

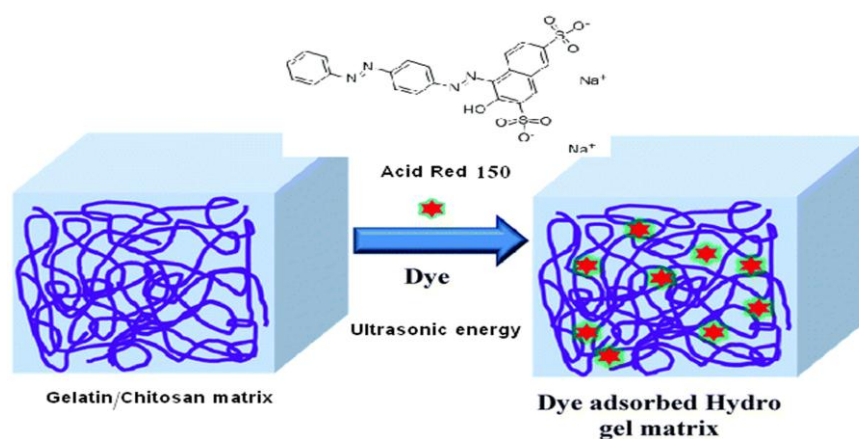
Kinetic Study of Gelatin/Chitosan Based Nanocomposites for Acid Red 150 Dye Adsorption Using Ultrasonic Energy

A.A. Haroun^{1*}, H.M. Mashaly², H.M. Helmy², M.M. Kamel²

¹Chemical Industries Research Division and ²Textile Research Division, National Research Centre, 12622 Dokki, Giza, Egypt.

THE EFFECT of acid red 150 dye (AR150) adsorption parameters onto the prepared gelatin/chitosan/laponite based nanocomposites (NC I, II, III and IV) such as dye concentration, contact time, pH, and sorbent amount using ultrasonic energy had been studied. The standard adsorption affinity through the adsorbent suitability was also carried out. The results revealed that high concentrations of AR150 and chitosan content led to an increasing in the adsorption rate. A comparison between adsorption kinetics of the prepared nanocomposites with conventional and ultrasonic techniques was investigated. The time/dye-uptake isotherms were declared the improvement of dye adsorption in the second phase. Adsorption rate, reaction half-time and standard affinities were also studied. The dye adsorption uptake using ultrasound technique was higher than that in case of the conventional one in case of all the used nanocomposites. On the other hand, in case of using ultrasonic technique, the isotherms showed high dye uptake. In other words, the best dye adsorption rate could be obtained at different nanocomposite concentrations 0.3 and 0.8 g/L in case NCIII and IV, respectively.

Keywords: Chitosan, laponite, dye removal, ultrasonic, acid red150 dye, Kinetic isotherm.



*Corresponding author email: haroun68_2000@yahoo.com

The textile industry is one of the largest producers of industrial waste water which including several pollution sources such as the different colours and pigments. This is a serious environmental problem due to its high toxicity. Consequently, this issue is currently of great interest. Because of the difficulty of treating such water by conventional methods, the needing to use of the modern techniques is urgent issue. Ultrasonic Power could enhance a great variation of physical and chemical processes, essentially because of marvel known as cavitation inside the fluid environment which developed and exploded a breakdown of microscopical bubbles. Unexpected and burst breakdown of these bubbles could create centralize high pressure, high temperature, and stun waves [1,2]. The influence of ultrasonic application on the dye adsorption can emphases in terms of regular diffusion of the dye molecules in the fluid, accelerating of the dispersion rate of dye within the composite and enhancing the reaction between composite and dye[3-11]. Recently, some attempts have been made to prepared and investigate novel nanocomposites for adsorption of some toxic materials. Moreover, the polymeric hydrogels play an important role in the treatment of waste water[12]. Because of the ultrasonic power has a high impact in the different industrial applications, this work was to investigate the ultrasonic energy effects on the adsorption kinetics of AR150 dye onto gelatin/chitosan/laponite based nanocomposites.

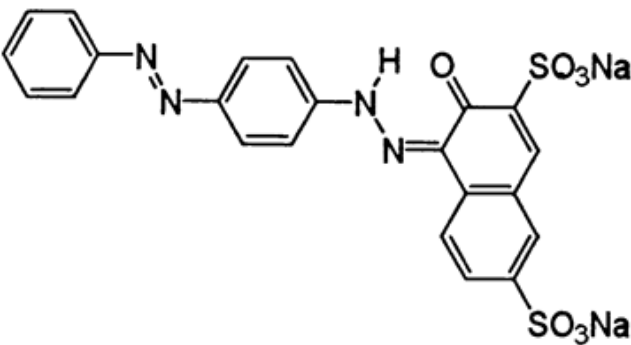
Experimental

Materials

Acid dye red 150 (Table 1) was purchased from Ciba Co. and used without further purification.

All other chemicals and reagents were used as received.

TABLE 1. Chemical structure of Acid Red 150 (AR 150).

