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Synthesis of a biodegradable interpenetrating polymer network of Av-cl-poly(AA-ipn-AAm) for malachite green dye removal: kinetics and thermodynamic studies

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This paper deals with the synthesis of a biodegradable interpenetrating polymer network (IPN) from the natural polysaccharide aloe vera (AV), acrylamide (AAm) and acrylic acid (AA), and its evaluation as a dye removal device. In the synthesis of Av-cl-poly(AA-ipn-AAm), ammonium persulfate (APS) was used as an initiator, N,N'-methylene bisacrylamide (MBA) as a cross-linker, AA and AAm as primary and secondary monomers, respectively. Soil burial and composting methods were used to study the biodegradability of the synthesized IPN and the results showed 94% degradation within 70 days using the composting method and 86% degradation within 77 days using the soil burial method. Biodegradation was confirmed by Fourier transform infrared spectroscopy (FTIR) and scanning electron microscopy (SEM) techniques. The synthesized IPN was used as a device for the removal of malachite green (MG) dve from agueous solution. The maximum MG removal capacity of the synthesized IPN was found to be 97.3% under the optimal conditions (i.e. time = 180 min., pH = 4.5, adsorbent dose = 5 g L^{-1}). The adsorption kinetics of malachite green molecules onto synthesized IPN was studied and compared using pseudo-first-order and pseudo-second-order models and we found that the adsorption process is better represented by the pseudo-second-order model. The different adsorption isotherm models like Langmuir, Freundlich, Dubinin-Radushkevich, Temkin, Redlich-Peterson and Sips isotherms were studied. The best-fitting isotherm model for the present experiment is the Langmuir model.

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Introduction

Different pollutants like noxious metal ions, dyes, phenols, insect repellents, which are used in agricultural practices and aromatic compounds present in wastewater affect our environment and have a detrimental effect on aquatic animals as well as human beings. The use of synthetic dyes is increasing with the advancement in industrialization and urbanization. Most of the dyes are harmful and oncogenic, causing a harsh ecological impact. Industrial wastewater treatment is therefore of utmost importance in saving our environment^{1,2} since these dyes not only have a negative effect on human beings and animals but also affect symbiotic processes by disturbing photosynthetic activity.³

Researchers all over the world are interested in the use of economical and eco-friendly sorbents for the removal of harmful dyes from aqueous solution. There are different methods like adsorption, coalescence, membrane separation, oxidation. photocatalytic dilapidation, electrochemical processes and deterioration by microorganisms in the presence or absence of oxygen, which have been employed for the removal of dyes from water. However, adsorption is the most operative and viable process for wastewater treatment because of its great efficacy, easy procedure and non-susceptibility to harmful pollutants.4 There are several adsorbents that are used for the removal of harmful dyes from the aqueous effluents, for example, silica, fly ash, waste banana pith, orange peel, palm fruit bunch, etc. However, no kinetic model has been used to study sorption processes in all these adsorbents, since the sorption process is complicated and is categorized into two main types: physisorption and chemisorption. Physisorption involves weak intermolecular forces like van der Waals and hydrogen bonding interactions between adsorbate and adsorbent. In the chemisorption process, the electrons are covalently bonded or exchanged between adsorbate and adsorbent.5

Hydrogels based on acrylic acid and sodium alginate in the presence of a crosslinker and initiator have been synthesized

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and used for removing methylene blue dye from aqueous solution.6 A guar gum, acrylic acid and aniline-based IPN was prepared and assessed for its conductivity, antibacterial and dye adsorption properties and also for the removal of methylene blue dye from wastewater. Hydrogels based on poly-(methacrylic acid), N,N-methylenebisacrylamide and 2,2-azobis-[2-(2-imidazolin-2-yl)propane]dihydrochloride were synthesized were used for the removal of basic yellow 28.8

Poly(acrylic acid-acrylamide) hydrogel was synthesized by the frontal polymerization technique and was used for the removal of crystal violet and basic magenta dyes.9 The removal of crystal violet dye from its aqueous solution with the help of sodium dodecyl sulphate-treated NiFe₂O₄ magnetic nanoparticles and its adsorption kinetics has been studied in accordance with the pseudo-second-order model.10

Cempedak durian peel was considered to be a worthy adsorbent for acid blue 25 and the effects of various parameters such as contact time, pH, ionic strength, initial concentration and temperature were investigated; the different functional groups present in the cempedak durian peel were characterized using FTIR.11

Artocarpus odoratissimus leaves were investigated and considered to be an active sorbent for the exclusion of noxious crystal violet, malachite green and methylene blue dyes, and analysis was carried out using the Langmuir, Freundlich, Temkin, Dubinin-Radushkevich, Redlich-Peterson and Sips isotherm models.12-15 It was reported that the peels of breadnut, (Artocarpus camansi) and bitter gourd (Momordica charantia) were used as potent adsorbents in the sorption of methylene blue and crystal violet dyes. 16,17 The biosorption of methylene blue and methyl violet 2B on breadfruit (Artocarpus altilis) peel and its core was studied and it exhibited 80% dye removal; the Langmuir adsorption isotherm model was best fitted for this. 18,19 Adsorption kinetics is significant in the treatment of wastewater as it provides information about the reaction pathways and adsorption mechanisms. Kinetics also describes the solution uptake rate, which indicates the residence time of adsorbate uptake at the interface of the solid and solution.20-22

IPNs are cross-linked, three-dimensional networks of polymeric materials with different functional groups and have high water-swelling characteristics.23 IPNs are superior to hydrogels as they are formed from two different polymeric chains having different characteristics, which enhance the swelling capacity, have better mechanical properties and are stable at high temperature and in a wide pH range. The functional groups present in IPN interact with metal ions/dye molecules, which proves them to be efficient adsorbents; the structure of IPN can be flexibly designed and the properties can be adjusted by altering the type and number of functional groups according to the usage and requirement.24 In the last few years, the development of novel sorbents capable of removing dyes, organic or heavy metal pollutants from wastewaters has been an intensive and popular research field. Various studies have been devoted to the adsorption of cationic dyes by anionic IPN containing weakly acidic or strongly charged groups. The combination of electrostatic interactions with hydrophobic interactions is usually the driving force in the case of cationic organic species.21

Aloe vera (Av) is a wonder drug used by tribal communities but is also used for food, cosmetics, Ayurveda, homeopathic and allopathic medicines.22 Av-based hydrogels have been prepared and characterized for application in the treatment of several types of wounds.23 It has also been evaluated as a natural flocculant24 as well as a drug carrier.25

The present study deals with the synthesis of an interpenetrating polymer network of Av, AA and AAm using APS as an initiator and MBA as a cross-linker. Aloe vera is used as the backbone instead of other polysaccharides because it has high water swelling efficiency, it can be easily grafted with different vinyl monomers and does not have any harsh impact on human health. The vinyl monomers, AA and AAm, enhance the adsorption efficiency of the synthesized IPN. The biodegradability of the synthesized IPN has been studied through the soil burial and composting method. This candidate polymer has been evaluated as a device for dye sorption and is one of the most eco-friendly methods for dye adsorption.

2. **Experimental**

Materials and methods

Av was purchased from Kshipra Biotech Pvt. Ltd. APS was received from Avantor Performance Material, India Ltd. MBA, MG dye, AA and AAm were purchased from (Loba Chemie laboratory reagents and fine chemicals).

2.2. Instrumental analysis

IPN synthesis and its biodegradability were confirmed by different techniques including SEM and FTIR. Thermal stability and crystallinity were determined by TGA and XRD, respectively. FTIR spectra of the backbone and synthesized IPN were recorded on a Perkin Elmer spectrophotometer using KBr pellets. Scanning electron micrographs (SEMs) of the backbone and IPN were obtained on JSM 6100. Weighing was done on an electronic balance (LIBROR AEG-220, Shimadzu). The pH of solutions was adjusted using HCl and NaOH.

2.3. Synthesis of Av-AA-co-AAm based IPN [Av-cl-poly(AA-ipn-AAm)]

AA was grafted onto Av in an aqueous medium using APS as an initiator and MBA as a cross-linker. To get the graft copolymer Av (1 g) was taken in a reaction flask containing 4 mL of distilled water and was stirred thoroughly. APS $(0.438 \times 10^{-2} \text{ mol L}^{-1})$ was added followed by the drop by drop addition of AA (1.45 imes 10^{-2} mol L^{-1}) and MBA (1.1 \times 10^{-2} mol L^{-1}) to the reaction mixture. The reaction was carried out at 50 °C and pH 7. Stirring was continued until a homogeneous mixture was obtained. The graft copolymer was extracted using slightly hot distilled water and was dried in a hot air oven at 60 °C. The synthesized product was then taken in a flask having a known amount of distilled water, then AAm and AP were added and the mixture kept for 24 h. After 24 h, MBA was added with constant stirring. The product was washed with distilled water and dried in a hot air oven. The sample of known weight was immersed in distilled water and after every 2 h interval, the weight gain of the swollen

sample was noted. The percentage swelling (P_s) of the IPN was calculated from the following equation:26,27

> $P_{\rm s} = \frac{W_{\rm s} - W_{\rm d}}{W_{\rm s}} \times 100$ (1)

$$P_{\rm s} = \frac{W_{\rm s} - W_{\rm d}}{W_{\rm d}} \times 100 \tag{1}$$

where W_s and W_d are the weights of the swollen and dry polymer, respectively.

