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Microstructure Investigation of Activated Carbon Prepared from Potato Peel

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Abstract

This research investigates how activated carbon (AC) was synthesized from potato peel waste (PPW). Different ACs were synthesized under the atmosphere's conditions during carbonation via two activation methods: first, chemical activation, and second, carbon dioxidephysical activation. The influence of the drying period on the preparation of the precursor and the methods of activation were investigated. The specific surface area and pore volume of the activated carbon were estimated using the Brunauer-Emmett-Teller method. The AC produced using physical activation had a surface area as high as 1210 m²/g with a pore volume of 0.37 cm³/g, whereas the chemical activation had a surface area of 1210 m²/g with a pore volume of 0.34 cm³/g. The main aim of this research is to produce activated carbon from natural materials and to prepare and characterize the elemental analysis, surface area, and morphological properties of ACs from potato peel waste using potassium hydroxide (KOH) AC-PPK and Carbon dioxide (CO₂) AC-PPC as activating agents. X-ray diffraction analysis showed the degree of crystallinity to be 35.03% in the case of AC-PPK, and AC-PPC showed a crystallinity of 35.46%. In both methods, the results showed that the crystallographic structure revealed that all the synthesized AC took on an amorphous state with low crystallinity. The atomic force microscopy (AFM) image of AC shows the presence of nanotips on the surface and shows that the maximum height was 1396 nm and 778 nm. The outer surfaces are full of cavities and highly irregular as a result of activation. The morphological analysis of the precursors was determined by scanning electron microscopy. The external surfaces are full of cavities and quite irregular as a result of activation. Also, activated carbon prepared from potato peel waste is a low-cost and effective adsorbent when compared with several activated carbon sources.

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1. Introduction

Activated carbon adsorbents have found numerous and diverse applications in recent times, frequently as a result

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of both a well-created polydisperse permeable structure with a measure of dispersion favorable for atomic sorption and a large inside surface region [1]. Any carbonaceous solid, natural or synthetic, is a carbon precursor [2]. In most cases, waste material is used as a precursor due to its availability, cheap price, and continuous need for recycling waste [3]. Examples of carbon precursors are; agricultural residues, forestry wastes, sewage sludge, used tires, and waste plastics [4]. The carbon precursor consists of (i) graphitic fabric (carbon), which is the main portion of the precursor, and (ii) impurities, which are non-graphitic fabrics such as inorganic matter (salts) [5]. As a result, it is critical to select appropriate precursor materials with high density and rising hardness, but the low inorganic structure, to produce a kindly AC [6]. Non-porous materials exist within the inorganic material structures; therefore, the adsorption ability will be reduced, in amount and accessibility. The cost of crude materials influences the ultimate price fetched. Particularly with significant masslessness in all activation treatments [7]. So, materials with a high carbon content are favorable for the generation of enacted carbon, as these materials don't require costly forms to deliver the enacted carbon [8]. Generally, brown wood, cellulosic materials, and a little bit of polymer are used as tall carbon roots to produce AC materials. [9]. Because of its accessibility and high content of cellulose, lignin, and hemicellulose, wood is the most commonly used fabric in actuated carbon generation. [10]. There has been surprising intrigue within the generation of AC due to its absorptive, natural, warm, electrical, and mechanical attributes [11]. Previous research on features such as porous structure and adsorption gain ability discovered that AC has a mechanical material structure [12]. As of now, there have been numerous studies on the auxiliary properties of active carbon. Ahmed I. Osman [13] studied activated carbon synthesized from potato peel waste (PPW) using XRD analysis and SEM, and the diffraction lines that refer to crystalline and amorphous cellulose phases constructed the precursor PPW. The fabric's surface region, pore volume, and included esteem were compared with the parameters of the precursor potato peel. The use of potato peels as feedstock not only results in a run of unused enacted carbon materials, but it may also be risky in terms of accessibility and feasibility [14]. The goal of this study is to plan and characterize the basic investigation, surface region, and morphological properties of ACs derived from potato peel waste (PPW). Chemical activation involves pyrolysis. However, in this case, the biomass is first activated with chemical agents such as H3PO4, ZnCl₂, or KOH [15]. These are the most common activators. ZnCl₂ is considered to be the most expensive activating agent, while the cheapest activating agent is KOH, and it has been extensively used in this investigation [13]. The pyrolysis of the activated biomass then occurs at much lower temperatures than that of the physical activation, typically at temperatures of approximately 500 °C [16]. Physical activations involve pyrolysis of the source material at 600°-800°C to produce charcoal [17]. Analysts detailed the impact of interfacial structure on the interfacial properties between actuated carbon fillers and the natural matrix of composites. Regardless, the high surface region and porosity are required for activated carbon [18].

