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# **Carboxylic Acid Assisted Synthesis of Crystalline Silicon Derived from Coal Fly-Ash for Li-Ion Batteries Anode Material**

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**Abstract**: Demand for lithium-ion batteries (LIB) in wireless electronics, power tools, electric vehicles, and reliable energy storage systems is increasing. To meet lithium-ion battery (LIB) requirements, silicon or Si has been considered a state-of-the-art anode material owing to its significantly different specific capacity compared to the other anode. In this study, we attempted to synthesize crystalline Si (cSi) from coal combustion fly ash. The Si was prepared by caustic extraction, acidic precipitation, and magnesiothermic reduction of the as-obtained silica. Carboxylic acids, i.e. citric acid, acetic acid, and lactic acid, were used as the gelling agents. The  $SiO<sub>2</sub>$  and Si were characterized thoroughly to ensure the end-product quality. Based on our findings, the asobtained  $SiO<sub>2</sub>$  samples are in an amorphous phase, while the as-obtained Si micro-powder has a highly crystalline structure. Based on the SEM-EDX analysis, the Si samples have high purity. In a full-cell LIB application, the pure Si anode coupled with  $LiNi<sub>0.6</sub>Co<sub>0.2</sub>Mn<sub>0.2</sub>O<sub>2</sub>$  (NCM) cathode delivered poor electrochemical performance; however, when graphite was mechanically introduced Si, the performance was significantly improved. The Li-ion storage mechanism of Si/Graphite was investigated. The coal fly ash-derived Si powder can be utilized and modified to obtain Li-ion batteries with excellent electrochemical performance.

Keywords: Anode; Battery; Fly-ash; Material; Silicon, Waste

# **1. Introduction**

Renewable energy sources have been widely researched because of the increasing environmental pollution and the rising number of energy needs as the number of people in the world grows. New and renewable energy generation largely depends on energy storage systems. Energy storage has been widely researched, especially in Lithium-Ion Batteries. The lithium-ion battery (LIB) was chosen because it is the most preferred power source for portable/wireless electronics, mobile applications, power tools, and energy storage systems. This is because LIBs have a distinctive high voltage quality, which results in considerable energy density and long cycle<sup>1)</sup>. The anode is an essential component in lithium-ion batteries. The anode that is used is graphite. Graphite was chosen because it is relatively cheap and easy to make electrodes. On the other hand, it has a considerably low of  $\sim$ 378 mAh/g (vs. Li/Li+), graphite was considered to be unable to meet the quality and increasing demands for high density, high safety, and long cycle-life energy storage; thus, new anode materials were  $developed<sup>2–5</sup>$ .

The silicon-based anode is expected to be the state-ofthe-art anode active material for the next generation of high capacity and energy density LIBs owing to a theoretical capacity of ~4200 mAh/g. Unfortunately, using Si anode material has several drawbacks, such as low electrical conductivity. During the insertion and deinsertion of the Li-ion process, Si material experiences significant volume expansion (>300%). This expansion can cause material passivation, mainly from deformation and isolation from the main electrode phenomena. Another study proposed the utilization of reduced  $SiO<sub>2</sub>$  or silicon suboxide (SiOx) to elevate the structural stability of Si-based anode. In fact,  $SiO_x$  (x < 2) has a lower expected storage capacity of 1961 mAh/g than Si, but

 $SiO<sub>x</sub>$  has less volume expansion, lower cost, and is environmentally friendly<sup>6,7)</sup>. However, implementing  $SiO<sub>x</sub>$  anode material also has disadvantages, such as relatively low conductivity and high irreversible coulombic efficiency during the initial cycle. Prelithiation of SiOx can be performed to improve the performance. However, cost-wise, this approach could be more beneficial<sup>8)</sup>

To successfully apply Si as an anode material for the commercialization of high-capacity Li-ion batteries, the focus is not only on its performance-enhancing approach but also on the sustainability of the Si. Many efforts have been developed to obtain Si with excellent electrochemical performance. Still, they also often sacrifice the industrialization aspect, such as expensive and rare Si precursors complex, non-scalable, and energyintensive processes<sup>9,10)</sup>. By this technique, the Si material is no longer attractive for the sustainability of energy storage. Developing high-quality and high-purity Si material from cheap and abundant resources using a simple and facile production process is necessary, considering the economic and environmental aspects.

