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1	Removal of Malachite Green and Crystal Violet Dyes from
2	Aqueous Solution with Bio-Materials: A Review
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#### 7 Abstract

8 Malachite Green and Crystal violet are among millions of dyes which are being used in every

<sup>9</sup> aspect of day to day life of a human being. Approximately 12

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11 Index terms— malachite green, crystal violet, dyes, toxic, biomaterials.

### <sup>12</sup> 1 Removal of Malachite Green and Crystal Violet

Dyes from Aqueous Solution with Bio-Materials: A Review Binod Kumar ? & Upendra Kumar ? Abstract-13 Malachite Green and Crystal violet are among millions of dyes which are being used in every aspect of day to 14 day life of a human being. Approximately 12% of synthetic dyes are lost during manufacturing and processing 15 operations and 20% of the resultant color enters the environment through effluents from industrial wastewater. 16 They are toxic and having extremely harmful consequences; hence many governmental and environmental agencies 17 have put in place very strict regulation and restriction on discharge of industrial waste water/effluent containing 18 dyes into the natural water bodies. There are various technique available for removal of dyes from waste water 19 but adsorption is the process of choice. Activated carbon is the best known adsorbent. But its use in treating 20 the industrial waste water especially in developing countries is restricted due to very high cost. This high cost of 21 activated carbon has forced the researchers to find out low cost and effective adsorbent which may be used as an 22 efficient alternative of activated carbon. In this paper an attempt has been made to compile the work of various 23 24 researchers on removal of crystal violet and malachite green dyes from aqueous solution by using biomaterials and agricultural waste during the last five years. 25

## 26 2 Introduction

yes in different form are being used in every aspect of day to day life of a human being. At present there are 27 more than 1,00,000 Commercial dyes with rough estimates of production range between 7x10 5 tons to 1x1028 6 tons per year [1] Among them Crystal Violet and malachite green are one of the widely used synthetic and 29 cationic dyes. The molecular formula of crystal violet and malachite green are C 25 N 3 H 30 Cl and C 23 N 30 2 H 25 Cl, the molecular mass are 407.979 g mol ?1 and 364.91 g mol ?1 and melting point are 205  $^{\circ}\mathrm{C}$  and159 31 °C respectively. Both the dyes are soluble in Water and Alcohol. They are Stable, Incompatible with strong 32 oxidizing agents, strong acids, Light-sensitive and Combustible [2]. Color Index Number (CI) of crystal violet 33 and malachite green are 42555 and 42000 respectively. When crystal violet is dissolved in water the dye has 34 a blue-violet colour with an absorbance maximum at 590 nm. Similarly when malachite green is dissolved in 35 36 water the dye has a green colour with an absorbance maximum at 617 nm. Both the dyes are extensively used 37 as a dye in paper and pulp industries, leather industries, and textile industries. They are also widely used for 38 various medicinal purposes like as a biological stain for microscopic analysis of cell biology and tissue samples. 39 Approximately 12% of synthetic dyes are lost during manufacturing and processing operations and 20% of the resultant color enters the environment through effluents from industrial wastewater [3]. The presence of very small 40 amounts of dyes in water (less than 1 ppm for some dyes) is highly visible and undesirable on aesthetic ground. 41 [4]. It also interferes with transmission of light and upset the biological metabolism process which causes the 42 destruction of aquatic communities present in Eco-system. Further, dye has the tendency to sequester metal 43 and may cause micro toxicity to fish and other organism. Due to these harmful consequences many countries 44

