles [FEC]) for 9 test conditions at 4C. Maximum cycle life of 4320 was obtained for T80-100, while the T100-100 condition delivered a minimum cycle life of 956. The onset of the capacity fade was observed for all test conditions at different FEC; afterward, a rapid capacity fade up to 80% of the BOL capacity was observed for all the cells. The onset of capacity fade was observed relatively earlier for FCFD (~400th cycle), whereas it occurred...
around 1000–1500 cycles and 1500–2500 cycles for FCPD and PCFD test conditions, respectively. In the case of PCPD, the onset capacity fade was identified after 1000 cycles. After the onset of capacity fade, a gradual increase in the cell temperature was observed (Figure 5) for all the cases. The measured BOL cell temperatures were found to be between 34°C and 37°C at the BOL, while the EOL temperatures were recorded between 43°C and 48°C. The cyclic stability and the temperature profiles for each test condition were distinct in nature.

The cell temperature of the T 100-100 cell increased to ~40°C up to 400 FEC (Figure 5B) when the capacity fade initiates in the cell. It is well accepted that cycling the cell at high SOC and high C-rate will cause significant mechanical strain on the graphite lattice of the anode electrode due to the gradient of lithium ions and volume changes (13%) of the graphite particle (Aurbach et al., 2002). These volumetric changes, together with the mechanical strain caused by the gradient of lithium ions, can fracture the graphite particle and raise the cell’s temperature. Herein, the mechanical strain could not exceed the fracture strength of the graphite materials during the initial cycles and hence could not result in anode cracking or loss of active material, as indicated by the fact that capacity degradation occurred only after 400 FEC. However, after 400 FEC, the cell temperature exceeded 40°C, and concurrent capacity fade was observed. It is commonly accepted that the formation of gas pockets between the electrodes, due to the cell temperature above 40°C, has a significant impact on its performance, resulting in a rapid capacity loss (Winter et al., 1998; Xiong et al., 2014). As a result of the cumulative effect of all of these events, the T 100-100 cell has the shortest cycle life of 956 cycles (Figure 5A).

The cycle life of the FCPD cells (T 100-80 and T 100-90) does not show a straightforward SOC dependence on the cycle life. Instead, a higher DOD caused faster capacity decay (Gu et al., 2014). For example, the cycle life of the T 100-90 cell is less than the T 100-80 cell (Figure 5C). The cycle life for T 100-90 and T 100-80 were 2000 and 3200 (second highest cycle life), respectively. Although the final cell temperature was found to be ~47°C for both the cells, the rate of increase in temperature was higher for the T 100-90 cell (Figure 5D). The T 80-100 cell of the PCFD group showed the highest cycle life (4320 cycles) (Figure 5E), while the T 90-100 cell had almost half of the cycle life (2300 cycles) as T 80-100. Comparing the FCPD and PCFD cells results, it could be well understood that lowering the SOC with high DOD levels (fully discharge) significantly enhances the cycle life. It also contradicts the results described in earlier reports where the cells cycled at 100% DOD led to shorter cycle life (Gu et al., 2014; Kupper et al., 2018). The cycle life has followed the order of T 80-100 > T 100-80 > T 90-100 > T 100-90 > T 100-100, signifying the fact that the electrode lithiation-delithiation level (SOC-DOD) had a prominent effect on the cycle life of lithium-ion cells at a constant high C-rate (4C) charge-discharge. Additionally, these findings indicate that storing excess lithium in the cathode or anode prolongs the cycle life of lithium-ion cells compared to complete lithium extraction from both electrodes. Due to the increased overpotentials at high C-rates, the electrodes cannot be delithiated or lithiated to the rated cell capacities, resulting in the inactivity of the active lithium stored in the electrodes and a significant reduction in the cycle life. The lithium loss might probably get replenished from the lithium stored in the cathode during the cycling due to the absence of a thick interface layer and several lithium-ion intercalation phases like that on the anode.

For all PCPD group cells (T 90-90, T 80-80, T 80-90, and T 90-80), the cycle life was found in the range of 1970–2650 (Figure 5G). The cycle life of the cells was in the order of T 80-90 > T 90-90 > T 80-80 > T 90-80. As discussed earlier, the effective C-rate on both the cathode and anode remains above 4C for these cells (Figure 3), resulting in faster capacity decay and relatively lower cycle life than the FCPD and PCFD cells. This group of cells also demonstrates that the amount of lithium cycling between cathode and anode does not influence the cycle life of the cell; rather, the SOC-DOD levels play a decisive role. The T 80-90 and T 90-80 cells, having 72% cycling lithium level (Figure 3), showed a cycle life of 2650 and 1970, respectively. While the T 90-90 and...
T80-80 cells had 81% and 64% cycling lithium levels exhibited moderate cycle life of 2500 and 2100, respectively.