Due to today's reliability in coal-powered energy generation, coal-derived fly ash is continuously generated in large amounts. With the increasing energy demand, the coal exploitation rate has significantly increased. Aside from its carbon footprint, fly ash waste is also potentially harmful to living organisms and the environment. Due to the significant content of  $SiO<sub>2</sub>$ , the fly ash can be utilized as the silica precursor during the production of silicon anode material<sup>11,12</sup>).

In this report, high-purity silicon was produced from coal fly ash (CFA) obtained from the Indonesian Power Plant. Si is the dominant element in the CFA; however, further purification steps are required to obtain highquality silicon powder: (i) caustic extraction of silica, (ii) organic acid-assisted precipitation, (iii) magnesiothermic reduction, and (iv) Si purification by acid leaching. The method does not use expensive chemicals and additives. The processing of CFA significantly improves its economic value and reduces its hazardous level. Carboxylic acids such as citric acid, acetic acid, and lactic acid are used as the gelling agent instead of mineral acids, further improving the safety and the eco-friendliness of the overall process<sup>13,14)</sup>. The electrochemical performance of Si is investigated, especially in full Li-ion cells, which is rarely performed in research papers. The overall process can be considered highly efficient, scalable, cost-effective, and straightforward, and it can be directly applied to commercializing high-capacity LIBs.

# **2. Material and Methods**

#### **2.1. Materials**

The CFA waste was obtained from the Paiton Power Plant, Probolinggo, Indonesia. NaOH flakes with 95% purity were obtained from PT Asahimas Chemical, Indonesia. Glacial acetic acid (Merck, Germany), 85% lactic acid (Merck, Germany), and citric acid monohydrate (Merck, Germany) were used as the gelling agent. Magnesium powder (Merck, Germany) was used as the reducing agent for  $SiO<sub>2</sub>$ . All of these materials were used without any purification step.

### **2.2 Methodology**

#### **2.2.1 Preparation of Silica (SiO2) Xerogel**

SiO2 content in CFA waste was extracted using a 4 M sodium hydroxide solution. The liquid per solid (L/S) ratio of coal extraction is 20 mL/g. The extraction process was performed in a 2 L beaker under vigorous stirring of 500 rpm at 60  $\degree$ C for 4 hours. The resulting suspension was aged for 24 hours at room temperature. The residue from extraction was precipitated, and the supernatant was easily obtained via decantation. A demineralized water diluted the supernatant until the volume reached 2 L. This cloudy solution was labeled sodium meta-silicate solution and then separated into three 1-L beakers with identical volumes. 1 M of acetic acid (AA), lactic acid (LA), and citric acid (CA) were introduced to each respective beaker until the pH of the final solution was 7-8. During the reaction, the solutions were mixed at 300 rpm. After the desired pH is achieved, the white gel is formed. The pH was kept by adding acid dropwise for 1 hour. Next, the gel was aged for a day, filtered, and washed several times using deionized water and one time using ethanol to obtain the  $SiO<sub>2</sub>$  xerogels. The xerogels were vacuum-dried. The as-obtained  $SiO<sub>2</sub>$  were labeled according to the gelling agent, i.e.,  $SiO<sub>2</sub> - AA$ ,  $SiO<sub>2</sub> - LA$ , and  $SiO<sub>2</sub> - CA$ .

#### **2.2.2. Preparation of Silicon (Si) Powder**

The as-prepared  $SiO<sub>2</sub>$  powders were ground and sieved using a 200-mesh screen. Each  $SiO<sub>2</sub>$  was mixed with magnesium (Mg) powder with a mass ratio of 4:5 or a molar ratio of 1:2. The solid-state mixing at lab-scale synthesis was performed using mortar and pestle until homogenous. In an alumina crucible, the mixture was fired in a box furnace at 750 ℃ for 2 hours under steady N2 flow (PT. Samator Indonesia). After being cooled down, the as-obtained powders were dissolved in their respective acids with an L/S ratio of 50. During the leaching process, effervescence occurred. The leaching was stopped until the bubbles no longer appeared. The suspension was aged for 1 hour, and then the solids were separated using a paper filter. The solid was washed several times and vacuum dried at 90 ℃. The dried powder samples were ground and sieved using a 200 mesh screen and labeled according to the gelling and leaching agent, i.e., Si-AA, Si-LA, Si-CA, and Si-MA.