Name of the dye	Chemical structure
Acid Red 150 (AR 150)	

Apparatus

Ultrasonic bench top cleaner bath, model 575 D was used. The used experimental setup was composed of an electrical generator operated at a frequency of 38.5 kHz and power range 100-500 W. The output power levels are from 100 up to 500 W and supplied by three transducers at the bottom of the industrial grade tank. Precise digital control of time (0–90 min), thermostatically controlled heater (ambient to 80°C) and power level and degas functions. The conventional method device is Julabo SW 20 Seelbach/Germany, V/Hz 230/50/60.

Batch adsorption experiments using ultrasonic energy

Table 2 shows the chemical composition of the prepared nanocomposites (hydrolyzed polyethylene terephthalate, gelatin and chitosan were blended in equimolar ratios in presence of different concentrations of laponite, as described previously in details [13]. The dye AR150 solution was prepared without pH adjustment (the initial dye concentration 277 mg L⁻¹). 0.3 g of the prepared NCs (I, II, III, and IV) and 50 mL of the standard solutions were stirred for a certain period of time (40 min). After filtration, the concentration of AR150 in the supernatant was analyzed by ultraviolet visible spectrophotometer (Schimadzu, Japan) with 2 nm resolution using calibration curve at λ 530 nm. The adsorption of AR150 was studied in pH range 3–9, through adjusting pH using either HCl or NaOH. The prepared NCs were equilibrated at a particular pH value for different time and temperature intervals (20, 40, 60, and 80) min, (40, 50, 60, and 70°C), and power level (100, 200, 300, 400 and 500 w) respectively. Adsorption equilibrium tests were conducted at optimum conditions. The adsorption capacity was calculated according to Lambert-Beer law and the adsorption kinetics Langmuir isotherm equations. The concentration of dye solutions was determined after reference to the respective calibration curve of acid dye.

TABLE 2. Chemical composition of the prepared Nanocomposites.

Nanocomposites	Chemical compositions (wt %)			
	Gelatin	HPET	Chitosan	Laponite
I	1	1	1	0
II	1	1	1	0.05
III	1	1	1	0.2
IV	1	1	2	0.05

Results and Discussion

Effect of pH value on AR150 adsorption ratio (%) using ultrasonic technique

Figure 1 demonstrates the influence of pH on adsorption ratio (%) of AR150 in various NCs. The pH influences essentially the adsorption limits of AR150 onto the various NCs. Generally, high uptake always happened in acidic medium rather than neutral or alkaline ones. As expected, at low pH, a lot of protons had been accessible to give protonated amino groups for chitosan and/or gelatin particles for forming $-(NH_3^+)$, that way augmenting electrostatic attraction of positive charge of adsorption positions and negative charge of dye anions which led to creating a higher adsorption of dye [14,15]. This clarification concurs with our information on pH affect. As described previously, the pH of hydrous solution assumes a critical part in acid dye adsorption on adsorbents [16-27]. (Fig 1).

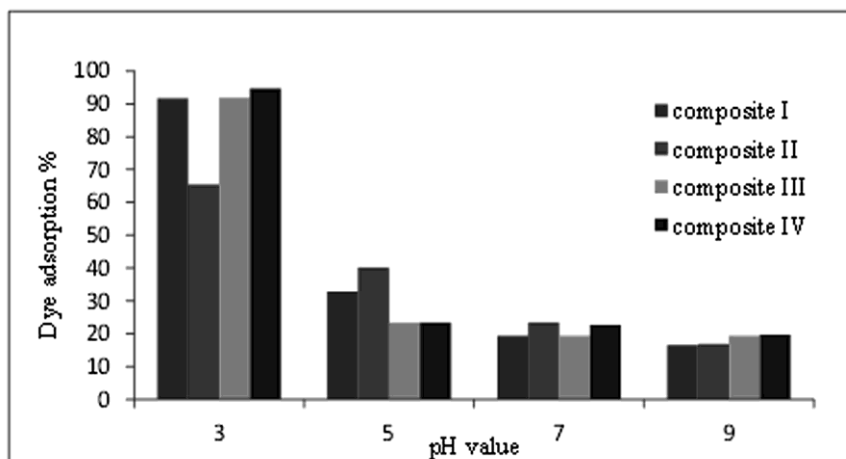


Fig.1. Effect of pH values on dyestuff sorption of prepared NCs using ultrasonic method.

Effect of time on AR150 adsorption ratio (%) using ultrasonic technique

Figure 2 illustrates the impact of time on adsorption ratio (%) of acid dye onto several NCs. The impact of time on dyestuff sorption of prepared NCs was investigated for different time periods (20–80 min). With all NCs (I, II, III and IV), the sorption ratio of dye progressively augmented within 20 to 80 min and the percentage adsorption of acid dye were stabilized at 60 min.

Effect of temperature on AR150 adsorption ratio (%) using ultrasonic technique

Figure 3 demonstrates the influence of temperature on adsorption of acid dye onto various NCs at pH 3 for 60 min using 500 mg L⁻¹ dye concentration utilizing ultrasonic technique with power level 400 Watts. The effect of temperature (40-70°C) exhibited an augmenting of dye uptake. After equilibrium, the augmentation of dye uptake showed endothermic type of adsorption [28,29].