have put in place very stringent regulation. Due to enforcement of very stringent regulation by the various 45 environmental and governmental agencies upon the industries ,the interest of researcher have moved towards to 46 find the cost effective methods for removal of dyes from aqueous solution. The waste water containing dye(s) 47 is difficult to treat since the dyes are very complex organic molecule, resistance to aerobic digestion and are 48 stable to light, heat and oxidizing agents. [5]. during the past three decades, several physical, chemical and 49 biological decolourization processes have been reported. They are Coagulation, Flocculation, Biodegradation, 50 Adsorption on activated carbons, Membrane separations, Ionexchange, Oxidation, Advanced oxidation process, 51 Biomass, Selective biosorbents. However, these methods have several disadvantages that include incomplete 52 dyes removal, high reagent and energy requirements and generation of toxic sludge or other waste products that 53 requires proper disposal and further treatment. These methods are very costly, making them uneconomical and 54 unviable. Amongst the numerous techniques of dye removal, adsorption is the procedure of choice and gives 55 the best results. [6] Commercially Activated carbon adsorption has been cited as one of the best dyes control 56 technology by the US Environmental protection Agency [7]. The very high capacity of Commercially Activated 57 carbon for removal of dye through adsorption is mainly due to their structural characteristics and their porous 58 texture which gives them a large surface area, and their chemical nature which can be easily modified by chemical 59 treatment in order to increase their properties. However, activated carbon presents several disadvantages. It is 60 61 quite expensive, the higher the quality, the greater the cost, non-selective and ineffective against disperse and 62 vat dyes. The regeneration of saturated carbon is also expensive, not straightforward, and results in loss of the 63 adsorbent [8]. Due to the problems mentioned above, research interest into the production of alternative sorbents to replace the costly activated carbon has intensified in recent years. Attention has focused on various natural 64 solid supports, which are able to remove pollutants from contaminated water at low cost. Cost is actually an 65 important parameter for comparing the adsorbent materials. According to Bailey et al. (1999) [10] a sorbent can 66 be considered low-cost if it requires little processing cost, is abundant in nature or is a byproduct or waste material 67 from another industry. Recently attention has been diverted towards the biomaterials which are byproducts or 68 the wastes from large scale industrial operations and agricultural waste materials. The major advantages of 69 biosorption over conventional treatment methods are: low cost, high efficiency, minimization of chemical or 70 biological sludge, requirement of additional nutrient, and regeneration of biosorbents, and possibility of dyes 71 recovery. Presently, agricultural waste materials have been proposed as economic and eco-friendly adsorbents. 72 Agricultural materials are particularly those containing cellulose which shows a high potential in dye biosorption 73 74 capacity. The adsorption process of dyes presently in aqueous solution by low-cost adsorbents from plant wastes 75 can be carried with or without chemical modifications. Generally, chemically modified plant wastes are exhibiting 76 the higher adsorption capacities than unmodified forms. More recently, the great efforts have been directed to develop a new adsorbent and improving the existing adsorbents to have an alternative to activated carbon. 77

In this paper an attempt has been made to compile the work of various researchers on removal of crystal violet and malachite green dyes from aqueous solution by using biomaterials and agricultural waste during the last five years.

#### <sup>81</sup> **3 II**.

#### <sup>82</sup> 4 Literature Review

On scanning the available literature on the subject matter it has been observed that researchers have investigated 83 the effectiveness of biomaterials and agricultural wastes in removal of crystal violet and malachite green dyes from 84 aqueous solution by adsorption in batch mode isotherm experiment. They have examined the effect of rotational 85 86 speed of the shaker, pH value of the solution, contact time, initial concentration of dyes in the solution, dose 87 of adsorbent, temperature of solution on the adsorption of dyes by biosorbents. Adsorption isotherm modeling, rate kinetic modeling, intra-particle diffusion analysis, thermodynamic analysis have been done by them on the 88 experimentally observed data. Wang et al (2008) investigated the removal of malachite green from aqueous 89 solution using agricultural by-products (wheat bran and rice bran). It was found that adsorption increases with 90 increase in solution pH and attained equilibrium at 90 min. The equilibrium data of malachite green fitted 91 better to Freundlich model. The kinetic data fitted better to pseudo-second order equation. While the external 92 diffusion was the rate controlling step of the initial fast adsorption (< 15 minutes) and in the next stage the 93 intraparticle diffusion dominated the mass transfer. The thermodynamic analysis revealed that the adsorption 94 of MG onto wheat bran and rice bran is spontaneous and favorable process [11]. Baseri et al (2012) investigated 95 the adsorption of basic dyes (malachite green, crystal violet and rhodamine B) from synthetic textile effluent by 96 97 activated carbon prepared from Thevetia peruviana. It was found that the amount of dye adsorbed increased 98 with increasing initial dye concentration. The process was endothermic in nature. Kinetic studies showed that 99 the adsorption of these dyes followed pseudo-second order model with multistep intra particle diffusion model. 100 The data obtained from adsorption isotherms are well fitted with Langmuir model, the negative I?"G O value 101 obtained suggested the process was spontaneous in nature [12].

Prasad et al (2012) studied the adsorption of cationic dyes (crystal violet and rhodamine B) from aqueous solution onto Acacia nilotica leaves as an eco-friendly adsorbent. The result showed that the optimum pH was 6 and the equilibrium time was 120 minutes. The removal efficiency increased with increase in agitation time and initial dye concentration. The equilibrium data were best described by Langmuir isotherm model with maximum

