Operational parameters affecting the removal and recycling of direct blue industrial dye from wastewater using bleached oil mill waste as alternative adsorbent material

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Abstract—In this work the ability of “bleached” oil mill solid waste to reduce the dyestuff content in industrial textile wastewater was studied. Bleaching treatment consists in a preliminary oil mill solid waste management with NaOH and NaClO₂ for obtaining cellulosic materials, mainly removing lignin from the waste surface. Thus, a novel biosorbent from agricultural residues, named bleached olive pomace (OP), was presented. Direct Blue 78 was studied as a model azoic dye. Experiments were planned to study the effect of different initial conditions on the adsorption processes: oil mill waste amount as grains and as a fine powder (OPₚ), solution temperature values, initial dye concentration, pH values and electrolytes influence. The results showed that the adsorption process using bleached oil mill waste determined an excellent degree of water color reduction, reaching the best work conditions when pH 2 and OPₚ were used. The presence of electrostatic interactions was also suggested. The adsorption appeared to be influenced by temperature values showing an endothermic character. Interestingly, to confirm the role of ionic interactions between dye and sorbent at pH 2, fashionable results were obtained. The adsorption process was verified also at pH 6 with 100% of dye removal in presence of both NaCl and Na₂SO₄ avoiding the aforementioned strong acid conditions. A very important aspect of this work is the recycle of both the dye and the adsorbent, with particular attention to the dye reuse for coloring cotton fabric.

Keywords—Adsorbent recycle, adsorption, bleaching process, Olive pomace, textile dyes.

1. INTRODUCTION

In this generation, environmental questions such as water contamination are becoming progressively important.[1] As reported recently by Ertugay et al.[2] more than 25% of the total world population suffers from health and hygienic problems related to pollutant inflowing water.[2] Indeed, with the human development and improvement of technologies, large amounts of wastes are discharged every day into water. The composition of these pollutants encompasses a variety of contaminants as heavy metals, dyes, and/or other undesirable chemical compounds. The problem get worst and becomes more serious when dyes from textile industries flow into the water.[1,3] The nature of dyestuffs is very large and they are usually classified in accordance with the dyeing processes: some interact by ionic bonds or form covalent bonds or interact by electrostatic forces, and others interact by hydrophobic forces. It is worth to mention that among these, azo dyes are the dominant class among commercial dyes and belong to the category of direct dyes, largely used to color cellulose fibers mainly interacting through hydrophobic forces.[4] Nonetheless, the release of azo dyes into environment is of great concern, due to both their highly visible color in water, and their toxicity, mutagenicity and carcinogenicity, it still continues to find new applications in high-technology areas.[5,6] Although the amount of discharged dyes in water is not exactly known, recently Pirkarami et al.[7] reported that more than 5000 tons of dyeing materials are drained into the environment every year, affecting the human life and the global ecosystem.[7] Indeed, dyes are stable to light and heat, they have a high organic content, and the most part of them have complex aromatic structures not biodegradable.[8] As a result, the removal of synthetic dyes from industrial effluents is a great actual challenge. In this field the Authors of this paper have years of experience presenting innovative methods and materials with great performances in.
The agricultural waste (olive pomace), a byproduct of oil extraction, is increasingly being considered for wastewater treatment. Indeed, this procedure is usually used as a simple method with several advantages: ease of operations, simplicity of design, high efficiency and low cost applications in water decoloration processes. Among adsorbent materials for wastewater treatment, the agricultural wastes are very interesting sorbent materials requiring little processing, but showing good adsorption capacity, selectivity, low cost, free availability and easy regeneration. However, as reported by Saygili et al.[8], the application of untreated agricultural or plant wastes as adsorbents can also bring several problems and then, with the aim of improving their performance, the wastes are modified with different reagents and methods.[8] Regarding to the adsorbent used in this paper, i.e. oil mill solid residue (olive pomace), a chemical treatment by means of acids, bases or H₂O₂ are generally proposed in literature.[13] As alternative, the thermal pretreatments is also suggested, with the most part of papers related to the achievement of active carbon from oil mill solid waste. In this context, as recently evidenced by ourselves,[9], a very deep survey of papers related to the olive pomace use in dye removal from water showed as the number of publications was not high, being the most part of them focused on the heavy metal removal or requiring strong pre-treatment conditions.[9] In the same paper, we presented the performance of the olive pomace in removing disperse dyes from wastewater, by previously adsorbent treatment in hot water at 100°C, showing also the recycle of both the adsorbent material and dyes.[9] In this article, we show the best conditions adoptable to eliminate another class of dyes, direct dyes, from water enabling also the possibility to recover and reuse the dye itself. For the first time, the olive pomace was treated with NaOH and NaClO₃/acetic acid (bleaching process) to gradually remove lignin and holocellulose, in turn restituting mainly cellulose characterized by a high affinity with azo dyes.[14] Indeed, this procedure is usually used to obtain cellulose material from natural plant fibers.[15] As for the adopted model dye, Direct Blue 78 (DB) received by Colorprint Fashion, a Spanish textile industry, was used to reduce the contamination of colored water used for dyeing processes. Overall, effluent treatments aiming to reduce the amount of dyes having a structure similar to DB are known in literature, as for example degradation methods through Fenton’s oxidation process.[2] The Direct Blue 78 photocatalytic degradation using TiO₂ nanoparticles immobilized on recycled wool-based nonwoven material, was presented by Markovic et al.[16] The DB 71 adsorption from aqueous solution onto pistachio hull waste as a low-cost adsorbent was also presented by Biglari et al.[17] Activated carbon and poly pyrrole polymer composite prepared from Thevetia Peruviana were presented as materials to adsorb DB 71. [18] Without doubt the list is not complete for describing the plethora of studies in literature about the azoic blue dye removal or its degradation in aqueous environment, however in the present paper we highlight the use of a handy material exhibiting very high capabilities in sequestering the direct dye within few minutes in appropriate conditions. Moreover the recycle of DB 78 by means of desorption was obtained, enabling the possibility to color again cotton fibers, with 3 an environmental friendly green cycle and an alternative use of oil mill waste. As a whole the environmental disorder was reduced.