2.4. Batch adsorption study

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Batch experiments were carried out to study the dye adsorption by adding 0.05 g of synthesized IPN and 20.0 mL of (100 ppm) MG dye solution to a 100 mL flask. This flask was agitated at 250 rpm in an orbital shaker. The effect of different parameters like adsorbent dosage, initial dye concentration, contact time, pH and temperature were studied. The dye removal was studied using a UV-Vis spectrophotometer at wavelength λ_{max} of 615 nm. All the dye removal experiments were carried out in triplicate with a blank sample to avoid handling errors. The amount of MG dye adsorbed by the synthesized IPN at equilibrium, q_e (mg g⁻¹) was calculated using the equation given

$$q_{\rm e} = \frac{(C_{\rm i} - C_{\rm e})V}{m} \tag{2}$$

where C_i is the initial dye concentration (mg L⁻¹) and C_e is the dye concentration (mg L^{-1}) at equilibrium, respectively; m is the mass of the adsorbent (g) taken and V is the volume of the dye solution (L).

The percentage removal (% R) of the dye was calculated using the equation as follows:

$$\% R = \frac{(C_{\rm i} - C_{\rm e}) \times 100}{C_{\rm i}}$$
 (3)

2.5. Error analysis

The values of the coefficient of determination (R^2) , the kinetics and adsorption isotherm models were obtained from the experimental data. The value of qe $(q_{e,cal})$ was predicted from the formulae of different kinetics or isotherm models. The experimental and predicted values of q_e were fitted into the equations of various error analysis functions and the smallest value showed the least error. Four types of error analysis (sum of absolute error, chi-square test, average relative error and Marquardt's percent deviation error) were calculated using the equation given below:28,29

Sum of absolute error (EABS) =
$$\sum_{i=1}^{N} [q_{e,exp} - q_{e,cal}]$$
 (4)

Chi-squared test
$$(\chi^2) = \sum_{i=1}^{N} \frac{(q_{e,exp} - q_{e,cal})^2}{q_{e,exp}}$$
 (5)

Average relative error (ARE) =
$$\frac{100}{n} \sum_{i=1}^{N} \left[\frac{q_{e,exp} - q_{e,cal}}{q_{e,exp}} \right]_{i}$$
 (6)

Marquardt's percent standard deviation (MPSD)

$$=100\sqrt{\frac{1}{N-p}\sum_{i=1}^{N}\left(\frac{q_{e,exp}-q_{e,cal}}{q_{e,exp}}\right)^{2}}$$
 (7)

N and p are the number of data points in the experiment and the number of parameters of the kinetics, respectively.

2.6. Biodegradation study of the synthesized interpenetrating polymer network

Biodegradation is governed by various factors including the availability of functional groups, which increases hydrophilicity, size, density and molecular weight of the polymer, the number of crystalline and amorphous regions, structural complexity, types of bonds, molecular composition and the form of polymer film, powder and pellet.30

The biodegradability of synthesized IPN was studied through the composting and soil burial method. Compost was collected from the industrial sewerage effluent discharge of NIT Jalandhar, and soil was collected from the garden of the NIT Jalandhar campus. The compost and soil were contained in separate pots. The synthesized IPN samples (1 g) were buried in 3 cm apart in the pot containing compost and soil. All the samples were properly labelled on the outside of the pots. Compost contains large amounts of microbial flora, so the microbial species in the compost were regularly fed with discharge wastewater effluent for the proper growth of the microorganisms and fresh water was used for the soil burial method. The aqueous levels of the pots were maintained such that water just reached the base of the pot and care was taken that effluent was not leached out from the pot. Wastewater effluent was supplemented daily to the dried compost due to evaporation. The pots were covered with aluminum foil and maintained at ambient temperature. The weight of each sample was taken by removing samples at an interval of 7 days, followed by rinsing with water and drying at 50 °C. Percentage degradation (% D) in the form of % wt. loss as a function of the number of days was determined as follows:

$$\% D = \frac{W_{\rm i} - W_{\rm f}}{W_{\rm f}} \times 100 \tag{8}$$

where W_i = initial weight of the sample; W_f = final weight of the sample.31

3. Results and discussion

Synthesis of Av-cl-poly(AA-ipn-AAm)

Av underwent graft copolymerization with AA in the presence of APS as an initiator. MBA was used as the cross-linker, leading to the formation of Av-cl-poly(AA). The different reaction parameters were pre-optimized with respect to percentage swelling and are given in Table 1. IPN Av-cl-poly(AA-ipn-AAm) was synthesized, and the solvent and secondary monomer (AAm) were optimized with respect to percentage swelling. The AAm concentration decides the rate of reaction as well as swelling efficiency. As the weight ratio of the AAm to Av-cl-poly(AA) increased, the interpenetration of the AAm to Av-cl-poly(AA) also increased, leading to enhanced swelling; when it attained

equilibrium maximum swelling was obtained. As shown in Fig. 1a, the maximum percentage swelling ($P_{\rm s}$ 427%) was obtained at a concentration of 2.81×10^{-2} mol L⁻¹ of AAm. On further increasing the AAm concentration, no penetration of the AAm occurred and the AAm molecules were homopolymerized and removed from the solution by a simple extraction method; thus, a further increase in the AAm concentration did not increase the percentage swelling and in the case of solvent, maximum $P_{\rm s}$ (578%) was observed with 25 mL. This can be explained on the basis that the OH $^-$ ion concentration was initially sufficient to propagate the polymerization reaction until the optimum concentration was reached, but with the further increase in the amount of solvent (Fig. 1b), the OH $^-$ concentration increased causing the termination of the polymerization reaction.

3.1.1. Mechanism of IPN synthesis. The mechanism of IPN synthesis is given in Scheme 1. Persulphate ions from

ammonium persulfate generated SO₄^{-*} anionic radicals, which assisted in the abstraction of hydrogen atoms from OH groups present on the backbone. These became active sites capable of radically initiating the polymerization of AA, resulting in graft copolymerization with AAm and thus leading to the formation of Av-cl-pol(AA-ipn-AAm).³¹

3.2. Characterization

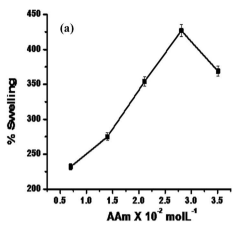
The synthesized IPN was characterized through different techniques like FTIR and SEM to study the morphological effects on Av after interpenetration and crosslinking. Thermal stability was studied using TGA.

3.2.1. SEM. The surface morphology of the AA and IPN were studied by the scanning electron microscopy (SEM) technique. SEM images of both samples showed morphological variations. Highly porous structures are seen in IPN, which represent the

Table 1 Optimization of different reaction parameters for the synthesis of Av-cl-poly(AA)

S. no	Av	Solvent (mL)	Acrylic acid (mol L ⁻¹)	Cross-linker (mol L^{-1})	Initiator (mol L^{-1})	Time (min)	рН	Temperature (°C)	Weight obtained (g)	Percentage swelling (P_s)
1	1 g	4 mL	10.9284	0.1622	0.0986	30	7	40	2.8537	717
2	1 g	4 mL	10.9284	0.1622	0.0986	60	7	40	3.1067	874
3	1 g	4 mL	10.9284	0.1622	0.0986	90	7	40	2.7789	812
4	1 g	4 mL	10.9284	0.1622	0.0986	120	7	40	2.7693	512
5	1 g	4 mL	10.9284	0.1622	0.0986	150	7	40	3.127	450
6	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	2.8261	918
7	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	3.1334	730
8	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	2.904	678
9	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	2.7801	615
10	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	2.8764	518
11	1 g	4 mL	10.9284	0.1622	0.0986	20	3	40	2.8632	498
12	1 g	4 mL	10.9284	0.1622	0.0986	20	5	40	3.078	570
13	1 g	4 mL	10.9284	0.1622	0.0986	20	7	40	3.0473	734
14	1 g	4 mL	10.9284	0.1622	0.0986	20	9	40	2.9605	529
15	1 g	4 mL	10.9284	0.1622	0.0986	20	11	40	2.9422	483
16	1 g	4 mL	10.9284	0.1622	0.0219	20	7	40	2.1776	559
17	1 g	4 mL	10.9284	0.1622	0.0438	20	7	40	2.1219	937
18	1 g	4 mL	10.9284	0.1622	0.0657	20	7	40	139.23	886
19	1 g	4 mL	10.9284	0.1622	0.0877	20	7	40	192.72	846
20	1 g	4 mL	10.9284	0.1622	0.1095	20	7	40	196.26	611
21	1 g	4 mL	10.9284	0.1622	0.0438	20	7	40	2.4177	1176
22	1 g	6 mL	10.9284	0.1622	0.0438	20	7	40	3.0445	1070
23	1 g	8 mL	10.9284	0.1622	0.0438	20	7	40	2.5787	918
24	1 g	10 mL	10.9284	0.1622	0.0438	20	7	40	3.0256	875
25	1 g	12 mL	10.9284	0.1622	0.0438	20	7	40	3.0435	573
26	1 g	4 mL	10.9284	0.0811	0.0438	20	7	40	2.438	388
27	1 g	4 mL	10.9284	0.1135	0.0438	20	7	40	2.7498	811
28	1 g	4 mL	10.9284	0.1459	0.0438	20	7	40	2.832	673
29	1 g	4 mL	10.9284	0.1784	0.0438	20	7	40	2.7463	530
30	1 g	4 mL	10.9284	0.2108	0.0438	20	7	40	2.8264	485
31	1 g	4 mL	6.9734	0.1135	0.0438	20	7	40	1.9735	676
32	1 g	4 mL	10.9284	0.1135	0.0438	20	7	40	2.6737	798
33	1 g	4 mL	14.5712	0.1135	0.0438	20	7	40	3.4709	852
34	1 g	4 mL	18.2139	0.1135	0.0438	20	7	40	3.6126	759
35	1 g	4 mL	21.8567	0.1135	0.0438	20	7	40	4.3639	633
36	1 g	4 mL	21.8567	0.1135	0.0438	20	7	30	2.3258	769
37	1 g	4 mL	21.8567	0.1135	0.0438	20	7	40	2.6734	839
38	1 g	4 mL	21.8567	0.1135	0.0438	20	7	50	2.9239	879
39	1 g	4 mL	21.8567	0.1135	0.0438	20	7	60	2.8632	841
40	1 g	4 mL	21.8567	0.1135	0.0438	20	7	70	2.6453	812

