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The beneficiation of asphalt waste through conversion into an efficient activated carbon adsorbent for diazinon pesticide, optimized through response surface methodology†

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This study reports on converting waste into an activated carbon material for the efficient removal of diazinon pesticide (DP). The asphalt waste obtained from the streets was converted into an activated carbon and was experimentally examined in a batch system. The prepared carbon adsorbents were characterized by energy-dispersive X-ray spectroscopy (EDX), X-ray diffraction (XRD), scanning electron microscopy (SEM), Fourier transform infrared spectroscopy (FTIR), Raman spectroscopy, and Brunauer–Emmett–Teller (BET) analysis. The results revealed that the activated carbon (AC) had an amorphous structure, high porosity, and a relatively high surface area of 788.33 m² g⁻¹. Additionally, functional groups such as –CH₂– and S=O were detected for the prepared adsorbent. The impact of DP sorption parameters, such as, sorbent dosage, initial concentration, and pH were modelled and optimized using central composite design (CCD) *via* response surface methodology (RSM). The optimal conditions obtained from the CCD were found to be 5.6, 30 mg, and 200 mg L⁻¹ for pH, sorbent dosage, and initial pesticide concentration, respectively, with adsorption capacity of 234.25 mg g⁻¹. The experimental data was fitted to the linear form of pseudo first (PFO) and second order (PSO) kinetic models and the data was well described by PSO kinetic models. Based on the thermodynamic parameters, the negative values of Gibbs free energy underscore the spontaneity of the adsorption process. Enthalpy change of 1.9037 kJ mol⁻¹ indicated the endothermic nature, while entropy change of 0.01751 kJ mol⁻¹ K⁻¹ indicated increased disorderliness at the adsorbent–solution interface. The study contributes to sustainable, economical solutions for pesticide contamination, emphasizing the potential of ACs derived from abundant waste materials.

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Sustainability spotlight

The study highlights sustainable conversion of asphalt waste into effective activated carbon for removing diazinon pesticide. Through innovative technology and optimization methods, high adsorption capacity is achieved, offering a sustainable solution to pesticide contamination. Repurposing abundant asphalt waste materials contributes to waste reduction and environmental protection. This approach not only mitigates environmental harm but also underscores the importance of resource efficiency and circular economy principles in addressing pollution challenges and promoting sustainable practices.

1 Introduction

Sustainable waste management (SWM) is a vital component of both the circular economy (CE) and broader sustainability efforts. Contemporary lifestyles have strayed from sustainable

paths, leading to what's termed a 'waste society' rather than a 'prosperous society', marked by excessive waste generation and a pervasive culture of waste.¹ Despite these challenges caused by excessive production of wastes, there are emerging strategies aimed at reversing these trends, including reintegrating municipal and industrial waste into production cycles through material recovery, fostering economies of scale, and embracing circular regenerative initiatives. The benefits of SWM recycling within CE frameworks are manifold, including increased market value of recycled products, reduction of waste disposal sites, and the sustainable sourcing of raw materials.

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However, a pressing question remains: what impact can SWM have on sustainability in the long run?²

The current study investigates the conversion of waste asphalt, recovered from discarded road debris, into activated carbon. This activated carbon is then utilized for the adsorption of diazinon pesticide. Diazinon, a widely used pesticide along with other organophosphorus pesticides (OPPs) has been categorized by the World Health Organization as “moderately harmful” class II due to its acute toxicity, which hinders the activity of the acetylcholinesterase enzyme.³ The mechanisms of action of these pesticides involve inhibiting enzyme activity within the nervous system, thereby leading to various neurological side effects.³ Even at very low concentrations (ng L^{-1} ranges) in drinking water, OPPs can still cause serious negative health effects to humans. For this reason, the European Union has established guidelines for the maximum permissible concentrations of pesticide in drinking water, with a threshold of $0.1 \mu\text{g L}^{-1}$ for individual pesticide compound and $0.5 \mu\text{g L}^{-1}$ for overall pesticides.⁴

To meet the guidelines, scientists, and researchers, have come up with several techniques to remove these pesticides and each of these techniques have its unique set of attractive

features as well as drawbacks (Table 1S, in the ESI†). Nevertheless, a common drawback among most of these treatment approaches is the necessity for high maintenance and elevated operational costs, potentially leading to the generation of by-products post-treatment.⁵ Consequently, the adsorption method is favored for its robust process development, simple design, and operational ease, promoting environmentally sustainable pesticide removal.⁴ Various adsorbents including iron modified montmorillonite,⁶ activated carbon,⁷ biochar,⁸ microbial system,⁹ clays,¹⁰ and zeolites,¹¹ have been developed as a solution for the removal of diazinon pesticide (DP) from the water bodies. One of the benefits of meso- and microporous AC materials, in comparison to alternative solid sorbents, lies in the economical nature of their raw materials. The cost-effectiveness at the industrial production stage is attributed to the abundance of diverse carbon sources, including biomass, wood, polymeric materials, pitch, coals, coke pitch, and various industrial byproducts.¹²

The raw material employed in this study is asphalt, a sticky, dark, and highly viscous liquid, or semi-solid substance. It is commonly found in crude petroleum and occurs in certain natural deposits.¹³ The general composition of asphalt is complex



Fig. 1 A schematic illustration of asphalt recovery process.



consisting of asphaltene as well as the component of crude asphalt soluble in benzene or toluene.^{14,15} Asphaltenes encompass nano-sized conjugated aromatic regions connected by minor aliphatic side chains featuring polar functional groups containing heteroatoms. This configuration facilitates the formation of porous asphaltene aggregates through van der Waals forces, charge-transfer interactions, and hydrogen bonding.¹² Some studies have reported the use of asphalt as an unconventional carbon source,¹⁶ for instance, Homam *et al.*¹⁷ produced AC from a blend of asphalt and polypropylene waste for the sorption of azo dye. Rabeea *et al.*¹⁸ produced AC from asphalt obtained from Biji Refinery. Wang *et al.*¹⁹ utilized magnetic asphalt-based AC for the sorption of methylene blue. Uddin *et al.*¹³ synthesized AC through pyrolysis while, despite several advantages of ACs, one of the major drawbacks that still limit their industrial application is high cost of production due to high energy consumption.²⁰ However, this work was conducted to evaluate the possibility of the use of waste asphalt from road debris as a precursor for AC synthesis. The asphalt used in this work was obtained from road debris sourced within the locality of Palapye, Botswana. Asphalt waste presents a readily available and abundant resource that can be repurposed into high-quality AC, providing a sustainable solution for both waste management and resource conservation. Despite containing trace amounts of hazardous substances like polycyclic aromatic hydrocarbons and heavy metals, proper processing techniques can mitigate these risks effectively. Moreover, the utilization of asphalt waste for AC production not only reduces environmental impact by diverting waste from landfills but also offers economic benefits by creating value-added products. By implementing stringent safety protocols, investing in state-of-the-art processing facilities, and ensuring regulatory compliance, the risks associated with asphalt-derived activated carbon can be minimized, allowing for the realization of its vast potential as a versatile and sustainable raw material.^{21–23}

The asphalt used as a starting material was recovered using rotary evaporation (Rotavap) process (Fig. 1). This process is recognized for its reduced issues related to residual solvent in the recovered asphalt, particularly for high-viscosity asphalt. Furthermore, it has demonstrated greater efficiency in handling larger sample sizes, simplicity, and reduced labor intensity compared to alternative methods.²⁴ The production of AC,

incorporating factors like activation temperature, time, and ratio, was systematically optimized using the Taguchi method for experimental design. The resulting effective carbon materials were then optimized for the removal process of DP from aqueous solutions. The parameters of sorbent dosage, concentration, and pH were methodically optimized through the application of CCD. This comprehensive methodology of using Taguchi and CCD was employed to enhance both the preparation of AC and its effectiveness in DP removal, respectively.

2 Materials and chemicals

Materials: The asphalt used in this work was recovered from road debris sourced within the locality of Palapye, Botswana. **Chemicals:** All the chemical reagents such as NaOH, KOH, HCl and other chemicals were utilized without any further modification because they were of analytical-grade quality. The DP (Table 1) utilized here was obtained from a local agrovet within Palapye, Botswana. Dissolution of 10.8 mL of diazinon (270,000 mg L⁻¹) in 5000 mL distilled water produced a stock solution of 5000 mg L⁻¹ concentration. Different desired concentrations applied in this study were obtained from the stock solution. UV-vis spectrophotometer (Evolution-201, Shimadzu, Japan) was used to detect diazinon at maximum wavelength of 246 nm.