II. MATERIALS AND METHODS

2.1 Chemicals. All the chemicals used were of analytical grade and samples were prepared using double distilled water. Direct Blue 78 (chemical formula: C₄₂H₃₂N₃Na₄O₁₃S₂, MW: 1055.1 g mol⁻¹) reported in Scheme 1, was received by Colorprint Fashion, S.L. and used without further purification. Dye stock solutions with a concentration of 1.0 × 10⁻⁴ M were prepared and dilutions were carried out with double distilled water in order to obtain different dye concentrations namely 5 × 10⁻⁵ M and 1.0 × 10⁻⁴ M. The pH of the various aqueous solutions was adjusted using concentrated HCl and NaOH solutions. NaOH and HCl were purchased from Sigma-Aldrich (Milan, Italy). The same commercial source was also adopted for the following chemicals: Acetic acid (99.9 %), NaClO₃, NaCl and Na₂SO₄.

![Scheme 1: Chemical structure of Direct Blue 78.](image)

2.2 Preparation of the biosorbent. The biosorbent material was the solid waste of oil mill named Olive Pomace (OP), obtained during the oil production. OP was obtained from a local oil mill settled in Bari, south of Italy. OP was treated exploiting a procedure generally used to bleach natural fibers for removing impurities, lignin and holocellulose.[14,15] Several experimental conditions were studied, however among these the following procedure was proposed in this paper:

a) Alkaline treatment: 26.00 g of OP were boiled, for 2 h, in 140 mL NaOH 3M with continuous magnetic stirring.

b) Bleaching: The OP was subsequently boiled, for 2h, in 70 mL of NaClO₃ 1.7% w/w and 70 mL of Acetic buffer, 0.5M.
The experiments were performed both using OP as obtained after the treatment and sieved (OPₚ) obtaining a fine powder.