CT images of the cells were acquired at BOL and EOL cycles to analyze the structural degradation and mechanical failures. Figure 6 shows the CT images of the lithium-ion cell at BOL and EOL cycles for the T100-100 test condition. The BOL cell showed a compact jelly roll with no gaps between the electrode and separator layers (Figures 6A and 6C). At EOL, gas pockets were observed throughout the cross-section of the jelly roll (Figure 6B) and close to the positive terminal for the cell (Figure 6D). The gas pockets were formed due to the rapid increase in the cell temperature per cycle (Figure 5B) and subsequent electrolyte decomposition. As a result, the gap between the jelly roll and the positive terminal increased from 2.27 mm (BOL) to 3.12 mm (EOL) (Figure 6D). Excess pressure generated due to the gas formation; the terminal-to-lid welding detached at EOL (Figure 6D). The deformation of electrodes was also observed near the hollow tube (Figure 6B). The magnified views of the jelly rolls shown in Figures 6E and 6F reveal the degradation of the LFP electrode and wrinkle formation in the anode electrode at EOL.

CT images of the cells were acquired at BOL and EOL cycles to analyze the structural degradation and mechanical failures. Figure 6 shows the CT images of the lithium-ion cell at BOL and EOL cycles for the T100-100 test condition. The BOL cell showed a compact jelly roll with no gaps between the electrode and separator layers (Figures 6A and 6C). At EOL, gas pockets were observed throughout the cross-section of the jelly roll (Figure 6B) and close to the positive terminal for the cell (Figure 6D). The gas pockets were formed due to the rapid increase in the cell temperature per cycle (Figure 5B) and subsequent electrolyte decomposition. As a result, the gap between the jelly roll and the positive terminal increased from 2.27 mm (BOL) to 3.12 mm (EOL) (Figure 6D). Excess pressure generated due to the gas formation; the terminal-to-lid welding detached at EOL (Figure 6D). The deformation of electrodes was also observed near the hollow tube (Figure 6B). The magnified views of the jelly rolls shown in Figures 6E and 6F reveal the degradation of the LFP electrode and wrinkle formation in the anode electrode at EOL.

CT cross-sectional images of the FCFD, FCPD, PCFD, and PCPD cells are compared in Figure 7. The deformation of the jelly roll was found to be more perceptible and randomly disseminated across the diameter of the jelly roll for the T80-80, T90-100, T90-80, and T100-100 cells compared to the T80-100 and T100-80 counterparts. For all the cells, electrode deformation was evident near the central hollow tube along with the sharp bends in the electrode layers directed toward the core of the jelly roll.
These deformations were observed up to the 15th and 17th layers for the T_80-80 and T_90-100 cells, respectively. The formation of gas pockets between the electrode layers resulted in strain in the spiral-wound jelly roll. This strain introduced sharp bends at the interior layers of the jelly roll for stress relaxation. Due to the flexible nature of the separator, the deformation was extended from the origin to the center of the jelly roll and deformed along with electrodes. It can be seen from Figure 7, T_80-100 test cell (Figure 7B) showed lower deformation, which extended up to ~10 layers. Whereas other test conditions showed more deformation, extended up to several layers. Due to the lower deformation of the jelly roll for the T_90-100 test condition, the cell exhibited the highest number of charge/discharge cycles (4320 cycles), T_100-80 test cell exhibited the second-highest number of cycles (3200 cycles). In addition, the formation of gas pockets between electrode layers leads to increased cell impedance (Figure S4), causes temperature rise over the charge-discharge cycles.

To further elucidate the effect of mechanical deformation and structural changes in the electrode on the cell cycle life, all nine cells were dismantled after the EOL cycle, and the failure analysis was conducted. Figure 8A shows the digital photographs of graphite electrodes after the EOL cycle for all 9 test conditions. Significant changes in the macroscopic features of graphite electrodes were observed upon visual inspection compared to the BOL electrode. The extensive delamination of the graphite electrode from copper foil was observed at the EOL cycle for the T_100-100 cell, which could be attributed to volume changes of the graphite particles during the full lithiation-delithiation of the graphite anode and binder degradation over the cycles. The graphite electrodes obtained from T_90-100 and T_100-90 cells show a thick and dense shiny gray-colored layer in the middle of the electrode. The color changes from shiny gray to maroon toward the electrode’s edges.

In contrast, for the T_90-90 graphite electrode, the shiny gray layer was observed toward the edges, and the maroon-colored layer was observed in the middle of the electrode. The graphite electrodes from the T_80-90 and T_90-80 cells show a thick golden color portion in the middle of the electrode. The golden color was observed at the electrode edges for the T_80-80 electrode. The different colors on the graphite electrodes
indicate different levels of lithiation (Kang et al., 2020; Finegan et al., 2020; Guo et al., 2016; Shellikeri et al.,
2017) induced by non-uniform charge distribution experienced by the electrode during long cycles. The
graphite electrode of the T80-100 and T100-80 cells did not reveal any color contrast, and they visually looked
similar to the BOL electrode (Figure S5).

