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# Removal of methyl violet 2B dye from aqueous solution using a magnetic composite as an adsorbent

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#### Abstract

Despite the important role of the textile industry in the global economy, its effluents generate deep concern with regard to treatment and disposal. Adsorption is a promising technique for the removal of textile color effluent at relatively lowcost and with satisfactory efficiency. This study aimed to evaluate the adsorption capacity of a halloysite-magnetite-based compositein the removal of methyl violet 2B cationic dye. Afterpreparation, the obtained composite was characterized by applying several instrumental techniques, includingX-ray diffraction, scanning electron microscopy, energy dispersive spectroscopy and determination of pH at the point of zero charge. The influences ofstirring rate, adsorbent mass, pH,initial concentration ofdyeand contact time on the adsorption process werealso studied. The adsorption capacityof the composite was then investigated in a temperature-controlled batch system. The experimental results showed that the adsorption kinetics were better described using a pseudosecond-order model. Regarding the adsorption equilibrium, the experimental results suggest that both the Langmuir and Freundlich models were applicable. The thermodynamic data showed that dye adsorption onto the composite was spontaneous and endothermic and occurred byphysisorption. The employed composite could also be regenerated at least four times using a 1.0 mol L<sup>-1</sup>solution of NaOH as an eluent and was shown to be a promisingadsorbentfor the removal of cationic dyes.

**Keywords:**halloysite-magnetite-based composite; methyl violet 2B;adsorption; kinetics; thermodynamics

# **1** Introduction

Thefuture of Earthandits biodiversityisa common concern foreveryone.Rapid technologicaland industrial developmentin recent decadeshas led to anumber of environmentalproblems, particularly with regardto the pollutionof water resources[1].Synthetic dyes areone of themajor classes ofpollutants responsiblefor imbalancesin aquatic ecosystems; their complexmolecularstructureskeeps themrelatively stable, making them difficultto remove[2].

Color is usually the first contaminant to be recognized in wastewater because a very small amount of synthetic dye in water (<  $1.0 \text{ mg L}^{-1}$ ) is typically highly visible, affecting the aesthetic merit, transparency and gas solubility of water bodies. These dyes absorb and reflect sunlight, thereby interfering with the growth of aquatic species and hindering photosynthesis. Additionally, they can have acute and/or chronic effects on organisms, depending on their concentration and exposure time[3].

Methyl violet (MV) is particularly important because of its broad applications in textiles, paints and print inks[4, 5].MV isoften usedin the dyeingofcotton, silk, paper, bamboo, strawand leather[6]. In biomedical fields, MV is the active ingredient in Gram's biological stain for bacteria classification[7-10]. It can sometimes also be used as a moderateclass disinfectant but has beenfound to be poisonous to most animals. The inhalation of MV may cause irritation to the respiratory tract, whereas ingestion typically causes irritation to the gastrointestinal tract[11].

Dyes can be classified according to their chemical structure or method of application. MV belongs to a class of intensely colored organic compounds called triphenylmethane dyes due to the presenceofthreearylgroups, each of which is bonded to a nitrogen atomthat

3

interacts with one or two methyl groups. Moreover, MV is a basic dyebecause of the presence of a positive charge on the aminogroup [12].

MVis a mixture of tetramethyl, pentamethyl and hexamethyl pararosaniline chlorides[13]. This last chemical compound is also known as *crystal violet* or MV 10B and is the primary representative of these three dyesreported in the literature[14-19]. Conversely, in MV 2B,there is a predominance of the pentamethylated compound[13]. Due to this difference, the particular shade of MV differs, depending on the amount of each component present.

**Fig. 1**showsthe chemical structures of MV 2B and 10B. The small structural difference observed(the presence of an additionalmethyl group in MV 10B) is responsible for the different physical and chemical properties of the two dyes, which include the absorption maximum the visible region, solubility, staining intensity and melting temperature [13].



Fig. 1.Chemical structures of MV (a)2B and (b)10B.

A wide range of technologies has been developed for the removal of synthetic dyes from waters and wastewaters to decrease their environmental impacts. These include physical methods, such as membrane filtration processes and sorption techniques;chemical methods, such as coagulation, electroflotation, advanced oxidation processes and irradiation; and biological methods, such as aerobic and anaerobic microbial degradation and the use of enzymes[3]. Although chemical and biological methods are effective at removing dyes, they require specialized equipment and are usually energy intensive; in addition, large amounts of by-products are often generated. Conversely, the physical methodsare generally effective at removing dyes without producing unwanted by-products [20, 21].

With regard to the physical methods, sorption of synthetic dyes on inexpensive and efficient solid supports has been considered a simple and economical process for the removal of dyes from water and wastewater, producing high quality water; this process is an attractive alternative for the treatment of contaminated waters, particularly where the sorbent is inexpensive and does not require a pre-treatment step before its application. Sorption is superior to other techniques for water reuse in terms of initial cost, flexibility and simplicity of design, and ease of operation [3].

Activated carbon is the most widely used sorbent, and it has excellent sorption properties for a considerable number of synthetic dyes. However, the preparation of carbon sorbents is generally energy intensive, making commercially available products relatively expensive. Because a large amount of carbon sorbent is needed to remove the dye from a large volume of effluent, high cost can hinder its application [22]. In addition, the technology for manufacturing good-quality activated carbon is not fully available in developing countries. This limitation has prompted increasing research interest in the production of low-cost alternatives to activated carbon from a range of carbonaceous and mineral precursors[3].

