

**Comparative Study of Physical and Chemical Activation of Coal Fly Ash as Potential Adsorbent Based on FTIR and XRD Analyses****Wayan Gracias^{1*}, Radho Al Kausar¹, Ermin Riskiani²**¹Department of Chemistry, Faculty of Mathematics and Natural Sciences, University of Lampung, Jl. Prof. Dr. Soemantri Brodjonegoro 1, Gedong Meneng, Bandar Lampung 35141²Department of Chemistry, Faculty of Mathematics and Natural Sciences, Universitas Gadjah Mada, Bulaksumur, Caturtunggal, Depok, Sleman, Yogyakarta 55281*Corresponding author: gracias.wayan@fmipa.unila.ac.id

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Abstrak

Coal fly ash is a combustion residue that demonstrates potential use as an affordable adsorbent. However, the adsorption capacity is still limited if no modification is made, so the activation stage is crucial. The present work investigates the differences between physical and chemical activation on changes in the structure of coal fly ash. Physical activation was carried out through a calcination process at 400 °C within 2 hours, while chemical activation was carried out through a reflux process using a 2 M HCl solution at 50 °C for 6 hours period. The samples were characterized using infrared spectroscopy (FTIR) and X-ray diffraction (XRD) techniques. The obtained results clearly exhibit that physical activation did not cause significant structural changes. In contrast, chemical activation induces partial dealumination in the aluminosilicate framework, which is characterized by a shift in the silanol (Si-O) band from 995 cm⁻¹ to 1057 cm⁻¹ in the FTIR spectrum, as well as a decrease in mullite intensity and an increase in quartz dominance based on the XRD diffractogram as indicated by the proportion of quartz relative intensity to mullite from 1,000 in CFA to 1,099 and 1,505 respectively in CFAP and CFAC. These changes result in a more silica-rich surface with a higher density of silanol groups, thereby increasing the number of active sites for the adsorption process.

Keywords: *Coal fly ash, Physical activation, Chemical activation, Structural modification, Potential adsorbent***1. Introduction**

The sustainability of Indonesia's electricity supply remains heavily reliant on coal, which is the primary fossil fuel. This commodity plays a crucial role, given its role as the primary driver of coal-fired power plant (PLTU) infrastructure. According to official data from the Indonesian Ministry of Energy and Mineral Resources (ESDM), total domestic coal supply is expected to escalate to 790 million tons by 2025, with exports accounting for 65.1% and domestic consumption accounting for 32%, largely absorbed by the electricity sector [1]. Coal-based energy production activities inherently produce considerable amounts of solid waste materials known as fly ash. Coal fly ash (CFA) is a combustion residue composed of fine particles transported along the flue gas stream and subsequently captured using emission control systems. The massive accumulation of this material poses a serious threat to environmental quality if not managed responsibly, as it contains heavy metals that have the potential to leach into the soil and contaminate water bodies through leaching [2].

In recent years, the increasing generation of industrial solid waste has become a major ecological concern worldwide, primarily in developing countries with increasing energy demands [3]. Among various industrial by-products, CFA has become among the most abundantly generated residues due to the continued dependence on coal combustion for electricity generation [4]. The improper disposal of CFA not merely fills up large landfill areas but also contributes to secondary environmental pollution through the release of fine particulates and hazardous inorganic constituents [5]. Therefore, sustainable CFA utilization and valorization strategies are required [6].

On the other hand, CFA has a promising mineralogical composition as an adsorbent candidate because it is rich in inorganic components, which are mainly composed of silica (SiO₂), alumina (Al₂O₃), and a number of other elements such as iron, calcium, and unburned carbon. The mineralogical structure of CFA is characterized by the presence of crystalline phases including quartz and mullite incorporated into an

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amorphous aluminosilicate matrix [7]. Despite its rich aluminosilicate content, the majority of CFA must be modified before being used as an adsorbent to open up its active sites that are still covered by impurities.

