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# Characterization and dye adsorption effectiveness of activated carbon synthesized from olive pomace

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#### ABSTRACT

Studies, about products obtained from agricultural wastes, have increased within the scope of zero waste studies. The olive pomace is produced as a result of olive oil production. In the present study, activated carbon was synthesized using the olive pomace taken from the olive pomace processing plant operating with a three-phase process. The synthesized activated carbon characterization was performed using Scanning Electron Microscope (SEM), Fourier-Transform Infrared Spectroscopy (FT-IR), Brunauer – Emmett – Teller (BET), and X-Ray Crystallography (XRD) devices. Olive pomace activated carbon (OPAC) was used for the adsorption of dye from an aqueous solution. The adsorption efficiency of the OPAC was investigated. The initial pH value of dye solution (6–9), the amount of activated carbon (0.5 and 1.0 g/L), and initial dye concentration (600–1200 mg/L) were optimized. Also, adsorption kinetic and isotherm calculations were evaluated. The optimum parameters were found as the original pH value (pH=8) of dye solutions, OPAC amount of 1.0 g/L and the initial concentration of 1000 mg/L. The Langmuir isotherm model and the pseudo-second-order kinetic model were found as the most suitable models. It can be said that the synthesized material can be used at dye removing from wastewater.

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# INTRODUCTION

The olive oil production is one of the important agro-industries [1]. There are 900 million olive trees on an area of approximately 10 million hectares in the world. According to the averages in last five seasons, olive oil production is around 2.91 million tons [2]. Olive oil production process generates wastewater and pomace, which are difficult to treat. The amount and characteristics of wastes vary according to the production method. Two different processes, classical and continuous, are applied in olive oil production. In the classical production process, oil is extracted using hydraulic presses. Olives are washed with water, crushed and kneaded by adding water. The olive paste is then pressed

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Published by Yıldız Technical University Press, İstanbul, Türkiye This is an open access article under the CC BY-NC license (http://creativecommons.org/licenses/by-nc/4.0/). and oil, wastewater (black water) and solid phase (olive pomace) are separated. The continuous production process is based on the separation of the oil from the olive paste by centrifugation. It consists of feeding, washing, crushing and olive paste preparation units. Depending on the centrifuge used during production, there are two different processes: three-phase and two-phase. In the three-phase process, water is used. After the process, three phases are formed: oil, black water and olive pomace. Since process water is added, three times more wastewater is generated than in the conventional process. In the two-phase process, no process water is added. After the process, two phases are formed as oil and olive pomace. Black water is not formed. Most of the black water is in the pomace [3].

Pomace consists of olive, seed, and pulp. Approximately, 270 and 250 kg of anhydrous pomace is produced per ton of olive in two-phase and three-phase processes, respectively [4]. An average of 3.3 million tons of pomace waste in twophase systems and 3.06 million tons in three-phase systems (at 5% humidity) are produced annually. The results show that every year a high amount of pomace is produced in the world. Pomace has been used to synthesize activated carbon after various processes to reuse. The effectiveness of olive pomace activated carbon (OPAC) in removing various pollutants such as cadmium II, chromium III, arsenic III, phenol, toluene, iodine, methylene blue, etc. from wastewater has been investigated [5]. Dyes are one of these pollutants. If the wastewater containing dye is discharged to the receiving environment without treatment, the dyes in the wastewater are also released to the receiving environment. Dyes are toxic to living organisms. Since it gives color to the water and prevents sunlight from entering the water, it hinders the vital activities of aquatic organisms. For this reason, discharging wastewater containing dye to receiving environments without treatment causes a critical environmental problem. Many studies have been carried out on treating wastewater containing dye [6-9]. Methods such as adsorption, sedimentation, filtration, coagulation, electrocoagulation, biodegradation, electrochemical oxidation, Fenton process, ozonation, sonolysis, wet air oxidation, electrical discharges, photocatalysis, photolysis were applied for the treatment of such wastewaters [6]. Among these methods, adsorption is an effective, simple, and frequently preferred purification technique [10]. The type of adsorbent and the method of synthesis of adsorbent affect the cost of adsorption processes. Activated carbon is generally used in adsorption processes for high porosity, sufficient pore size, large specific surface area, and high mechanical strength. These properties of activated carbon make it one of the reliable adsorbents since it provides a high-efficiency treatment of many organic and inorganic pollutants [11–14]. However, the high price of commercial activated carbons increases the cost of treatment. Therefore, agricultural wastes are used to synthesize the activated carbon for