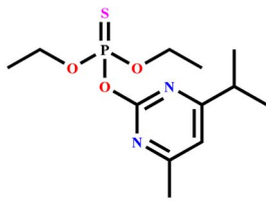
2.1 Recovery of asphalt

The rotary evaporator method was used for the recovery of the asphalt from the discarded road debris according to Fig. 1. In a typical recovery process, ~1 kg of the discarded road debris was dissolved in ~1 L of dichloromethane (DCM) solvent. The mixture was left to stand for about 2 h to allow all the asphalt to dissolve, thereafter it was filtered using vacuum pump. The viscous asphalt was recovered from the filtrate (DCM + asphalt) using rotary evaporation procedure. The recovered asphalt was then kept in a container for further use.

2.2 Taguchi experimental design

Taguchi method constitutes an approach that facilitates the utilization of experimental techniques capable of producing relevant data for efficient statistical examination. The use of

Table 1 Properties of diazinon pesticide^a

IUPAC name	Structure	Molecular weight (g mol ⁻¹)	λ_{\max} (nm)	Solubility
O,O-Diethyl O-[4-methyl-6-(propan-2-yl)pyrimidin-2-yl] phosphorothioate		304.35	246	40 mg L ⁻¹ at 25 °C
	C ₁₂ H ₂₁ N ₂ O ₃ PS			

^a Source: PubChem.



design of experiment is a robust route applied to attain optimal conditions with reduced number of experimental runs.²⁵ In the typical utilization of the Taguchi design technique, the initial stage involves the selection of the response variable under examination. Subsequently, the variables that exert an impact on the response variable are categorized in the second step. The third stage entails the determination of the levels associated with these factors. Following this, the sensitivity of the factor levels is tested through an experimental analysis utilizing an orthogonal array.²⁶

The conventional approach for optimizing the preparation of AC and its application in either energy storage or environmental remediation involves the sequential change of individual factors through “one factor at a time” method.²⁷ However, this traditional methodology has a significant drawback, primarily attributed to the extensive number of experiments it necessitates. The classical ‘changing one factor at a time’ method is notably inefficient, demanding a substantial investment of both labor and time.^{28,29} In contrast, the design of experiments emerges as a highly efficient alternative to traditional experimental approaches.²⁶ This streamlined approach not only reduces the burden of experimentation but also is a more time-effective strategy in optimizing the preparation of AC and its application in the removal of DP.

2.2.1 Analysis of variance. Analysis of variance (ANOVA) serves as a tool for assessing the relative importance of factors within a given context. The ANOVA table incorporates key statistical metrics, including sum of squares (SS), degrees of freedom (DF), variance, and contribution percentage. Within this table, the factor exhibiting the greatest contribution percentage is deemed important in relation to relative significance of all factors. Furthermore, this factor plays a substantial role in influencing the overall response. The *p*-value acts as a measure of the statistical significance opposing the null hypothesis. A statistically significant relationship between the response variable and a specific factor is noted when the *p*-value is lower than the predetermined significance threshold of 0.05.³⁰

2.2.2 Preparation of AC. The experimentation involved the utilization of the Taguchi L₉ orthogonal array, featuring three levels for each of the three factors, as illustrated in Table 2S, in the ESI.† The determination of factors and their respective levels was based on insights reported from existing literature.^{12,17–19} In

the pursuit of optimization, the DP removal efficiency was chosen as the output to be optimized. The calculation of signal to noise (S/N) ratios for the factors was executed by Minitab®19 software. To discern the optimal levels of the variables, both response and ANOVA tables were employed. Fig. S1 in the ESI† outlines the key steps of the process as designed by Taguchi method for the preparation of AC.

The recovered asphalt (~185 g) prior to activation was first carbonized at 400 °C for 180 min in a tube furnace (Model: OTF-1200X, MTI Corporation, USA) with a 10 °C min⁻¹ heating rate under a N₂ gas flow (20 cm³ min⁻¹). The resulting carbon hereafter referred to as AsC was then mixed with KOH at various activation mass ratios as denoted in Table 2. For comparison, the AsC was also activated using NaOH under optimal conditions. The carbon materials activated using sodium and potassium are hereafter referred to as NaAsAC, and KasAC. The resulting AC was dispersed in 50 mL of 1 M HCl solution and stirred for 1 h followed by washing with distilled water until the solution attained a pH range of 6.6–7.0. Subsequently, the AC samples were dried at 100 °C for 24 h and stored for further applications.

2.2.2.1 Adsorption experiments. All the DP sorption batch experiments onto the sorbents were performed in 250 mL Erlenmeyer flasks. Adsorption variables, including initial DP concentration, sorbent dosage, and pH were varied based on the levels outlined in Table 3S in the ESI.† The experimental runs were conducted according to the design outlined in Table 4S, in the ESI.† After 90 min of adsorption time, the solution was centrifuged at 13 000 rpm for 7 min, the remaining DP concentration in the solution was determined using a Shimadzu Evolution 201 UV-vis spectrophotometer at 246 nm wavelength. Subsequently, the efficiency of DP removal and adsorption capacity by the AsC, NaAsAC, and KAsAC adsorbent were calculated. The adsorption procedure was optimized using RSM *via* CCD facilitated by Design Expert version 13.0.5.0. The process variables affecting removal efficiency *viz* initial diazinon concentration, pH, and KAsAC dosage, were each systematically explored across defined ranges comprising five levels (Table 3S, in the ESI†). Consequently, this defined experimental domain resulted in a comprehensive matrix of 20 distinct experimental runs, each characterized by unique configurations of process variables (Table 4S, in the ESI†). Percentage removal (%) and

Table 2 L₉ orthogonal array test layout

S/N	Coded values			Actual values		
	Activation temperature (°C)	Activation time (min)	Activation ratio	Activation temperature (°C)	Activation time (min)	Activation ratio
1	1	1	1	500	45	1
2	1	2	2	500	60	2
3	1	3	3	500	75	3
4	2	1	2	600	45	2
5	2	2	3	600	60	3
6	2	3	1	600	75	1
7	3	1	3	700	45	3
8	3	2	1	700	60	1
9	3	3	2	700	75	2



adsorption capacity (q_e) were determined according to the following equations.

$$\% = \frac{C_0 - C_t}{C_0} \times 100 \quad (1)$$

$$q_e = \frac{C_0 - C_e}{w} \times V \quad (2)$$

where C_0 , C_t , and C_e , correspond to initial concentration, concentration at time intervals, and concentration at equilibrium while q_e , w , and V corresponds to adsorption capacity, sorbent dosage and the volume of the aqueous solution.

2.2.2.2 Kinetics and thermodynamics studies. Exploring kinetic models of the adsorption provides invaluable insights into the underlying mechanisms governing the diazinon adsorption onto AsC, NaAsAC, and KAsAC. The obtained experimental data from the sorption by these prepared carbon materials were meticulously fitted against PFO and PSO kinetic models, as displayed in Table 5S.† Recognizing the pivotal role of temperature in influencing the adsorption phenomena, experimental analysis was systematically conducted across a range of temperatures (298 to 328 K). These temperature variations served as a crucial parameter in comprehending the efficacy of KAsAC as an adsorbent for diazinon removal. Moreover, to discern the thermodynamic implications of temperature on diazinon adsorption, fundamental parameters like change in entropy (S°), change in enthalpy (ΔH°), and change in Gibbs energy (ΔG°) were methodically evaluated. The methodology for calculating these parameters is explained in the ESI.†

3 Results and discussions

3.1 Influence of control factors and noise factors on preparation of AC

Nine different samples of ACs were tested for their percentage removal of DP according to the L_9 array of the Taguchi method (Table 2). Additionally, noise factors were incorporated into the experimental design. The experimental and predicted percentage removal of these AC samples, along with their corresponding S/N ratios, are recorded in Table 3.

Subsequently, the S/N ratio was subjected to ANOVA to ascertain the relevance of the obtained S/N data. The ANOVA

Table 4 ANOVA for the S/N ratios

Source	DF	Adj SS	Adj MS	F-value	P-value
Regression	3	11.493	3.8308	7.20	0.029
Activation temperature	1	5.831	5.8314	10.96	0.021
Activation time	1	3.054	3.0540	5.74	0.062
Activation ratio	1	2.607	2.6071	4.90	0.078
Error	5	2.659	0.5318		
Total	8	14.152			

results pertaining to the S/N ratio are presented in Table 4. Kirby³¹ reported that the F -value serves as a valuable indicator of the influence exerted by control factors on the obtained results. The F -value below one indicates the variable effect is not significant whereas a value closer to two implies the effect of the variable is moderate. Therefore, it can be observed from Table 4 that all the three parameters notably influenced the preparation of ACs.