According to Bujdák[23], future investigations could be directed towards the optimization of wastewater treatment processes using suitable clay templatesand nanomaterials. In this way, adsorbents with magnetic properties have been extensively studied by many researchers. Zhao et al. [24] have reported, for example, a novel magnetic

hydroxamic acid modified polyacrylamide/Fe<sub>3</sub>O<sub>4</sub> adsorbent (M-PAM-HA) for the removal of Cd(II), Pb(II), Co(II) and Ni(II) ions from aqueous solutions. Ai et al. [25] and Sun et al. [26] have developeda xylan/poly(acrylic acid) magnetic nanocomposite hydrogel adsorbent and montmorillonite/CoFe<sub>2</sub>O<sub>4</sub> magnetic composite, respectively, for removalof methylene blue from wastewater.

The novelty brought by these adsorbents the improvement in accelerating separation speed, thereby enhancing water treatment efficiency. As a result, solid phase can be easily recovered by the application of an external magnetic field[27-29]. Moreover, adsorbents with magnetic properties produce no contaminants during wastewater treatment. As main disadvantage, most commercially magnetic particles are expensive and cannot be applied to large-scale processes, but magnetic modification of low cost adsorbents could lead to materials suitable for biotechnology and environmental applications[29]. Research hashown that it is possible to produce promising adsorbents in this field, creating new options for textile industry effluent mitigation[30-36].

Therefore, the aim of this work was to prepare, characterize and evaluate the removal of MV 2B from aqueous solution using a halloysite-magnetite-based composite (HNT–Fe<sub>3</sub>O<sub>4</sub>) as an adsorbent. The adsorption kinetics, equilibrium and thermodynamics of MV 2B onto this composite were also investigated.

# **2** Experimental

#### 2.1 Materials and cleaning procedure

MV 2B ( $C_{24}H_{28}N_3Cl$ , Color Index 42535) was purchased from VetecQuímica Fina Ltda (Rio de Janeiro, RJ, Brazil). HNT and iron salts (FeCl<sub>3</sub> . 6 H<sub>2</sub>O and FeSO<sub>4</sub> . 7 H<sub>2</sub>O) were

acquired from Sigma-Aldrich (São Paulo, Brazil). The chemicals used in this work were of analytical quality and were used without further purification. All aqueous solutions, including solutions of KNO<sub>3</sub>, KOH, HNO<sub>3</sub>, HCl and NaOH (Merck, São Paulo, SP, Brazil), were prepared using deionized water (18.2 M $\Omega$  cm resistivity) obtained from a Millipore Milli-Q UV Direct-Q 3 UV system (Darmstadt, Germany).

All laboratory glassware used in the preparation of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite and in the adsorption experimentswas first washed with tap water and then soaked in an alkaline Extran solution (15% v/v) for 24 h. After this period, they were washed thoroughly with tap water and then soaked in a solution of HNO<sub>3</sub> (5% v/v) for 24 h. Finally, the material was rinsed with deionized water and left to dry at room temperature.

#### 2.2 Preparation of HNT-Fe<sub>3</sub>O<sub>4</sub> composite

The HNT-Fe<sub>3</sub>O<sub>4</sub> composite was prepared using the chemical precipitation method [31]. Initially, 6.25 g of HNT was added into a 500 mL solution of 7.275 g of FeCl<sub>3</sub> . 6 H<sub>2</sub>O and 3.75 g of FeSO<sub>4</sub> . 7 H<sub>2</sub>O. The suspension was thenrefluxed for 3hin aglycerin bathat 120°C undernitrogen gas flow. When the mixed suspension was cooled to 50°C, NaOH solution (4.0 molL<sup>-1</sup>) was addeddropwise with vigorousmagnetic stirring. ThepH of the final suspension was controlled to be in the range of 9.0–10.0. The mixture was aged at 50°C for 2.5 h and then filtered and washed with deionized water repeatedly. At the end of this procedure, the obtained composite was dried at 60°C for 24 h.

#### 2.3 Characterization of HNT-Fe<sub>3</sub>O<sub>4</sub> composite

The crystalline phases in the HNT and HNT-Fe<sub>3</sub>O<sub>4</sub> composite were characterized by X-ray powder diffraction using a Shimadzu XRD-6000 diffractometer (Tokyo, Japan). The samples were scanned at room temperature in reflection mode using incident CuK<sub> $\alpha$ </sub> radiation ( $\lambda = 1.5405$  Å) at a step width of 0.05° min<sup>-1</sup> from 2 $\theta = 2^{\circ}$  to 80°.

The morphology of the HNT and HNT-Fe<sub>3</sub>O<sub>4</sub> composite was examined by scanning electron microscopy (SEM) using a ShimadzuSSX-550 (Tokyo, Japan) operating at an accelerating voltage of 10 kV. Before analysis, the samples were sputter-coated with a thin gold layer for 2.5 min. The qualitative analysis of iron in the HNT-Fe<sub>3</sub>O<sub>4</sub> compositewas performed by energy dispersive spectroscopy (EDS) of the same sample used in the SEM analysis and in the same microscope, operating at an accelerating voltage of 15 kV.

Nitrogen adsorption-desorption experiments were performed in order to obtain information on the specific surface area of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite. The measurements were carried out at 77 K using a Quantachrome Instruments Nova 2200e Surface Area Analyzer (Boynton Beach, FL, United States of America). About 100 mg of the sample was outgassed at 383 K for 24 h under a residual pressure of 0.01 Pa. Data were recorded for relative vapour pressures from 0.05 to 0.99. The specific surface area was determined using the Brunauer-Emmet-Teller (BET) equation based upon the cross-sectional area of nitrogen  $(0.163 \text{ nm}^2)$  at 77 K.