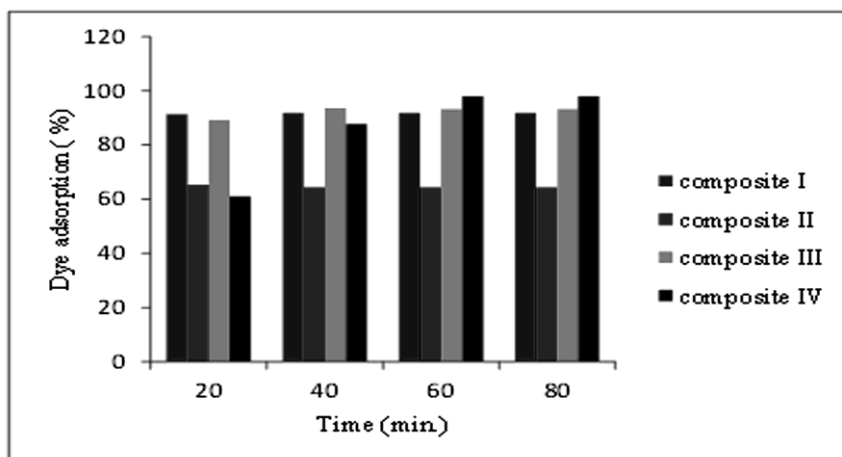


Fig. 2. Effect of time on dyestuff sorption of prepared NCs using ultrasonic method.

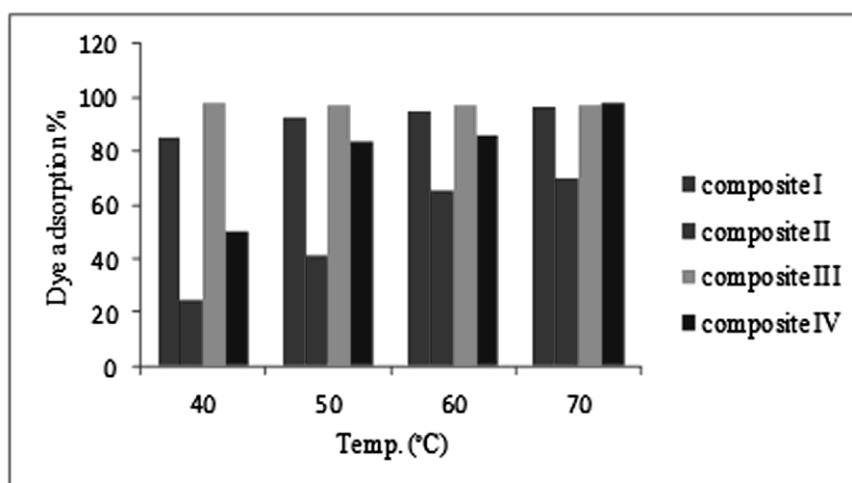


Fig.3. Effect of temperature on dye adsorption of prepared NCs using ultrasonic method.

Effect of nanocomposites dose using ultrasonic technique

Figure 4 demonstrates the adsorption ratio (%) of AR150 dye onto various prepared NCs using pH 3 at 60°C for 60 min utilizing ultrasonic technique with power level 400 Watt. The ratio augmented essentially with rising of adsorbent dose from 0.1 till 0.8g. Increasing of the adsorbent dose using constant concentration of AR150 gave more accessible adsorption positions to acid dye and along these lines augmented the range of acid dye uptake. NCs III afforded better results at 0.3 g/L, while IV afforded better results at 0.8 g/L adsorbent concentration.

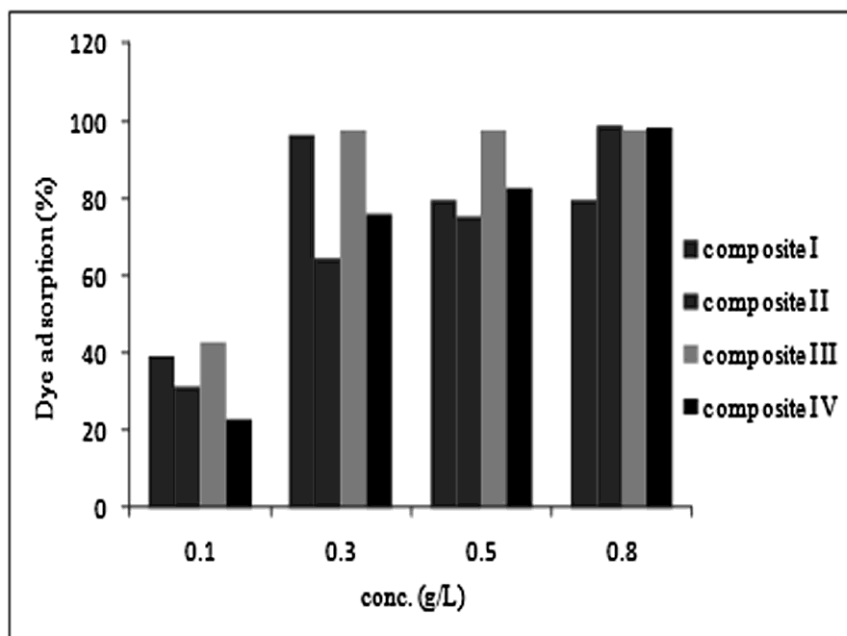


Fig.4. Effect of nano composite concentration using ultrasonic method.