2. Experimental Procedure

2.1 Preparing Precursor

Potato peel waste (PPW) has been used as the carbon precursor in this study. They do a preparatory washing well several times before the process of obtaining potato peels to remove mud or impurities using distilled water. The precursor sample dried in the environment at around 100 5 °C, as shown in Table 1. The drying was carried out during different periods to study the effect of this on the spectroscopic PPW. Where Wo and W it indicated, they indicate the weight before and after the drying process, respectively.

Samples W_{o} W **Dried Time** PPW1 52.7g 12 Hr 11.6g PPW2 51.3g 10.4g 24 Hr PPW3 52.1g 8.5g48 Hr

Table 1: Potato peel waste precursors weights before and after different drying periods.

2.2 Synthesis of Activated Carbons

Herein, activated carbons from PPW have been synthesized by chemical activation with KOH (86%), obtained from Riedel-de Haën AG, and physical activation with CO2. The washing process for these resulting carbons to remove the residuals of reactants and inorganic matter (ash) from the precursor makes the processing time- and energy-consuming, expensive, and environmentally unfriendly [18]. On the other hand, physical activation

presents some disadvantages: the ACs are obtained in two steps, with higher temperatures of activation and poorer control of the porosity [19].

2.3 Chemical Activation

Chemical Impregnation Step: The chemical agent KOH was used for impregnating potato peel waste (PPW). This is done by mixing PPWs, as shown in Table 1, with potassium hydroxide and deionized water at a ratio of 30%, as shown in the table below. At a temperature of 100 °C for a period of one hour, the blend was stirred and left at room temperature for 24 hours to dry, as shown in Table 2. The samples were assigned as PPK

Sample	PPW	КОН	Deionized water	Time Impregnation
PPK1	1.12g	0.338g	5ml	24Hr
PPK2	1.11g	0.320g	5ml	24Hr
PPK3	1 12σ	0 389σ	5ml	24Hr

Table 2: Shown the impregnation PPW in KOH.

Activation has been achieved by heating the PPKs in an oven (MTI Corp.) at around 500 ± 5 °C (with a heating rate of 5 °C per minute for 30 minutes). Then the samples have been left to cool at room temperature, during the activation phase, impurities may be generated and must be removed by washing with deionized water and HCL with a ratio of 1:1 several times to get rid of any traces of activation and to neutralize the pH. The clean activated carbon was dried at 100 ± 5 °C for 24 hours before being saved in containers for further study, as shown in Figure 1.



Figure 1: PPK Chemical Activation Samples.

2.4 Physical Activation

Physical activation is a two-step process in which the Potato Peel Waste (PPW) is first carbonized at 600 °C, followed by activation of the resulting char with oxidant gases such as CO₂. The samples were assigned as PPC in Table 3.

Samples	PPC	GAS	Temp	Time
PPC1	1.10g	CO_2	600°C	30min
PPC2	1.09g	CO_2	600°C	60min
PPC3	1.06g	CO_2	600°C	90min

Table 3: The physical activation samples of PPC.

Carbonization step; in this study, precursor PPWs were used for carbonization experiments in the environment's atmosphere, as shown in Table 1. Table 3 shows the ratio of PPCs after the carbonization process.

The 80 cm quartz tube with a 40 mm diameter is placed horizontally in the furnace with a CO_2 (1 mL/min) flow [19]. A small crucible (10 mm in diameter) was used and inserted into the tube at a temperature of 600 ± 5 °C

with a heating rate of 5 °C per minute to facilitate the handling of the material (powder) after the activation process. Then the PPCs were left to cool at room temperature and saved in containers for study, as shown below in Figure 2.