The point of zero charge  $(pH_{PZC})$  of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite was determined by adding 50 mL of 0.01 mol L<sup>-1</sup> KNO<sub>3</sub> with a previously adjusted initial pH (i.e., pH<sub>i</sub> values of the solutions were adjusted from 2.0 to 11.0 with 0.10 mol L<sup>-1</sup> of KOH or HNO<sub>3</sub>) to several 150 mL Erlenmeyer flasks. Each Erlenmeyer contained 0.10 g of the magnetic composite and was securely capped immediately with a sheet of aluminum foil. Then, the Erlenmeyers were shaken at a stirring rate of 400 rpm using a thermostated water bath shaker at a constant temperature of 25°C; the samples were then allowed to equilibrate for 24 h. At the end of this experimental procedure, the suspensions were filtered, and the final pH (pH<sub>f</sub>) values of the solutions were recorded using a Digimed DM-20 pHmeter (São Paulo, SP, Brazil). The value of  $pH_{PZC}$  is the point where the curve of  $\Delta pH$  (pH<sub>i</sub> – pH<sub>f</sub>) versus pH<sub>i</sub> crosses the line equal to zero[37].

# 2.4 Adsorption experiments

The adsorption experiments were performed in a Dist multiple spindle stirrer (Florianópolis, SC, Brazil) using 25 mL of dye solution at 25°C for 6 h. Subsequently, the Erlenmeyers were exposed to a magnetic field using a 3000 G magnet for 5 min to separate the magnetic composite from the aqueous solutions.

The initial and final concentrations of the MV 2B remaining in the solutions were determined by visible spectrophotometry using a Thermo Scientific Evolution 60 spectrophotometer (Waltham, MA, United States of America) fitted with a quartz cellwith a path length of 1.0 cm. Absorbance measurements were made at 582 nm (i.e., the maximum wavelength of MV 2B dye). A calibration curve was constructed withdye concentrations ranging from 1.0 to 15.0 mg  $L^{-1}$ .

The removal percentage (%*R*, %), the amount of MV 2B adsorbed at time *t* ( $q_t$ , mg g<sup>-1</sup>), and the adsorption capacity at equilibrium ( $q_e$ , mg g<sup>-1</sup>) were calculated by applying Equations (1), (2) and(3), respectively[31, 33]:

$$\mathscr{H} = \frac{(C_0 - C_t)}{C_0} \times 100$$
(1)  
$$q_t = \frac{(C_0 - C_t)}{m} \times V(2)$$

$$q_e = \frac{(C_0 - C_e)}{m} \times V(3)$$

where  $C_0$ ,  $C_t$  and  $C_e$  are the concentrations of MV 2B initially, at time *t* and at equilibrium, respectively (mg L<sup>-1</sup>);*m* is the weight of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite (g); and*V* is the volume of the MV 2B solution (L).

# **3** Results and discussion

3.1 Preparation and characterization of HNT-Fe<sub>3</sub>O<sub>4</sub> composite

# 3.1.1 X-ray diffraction (XRD)

The XRD patterns of the HNT and HNT-Fe<sub>3</sub>O<sub>4</sub> composite are shown in **Fig. 2**. The new diffraction peaks at 35.45° (3 1 1), 43.20° (4 0 0), 57.15° (5 1 1) and 62.75° (4 4 0) can be identified as Fe<sub>3</sub>O<sub>4</sub>, while the other diffraction peaks at 12.29° (0 0 1), 19.78° (0 2 0) and 24.99° (0 0 2) can be indexed to HNT[31, 33, 38]. These results indicated that magnetic particles were successfully anchored onto the surface of the clay.



Fig. 2. XRD patterns of the HNT and HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

# 3.1.2Scanning electron microscopy (SEM) and energy dispersive spectrometry (EDS)

Fig. 3shows the SEM images and EDS spectra for the HNT and the HNT-Fe<sub>3</sub>O<sub>4</sub> composite. As shown in the Fig. 3(a), HNT nanotubes are of different sizes with open ends. Conversely, the SEM image of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite (Fig. 3(b))shows that the Fe<sub>3</sub>O<sub>4</sub> nanoparticles are adhered to the surface of the HNT nanotubes; this is corroborated by the XRD analysis.

EDSspectrafor both materials(**Figs.** 3(c)and3(d)) confirmed the presenceofaluminum, oxygen and siliconelements, whichare characteristic of aluminosilicates, as well as ironin the compositedue to the covering of Fe<sub>3</sub>O<sub>4</sub>nanoparticles. Inboth samples, goldwasalsodetecteddue to the thin layer deposited to perform these analyses.



**Fig. 3.(a)** SEM image of HNT; **(b)** SEM image of HNT-Fe<sub>3</sub>O<sub>4</sub> composite; **(c)**EDS spectrum of HNT; and(d) EDS spectrum of HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

# 3.1.3 BET analysis

The adsorption and desorption isotherms of nitrogen gas revealed classical features for both non swelling clay mineral and iron oxides. The BET calculation applied to the nitrogen desorption isotherm gave a total specific surface area for the HNT-Fe<sub>3</sub>O<sub>4</sub> composite of 46 m<sup>2</sup> g<sup>-1</sup>, an intermediate value between the starting materials for which their specific surface area was reported to be in the range of 50-60 and almost  $9m^2$  g<sup>-1</sup>for kaolinitehalloysite clay minerals [39]and magnetite particles[40], respectively.