#### **2.2.3. Material Characterization**

To assure the quality of the  $SiO<sub>2</sub>$  and Si material, each sample characteristic was deeply investigated. FTIR spectroscope (IR-Spirit, Shimadzu, Japan) was used to analyze the surface functional group of the samples. The

structural properties and the purity of the samples were evaluated using XRD (MD10, Mini-diffractometer, MTI, USA). The morphological feature and element composition of the samples were investigated using a Scanning Electron Microscope- Energy Dispersive X-ray or SEM-EDX (JCM7000, Jeol, Japan). Further morphological characteristic analyses, such as surface area and pore dimension evaluation, were performed using the  $N_2$  sorption isotherms technique at 77.35 K using a surface area analyzer (Quantachrome, USA). The thermogravimetric analysis was performed using a DTG-60 Thermogravimetric Analyzer (Shimadzu, Japan) under air atmosphere.

#### **2.2.4. LiBs Cells Assembly and Testing**

The silicon-based materials were applied as an anode material for Li-ion full batteries. Pure Si and 50% Si/G composite were separately coated on each side of a copper foil (MTI, USA). In the full-cell analysis,  $LiNi<sub>0.6</sub>Co<sub>0.2</sub>Mn<sub>0.2</sub>O<sub>2</sub>$  or NCM was used as the counter cathode material. The electrode formulae and the controlled mass loading can be seen in Table 1. The overall assembly processes are similar to those of our previous study<sup>11,15)</sup>. The fabricated cells were stored in a vacuum oven before the electrolyte filling (1M LiPF6 dissolved in ethylene carbonate-diethyl carbonate solvent). The cells were aged for a day before the electrochemical testing. The formation of the cells was performed using an Arbin Battery Analyzer (Arbin Instruments, USA) at a current density of 0.1 C (1C = 200 mA/gNCM) or 0.5 mA/cm2 and a voltage window of 2.5-4.3 V. The cycle test was performed at 0.2 C for 50 times.





# **3. Result and Discussion**

To obtain the silica from fly ash, it is necessary to investigate the characteristics of the CFA waste. Figure 1 (a) displays fly ash samples' TG/DTA curve. The CFA has a low mass loss of less than 0.2% as the temperature increases. This shows that the sample has a low loss of ignition value and low fixed carbon content. SEM image of the CFA depicted in Fig. 1(b) presents a micron-size particle with a near-perfect spherical shape. This phenomenon is consistent with previous studies, which also have reported a micron-sized spherical particle of coal-fly ash  $^{16,17}$ . The composition of the coal-fly ash as EDX analysis results is listed in Table 2. Based on the result, Si, Al, Mg, Ca, and Fe have the highest composition based on weight or mol. The low content of Ca confirms that the fly ash is type F, which is effective for alkalinebased reactions or extraction<sup>16,18</sup>).

Table 2. The EDX analysis result of coal-fly ash

<b>Elements</b>	Composition $(wt\%)$ (%)	Composition $(mol\%)$
O	47.50	63.67
Mg	4.32	3.81
A <sub>1</sub>	12.90	10.25
Si	17.34	13.24
Ca	14.19	7.59
Fe	3.74	1.44
Sigma	100.00	100.00





**Fig. 1:** (a) TG/DTA curve and (b) SEM Image of CFA

From XRD patterns presented in Fig. 2. a,  $SiO<sub>2</sub> AA$ ,  $SiO<sub>2</sub>-LA$ , and  $SiO<sub>2</sub>-CA$  samples have a widening peak detected at a diffraction angle of 15–30◦; this explains that