Figure 2: The PPW samples coal under a 600 °C tube furnace.

2.5 Material Characterization

The X-ray diffraction (XRD) technique has been used to determine the crystallographic structure of PPW, PPK, and PPC. The analysis was carried out using XRD SSC with a copper K- X-ray tube (wavelength @ 0.15406) and a power source of 30KV/20mA. The data on diffracted radiation were obtained using a variable angle (2) range of 10 to 80. The degree of crystallinity of the precursor was estimated as follows [20]:

$$\% X_C = \frac{A_C}{(A_C + A_a)} X 100 \tag{1}$$

Where: % Xc refers to a degree of crystallinity, Ac refers to the crystallized area on the X-ray diffractogram; and Aa refers to the amorphous area on the X-ray diffractogram.

AFM estimations of arranged samples of potato peels for prepared actuated carbon with various connected conditions were performed with an AFM (NaioAFM Naosurf Switzerland) to get a clear quantitative picture of grain size and surface harshness for precursors PPWs, PPKs, and PPCs. The morphology studies of the precursors were conducted using the ultra-high resolution Field Emission Scanning Electron Microscope (FE-SEM) images collected on the ARYA Electron Optic MIRA3 FE-SEM Microscope. The precursors were mounted onto an SEM holder with double-sided electrically-conducting carbon adhesive tabs to block the surface of the specimens when exposed to the electron beam. The chambers use a 5-axis, fully motorized, compucentric stage that provides superior specimen handling and has an ideal geometry with ultra-high resolution at low beam energy: 1 nm at 1 keV and 0.7 nm at 15 keV.

3. Results and Discussion

3.1 Characterization of Precursor

As shown in Figure 3, the Potato Peel Waste (PPW) was obtained from a local market throughout the experiments. The crystallographic structure, grain size, surface harshness, and morphological and elemental information of the potato peel were explored through AFM, XRD, and FE-SEM, analysis to prepare activation carbon.



Figure 3: Prepare activation carbon precursor.

The X-ray diffraction patterns (XRD) of the three precursors (PPW1, PPW2, and PPW3), the product of activation carbon AC using the chemical process (PPK1, PPK2, and PPK3), and via physical activation (PPC1, PPC2, and PPC3), were recorded as shown in Figures 4-6 to see the nanoparticle measure and crystalline and undefined materials. The presence of a broad diffraction foundation and the absence of peaks for the precursors The influence of the drying period on the crystallographic structure The influence of the drying period on the crystallographic structure was revealed as the amorphous state with low crystallinity for the synthesized AC (PPK and PPC), as shown in Figures 5–6.

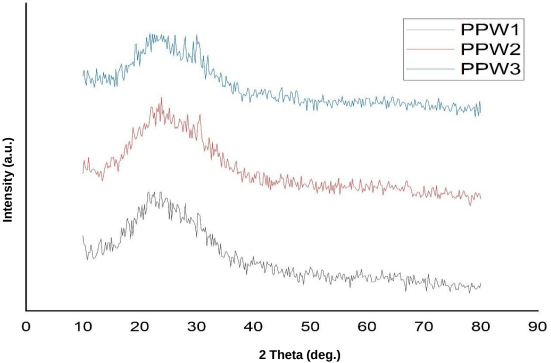


Figure 4: XRD patterns for PPW precursors.

Analysts who worked on X-ray diffractogram investigations of actuated carbons concluded that overwhelmingly amorphous solids have an expansive inner surface zone and pore volume. The effects of activation time on pore development, specific surface area, and pore volume of the prepared activated carbon were evaluated through Brunauer–Emmett–Teller (BET) and N2 adsorption testing measured using Micromeritics TriStar II Plus (USA). Table 4 shows the BET surface area of activated carbon prepared from potato peel waste for AC precursors and products of both methods. In the present study, using KOH and CO₂, most surface areas were greater than 800 m2/g, which is comparable to commercial activated carbon [21].

Table 4: Characteristics of potato peel waste (PPW) along with the ACs (PPCs and PPKs).