Effect of AR150 concentration on the dye adsorption using ultrasonic technique

Figure 5b demonstrates the influence of the different AR150 concentrations on the adsorption rate of various NCs at pH 3, 60 °C for one hour, and power level 400 Watt. With NC II and III, the increasing of AR150 concentration caused an augmentation in the adsorption quantity of acid dye on the NC. Contrary, using NC I and IV, the adsorption rate exhibited an increasing in the dye concentration and NC IV gave better result at 0.2 g/L. From Fig. 5a and 5b, it was observed that the dye adsorption rate of various NCs using ultrasonic technique was higher than that in case of using the conventional one, except NC III and NC IV at 0.5 g/L dye concentration. This indicates that both of the dye concentration and the composition of NC assumed a significant aspect in the adsorption quantity of acid dye onto various NCs. Moreover, the dye particles might transfer from outer surface to internal lamellar area and bringing about the disaggregation of the whole load. With high capacity rate of acid dye, agglomeration is predictable to be predominating, but monomers in addition to dimmers are basically missing within acid dye-NC complexes [30-37]. From Table 3, it can be concluded that in case of the nanocomposites I and II, the dye adsorption rate (%) was increased by increasing the dye concentration and the best dye adsorption rate can be obtained at 0.5 g/L dye conc. But for the nanocomposites III and IV, the dye adsorption rate (%) was decreased by increasing the dye concentration and the best dye adsorption rate can be obtained at 0.2 dye concentration.

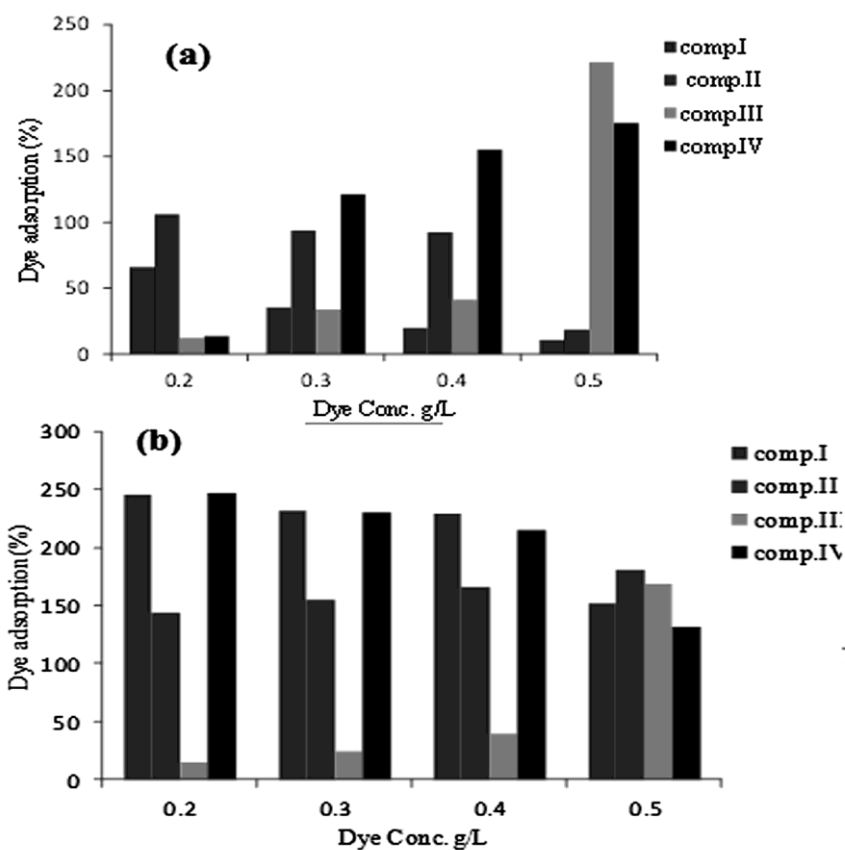


Fig. 5a. Effect of conventional dye concentration, b: Effect of ultrasonic dye concentration.

TABLE 3. Effect of the dye exhaustion rate (%) using ultrasonic method.

Dye conc. g/L	Exhaustion rate (%)			
	$E \% = (\text{Abs. before} - \text{Abs. after}) / \text{Abs. before} \times 100$			
	NC I	NC II	NC III	NC IV
0.2	61.7	76.1	95.5	95
0.3	73.8	91.2	93.4	69.9
0.4	81.7	96.1	89.7	69.7
0.5	96.7	98.1	60.1	68.4

Effect of ultrasonic power on the dye adsorption

Figure 6 demonstrated the influence of power of ultrasound on the adsorption of the acid dye on the NCs surface conducted at various power levels (100–500 watt). The acid dye adsorption on the various NCs was directly increased proportionally with an increasing of the power level supplied. Besides, the highest values of acid dye adsorption of the various NCs can be obtained at power level of 500 watt. This may be attributed to the effect of cavitation phenomena.