monolayer capacity of 33 and 37 mgg-1 for crystal violet and rhodamine B respectively. The adsorption kinetics 106 can be successfully fitted to pseudo-second order kinetic model. The result of the intraparticle diffusion model 107 suggested that intraparticle diffusion might not be the only rate controlling step. Desorption study revealed that 108 the recovery of the dye from adsorption was possible [13]. Tahir et al. [2009] investigated the removal of malachite 109 green by using household used black tea as an adsorbent. The adsorbent was used in raw form and impregnated 110 form. The equilibrium time of adsorption came out to be 10 minutes. Also, the data showed that there was 111 decrease in the amount of the dyes adsorption with increase in temperature. The thermodynamic parameter 112 showed that the process was exothermic and spontaneous. Furthermore, the result showed that the raw form of 113 the adsorbent showed better adsorption capacity compared to its impregnated form [14]. Kumar et al. (2005) 114 investigated the adsorption of malachite green onto Pithophora sp., fresh water algae. The algae were used raw 115 and some were thermally activated at 300 O C for 50 minutes. The equilibrium data was very well represented 116 by Redlich-Peterson isotherm. It was found that the thermally activated adsorbent possess a higher sorption 117 capacity of 117.647mg/g than the raw one with 64.4mg/g. The kinetic data followed the pseudo-second order 118 model closely and the negative value of Gibb's free energy change (Î?"G?) indicated that the sorption process 119 was spontaneous [15]. Remenarova et al. [2009] studied the sorption of Malachite Green from aqueous solutions 120 by moss Rhytidiadelphus squarrosus. The results showed that the equilibrium was reached within 1-2 hours. 121 122 The equilibrium data best fitted the Freundlich isotherm model [16]. Uma et al. (2013) used Sawdust a timber 123 waste as a bio-sorbent for the removal of a cationic dye, malachite green from aqueous solutions. Point zero 124 charge (pHzpc) of raw sawdust was calculated by titrimetric method and that was 7.37. FTIR was performed to determine various functional groups attached to the adsorbent surface. The experimental results indicate 125 that 5g/L g of sawdust was able to remove 93% of dye from an initial concentration of 20 mg/ L. Equilibrium 126 was achieved in 100 min and the maximum adsorption of malachite green occurred in the higher (alkaline) pH 127 range. The data was better fitted in Freundlich isotherm model. It was observed that dye removal decrease with 128 increasing concentrations of dye solution. The maximum adsorption capacity of adsorbent was found to be 13.87 129 mgg?1 at 298 K. The dye adsorption followed the pseudo second-order kinetics. Removal process was endothermic 130 in nature [17]. Jiao ?? i et al. (2013) investigated the removal of malachite green (MG) from aqueous solution 131 using modified peanut shell (MPS). The SEM and FTIR showed that some components of raw peanut shell (RPS) 132 had been removed during the chemical modification, and many cavities of various dimensions were clearly evident 133 on the surface of MPS. Adsorption experiments showed that MG adsorption uptake was increased with an increase 134 in initial concentration, contact time, solution temperature, and adsorbent dosage. Furthermore, neutral pH was 135 optimum for the removal of MG. The adsorption of MG onto MPS agreed well with the nonlinear Langmuir and 136 Sips isotherm models. The monolayer capacity (Q max ) was 32.73 or 35.85mg/g as calculated from Langmuir or 137 Sips isotherm models, respectively. The adsorption kinetic studies showed that the adsorption process followed 138 the pseudo-second-order kinetic model with a multi-step diffusion process ??18]. Chen Z et al. (2014) used the 139 Pleurotus ostreatus (a macro-fungus) as a new biosorbents to study the biosorption of hazardous malachite green 140 (MG) from aqueous solutions. The equilibrium data best fitted with Freundlich isotherm model. The biosorption 141 process followed the pseudosecond-order kinetic model. The Fourier transform infrared spectroscopy (FTIR) 142 showed the presence of the functional groups such as, carboxyl, hydroxyl, amino and phosphonate groups on the 143 biosorbents surface which could be the potential adsorption sites for MG biosorption ??19]. 144

Ashish S. Sartape et al. (??013) investigated successfully Wood apple shell (WAS) alternative adsorbent for 145 the removal of hazardous dye malachite green. The removal of MG dye was found to be 98.87% with initial 146 concentration 100 mg/L at pH 7-9 in 3.30 h by shaking at 150 rpm at 299  $\pm$  2 K. The shifting of peaks in 147 FTIR spectrum confirmed the MG dye adsorption onto WAS. The SEM study also made support to it by 148 observing difference in surface morphology of adsorbent before and after adsorption of MG. The adsorption 149 equilibrium data showed good fit to the Langmuir isotherm model as compared to the Freundlich isotherm 150 model. The adsorption capacity for WAS was increased from 12.35 to 80.645 mg/g as the MG concentration in 151 the test solution was increased from 100 to 700 mg/L. The adsorption kinetics followed pseudo first-order kinetic 152 equation for sorption of MG onto WAS. Thermodynamic study demonstrates the spontaneous and endothermic 153 nature of biosorption process due to negative values of free energy change  $\hat{I}$ ?"G? and positive value of enthalpy 154 change I?"H?, respectively. ??20] Mi-Hwa Baek et al. (2010) studied the degreased coffee beans as an adsorbent 155 for removal of MG from aqueous solutions. The adsorption was highly dependent on initial dye concentration, 156 temperature and pH. MG adsorption onto DCB reached almost equilibrium in about 4 h MG removal by RCB 157 reached equilibrium in about 8h. The optimal pH was 10-12. The result of the present investigations showed that 158 degreased coffee beans (DCB) have higher adsorption efficiency than raw coffee beans. The adsorbed amount 159 of MG increased as initial MG concentration increase. The kinetic data best fitted the pseudo second-order 160 kinetic model though the correlations coefficients from the pseudo first-order kinetic were as well relatively high 161 for the range of concentrations studied. On the basis of Correlation coefficients (R2) values it is revealed that 162 the sorption of MG onto DCB followed both Freundlich and Langmuir models. Freundlich parameter, n value 163 at equilibrium was 0.51, indicating a chemosorption adsorption of MG onto DCB. The maximum adsorption 164 (monolayer) capacity of DCB for MG is 55. 3 NaoH) and finally with acetone. t The optimal pH was found 165 in the range of 5 to 9. The adsorption process reached the equilibrium after 90 and 105 minute at initial 166 concentration of malachite green solution 4.9 and 7 mg/l, respectively. The process of removal follows pseudo 167 second order kinetics. Equilibrium data fitted well in Langmuir isotherm model. The maximum adsorption 168