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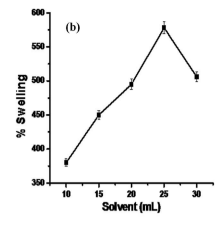


Fig. 1 (a) The effects of AAm concentration and (b) solvent on the percentage swelling (P_s) of the synthesized IPN.

regions of water permeation and interaction sites of external stimuli with the hydrophilic groups of the grafted polymer.

Additional irregular and coarse surfaces were observed in Avcl-poly(AA-ipn-AAm) in SEM results. These certainly reveal the morphological dissimilarities brought about by the chemical alteration of the backbone by grafting, crosslinking and interpenetration of the polymeric chain in Av as shown in Fig. 2a and 2b, due to the incorporation of covalent bonds between different polymeric chains on crosslinking with N,N-methylene bis acrylamide. The synthesized IPN had a porous structure. These pores might be the regions of water permeation and interaction sites of external stimuli.

3.2.2. FTIR (Fourier transform infrared spectroscopy). The broad peak observed in the IR spectrum of Av at about 3325 cm⁻¹ is due to the O-H stretching of the intermolecular hydrogen-bonded OH group. The peak at 2915 cm⁻¹ is due to the C-H stretching of the methyl group, that at 2101 cm⁻¹ is due to the C=C stretching of the monosubstituted group, and that at 1237 cm⁻¹ is due to the C-C deformation in the methyl group (Fig. 3a). On the other hand, the IR spectrum of Av-cl-poly(AAipn-AAm) showed a peak at 3327 cm⁻¹ due to the intermolecular hydrogen-bonded OH group. The intense absorption band at 2942 cm⁻¹ is due to the C-H stretching of the CH₂ group, the peak at 2323 cm⁻¹ is due to the N-H stretching of the secondary amide group, 1651 cm⁻¹ is due to the C=O stretching of tertiary amides, 1418 cm⁻¹ is due to the C-H deformation of the methyl group and 1110 cm⁻¹ is due to C-C stretching (Fig. 3b).

3.2.3. TGA of Av and Av-cl-poly(AA-ipn-AAm). Thermogravimetric analysis was used to measure the effect of temperature on synthesized IPN, it represents the stoichiometric decomposition reaction of particular substances exhibiting the mass of sample lost at different stages. It involves the burning of the sample, thereby forming combustion products. TGA of Av was conducted by taking 5.2 mg of sample. Av melts up to 310 °C and the decomposition takes place in several steps. In the first step, the volatile components like moisture, solvent or monomer are lost at 110 °C. This stage is followed by the pyrolysis of the polymer at 205 $^{\circ}$ C. At 280 $^{\circ}$ C, the atmosphere was switched from N2 to O2, which resulted in the combustion of carbon black additives. Some of the inorganic components remained

behind as residue at 310 °C. The evaluation of the pyrolysis weight loss of polymer content in various steps is as follows: first step, 24.4%; second step, 46.2%; third step, 28.9%.

TGA of Av-cl-poly(AA-ipn-AAm) was done by taking 5.0 mg of the synthesized IPN. The decomposition of IPN occurred up to 390 °C. In the first stage of the TGA of IPN, the volatile components like moisture, solvent or monomer were lost up to 110 °C, followed by the second step involving the pyrolysis of polymer up to 240 °C, at which decomposition takes place to release carbon monoxide. In the third step, further combustion of carbon takes place up to 370 °C. At 370 °C, the atmosphere was switched from N₂ to O₂, which resulted in the combustion of carbon black additives. Some of the inorganic components remained behind as residue up to 390 °C. The evaluation of the pyrolysis weight loss of the polymer content in various steps is as follows: first step, 25.4%; second step, 22.4%; third step, 48.6% (Table 2, Fig. 4a and b). This shows that grafting and crosslinking of Av with (AAm)-MBA increases the thermal stability of the synthesized IPN.32,33

3.2.4. XRD of Av and Av-cl-poly(AA-ipn-AAm). It is evident from Fig. 5a and b that compared to Av, the Av-cl-poly(AA-ipn-AAm) showed higher intensity peaks and the coherence length was found to be 0.986 Å, whereas it was 0.542 Å for Av. Higher coherence length indicates an increase in anisotropy with the alignment of different polysaccharide chains, poly(AA) chains and poly(AAm) chains along with cross-linking and interactions in comparison to the backbone.33

The coherence length of the samples was calculated by using the Scherrer equation:

$$L = 0.9 \lambda / \beta^{1/2} X \cos \theta$$

where λ = wavelength, θ = diffraction angle, L = coherence length and $\beta^{1/2}$ = full width at half maximum.

3.3. Biodegradation study using the composting and soil burial method

The biodegradability of the synthesized IPN Av-cl-poly(AA-ipn-AAm) and Av was investigated through the composting and soil burial method. The evaluation of the biodegradability of the

A Chain Initiation
$$S_2O_3^{2^+}$$
 $S_2O_4^{2^+}$ $S_2O_4^{2^+}$

Scheme 1 Representation of the mechanism of the synthesis of Av-cl-poly(AA-ipn-AAm).

Av-cl-poly(AA-ipn-AAm) and Av was conducted by considering three different stages of the biodegradation. The end of the first week was considered the first stage of biodegradation. The end of the sixth week was considered the second stage of biodegradation and the third stage was after the completion of the 11

weeks. It was found that Av was completely degraded in the 2nd and 3rd weeks in the composting and soil burial methods, respectively. Initially, in both cases, biodegradation occurred slowly but was later increased in composting as compared to soil because in the initial stage, the microflora acted on the

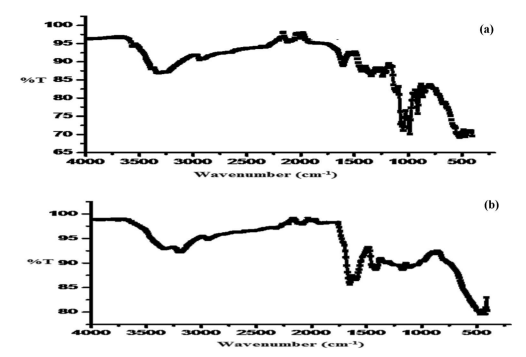


Fig. 2 (a) FTIR spectra of Av and (b) Av-cl-poly(AA-ipn-AAm).

surface of the synthesized IPN; when the surface was eroded then degradation was increased to a greater pace. The results showed 94% degradation of the IPN within 60–80 days in the composting method and 86% degradation was seen in the case of the soil burial method after 11 weeks (Table 3). The biodegradation may occur due to the presence of different strains of the microorganism in the compost and soil. This enzymatic, chemical and physical degradation may lead to the alteration of the physicochemical properties of the IPN. The biodegradation was slower in the case of the soil burial method as compared to the composting method. This could be due to the fact that soil has less microbial colonies as compared to compost.¹⁵

Morphological and chemical modification of the synthesized IPN due to biodegradation was studied using SEM and FTIR.³¹⁻³⁴

3.3.1. SEM of different biodegradation stages under the composting and soil burial methods. The changes at the morphological level in different stages of the biodegradation of the synthesized IPN were revealed through the SEM studies. SEM images of different biodegradation stages of Av-cl-poly(AA-ipn-AAm) through the soil burial and composting methods are shown in Fig. 6a–g. Different stages of the biodegradation were reasonably manifested from the SEM results. Fig. 6a shows the SEM image of the synthesized IPN.

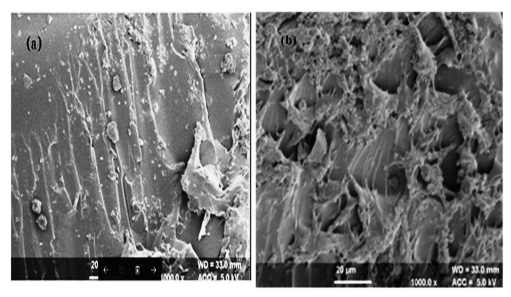


Fig. 3 (a) SEM of Av and (b) Av-cl-poly(AA-ipn-AAm).