#### $3.1.4 pH_{PZC}$ determination

The  $pH_{PZC}$  curve of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite is shown in **Fig. 4**. The  $pH_{PZC}$  is the pH value where positive and negative charges are equal on the surface of a material, which makes it possible to describe the properties of the resulting electrical double layer interfaces [41].

For pH values lower than  $pH_{PZC}$ , the adsorbent presents a positive surface charge that favors the adsorption of negatively charged compounds, such as anionic dyes. In the situation when the pH is greater than  $pH_{PZC}$ , the adsorbent presents a negative surface charge that favors the adsorption of positively charged compounds, such as MV 2B [37]. Thus, it can be concluded that the adsorption of this dye by HNT-Fe<sub>3</sub>O<sub>4</sub> composite should be favored in media with a pH greater than 5.02.



**Fig. 4.**Curve of the point of zero charge  $(pH_{PZC})$  of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite. Note: the horizontal dotted line shown is used only as a visual guide.

#### 3.2 Effect of experimental conditions on the adsorption process

To evaluate the effect of the experimental conditions on the adsorption process (e.g., stirring rate, adsorbent dosage, initial pH, initial dye concentration and contact time), the results were compared in terms of % R and  $q_t$ , which were calculated using Equations (1) and (2), respectively. For this calculation, it was necessary to know the value of the final dye concentration at the endof each experiment as obtained using acalibration curve (r=0.9997).

#### 3.2.1 Stirring rate

The experiments used to evaluate the influence of the stirring rate on the adsorption processwere conducted in a Dist multiple stirrers pindle at 25°C with an initial weight of 0.15 gof composite in 25mL of 90 mgL<sup>-1</sup> of MV2B(pH~4.2) for 6 h. The effect of the stirring rate on the dye removal percentage is shown in **Fig. 5**.



Fig. 5. Effect of the stirring rate on the MV 2B removal percentage.

As shown, %*R* increases as the stirring rate of the system increases, indicating the occurrence of a stronger interaction between the composite and the dye. Furthermore, a higher removal percentage occurs when the stirring rate of the system is equal to 700 rpm. For higher values, the %*R* begins to decrease; the associated high rate of turbulence in this case might promote the entrainment of the MV 2B particles before they can interact with the composite. The decrease in percentage removal may also be explained by the fluid dragging HNT-Fe<sub>3</sub>O<sub>4</sub> particles due to the high rate of the system, which also affects the interaction of the composite with the dye[42].

# 3.2.2 Adsorbent dosage

The experiments performed to evaluate the influence of the adsorbent mass on the adsorption process were also conducted in a Dist multiple stirrers pindle at 25°C with 0.025, 0.05, 0.10, 0.15, 0.20 and 0.25 g of composite. A volume of 25mL of MV2B at a concentration of 90 mg L<sup>-1</sup> was used, with the stirring rate previously optimized at apH value of approximately 4.2 for 6 h. The adsorbent dosage effect on the MV 2B removal percentage is shown in **Fig. 6**.



Fig. 6.Effect of the adsorbent dosage on the MV 2B removal percentage.

The results show that %*R* increases as the adsorbent dosage increases. It was found that the MV 2B removal percentage increased from 18.3% to 94.7% when the adsorbent dosage was increased from 0.025 to 0.25 g. This increase likely occurreddue the increment of the number of adsorption sites available for adsorption [43]. However, *q<sub>t</sub>* shows the opposite behavior of the removal percentage, likely due to the increase in the adsorbent mass at a determined dye concentration; fixed volume also leads to unsaturated adsorption sites during the adsorption process. Additionally, a high adsorbent mass may be favorable to particle aggregation, which would lead a decrease in total surface area of the adsorbent and an increase in diffusional path length [43]. In addition, this reduction of *q<sub>t</sub>* can be mathematically explained by combining Equations (1) and (2), as shown in Equation (4):

$$q_t = \frac{\% R C_0 V}{100 \ m} \tag{4}$$

where %R is the removal percentage (%),  $C_0$  is initial concentration of MV 2B (mg L<sup>-1</sup>), *m* is the weight of HNT-Fe<sub>3</sub>O<sub>4</sub> composite (g), *V* is the volume of MV 2B solution (L), and  $q_t$  is the amount of MV 2B adsorbed at time *t* (mg g<sup>-1</sup>).

According toEquation (4), the amount of dye adsorbed at a time  $t(q_t)$  and the mass of adsorbent (*m*) are inversely proportional. For fixed values of the dye percentage removal (%*R*), the treated volume of dye solution (*V*) and the initial dye concentration (*C*<sub>0</sub>), the increase of adsorbent mass leads to a decrease in  $q_t$  values [44].

Based on these results, the value of adsorbent mass should be chosen based on the intersection of the curves %R and  $q_t$  with the justification that this location will produce the most cost-effective system [31]. However, for the adsorption experiments, an adsorbent amount of 0.15 gwas chosen because this dosage results in a satisfactory removal percentage (i.e., above 85%) and  $q_t$  values (e.g., 12.18 mg g<sup>-1</sup>), particularly when compared to 0.10 g, which typically yields a removal percentage of approximately 60% and a  $q_t$  of 12.61 mg g<sup>-1</sup>. The use of an amount greater than 0.15 g does not show significant advantages in terms of percent removal with an increase of approximately 5%.