To optimize the adsorption performance of CFA, activation requires two main approaches: physical and chemical activation. Physical activation relies on thermal and mechanical treatments to promote higher surface area and enhance the porosity of the material [8], meanwhile, chemical activation utilizes acid or base solutions with the aim of dissolving the impurity phase, opening active sites, and forming functional groups that have higher reactivity towards the adsorbate. [9]. The difference in mechanism between these two approaches directly determines the extent to which the aluminosilicate framework undergoes structural modification.

CFA has been extensively studied as a material for removing heavy metals and dyes from aqueous systems. Several studies have demonstrated the success of chemically treated CFA in adsorbing various contaminants with high efficiency. Hussain [10] reported that acid-base treatment with HCl and NaOH promoted higher surface area in CFA up to three times compared to unmodified CFA, with dye removal efficiency reaching 98%. Nadeem [11] showed that acid-base activated surface enhanced coal fly ash (SECFA) achieved highly effective crystal violet adsorption with 97.52% efficiency, far exceeding unmodified CFA which only reached 81.52% under the same conditions. Eteba [12] also reported that hydrochloric acid treatment of CFA significantly reduced particle size and increased zeta potential, resulting in dye removal efficiencies of up to 99.7%. In general, these studies confirmed that chemical activation with acidic solutions, especially HCl, resulted in more effective surface modification than untreated CFA.

Despite extensive investigations of CFA in adsorption studies, comparative studies on the effects of physical and chemical activation on changes in the structure and functional groups of the surface, especially on Indonesian CFA, are still limited. [13,14]. Therefore, this study focuses on the comparison of physical activation (CFAP) and chemical activation (CFAC) through the analysis of crystal structure and functional groups using XRD and FTIR [15]. This research does not include adsorption performance testing, but rather focuses on material characterization as a basis for developing CFA-based adsorbents.

2. Research Methodology

2.1. Equipments and Materials

Equipments: This research utilized a number of laboratory equipment, including glassware, an analytical balance (Kern), a hot plate stirrer (ThermoScientific), an oven (InnotechLab), and a muffle furnace (Nabertherm). The material characterization instruments used included an X-ray diffractometer (Xpert PANalytical) and an infrared spectrophotometer (Shimadzu model FT-IR/8201 PC).

Materials: The CFA employed in the present study were derived from coal combustion residues at the Paiton Steam Power Plant (PLTU) in East Java. Supporting chemicals included hydrochloric acid (HCl, Merck), a universal indicator (Merck), and distilled water to wash the samples to a neutral pH. All reagents were used as received without additional purification.

2.2. Experimental Procedure

2.2.1 Physical Activation of CFA

Physical activation of CFA was carried out through thermal treatment using a muffle furnace. A 25 gram sample of fly ash was heated at 400°C for two hours using an activation furnace. This temperature was chosen because it is within the optimum temperature range for modifying the characteristics of CFA [16,17]. After the heating stage is complete, the material was permitted to cool down to ambient temperature in a desiccator to prevent adsorption of water vapor from the environment. Physically activated samples are designated CFAP (physically activated CFA).

2.2.2 Chemical Activation of CFA

CFA was chemically activated through a reflux process with acid treatment using 2 M HCl solution at an operating temperature of 50 °C for 6 hours with constant stirring. The resulting suspension was then filtered and rinsed with distilled water until neutral pH was achieved [18].

To remove residual moisture, the resulting material was subsequently dried at 105 °C for 6 h and then stored in an airtight condition using a desiccator. The chemically activated sample was designated as CFAC (chemically activated CFA).

2.2.3 Material Characterization

All samples of pre-activated fly ash (CFA), physically activated fly ash (CFAP) and chemically activated fly ash (CFAC) were characterized using infrared spectrophotometer (FTIR) to identify functional groups and structural changes that occurred during activation, as well as X-ray diffraction (XRD) to identify the crystalline phase of the material.