capacity (monolayer coverage) was 303.03 (mg/g) at temperature 30 0 C. The intra particle studies suggested the intra particle diffusion as rate controlling process. The negative values of  $\hat{I}$ ?"G 0 and  $\hat{I}$ ?"H 0 obtained from thermo dynamical analysis of the experimental data indicated the spontaneous and exothermic nature of the adsorption.

[23] T. Santhi et al. (??009) studied a sample of raw (PR-Raw) and activated carbon (PR-Carbon) from 173 prawn waste as an adsorbent for removal of malachite green (MG)) from aqueous solution at 28 0 C. It was 174 found that the amount of MG adsorbed increased as the sorbent particle size decreased. The percentage of 175 adsorption increased as the adsorbent concentration increased. The value of initial pH 7 for PR-Carbon and pH 176 8 for PR-Raw, were found optimal for removal of MG effectively. When the dye concentration was increased 177 the percentage of dye adsorbed decreases. The equilibrium adsorption capacity of MG onto the PR-Carbon is 178 significantly larger than those onto the PR-Raw. From the values of R 2 it was concluded that the equilibrium 179 data best fitted the Freundlich isotherm model. The biosorption processes followed the Pseudo -first order rate 180 kinetics. Intra particle diffusion modeling indicating that Surface adsorption and intra particle diffusion were 181 likely to take place simultaneously. ??24]. Le Phan Linh et al.(2012) studied the removal of malachite green 182 from aqueous solutions by adsorption onto rubber wood (Hevea Brasiliensis) sawdust in batch kinetic experiment. 183 The equilibrium study showed that adsorption process of malachite green reached equilibrium after 2.5 hours 184 185 and optimal pH was 5. As the temperature increases the adsorption amount increased. Equilibrium data fitted 186 well in Langmuir isotherm model (R 2 = 0.9902) in compare to Freundlich isotherm model (R 2 = 0.7776). The 187 maximum adsorption capacity (Q 0) (monolayer coverage) was found 27.4 (mg/g) at temperature 40 0 C.

[25] Shabudeen P.S. syed (2011) studied Kapok hull activated carbon (KHAC) as an adsorbent for removal 188 of malachite green from aqueous solution. Here they used sulfuric acid (1:1) to activate the adsorbent. The 189 adsorption was found independent of pH of the medium. It was also observed that the adsorption increased as 190 temperature increases from 300 to 318K and further the increase in temperature did not have any influence upon 191 adsorption. It was found that the data fitted well in Langmuir adsorption isotherm model as well as Freundlich 192 adsorption isotherm model. Further, it was revealed that R L values lay between 0 and 1 and the value of 193 Freundlich parameter n lay between 2 to 10 at various temperature and particle sizes showing the adsorption of 194 MG onto KHAC was fovourable. Adsorption followed the pseudo first order rate equation and the intra particle 195 diffusion model indicated that more than one mode of sorption mechanism was in operation. From the thermo 196 dynamic analysis of the experimental data the value of  $\hat{I}$ ?"G°,  $\hat{I}$ ?"H° and  $\hat{I}$ ?"S° were found which come out to 197 be negative, positive and positive indicating the process as spontaneous, endothermic and there was a decrease 198 in randomness at the surface of adsorbent after adsorption. It was observed that the  $\hat{I}?"H^\circ$  and  $\hat{I}?"S^\circ$  values 199 increased with decrease of particle size. The positive and increased Î?"S° values for smaller particle size indicated 200 that the KHAC showed greater affinity towards the dye. It was observed that, the value of energy of activation 201 for malachite green by KHAC was in between 5 to 20 kJ/mol confirms the activated adsorption. 202