# 3.2.3 Initial pH

To optimize pH, experiments were performed with a range of values between 3.0 and 11.0 in increments of unity, adjusted by adding HCl or NaOH solutions. The concentration of the MV 2B solution was set to 90 mg  $L^{-1}$  by applying the stirring rate and adsorbent dosage previously determined to be optimal (i.e., 700 rpm and 0.15 g, respectively)for 6 h at 25°C.The effect of the initial pH on the MV 2B removal percentage is shown in **Fig. 7**.



Fig. 7.Effect of initial pH on the MV 2B removal percentage.

As shown, %*R* increases as pH increases. In an acidic medium, little interaction occurs between the adsorbent and adsorbate. This behavior is expected because the pH<sub>PZC</sub> value of the adsorbent is lower than 7.0. Under these conditions, the HNT-Fe<sub>3</sub>O<sub>4</sub> composite is positively charged (i.e., with the same net charge as the dye), resulting in electrostatic repulsion between the composite and the dye. For pH values lower than 3.2, MV 2B begins to change color towards yellow, which ultimately changes the maximum absorption in the visible region. All of these factors result in a non-optimal scenario for adsorption.

In systems with higher pH(e.g., 9.0 to 11.0), there is a large increase in %*R*,includingvalues near100% at pH11.0. According toDuynsteeandGrunwald[45], in the pH rangebetween9.0and 13.0,a reactionbetweenthe dyeandOH<sup>-</sup>ions occurs. These ions attack the central carbon atom of the MV 2B molecule, forming a base carbinol with a different molecular geometry.Theresulting compoundhas nocolor,which explains the high %*R*values in thispH range. Despite thenearly complete removal ofcolor, as shownby the high%*R* value, the dye still remains in solution. When these carbinol bases form, the absorption peakshifts to the UV region, making it difficult to study the adsorption process of the composite because the calibration curveused to quantify the dyewas constructed using the maximum absorption of the dyeat 582nm. Finally, the OH<sup>--</sup>ion scompete with the adsorbent in the interaction with MV2B molecules.

Based on these observations, the best conditions for adsorption are in the range of pH values between 5.0 and 9.0; this is the range where removal is improved without a loss of colordue to chemical reactions. Because of the differences in percentage removal vary by only 5%, an eutral pH value was adopted for this study. Furthermore, at this pH value (which is higher than the  $pH_{PZC}$  of the HNT-Fe<sub>3</sub>O<sub>4</sub> composite) the adsorbent surface is now completely negatively charged and, as a result, it can better interact with the MV 2B molecules (positively charged) by electrostatic interactions favoring the adsorption process (**Fig. 8**).



Fig. 8.Electrostatic interactions between MV 2B molecules and HNT-Fe<sub>3</sub>O<sub>4</sub> composite at pH values higher than the  $pH_{PZC}$  of the adsorbent.

#### 3.2.4 Effect of initial dye concentration and contact time

To optimize the initial concentration of dye and contact time on the adsorption process, experiments wereconducted with 25 mLofMV2Bat concentrations of 60, 90,120, 150 and 180 mgL<sup>-1</sup> over time intervalsof 10, 20, 40, 60, 90, 120, 150, 180, 240 and 360 min. The stirring rate, adsorbent dosage and initial pH used were those previously optimized at 25°C.

As shown in Fig. 9,  $q_t$  increases as the initial concentration of MV 2B in solution increases; however, these values tend to be constant after a certain period of time. The increase in  $q_t$  in the experiments that contained higher initial concentrations of dye can be explained by the existence of a higher concentration gradient of MV 2B, which increases the diffusive contribution of the mass transfer process [31, 46]. This increase in the values of  $q_t$  at equilibriumdoes follow aproportionality, presenting shorter not intervalsas theinitial concentration of MV2Bincreases, particularlyat higherconcentrations. According toGusmãoet al.[47], this is an indication that theremoval percentage decreases as the initial concentration of MV2Bincreasesdue to the greateramount ofdye presentin the bulk solution. The mass of HNT-Fe<sub>3</sub>O<sub>4</sub>composite used (i.e., 0.15 g)islikely not sufficient to promotea highpercentageremovalforhigher concentrations, as discussed in section 3.2.2; this is alsoin withHameed[48].Theadsorbeddye moleculeslikely accordance exert anelectrostatic repulsiononthose stillin solution, increasing resistance to mass transfer.



Fig. 9.Effect of the initial dye concentration and contact time on the adsorption process.

Regarding the influenceof thecontact time,  $q_i$  is shown to increase as the process evolvesuntil equilibrium is reacheddue to the longer period, so that the interaction between the soluteand composite occurs, increasing the amount of adsorbed MV2B [31]. However, the curves shown in **Fig. 9** show that the removal is faster in the initial minutes of the process because more adsorption sites are available during this period. As these sites become progressively occupied, adsorption becomes slower and less efficient [37]. Another important aspect to note is that as the initial concentration of MV2B increases, more time is required for equilibrium to be reached. The reason for this phenomenon is that a higher concentration of adsorbate in the solution increases the electrostatic repulsion between molecules present in the medium, increasing the diffusive resistance to mass transfer with in the solution but not in the interface as previously discussed; this results in a slower process [48]. To identify the mechanism of adsorption, the potential removal rate and the control of the steps involved in the process, two kinetic models (i.e., the pseudo-first- and pseudo-second-order) can be adopted [46]. Kinetics experiments were performed at the same conditions as described in the section 3.2.4. The pseudo-first- and pseudo-second-order kinetic models are shown in **Fig.10**.