Table 2 TGA of Av and Av-cl-poly(AA-ipn-AAm)

	Thermogravimetric analysis						
Sample Code	First stage decomposition (°C), pyrolysis weight loss of polymer content (%)	Second stage decomposition (°C), pyrolysis weight loss of polymer content (%)	Third stage decomposition (°C), pyrolysis weight loss of polymer content (%)				
Av	110 °C, 24.4%	110-205 °C, 46.2%	205–280 °C, 28.9%				
IPN	110 °C, 23.4%	110–205 °C, 22.4%	205–280 °C, 28.9%				

In the first stage of the biodegradation, there were external cracks in both the composting and soil burial samples (Fig. 6b and e). The size of these cracks increased as the time of biodegradation increased. Large cracks and deep pits are clearly seen in the second stage of biodegradation (Fig. 6c and f). The results showed the complete disintegration of the samples in the third stage of biodegradation (Fig. 6d and g) in both composting and soil burial methods. Thus, SEM results clearly revealed that the synthesized IPN is biodegradable. 32,33

3.3.2. FTIR analysis of different biodegradation stages under the composting and soil burial method. FT-IR spectra of different biodegradation stages of Av-cl-poly (AA-ipn-AAm) were recorded (Fig. 5a-f). Av-cl-poly(AA-ipn-AAm) (Fig. 7a) had peaks at 3327 cm⁻¹ (OH stretching of the intermolecular H bonded OH group), 3187 cm⁻¹ (due to the broad O-H stretching of the COOH group), 2942 cm⁻¹ (C-H stretching of the methyl group), 1651 cm⁻¹ (C=O stretching of the tertiary amide group), 1418 cm⁻¹ (due to C-H deformation of the methyl group) and

463 cm⁻¹ due to the presence of the pyranose ring. There was a decline in the intensity of the peaks of the degraded samples as compared to the synthesized IPN and there was also a loss of some peaks in the degraded IPN. The intensities of the peaks initially observed at 3327 cm⁻¹, 2942 cm⁻¹ and 1651 cm⁻¹ were found to be decreased at biodegradation stage-I (Fig. 7b and e), due to the beginning of the breaking down of the cross-linked network. However, the degradation of individual compounds like Av, poly(AA-ipn-AAm) and MBA started in biodegradation stage-II (Fig. 7c and f). In the third stage of biodegradation, most of the FTIR peaks initially observed in the FT-IR spectrum of Av-cl-poly (AA-ipn-AAm) were shifted (Fig. 7d and g). Thus, FT-IR confirmed that the grafted chains of poly (AA-ipn-AAm) and Av were completely biodegraded within three weeks.33,35

3.4. The effects of different parameters on dye removal

3.4.1. The effect of synthesized IPN dosage on dye removal. The effect of synthesized IPN dosage on MG dye (10 ppm)

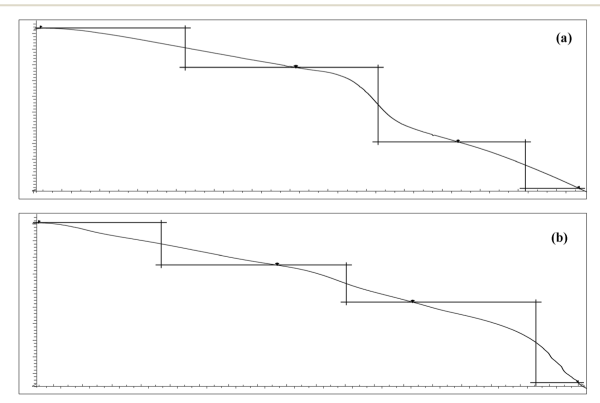


Fig. 4 TGA of (a) Av and (b) Av-cl-poly(AA-ipn-AAm).



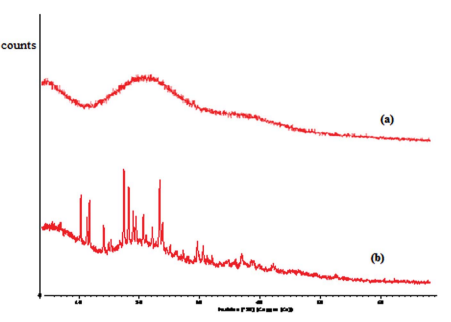


Fig. 5 XRD of (a) Av and (b) Av-cl-poly(AA-ipn-AAm).

removal was studied with different doses (1–7 g $\rm L^{-1}$) of IPN, as presented in Fig. 8. The results showed that dye removal increased with the increase in the dosage level. The maximum dye removal was found to be 97.1% with 5 g $\rm L^{-1}$ and remained constant with further increase in the adsorbent dose (Fig. 8). This is because as the adsorbent dose increased, there was an increase in the surface area for the adsorption of the MG dye from the solution, thus an increase in the dye removal.³⁶

3.4.2. The effects of pH and point of zero charge. It is important to know the range of pH that did not degrade or lead to the loss of the colour intensity of the MG dye. The solution of MG dye (10 ppm) was prepared and adjusted from pH 2.0–10.^{37,38}

pH plays an important role in adsorption studies because the pH of the aqueous solution can lead to the protonation or deprotonation of the functional groups on the adsorption surface. pH also plays an important role in the electrostatic interaction between the dye molecules and adsorbent. The maximum dye removal was observed at pH 4.5 (Fig. 9), which may be due to the fact that MG dye molecules exist in monomeric form at pH 4.5, leading to the easy penetration of the adsorbent by the dye molecules. The dye removal was less at higher pH, which may be due to the fact that MG molecules exist in zwitterionic form and this leads to the aggregation and formation of large molecules. The entry of large molecules is

restricted in the small pore size of the adsorbent and thus there is a decrease in dye adsorption. The point of zero charge for synthesized IPN is pH 6.2, while it is (predominantly) positively charged if the solution pH is less than 6.2 and negatively charged if the pH is higher than 6.2. At pH greater than 5, the MG molecules exist in cationic form, which leads to the lower adsorption due to electrostatic repulsion forces.³³⁻⁴¹ The pH of dye solution after adsorption was found to be 6.4.

3.4.3. The effect of contact time. It is important to ensure complete equilibrium between the dye molecules and the adsorbate surface, so the contact time for maximum dye adsorption (97.84%) was optimized. It was found from the results that the dye removal was initially slow and later it rapidly increased; equilibrium was finally attained at 180 min (97.24%) (Fig. 10).

The slow adsorption of the dye in the initial stage (within 2 h) was due to the slow swelling of the adsorbent at this stage, which further led to slower adsorption. The high availability of the adsorption sites increased the adsorption (after 2 h) and subsequently, the adsorption slowed down due to the attainment of equilibrium and the depletion of the adsorption sites.⁴¹ For complete equilibrium between the dye molecules and adsorbent surface, the contact time was optimized to get the maximum dye adsorption. It was found that dye removal was initially slow, then the adsorption rapidly increased and equilibrium was finally attained at 180 min (Fig. 10).

 Table 3
 Biodegradation studies of IPN using the composting and soil burial methods

Biodegradation studies of IPN using compositing and soil burial methods										
Sample code Percentage lost at different time interval (days)										
Av-cl-poly(AA-IPN-AM) compositing	7	14	21	28	35	42	49	56	63	70
method	7	15	23	30	41	48	59	66	81	94
Av-cl-poly(AA-IPN-AM) soil burial method	5	11	20	29	36	45	55	64	75	86

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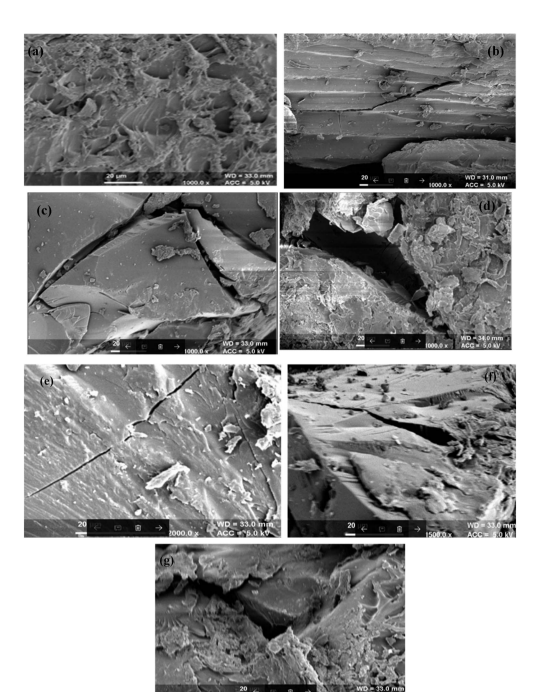


Fig. 6 SEM of (a) Av-cl-poly(AA-ipn-AAm), (b) Av-cl-poly(AA-ipn-AAm) biodegradation-I, (c) Av-cl-poly(AA-ipn-AAm) biodegradation-II, (d) Av-cl-poly(AA-ipn-AAm) biodegradation-III using the composting method, (e) Av-cl-poly(AA-ipn-AAm) biodegradation-I, (f) Av-cl-poly(AA-ipn-AAm) biodegradation-II and (g) Av-cl-poly(AA-ipn-AAm) biodegradation-III using the soil burial method.