The impact of control factors on the S/N ratio for DP removal onto the prepared AC samples are shown in Fig. 2. Increasing the activation temperature from 500 to 600 °C resulted in a higher removal capacity, however, at 700 °C there was a decline in S/N ratio of AC. This initial enhancement in removal capacity may be as a result of the creation of additional pores, improved pore structure and surface area. Conversely, at higher activation temperature, the pores are potentially destroyed leading to a loss of sorption sites, thereby reducing the removal capacity.³² The sorption capacity of DP was noted to sharply increase with the rise in activation time and ratio. This is attributed to increase pore formation as a result of prolonged exposure and increasing the amount of activating agent.³³ The optimum parameters for the production of AC were 600 °C, 75 min and activation ratio of 1 : 3.

3.2 X-ray diffraction analysis

The XRD patterns of recovered asphalt, AsC, NaAsAC, and KAsAC are illustrated in Fig. 3. The recovered asphalt exhibited a sharp peak at 12.7° and broader peak between 15.0 and 25.0° which corresponds to γ -band and graphene (002) band.¹⁹ Since it was a recovered asphalt, other peaks at 26.8, 27.6, and 28.0 which contains crystalline components found in asphalt could be ascribed to minerals like quartz, calcite, and various clay minerals. The presence of the γ -band is associated with the

Table 3 Experimental design matrix, response, and the predicted response

Run	Activation temperature	Activation time	Activation ratio	Yield (%)	S/N	Response (% removal)	
						Experimental	Predicted
1	500	45	1	58.88	39.55	94.91	95.51
2	500	60	2	43.45	39.71	96.67	96.88
3	500	75	3	45.34	39.83	98.04	98.26
4	600	45	2	40.86	39.82	97.99	97.16
5	600	60	3	35.90	39.89	98.77	98.53
6	600	75	1	35.46	39.90	98.90	97.92
7	700	45	3	25.78	39.89	98.75	98.81
8	700	60	1	20.88	39.81	97.80	98.20
9	700	75	2	25.38	39.91	98.99	99.57



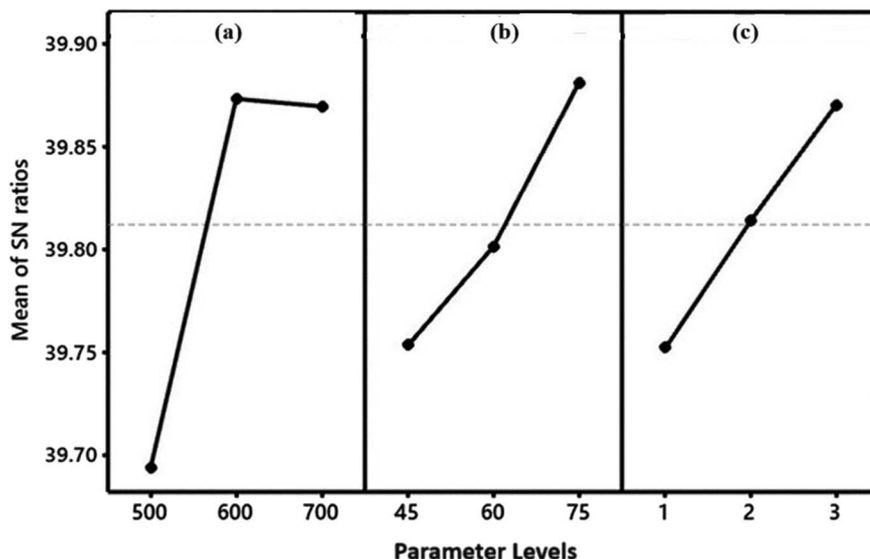


Fig. 2 Influence of activation (a) temperature, (b) time, and (c) ratio preparation parameters on S/N ratio.

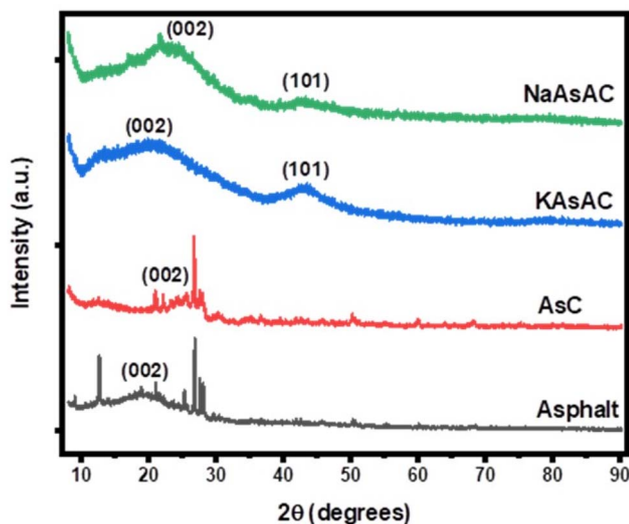


Fig. 3 XRD pattern of asphalt and the prepared carbon samples.

existence of aliphatic hydrocarbon chains, signifying the presence of amorphous disordered structure.³⁴ Graphene layers (002) originate from planar arrangement of aromatic structure.³⁵ Unlike the recovered untreated asphalt, the prepared AC samples (KAsAC and NaAsAC) showed no γ peaks, indicative of the removal of aliphatic hydrocarbon chains during the process of activation. The broad diffraction peaks of KAsAC and NaAsAC at 20.5 and 43.1° correspond to (002) and (101) crystal plane of disordered and graphitic carbon, correspondingly.

3.3 Raman and FTIR analysis

Raman analysis serves as a versatile method for discerning structural alterations within carbonaceous materials. In the pursuit of unveiling the properties and disorder present in activated carbons, Raman analysis was conducted, and the

corresponding spectra can be found in Fig. 4a. Two distinct peaks manifested in the spectra. The G band, positioned around 1585 cm^{-1} , primarily corresponds to the E_{2g} phonon mode, which involves in-plane stretching vibrations of sp^2 carbon atoms in the graphite-like structure. Conversely, the D band, observed at approximately 1325 cm^{-1} , is associated with disorder-induced defects, such as vacancies, dislocations, or grain boundaries, leading to the disruption of the carbon lattice. These observations offers insights into the degree of structural order, providing valuable information about the material's arrangement and organization.³⁶ It is noteworthy to highlight a distinct contrast in the intensities of the D-bands observed across the prepared carbon samples. Comprehensive details, including the Raman shift and intensity ratio across all carbon samples, are presented in Table 5. A noticeable shift was evident in the I_D/I_G ratio, a change that is intricately linked to the level of order within the structure. The D band, associated with structural disorder, indicated a higher intensity, serving as a key indicator of a profoundly carbon disordered structure.^{37,38}

The IR spectra of recovered asphalt material, pretreated material AsC, KasAC, and NaAsAC are given in Fig. 4b. The AsC bands at 2915 and 2850 cm^{-1} were attributed to the stretching vibration of the methylene group, whereas the bending vibrations of $-\text{CH}_2-$ are represented by 1465 and 1375 cm^{-1} bands. The observed peaks at 1590 and 1000 cm^{-1} are indicative of pyridine clusters and S=O vibration functional, respectively. The spectral bands detected in the region of 1600–1400 cm^{-1} signify the existence of the skeletal aromatic structure.^{19,39} Fig. 4b illustrates chemical and structural transformation of the recovered untreated asphalt to AsC and AC (KAsAC and NaAsAC). In the spectrum of AsC, the intensity of the bands associated with the bending and stretching vibrations of the methylene group was notably diminished compared to those observed recovered asphalt. Moreover, no vibration bands attributed to $-\text{CH}_2-$ were detected in KAsAC and NaAsAC. This



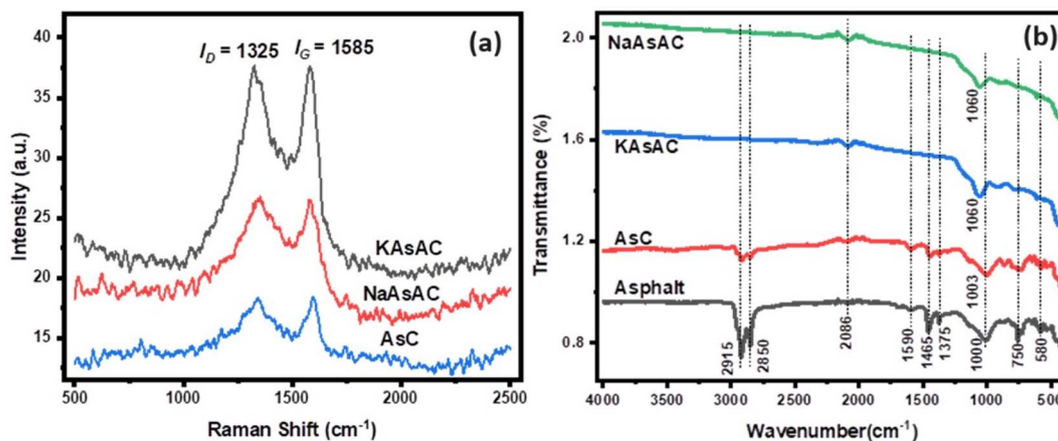


Fig. 4 (a) Raman spectra of the prepared samples from asphalt and (b) FTIR spectra for asphalt and the prepared samples.