3. Results and Discussion

3.1. Identification of Functional Groups Using FTIR

To observe structural changes in coal fly ash due to physical and chemical activation treatment, characterization was carried out using FTIR. Overall, all three spectra exhibit similar absorption bands in the wavenumber regions typical for aluminosilicate materials, particularly about $\sim 3400\text{ cm}^{-1}$, $\sim 1057\text{--}995\text{ cm}^{-1}$, and $\sim 680\text{--}465\text{ cm}^{-1}$ which indicate the presence of hydroxyl groups from water, silanol or aluminum, respectively from the ash fly. The uniformity of this pattern confirms that the basic framework of aluminosilicate is preserved, despite the possibility of local changes in the surface chemical environment due to activation treatment

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[19,20]. The FTIR spectra of CFA, CFAP, and CFAC is presented in Figure 1.

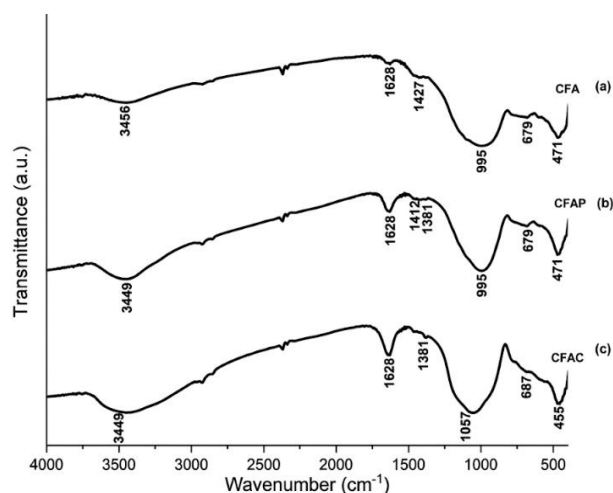


Figure 1: FTIR spectra of (a) CFA; (b) CFAP; (c) CFAC

However, pronounced differences were observed in the peak intensity and peak shifts in CFAC compared to CFA and CFAP. This difference indicates that chemical activation using HCl 2 M produced a more significant structural modification impact than simply providing thermal energy through a furnace. Table 1 shows a summary of absorption interpretations on CFA, CFAP and CFAC.

The absorption band observed in the spectral region around 3400 cm^{-1} corresponds to O–H stretching vibrations, indicating the presence of hydroxyl groups originating from free water molecules. This absorption band generally overlaps with the O–H stretching vibrations of silanol groups present on the surface of the fly ash. [11,13]. After activation, the absorption peak shifted to 3449 cm^{-1} in both CFAP and CFAC. This shift indicates that, in CFAP, furnace heating promoted the release of bound water molecules, resulting in the elongation of hydroxyl bonds and facilitating silanol condensation as well as the rearrangement of hydroxyl groups due to changes in the hydrogen-bonding pattern.

In contrast, in CFAC, treatment with HCl induced protonation of surface hydroxyl groups, making them more prone to interact with water molecules and other surrounding groups. Consequently, the O–H bonds became weaker, and the resulting spectrum broadened due to variations in hydrogen-bonding strength. [18]. The substantial increase in the absorption intensity of hydroxyl groups observed in CFAC indicates the enhancement of active sites in the form of silanol groups following acid activation [14]. The increased density of silanol groups contributes to the enhanced adsorption potential of CFAC, as Si–OH groups function as the primary active sites in surface complexation and ion-exchange mechanisms involving heavy metal cations.

Table 1: Interpretation of the infrared spectrum from CFA, CFAP and CFAC

Wavenumber (cm^{-1})			Interpretation
CFA	CFAP	CFAC	
3456	3449	3449	O–H stretching vibrations associated with water molecules and silanol groups (Si–OH) [19,21].
1628	1628	1628	H–O–H bending vibrations [22]
1427	1412	1381	Stretching vibrations of carbonate groups or C–O, C–C, and C–H bonds [20,23]
995	995	1057	Asymmetric stretching vibrations of Si–O–T bonds (T = Si or Al) [20,24]
679	679	687	Symmetric bending vibrations of Si–O–Si or Si–O–Al bonds [24]
471	471	465	Symmetric bending vibrations of Si–O–Si bonds [25]