The SEM images of the graphite and LFP electrodes (after the EOL cycle) for T80-80, T80-100, T90-80, and
T90-100 are shown in Figures 8B–8E. The low magnification image of the T90-80 graphite electrode showed
thick SEI-covered particles and micro cracks on the electrode surface (Figure 8B). The high-magnification
image of the golden color area on the electrode showed fewer SEI-covered graphite particles with micro-
cracks. The electrode’s high-magnification image of the middle area (black color) showed thick
SEI-covered graphite particles, indicating an increase in the local current density (A/cm²), SEI break-
down-reformation, and gas evolution. This, in turn, led to lower utilization of the graphite material and
lower cycle life of 2100. Microcracks with longer lengths were observed on the LFP electrode (Figure 8B).
The T80-100 graphite electrode does not show cracks and thick SEI-covered graphite particles (Figure 8C).
No cracks were visible on the LFP electrode too. These observations go in accordance with the highest cy-
cle life (4320 cycles) obtained for the T80-100 cell. The SEM images from the golden color middle portion of
Figure 8. (A) Digital photographs of the graphite electrode after EOL cycle. (B–E) SEM images of the graphite and LFP electrodes after EOL cycle for T80-80, T80-100, T90-80, and T90-100.
the T90-80 graphite electrode showed few microcracks and non-uniform SEI-covered graphite particles (Figure 8D). Whereas at the black color edges, thick SEI-covered graphite particles were evident. The LFP electrode, in this case, was crack-free. The shiny gray color area at the center of the T90-100 graphite electrode showed thicker SEI-covered particles than the maroon color edge (Figure 8E). The high-magnification SEM images of the LFP electrode exhibited deposits of lithium compounds on the surface.

The thick SEI deposits observed on the aged graphite electrodes in Figures 8B–8D and 8E establish that continuous formation of SEI and associated gas evolution have taken place on the graphite electrodes due to high electrochemical activity caused electrode deformation in the jelly roll (Figure 7). Decomposing the electrolyte at 4C (to form the SEI layer) might have caused inadequate electrolyte wetting across the electrode surface and accompanying structural changes in the active material. The XRD patterns acquired from the center and the edge portions of the graphite electrodes after cell discharge for T80-80, T80-100, T90-80, and T90-100 cells are shown in Figure 9A. The peaks corresponding to LiC₆, LiC₁₂, LiC₂₄, and LiC₃₆ were identified along with graphite (002). The T80-100 graphite electrode showed a small peak corresponding to the LiC₂₄ phase and graphite (002) peak after the complete delithiation, which indicates that the electrode was almost delithiated at EOL’s highest cycle life. The T90-100 graphite electrode showed LiC₆, LiC₁₂, LiC₂₄, and LiC₃₆ phases, indicating incomplete delithiation due to built-in overpotential and lower cycle life than T80-100. The T80-80 and T90-80 cells showed similar phases as T90-100 due to electrode overpotential or Li leftover/lithiation, leading to low cycle life. The complementary results were observed in the XRD patterns of the LFP electrodes after EOL (Figure 9B). The peaks corresponding to FePO₄ (FP) were observed in all the electrodes except T90-100, which indicates that all the other electrodes suffered irreversible lithium loss during cycling, which led to lower cycle life than T80-100. There was no evidence of other secondary phases that ascertain the LFP electrode’s structural integrity.

Additionally, to identify the distribution of the Li-graphite intercalation phase on the surface of the graphite electrode corresponding to the gray color portion, Raman spectra (Figure 9C) and 2D mapping (Figure 9D) were acquired from a 25 × 25 µm region on the electrode. The peak position in the spectra was used to
represent the local Li-graphite intercalation phases. The Raman spectrum of graphite contains a prominent peak at 1583 cm\(^{-1}\) attributed to the \(E_{2g2}\) vibrational mode (G band). The 1588 and 1600 cm\(^{-1}\) peaks are attributed to the solid Li\(_{12}\) and dilute Li\(_{18}\) phases, respectively (Migge et al., 2005). The Raman mapping showed the distribution of graphite, Li\(_{12}\), and Li\(_{18}\) phases corresponding to the color of the spectra in Figure 9C. The \(T_{80-80}\) and \(T_{90-80}\) revealed a higher concentration of Li\(_{12}\) and Li\(_{18}\) phases than the \(T_{80-100}\) and \(T_{90-100}\). It is to be noted that these two phases were not identified in the XRD pattern of graphite for \(T_{80-100}\).