Table 5 Raman spectral shift and the ratio of band intensities of AC samples

	Samples		
	AsC	KAsAC	NaAsAC
D band (cm^{-1})	1342	1325	1350
G band (cm^{-1})	1595	1585	1578
I_D/I_G	0.98	1.03	1.00

occurrence is linked to the dehydrogenation of AsC at elevated temperatures. The thermal treatment at elevated temperatures facilitated the removal of hydrogenous organic compounds and induced the conversion of secondary carbon species into graphitic carbon. The peak at 1590 cm^{-1} was absent in the AC samples showing the absence of pyridine functional groups on the prepared ACs. No significant change was observed in the intensity of the band at 1000 cm^{-1} which indicates the presence of S=O group before and after treatment. The bands between 1600 and 1400 cm^{-1} disappeared in KAsAC and NaAsAC

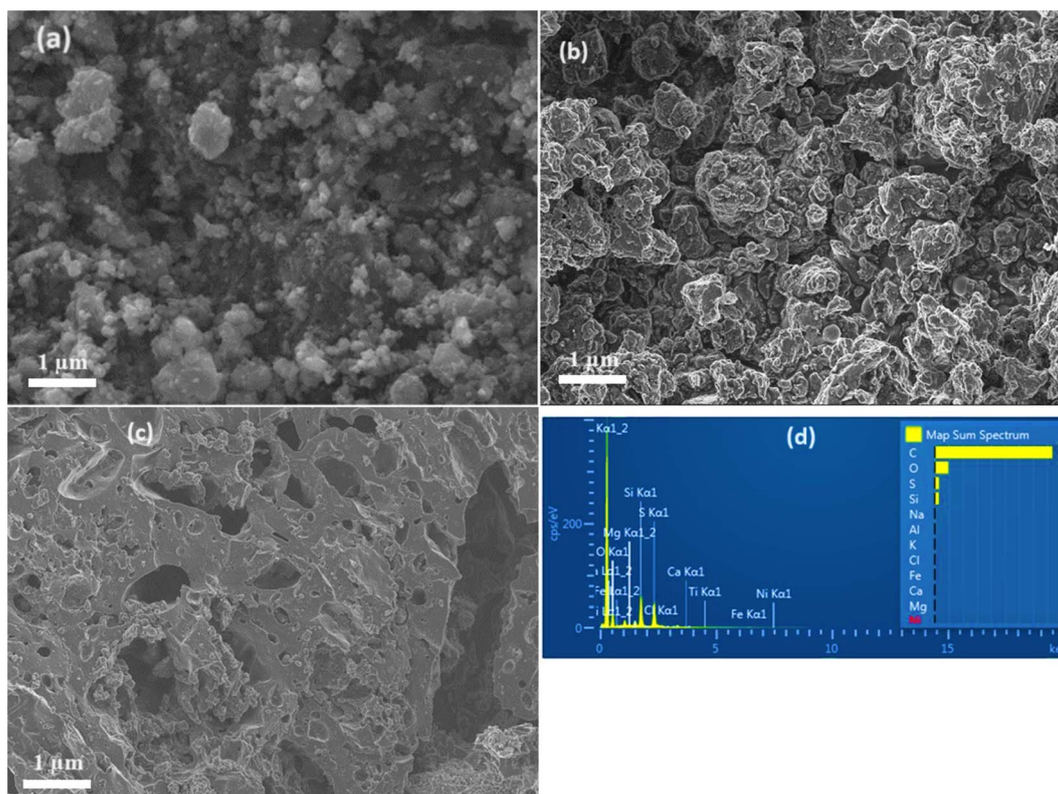


Fig. 5 SEM images and the EDS of the prepared carbon materials.



samples. This suggests that the aromatic skeletal structure underwent cross-linking and gradually transformed into a graphite carbon structure.⁴⁰

3.4 Scanning electron microscopy

The SEM micrographs of the prepared carbon samples and the EDX of KAsAC are shown in Fig. 5. In Fig. 5a the micrograph depicts the morphology of AsC which appears irregular and somewhat clumped, lacking well-defined pores or surface features. Fig. 5b is AsC activated by NaOH (NaAsAC). NaOH activation enhanced the material's structures resulting in clearer grayscale contrast, more interparticle porosity and agglomerated particles. Fig. 5c for AsC activated by KOH (KAsAC), showed a more improved porosity compared to NaAsAC. KOH activation led to a highly porous structure with higher surface area. Both the NaOH and KOH activation introduced functional groups and created a more porous network within the carbon material improving their adsorptive capabilities. Fig. 5d displays EDX analysis of KAsAC which shows it is mainly composed of carbon. Various trace elements like O, S, Si, Na, Al, K, Cl, Fe, Ca, Mg, and Ni in the prepared AC from reclaimed asphalt originate from oxidation, asphalt mix, aggregate minerals, and environmental sources.

3.5 Thermogravimetric analysis

The TG curves for the AsC, KAsAC, and NaAsAC are shown in Fig. 6. As depicted in Fig. 6, a noticeable weight reduction for the carbonized asphalt begins at ~400 °C and reaches around 35% of the initial weight at 750 °C. The initial weight loss predominantly results from the volatilization of organic oil residues with lower molecular weights. The AsC exhibited higher thermal stability due to its relatively higher degree of graphitization and higher inorganic impurities compared to KAsAC and NaAsAC. During the carbonization process, asphalt undergoes pyrolysis, leading to the elimination of volatile components and the formation of a more ordered carbon structure. This enhanced graphitization

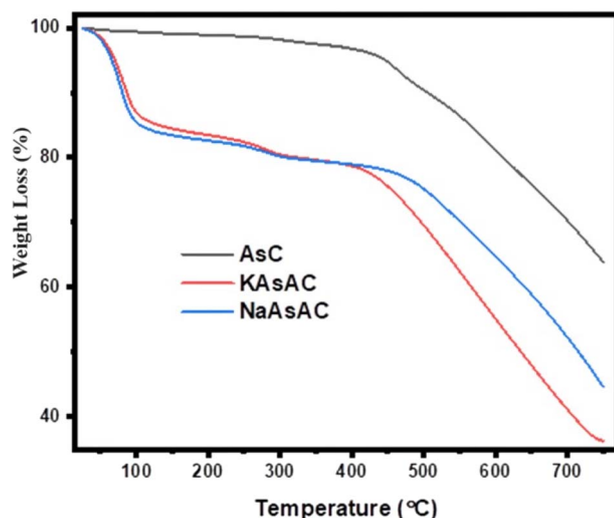


Fig. 6 TG curves for AsC, KAsAC, and NaAsAC.

results in stronger intermolecular interactions and a higher thermal decomposition temperature. Additionally, the gradual weight loss observed during decomposition indicates a slower rate of bond cleavage, which is characteristic of a more stable carbonaceous material.⁴¹

The treatment of AsC with KOH and NaOH resulted in the incorporation of additional functional groups and porosity which lowered the thermal stability of KAsAC and NaAsAC. The defects and functional groups in the KAsAC and NaAsAC carbon probably act as initiation sites for decomposition reactions at lower temperatures below 200 °C. Furthermore, the existence of highly reactive oxygen-containing functional groups like hydroxyl and carboxyl groups, can facilitate oxidative decomposition processes, leading to a more pronounced weight loss at lower temperatures. The differences in thermal stability between KAsAC and NaAsAC can be attributed to the difference in the etching ability of the activating agents. NaAsAC exhibited slightly better thermal stability compared to KAsAC, possibly due to differences in the extent of functional group formation and pore structure development.

3.6 Surface area analysis

Fig. 7 shows the N₂ adsorption–desorption isotherms and the distribution of the pore sizes of AsC, KAsAC, and NaAsAC. The isotherms shown in Fig. 7a describe the correlation between the quantity of gas or vapor adsorbed by a material and the relative pressure (P/P_0) at a constant temperature while the pore distribution (Fig. 7b) represents the distribution of different pores sizes within the material. As depicted in Fig. 7b, AsC, KAsAC, and NaAsAC materials exhibited a combination of micro and mesopores within their structure. According to IUPAC classification, KAsAC shows type I isotherm characterized by monolayer adsorption on a uniform surface which is shown by a steep increase in adsorption of N₂. This is suggestive of monolayer adsorption at specific sites which mainly composed of micropores.^{42,43} NaAsAC shows type IV isotherm signifying a multilayer adsorption on a heterogeneous surface. This is explained by a broader distribution in NaAsAC indicating the formation of multiple layers of adsorbate molecules.⁴⁴ The nitrogen adsorption isotherm for AsC initiates at a lower volume of nitrogen adsorbed and steadily rises as the relative pressure increases. The curve exhibits a smooth and consistent trajectory without any discernible inflection points. This characteristic suggests that the material possesses moderate porosity, indicative of its ability to accommodate nitrogen molecules within its structure without abrupt changes in adsorption behavior. The BET surface area of AsC, KAsAC, and NaAsAC was 1.02, 788.33, and 58.74 m² g⁻¹, respectively, while the total pore volume of AsC, KAsAC, and NaAsAC was 0.0021, 0.4451, and 0.0521 cm³ g⁻¹, respectively.