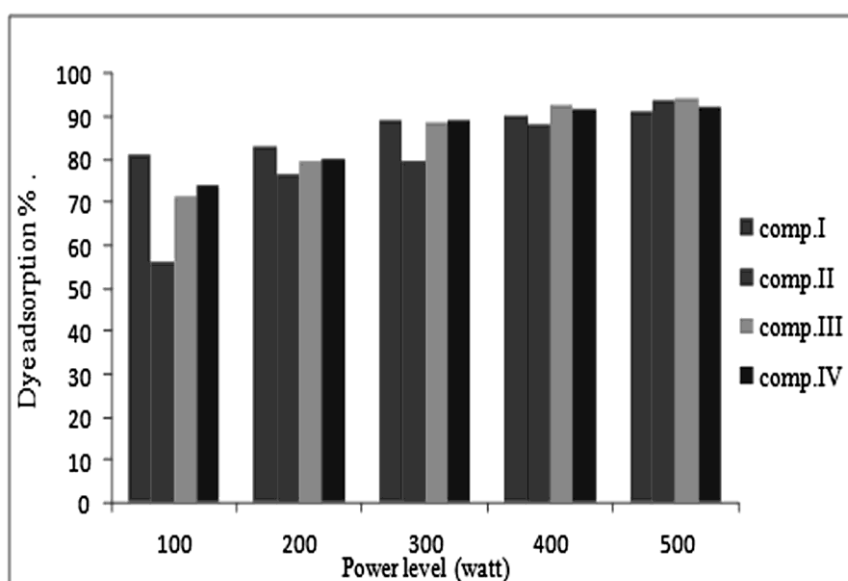


Fig. 6. Effect of power level of ultrasonic power.

Adsorption kinetics

It is realized that the rate of any procedure implies an alteration in any of the starting substrates which happens in the procedure or the item that gained for each time unit. Stratifying this clarification in the adsorption procedure could be seen as an adjustment in dye uptake for each duration unit [37, 38]. Duration of dye uptake isotherms for the composites adsorbed by acid red150 using conventional and ultrasonic heating methods are presented in Figs 7 a, b, c and d. These figures explains that dye uptake percentages for the adsorbed nanocomposites using ultrasound technique. It was observed that in the most part the adsorption rate was lower than those got from the conventional technique. Furthermore, the isotherms for the two techniques were separated against each another and showed lower dye uptake in case of the ultrasonic technique relative to the conventional one(39). Data in Fig. 7 a, b, c and d could be evaluated via available common type of the primary order rate for the adsorption of NCs I, II, III and IV to the acid dye red150.

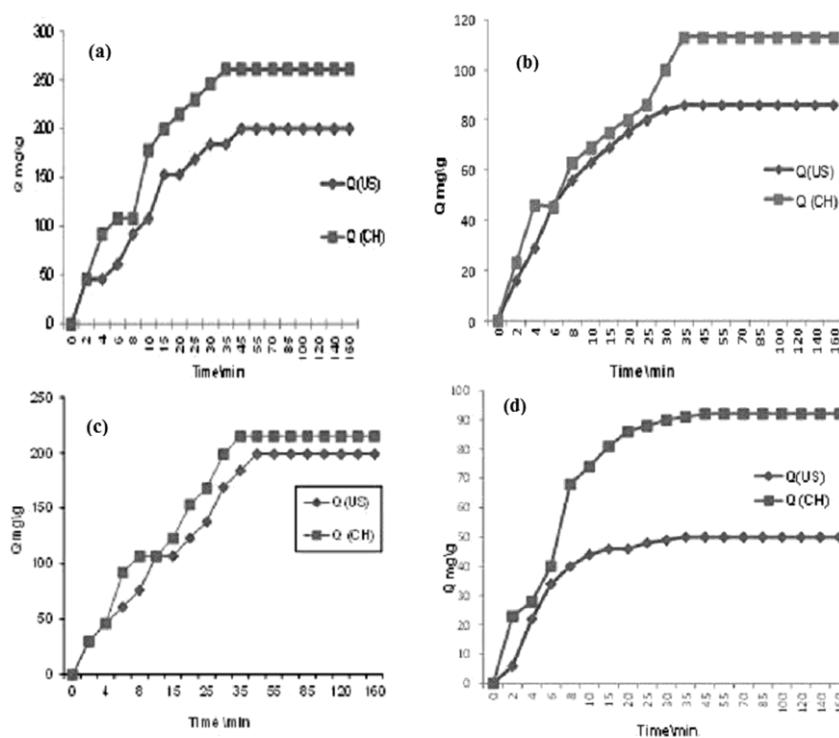


Fig. 7a. Ultrasonic and conv. adsorption rate of composite I, **b:** ultrasonic and conv. adsorption rate of composite II, **c:** ultrasonic and conv. adsorption rate of composite III, **d:** ultrasonic and conv. adsorption rate of composite IV

Standard affinity ($-\Delta\mu$)

The information of dyeing equilibrium are usually announced as the standard affinity of adsorption [40], therefore, the standard affinity can be mentioned in the experimental part. Fig. 8a demonstrates a plot for $\ln|Q_t - Q_f|$ as a parameter for duration of adsorption of NCI utilizing both conventional and ultrasonic techniques. Figure 8b demonstrates a plot for $\ln|Q_t - Q_f|$ as a parameter for duration of adsorption of NCII using both conventional and ultrasonic techniques. Figure 8c demonstrates a plot for $\ln|Q_t - Q_f|$ as a factor for duration of adsorption of NCIII using both conventional and ultrasonic techniques. Fig. 8d demonstrates a plot for $\ln|Q_t - Q_f|$ as a factor for duration of adsorption of NCIV using both conventional and ultrasonic techniques. As expected, the linear suitability holds without a doubt and the amounts of the adsorption constants can be calculated and recorded in Table 4.