Samples		PPW1	PPW2	PPW3	PPC1	PPC2	PPC3	PPK1	PPK2	PPK3
BET	SBET(m ² /g)	1.8	2.4	775	775	1210	886	544	966	665
results	Pore volume (cm ³ /g)	0.019	0.041	0.22	0.22	0.37	0.26	0.2	0.34	0.29

In previous studies, the pore development process through the activation process was analyzed. At a temperature of 1000°C, it was found to be 30.4% by XRD analysis [22]. In the present study, changes in the structural parameters (based on the XRD results) during the activation process were analyzed within the same stage classification as used in the previous study. Table 5 shows the degree of crystallinity within the enacted carbon powder. Figures 5-6 show that the AC-PPK with a KOH impregnation period of 24 h has a percent crystallinity is 35.03%, and the amorphous is 64.97% for PPK2, while the AC-PPC with a CO₂ activation period of 1 hour has a crystallinity of 35.46% with amorphous 64.54%. The reason that such an immoderate activation led to a decrease in the particle size and an increase in the degree of crystallinity was considered to be due to the

movement of the carbon atoms and the reordering of the lattice layers. In all other cases, AC treated with KOHCO2 appeared to have less crystallinity, which implies that the tests are semi-crystalline or undefined. In the case of enacted carbon, a few analysts detailed enacted carbon as being a shapeless material. Enacted carbon, which is generally regarded as a shapeless carbon, has a large surface region and porosity. Previous studies have shown that the synthesized activated carbon is in a highly amorphous state of 64.8% and a low crystallinity of 35.2%, the results achieved in this study as agree with previous studies [23-25].

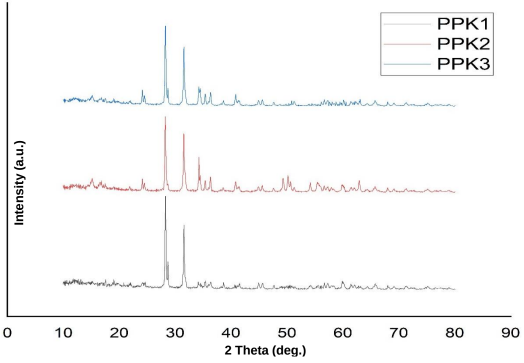


Figure 5: XRD patterns for chemical activation with KOH.

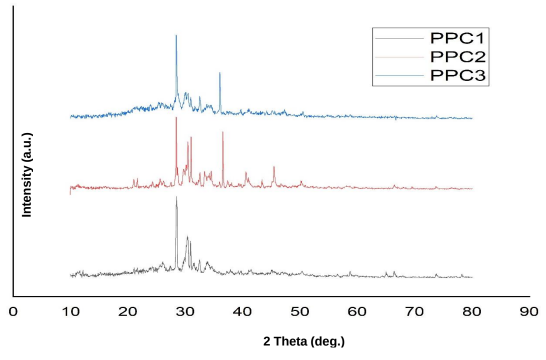


Figure 6: XRD patterns for Physical activation with CO₂.

Table 5: V	Variation	of percentage	crystallinity	of the AC.
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Samples	AC by KOH, %	Samples	AC by CO ₂ , %
PPK1	24.87	PPC1	18.76
PPK2	35.03	PPC2	35.46
PPK3	27.73	PPC3	19.18

The results of AFM analysis on the surface of the precursor PPWs are shown in Figures 7 (a–c), which depict a three-dimensional picture of the potato peel waste. It can be shown from these figures' numerous acute projections, the most noteworthy of which might reach up to 23 nm, 82 nm, and 29 nm with a scanning area of $1.00\mu m \times 1.00\mu m$. The enhancement within the drying period can be related to the surface morphology of the AC observed within the AFM picture, as shown in Figures 8 and 9, respectively. Which gives a high-elevation surface region for the carbon at 1396 nm (chemical activation) and 778 nm (physical activation) within a scanning area of $5.00 \ \mu m \times 5.00 \ \mu m$.