2.3 Experimental procedures. The experiments were conducted in 10 mL glass beakers containing known concentrations of dye solutions. The effects of the biosorbent and dye dosages on dye removal from wastewater, were assessed changing the amount of OP/OPₚ from 0.10 to 1.00 g, at 5×10⁻⁵ M and 1×10⁻⁵ M of dye concentration. The effect of both temperature and salt concentration were also evaluated in the range 25 ± 70°C and 5×10⁻³ M ÷ 1M, respectively. The mixtures were stirred at 140 rpm for different contact times using a digitally controlled magnetic stirrer. The adsorption process was studied following the DB absorption spectrum evolution at 600 nm when the solution was in contact with the pomace. In accordance with papers reported in literature[9] the adsorption capacity qₑ (mgxg⁻¹) at time t of dye, was inferred by applying the following equation (1):

\[ qₑ = \frac{C₀ - Cₜ}{W} \]  
where V represents the adopted total volume of solution (herein 10 mL), W is the weight of the dry adsorbent material (g), C₀ and Cₜ represent the initial concentration and the concentration at time t of the dye (mgxL⁻¹).

2.4 Desorption studies. The DB desorption studies from pomace surface were carried out by using a basic solution at pH 12. The olive pomace was loaded with dye’s initial concentration (5×10⁻⁵ M in 10 mL at pH 2) for 15 minutes. The dye loaded pomace samples were separated from the initial dye solutions, then washed with distilled water for the removal of unadsorbed dye and placed in contact with 10 mL of a 0.01 M of NaOH solution. The efficiency of desorption was calculated by using the following equation: where mₐ is the amount of dye desorbed and mₑ is the amount of dye adsorbed.

\[ E \% = \frac{mₐ}{mₑ} \times 100 \]  
The same Equation was used to calculate also the efficiency of adsorbed dye. In that case mₐ is the amount of dye adsorbed and mₑ is the initial amount of dye.

2.5 Dyeing experiments. The dyeing experiments were performed dyeing cotton pieces (1cmx1cm) for 60 minutes at 95°C in presence of increasing amounts of sodium sulfate to promote the dye exhaustion, that is the process of dye transferring from the water to cotton fibers.

2.6 Visible and FTIR-ATR spectroscopic measurements. Visible absorption spectra were recorded using a Varian CARY 5 UV-Vis-NIR spectrophotometer (Varian Inc., now Agilent Technologies Inc., Santa Clara, CA, USA). FTIR-ATR spectra were recorded within the 600–4000 cm⁻¹ range using an Fourier Transform Infrared spectrometer 670-IR (Varian Inc., now Agilent Technologies Inc., Santa Clara, CA, USA), whose resolution was set to 4 cm⁻¹. 32 scans were summed for each acquisition.

2.7 Scanning Electron Microscopy (SEM). In the case of SEM analysis, an electron microscope FESEM-EDX Carl Zeiss Sigma 300 VP was used. The samples were fixed on aluminum stubs and then sputtered with graphite by the use of a Sutter Quorum Q150.

III. RESULTS AND DISCUSSIONS

Fig. 1 (panels A and B) reports the SEM images of the adsorbent material. The morphology investigation shows the presence of irregular domains that conferred to OP the important features to be a material having a high porous character. In detail, Fig. 1A and Fig. 1B show OP before and after the adopted bleaching process, respectively. Interestingly, as it can be seen directly from Fig. 1B, the surface of the biosorbent occurred not affected by the bleaching process, maintaining the porous surface already observed in Figure 1A, which can be better appreciated in Fig. 1C. Indeed, changing the magnification ratio, the presence of cavities and irregular islands are better evidenced in that figure enabling OP to host dye molecules from aqueous solutions.

![Fig. 1: SEM images of olive pomace before (A) and after the treatments with a scale bar of 20 μm (B) and 1 μm (C).](image-url)
OP was recently presented in literature by ourselves for the removal and recover of a disperse dye from wastewater, using OP simply washed with hot water in order to remove impurities.[9] Using the OP treated in the same way for removing direct azo dyes (data not shown), it is possible to obtain the dye removal, but not its recovery. This led us to change the treatment procedure of biosorbent, obtaining also a quick dye removal using suitable conditions of work. In detail, the OP was subjected to an alkaline treatment followed by a bleaching procedure. The changes induced by these new processes can be appreciated observing the FTIR-ATR spectra reported in Fig. 2A and Fig. 2B. Before the treatment (Fig. 2A), OP showed typical bands indicating the presence of lignin, cellulose and cellulose-like structures as main components.[9]

Along with bands at 2920 cm$^{-1}$ and 2840 cm$^{-1}$ evidencing the presence of lignin and carbohydrates, signals in the region 1520-1540 cm$^{-1}$ indicated the presence of esters in the lignin structure. The bending vibrations of aliphatic -CH were also observed at 1366–1320 cm$^{-1}$. Bands at 1540 cm$^{-1}$ and 1630 cm$^{-1}$ suggested the presence of amino and carboxyl groups, respectively.