reducing the cost [11]. There are studies in which agricultural wastes are used as activated carbon after various processes. Rice husk, citrus peel, eggshells, sawdust, cigarette waste, alfalfa, by-products from industries (e.g., sugar cane, paper, tea leaves, and others), palm tree cob, plum kernels, nutshell, wood, corn cob cottonseed shell, rubber seed shell, almond shell, coconut shell, bamboo powder, sunflower seed shell, peach kernel, and olive pomace can be given as examples of these agricultural wastes [11, 15]. These wastes are used as adsorbent to treat wastewater.

The aim of this study was to demonstrate the effectiveness of reusing an agricultural waste as an adsorbent in removing dye from an aqueous solution. OPAC synthesized from pomace was used as an adsorbent. The novelty of this study is that commercial dyes used at paper industry were studied.

# EXPERIMENTAL

#### **Carbonization Experiments**

The olive pomace used in this study was taken from the three-phase pomace processing plant located in Mersin, Türkiye. The moisture content of olive pomace was calculated from the weight loss after drying at 103–105 °C for 3 hours and determined as 3.7%. Next, dried pomace (49.83 gr) was calcined in a specially produced metal screwed crucible at 800 °C for 1440 minutes using a muffle-type furnace.

The flow chart of the OPAC synthesis was given in Figure 1. It was determined that there was a weight loss of 21.41% in the pomace after the calcination process. Before the calcination process, as a result of the carbon-sulfur (Eltra CS-580) analysis, it was determined that the pomace contained 52.01% carbon and 0.025% sulfur. It was determined that the calcined pomace had 88.98% carbon and 0.038% sulfur.

#### **Adsorption Experiments**

Three different water-based dyes (DS-00102:WB dispensing yellow base: JR05 (DS-00102); DS-00203:WB dispensing orange base: JR05 (DS-00203); DS-00502:WB dispensing blue base: JR05 (DS-00502)) and the mixture of three water-based dyes solution (V: V: V; 1: 1: 1) were used. The dyes were taken from a cardboard factory in Mersin, Türkiye.

The pH, OPAC amount, and initial dye concentration effect were studied (Table 1). The sample volume was 200 mL. The original pH level of the aqueous solutions of the dyes was approximately 8. The pH values of solutions were adjusted using 0,1 and 0,01 M  $H_2SO_4$  and NaOH solutions. OPAC was added to the samples after pH adjustment. The magnetic stirrer (at 150 rpm) was used for the mixing process, and samples were taken for color measurement at intervals of 15 minutes. Collected samples were centrifuged at 6000 rpm for 5 minutes for the color measurement of the supernatant. The color measurement was carried out using Hach Lange, DR3900 model spectrophotometer.



Figure 1. OPAC synthesis flow chart.

Table 1. Adsorption experiments conditions

	The effect of pH	The effect of OPAC amount
рН	6, 8*, 9	Optimum pH
AC amount (g AC/200 mL)	0,5	0.5, 1.0

\*: Original pH value of aqueous solutions of dyes.

The dye removal efficiency was calculated using Equation 1. C = C

 $Removal \% = \frac{c_0 - c_t}{c_0} \times 100 \tag{1}$ 

where  $C_t$  (mg/L) is the concentration at t time, and  $C_0$  (mg/L) is the initial concentration.

Isotherm (Langmuir and Freundlich) and kinetic models (pseudo-first and second order, and intraparticle diffusion) were calculated.

## **RESULTS AND DISCUSSION**

## Characterization of the OPAC

Structural properties of OPAC were analyzed with a Scanning Electron Microscope (SEM) (FEI-Quanta 650). Surface functional groups were investigated with Fourier Transform Infrared Spectrometer (FT-IR) (Jasco-6700). The surface area and pore volume of the OPAC were determined using The Brunauer – Emmett – Teller (BET) method by N<sub>2</sub> adsorption at -196 °C with surface area-pore size analyzer (Micrometrics Surface Area and Porosity-TriStar II). The structural feature of OPAC was examined by X-Ray Diffraction Measurement (XRD) (PANalytical-EMPYRE-AN) using Cu K $\alpha$  as a radiation source.