Response surface method using Box-Behnken design of experiments was adopted and gives a mathematical 203 model for the adsorption of dye stuff. The results obtained by adapting Box-Behnken model in the study of 204 absorption of MG dyes on KHAC proves, absorption of dye depends only upon the particle size of absorbent and 205 it was not influenced by pH or temperature. The experimental values and the predicted values of Box-Behnken 206 design model were in close agreement with quadratic regression >98%. ??26]. Jagdish Singh et al. (2013) studied 207 the removal of malachite green from aqueous solutions by adsorption onto agricultural waste rice straw in batch 208 kinetic experiment. The optimal pH of the solution was found to be 8 and the equilibrium time was about 15 min. 209 The removal of MG was found increases as the initial concentration of the MG increases. The adsorption of MG 210 211 increased with increase in temperature. The adsorption was found to follow the Pseudo -first order rate kinetics. The equilibrium data best fitted the Freundlich isotherm model. The Freundlich parameters n and K F were 212 obtained as 9.3, 10.3, 11.3 and 87, 90 and 92 mg/gm respectively at different temperature 25,30 and 35 0 C. The 213 values of Î?"G 0, Î?"H 0 and Î?"S 0 obtained from thermo dynamical analysis of the experimental data were found 214 negative, positive and positive respectively. [27] S. Sivamani et al. (??009) investigated the removal of malachite 215 green from its aqueous solution by Pithophora sp., fresh water algae by means of a batch system. The optimum pH 216 was found to be 5 and equilibrium time as 10 min and the maximum dye removal rate was found to be 94.35%. It 217 may be that the zero point charge for the prepared adsorbent could be found at a pH of 5. [28]. Anna Jasi?ska et 218 al. (??013) studied the waste of rapeseed press cake (WRPC), obtained after the preparation of microbial culture 219 medium, in a batch system as an adsorbent of malachite green (MG). The highest sorption of MG was observed 220 after 180min in solution containing 50mg/L of MG and 2.5mg/L of WRPC at pH 6.5. The equilibrium data 221 best fitted with the Langmuir isotherm model. MG sorption followed the pseudo-second-order rate kinetics. MG 222 desorption efficiency (94.5%) from WRPC with the use of 0.1M NaoH solution was determined. [29]. Chowdhury 223 .S et al. (2013) investigated thermo chemically modified Wheat bran with citric acid as a potential adsorbent for 224 removal of Malachite Green (MG) from aqueous solutions in batch experiment. It was found that operational 225 parameters, such as solution pH, adsorbent dose, initial adsorbate concentration as well as temperature, greatly 226 influenced the adsorption efficiency of the adsorbent. The Langmuir isotherm model showed excellent fit to the 227 adsorption data of MG. The sorption processes followed the pseudosecond-order rate kinetics. Thermodynamic 228 study showed spontaneous and exothermic nature of the sorption processes ??30]. Jia Tan et al (2012) used 229 Waste newspaper fiber (WNF), separated and deinked it(denoted by DWNF) for use as an absorbent for removal 230 of Malachite Green (MG) from aqueous solutions in batch kinetic experiment. The equilibrium study showed that 231