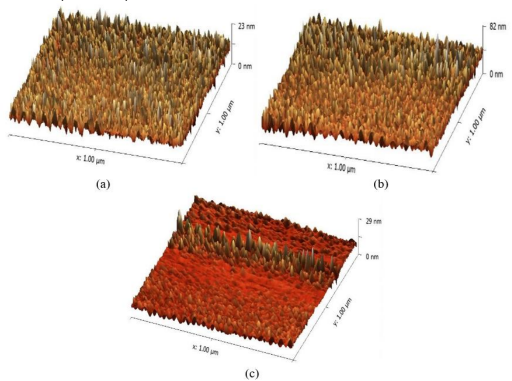


Figure 7: AFM at a magnification (1.00 μm x 1.00 μm) of precursor under the environment atmosphere (a) PPW1, (b) PPW2, and (c) PPW3

The surface morphology of precursor PPW, activated carbon PPK, and PPC was characterized using a FE-SEM test. Figures. 10 (a-c) refers to the nanostructure image for drying at 100°C within 12, 24, and 48 h, respectively. SEM macrographs Figures 11(a) and 12(a) show the external morphology of activated carbon by KOH activation via a chemical process under an atmosphere environment with a carbonization temperature of 500 °C and more intact compared to other activated carbons. The surface of the samples was found to be relatively organized without any pores except for some occasional cracks as compared to raw materials [13]. There are many clear, fine pores visible within the microstructure of activated carbon. The SEM pictures of the two activated carbons, PPK and PPC, indicated that the outside surfaces of these carbons were full of cavities and very unpredictable as a result of activation. There are many cracks, various splits, and small pits distributed over the surface, as well as little cavities dispersed over the surface, showing the extreme interaction of the KOH and CO2 with PPW, which explains the difference between raw materials from agricultural wastes in previous

studies [26, 27].

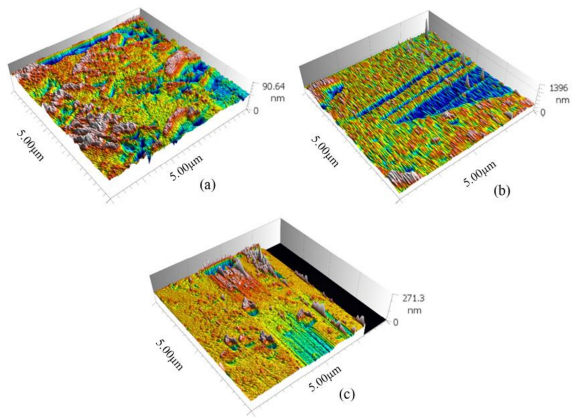


Figure 8: AFM at a magnification (5.00 μ m x 5.00 μ m) of precursor after chemical activation (a) PPK1, (b) PPK2, and (c) PPK3.

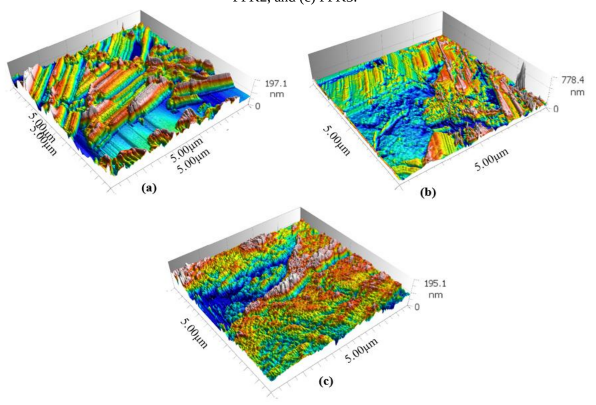


Figure 9: AFM at a magnification (5.00 μ m x 5.00 μ m) of precursor after chemical activation (a)PPC1, (b) PPC2, and (c) PPC3.