3.4.4. The effect of dye concentration. The initial concentration of the dye also impacted the removal of the dye; different concentrations of dye were taken (10–80 ppm) to study the effect on percentage dye removal. It is clear from the results that maximum dye removal was found (97.3%) using 10 ppm (Fig. 11).

3.5. Adsorption kinetics

The kinetics of the adsorbent for the treatment of dye was studied. There are different types of kinetic models explaining the mechanism by which dyes are adsorbed. The kinetics of the dye adsorption is important in determining the efficiency of the process. The pseudo-first-order, pseudo-second-order, Weber–Morris intraparticle diffusion and Boyd models were used for characterizing the kinetics data. The pseudo-first-order equation is expressed as follows:

$$\log(q_{\rm e} - q_{\rm t}) = \log q_{\rm e} - \frac{t}{2.303} k_1 \tag{9}$$

where q_e and q_t are the amount of dye adsorbed (mg g⁻¹) at initial time t and at equilibrium, respectively. k_1 is the pseudo-

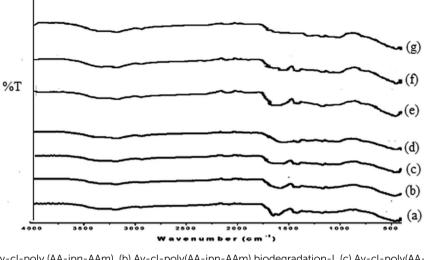
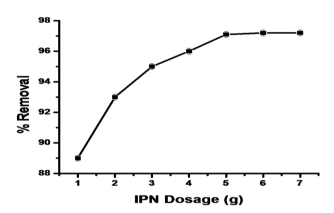


Fig. 7 FTIR spectra of (a) Av-cl-poly (AA-ipn-AAm), (b) Av-cl-poly(AA-ipn-AAm) biodegradation-I, (c) Av-cl-poly(AA-ipn-AAm) biodegradation-II, (d) Av-cl-poly(AA) biodegradation-III using the soil burial method; (e) Av-cl-poly(AA-ipn-AAm) biodegradation-I, (f) Av-cl-poly(AA-ipn-AAm) biodegradation-II and (g) Av-cl-poly(AA); biodegradation-III using the composting method.



The effect of IPN dosage on MG dye removal.

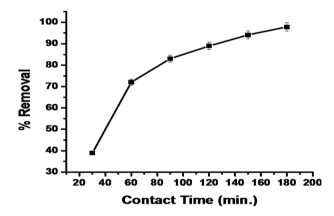


Fig. 10 The effect of contact time on MG dye adsorption by synthe-

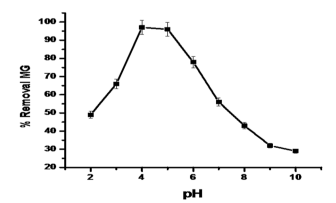


Fig. 9 The effect of pH on MG dye adsorption by synthesized IPN.

first-order rate constant (\min^{-1}) and t is the time taken (\min) . The values are derived from the linear plot of $ln(q_e - q_t)$ vs. t.

The pseudo-first-order equation differs from the actual equation in two ways: q_e does not represent the actual number of available sites and $\log q_e$ is not always equal to the intercept of the plot (log $q_e - q_t$) vs. t.

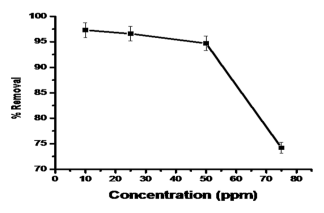


Fig. 11 The effect of initial dye concentration on MG dye adsorption by synthesized IPN.

Table 4 Pseudo first order and pseudo second order kinetics, intra particle diffusion constants and Boyd model

Pseudo first orde	er							
Conc. (ppm)	$K_1 \left(\min^{-1} \right)$	$q_{ m e,cal}~({ m mg~g^{-1}})$	$q_{\mathrm{exp}} (\mathrm{mg \ g^{-1}})$	R^2	χ^2	EABS	ARE	MSD
25	0.0403	0.54	0.61	0.953	63	96	46	56
50	0.0567	0.89	0.82	0.962	92	143	37	68
75	0.0539	0.32	0.25	0.968	126	187	42	62
Pseudo second o	rder kinetics							
Conc. (ppm)	$k_2 ({\rm g \ mg^{-1} \ h^{-1}})$	$q_{ m e,cal} ({ m mg \ g^{-1}})$	$q_{ m exp}~({ m mg~g^{-1}})$	R^2	χ^2	EABS	ARE	MSDF
25	0.558	0.68	0.56	0.982	12	32	14	21
50	0.776	0.94	0.87	0.991	7	28	8	14
75	0.501	0.39	0.26	0.987	6	21	6	16
Intra-particle dif	fusion							
Concentration								
(ppm)		$K_{\rm p}~({\rm mg~g}^{-1})$	$h^{-1/2}$)		C			R^2
25		0.469			0.41			0.968
50		0.562			0.62			0.986
75		0.332			0.34			0.964
Boyd model								
Concentration								
(ppm)		Slope		Int	tercept			R^2
25		0.016		0.0)72			0.942
50		0.014		0.1	124			0.951
75		0.015		0.1	179			0.959

The pseudo-second-order equation for the chemisorption kinetics is represented as

$$\frac{t}{q_t} = \frac{1}{q_e^2 k_2} + \frac{t}{q_e} \tag{10}$$

where k_2 is the pseudo-second-order rate constant (g mg⁻¹ min⁻¹) and it is calculated from the plot of t/q_t vs. t.

The parameters of the kinetics are summarised in Table 4. The most important determinant that defines the best fitting model for the experimental data is R^2 . It is quite clear from the

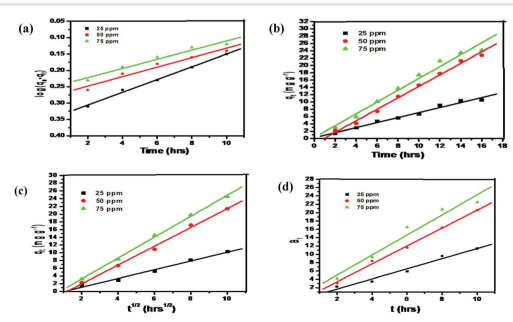


Fig. 12 (a) Pseudo first order, (b) pseudo second order, (c) intra particle diffusion and (d) Boyd plot.

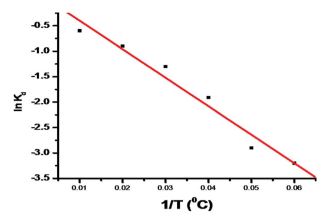


Fig. 13 The plot of $\ln K_d$ vs. 1/T for the evaluation of standard enthalpy and entropy changes.

results that the value of \mathbb{R}^2 for pseudo-second-order (0.991, Fig. 10b) is higher as compared to the pseudo-first order (0.968, Fig. 12a) and the values of error functions like χ^2 , EABS, ARE and MSDP were lower for the pseudo-second-order model as compared to the pseudo-first-order model, which indicated the lower experimental error. The calculated and experimental values of adsorption were very close for the pseudo-second-order. Therefore, based on this forgone discussion, it can be concluded that the pseudo-second-order model was best fit to the experimental data, thus the adsorption process may be of the chemisorption type.

The diffusion mechanism can be described by the Weber– Morris intra-particle diffusion and it is expressed by the equation given below:

$$q_t = k_3 t^{1/2} + C (11)$$

where k_3 and C are the intra-particle diffusion rate constant (mg g⁻¹ min^{-1/2}) and intercept, respectively. The C and k_3 parameters were determined from the linear plot of q_t vs. $t^{1/2}$.

The diffusion process may be of three different types: fast external diffusion, steady surface adsorption and slow equilibrium. At the initial stage of adsorption, there is fast external diffusion, which was not observed in Fig. 12b. This is followed by the surface adsorption intra-particle diffusion step, shown as the first linear region in the figure, which is followed by the constant equilibrium step, shown as the second linear region in the Fig. 12c. As per the Weber–Morris model, if the diffusion mechanism is controlled by the intraparticle diffusion, the intercept *C* should cross the origin. It is quite clear from the result (Table 4) that intercepts are non-zero, meaning that intraparticle diffusion is not the rate limiting step. 41-44

The diffusion mechanism was studied using the Boyd model and the equation is expressed as follows:

$$B_t = -0.4977 - \ln(1 - F) \tag{12}$$

 B_t is the mathematical function of F and is equivalent to the $q_t/q_{\rm e}$.