3.7 Adsorption studies

3.7.1 Optimization of the DP removal parameters

3.7.1.1 Effect of adsorption time. The influence of adsorption time on the removal of DP was studied at a fixed adsorbent dosage (30 mg), DP initial concentration (150 mg L⁻¹), and



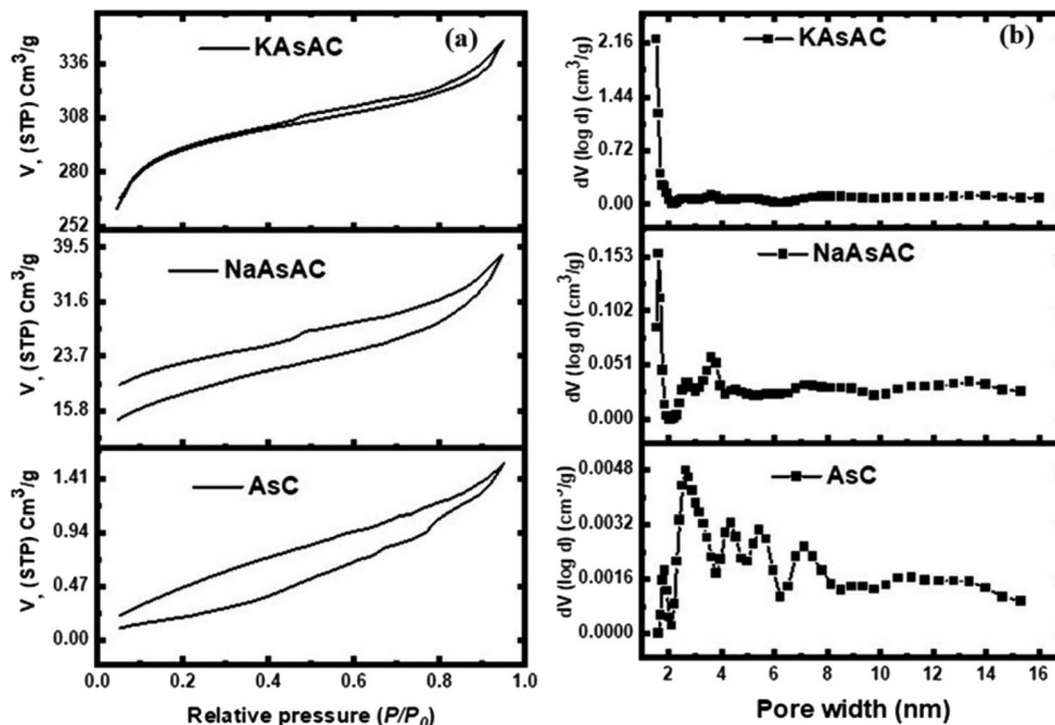


Fig. 7 N_2 adsorption–desorption isotherms and pore width distribution of AsC, NaAsAC, and KAsAC samples.

natural pH of the DP solution indicate the actual value of pH. The sorption uptake was monitored from 10 to 90 min. The impact of adsorption time on the sorption uptake of AsC, NaAsAC, and KAsAC is shown in Fig. 8. Fig. 8 demonstrates that the sorption of DP increases with time attaining saturation around 20 min for AsC and NaAsAC while KAsAC attained saturation after 50 min of contact time. The high initial uptake capacity was due to high number of available adsorption sites which were responsible for the adsorption of DP the prepared carbon sites. After the saturation of the adsorption sites, no

observable change was seen on the removal of DP. The adsorbate removal exhibited rapid initial kinetics, followed by a gradual decline until reaching equilibrium.^{45,46} After 90 min of adsorption time, the adsorption capacity of AsC, NaAsAC, and KAsAC was 147.40, 159.33, and 178.70 $mg\ g^{-1}$, respectively.

3.7.2 Optimization by CCD statistical analysis and model fitting. From the CCD, a connection between the process variables (adsorbent dosage, DP initial concentration, and pH) and output response (adsorption capacity) can be formulated by the quadratic model. Table 6 shows experimental and the predicted adsorption capacities obtained from the design. Based on the ANOVA approach, the following quadratic expression was obtained.

$$\begin{aligned} \text{Adsorption capacity} = & 201.51 + 10.49 \times X_1 + 1.66 \times X_2 \\ & + 1.51 \times X_3 - 0.018 \times X_1 \cdot X_2 + 0.063 \times X_1 \cdot X_3 - 0.00102 \\ & \times X_2 \cdot X_3 + 0.012 \times X_1^2 + 0.00015 \times X_2^2 - 0.29 \times X_3^2 \quad (3) \end{aligned}$$

Table 7 summarizes the information related to the ANOVA for Quadratic model level response. The ANOVA for the response surface quadratic model applied to the sorption of DP by KAsAC indicated a substantial model significance, as evidenced by the model's F -value of 121.59. The p -values below 0.05 further established the relevance of the model terms. In this instance, the significant model terms were X_1 , X_2 , and X_1^2 . The predicted R^2 value of 0.9366 shows reasonable agreement with the Adjusted R^2 of 0.9828, with a difference of less than 0.2. Adequate precision, which measures the S/N, is desirable above 4; here, the ratio of 38.321 indicates an adequate signal.

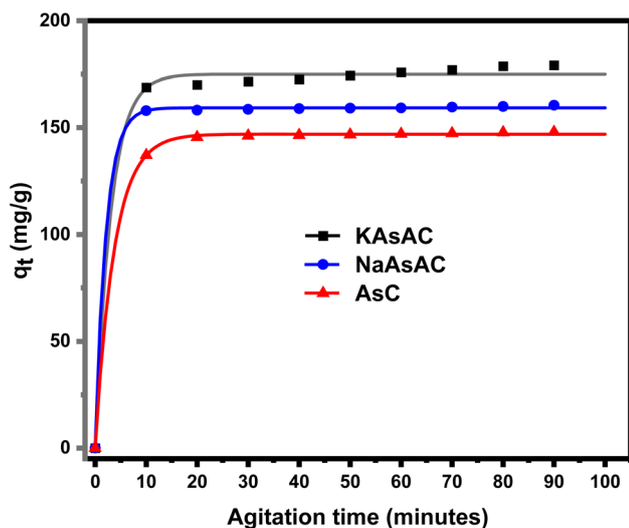


Fig. 8 Effect of adsorption time on DP over AsC, NaAsAC, and KAsAC sorbents.



Table 6 The actual and predicted adsorption capacity from the experimental matrix

Run	Parameters			Response (adsorption capacity, mg g ⁻¹)	
	Adsorbent dosage, X ₁ (mg)	DP initial concentration, X ₂ (mg L ⁻¹)	pH, X ₃	Experimental	Predicted
1	40	150	1.2	113.48	119.76
2	30	100	3	74.86	113.48
3	40	66	5.5	236.99	53.96
4	50	100	8	154.92	77.31
5	40	150	5.5	116.66	131.88
6	23	150	5.5	77.31	222.09
7	50	200	3	232.65	154.92
8	50	200	8	163.87	163.87
9	30	200	8	222.09	232.60
10	40	150	5.5	105.22	134.10
11	40	150	9.7	53.96	128.40
12	40	150	5.5	206.46	129.45
13	40	150	5.5	119.76	135.40
14	40	150	5.5	128.43	130.32
15	40	234	5.5	137.49	206.46
16	30	100	8	129.45	116.66
17	30	200	3	130.32	236.99
18	50	100	3	134.17	74.86
19	57	150	5.5	135.40	105.22
20	40	150	5.5	131.88	137.49

Therefore, this model is suitable for exploring the design parameters.

The R^2 for this model was 0.9909, indicating the model's significant capacity to predict variations in response. The residuals cluster closely around or align with the 45° line, affirming the normal distribution of errors (Fig. S2a†). Additionally, the scatter plot of residuals against predicted values (Fig. S2b†) demonstrates their random dispersion around the reference line, with no discernible pattern. According to eqn (6), the estimated efficiency of DP removal by the KAsAC, and the comparison between model predictions and experimental data

is outlined in Fig. 9a. These projections from the RSM exhibited a strong alignment with the experimental indicating a robust fit.