The SEM images of OPAC was given in Figure 2. The surface of the OPAC was smooth. This is the reason that total surface area was low as mentioned below. The sizes of the chars changed at wide range. Carbonization parameters such as the temperature, time and pressure affect the textur-



Figure 2. SEM images of OPAC.

al properties of synthesized material [16, 17]. At this study carbonization method was not optimized. The optimization of carbonization conditions can develop surface properties and adsorption capacity of material.

The FT-IR spectra for OPAC were shown in Figure 3. For the O-H bond of most carboxylic acids, a characteristic broad property is observed in the range 3300–2500 cm<sup>-1</sup>, with a secondary absorption close to 2600 cm<sup>-1</sup> and overlapping the C-H stretching region [18]. Highly non-absorbent peaks above the region between 2700 and 2400 are not associated with any compound has multiple bounds. Hydride vibrations such as thiols and sulfides (SH), boranes (BH), phosphines (PH), silanes (SiH) and arsines (AsH) often cause these absorptions [18]. The peak at 2663 cm<sup>-1</sup> is indicative for O-H stretching [19]. The peak at 2330 cm<sup>-1</sup> can represent CO<sub>2</sub> [19]. The religion between 2260–2100 cm<sup>-1</sup> (2113 cm<sup>-1</sup>) represents C≡C stretch [18]. The peak at 1995 cm<sup>-1</sup> may represent transition metal carbonyl that has the bands between 2100-1800 cm<sup>-1</sup>. The multi-bonded CO group gives an intense absorption band



Figure 4. XRD plot of OPAC.

in regions close to 2000 cm<sup>-1</sup>. The complexity and actual location of bands depend on the compounds nature [18]. C=O stretching vibrations at 1769 cm<sup>-1</sup> peak represents of aldehydes, ketones, lactones or carboxyl groups [20]. Between 2000–1660 cm<sup>-1</sup> is the aromatic combination in general [18]. Bands range between 1650 and 1750 cm<sup>-1</sup> represent esterified and free carboxyl groups that may be convenient to identify pectins [21].

BET analysis results were given in Table 2. When the results were compared with other studies, it was seen that the calculated values for OPAC were lower [11, 12, 22]. It was thought that this situation was caused by the carbonization method applied. Temperature change, carbonization time, and used chemicals create differences in the surface properties of the obtained activated carbon [22].

XRD graph of OPAC was given in Figure 4. In the XRD graph, strong and weak peaks were observed at  $2\theta=25^{\circ}$  and  $2\theta=45^{\circ}$  respectively. The pores formed by the decomposition of carbon along the direction of the graphic structures form the peak at  $2\theta=25^{\circ}$ . This structure produces an more stable aromatic carbon than amorphous carbon [23, 24].

Table 2. Properties of OPAC

Total pore volume	Total pore area	Surface area BET
(cm <sup>3</sup> /g)	(m²/g)	(m²/g)
0,04976	26,962	17,5958

#### **Adsorption Experiments**

# The Effect of Initial pH

One of the essential parameters in adsorption experiments is the pH value of solution [25]. The ionization degree of the pollutant compound and the adsorbent surface charge are affected by the pH of the solution [12, 26]. In addition, hydronium ( $H_3O^+$ ) and hydroxide ( $OH^-$ ) ions in the solution are adsorbed by the adsorbent, which affects the removal capacity of pollutants [27]. Researchers have studied the adsorption of various pollutants at pH ranging between 2–10.9 in different studies [5]. In this study, three initial pH values (6, 8, and 9) were studied. The experiments were performed at initial concentrations of 912, 820, and 996 mg/L for DS-00102, DS-00203, and DS-00502 dyes, respectively. Color removal findings for each sample were given in Figure 5.

Dye removal efficiencies of pH 6, 8, and 9 were close to each other for each dye. The maximum dye removal was carried out in the first 15 minutes. In the following minutes, the amount of dye remained at approximately the same values. The highest removal efficiencies for DS-00102, DS-00203, and DS-00502 dye solutions were calculated as 32, 37, and 55%, respectively.