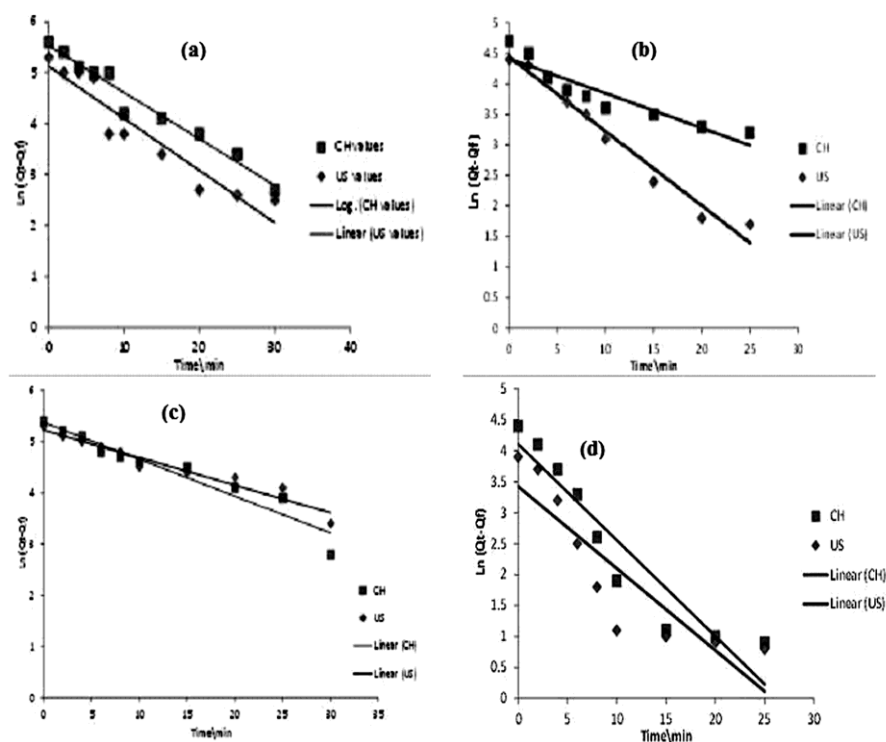


Fig. 8a. Relation between $\ln(Q_t - Q_f)$ and time using conventional and ultrasonic conditions for composite I, b: Relation between $\ln(Q_t - Q_f)$ and time using conventional and ultrasonic conditions for composite II, c: Relation between $\ln(Q_t - Q_f)$ and time using conventional and ultrasonic conditions for composite III, d: Relation between $\ln(Q_t - Q_f)$ and time using conventional and ultrasonic conditions for composite IV

TABLE 4. Adsorption percentage (%) of constant K, time for half reaction $t_{1/2}$, standard affinity $-\Delta\mu$ and quantity for final dyestuff uptake using NC I, II, III and IV.

NCs	$K \times 100$ (min^{-1})		$-\Delta\mu$ (kJ/mol)		$t_{1/2}$ (min)		Q_f (mg/g)	
	US	Conv.	US	Conv.	US	Conv.	US	Conv.
NC I	7	10	18	19.1	10.1	7.2	200	250
NC II	10	11.6	16.9	18.7	11.6	10	110	225
NC III	10	13.2	16.7	17.1	14	11.6	165	210
NC IV	8.6	9	16	17.7	10	8	60	136

Conclusion

The data confirmed that the adsorption rate of acid red 150 dye onto the prepared gelatin/chitosan/laponite based nanocomposites was improved using ultrasonic technique relative to the conventional one. On the other hand, in case of high content of laponite (2 wt%), the maximum adsorption rate could be obtained at low dose of the nanocomposite. In other words, both chitosan and laponite concentrations played an important role for enhancing the adsorption capacity.

References

1. **Kamel, M.M., El-Shishtawy, R.M., Hanna, H.L. and Ahmed, N.S.E.** Ultrasonic-assisted dyeing: I. Nylon dyeability with reactive dyes. *Polym Inter.* **52**, 373(2003).
2. **Kamel, M.M., M.M. El Zawahry, N.S.E. Ahmed and F. Abdelghaffar.** Ultrasonic dyeing of cationized cotton fabric with natural dye. Part 1: Cationization of cotton using Solfix E. *Ultrasonics Sonochemistry*, **16**, p.243(2009).
3. **Lee, K.W., Chung, Y.-S. and Kim, J.-P.** Effects of ultrasound treatment and dye crystalline properties on particle size distribution. *Textile Research, J.* **71**, 976 (2001).
4. **Kamel, M.M., El-Shishtawy, R.M., Yussef, B. and Mashaly, H.** Ultrasonic assisted dyeing: III. Dyeing of wool with lac as a natural dye. *Dyes and Pigments*, **65**, 103(2005).
5. **Ince, N.H., Tezcanli, G., Belen, R.K. and Apikyan, I.G.** Ultrasound as a catalyzer of aqueous reaction systems: the state of the art and environmental applications. *Applied Catalysis B: Environmental*, **29**, 167(2001).
6. **Sekar, N.,** ITMF: Sudhir Thackersey, new senior vice president. *Colourage*, **48**, 42(2001).
7. **Kamel, M.M., Helmy, H., Mashaly, H. and Kafafy, H.** Ultrasonic assisted dyeing: Dyeing of acrylic fabrics CI Astrazon Basic Red 5BL 200%. *Ultrasonics Sonochemistry*, **17**, 92 (2010).
8. **Kamel M.M., El-Hossamy M., Helmy H. and El-Hawary N.** Some studies on dyeing properties of cotton fabrics with Curcuma Longa (turmeric)(roots) using ultrasonic method. *Polish J. Applied Chem* 101(2008).
9. **Kamel M.M., Helmy H. and El Hawary N.** Some studies on dyeing properties of cotton fabrics with crocus sativus (Saffron flowers) using an ultrasonic method. *J. Natural Fibers* **6**, p.151(2009).
10. **Vankar, P.** Dyeing of cotton with Eucalyptus bark. *Asian Textile J.-Bombay* **9**, 30(2000).