The impacts of the primary variables (−1 to +1 levels) on diazinon adsorption capacity are outlined in Fig. 9b. It is imperative to highlight that in Fig. 9b, when describing the influence of one variable, other variables are kept at the zero level. For instance, it is observed that for the sorbent dosage variable (X_1) when increased from −1 (30 mg) to +1 (50 mg), the remaining two variables—diazinon concentrations (X_2) set at 150 mg L⁻¹ and pH (X_3) at 5.5 remain at the zero level. Optimal values for sorbent dosage, DP initial concentration, and pH

Table 7 ANOVA for response surface quadratic model in adsorption of DP by KAsAC

Source	Sum of squares	D_f	Mean square	F -value	P -value
Model	48 405.63	9	5378.40	121.59	<0.0001
X ₁ -dosage	13 249.16	1	13 249.16	299.53	<0.0001
X ₂ -initial concentration	32 147.55	1	32 147.55	726.78	<0.0001
X ₃ -pH	45.11	1	45.11	1.02	0.3364
X ₁ X ₂	663.88	1	663.88	15.01	0.0031
X ₁ X ₃	19.72	1	19.72	0.4459	0.5194
X ₂ X ₃	0.1314	1	0.1314	0.0030	0.9576
X ₁ ²	2144.58	1	2144.58	48.48	<0.0001
X ₂ ²	2.03	1	2.03	0.0460	0.8345
X ₃ ²	46.01	1	46.01	1.04	0.3318
Residual	442.33	10	44.23		
Lack of fit	394.06	5	78.81	3.16	0.189
Pure error	48.27	5	9.65		
Cor total	48 847.96	19			
	R^2	0.9909			
	Adj. R^2	0.9828			
	Pred. R^2	0.9366			
	Adeq precision	38.32			



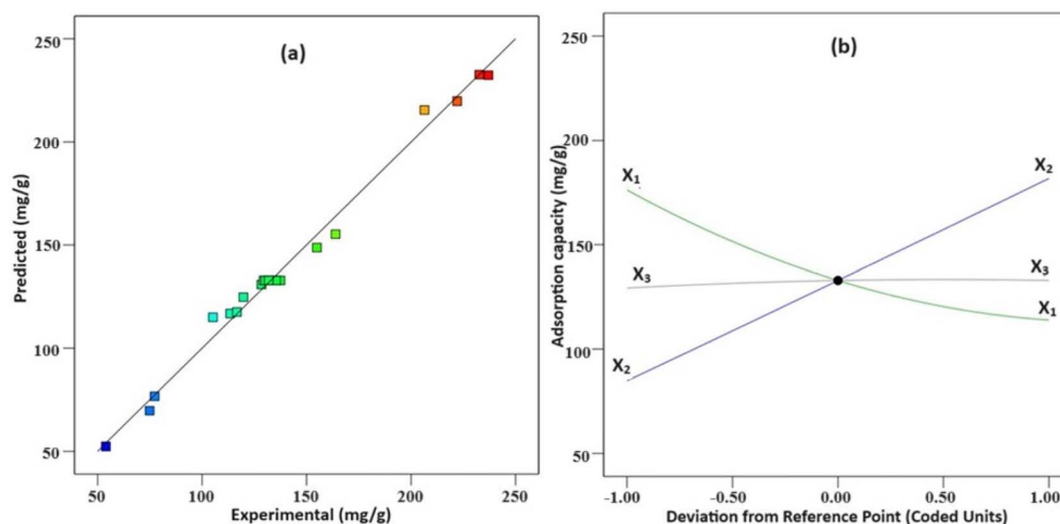


Fig. 9 (a) Experimental and predicted plot obtained from the design, (b) response surface (perturbation) illustrating the impact of primary variables (X_1 , X_2 , and X_3) on diazinon removal efficiency.

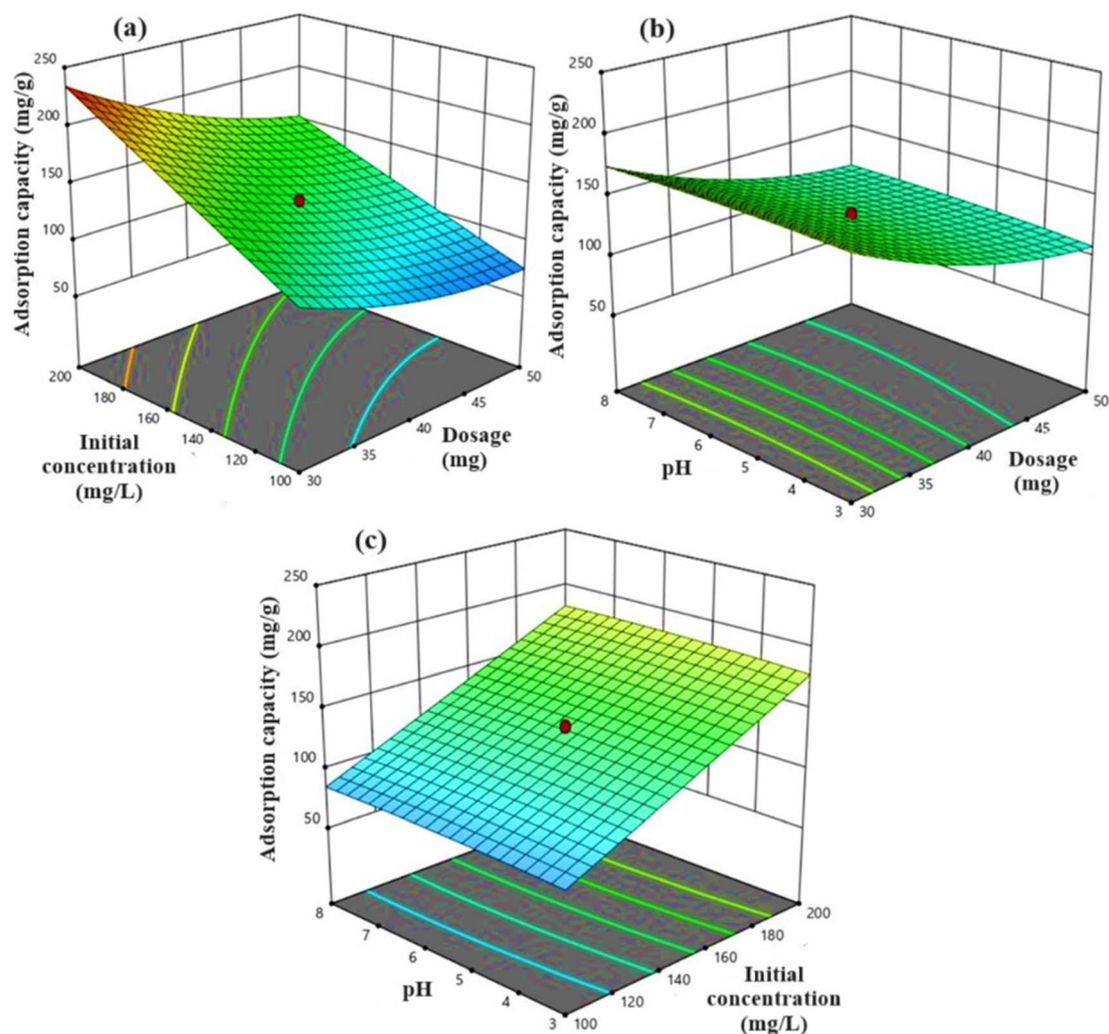


Fig. 10 The 3D plots illustrating the interactions between (a) initial concentration and dosage, (b) pH and dosage, and (c) pH and initial concentration for DP adsorption.



were determined to be 30 mg, 200 mg L⁻¹, respectively. Under these conditions, the predicted diazinon adsorption capacity was 234.25 mg g⁻¹, with a desirability score of 0.985. Conducting similar experiments under these optimal conditions showed the high repeatability of the method in predicting the actual adsorption capacity, with a low relative deviation. Under these optimal conditions, the experiment was repeated, demonstrating a consistent adsorption capacity of 231.97 ± 1.61 mg g⁻¹, confirming the accuracy of the model and its suitability for the study.