The removal efficiencies obtained at pH 6, 8, and 9 indicate that dye removal can be done in this range. The different removal efficiencies for each dye were attributed to the differences in the dye surface load. Since pH 8 was the original pH, it was determined as the optimum pH value. Similar pH values (8.10–8.53) were studied by Pala et al. [28] for color removal. In addition, there are studies using similar [29, 30], acidic [12, 31], and basic [27, 30] pH values in adsorption studies with activated carbon obtained from olive pomace. The difference in pH is attributed to the difference in pollutants and carbonization methods.



Figure 5. pH value optimization for color removal from DS-00102, DS-00203, and DS-00502 dye solutions (0.5 g OPAC/200 mL).



Figure 6. OPAC amount optimisation for colour removal from DS-00102, DS-00203, DS-00502, and mix dye solutions (pH=8).

The pH change was monitored during the experiment. After 60 minutes, the pH increased from 6 and 8 to 8.3 and 9.6, respectively.

#### The Effect of OPAC Amount

The experiments investigating the effect of OPAC amount (0.5 and 1.0 g/200 mL) were carried out at pH 8. The initial dye concentrations were 912, 820, 996, and 892 mg/L for the DS-00102, DS-00203, DS-00502, and mix samples (V: V: V; 1: 1: 1), respectively. Color removal efficiencies for different amounts of OPAC (0.5 and 1.0 g/200 mL dye solution) for each dye were given in Figure 6.

As the OPAC amount was increased from 0.5 g to 1.0 g, the color removal efficiency increased. The highest removal efficiencies with 1.0 g OPAC for DS-00102, DS-00203, DS-00502, and mixture samples were calculated as 37, 46, 92, and 63%, respectively. Among all dye solutions, the highest removal was achieved in DS-00502 (92%). Increasing the OPAC amount from 0.5 to 1.0 g increased the removing efficiency of the DS-00102 and DS-00203 dyes by 5 and 9%, re-

spectively. However, the increasing amount of OPAC in the DS-00502 dye solution increased the removal from 55% to 92%. Compared to previous studies, it was seen that higher removal efficiency was performed in a shorter time [5].

#### The Effect of Initial Dye Concentration

To examine the effect of initial dye concentration, DS-00502 dye solutions were prepared as 600, 800, 1000, and 1200 mg/L concentrations. OPAC (1.0 g/200 mL) was added to the solutions. Removal of 97, 93, 92 and 85% was obtained for 600, 800, 1000 and 1200 mg/L, respectively. When the initial dye concentration increased from 600 mg/L to 1200 mg/L, the adsorption capacity of OPAC increased from 110 to 207 mg/g. The highest dye removal was achieved at 1200 mg/L (1020 mg/L). If the remaining dye amount considered, the least amount of dye (18 mg/L) remained from initial concentration of 600 mg/L. However, the most effective removal was achieved with 1000 mg/L. In all experiments, 15 minutes was enough for the system to reach equilibrium. Time-dependent removal results were given in Figure 7.



**Figure 7**. Initial dye concentration optimization for color removal from DS-00502 (pH=8 and 1.0 g OPAC /200 mL).



Figure 8. Langmuir isotherm curve.

For isotherm and kinetic calculations, DS-00502 dye solutions in four different initial concentrations (600, 800, 1000, and 1200 mg/L) were used. Experiment conditions were determined as pH 8, 1.0 g/200 mL OPAC and a reaction time of 120 minutes. The capacity of OPAC at equilibrium was calculated using Equation 2.

$$q_e = (C_0 - C_e) V/W \tag{2}$$

where  $C_e$  (mg/L) is the equilibrium concentration and  $C_0$  is the initial concentration, W (g) OPAC amount and V (L) the sample volume. The capacity of OPAC at time t was calculated using Equation 3.

$$q_t = (C_0 - C_t) V / W \tag{3}$$

where  $C_t$  (mg/L) is the concentration at time t.

## **Adsorption Isotherms**

Langmuir [32] and Freundlich [33] isotherms were calculated in this study.