adsorption process of malachite green reaches equilibrium after one hour but 88% of removal took place within 232 10min. the optimal pH of the solution was found to be 7. Equilibrium data was found fitted well in Langmuir 233 adsorption isotherm model. The maximum adsorption capacity  $(Q \ 0)$  (monolayer coverage) was found 85.25 234 235 (mg/g) at temperature 40 0 C. Sorption kinetic analysis revealed that adsorption of MG onto DWNF followed the pseudo second order rate model. From thermodynamic analysis they obtained value of Î?"G° Gibbs free energy, 236 î?"H° change in enthalpy, and î?"S° change in entropy were found negative, negative and positive respectively. . 237 Sorption kinetic analysis revealed that adsorption of MG & MB binary mixture onto CAC and TKP followed the 238 pseudo first order rate kinetic expression. The rate constant value is higher in TKP than in CAC. This shows that 239 rate of adsorption is higher in TKP. This gives support for the efficiency of TKP. Adsorption data obeyed both 240 Freundlich and Langmuir adsorption isotherms. Intra particle diffusion plots for the removal of dyes from their 241 binary mixture by adsorption revealed the intraparticle diffusion was found to be rate determining step. ??32]. 242 K. RajasekharJia (2014)) used corn cob as an absorbent for removal of Malachite Green (MG) from aqueous 243 solutions in batch kinetic experiment. The contact time needed for dye solution to reach equilibrium was found 244 100min. The result shows that there was no significant change in the present removal of dye over the entire pH 245 range. The study reveals that percentage of adsorption increases with increasing the adsorbent dosage. IT was 246 found that as the initial concentration of dyes increased the removal of % of dyes decreased. ??33] Sharma et al. 247 248 (??013) studied the adsorption potential of agricultural waste material sugarcane baggase to remove malachite 249 green dye from aqueous solution. The adsorbent was characterized by BET surface area measurement and FTIR 250 analysis. It was observed that more than 95% removal efficiency was obtained within 120 min at adsorbent dose of 1 g/L for initial dye concentration of 50 mg/L. It was observed that with increase in temperature, adsorption 251 capacity decreases indicating the adsorption is exothermic in nature. The high values of R 2 (~1) and good 252 agreement between two q e Values indicate that the adsorption system followed pseudo-second-order kinetic 253 model and hence the process is chemisorptions controlled. From thermodynamic analysis the value of  $\hat{I}$ ?"G o , 254 î?"H o , î?"S o were found which came out to be negative, negative, negative respectively at every temperature 255 under study. Adsorption mechanisms were investigated with intraparticle diffusion model, Furusawa and Smith 256 model and Boyd's model which shows that both film diffusion and intra particle diffusion were simultaneously 257 occurring during the adsorption. Langmuir isotherm model was fitted the best for the adsorption system with 258 an adsorption capacity of 190 mg/g of adsorbent. ??34]. Makeswari et al. (??013) studied and analysed the 259 Competitive adsorption of malachite green (MG) in single and binary system on microwave activated epicarp of 260 Ricinus communis (MRC) and microwave assisted zinc chloride activated epicarp of Ricinus communis (ZRC). It 261 had indicated that ZnCl2 was a suitable activating agent for the preparation of activated carbon from epicarp of 262 Ricinus communis by microwave radiation. SEM micrographs showed that the external surface of the chemically 263 activated car-bon was full of cavities compared with untreated Ricinus communis. The activated carbon prepared 264 could effectively used as adsorbent for the removal of basic dye from aqueous solutions. 5 was the optimum pH 265 value observed for the adsorption of MG onto MRC and ZRC. It was observed that the percentage of adsorption 266 increases with increase in adsorbent dose from 0.2 g to 1 g in MG with the concentration of dye solution of 100 267 mg/L. It was seen that the percentage removal decreased with the increase in initial concentration. Adsorption 268 Isotherm Studies for adsorption of MG onto MRC (S), MRC (B), ZRC (S), ZRC (B) revealed that the obtained 269 data was best fitted in Langmuir isotherm model compare to Freundlich isotherm model, Dubinin-Radushkevich 270 model, Temkin model. The maximum adsorption capacity (monolayer coverage) for adsorption of MG onto MRC 271 (S), MRC (B), ZRC (S), ZRC (B) were found 12.6500, 11.7647, 24.3900 and 20.4081 mg/gm respectively. The 272 higher R 2 values confirm that the sorption process of dyes onto MRC and ZRC follow a pseudo-second-order 273 kinetic model. From Intra Particle Diffusion Model it was observed that the plot between q e Vs t 1/2 did not 274 pass through the origin for both the single and binary system which indicated that surface adsorption and intra-275 particle diffusion were concurrently operating during the MRC and ZRC interactions. MG adsorption rate onto 276 MRC and ZRC was greater in single system (S) than in binary system (B) due to the competitive adsorption of 277 dye onto the active site of the activated carbon. Among MRC and ZRC, ZRC shows most adsorption ability than 278 MRC in single and binary system. ??35] Rajeshkannan et al. (2010) studied the Hydrilla verticillata, a cheap 279 and widely available biomass as a potential adsorbent to remove malachite green from the aqueous solutions. 280 The effects of operating parameters such as temperature, adsorbent dosage, contact time, adsorbent size, and 281 agitation speed on the sorption of Malachite green were analyzed using response surface methodology (RSM). 282 The proposed quadratic model for central composite design (CCD) fitted very well to the experimental data that 283 it could be used to navigate the design space according to ANOVA results. The optimum sorption conditions were 284 determined as temperature -43.5 o C, adsorbent dosage -0.26g, contact time -200min, adsorbent size -0.205mm 285 (65mesh), and agitation speed -230rpm. Freundlich isotherm fits the data better with R 2 0.991 than Langmuir 286 isotherm with R 2 0.963. Freundlich parameters are K f -3.17, n-1.299. The maximum adsorption capacity 287 was obtained from Langmuir isotherm (91.97 mg/g) at a solution pH ~8.0. From the kinetic and equilibrium 288 studies it was found that pseudo second order kinetics and Freundlich isotherm fits the data well respectively. 289 Intra particle diffusion models analysis revealed that both the external diffusion as well as intra particle diffusion 290 contributes to the actual sorption process. ??36]. ??ubbareddy Y et.al.(2012) investigated the potential use 291 of a low-cost Fuller's Earth (FE) for removal of Malachite Green oxalate (MG) dye from an aqueous solution. 292 The experimental equilibrium data were found best fitted with the Langmuir model. The maximum adsorption 293 capacity of FE was found to be 1.96 x10-4mol/g for MG at room temperature. The energy of adsorption was 294