Table 5 Parameters of isotherm models for the adsorption of malachite green dye using synthesized IPN

Langmuir isotherm			
Temperature (°C)	$k_{\mathrm{L}} \left(\mathrm{L~mg}^{-1} \right)$	$q_{ m max~cal}~({ m mg~g}^{-1})$	R^2
25	0.862	1.21	0.994
50	0.955	1.12	0.997
75	0.767	1.54	0.982
Freundlich isotherm			
Temperature (°C)	$k_{\rm F} ({ m mg g}^{-1})$	N	R^2
25	1.189	1.14	0.991
50	1.316	1.36	0.989
75	1.376	1.47	0.978
Dubinin–Radushkevi	ch isotherm		
		E	
Temperature (°C)	$k_{\mathrm{D}} (\mathrm{mol^2 kJ^{-2}})$	(kJ mol ⁻¹)	R^2
25	2.561	0.389	0.783
50	3.421	0.289	0.672
75	4.762	0.326	0.746
Temkin isotherm			
Temperature (°C)	$b ext{ (kJ mol}^{-1})$	$k_{\mathrm{T}} \left(\mathrm{L} \; \mathrm{g}^{-1} \right)$	R^2
25	98.4	0.464	0.89
50	81.8	0.578	0.92
75	78.2	0.325	0.95
Redlich-Peterson iso	therm		
Temperature (°C)	$a_{\rm R} ({\rm L \ mg^{-1}})$	β	R^2
25	0.49	0.66	0.93
50	0.42	0.75	0.94
75	1.18	0.68	0.89
Sips isotherm model			
Temperature (°C)	$k_{\rm S} ({\rm L \ mg^{-1}})$	$q_{ m m} ({ m mg g}^{-1})$	R^2
25	0.872	1.36	0.962
	0.986	1.18	0.965
50	0.980	1.10	0.505

In the Boyd model, if the linear plot of B_t vs. t crosses the origin, then the diffusion process is controlled by particle diffusion and if it does not cross the origin, then the diffusion process is by film diffusion. It is clear from the results (Fig. 12d and Table 4) that the intercepts are non-zero, thus suggesting that the diffusion process is governed by film diffusion.^{41–50}

3.6. Adsorption isotherm models

The adsorption isotherm is a curve that describes the retention phenomena and mobility of substances from the porous medium to aquatic environments in a solid phase at constant pH and temperature. Adsorption equilibrium is achieved when

Table 6 Thermodynamic analysis at different temperatures^a

Temperature (°C)	$\ln k_{ m d}$	ΔG° (kJ mol ⁻¹)	ΔH° (kJ mol ⁻¹)	$\Delta \textit{S}^{\circ} \left(J \text{ mol}^{-1} \text{ K}^{-1} \right) \times 10^{-2}$	R^2
20	-0.49	1.24	-3.62	-1.64	0.982
30	-0.56	1.33			0.986
40	-0.63	1.49			0.988
50	-0.71	1.64			0.979
60	-0.77	1.75			0.983
70	-0.82	1.86			0.986

^a Where, k_d = thermodynamics equilibrium constant; ΔG° = Gibbs function; ΔH° = standard enthalpy change and ΔS° = standard entropy change.

the phase containing the adsorbate has been in contact with the adsorbent for an adequate time, with the concentration of the adsorbate in the bulk solution changing with the interface concentration. The mathematical correlation, like modelling analysis and operational design, is depicted by graphically expressing the solid-phase against its residual concentration. Physicochemical parameters with thermodynamic assumptions give an idea about the adsorption mechanism, the degree of affinity of the adsorbent and surface properties (Fig. 13).

There are many equilibrium isotherm models (Langmuir, Freundlich, Temkin, Toth, Brunauer-Emmett-Teller, Koble-Corrigan, Redlich-Peterson, Sips, Hill, Flory-Huggins, Khan and Radke-Prausnitz isotherm) that are used for kinetics considerations and adsorption equilibrium.⁴²⁻⁵¹

3.6.1. Two parameter isotherms

Langmuir isotherm model. This model was originally developed to study the gas-solid-phase adsorption onto activated carbon and this empirical model assumes monolayer adsorption on a finite number of definite sites, which are equivalent, with no lateral interactions and steric hindrance. The Langmuir isotherm generally refers to the homogeneous adsorption with no transmigration of the adsorbate. It is characterized graphically by a plateau and shows that when equilibrium is attained, once a molecule occupies a site, no further adsorption can take place. Langmuir theory also indicates that with an increase in distance, there is a rapid decrease in the attractive forces between the inter-molecules. The Langmuir equation is expressed as follows:

$$\frac{C_{\rm e}}{q_{\rm e}} = \frac{1}{k_{\rm L}q_{\rm m}} + \frac{C_{\rm e}}{q_{\rm m}} \tag{13}$$

where $k_{\rm L}$ is the Langmuir adsorption constant (L mg⁻¹) and $q_{\rm m}$ is the maximum monolayer adsorption capacity of the adsorbent (mg g⁻¹).

The essential characteristic of the Langmuir model is the separation factor ($R_{\rm L}$), which is a dimensionless constant. The separation factor indicates that if the isotherm is favourable, (0 < $R_{\rm L}$ < 1), linear ($R_{\rm L}$ = 1), unfavourable ($R_{\rm L}$ > 1) or irreversible ($R_{\rm L}$ = 0).

$$R_{\rm L} = \frac{1}{(1 + k_{\rm L} C_{\rm o})} \tag{14}$$

where C_0 is the highest initial dye concentration.

Freundlich isotherm model. This is the earliest known model, which describes the non-ideal and reversible adsorptions that are not restricted to the formation of a monolayer. This model assumes multilayer adsorption with the non-uniform distribution of adsorption heat and affinities over the heterogeneous surface. The Freundlich equation is expressed as follows:

$$\ln q_{\rm e} = \frac{1}{n_{\rm F}} \ln C_{\rm e} + \ln k_{\rm F} \tag{15}$$

where $n_{\rm F}$ is the Freundlich constant indicating the favourability of the adsorption process, which is favourable if $1 < n_{\rm F} < 10$. $k_{\rm F} \ ({\rm mg}^{1-1/n}\ {\rm L}^{1/n}\ {\rm g}^{-1})$ is the adsorption capacity of the adsorbent.

Dubinin–Radushkevich (D–R) isotherm model. D–R is an empirical model, which typically considers the adsorption of subcritical vapours onto the porous solid. It is used to express the adsorption mechanism with a Gaussian energy distribution on a heterogeneous surface. The D–R isotherm is a temperature dependent isotherm model and assumes that the adsorbent surface is heterogeneous and the equation is expressed as follows:

$$\ln q_{\rm e} = \ln q_{\rm m} - k_{\rm D} \varepsilon^2 \tag{16}$$

$$\varepsilon = RT \ln 1 + \frac{1}{C_c} \tag{17}$$

where $k_{\rm D}$ (mol² kJ⁻²), ε and R (8.314 \times 10⁻³ kJ mol⁻¹ K⁻¹) are the D–R constant and D–R isotherm constant and gas constant, respectively. $q_{\rm m}$ (mg g⁻¹) and T (K) are the saturation capacity and temperature, respectively. The $k_{\rm D}$ is used to obtain the mean free energy, *i.e.* E (kJ mol⁻¹) using the equation given below:

$$E = \sqrt{\frac{1}{2k_{\rm D}}} \tag{18}$$

Temkin isotherm model. This isotherm is the early model, which describes the hydrogen adsorption onto platinum electrodes in an acidic solution. This isotherm tells about the interaction between adsorbate and adsorbent. It is a two-parameter model, which considers the uniform distribution of energy. It is assumed that in the layer, the heat of adsorption of all the molecules decreases linearly rather than logarithmically. The Temkin equation is expressed as follows:

$$q_{\rm e} = \frac{RT}{b} \ln k_{\rm T} C_{\rm e} \tag{19}$$

where $k_{\rm T}$ is the equilibrium binding constant and b (kJ mol⁻¹) is the Temkin isotherm constant.

3.6.2. Three parameter isotherms

Redlich-Peterson (R-P) isotherm model. The R-P isotherm is attributed to both Langmuir and Freundlich isotherms, which incorporate three parameters into an empirical equation. To represent the adsorption equilibria over a wide range of concentrations, this model is dependent on the concentration in the numerator and an exponential function in the denominator to represent the adsorption equilibria over a large range of pH. This model is applicable to both homogeneous and heterogeneous systems. The R-P equation is expressed as follows:

$$q_{\rm e} = \frac{k_{\rm R} C_{\rm e}}{1 + a_{\rm R} C_{\rm e}^{\beta}} \tag{20}$$

where $k_{\rm R}$ (L g⁻¹) and $a_{\rm R}$ (L mg⁻¹) are the R-P constants. β is the R-P exponent.

Sips isotherm model. This model is a combination of the Langmuir and Freundlich expressions, used for the heterogeneous adsorption system. This isotherm overcomes the limitation of an increase in the concentration of adsorbate associated with the Freundlich isotherm. It predicts a monolayer adsorption capacity characteristic of the Langmuir isotherm at high concentration, and at low adsorbate concentration it is reduced to the Freundlich isotherm. The equation parameters are generally governed by operating conditions like temperature, pH and concentration. The equation is expressed as follows:

$$q_{\rm e} = \frac{q_{\rm m}k_{\rm S}C_{\rm e}^{k_{\rm LF}}}{1 + k_{\rm S}C_{\rm e}^{k_{\rm LF}}} \tag{21}$$

where $k_{\rm S}$ (Lg⁻¹) and $k_{\rm LF}$ are the Sips constant and Sips exponent, respectively.