11. **Didenko, Y.T. and Suslick K.S.**, The energy efficiency of formation of photons, radicals and ions during single-bubble cavitation. *Nature* , **418**, p.394(2002).
12. **Siddaramaiah, P.L., Kim NH., Heo, SB. and Lee J.H.** Novel PAAm/laponite clay nanocomposite hydrogels with improved cationic dye adsorption behaviour. *Composite B* **31**, 756 (2008).
13. **Haroun, A.A., Mashaly H. and El-Sayed N.** Novel nanocomposites based on gelatin/HPET/chitosan with high performance acid red 150 dye adsorption. *Clean Techn Environ Policy*, **15**, 367(2013).
14. **Ravi Kumar, M.N.V.** A review of chitin and chitosan applications. *Reactive and Funct. Polym.* **46**, 1(2000).
15. **Manoj Kumar, P.P.N.** Grafting Vinyl Monomers onto Chitosan:IV:Graft copolymerized of acrylic acid onto chitosan using ceric ammonium nitrate as the initiator characterization and antimicrobial activities. *Mater Sci and Applications* **2**, 1741(2011).
16. **Haroun, A.A and El-Halawany N.R.** Preparation and evaluation of novel interpenetrating polymer network-based on newspaper pulp for removal of copper ions. *Polymer-Plastics Technol. Eng.* **50**, 232 (2011).
17. **Haroun, A.A., Taleb E.M.A. and Abd El-Ghaffar M.A.** Synthesis and characterization of novel thermoplastic films for removal of heavy metal ions. *Polymer-Plastics Technol. Eng.* **49**, 454(2010).
18. **Elhalawany, N., Haroun, A., Abbas, H. and Hammad, F.** Conducting chelating polymer composites based on grafted waste polystyrene for removal of toxic copper ions. *J. Elastomers and Plastics*, **46**, 553(2014).
19. **Azlan, K., Wan Saime W.N. and Lai Ken L.** Chitosan and chemically modified chitosan beads for acid dyes sorption. *J. Environmental Sci.* **21**, 296(2009).
20. **Cestari, A.R., Vieira, E.F.S., Tavares A.M.G. and Bruns R.E.** The removal of the indigo carmine dye from aqueous solutions using cross-linked chitosan-Evaluation of adsorption thermodynamics using a full factorial design. *J. Hazardous Mater.* **153**, 566 (2008).
21. **Cheung, W.H., Szeto, Y.S. and McKay G.** Intraparticle diffusion processes during acid dye adsorption onto chitosan. *Bioresource Technol.* **98**, 2897(2007).
22. **Zubieta, C.E., Messina, P.V., Luengo, C., Dennehy, M., Pieroni, O. and Schulz P.C.** Reactive dyes remotion by porous TiO₂-chitosan materials. *J. Hazardous Mater.* **152**, 765(2008).
23. **Wang, L. and Wang, A.** Adsorption properties of congo red from aqueous solution onto N,O-carboxymethyl-chitosan. *Bioresource Technol.* **99**,1403(2008).
24. **Das, S.K., Bhowal, J. Das, A.R. and Guha, A.K.** *Langmuir* **22**, 7265(2006).