The alteration of O–H groups is further supported by the appearance of an infrared absorption band at 1628 cm^{-1} , which corresponds to the bending vibrations of H–O–H molecules, indicating the presence of adsorbed water molecules on the surface of all CFA samples [22]. The absorption band consistently observed at 1628 cm^{-1} in all samples corresponds to the H–O–H bending vibrations of water molecules trapped on the material surface. In the CFA sample, this band was already present and remained at the same position in CFAP. However, in CFAC, the band became sharper while also exhibiting noticeable broadening. Although no shift in peak position was observed, the change in band profile indicates alterations in the condition and environment of water molecules on the surface. The increased sharpness suggests a more ordered water structure resulting from interactions with newly formed surface sites, whereas the band broadening reflects variations in the molecular environment, likely caused by modifications in the silica network structure during chemical activation [26]. The simultaneous changes observed at 3449 cm^{-1} and 1628 cm^{-1} collectively confirm that chemical activation not only increased the number of silanol groups but also fundamentally altered the interaction characteristics between water molecules and the material surface, ultimately contributing to the enhancement of adsorption active sites in CFAC.

The absorption band in the $1350\text{--}1500\text{ cm}^{-1}$ region represents the stretching vibrations of

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carbonate groups or organic constituents, including C–O, C–C, –CH₂, and –CH bonds [20,23]. In this region, significant differences in the absorption profiles were observed among the three samples. CFA exhibited a peak at 1427 cm⁻¹ attributed to the asymmetric stretching vibrations of carbonate groups and organic residues, indicating that the initial surface was still covered by minerals such as calcite [20]. Following physical activation (CFAP), the peak shifted to 1412 cm⁻¹ while the absorption intensity remained observable, indicating partial decomposition without complete elimination of impurity phases [13]. In contrast, CFAC exhibited a substantial decrease in absorption intensity accompanied by a peak shift to 1381 cm⁻¹, demonstrating that 2 M HCl was considerably more effective through an acid-leaching mechanism. Hydrochloric acid successfully dissolved free carbonates and metal oxides into water-soluble chloride salts, thereby exposing active sites that had previously been blocked. [27].

The most significant structural transformation was observed from the shift of the main peak in the 995–1057 cm⁻¹ region, which represents the asymmetric stretching vibrations of Si–O–T bonds (T = Si or Al) within the tetrahedral framework structure of fly ash [20,24]. Both CFA and CFAP exhibited absorption peaks at 995 cm⁻¹, whereas CFAC showed a blue shift to 1057 cm⁻¹. This shift indicates the occurrence of a dealumination process, in which HCl facilitated the dissolution of aluminum atoms from the aluminosilicate framework, thereby increasing the Si/Al ratio. Since the force constant of siloxane bonds (Si–O–Si) is greater than that of Si–O–Al bonds, the vibrational energy of the framework increased, resulting in a higher wavenumber. This shift also reflects a reduction in Si–O–Al linkages and a relative increase in Si–O–Si bonds, which is consistent with partial dealumination during acid treatment [14]. This finding is further supported by complementary XRD analysis results, as discussed in the following subsection.

3.2. Identification of Mineral Phases and Structural Changes Through XRD Analysis

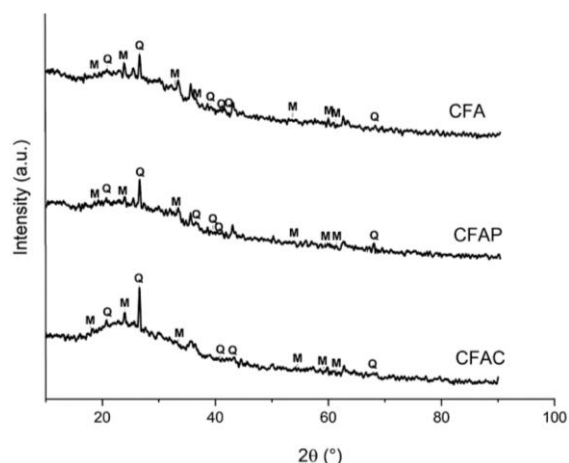


Figure 2. XRD diffractograms of CFA, CFAP, and CFAC (Note: M = Mullite, Q = quartz).