The aging of graphite electrodes was responsible for both electrode jelly roll deformation and a reduction in overall cell capacity. On the other hand, the LFP cathode is a robust material with a high cycling rate capability and no structural deterioration. Despite the LFP electrode’s excellent performance, the degree of lithiation of the LFP electrode dropped considerably with graphite electrode aging. LFP electrodes may deteriorate if subjected to high temperatures due to gas pockets developing between electrodes at high SOC (Figure 6). The SEI layer grew progressively in the electrochemically active region of the graphite electrode. Thus, it was observed that the continuous growth of the SEI layer on the graphite surface reduced the active graphite anode surface, resulting in an unequal charge distribution over the graphite electrode surface (Figure 9). According to the XRD data, it was not a result of inhomogeneous deterioration of the LFP electrode itself (Figure 6). These values are comparable since all cells were discharged completely at the same rate prior to dismantling.

In conclusion, it is important to understand the effect of SOC-DOD operating range on key lithium-ion cell parameters such as cycle life, charging time, discharge capacity, energy density, and cell temperature during fast charging for assorted cell geometry and chemistry. A schematic image of this work comparing cell parameters and various cell operating voltage window is shown in Figure 10.

**Conclusions**

The influence of fast charging (4C-rate) with different combinations of SOC and DOD on the cycle life of the 2.5 Ah commercial LFP/graphite cell is systematically studied and summarized in Figure 10. The test matrix was formulated with the primary objective of determining the cycle life of the cells at room temperature (23°C). The rise in cell temperature lowered the cycle life of the cells. The lithium levels on the anode and cathode played a decisive factor in the cell cycle life. Our experimental investigations showed that the combination of 80% SOC-100% DOD exhibited the highest cycle life (4200 cycles) with a short charging time (12 min). To the best of the authors’ knowledge, this is the highest cycle life reported for the LFP/graphite cell at 4C. The comprehensive CT image analysis revealed gas pocket formation and electrode deformation for the 100% SOC-100% DOD cell resulting in the lowest cycle life of 956 cycles. This study
highlights that cycle life can vary non-trivially due to the associated SOC-DOD combination. Therefore, the SOC-DOD combination becomes an important parameter while maximizing the cycle life and energy utilization of LIBs at fast charging for EV applications. This study could lead to shorter charging times, longer driving range, reduced battery pack size, lower pack cost, and high safety for EV applications.

Limitations of the study
The study is limited to high-power 26650-type cylindrical LiFePO4/graphite lithium-ion cells.

STAR METHODS
Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
  - Lead contact
  - Materials availability
  - Data and code availability
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
  - Test method
  - Capacity confirmation test (CCT): Determination of BOL and EOL
  - Test matrix
  - Non-destructive cell characterization using X-Ray computed tomography
  - Structural characterization of electrodes

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.104831.

ACKNOWLEDGMENTS
The authors would like to acknowledge the financial support from the Technical Research Centre (TRC), Department of Science & Technology (TRC project: AI/1/65/ARCI/2014), Government of India to complete this work. The authors also thank the late Dr. G. Padmanabham, former Director, ARCI, for his constant encouragement and support for this work. Again, the authors thank Prof. K. Balasubraman-yan, Department of Mechanical Engineering, IIT Madras, for providing the X-ray computed tomography characterization facility. We want to acknowledge M. Mrinalini from ARCI for her help in experimental work.

AUTHOR CONTRIBUTIONS
V.R.R. and A.C. designed and performed the experiments. S.R.S. and V.R.R. wrote the manuscript with support from A.C. and G.S. G.S. and R.P. aided in interpreting the results. R.G. supervised the project and helped the financial support from the Technical Research Centre (TRC). All authors discussed the results and commented on the manuscript.

DECLARATION OF INTERESTS
The authors declare no conflict of interest.

Received: January 31, 2022
Revised: June 30, 2022
Accepted: July 20, 2022
Published: September 16, 2022

REFERENCES
Asenbauer, J., Eisenmann, T., Kuenzel, M., Kazzazi, A., Chen, Z., and Bresser, D. (2020). The success story of graphite as a lithium-ion anode material – fundamentals, remaining challenges, and recent developments including silicon (oxide) composites. Sustain. Energy Fuels 4, 5387–5416. https://doi.org/10.1039/D0SE00175A.

Aurbach, D., Zinigrad, E., Cohen, Y., and Teller, H. (2002). A short review of failure mechanisms of lithium metal and lithiated graphite anodes in liquid electrolyte solutions. Solid State Ion. 148, 405–416. https://doi.org/10.1016/S0167-2738(02)00080-2.