Fig. 10.Kinetics experimental results: (a) pseudo-first- and (b) pseudo-second-order.

Thepseudo-first order kinetic modelcanbe expressedbyEquation(5)[49]:

$$\ln(q_e - q_t) = \ln q_e - k_1 t(5)$$

where  $k_1$  is the pseudo-first-order rate constant (min<sup>-1</sup>). The  $k_1$  and  $q_e$  parameters can be determined from the slopeandintersection of the linesoflog ( $q_e$ - $q_t$ ) versus t. These values are summarized in Table 1.

The coefficient of correlation (r) for the pseudo-first-order model ranged from 0.8734 to 0.9566forinitial concentrationsof MV2Bof60, 90, 120, 150 and 180 mg L<sup>-1</sup>. The experimental values of  $q_e(q_{eexp}, \text{mg g}^{-1})$  diverged from the calculated values  $(q_{e cal}, \text{mg g}^{-1})$ , as shown in Table 1. This result indicates that he adsorption of the MV 2B by the compositedoes not follow thepseudo-first-orderkinetic model(i.e., the process is nota phenomenon controlled bydiffusion;othersteps areresponsible controlling for thekinetics observed).Therate constant  $k_1$  can be interpreted as a temporal scale factor, which is intended to indicate therate at which the system reaches equilibrium. Higher values of  $k_1$  indicate less time to reach that condition and vice versa. This parametermay or may notbe dependent on the experimental conditions of the process[49]. As shown in Table1, the adsorption process does not reachequilibriumin a short periodaccording to the pseudo-first-order model because the values  $ofk_1$  are on the order of  $10^{-2}$ , with the greatest difference occurring in the experiments in which the initialdye concentrations are 90 and 120mg L<sup>-1</sup>. This finding also indicates that  $k_1$  does not depend on the initial concentration of MV 2B parameter.

The pseudo-second-order kinetic model can be expressed by Equation (6)[50]:

$$\frac{t}{q_t} = \frac{1}{k_2 q_e^2} + \frac{1}{q_e} t(6)$$

where  $k_2$  is the pseudo-second-order rate constant (g mg<sup>-1</sup> min<sup>-1</sup>). Analogous to thepseudofirst-orderkinetic model, the  $k_2$  and  $q_e$  parameters were determined from the linear and angular coefficients of the equations formed by regressing  $t/q_t$  versus t, respectively. The real value at different concentrations of the linear portions greater than 0.9997, which suggests that the adsorption of the MV 2B follows the pseudo-second-order kinetic model wherein the rate of chemical interactions between the dye and the sites of adsorbent is the step that controls the overall kinetics of the process. Furthermore, the calculated data ( $q_{e,cal}$ ) are consistent with the experimental data ( $q_{e,exp}$ ), as shown in Table 1.

#### Table 1

	Initial dye concentration (mg L <sup>-1</sup> )					
	60	90	120	150	180	
$q_{e, exp} (\mathrm{mg g}^{-1})$	9.12	12.18	15.05	16.72	18.23	
Pseudo-first-order						
$q_{e, cal} (\mathrm{mg \ g}^{-1})$	1.39	3.65	5.79	4.56	3.81	
$k_{l} \times 10^{2} ({\rm min}^{-1})$	1.57	1.02	2.09	1.34	1.46	
r	0.8734	0.9021	0.9566	0.9315	0.9079	
Pseudo-second-order						
$q_{e, cal} (\mathrm{mg \ g}^{-1})$	9.16	12.29	15.29	16.69	18.37	
$k_2 \times 10^2 \text{ (g mg}^{-1} \min^{-1}\text{)}$	5.24	1.11	1.24	1.10	1.55	
r	0.9999	0.9997	0.9999	0.9999	0.9999	

Kinetics parameters for MV 2B adsorption by HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

Thekinetic constant $k_2$  has the same physical meaning  $k_1$  and mayor may not also depend on the experimental conditions of the adsorption process[49]. In this case, the results shown in Table 1 demonstrate that  $k_2$  does not depend on the initial concentration of MV 2B parameter either. The results of  $k_2$  are also on theorder of  $10^{-2}$ , which indicates that adsorption did not reachequilibrium within a short time period, according to the pseudo-second-

order model.Unlike whatwaspreviouslyobserved, the value of  $k_2$  differs slightlyin experiments with lowerMV2B concentrations ( $C_0 = 60 \text{ mg L}^{-1}$ ) than for the other experiments, which show a valueapproximately 4 times higher, indicating that equilibrium is attained more rapidly under these conditions.

#### 3.4 Adsorption equilibrium

Adsorption isotherm modelsare fundamentalto describing the interactive behavior between theadsorbateandadsorbent and are also important for investigating mechanisms of adsorption. In this study, equilibrium data were analyzed using the Freundlich and Langmuir isotherms. It is important to emphasize that the equilibrium experiments were performed at the same conditions of adsorption kinetics.

To evaluate which model was best suited to describe these processes, r and the error function( $F_{error}$ ) were analyzed; a lower result from the error function indicated a smaller difference between adsorption capacity calculated by the model ( $q_{cal}$ ) and the experimental ( $q_{exp}$ ).  $F_{error}$  can be expressed according to Equation (7)[51]:

$$F_{error} = \sum_{i}^{P} \left( \frac{q_{i\,cal} - q_{i\,exp}}{q_{i\,exp}} \right)^2 \tag{7}$$

where  $q_{i \ cal}$  is a value of q predicted by the fitted model; $q_{i \ exp}$  is a value of q measured experimentally; *i* indicates the values of the initial dye concentration of the experiments (60, 90, 120, 150 and 180 mg L<sup>-1</sup>); and *P* is the number of experiments performed.