Table 3. Param	eters of isoth	erm models
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Langmuir isotherm			Freu	Freundlich isotherm			
q <sub>max</sub> R <sup>2</sup> (mg/g)		K <sub>L</sub> (L/mg)	n	R <sup>2</sup>	K <sub>F</sub>		
238.09	0.9886	0.0382	3.4530	0.8877	47.5663		

**Table 4.** The comparison of maximum adsorption capacity with previous studies

Activated carbon material	Removed pollutant	Max. adsorption capacity (mg/g)	References	
This study	Dye	238.09		
Olive Stones	Phenol	58.8	[12]	
Clay	Acid blue 29	104.83		
	Methylene blue	178.64	[17]	
Olive Stone	Phenol	51	[31]	
	Methylene blue	714		
Olive-seed waste residue	Methylene blue	263	[41]	
Acorns		127		
Olive seeds	Methylene blue	115	[42]	
Olive-waste cakes	Iodine	1495	[43]	
	Methylene blue	490		

Langmuir isotherm states that the surface of the adsorbent has a certain number of active sites with the same energy, and the adsorption is reversible. According to this model, adsorption is limited to a single molecular layer [12]. Equilibrium is reached when the adsorption rate is equal to the desorption rate. The Langmuir isotherm accepts adsorption in a single layer, and the surface is homogeneous. It fills the homogeneous surface until the equilibrium moment. At equilibrium, the maximum amount of adsorption is reached [12, 34].

The Langmuir equation was given in Equation 4.

$$C_e' q_e = 1/(q_{\max} K_L) + C_e' q_{\max}$$
(4)

where  $K_L$  is the Langmuir constant and  $q_m$  (mg/g) is the maximum adsorption capacity of OPAC [35].

The graph of  $C_e$  versus  $C_e/q_e$  was given in Figure 8.

 $1/q_{max}$  obtained from Figure 8 gives the slope [36]. Langmuir parameters were shown in Table 3.

Freundlich isotherm is applied to determine adsorption characteristics on heterogeneous surfaces [36]. Equations were given in Equations 5 and 6.

$$q_e = K_F C^{\nu_n} \tag{5}$$

$$Logq_e = LogK_F + 1/n \ LogC_e \tag{6}$$



Figure 9. Freundlich isotherm curve.



Figure 10. Adsorption capacities of OPAC.

where 1/n is heterogeneity factor and  $K_F(L/g)$  is the Freundlich constant [36].

The logarithmic form of Freundlich equation can be expressed as a line equation whose slope is 1/n and where it intersects the axis Log K<sub>f</sub> [12]. The graph of Log C<sub>e</sub> versus Log q<sub>e</sub> was given in Figure 9, and the parameters of the Freundlich isotherm were shown in Table 3.

The  $R^2$  value obtained with the Langmuir isotherm is higher ( $R^2$ =0.9886) than Freundlich isotherm (Table 1). Adsorption occurs equally on the active sides of adsorbent, according to the Langmuir isotherm [37]. Similar findings were obtained in the color removal study of Pala et al. [28] and the phenol removal study of Bohli et al. [12].

The adsorption capacity of OPAC according to initial dye concentration was given in Figure 10. The maximum adsorption capacities were found as 110.4, 142.4, 185.6 and 207.4 for 600, 800, 1000 and 1200 mg/L initial dye concentrations, respectively. The results showed that the dye adsorption capacity in-



Figure 11. Pseudo-first-order curve.



Figure 12. Pseudo-second-order curve.



Figure 13. Intraparticle diffusion curve.

creased with increasing initial dye concentration. In Langmuir isotherm calculations, maximum adsorption capacity q<sub>max</sub> was

Tuble 5. 1 drameters of kinetic models										
Initial concentration (mg/L)	Pseudo-first-order			Pseudo-second-order				Intra particle diffusion		
	q <sub>e</sub> (mg/g) cal.	q <sub>e</sub> (mg/g) exp.	k <sub>1</sub> (min <sup>-1</sup> )	R <sup>2</sup>	q <sub>e</sub> (mg/g) cal.	q <sub>e</sub> (mg/g) exp.	k <sub>2</sub> (min <sup>-1</sup> )	R <sup>2</sup>	k <sub>d</sub> (mg/g.min)	R <sup>2</sup>
600	29.11	110.4	0.0237	0.9178	113.63	110.4	0.0015	0.9988	2.9485	0.9262
800	66.51	142.4	0.0382	0.9644	149.25	142.4	0.001	0.9997	4.3481	0.9506
1000	102.92	185.6	0.0568	0.9919	192.3	185.6	0.0011	0.9997	5.4924	0.7553
1200	63.22	207.4	0.0262	0.9492	217.39	207.4	0.0007	0.9991	5.5461	0.9810