Bourzac, K. (2015). Batteries: 4 big questions. Nature 526, 510S. https://doi.org/10.1038/526S10S.
Choi, J.W., and Aurbach, D. (2016). Promise and reality of post-lithium-ion batteries with high energy densities. Nat. Rev. Mater. 1, 1603. https://doi.org/10.1038/natrevmats.2016.13.

de Vries, H., Nguyen, T.T., and Op het Veld, B. (2015). Increasing the cycle life of lithium-ion cells by partial state of charge cycling. Microelectron. Reliab. 55, 2247–2253. https://doi.org/10.1016/j.microrel.2015.08.014.

Ecker, M., Shafie Sabet, P., and Sauer, D.U. (2017). Influence of operational condition on lithium plating for commercial lithium-ion batteries – electrochemical experiments and post-mortem-analysis. Appl. Energy 206, 934–946. https://doi.org/10.1016/j.apenergy.2017.08.034.

Edge, J.S., O’Kane, S., Prosser, R., Kirkaldy, N.D., Patel, A.N., Hales, A., Ghosh, A., Ai, W., Chen, J., Yang, J., et al. (2021). Lithium ion battery degradation: what you need to know. Phys. Chem. Chem. Phys. 23, 8200–8221. https://doi.org/10.1039/D1CP00359C.

Finegan, D.P., Quinn, A., Wragg, D.S., Colclasure, A.M., Lu, X., Tan, C., Heenan, T.M.M., Jervis, R., Brett, D.J.L., Dai, S., et al. (2020). Spatial dynamics of lithium and lithium plating during high-rate operation of graphite electrodes. Energy Environ. Sci. 13, 2570–2584. https://doi.org/10.1039/D0EE01191F.

Finegan, D.P., Schell, M., Robinson, J.B., Tjaden, B., Hunt, I., Mason, T.J., Millichamp, J., Di Michel, M., Offer, G.J., Hinds, G., et al. (2015). In-operando high-speed tomography of lithium-ion batteries during thermal runaway. Nat. Commun. 6, 6924. https://doi.org/10.1038/ncomms7924.

Gao, Y., Jiang, J., Zhang, C., Zhang, W., and Jiang, Y. (2018). Aging mechanisms under different state-of-charge ranges and the multi-indicators system of state-of-health for lithium-ion batteries. J. Electrochem. Soc. 165, A3486–A3480. https://doi.org/10.1149/2.0941814jes.

Leverenz, M., Warnecke, A., and Sauer, D.U. (2017). Post-mortem analysis on LiFePO4/Graphite cells describing the evolution & composition of covering layer on anode and their impact on cell performance. J. Power Sources 369, 122–132. https://doi.org/10.1016/j.jpowsour.2017.10.003.

Li, D., Danilov, D.L., Zwikirsch, B., Fichtner, M., Yang, Y., Eichel, R.A., and Notton, P.H. (2018). Modeling the degradation mechanisms of C6/LiFePO4 batteries. J. Power Sources 375, 106–117. https://doi.org/10.1016/j.jpowsour.2017.11.049.

Li, Y., El Gabaly, F., Ferguson, T.R., Smith, R.B., Bartelt, N.C., Sugar, J.D., Fenton, K.R., Cogswell, D.A., Kilcoyne, A.L.D., Tyliszczak, T., et al. (2014). Current-induced transition from particle-by-particle to concurrent intercalation in phase-separating battery electrodes. Nat. Mater. 13, 1149–1156. https://doi.org/10.1038/nmat4084.

Li, S., Liu, Y., and Ma, Q. (2019). Challenges and opportunities towards fast-charging battery materials. Talanta 202, 540–545. https://doi.org/10.1016/j.talanta.2018.10.010.

Lutz, B., Yan, Z., Gerschler, J.B., and Sauer, D.U. (2012). Influence of plug-in hybrid electric vehicle charging strategies on charging and battery degradation costs. Energy Pol. 46, 511–519. https://doi.org/10.1016/j.enpol.2012.04.017.

Migge, S., Sandmann, G., Rahner, D., Dietz, H., and Plie, W. (2005). Studying lithium intercalation into graphite particles via in situ Raman spectroscopy and confocal microscopy. J. Solid State Electrochem. 9, 132–137. https://doi.org/10.1007/s10008-004-0563-4.

Palacin, M.R. (2018). Understanding ageing in lithium-ion batteries: a chemical issue. Chem. Soc. Rev. 47, 4924–4933. https://doi.org/10.1039/C7CS00889A.

Pregler, Y., Barkholtz, H.M., Fresquez, A., Campbell, D.L., Juba, B.W., Roman-Kustas, J., Ferrera, S.R., and Chalamala, B. (2020). Degradation of commercial lithium-ion cells as a function of chemistry and cycling conditions. J. Electrochem. Soc. 167, 120532. https://doi.org/10.1149/1945-7117/abea37.

Rohan, J.F., Hasan, M., Patil, S., Casey, D.P., and Clancy, T. (2014). Energy Storage: Battery Materials and Architectures at the Nanoscale (ICOT - Energy - Concepts Towards Zero - Power Information and Communication Technology). https://doi.org/10.5772/57139.