three samples show an amorphous phase. In the three sample variations, no impurities were detected. In addition, all samples are appropriately indexed and are compliant with  $SiO<sub>2</sub>$  reference ICSD Card no. 061662<sup>19,20</sup>). These results prove that the variation in the use of different acid additions in the sample has no significant effect on the SiO<sub>2</sub> structure. FTIR spectra of SiO<sub>2</sub>-AA, SiO<sub>2</sub>-LA, and  $SiO<sub>2</sub>-CA$  samples are shown in Fig. 2(b). As seen in the figure, the absorption at wavenumber  $1110 \text{ cm}^{-1}$ , 950 cm<sup>-1</sup>  $1$ , and 790 cm<sup>-1</sup> show Si-O-Si, Si-OH, and O-Si-O stretch vibration, respectively. Of all the types of vibrational bands formed from the sample, it is typically amorphous  $SiO<sub>2</sub><sup>21,22</sup>$ . Figure 2(c) presents the TG/DTA curve of the  $SiO<sub>2</sub>$  sample obtained using acetic acid, lactic acid, and citric acid. The TG curve confirms that there is a significant loss of mass of 13-15%. Based on the DTA analysis, the mass loss at less than 200 ℃ can be assigned to the endothermic evaporation of trapped water molecules in  $SiO<sub>2</sub>$  samples. ]The following mass loss can be attributed to the exothermic decomposition of a residual organic compound, i.e., ethanol. Overall, the analysis confirmed that each  $SiO<sub>2</sub>$  sample has hydrophilicity, which attracts moisture from the atmosphere. To maintain the quality of  $SiO<sub>2</sub>$ , the  $SiO<sub>2</sub>$ should be stored in a vacuum oven before the solid-state reduction reaction.





**Fig. 2:** (a) X-ray diffractogram, (b) FTIR-Spectra, and (c) TG/DTA of  $SiO<sub>2</sub>$  obtained using various acids

SEM Images of  $SiO<sub>2</sub>$  samples can be seen in Fig. 3. The images confirm the presence of micron-sized silica secondary particles that consist of submicron-sized primary particles. The formation of secondary particles might be affected by the Ostwald ripening effects. The primary particles formed through the densification of Si-OH nuclei, which increases as the reaction time is prolonged. The mapping of  $SiO<sub>2</sub>$  confirms the silicon dioxide component of the sample with no observable impurities.  $SiO<sub>2</sub>$  prepared using citric acid produces denser particles with smoother surfaces compared to the model prepared using acetic acid and lactic acid, in which the adherents of small particles are observed. Based on previous studies, gelation of  $SiO<sub>2</sub>$  by organic acids such as HCl,  $H_2SO_4$ , or  $HNO_3$  often results in agglomerated submicron particles into polymorphs of secondary SiO<sub>2</sub>. Unique-shaped  $SiO<sub>2</sub>$  particles such as spherical, nano-rods, nano-tubes, and nano-flowers are usually prepared using organo-silicon precursors such as tetra-ortho silicate (TEOS) or using advanced and complex procedures<sup>12,23,24)</sup>. The formation of uniquely shaped  $SiO<sub>2</sub>$  from silicacontaining waste is an interesting topic that will be developed in the future.



Fig. 3: SEM images and EDX Mappings of SiO<sub>2</sub> obtained using various acids: (a) acetic acid, (b) lactic acid, and (c) citric acid

Figure 4 shows an XRD analysis of silicon with various variations in acid addition. Silicon peaks formed on the curve are by the reference. Therefore, the magnesiothermic reduction of amorphous silica into crystalline silicon (cSi) is successfully conducted. In addition, silicon purification is also successful due to the dominant peaks of silicon in the diffractogram. However, there is a slight impurity phase assigned to magnesium silicate  $(MgSiO<sub>3</sub>)$  in all of the samples, which is a side reaction during the magnesiothermic reduction. The crystalline silicon has a cubic diamond crystal structure with the dominant lattice point at the peak (111) at a diffraction angle of 28◦ of 2 theta formed in all three variations<sup>25)</sup>. Overall, this shows that adding various types of acids also has no significant effect on the silicon structure. As seen in Fig. 4(b), the FT-IR spectrum of Si-CA, Si-LA, and Si-AA has formed peaks at wavenumber 860 cm-1 and 915 cm-1, which can be assigned to vibrations of silicon. In addition, the Si-O-Si stretching mode also found peaks at a wave number of  $1070 \text{ cm}^{-1}$ , indicating oxygen's presence from an incomplete reduction process<sup>26)</sup>. The oxygen appearance on the material's surface is considered usual since, in a previous study, there were additional washing steps using HF. Nevertheless, in this study, we decided to avoid harmful  $HF^{14,27,28}$ .