The parameters of the Langmuir, Freundlich, D–R, Temkin, R–P and Sips isotherm models were obtained from the linear plots of C_e/q_e vs. C_e , $\ln q_e$ vs. $\ln C_e$, $\ln q_e$ vs. ε^2 , q_e vs. $\ln C_e$, $\ln [(k_R(C_e/q_e) - 1]$ vs. $\ln C_e$ and $\ln [q_e/(q_m - q_e)]$ vs. $\ln C_e$, respectively^{46–50} (Table 5).

The value of \mathbb{R}^2 was compared for all the models and it was observed that the Langmuir model had the highest \mathbb{R}^2 value, suggesting that the experimental data is best represented by the Langmuir model. The results from the Langmuir model specify that on the surface of the adsorbent, there is a homogeneous distribution of adsorption sites and therefore, there is the formation of a single layer of dye molecules on the IPN surface.

3.7. Thermodynamics studies and activation energy

The thermodynamics studies provide information about the effect of temperature on the adsorption process and were conducted at temperatures ranging from 20 to 70 $^{\circ}$ C. The following equations were used for obtaining the thermodynamic parameters. $^{51-53}$

$$\Delta G^{\circ} = \Delta H^{\circ} - T \Delta S^{\circ} \tag{22}$$

$$\Delta G^{\circ} = -RT \ln k_{\rm D} \tag{23}$$

$$k_{\rm D} = \frac{q_{\rm e}}{C_{\rm o}} \tag{24}$$

The temperature effect on the thermodynamic constant can be calculated using the following equation:

$$\frac{\mathrm{dln}\,k_{\mathrm{D}}}{\mathrm{d}T} = \frac{\Delta H^{\circ}}{RT^{2}}\tag{25}$$

On integration and rearrangement, the equation becomes

$$\ln K_{\rm d} = -\frac{\Delta H^{\circ}}{RT} + \frac{\Delta S^{\circ}}{R} \tag{26}$$

where ΔH° , ΔG° and ΔS° are the change in enthalpy, Gibbs free energy and the change in entropy, respectively. k_D and T(K) are the distribution coefficient for adsorption and temperature, respectively. q_e and C_e (mg L⁻¹) are the concentration of dye adsorbed by the adsorbent and dye remaining in the solution at equilibrium, respectively. R (J mol⁻¹ K⁻¹) is the gas constant. K_d was obtained from dye adsorption at different concentrations at different temperatures. The results of $\ln K_d$ vs. 1/T (Fig. 11) prove that the dye removal decreases as the temperature increases, thus indicating the exothermic nature of dye removal. The value of ΔG° is positive at all temperatures and it decreases with increasing temperature (Table 6). This shows that the dye removal process is not spontaneous and the dye removal is higher at low temperatures. The negative value of ΔH° confirms that the dye removal process is exothermic in nature.52-55

4. Conclusion

A noble IPN was synthesized using the natural polysaccharide, Av, as the backbone, APS as the initiator, MBA as the crosslinker, AA and AAm as primary and secondary monomers, respectively. The biodegradability of the synthesized IPN was investigated through soil burial and composting methods and it is clear from the results that the synthesized IPN is biodegradable by natural microorganisms. The synthesized IPN was also used for the removal of MG dye from aqueous solution and was found to remove 97.3% of MG dye. The synthesized device is very effective for MG dye adsorption from aqueous solution. The adsorption of MG dye on the synthesized sample followed the Langmuir and Freundlich isotherms. The kinetics of adsorption followed a pseudo-second-order rate equation. The Langmuir isotherm model was found to be the best-fitting model for the present experimental data. MG dye removal decreased as the temperature increased, suggesting that the adsorption process was exothermic. Thus, the synthesized IPN was found to be an efficient, cost-effective and environmentally friendly adsorbent for the removal of MG dye from aqueous solution.

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Conflicts of interest

There are no conflicts to declare.

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References

- 1 Saruchi and V. Kumar, Adsorption kinetics and isotherms for the removal of rhodamine B dve and Pb2+ ions from aqueous solutions by a hybrid ion-exchanger, Arabian J. Chem., 2016, DOI: 10.1016/j.arabjc.2016.11.009.
- 2 V. P. Mahida and M. P. Patel, Removal of some most hazardous cationic dyes using novel poly (NIPAAm/AA/Nallylisatin) nanohydrogel, Arabian J. Chem., 2016, 9, 430-442.
- 3 S. Thakura, S. Pandeya and O. Arotibaa, Sol-gel derived xanthan gum/silica nanocomposite—a highly efficient cationic dyes adsorbent in aqueous system, Int. J. Biol. Macromol., 2017, 103, 596-604.
- 4 B. Mua and A. Wanga, Adsorption of dyes onto palygorskite and its composites, J. Environ. Chem. Eng., 2016, 4, 1274-
- 5 V. Bekiari and P. Lianos, Poly (sodium acrylate) hydrogels as potential pH-sensitive sorbents for the removal of organic and inorganic pollutant from water, Global NEST J., 2010, 12, 262-269.
- 6 S. Thakura, S. Pandeya and O. Arotibaa, Development of a sodium alginate-based organic/inorganic superabsorbent composite hydrogel for adsorption of methylene blue, Carbohydr. Polym., 2016, 153, 34-46.
- 7 R. Sharma, B. S. Kaith, S. Kalia, D. Pathania, A. Kumar, N. Sharma, R. Street and C. Schauer, Biodegradable and conducting hydrogels based on Guar gum polysaccharide for antibacterial and dye removal applications, J. Environ. Manage., 2015, 162, 37-45.
- 8 V. Panic, Z. Madzarevic, T. Husovic and S. Velickovic, Poly(methacrylic acid) based hydrogels as sorbents for removal of cationic dye basic yellow 28: Kinetics, equilibrium study and image analysis, Chem. Eng. J., 2013, 217, 192-204.
- 9 S. Li, H. Zhang, J. Feng, R. Xu and X. Liu, Facile preparation of poly (acrylic acid-acrylamide) hydrogels by frontal polymerization and their use in removal of cationic dyes from aqueous solution, Desalination, 2011, 280, 95-102.
- 10 N. Alizadeh and M. Mahjoub M, Removal of crystal violet dye from aqueous solution using surfactant modified NiFe₂O₄ as nanoadsorbent, isotherms, thermodynamics and kinetics studies, I. Nanoanalysis, 2017, 4, 8-19.
- 11 M. Dahri, L. Lim, N. Priyantha and C. Chan, Removal of Acid Blue 25 Using Cempedak Durian Peel From Aqueous

- Medium: Isotherm, Kinetics and Thermodynamics Studies, Int. Food Res. J., 2016, 23, 1154-1163.
- 12 L. Lim, N. Priyantha, H. Cheng and N. Zaidi, Adsorption characteristics of Artocarpus odoratissimus leaf toward toxic crystal violet dye: thermodynamics and regeneration studies, Journal of Environment and Biotechnology Research, 2016, 4, 32-40.
- 13 N. Zaidi, L. Lim, A. Usman and M. Kooh M, Efficient adsorption of malachite green dye using Artocarpus odoratissimus leaves with artificial neural network modelling, Desalin. Water Treat., 2018, 101, 313-324.
- 14 L. Lim, N. Priyantha, C. Ing, M. Dahri, D. Tennakoon, T. Zehra and M. Suklueng, Artocarpus odoratissimus skin as a potential low-cost biosorbent for the removal of methylene blue and methyl violet 2B, Desalin. Water Treat., 2015, 53, 1-13.
- 15 H. Chieng, L. Lim and N. Priyantha, Enhancement of crystal violet dye adsorption on Artocarpus camansi peel through sodium hydroxide treatment, Desalin. Water Treat., 2017, 58, 320-331.
- 16 L. Lim and N. Priyantha, Potential use of Momordica charantia (bitter gourd) waste as a low-cost adsorbent to remove toxic crystal violet dye, Desalin. Water Treat., 2017, 82, 121-130.
- 17 N. Priyantha, L. Lim, D. Tennakoon, E. Liaw, C. Ing and A. Liyandeniya, Biosorption of cationic dyes on breadfruit (Artocarpus altilis) peel and core, Appl. Water Sci., 2018, 8, 37.
- 18 L. Lim, N. Priyantha, Y. Chen and N. Zaidi, Effective removal of methyl violet dye using pomelo leaves as a new low-cost adsorbent, Desalin. Water Treat., 2018, 110, 264-274.
- 19 P. K. Sahu, D. D. Giri, R. Singh, P. Pandey, S. Gupta, A. K. Shrivastava, A. Kumar and K. D. Pandey, Pharmacol. Pharm., 2013, 4, 599-610.
- 20 R. F. Pereira, A. Mendes and P. J. Bartolo, Novel Alginate/Aloe Vera Hydrogel Blends as Wound Dressings for the Treatment of Several Types of Wounds, Chem. Eng. Trans., 2013, 32, 1009-1014.
- 21 N. Yewegnon, E. A. Irma, S. Philippe, A. Abdoukarim, Y. A. K. Alassane, A. C. Pascal, M. Daouda and S. K. C. Dominique, Evaluation of Aloe Vera Leaf Gel as a Natural Flocculant: Phytochemical Screening and Turbidity Removal Trials of Water by Coagulation Flocculation, Res. J. Recent Sci., 2016, 5, 9-15.
- 22 O. Khyati, S. Vranda, S. Gupta and S. R. Suseem, Formulation and evaluation of hydrogel with ascorbic acid using aloe vera gel powder as a drug carrier, Innovare Journal of sciences, 2013, 1, 18-20.
- 23 Y. S. Ho, C. C. Chiang and Y. C. Hsu, Sorption kinetics for dye removal from aqueous solution using activated clay, Sep. Sci. Technol., 2001, 36(11), 2473-2488.
- 24 T. R. Hoare and D. S. Kohane, Hydrogels in drug delivery: progress and challenge, Polymer, 2008, 49, 1993-2007.
- 25 Y. Wang, W. W. Wanga and A. AiqinWanga, Efficient adsorption of methylene blue on an alginate-based nanocomposite hydrogel enhanced by organo-illite/ smectite clay, Chem. Eng. J., 2013, 228, 132-139.