Sarasaka-Zabala, E., Gandiga, I., Martinez-Lasema, E., Rodriguez-Martinez, L., and Villarreal, I. (2015). Cycle ageing analysis of a LiFePO4/graphite cell with dynamic model validations: towards realistic lifetime predictions. J. Power Sources 275, 573–587. https://doi.org/10.1016/j.jpowsour.2014.10.153.

Saxena, S., Hendricks, C., and Pecht, M. (2016). Cycle life testing and modeling of graphite/ LiCoO2 cells under different state of charge ranges. J. Power Sources 327, 394–400. https://doi.org/10.1016/j.jpowsour.2016.07.057.

Shellikeri, A., Watson, V., Adams, D., Kalu, E.E., Reed, J.A., Iow, T.R., Zheng, J.S., and Zheng, J.P. (2017). Investigation of pre-lithiation in graphite and hard-carbon anodes using different lithium source structures. J. Electrochem. Soc. 164, A3914–A3924. https://doi.org/10.1149/2.1511714jes.

Shen, X., Zhang, X.Q., Ding, F., Huang, J.Q., Xu, R., Chen, X., Yan, C., Su, F.Y., Chen, C.M., Liu, X., and Zhang, Q. (2021). Advanced electrode materials in lithium batteries: retrospect and prospect. Energy Mater. Adv. 2021, 1–15. https://doi.org/10.1039/D1EM00324A.

Son, B., Ryoo, M.H., Choi, J., Kim, S.H., Ko, J.M., and Lee, Y.M. (2013). Effect of cathode/anode area ratio on electrochemical performance of lithium-ion batteries. J. Power Sources 243, 641–647. https://doi.org/10.1016/j.jpowsour.2013.06.062.

Tarascon, J.M., and Armand, M. (2001). Issues and challenges facing rechargeable lithium batteries. Nature 414, 359–367. https://doi.org/10.1038/35104644.

Tomaszewski, A., Chu, Z., Feng, X., O’Kane, S., Liu, X., Chen, J., Ji, C., Endler, E., Li, R., Liu, L., et al. (2019). Lithium-ion battery fast charging: a review. eTransportation 1, 100011. https://doi.org/10.1016/j.etran.2019.100011.

Wang, J., Liu, P., Hicks-Garner, J., Sherman, E., Soukiazian, S., Verbrugge, M., Tataria, H., Musser, J., and Finanmore, P. (2011). Cycle-life model for graphite/LiFePO4 cells. J. Power Sources 196, 3942–3948. https://doi.org/10.1016/j.jpowsour.2010.11.134.

Wang, M., Li, J., He, X., Wu, H., and Wan, C. (2018). Effect of local current density on electrode design for lithium-ion batteries. J. Power Sources 207, 127–133. https://doi.org/10.1016/j.jpowsour.2011.12.063.

Wilker, E., Björklund, E., Fridner, J., Brandell, D., and Thiringer, T. (2021). How the utilised SOC window in commercial Li-ion pouch cells influence battery ageing. J. Power Sourc. Adv. 8, 100054. https://doi.org/10.1016/j.powers2021.100054.

Winters, M., Besenhard, J.O., Spahr, M.E., and Novak, P. (1998). Insertion electrode materials for rechargeable lithium batteries. Adv. Mater. 10, 725–763. https://doi.org/10.1002/(SICI)1097-4652(199807)10:7<725::AID-ADMA725>3.0.CO;2-Z.
Woody, M., Arbabzadeh, M., Lewis, G.M., Keoleian, G.A., and Stefanopoulou, A. (2020). Strategies to limit degradation and maximize Li-ion battery service lifetime - critical review and guidance for stakeholders. J. Energy Storage 28, 101231. https://doi.org/10.1016/j.est.2020.101231.

Xiong, R., Sun, F., Gong, X., and Gao, C. (2014). A data-driven based adaptive state of charge estimator of lithium-ion polymer battery used in electric vehicles. Appl. Energy 113, 1421–1433. https://doi.org/10.1016/j.apenergy.2013.09.006.

Xu, J., Deshpande, R.D., Pan, J., Cheng, Y.T., and Battaglia, V.S. (2015). Electrode side reactions, capacity loss and mechanical degradation in lithium-ion batteries. J. Electrochem. Soc. 162, A2026–A2035. https://doi.org/10.1149/2.0291510jes.

Yang, X.G., and Wang, C.Y. (2018). Understanding the trilemma of fast charging, energy density and cycle life of lithium-ion batteries. J. Power Sources 402, 489–498. https://doi.org/10.1016/j.jpowsour.2018.09.069.