The Freundlich and Langmuir isotherm linearized models are shown in Fig. 11.



Fig. 11.(a) Freundlich and (b) Langmuir isotherm linearized models.

The empirical Freundlich equation is applicable to adsorption on heterogeneous surfaces, where the interaction between the adsorbed molecules is not limited to the formation of a monolayer. The Freundlich constant ( $k_F$ ) is related to the adsorption capacity of the adsorbent: the higher the value, the greater the affinity for the adsorbate. The empirical parameter 1/n is related to the strength of adsorption, which varies with the heterogeneity of the material. When the values of 1/n are between 0.1 and 1.0, the adsorption process is considered favorable [46]. The linearized form of the Freundlich equation is expressed according to Equation (8)[52]:

$$\ln q_e = \ln k_F + \frac{1}{n} \ln C_e (8)$$

where  $k_F$  is the Freundlich constant (L g<sup>-1</sup>) and 1/n is a dimensionless empirical parameter. The  $k_F$  and 1/n values were determined from the linear and angular coefficients of the equations formed by regressing ln  $q_e$  as a function of ln  $C_e$ , respectively. These values, along with *r* and  $F_{error}$ , are shown in Table 2. In general, the experimental data are shown to be well adjusted to the Freundlich model because the *r* value is greater than 0.98, and  $F_{error}$  is near 0.0022. The value of 0.2696 for the parameter 1/n is lower than 1.0, suggesting that the adsorption is favorable under the conditions proposed by the Freundlich model.

The Langmuir model is based on the assumption of monolayer adsorption on a structurally homogeneous adsorbent, where all sorption sites are identical and energetically equivalent [42]. The linearform of theLangmuirequation canbe expressedaccording toEquation (9)[52]:

$$\frac{C_e}{q_e} = \frac{C_e}{q_m} + \frac{1}{q_m k_L}(9)$$

where  $q_m$  is the theoretical maximum adsorption capacity corresponding to monolayer coverage (mg g<sup>-1</sup>), and  $k_L$  is the Langmuir constant (L mg<sup>-1</sup>). The  $k_L$  and  $q_m$ weredetermined from the linear and angular coefficients of the equations formed by regressing  $C_e/q_e$  as a function of  $C_e$ , respectively.

To determine whether the adsorption process is favorable, a dimensionless constant separation factor  $R_L$  is defined. The adsorption process is irreversible when  $R_L$  is 0.0, favorable when  $R_L$  is between 0.0 and 1.0, linear when  $R_L$  is equal to 1.0, and unfavorable when  $R_L$  is greater than 1.0. The  $R_L$  parameter can be defined based on Equation (10)[52]:

$$R_L = \frac{1}{1 + k_L C_0} (10)$$

where  $C_0$  is the initial dye concentration (mg L<sup>-1</sup>).

The isotherm parameters for MV 2B adsorption by the HNT-Fe<sub>3</sub>O<sub>4</sub> composite are shown in Table 2. As shown, the experimental data are well adjusted to the Langmuir modelas well. The  $R_L$  values are between 0.130–0.048, confirming that the adsorption process is also favorable. Although the *r* value is greater than 0.99, its  $F_{error}$  is higher than 0.04 and, thus, higher than the one obtained for the Freundlich model. It can be inferred that analyzing the correlation coefficient of the fitting is not the only parameter that should be account for to verify the best fitting isotherm model. Thus, thesedata suggest that both models were suitably fitted [53]. Similar results were also observed by Lima et al. [51], Namasivayam and Sureshkumar [54], and Kalavathy and Miranda [55]. According to these authors, the adsorption mechanism by both monolayer adsorption and heterogeneous energetic distribution of active sites on the adsorbent surface is possible [54].

#### Table 2

Isotherm		Temperature (25°C)
	$k_F$ (L g <sup>-1</sup> )	5.835
	1/n	0.2696
Freundlich	r	0.9892
	$F_{error}$	0.0022
	$q_m(\text{mg g}^{-1})$	20.04
	$k_L$ (L mg <sup>-1</sup> )	0.112
Langmuir	$R_L$	0.130-0.048
	r	0.9955
	F <sub>error</sub>	0.0417

Isotherm parameters for MV 2B adsorption by HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

A comparison of the  $q_m$  of a few adsorbents available in the literature for removal of MV 2B in aqueous solution is given in Table 3.It is clear shown that HNT-Fe<sub>3</sub>O<sub>4</sub> composite used in this work had a moderate adsorption capacity, despite of its relatively small surface area.

# Table 3

Comparison of maximum adsorption capacity(q<sub>m</sub>) of other adsorbents for removal of MV 2B.

Adsorbent	$q_m (mg g^{-1})$	Reference
Modified cation exchange membrane	10.1	[56]
Cellulose-based wastes	10.5	[57]
Mansonia wood sawdust	16.11	[58]
HNT-Fe <sub>3</sub> O <sub>4</sub> composite	20.04	This study
Sunflower seed hull	92.59	[48]

# 3.5 Thermodynamics studies

Thermodynamics experiments were performed at 283, 293 and 303 K at an initial dye concentration of 90 mg  $L^{-1}$ . This initial concentrationwas chosenbecause it provided the best opportunity to monitor adsorption process. The other conditions used were the same as those used in the kinetics and equilibrium studies.