X-ray diffraction characterization was conducted to identify and compare the mineral phases present in CFA, CFAP, and CFAC, as well as to examine the structural changes induced by the activation treatments in a complementary manner to the FTIR analysis findings. The XRD diffractograms of CFA, CFAP, and CFAC are presented in Figure 2, whereas the data on 2θ peak positions, d-spacing values, and phase identification are summarized in Table 2.

Table 2. Diffraction peak positions (2θ), interplanar spacing (d-spacing), and mineral phase identification of CFA, CFAP, and CFAC.

Sample	2θ	d-spacing	Mineral Phase
CFA	18,6004	4,7704	Mullitee
	20,8323	4,2641	Quartz
	23,7047	3,7535	Mullite
	26,5387	3,3599	Quartz
	32,8356	2,7254	Mullite
	36,5917	2,4538	Mullite + Quartz
	38,9612	2,3098	Quartz
	40,1257	2,2454	Mullite
	42,1618	2,1416	Mullite
	53,4220	1,7137	Mullite
	59,6686	1,5484	Mullite
	60,5407	1,5281	Mullite
	68,0216	1,3771	Quartz
	CFAP	18,6925	4,7471
20,6571		4,2998	Quartz
23,5592		3,7764	Mullite
26,5673		3,3552	Quartz
32,9794		2,7138	Mullite
36,2120		2,4786	Quartz + Mullite
39,3239		2,2893	Quartz
40,1033		2,2466	Mullite
53,5785		1,7091	Mullite
59,6945		1,5477	Mullite
CFAC	60,7198	1,5278	Mullite
	67,8427	1,3803	Quartz
	18,5706	4,7780	Mullite
	20,7289	4,2851	Quartz
	23,9215	3,7169	Mullite
	26,4986	3,3637	Quartz
	32,8727	2,7224	Mullite
	36,4709	2,4616	Mullite + Quartz
39,3856	2,2859	Quartz	
40,0376	2,2502	Mullite	
42,2761	2,1414	Mullite	
53,7956	1,7027	Mullite	
59,8546	1,5439	Mullite	
60,7591	1,5231	Mullite	
67,8383	1,3804	Quartz	

Based on the XRD analysis results, the crystalline phases of all samples were predominantly composed of Mullite (3Al₂O₃·2SiO₂, JCPDS No. 15-0776), characterized by an orthorhombic crystal system with lattice parameters of a = 0.7543 nm, b = 0.7693 nm, c =

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0.2885 nm, and space group Pbam, as well as Quartz (SiO₂, JCPDS No. 46-1045). In addition, a broad hump observed at $2\theta = 15\text{--}35^\circ$ indicates the presence of a significant amount of amorphous aluminosilicate matrix. [7,28]. The presence of these two crystalline phases is a typical mineralogical characteristic of Class F fly ash formed from the high-temperature combustion of bituminous coal and is also consistent with the ASTM C618 classification standard [7].

The diffractogram of CFAP was nearly identical to that of CFA, with no indication of new phase formation, confirming that physical activation at 400 °C did not significantly alter the crystalline phase composition. This is attributed to the activation temperature being considerably lower than the degradation temperature of mullite (>1000 °C) and the transformation temperature of quartz to cristobalite (>870 °C) [13]. In contrast, the diffractogram of CFAC exhibited distinct changes, including an increase in the relative intensity of the quartz peak at $2\theta \approx 26.50^\circ$, which became sharper and more dominant, accompanied by a decrease in the relative intensity of mullite peaks and alterations in the characteristics of the amorphous hump, indicating modification of the aluminosilicate glass phase due to acid attack. These observations were further supported by semi-quantitative calculations based on the XRD data of all three samples, as presented in Table 3.

Table 3. Comparison of the relative intensities of XRD diffraction peaks in CFA, CFAP, and CFAC.