Yang, X.G., Liu, T., Gao, Y., Ge, S., Leng, Y., Wang, D., and Wang, C.Y. (2019). Asymmetric temperature modulation for extreme fast charging of lithium-ion batteries. Joule 3, 3002–3019. https://doi.org/10.1016/j.joule.2019.09.021.

Yang, X.G., Zhang, G., Ge, S., and Wang, C.Y. (2018). Fast charging of lithium-ion batteries at all temperatures. Proc. Natl. Acad. Sci. USA 115, 7266–7271. https://doi.org/10.1073/pnas.1807115115.

Zhan, C., Wu, T., Lu, J., and Amine, K. (2018). Dissolution, migration, and deposition of transition metal ions in Li-ion batteries exemplified by Mn-based cathodes – a critical review. Energy Environ. Sci. 11, 243–257. https://doi.org/10.1039/C7EE03122J.
**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Lithium ion cells | A123 systems, Endrich Co., Ltd. | Order No#20160406000000088 |
| 3.2V-2.5Ah cylindrical 26650 LiFePO4-graphite lithium-ion cells (100 numbers with Tabs) | Bitrode corporation | LCV8 60-100 |
| Life cycle tester (Input power: 3 Phase, 400 V, 50 Hz +/- 10% input Power variation) | VisualCN Lab Client Software |  |
| 16 Ch - RIO, CVM, 8.000V, 20 ft leads | Bitrode corporation |  |
| 16 Ch RIO, temperature monitoring | VisuaLCN Lab Client Software |  |
| Electrochemical impedance spectroscopy | Ametek scientific instruments | PARSTAT MC 8 Channels |
| Field emission SEM equipped with energy-dispersive X-ray spectroscopy (EDS) unit. | Zeiss Merlin |  |
| Raman Analaysis | Horiba Jobin Yvon | Lab RAM HR-800 system (argon laser of excitation wavelength 532 nm) |
| XRD instrument | Rigaku | Cu Kα radiation, λ = 1.5418 Å, |
| X-ray Tomography analysis | GE | Phoenix Vtomex; 240kV/320W |

**RESOURCE AVAILABILITY**

**Lead contact**

Further information should be directed to and will be fulfilled by the lead contact, R. Gopalan (gopy@arci.res.in).

**Materials availability**

This study did not generate new materials.

**Data and code availability**

All the data supporting this study have been shown in the article and Supporting information. Other related data are available from the corresponding author upon reasonable request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

In the present study, a single production batch of 26650-type commercial LiFePO4/graphite lithium-ion cells (A123 systems) were used to determine the cycle life. Each cell had a nominal capacity of 2.5 Ah and a nominal voltage of 3.2 V and was cycled at a constant 4C-rate with different predetermined combinations of SOC and DOD. A test matrix was designed to determine the optimum SOC-DOD combination for obtaining the highest cycle life. The cycle life studies were conducted sequentially to identify the optimum test condition for obtaining the highest cycle life for each test condition. In order to ensure the consistency and reproducibility of the test results, two cells for each of the test conditions were evaluated. Herein, the cycle life experiments were carried out at room temperature (23°C).

**Test method**

**Overview of the test sequence**

Figure 4 illustrates an overview of the test sequence for the cycle life study. Initially, capacity confirmation tests (CCT) were carried out at different C-rates to determine the cells’ beginning-of-life (BOL) capacity. This was followed by forming a test matrix with nine different SOC-DOD combinations. The cell cycling aging tests were carried out using the test matrix until 20% of its initial capacity fade (80% capacity retention), end-of-life (EOL).
Capacity confirmation test (CCT): Determination of BOL and EOL

Before cycle aging studies, the CCT has performed in the constant current (CC) method for five subsequent charge-discharge cycles at 0.5, 1, 2, and 4C-rates. The cells showed a discharge capacity of 2.5 Ah at all C-rates, which was considered the nominal cell capacity (Figure S1). As per the industry standards, a pristine cell’s discharge capacity is denoted as the cell capacity (BOL capacity). The BOL capacity and discharge voltage depend on the operating SOC and DOD range. In general, the SOC of a cell is defined based on the lithiation state of the anode. For example, 100% SOC implies that the anode is fully lithiated. Similarly, the DOD is defined in terms of the delithiation of the anode. Three different situations arise, as illustrated in Figure 2. If the anode is fully charged (lithiated) and discharged (delithiated) (Figure 2A), then the cycling charge between the cathode and anode is equal to the total charge (Q Ah) of the cathode. In case, amount of charge is transferred from cathode to anode during the first charge (Figure 2B); we write

\[
\text{SOC} = \frac{Q^I}{Q} \quad \text{(Equation 1)}
\]