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26 J. Arutchelvi, M. Sudhakar, A. Arkatkar, M. Doble, S. Bhaduri and P. V. Uppara, Biodegradation of polyethylene and polypropylene, *Indian J. Biotechnol.*, 2008, 7, 9–22.

- 27 Saruchi, B. S. Kaith, V. Kumar and R. Jindal, Biodegradation study of enzymatically catalyzed interpenetrating polymer network: evaluation of agrochemical release and impact on soil fertility, *Biotechnol. Rep.*, 2016, **9**, 74–81.
- 28 M. Chigondo, H. K. Paumo, M. Bhaumik, K. Pillay and A. Maity, Rapid high adsorption performance of hydrous cerium magnesium oxides for removal of fluoride from water, *J. Mol. Liq.*, 2018, DOI: 10.1016/j.molliq.2018.06.015.
- 29 A. Alvarez, L. H. Velazquez-Jimenez, L. P. Chazaro-Ruiz, P. E. Diaz-Flores and J. R. Rangel-Mendez, F⁻ removal in water by a hybrid adsorbent lanthanum-carbon, *J. Colloid Interface Sci.*, 2015, 455, 194–202.
- 30 A. Albertsson, C. Anderson and S. Karlsson, The mechanism of biodegradation of polyethylene, *Polym. Degrad. Stab.*, 1987, **18**, 73–87.
- 31 A. S. Sarc, Redox polymerization, *Prog. Polym. Sci.*, 1999, 24, 1149–1204.
- 32 Saruchi, B. S. Kaith, R. Jindal and V. Kumar, Biodegradation of gum tragacanth acrylic acid based hydrogel and its impact on soil fertility, *Polym. Degrad. Stab.*, 2015, **115**, 24–31.
- 33 Saruchi, B. S. Kaith, R. Jindal, G. S. Kapur and V. Kumar, Synthesis, characterization and evaluation of gum tragacanth and acrylic acid based hydrogel for sustained calcium chloride release enhancement of water-holding capacity of soil, *J. Chin. Adv. Mater. Soc.*, 2014, 2(1), 40–52.
- 34 H. Mittal, V. Kumar, Saruchi and S. S. Ray, Adsorption of methyl violet from aqueous solution using gum xanthan/ Fe₃O₄ based nanocomposite hydrogel, *Int. J. Biol. Macromol.*, 2016, 89, 1–11.
- 35 M. Ghaedi, N. Zeinali, A. M. Ghaedi, M. Teimuori and J. Tashkhourian, Artificial neural network-genetic algorithm based optimization for the adsorption of methylene blue and brilliant green from aqueous solution by graphite oxide nanoparticle, *Spectrochim. Acta, Part A*, 2014, **125**, 264–277.
- 36 M. Ghaedi, A. M. Ghaedi, M. Hossainpour, A. Ansari, M. H. Habibi and A. R. Asghari, Least square-support vector (LS-SVM) method for modeling of methylene blue dye adsorption using copper oxide loaded on activated carbon: kinetic and isotherm study, *J. Ind. Eng. Chem.*, 2014, 20(4), 1641–1649.
- 37 H. Mittal, A. Maity and S. S. Ray, The adsorption of Pb²⁺ and Cu²⁺ onto gum ghatti-grafted poly(acrylamide-co-acrylonitrile) biodegradable hydrogel: isotherms and kinetic models, *J. Phys. Chem. B*, 2015, **119**(5), 2026–2039.
- 38 S. Ghorai, A. Sarkar, M. Raoufi, A. B. Panda, H. Schonherr and S. Pal, Enhanced Removal of Methylene Blue and Methyl Violet Dyes from Aqueous Solution Using a Nanocomposite of Hydrolyzed Polyacrylamide Grafted Xanthan Gum and Incorporated Nanosilica, ACS Appl. Mater. Interfaces, 2014, 6(7), 4766-4777.
- 39 B. S. Kaith, Saruchi, R. Jindal and M. S. Bhatti, Screening and RSM optimization for synthesis of a gum tragacanth–acrylic

- acid based device for *in situ* controlled cetirizine dihydrochloride release, *Soft Matter*, 2012, **8**, 2286–2293.
- 40 M. Ghaedi, A. Ansari, M. H. Habibi and A. R. Asghari, Removal of malachite green from aqueous solution by zinc oxide nanoparticle loaded on activated carbon: kinetics and isotherm study, J. Ind. Eng. Chem., 2014, 20(1), 17–28.
- 41 V. K. Gupta, D. Patania, S. Agarwal and S. Sharma, Removal of Cr(VI) onto *Ficus carica* biosorbent from water, *Environ. Sci. Pollut. Res.*, 2013, 20, 2632–2644.
- 42 M. K. Dahri, M. R. R. Kooh and L. B. L. Lim, Water remediation using low cost adsorbent walnut shell for removal of malachite green: equilibrium, kinetics, thermodynamic and regeneration studies, *J. Environ. Chem. Eng.*, 2014, 2(3), 1434–1444.
- 43 V. K. Gupta, D. Patania, S. Agarwal and S. Sharma, Removal of Cr(VI) onto *Ficus carica* biosorbent from water, *Environ. Sci. Pollut. Res.*, 2013, 20, 2632–2644.
- 44 I. Langmuir, The adsorption of gases on plane surfaces of glass, mica and platinum, *J. Am. Chem. Soc.*, 1918, **40**, 1361–1403.
- 45 D. G. Kinniburgh, General purpose adsorption isotherms, *Environ. Sci. Technol.*, 1986, **20**, 895–904.
- 46 M. Dogan, M. Alkan, A. Turkyilmaz and Y. Ozdemir, Kinetics and mechanism of removal of methylene blue by adsorption onto perlite, *J. Hazard. Mater.*, 2004, 109, 141–148.
- 47 M. Roosta, M. Ghaedi, A. Daneshfar, R. Sahraei and A. Asghari, Optimization of the ultrasonic assisted removal of methylene blue by gold nanoparticles loaded on activated carbon using experimental design methodology, *Ultrason. Sonochem.*, 2014, 21(1), 242–252.
- 48 M. Ghaedi, A. M. Ghaedi, E. Negintaji, A. Ansari and F. Mohammadi, Artificial neural network imperialist competitive algorithm based optimization for removal of sunset yellow using Zn(OH)₂ nanoparticles-activated carbon, *J. Ind. Eng. Chem.*, 2014, **20**(6), 4332–4343.
- 49 H. L. Chieng, L. B. L. Lim and N. Priyantha, Sorption characteristics of peat from Brunei Darussalam for the removal of rhodamine B dye from aqueous solution: adsorption isotherms, thermodynamics, kinetics and regeneration studies, *Desalin. Water Treat.*, 2015, 55, 664– 677.
- 50 O. Hamdaoui and E. Naffrechoux, Modeling of adsorption isotherms of phenol and chlorophenols onto granular activated carbon part II. Models with more than two parameter, *J. Hazard. Mater.*, 2007, 147, 401–411.
- 51 M. S. Chiou and H. Y. Li, Equilibrium and kinetic modeling of adsorption of reactive dye on cross-linked chitosan beads, *J. Hazard. Mater.*, 2002, **93**, 233–248.
- 52 H. I. Chieng, L. B. L. Lim and N. Priyantha, Enhancing adsorption capacity of toxic malachite green dye through chemically modified breadnut peel: equilibrium, thermodynamics, kinetics and regeneration studies, *Environ. Technol.*, 2015, **36**(1), 86–97.
- 53 O. S. Bello, M. A. Ahmad and B. Semire, Scavenging malachite green dye from aqueous solutions using pomelo (*Citrus grandis*) peels: kinetic, equilibrium and

- thermodynamic studies, *Desalin. Water Treat.*, 2015, 56(2), 647–658.
- 54 L. B. L. Lim, N. Priyantha and N. H. M. Mansor, Utilizing *Artocarpus altilis*(breadfruit) skin for the removal of malachite green: isotherm, kinetics, regeneration, and
- column studies, *Desalin. Water Treat.*, 2016, 57(35), 16601–16610.
- 55 M. R. R. Kooh, M. K. Dahri and L. B. L. Lim, The removal of rhodamine B dye from aqueous solution using *Casuarina equisetifolia* needles as adsorbent, *Cogent Environ. Sci.*, 2016, 2457, 1–14.