Not surprisingly, an intense band was detected at 3313 cm$^{-1}$ and ascribable to the hydroxyl and amino group stretching. The broad bands at 1160–1000 cm$^{-1}$ represented the characteristic C-O-C and OH vibrations of polysaccharides and, among them, of cellulose. When the bleaching process was applied to OP, the variation on the material surface, ascribable to changes of natural polysaccharides, modified the corresponding FTIR-ATR spectrum (Fig. 2B). Indeed, after the treatment, significant variations were observed in the fingerprint region affecting mainly the signals of lignin. In particular, the bands at about 1630 cm$^{-1}$ and 1032 cm$^{-1}$ (Fig. 2A) changed their ratio with the latter signal that moved to 1025 cm$^{-1}$ (Fig. 2B). Along with these variations the signals detected at 3313 cm$^{-1}$ before the treatment (Fig. 2A) shifted at about 3342 cm$^{-1}$ indicating the main presence of O–H stretching modes of wood fibers (Fig. 2B).[19-23] Moreover, as indicated by Kondo, this signal can be ascribed to a peculiar intramolecular hydrogen bond in cellulose structure.[24] As suggested by Poletto et al.[25] the band at about 2890 cm$^{-1}$ (Fig. 2B) indicated the stretching vibration mode of methyl and methylene groups present in the spectra of all of the fiber components, but mainly in the spectrum of cellulose.[25] Bands detected at 1600 cm$^{-1}$ and below this region, when the treated sample is considered (Fig. 2B), were assigned to C–H and C–O deformation, i.e. bending or stretching vibrations of carbohydrates. Not surprisingly, the deformation or stretching vibrations, observed at 1163 and 1025 cm$^{-1}$, confirm the presence of C–O–C and C–O groups.[25] Overall results suggested that lignin was the main component removed during the treatment, with the remaining adsorbent material largely composed by cellulose and cellulose-like structures with traces of lignin.[9,14,15]

Fig. 2: Comparison between detailed views (wavenumbers range: 500–4000 cm$^{-1}$) of FTIR–ATR spectra of olive pomace before (A) and after the treatments (B).
The removal of lignin was also confirmed by observing Fig. 3A and Fig. 3B that show as the color of the adsorbent changed after the treatments: olive pomace grains altered their color from dark brown to yellow, when the bleaching process was performed (Fig. 3B). The removal of lignin could be considered as the main factor inducing these chromatic changes. As well as to SEM images (Fig. 1), these camera pictures offer a macroscopic view of the pomace aspect, confirming the irregular character of the material offering active sites to host dye molecules. The dye used in this paper, as an example of direct ionic azo dyes, was Direct Blue 78 (DB78, See Scheme 1). The dye was removed from water with an excellent performance by means of adsorption process on treated OP surface. However, particular conditions of work must be adopted and will presented in the next section of the manuscript. As it can be seen macroscopically, by observing the OP camera pictures in Fig. 3C, the OP grains in contact with DB78 solutions changed their color from yellow to blue, indicating the uptake of dye molecules from wastewater. The dye recovery was also obtained offering the possibility to open a green virtuous cycle in which both the adsorbent and the dye can be re-used.