 Table 5. Parameters of kinetic models

found as 238.09 (Table 3). Adsorption isotherm models give the theoretical maximum adsorption capacity [38, 39]. OPAC did not reach the theoretical maximum adsorption capacity. It means that OPAC was not fully covered by dye and still has surface area for more adsorption [38, 40].

There are dye removal studies with similar  $q_{max}$  values [41]. In dye removal studies, methylene blue removal has been mostly studied. The maximum adsorption capacities of other studies were shown Table 4. The removal capacities of activated carbon can be different according to some properties or conditions such as used material, carbonization temperature, carbonization time, removed pollutant, initial dye concentration, initial pH, surface area of activated carbon, the amount of activated carbon, etc.

#### **Adsorption Kinetics**

Pseudo-first and second-order, and intraparticle diffusion model kinetic parameters were calculated.

According to Lagergren, the rate of adsorption is directly proportional to the number of pores on the adsorbent surface. The equation that gives the relation between adsorption capacity and contact time was given in Equation 7 [44].

$$Log(q_{e}-q_{t}) = Log q_{e} - k_{t} \times t/2.303$$
 (7)

where  $k_1$  (min<sup>-1</sup>) is the pseudo-first-order rate constant. The parameters were calculated using the graph of Log  $(q_e-q_t)$  versus t [34]. The pseudo-first-order curve was given in Figure 11. The parameters were shown in Table 5.

According to pseudo-second-order model, valence forces are formed by exchanging electrons or joint use of electrons between adsorbent and adsorbate during adsorption [34, 45].

Calculations were made according to Equation 8.

$$t/q = 1/K_2 q_e^2 + (1/q_e) t \tag{8}$$

where  $K_2$  (min<sup>-1</sup>) refers to the pseudo-second-order rate constant. The parameters were calculated using the graph of t/q, versus t (Fig. 12) and shown in Table 5.

The kinetic model defined by Weber was applied to evaluate

the diffusion mechanism and speed control steps [46]. The intraparticle diffusion kinetic equation was given in Equation 9.

$$q_t = k_d t^{1/2} + C \tag{9}$$

where C (mg/g) is the constant depending on the boundary layer thickness and  $k_d$  (mg/g.min) is the intraparticle diffusion rate constant [47].

Boundary layer diffusion occurs in the first few minutes, and intraparticle diffusion occurs in the time left. The step determining the adsorption rate is intraparticle diffusion [48]. The qt versus t<sup>1/2</sup> was drawn to determine the effect of intraparticle diffusion (Fig. 13). The slope of the obtained equation gives the velocity constant [49, 50]. An intraparticle diffusion plot was drawn using the data obtained after the 15<sup>th</sup> minute.

Results showed that R<sup>2</sup> value closest to 1 were obtained with pseudo-second-order kinetic. Therefore, the adsorption was carried out by the pseudo-second-order kinetic.

#### CONCLUSIONS

Activated carbon was synthesized from the pomace obtained by the three-phase process. Synthesized activated carbon was used to adsorption four different dyes (DS-00102, DS-00203, DS-00502, and mix) from the aqueous solution. The pH value of dye solution, the amount of activated carbon, and the initial dye concentration parameters were optimized. Adsorption kinetics and isotherm models were compared. The most effective removal (92%) was obtained in DS-00502 dye. The optimum parameters were the original pH value (pH=8), 1.0 g/L OPAC, 1000 mg/L initial concentration. The Langmuir isotherm and pseudo-second-order kinetic have been found as the most suitable models. As a result, it can be said that the synthesized material can be used at dye removing from wastewater.

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# DATA AVAILABILITY STATEMENT

The authors confirm that the data that supports the findings of this study are available within the article. Raw data that support the finding of this study are available from the corresponding author, upon reasonable request.

# CONFLICT OF INTEREST

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

## ETHICS

There are no ethical issues with the publication of this manuscript.

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