Thus, Q^I and Q < Q denotes full and partial charging, respectively. Thereafter, if the charge Q^II is transferred between anode and cathode in subsequent cycles (Figure 2C), causing the charge level at the cathode to alternate between Q – Q^I and (Q – Q^I) + Q^II. Thus, the DOD is calculated as the ratio of the amount of charge Q^II transferred to the cathode from the total charge of the anode (Q^I). We write

\[
\text{DOD} = \frac{Q^II}{Q^I} \quad \text{(Equation 2)}
\]

Q^II = Q^I and Q^II < Q^I results in a full and partial discharge, respectively. From Equations (1) and (2), the BOL capacity for each test condition is calculated as

\[
\text{BOL capacity} = Q^I = Q \times \text{SOC} \times \text{DOD} \quad \text{(Equation 3)}
\]

Denoting Q^II as the cell BOL capacity, the end-of-life (EOL capacity) is calculated as

\[
\text{EOL capacity} = Q^I \times 80\% \quad \text{(Equation 4)}
\]

In Equation (4), the cycle number at EOL capacity denotes the cell cycle life. It indicates that the cycling charge (Q^II) or the operating range of SOC-DOD influences the cell cycle life. Only a few reports are available on the cell cycle life studies under various SOC-DOD combinations, and in many of these reports, the cycle life studies were performed with either variable SOC and fixed DOD levels or fixed SOC and variable DOD levels (Lunz et al., 2012; Sarasketa-Zabala et al., 2015; Wang et al., 2011; Gao et al., 2018). Moreover, the reported optimal SOC level to achieve a high cell cycle life was below 80%, which would be inappropriate for EVs to obtain the desired driving range per charge. Hence, the present study aims to identify the optimum test condition for obtaining higher cell cycle life at maximum utilization of cell energy (≥80% SOC charge) at a 4C charge-discharge rate.

Test matrix

A test matrix was designed with three different SOC and DOD levels, between 80% to 100% (Table 1). Each test condition is denoted by T_{SOC-DOD}. The discharge voltage, charge time/capacity, and discharge time/capacity for each test condition were measured from initial charge-discharge cycles at a 4C rate and are tabulated in Table 1. In the present study, charge and discharge cycles were performed under the constant current (CC) method. The amount of charge Q^II cycling between cathode and anode is the same for the charging and discharging; thus, the cell charge and discharge times are equal for each test condition. The cycling test continued until the EOL cycle (20% capacity fade) for all test conditions without any rest period given between cycle-to-cycle. In the present study, the discharge voltage, the cycling charge Q^II and the charge/discharge times were set as operating cut-off limits for cycling experiments to identify the EOL cycle (Table 1). On reaching one of the cut-off limits during the discharge, the cell will proceed to the next cycle. A Bitrode life cycle tester (model LCV 100–60) was used for the life cycle tests. The cell aging at 4C for each test condition is followed by the reference performance test (RPT), as shown in Figure 4. The RPT is performed.

Non-destructive cell characterization using X-Ray computed tomography

X-ray computed tomography (X-ray CT) characterizations were used to monitor the cell degradation after EOL and compared to the BOL. X-ray CT technique, owing to its non-destructive nature, is a well-adopted measurement to inspect the cell degradation mechanisms beyond the electrochemical diagnosis by
visually identifying the microscopic structural changes of the cell and its components (Finegan et al., 2015). The cell imaging was carried out using an industrial X-ray 3D CT system (GE Phoenix Vtomex; 240kV/320W). The lithium-ion cell was mounted vertically to the sample rotating platform of the CT system and rotated between 0 and 360 degrees during the CT scan. The CT scan took an hour to complete. The image resolution was ~25–30 μm, and the contrast was enhanced using Volume Graphics Studio MAX 2.0. Approximately 500 tomographic images for each cell were processed at distances in equal interval distances to detect the structural defects in the electrodes jelly roll. In the present study, all images were analyzed using the open-source Image-J software.

**Structural characterization of electrodes**

After the EOL cycle had reached each test condition, all cells were dissembled inside an argon-filled glovebox in the discharged cutoff voltage state for each test condition (oxygen and moisture content <1 ppm). The electrodes were rinsed carefully with dimethyl carbonate and dried inside an argon-filled glovebox for 24 h. The crystal structure of the aged graphite and LFP electrodes was studied using a Rigaku XRD instrument (Cu Kα radiation, λ = 1.5418 Å, scattering angles of 20–40° and step size of 0.5°). 2D Raman mapping of the electrodes was recorded at room temperature on a Horiba Jobin Yvon Lab RAM HR-800 system (argon laser of excitation wavelength 532 nm). The acquisition time was 1h, and the spectral resolution was 1 cm⁻¹. The digital photographs were taken using a high-quality digital camera. The morphology of the samples was characterized using a Zeiss Merlin field emission SEM (accelerating voltage of 20 kV, working distance 8 mm) equipped with energy-dispersive X-ray spectroscopy (EDS) unit.