3.1 Adsorption experiments.
3.1.1 Effect of pH. As a first step of this study, 1.00 g of OP as grains was placed in wastewater containing DB dye and the UV-Vis absorption spectra were collected each 30 minutes monitoring the DB78 absorption maximum at 600 nm (Fig. 4).
Before the measurements, the sample at pH 6 was centrifugated and the supernatant was subject to the spectroscopic analysis. The absence of results (Fig. 4a) induced us to change the solution pH. Interestingly, when the pH was decreased to pH 2, important results were obtained with the 50% of dye molecules removed from water in 90 minutes, if OP was used as grains (Fig. 4B). Interestingly, by using the same amount of pomace, but crushed in fine powder, additional results were observed: only 15 minutes were necessary to completely remove the dye from the solution (Fig. 4C). The efficiency of 100% was thus presented. For that reason, the study was focused on the pomace reduced in fine powder (OP) as biosorbent and the effect of solution pH values was evaluated in the range 2-12. The amount of pomace and dye were settled at 0.25 g and 5×10⁻⁵ M in 10 mL, respectively.

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Table 1: Experiments of adsorption/desorption of DB78 at concentration of 5×10⁻⁵ M (Volume: 10 mL), at a fixed amount of OP (0.25 g), on the contact time enough to obtain the reported uptake or the recovery of dye molecules from aqueous solutions at pH 12 (Volume: 10 mL).

In order to better evidence the effect of various pHs affecting the DB78 removal from water, the q values (using (1)) were calculated and compared in Fig. 5. Clearly, it is evident that the dye removal from wastewater was obtained only at pH 2, with slight and not important variations at the other pH values ranging from 4 to 12. To better explain this finding, it is noteworthy that DB78 is an ionic negatively charged dye that in aqueous solution carries a net negative charge due to the presence of sulfonate groups. The UV-Vis absorption spectra of DB78 reported in Fig. 4 suggested as the dye was almost insensitive to the acidity of the solution. The typical absorption band, located at about 550-600 nm, attributable to the characteristic chromophoric part of the dyes comprising the azo-groups (–N=N–) interacting with the adjacent aromatic moieties, was detected. The lack of spectral modifications in the wavelength position of the dye aqueous solutions at different pHs, suggested the π→π* nature of that band. While, the change in the signal intensity could be attributed to the protonation of the central secondary amino group (pKₐ< 4). The sulfonate groups, can be considered deprotonated in the adopted range of pH values, due to their pKₐ< 2.[26,27] These results were in excellent agreement with the considerations reported by Abbott et al.[5] about the role of pH values on the pH-dependent equilibrium forms of DB78.[5] Consequently, in our condition, as well described by Saygili et al.[8] in their studies related to the chemical modification of a cellulose-based material to improve its adsorption capacity for anionic dyes, at the increasing of the dye solution pH, the adsorptions of the dye decreased due to the electrostatic repulsion between the negatively charged pomace surface and DB78 charges.[8] Moreover, in presence of OH⁻ ions in excess, these efficiently competed with dye anions for the adsorption sites onto the sorbent surface. While, Safa et al.[29] described that, lowering the pH values, the concentration of H⁺ ions increased, positively charging the surface of the biomass. In that condition electrostatic interactions between positively charged biomass surface and negatively charged dye molecules could be taken in account.[29] The nature of electrostatic interactions was confirmed when OP was loaded with DB78 dye (at 0.25 g of pomace and 5×10⁻⁵ M of DB78 in 10 mL) was placed in contact with a basic solution containing NaOH (0.01 M), at pH 12. Preliminary the OP loaded with DB78 was carefully washed with water in order remove unadsorbed dye molecules and then was placed in a solution at pH 12. By using (2), the efficiency of the dye desorption was evaluated: the 70% of the adsorbed dye was recovered after 1 h (see Table 1, the first cycle of adsorption/desorption).

Fig. 5: Effect of solution pH values, ranging from 2 to 12, on the adsorption capacity q at (mg×g⁻¹) of DB78 removal from aqueous (5×10⁻⁵ M) solutions, at fixed OPP amount (0.25 g).
The obtained results suggested that the adsorbed dye cannot be desorbed by simple water, but it is necessary the use of a strong base, as NaOH, indicating also that the attachment of the dye onto the adsorbent occurred through electrostatic interaction.[8] The excellent performance of the presented material was highlighted by the possibility of re-use of both the adsorbent and dye, for others cycles of adsorption/desorption. Table 1 reports the obtained results after 3 consecutive cycles. From the data in Table 1, it is possible to observe as the DB78 adsorption maintained almost unaltered the performance obtaining the complete removal of the dye from water within 30 minutes. While, the recovery of dye was unfortunately not complete, although efficiencies of 80% and 90% were obtained in the first 60 minutes for the 2nd and 3rd cycle, respectively. Further, this paper acquire a great importance considering the real re-use of desorbed DB778 in dying cotton fibers, however this aspect will be discussed at the end of the paper.