Fig. 4: (a) XRD of SiO<sub>2</sub> obtained using various acids: (a) acetic acid, (b) lactic acid, and (c) citric acid and (b) FTIR-Spectra of SiO2 obtained using various acids: (a) acetic acid, (b) lactic acid, and (c) citric acid

Figure 5(a) displays the SEM-EDX analysis of the asprepared silicon powders from the  $SiO<sub>2</sub>$  samples. Inhomogeneous-shaped micro-sized particles with rough surfaces can be observed for all silicon samples. The Si-AA and Si-LA prepared from  $SiO_2$ -AA and  $SiO_2$ -LA, respectively, have sub-micron-sized primary particles that can be seen. Meanwhile, the Si-CA sample has a denser and smoother morphology than the others. This proves that the magnesiothermic reduction process does not significantly alter the particle of  $SiO<sub>2</sub>$  precursor<sup>29</sup>. The presence of small oxygen in the EDX-Map of Si samples also proved that magnesium silicate exists on the surface of the particle. This was confirmed by the quantitative analysis using EDX, which is presented in Table 3. Si-AA exhibits high O and Mg content, which is consistent with the XRD result. The presence of magnesium silicate can be neglected. Based on these findings, the Si preparation can be summarized in the equations below:

 $SiO<sub>2</sub> + 2Mg \rightarrow Si \cdot MgO$  (1)<br>  $Si \cdot MgO + 2HA \rightarrow Si + MgA + H<sub>2</sub>O$  (2)  $Si \cdot MgO + 2HA \rightarrow Si + MgA + H<sub>2</sub>O$ 

Based on the reactions, the organic acid reacted with

MgO and yielded soluble Mg-salt. We expect the reaction of AA with MgO to be slower due to the high pKa value of AA, causing the residual MgO in the sample. Meanwhile, LA and CA have higher pKa, which means stronger acidity compared with AA. Meanwhile, the pKa of CA is larger than the pKa of LA. Based on our data, the Mg content in Si-LA is lower. We predict that at the same concentration, the CA solution has higher viscosity due to the large molecular weight of CA. The viscous CA solution reduces the mass transfer during the reaction, which lowers the overall reaction kinetics.

The surface area analysis result shown in Fig. 5(b) is performed for LA samples. The surface area calculated using the BET method for  $SiO_2$  - LA is 106.97 m<sup>2</sup>/g with 8.5 nm, while after being converted into silicon powder, the surface area of  $Si - LA$  is reduced to 28.7 m<sup>2</sup>/g with a pore diameter of 4.1 nm. Therefore, both  $SiO_2$  - LA and Si – LA can be considered mesoporous with only a difference in the surface area value. This shows that during the magnesiothermic reduction, the particles

become denser, which is followed by a reduction in surface area. However, the densification also promotes pore shrinkage, which can be beneficial for energy storage applications. The adsorption and desorption of  $N_2$  curves of SiO2 and Si samples have type-IV shape curves, which confirm the presence of mesoporous materials $28,30$ .

Table 3. EDX Result of Si samples

Element	$Si-AA (mol%)$	$Si-LA(mol\%)$	$Si-CA(mol\%)$
Si	67.74	85.57	79.23
O	23.36	11.98	17.21
Mg	9.1	2.45	3.56



**Fig. 5: (a)** SEM images of SiO2 obtained using various acids and (b)  $N_2$  isotherm adsorption-desorption analysis of  $SiO_2$  – LA and Si – LA

The charge-discharge analysis is conducted to evaluate the electrochemical performance of the as-prepared samples. Initially, we performed an analysis for pure samples. The charge-discharge and dQ/dV curves can be seen in Fig. 6(a) and (b), as we can see that the as-prepared Si material is able to operate similarly to commercial anodes such as graphite and other carbon-based materials. In theory, the commercial graphite has a lithiation and delithiation potential of 0.07-0.19 V and 0.1-0.23 V, respectively. Meanwhile, the lithiation and delithiation potential of Si is 0.05-0.21 and 0.31-0.47 V, respectively, almost similar to graphite. Therefore, the charge and discharge curve of Si-anode in LIBs full cell will be similar to a cell with graphite anode. When NCM622 is the cathode, the material is expected to operate at a voltage window of 2.5-4.3  $V^{31}$ . The charge capacity is 3.7 Ah/g, about ten times larger than graphite's. However, massive capacity loss occurs during discharge with a deliverable capacity of 1.17 Ah/g. The initial coulombic efficiency (ICE) of pure  $Si - LA$  is 31.6%, which is considered very poor. This phenomenon can be assigned to the electrochemical property of Si material, which often experiences expansion during the charging process. Based on the previous report, the expansion of Si material can achieve up to  $270\%$  from its initial volume<sup>31,32)</sup>. The expansion can have a major effect on the battery system, such as pulverization of silicon during the shrinking process, which causes contact loss or isolation of material from the current collector or loss of electrical contact. This can cause significant capacity loss during cycling. Therefore, blending Si powder with commercial graphite can overcome this issue. The dQ/dV curve also confirms major phase alteration of NCM cathode material, which is also responsible for a capacity drop during the initial  $cycle^{10,33}$ .