The values of the standard Gibbs free energy change  $(\Delta G^{\circ})$  for each temperature were calculated from the Equation(11)[59]:

 $\Delta G^o = -RT \ln k_c \qquad (11)$ 

where *R* is the universal gas constant (8.314 J mol<sup>-1</sup> K<sup>-1</sup>), *T* is temperature (K) and  $k_c$  is the equilibrium stability constant, which was calculated at each temperature by the Equation (12)[59]:

$$k_c = \frac{C_s}{C_e}(12)$$

where  $C_s$  and  $C_e$  are the equilibrium concentrations on the adsorbent and on the aqueous phase, respectively.

The enthalpy change  $(\Delta H^{\circ})$  and entropy change  $(\Delta S^{\circ})$  are calculated from the slope and intercept of the plot of  $\ln k_c \text{versus} 1/T$ , based on the Van't Hoff equation (Equation 13)[59]:

$$\ln k_c = \left(\frac{\Delta S^{\circ}}{R}\right) - \left(\frac{\Delta H^{\circ}}{R}\right) \frac{1}{T} (13)$$

The results of the thermodynamics experiments are shown in Table 4. The negative values of  $\Delta G^{\circ}$  describe the spontaneous nature of the adsorption process. As the temperature increases,  $\Delta G^{\circ}$  increase as well, indicating less driving force and hence slower adsorption capacity at higher temperatures [60].

The negative value of  $\Delta H^{\circ}$  indicates that the adsorption process is exothermic. This magnitude is related to adsorb tein teractions with the adsorb entandre arrangement of the surface during the adsorption process. It is also be served experimentally as the fraction of the surface that is covered by adsorb ate. Enthalpy change data is useful for distinguishing physisorption and chemisorption. The typical value for physisorption usually lies below 84 kJ 30

mol<sup>-1</sup>, which agrees with the value obtained for MV 2B adsorption onto HNT-Fe<sub>3</sub>O<sub>4</sub>; for chemisorption, bond strengths lie between 84 and 420 kJ mol<sup>-1</sup>. Finally, the results of  $\Delta S^{\circ}$  indicate decreased randomness at the solid/solute interface during the adsorption of MV 2B onto the HNT-Fe<sub>3</sub>O<sub>4</sub> composite [60]. This behavior (negative values of  $\Delta G^{\circ}$ ,  $\Delta H^{\circ}$  and  $\Delta S^{\circ}$ ; slower adsorption capacity at higher temperatures and physisorption) wasalso observed by others researchers thatemployed magnetic composites for removal of dyes[61-63].

# Table 4

Temperature	Thermodynamics			
(K)	$\Delta G^{\circ}$ (kJ mol <sup>-1</sup> )	$\Delta H^{\circ}$ (kJ mol <sup>-1</sup> )	$\Delta S^{\circ} (\mathrm{J} \mathrm{mol}^{-1} \mathrm{K}^{-1})$	r
283	- 4.09			
293	- 3.75	- 13.23	- 32.32	0.9744
303	- 3.44			

Thermodynamics parameters for the adsorption of MV 2B onto HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

#### 3.6 Reusability of HNT-Fe<sub>3</sub>O<sub>4</sub> composite

It is crucial to investigate the reusability of adsorbents for practical applications to address ecological and economic demands for sustainability[64]. Adsorption experiments were performed at a 700 rpm stirring rate and a pH equal to 7.0 with 0.15 g of composite in25mL of a 90 mg  $L^{-1}$  solution of MV2B at25°C for 6 h. After each adsorption experiment, the adsorbent was collected by external magnet force(3000 G) and recovered by a 1.0 mol  $L^{-1}$  solution of NaOH at 250 rpm for 3 h. Then, the HNT-Fe<sub>3</sub>O<sub>4</sub> composite was removed from the alkaline solution and placed in a 0.5% v/v acetic acid solution at the same stirring rate and

contact time. Finally, it was washed several times with distilled water, dried and reused again. The effect of recycling times of HNT- $Fe_3O_4$  composite on MV 2B adsorption performance was repeated four times.

Fig. 12 showed that the removal percentage of MV 2B was still near80% after four cycles, indicating that HNT-Fe<sub>3</sub>O<sub>4</sub>can be reused. The decrease in %*R*might be due to irreversible adsorption of the adsorbent during the regeneration process[64].



Fig. 12. Reusability cycles of HNT-Fe<sub>3</sub>O<sub>4</sub> composite.

# **4** Conclusion

In general, the method used to prepare the HNT- $Fe_3O_4$  composite was efficient, as shown by the characterization analysis, which verified that the magnetite nanoparticles are anchored onto the surface of the HNT.

The kinetics of the adsorption process wasshown to be better described by a pseudosecond-order model. The experimental results also suggest that both the Langmuir and Freundlich models were suitable to predict adsorption equilibrium. The experimental data concerning the Langmuir constant  $(k_L)$ , maximum adsorption capacity  $(q_m)$ , Freundlich constant  $(k_F)$ , and 1/n were 0.112 mg L<sup>-1</sup>, 20.04 mg g<sup>-1</sup>, 5.835 L g<sup>-1</sup> and 0.2696, respectively. Moreover, the thermodynamic data have shown that the adsorption of MV 2B onto HNT-Fe<sub>3</sub>O<sub>4</sub> composite is a spontaneous and exothermal process by physisorption.

Finally, the employed composite could also be regenerated at least four times using a  $1.0 \text{ mol } \text{L}^{-1}$  solution of NaOH as an eluent and was shown to be a promising adsorbent for the removal of MV 2B cationic dye.

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