The adsorption capacity shown in Fig. 10a was reduced from 176.24 to 113.99 mg g⁻¹ with an increase in KAsAC dosage from 30 to 50 mg. At high adsorbent dosages, interparticle interactions become more pronounced and the adsorbent particles start blocking each other's active sites and thus increasing diffusional resistance thereby decreasing the adsorption of the diazinon molecules. Also, interparticle interactions reduce the effective surface area available for adsorption.^{47,48}

The increase in adsorption of diazinon onto KAsAC with increase in the initial concentration from 100 to 200 mg L⁻¹ can be explained by several factors. Firstly, higher initial concentrations of diazinon in the solution provide a greater driving force for adsorption onto the KAsAC surface. This means that as the concentration of diazinon increases, more diazinon molecules are available to interact with the adsorption sites on the KAsAC, leading to an overall increase in adsorption capacity as shown in Fig. 10b. Moreover, increased initial concentrations of diazinon can also result in a higher concentration gradient between the solution phase and the activated carbon surface. This concentration gradient facilitates the movement of the molecules of diazinon from the solution onto the activated carbon surface, promoting greater adsorption efficiency especially when multilayer adsorption is involved.⁴⁹

The influence of pH on adsorption is displayed in Fig. 10c. The slight increase in DP uptake onto KAsAC, from 129.28 to 132.93 mg g⁻¹, with a change in pH from 3 to 8 could be associated with the existence of -CH₂- and S=O functional groups on KAsAC surface. At lower pH values (pH 3), the KAsAC surface tends to be more positively charged because of protonation of -CH₂- groups and the presence of positively

charged species on the S=O groups. This positive surface charge can enhance electrostatic interactions with the negatively charged diazinon molecules, promoting adsorption. Additionally, at lower pH, diazinon molecules may exist predominantly in their protonated form, increasing their affinity for the activated carbon surface through hydrogen bonding and other interactions with functional groups.⁵⁰

3.7.2.1 Effect of temperature. The impact of temperature on the sorption uptake of DP was evaluated at temperature ranging from 298 to 328 K, with DP initial concentration of 200 mg L⁻¹, 90 min duration, pH of 5.6, and sorbent dosage of 1 g L⁻¹ using KAsAC. Fig. 11a shows that adsorption of DP onto KAsAC increases with increase in temperature from 298 to 328 K. The higher temperatures typically led to greater molecular motion and kinetic energy within the system. This increased thermal energy enhances the collision frequency between DP molecules in the solution and the adsorption sites on the KAsAC surface, promoting more effective adsorption.^{45,49,51} The results of the impact of temperature were further examined with the thermodynamics parameters ΔG° (J mol⁻¹), ΔH° (J mol⁻¹), and ΔS° (J K⁻¹ mol⁻¹).⁴⁵ These parameters were determined according to the following expressions:

$$\Delta G^\circ = T\Delta S^\circ - \Delta H^\circ \quad (4)$$

$$\Delta G^\circ = -RT \ln K^\circ \quad (5)$$

$$K^\circ = \frac{q_e}{C_e} \quad (6)$$

$$\ln K^\circ = \frac{-\Delta H^\circ}{RT} + \frac{\Delta S^\circ}{R} \quad (7)$$

where K° denotes equilibrium constant, T denotes absolute temperature (K), and R denotes the universal gas constant (0.008314 kJ mol⁻¹ K⁻¹). ΔS° and ΔH° were determined from the intercept slope of $\ln K^\circ$ vs. $1/T$ plot. The sorption data was subjected to further analysis through the van't Hoff plot as shown in Fig. 11b while Table 8 summarizes the thermodynamic information. The positive value of ΔH° indicates that the sorption of diazinon onto KAsAC is endothermic chemisorption.⁵² As shown in Table 8, the Gibbs free energy change (ΔG°)

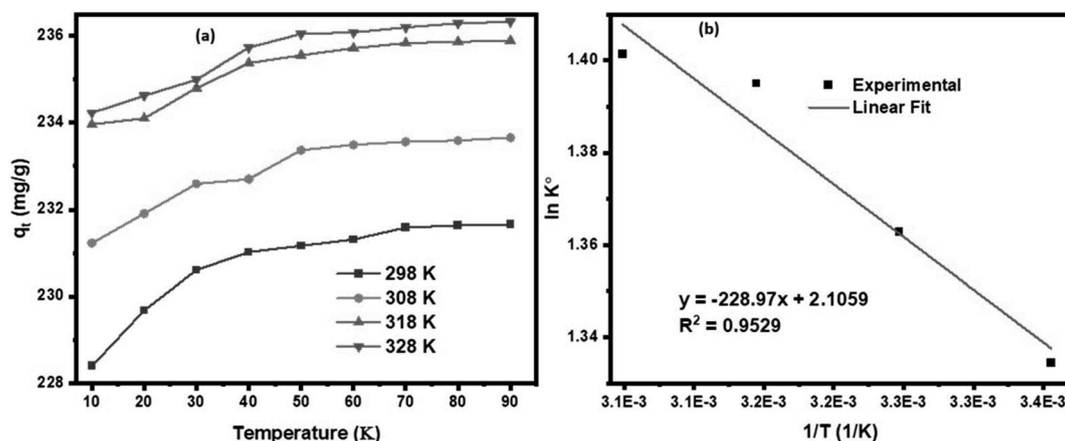


Fig. 11 (a) Effects of temperature on the DP over KAsAC, (b) van't Hoff plot for DP sorption onto KAsAC.



Table 8 Diazinon sorption over KAsAC thermodynamic parameters

Temperature (K)	K°	$\ln(K^{\circ})$	ΔG° (kJ mol ⁻¹)	ΔH° (kJ mol ⁻¹)	ΔS° (kJ mol ⁻¹ K ⁻¹)
298	3.7977	1.3344	-3.3061	1.9037	0.01751
308	3.9074	1.3629	-3.4899		
318	4.0351	1.3950	-3.6882		
328	4.0607	1.4013	-3.8215		

exhibits negative values across all tested temperatures, indicating the spontaneous nature of diazinon adsorption on KAsAC. The negative values ΔG° signify that solutes favor retention in the stationary phase over migration into the mobile phase. The transfer of solutes from the mobile phase to the stationary phase is thermodynamically favored in terms of enthalpy. The ΔS° value was positive suggesting an increasing randomness or disorderliness within the system during the process of adsorption process. This could occur if the sorption process leads to a more disordered arrangement of molecules on the activated carbon surface or if there is an increase in the degree of freedom of the molecules involved.

3.7.2.2 Kinetics models. The investigation into factors influencing adsorption rate involved a study on sorption kinetics. The study employed PFO and PSO equations to determine the kinetic model that best describes the process. Table 10 represents the results, including the computed parameters for each model while Fig. 12 displays the kinetic model plots. The greater R^2 values (Table 9) shows the sorption process was best described by PSO model. The model also showed a precise fit for the experimental and calculated data for the AsC, NaAsAC, and KAsAC adsorbents. Furthermore, it was observed that the calculated equilibrium capacity ($q_{e,cal}$) determined from PSO equation were close to experimental equilibrium capacity ($q_{e,exp}$). This observation indicates that the sorption rate is primarily influenced by the number of vacant sites on the sorbent.

The possible adsorption mechanism of DP adsorption onto the prepared carbon materials is shown in Fig. 13. According to

the figure, surface of the prepared ACs interacted with diazinon through several mechanisms. The nonpolar methylene group ($-\text{CH}_2-$) facilitated weak van der Waals interactions with diazinon's nonpolar regions. Pyridine clusters, with their π -electron-rich aromatic structure, engaged in π - π stacking interactions with diazinon's π -electron system, and hydrogen bonding could occur between the nitrogen in pyridine and the hydrogen or oxygen in diazinon. The sulfone group ($\text{S}=\text{O}$) formed hydrogen bonds, with the oxygen acting as a hydrogen bond acceptor. Additionally, the aromatic skeletal structure of the activated carbon allowed for π - π interactions with diazinon's aromatic rings.

3.7.2.3 Comparison with the reported literature. This study aimed at evaluating the efficacy of KAsAC in adsorbing diazinon, a common pesticide, by comparing its performance with previously reported data from the literature. The study aimed to enhance understanding of the adsorption capacities of the

Table 9 Kinetic model parameters of the prepared sorbents

Model	Parameters	Sorbents		
		AsC	NaAsAC	KAsAC
PFO	$q_{e,exp}$ (mg g ⁻¹)	147.40	159.33	178.70
	$q_{e,cal}$ (mg g ⁻¹)	143.79	157.39	173.73
	K_1 (L min ⁻¹)	-0.0359	-0.0235	-0.03
	R^2	0.9359	0.8851	0.8966
PSO	$q_{e,cal}$ (mg g ⁻¹)	148.59	160.00	181.16
	K_2 (g mg ⁻¹ min ⁻¹)	0.00673	0.00625	0.00552
	R^2	0.9998	0.9998	0.9998

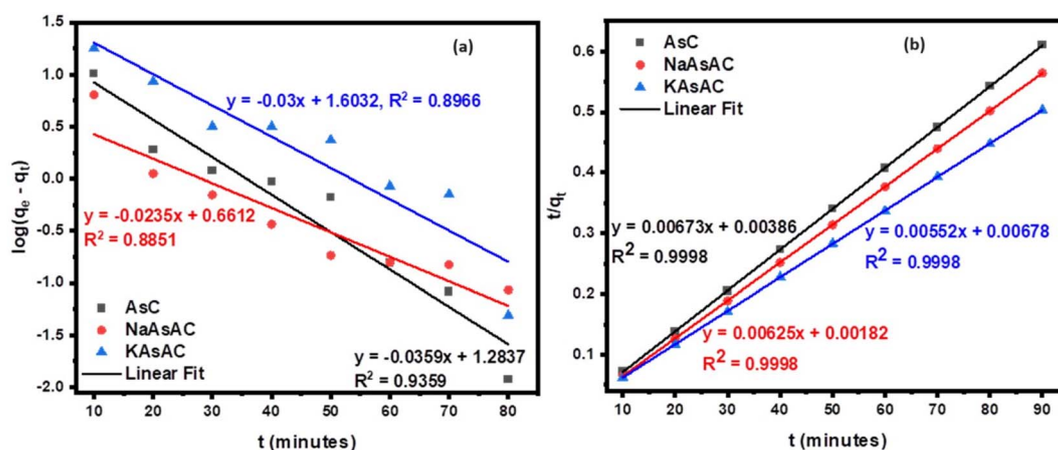


Fig. 12 The (a) PFO and (b) PSO plots for AsC, NaAsAC, and KAsAC samples.