25. **Robinson, T., McMullan, G., Marchant, R. and Nigam P.**, Remediation of dyes in textile effluent: a critical review on current treatment technologies with a proposed alternative. *Bioresource Technol.* **77**, 247(2001).
26. **Rauf, M.A., Bukallah, S.B., Hamadi, A., Sulaiman, A. and Hammadi F.**, The effect of operational parameters on the photoinduced decoloration of dyes using a hybrid catalyst V2O5/TiO2. *Chemical Eng J.* **129**, 167(2007).
27. **Gürses, A. Ç. Doğar, M. Yalçın, M. Açıkyıldız, R. and Bayrak, S. Karaca**, The adsorption kinetics of the cationic dye, methylene blue, onto clay. *J. Hazardous Mater.* **131**, 217(2006).
28. **Behnajady, M. N., Modirshahla, N. Daneshvar, M. Rabbani**, Photocatalytic degradation of an azo dye in a tubular continuous-flow photoreactor with immobilized TiO₂ on glass plates. *Chemical Engineering J.* **127**, 167(2007).
29. **Demirbas A.**, Agricultural based activated carbons for the removal of dyes from aqueous solutions: A review. *J. Hazardous Mater.* **167**, 1(2009).
30. **Ekici, S., Işıkver, Y. and Saraydın D.** Poly(acrylamide-sepiolite) composite hydrogels: preparation, swelling and dye adsorption properties. *Polym. Bull.* **57**, 231(2006).
31. **Gupta, V.K.**, Suhas, Application of low-cost adsorbents for dye removal-A review. *J. Environmental Management* , **90**, 2313(2009).
32. **Li, P., Siddaramaiah, Kim N.H., Heo S.-B., Lee J.-H.**, Novel PAAm/laponite clay nanocomposite hydrogels with improved cationic dye adsorption behavior. *Composites Part B: Engineering*, **39**, 756(2008).
33. **Wong, Y.C., Szeto, Y.S., Cheung, W.H. and McKay, G.**, Adsorption of acid dyes on chitosan-equilibrium isotherm analyses. *Process Biochemistry*, **39**, 695(2004).
34. **Luo, X. and Zhang L.**, High effective adsorption of organic dyes on magnetic cellulose beads entrapping activated carbon. *J. Hazardous Mater* **171**, 340(2009).
35. **Suresh, S.** Adsorption of acid red 18 from aqueous solution onto activated carbon prepared from *Murraya koenigii* (curry tree) seeds. *Asian J Chem* **23**, 219(2011).
36. **de Carvalho, G.M., E.C. Muniz, and A.F. Rubira**, Hydrolysis of post-consume poly(ethylene terephthalate) with sulfuric acid and product characterization by WAXD, 13C NMR and DSC. *Polym Degrad Stab* **91**, 1326(2006).
37. **Fatyeyeva, K., J. Bigarré, B. Blondel, H. Galiano, D. Gaud, M. Lecardeur and F. Poncin-Epaillard**, Grafting of p-styrene sulfonate and 1,3-propane sultone onto Laponite for proton exchange membrane fuel cell application. *J. Membrane Sci.* **366**, 33(2011).
38. **Annadurai, G., L.Y. Ling and J.F. Lee**, Adsorption of reactive dye from an aqueous solution by chitosan: isotherm, kinetic and thermodynamic analysis. *J. Hazard Mater*, **152**, 337(2008).

39. **Zemaitaitiene, R.J., Zliobaite E., Klimaviciute, R. and Zemaitaitis A.**, The role of anionic substances in removal of textile dyes from solutions using cationic flocculant. *Colloids and Surfaces A: Physicochem Eng Aspects* **214**, 37(2003).
40. **Mittal, A., Mittal, J. and Kurup L.**, Adsorption isotherms, kinetics and column operations for the removal of hazardous dye, Tartrazine from aqueous solutions using waste materials-Bottom Ash and De-Oiled Soya, as adsorbents. *J. Hazardous Mater* **136**, 567(2006).

(Received 27/1/2017;
Accepted 5/3/2017)

دراسة حركية متراكبات متناهية الصغر مبنية علي الجيلاتين مع الكيتوزان لإمتزاز الصبغة الحمضية الحمراء 150 باستخدام طاقه الموجات فوق الصوتيه

أحمد عوض هارون¹، حماده مشالي²، هاني حلمي²، مني كامل²
¹شعبة بحوث الصناعات الكيمايئيه،²شعبة بحوث الصناعات النسيجه، المركز القومي للبحوث، شارع
البحوث، 12622 الدقي، جيزه، جمهورية مصر العربيه.

يهدف هذا البحث إلي دراسة تأثير مختلف عوامل إمتزاز الصبغة الحمضية الحمراء 150 علي متراكبات متناهية الصغر مبنية علي الجيلاتين مع الكيتوزان في وجود طفلة لابونيت باستخدام طاقه الموجات فوق الصوتيه مثل: تركيز الصبغه، زمن الإمتزاز ، درجة الاس الهيدروجيني، تركيز المتراكبات المستخدمه.
بالإضافه إلي المقارنه بين إستخدام طاقه الموجات فوق الصوتيه والطاقه الحراريه التقليديه في توضيح مدي التأثير في قابليه إمتزاز المتراكبات المحضره للصبغه الحمراء. علاوه علي دراسة ديناميكية المواد المحضره للإمتزاز.
أثبتت النتائج انه كلما زادت نسبة الصبغه وكذلك نسبة الكيتوزان زاد معدل الإمتزاز. كذلك كلما زادت قوة طاقه الموجات فوق الصوتيه زاد معدل الإمتزاز لكل المتراكبات المستخدمه. ومن ناحيه اخري كان افضل معدل إمتزاز عند إستخدام تركيز للصبغه 277 مللجرام/لتر وتركيزات للمتراكبات تتراوح من 0.3 إلي 0.8 جرام حسب التركيب الكيمايئي للمتراكب المحضر.
يمكننا القول بأن كل من طفلة لابونيت والكيتوزان يلعبان دورا مهما في زيادة معدل الإمتزاز عند إستخدام طاقه الموجات فوق الصوتيه.