25.828 kJ/mol indicating chemisorptions. Adsorption data of MG onto FE was fitted well by the pseudo second 295 order model. From thermodynamic analysis Î?"G° and Î?"H 0 were found negative and positive respectively 296 suggesting the process of removal of MG by FE is a spontaneous and endothermic nature. ??37]. Madrakian .T 297 et al. (2012) investigated the removal of crystal violet, CV, onto the Magnetite nanoparticles loaded tea waste 298 (MNLTW) obtained from treatment of tea waste as naturally occurring waste by FeCl3\_6H2O, FeCl2\_4H2O and 299 NH 3 solution. It was observed that the dye became adsorbed after 35 min and under the optimized conditions, 300 up to 98% of dyes can be removed from the solution onto the MNLTW surface and the optimal pH was found 301 10. The equilibrium data of adsorption of CV onto MNLTW best fitted with Langmuir model with R L lying 302 between 0 to 1(0 < RL < 1). Maximum adsorption capacities at 25 0 C for removal of CV were found to be 303 129.87 mg/g. It was observed that that adsorption data followed the pseudo second-order kinetic model. ??38]. 304 Karla Aparecida Guimarães Gusmãoa et al. (2012) have investigated the adsorption of crystal violet onto the 305 Succinylated sugarcane bagasse (SCB 2) prepared from sugarcane bagasse, an important agricultural waste, after 306 only one chemical modification under several operating conditions. Equilibrium adsorption times was found to 307 be 20 hours for CV and optimum pH for removal of CV was found to be equal to 8.0. The experimental data 308 fitted very well to the Langmuir model. Maximum adsorption capacities for removal of CV were found to be 309 1273.2 mg/g. The adsorption process was well described by pseudo-second-order model; however the intraparticle 310 311 diffusion model yielded three linear regions suggesting multiple sorption rates. [39] Prasad and Santhi (2012) 312 studied the adsorption of cationic dyes (crystal violet and rhodamine B) from aqueous solution onto Acacia 313 nilotica leaves as an eco-friendly adsorbent. The equilibrium adsorption was practically achieved in 120 minutes. The highest removal of dyes was obtained at pH 6. The removal efficiency increased with increase in agitation 314 time and initial dye concentration. The equilibrium data were best described by Langmuir isotherm model with 315 maximum monolayer capacity of 33 and 37 mg g-1 for crystal violet and rhodamine B respectively. The adsorption 316 kinetics can be successfully fitted to pseudo-second order kinetic model. The result of the intraparticle diffusion 317 model suggested that intraparticle diffusion might not be the only rate controlling step. Desorption study revealed 318 that the recovery of the dye from adsorption was possible. [40]. M. M. EI. Jamal et al. (2011) investigated the 319 adsorption of crystal violet onto the Chaetophora Elegans Alga under several operating conditions. Equilibrium 320 adsorption times was found to be 20 minutes for CV and optimum pH for removal of CV was found to be 321 greater than 8.0. The experimental data fitted very well to the combined Langmuir-Freundlich model. Maximum 322 adsorption capacities for removal of CV were found to be 158.7 mg/g at temperature 25 0 C. The adsorption 323 process was well described by pseudosecond-order model; besides the thermodynamic analysis revealed that the 324 present adsorption process is endothermic and spontaneous as  $\hat{I}$ ?"H 0 and  $\hat{I}$ ?"G° were found positive and negative 325 respectively. [41] Satish Patil et al. (??011) have investigated the naturally available materials viz. the Mangrove 326 plant (Sonneratia Apetala) leaf powder (MPLP), Mangrove plant (Sonneratia Apetala) fruit powder (MPFP), 327 Mango (Mangifera Indica) leaf powder (MLP), Tamarind (Tamarindus indica) fruit shell powder (TFSP), Teak 328 tree (Tectona Grandis) bark powder (TTBP), Almond tree (Terminialia cattapa) bark powder (ATBP) as an 329 adsorbent for removal of crystal violet from aqueous solution. The monolayer (maximum) adsorption capacities 330 (Q 0) of MPLP, MPFP, MLP, TFSP, TTBP, ATBP were found to be 200, 250, 200, 142.857, 200, 166.667 331 mg/g respectably. The pseudo second order model best fits the kinetics of adsorption. Intra particle diffusion 332 plot showed boundary layer effect and larger intercepts indicates greater contribution of surface sorption in rate 333 determining step. Adsorption was found to increase on increasing pH, increasing temperature and decreasing 334 particle size. Î?"G, Î?"H and Î?"S values showed favorable, spontaneous, endothermic physical adsorption with 335 increased disorder and randomness at the solid solution interface of CV with biosorbents. Adsorption capacities 336 of different adsorbents towards CV were found to be of the order of MPLP > MPFP > TTBP > MLP > ATBP 337 > TFSP. ??42]. Nagda, G. K et al (2008), Studied Raw tendu waste (TLR), sulfuric acid carbonized tendu 338 waste (TLR-CM) and tendu waste treated with dilute sulfuric acid (TLR-2N) as sorbent for uptake of crystal 339 violet from aqueous solutions. The experimental result showed that it followed the pseudosecond-order kinetics 340 and followed the Langmuir adsorption isotherm. The maximum adsorption capacities for crystal violet for TLR-341 2N, TLR and TLR-CM are 67.57, 42.92 and 22.47 mg/g respectively. Interestingly, milder acid treatment of 342 the tendu waste enhanced biosorption, whereas drastic acid carbonization of tendu waste resulted in reduced 343 adsorption of dye. [43] Verma and Mishra (2010) used rice husk carbon to adsorb dyes (crystal violet, direct 344 orange and magenta). It was found that the optimum time was 45 minutes and there was decrease in adsorption 345 capacity in the low pH region. Also, the removal of dyes increased with increase in temperature and there was 346 increase in removal of dyes with increasing adsorbent dose. The removal percentage decreased with increase in 347 initial concentration of dyes. [44]. Bharathi K. S. et al. (2012) used Citrullus lanatus (Watermelon) rind, an 348 agricultural solid waste as a bio-sorbent for the removal of crystal violet from aqueous solutions. The maximum 349 amount of CV adsorbed corresponding to the equilibrium time of 180 min was found to be 87% for a dose of 350 1.0 gm/lit of the adsorbent. The optimal pH value for the sorption of CV was found to be in the range of 351 8.0-12.0. The extent of adsorption of CV was found to increase with increase in temperature in the range of 352 30-500C, indicating the process to be endothermic in nature. The experimental equilibrium data best fitted to 353 the Freundlich isotherm model. The adsorption capacity was found to be 4.82 mg/g at 30 o C. The kinetics of 354 adsorption was found to follow pseudo-second-order kinetic model. From thermodynamic analysis the value of 355 Î?"G 0, Î?"H 0 and Î?"S 0 were found negative, positive and positive respectively. [45]. Sagnik Chakraborty 356 et al. ??2011) studied, equilibrium, kinetics and thermodynamics of Crystal Violet (CV) adsorption onto NaoH 357