Parameter	CFA	CFAP	CFAC
Quartz peak 2θ (°)	26,56	26,56	26,58
Mullite peak 2θ (°)	33,26	33,22	32,86
Relative Quartz peak intensity with respect to CFA (I_Q)(%)	100,0	100,6	144,2
Relative Mullite peak intensity with respect to CFA (I_M) (%)	100,0	91,5	95,8
Quartz/Mullite intensity ratio (I_Q/I_M)	1,000	1,099	1,505
Relative ratio change with respect to CFA (%)	-	+9,9	+50,5

As presented in Table 3, the relative intensity ratio of quartz to mullite peaks (I_Q/I_M) indicates changes in the relative composition of the crystalline phases after treatment. The ratio increased from 1.000 in CFA to 1.099 in CFAP, corresponding to an increase of 9.9%. This relatively minor increase is consistent with previous studies reporting that, in fly ash, significant changes in crystalline phases and enhancement of crystallinity generally become more evident at higher activation temperatures, whereas the presence of mullite and aluminosilicate phases is associated with relatively low material reactivity [29,30].

In contrast, the CFAC sample exhibited an increase in the relative intensity ratio to 1.505, representing a 50.5% increase compared with CFA.

This result is consistent with the enhanced sharpness of the quartz peak at $2\theta \approx 26.5^\circ$ and the relative decrease in mullite peak intensity. These changes indicate that acid treatment modified the amorphous aluminosilicate matrix, causing the quartz phase to become relatively more dominant than the other phases. [31,32].

Furthermore, the increased relative dominance of the quartz phase in CFAC does not indicate the formation of new quartz, but rather results from the partial degradation of Mullite caused by selective dealumination by HCl through the reaction $\text{Al}_2\text{O}_3 + 6\text{HCl} \rightarrow 2\text{AlCl}_3 + 3\text{H}_2\text{O}$. This process produced a silica-rich amorphous surface with a higher density of silanol groups (Si-OH), which function as active adsorption sites through ion-exchange and surface-complexation mechanisms [27,33]. Chemical activation also effectively eliminated secondary impurity phases through carbonate leaching reactions ($\text{CaCO}_3 + 2\text{HCl} \rightarrow \text{CaCl}_2 + \text{H}_2\text{O} + \text{CO}_2$), thereby exposing surface active sites that had previously been blocked, which is consistent with the decrease in the carbonate peak at 1381 cm^{-1} in the FTIR spectrum of CFAC [10].

The shift in the Mullite peak position in CFAC, such as the peak at $2\theta = 36.47^\circ$ ($d = 2.4616\text{ \AA}$) compared with 36.59° ($d = 2.4538\text{ \AA}$) in CFA, indicates the occurrence of crystalline lattice relaxation due to the extraction of Al^{3+} from the Mullite framework, which was subsequently replaced by silanol groups. This observation is consistent with the blue shift of the Si-O-T peak from 995 cm^{-1} to 1057 cm^{-1} in the infrared spectrum [14]. Overall, the XRD and FTIR analyses indicate that physical activation did not significantly alter the aluminosilicate framework, whereas chemical activation induced selective dealumination, as evidenced by changes in the quartz/mullite peak intensity ratio, shifts in lattice parameters, and an increase in silanol group density. These modifications suggest that CFAC is the most promising adsorbent candidate among the three investigated samples.

4. Conclusions

This study demonstrates that chemical activation of coal fly ash produces far more fundamental structural modifications than physical activation. Physical activation at 400 °C did not induce significant changes in the crystalline phases, whereas chemical activation using 2 M HCl promoted selective dealumination of the Mullite framework, as confirmed by the increased dominance of the Quartz peak, the blue shift of the Si-O-T peak from 995 cm^{-1} to 1057 cm^{-1} , and the increased density of silanol groups. Semi-quantitative XRD calculations showed that the Quartz/Mullite intensity ratio increased from 1.000 in CFA to 1.505 in CFAC, corresponding to an increase of 50.5%, indicating changes in the crystalline phase distribution after acid treatment. Overall, the characterization results revealed that CFAC possessed a more silica-rich surface with a higher density of silanol groups, highlighting its

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potential for further development as an adsorbent material.

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