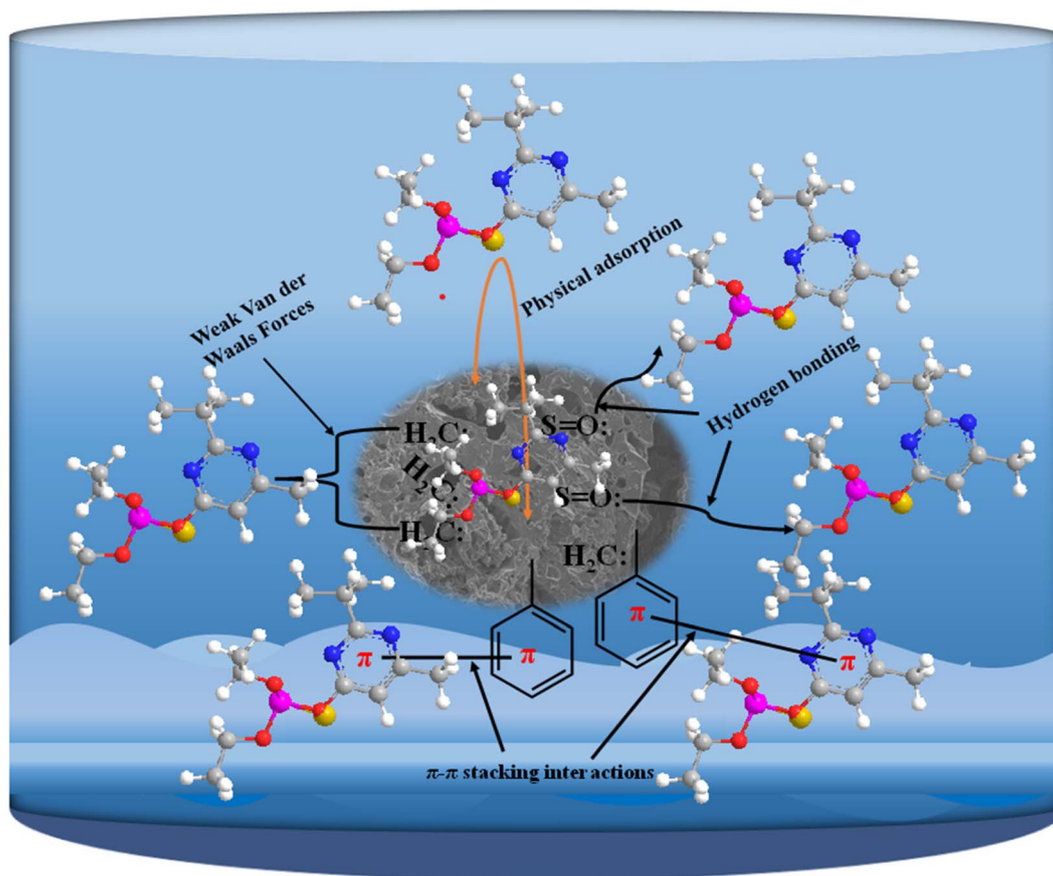


Fig. 13 A schematic illustration of adsorption mechanism of diazinon pesticide onto the prepared AC.

prepared AC, offering insights into its potential applications in environmental remediation efforts. As shown in Table 10, KAsAC sorbent showed adequate sorption uptake for diazinon removal from aqueous solution. Numerous attributes such as

cost-effectiveness, abundance, accessibility, and notable adsorption efficacy underscore the advantage of the asphalt based activated carbon over several sorbents for the removal of diazinon from aqueous solutions.

Table 10 Comparison of diazinon sorption uptake by KAsAC with other reported materials^a

Material	Initial concentration (mg L ⁻¹)	Dosage	Contact time (min)	pH	Percentage removal (%)	Adsorption capacity (mg g ⁻¹)	Ref.
Treated zeolites (MZ & ATZ)	50	0.3 g	80	6.00	NR	61.73 & 15.10	3
Coconut shell-modified biochar	1	2.0 g L ⁻¹	120	7.00	NR	10.33	4
Fe/Ni bimetallic nanoparticles	1.4	0.75 g	16	7.00	96%	0.202 045	45
Pumice	6.288	4 g L ⁻¹	30	3.00	76.3	20.65	49
Microorganism (<i>Saccharomyces cerevisiae</i>)	2.5	3.88%	1365	5.50	96%	NR	50
NH ₄ Cl-induced activated carbon	20	0.3 g L ⁻¹	30	7.00	97.5%	NR	51
Zero-valent iron supported on biopolymer chitosan	100	0.05 g	20	4.60	96.95	NR	53
Microorganism (<i>Escherichia coli</i> LMG 15862)	NR	775 mg L ⁻¹	65	6.75	96.06%	NR	54
Photocatalyst (C-TiO ₂ /g-C ₃ N ₄ nanocomposite)	15	0.1 g L ⁻¹	30	5.50	98.34%	NR	55
Zinc oxide nanoparticles loaded activated carbon	5	0.83	55	5.00	100%	NR	56
Asphalt based AC (KAsAC) Present study	200	1 g L ⁻¹	90	5.6		234.25	

^a NR = not reported.



4 Conclusion, limitations, and future research

The cost-effective and abundantly available asphalt was utilized as a starting material for the synthesis of carbon samples (AsC, NaAsAC, and KAsAC) for the adsorption of diazinon from water sources. The preparation process was optimized by Taguchi method with optimal conditions found as 600 °C activation temperature, 75 min activation time, and 1:3 activation ratio. The synthesized carbon samples were characterized using XRD, SEM, BET, FTIR, and Raman. The prepared AC samples were predominantly amorphous, porous, with improved surface area after activation, and $-\text{CH}_2-$, $\text{S}=\text{O}$ functional groups. The prepared carbon samples were employed in the removal of diazinon pesticide with removal capacities of 147.40, 159.33, and 178.70 mg g^{-1} for AsC, NaAsAC, and KAsAC, respectively. The sample with high adsorption uptake (KAsAC) was used in a systematic statistical approach, to optimize the impact of independent variables including pH, DP initial concentration, and KAsAC sorbent dosage using CCD. A quadratic model formulated using ANOVA in RSM predicted diazinon sorption capacity under varying process conditions, with model validation indicating a robust fit with an R^2 value of 0.9909 and a significantly low p -value (<0.001). An Adsorption capacity of 234.25 mg g^{-1} was achieved at initial diazinon concentration = 200 mg L^{-1} , pH = 5.6, KAsAC dosage = 1 g L^{-1} , and adsorption time = 90 min. Exploring kinetic models showed that adsorption of diazinon was well described by PSO model. Overall, the findings underscored the favorable, spontaneous, and endothermic nature of the diazinon adsorption process by KAsAC, positioning it as a potentially effective sorbent for the removal of pesticide from aqueous solutions.

While this study successfully demonstrated the conversion of waste asphalt into an AC material for the efficient removal of DP, there were some limitations encountered. These include the scalability of the production process, the variability in asphalt composition due to factors such as road type, traffic load, climate conditions, and maintenance history may lead to inconsistent quality and performance of the activated carbon. Therefore, future research could explore the economic viability of the process such as cost of asphalt recovery, purification, activation, and product testing. Additionally, multiple applications for various contaminants and the long-term stability and reusability of the AC should be established for large-scale implementation. Addressing these aspects will further enhance the applicability and effectiveness of the developed technology.

Author contributions

Robert O. Gembo: conceptualization, methodology, data collection, data analysis, data curation, writing – original draft; Sebusi Odisitse: project supervision and writing – review & editing; Titus A. M. Msagati: writing – review & editing; Cecil K. King'onde: conceptualization, project supervision, funding acquisition, resources, and writing – review & editing.

Conflicts of interest

The authors affirm that they do not have any discernible competing financial interests or personal relationships that might have potentially impacted the integrity of the research presented in this manuscript.

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