modified rice husk (NMRH). The adsorption was favored at higher pHs and lower temperatures. Adsorption data were well described by the Freundlich model, although they could be modeled by the Langmuir model as well. The adsorption process followed the pseudo-second order kinetic model. It was found that intraparticle diffusion was not the sole rate controlling step. The activation energy (Ea) of the system was calculated as 50.51 kJ mol?1. Thermodynamic parameters suggest that the adsorption is a typical chemical process, spontaneous, and exothermic in nature. (Sagnik Chakraborty, Shamik Chowdhury, Papita Das Saha, Adsorption of Crystal Violet from aqueous solution onto NaoH-modified rice husk, Carbohydrate Polymers 86 (2011) 1533-1541).

Rice husk carbon was used to adsorb dyes (crystal violet, direct orange and magenta) by . It was found that the equilibrium time was 45 minutes and there was decrease in adsorption capacity in the low pH region. Also, the removal of dyes increased with increase in temperature and there was increase in removal of dyes with increasing adsorbent dose. The removal percentage decreased with increase in initial concentration of dyes. **??**46] III.

## <sup>369</sup> 5 Further Scope of Study

From the above review it can be observed that a number of biomaterials have been tested as an adsorbent for 370 the removal of CV and MG in batch mode experiment. These studies will be proved useful only when suitable 371 technology is being evolved for their actual use in removal of these dyes from industrial effluents containing CV 372 and MG. Though lots of works have been done to find out the effectiveness of biomaterials in removal of CV 373 and MG by means of adsorption but very little work has been done to understand the actual mechanism of 374 adsorption of these dyes onto the various biomaterials. This is one of the areas where researchers have to put 375 their attention. There is the need to develop mathematical model to predict the effect of various parameters like 376 the rotational speed, pH value of solution, initial concentration of dye, dose of adsorbent, temperature. Response 377 surface methodology (RSM) is an attempt in this direction but still much more is required to evolve mathematical 378 models which fully explain the various factors. During the last five years a number of biomaterials/agricultural 379 waste have been tested as an adsorbent for the removal of the crystal violet and malachite green dye from its 380 aqueous solution by researchers. These biosorbents shows excellent adsorption capacity for the CV and MG dyes 381 and can be used as an alternative of activated carbon.  $^{1}$ 



Figure 1:

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