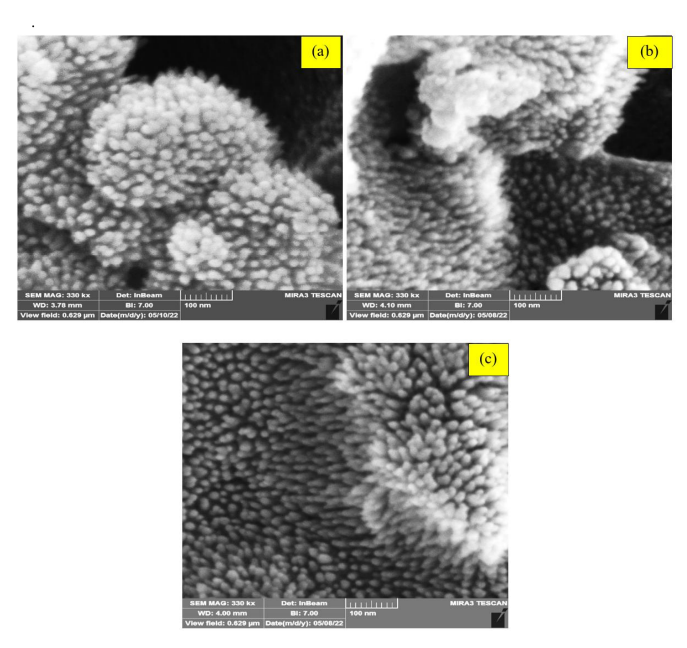


Figure 10: SEM at a magnification $(1\times10.00\text{kx})$ of precursor under the environment atmosphere (a) PPW1, (b) PPW2, and (c) PPW3.

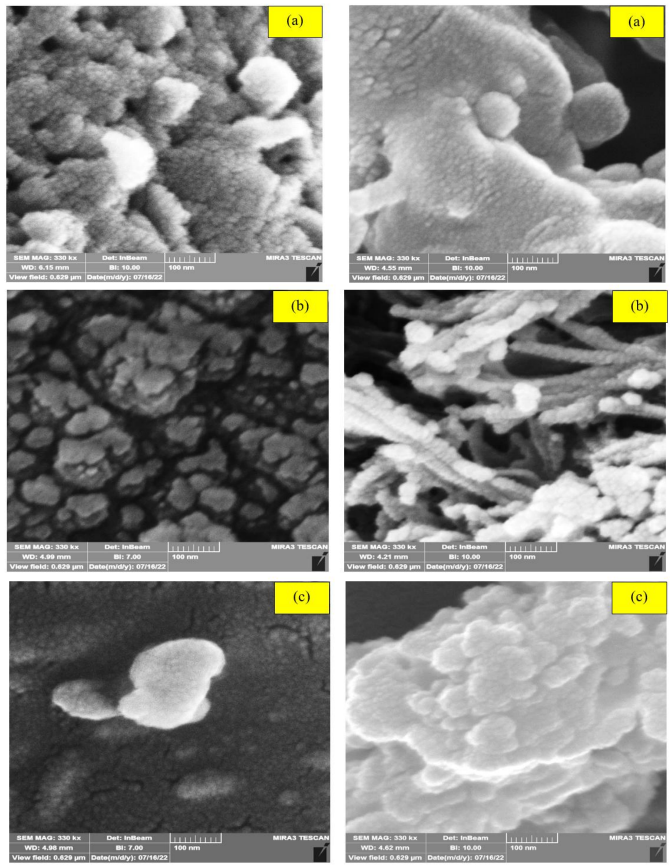


Figure 11: SEM at a magnification (1×25.0KX) of precursor after chemical activation (a) PPK1, (b) PPK2, and (c) PPK3

Figure 12: SEM at a magnification (1×25.0 kx) of precursor after physical activation (a) PPC1, (b) PPC2, and (c) PPC3

4. Conclusions

Actuated carbon adsorbents have numerous applications and have changed applications in later times. Typically, the result of both a well-designed polydisperse permeable structure with a high degree of dispersion for atomic sorption and an expansive inside surface zone.

The AFM image of AC shows the presence of nanotips on the surface. PPK and PPC XRD results confirmed that the pore structure and lack of peak appearance were consistent. The XRD results confirmed that AC-treated KOH and CO₂ had a less crystalline appearance.

The activation of 3-FE-SEM-activated carbon photos derived from PPW by KOH is removed. Also, it appears that the outside surfaces of these carbons are full of cavities and very unpredictable as a result of being activated. This has been affirmed by the well-developed surface area that can be seen on the surface of the potato peel as it appears in this work.

Surface advancement within the potato peel amid pyrolysis was critical, as this would improve the surface area and pore volume of the PPK versus the PPC by accelerating the dissemination of KOH atoms into the pores and subsequently expanding the KOH carbon response through corrosive hydrolysis forms, which would then make more pores.

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Conflict of Interest

The authors declare that they have no conflict of interest.

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