**Fig. 6:** Full Cells analysis of the as-prepared Si powders using Arbin Battery Analyzer: (A) Pure Si – LA chargedischarge analysis, (B) dQ/dV of pure Si – LA

To improve the ICE level of Si-based material, the asprepared Si samples are composited with commercial graphite with a 50:50 composition. The composites were mixed using mortar and pestle for 30 minutes. The chargedischarge analysis is depicted in Fig. 7. The chargedischarge capacity of Si – AA, Si – LA, and Si - CA are 1.64 Ah/g /1.20 Ah/g, 1.71 Ah/g /1.22 Ah/g, and 1.67 Ah/g /1.15 Ah/g, respectively. Thus, the ICE of Si – AA, Si – LA, and Si – CA are 73.1%, 71.3%, and 69.2%, respectively. Si-LA achieves the highest discharge capacity; however, there are no significant differences between each sample. Based on the ICE, the discharge capacity and the dQ/dV value show significant improvements compared to the cell with pure Si powder as an anode. The effect of graphite in the composite is predicted to give better structural stability and compensate for the significant volume expansion of  $Si<sup>34,35</sup>$ . The dQ/dV vs. V curve confirms initial capacity loss due to the phase transformation of NCM cathode material $36-38$ .



**Fig. 7:** Full Cells analysis using Arbin Battery Analyzer of (a-c) Si-AA/G, (d-f) Si-CA/G, and (g-h) Si-LA/G

The cycle performance of each cell is depicted in Fig. 8(a-c). The charge and discharge current were 0.2 C/ 0.2C (1C=200 mA/g). After 50 cycles, the capacity retention of Si-AA, Si-LA, and Si-CA are 20.11%, 24.92%, and 23.04% (Fig.8d), respectively. This means that the 50 w% Si is still large enough to be present as an anode, resulting in similar behavior to pure Si only at a prolonged cycle. However, the average coulombic efficiency is above 95%. The optimization of Si content in the composite is recommended to be investigated in future research.



**Fig. 8:** Cycle Performance of (a) Si-AA/G, (b) Si-CA/G, and (c) Si-LA/G, and (d) Capacity Retention of Si-AA/G, Si-CA/G, and Si-LA/G

# **4. Conclusion**

In this study, the synthesis of crystalline silicon (cSi) from alkaline extraction of silica followed by magnesiothermic reduction of F-type coal fly ash derivedsilica is successfully performed. The extracted silica as the intermediate product has an amorphous structure and high purity, even derived from coal fly ash. Organic carboxylic acids successfully precipitate the silica xerogel, which improves the Eco-friendliness of the overall process. Based on the study, the carboxylic acids, i.e., acetic acid, lactic acid, and citric acid, do not significantly influence the structural properties of both  $SiO<sub>2</sub>$  and the final Si product. However, the particle morphology obtained using citric acid has the densest morphology and smooth surface. The EDX mapping of samples proved the dominance of the crystalline silicon (cSi) material. The as-prepared silicon powders were applied as the anode material of the NCM battery and could operate at a voltage range of 2.5- 4.3 V based on the charge-discharge analysis with poor electrochemical performance. However, by compositing Si and commercial graphite (G), the cell achieved the highest discharge capacity of 1.22 Ah/g<sub>Si-LA/G</sub> with an initial coulombic efficiency of 71.3%. In the dQ/dV curve, a phase transformation of the NCM cathode can be detected as the reason for low coulombic efficiency. In the end, the study promises to be considered for the coal fly ash waste processing technique and polycrystalline silicon preparation.

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