3.1.2 Effect of salts. To better understand the mechanism of DB78 molecule adsorption on OPF, the process was further studied changing the ionic strength of the solutions. It is worth to mention that textile dyeing processes are generally performed using large amount of salts[9]; as a results this evaluation becomes important for industrial applications. Surprising results were obtained. Since an amount of 0.25 g of OPF resulted to be so efficient in removing the dye to use 15 min for a 100% removal, these experiments were carried out with the amount of OPF lowered at 0.10 g in order to appreciate the spectral variations. Indeed, if the same experimental conditions were adopted (this aspect is discussed later in the paper), decreasing the weight of adsorbent, the efficiency of dye uptake from water decreased, enabling to follow the absorption spectrum of DB78 at several contact time. Overall, adopting pH 2 as pH of work, 5×10⁻³ M as dye concentration in 10 mL and 0.10 g of pomace, Fig. 6A shows the obtained results.

Starting from the results related to NaCl, by adding an appropriate amount of salt, such as 5×10⁻² M, 7×10⁻³ M and 1×10⁻³ M, in the mixture of dye and pomace, the adsorption process was improved. On the other hand, increasing or decreasing the amount of salt around these values, slight variations were obtained (see also Fig. S1A to better appreciate variations). A similar trend was obtained also change the nature of salt, i.e. Na₂SO₄. As an example, Fig. S1A and Fig. S1B report the effect on the adsorption process when Na₂SO₄ was used at concentration of 1×10⁻¹ M and 1 M. From these results, along with the role of the electrolyte concentration on the adsorption process, the effect of salts having different nature was also evidenced. The process resulted to be more sensitive by using Na₂SO₄.

A high ionic strength was induced under these experimental conditions. Overall, several reasons can be considered to characterize the behavior observed in presence of salts: (i) the increase of ionic strength induced the compression of the diffuse double layer on the electrostatic attraction and consequently contributed to the adsorption process. Further, the osmotic pressure of solution increased with salt concentration and consequently a concentration polarization layer will be built up by the salt:[30] (ii) the addition of salt rendered the dye molecules more hydrophobic favoring the adsorption process. Indeed, in the presence of counter-ions, the charges associated to DB78 molecules were screened inducing the presence of novel interactions affecting the adsorption process.[4]

![Fig. 6: The effect of salts on the adsorption capacity q<sub>0</sub> (mg×g⁻¹) for DB removal from aqueous solutions (5 × 10⁻³ M DB78, 0.10 g OPF in 10 mL) at pH 2 evaluating the concentration of NaCl in the range 5×10⁻¹ M – 1 M (A); at pH 6 evaluating the concentration of NaCl (B) and Na₂SO₄(C) in the range 1×10⁻² M – 1 M.](http://dx.doi.org/10.22161/ijeab/2.4.15)
Accordingly, both the charges of the pomace and DB78 were reduced inducing specific dipole–dipole and hydrogen bonding interactions as well as non-specific induction and dispersion interactions; hydrophobic interactions may also be important. Miyamoto et al. suggested that direct ionic dyes interrupt hydrophobic stacking between cellulose polymer forming hydrogen bonds. Moreover, the planes of glucose rings interacted with the dye aromatic moieties, and the sulfonate groups of the dye molecules interacted with the cellulose hydroxyl groups. In addition, the CH groups of glucose rings and aromatic moieties of dyes (e.g., naphthalene and biphenyl moieties) interacted weakly. As reported by Porter, the cellulose substrate was considered as heterogeneous material with several accessible regions in which dye molecules can be hosted. Surprisingly, if on one hand by using OP in neutral medium the adsorption did not take place, on the other hand the use of electrolytes as NaCl and Na₂SO₄ favor the adsorption process also at pH 6. Fig. 6B and Fig. 6C report the obtained results. In these conditions a high amount of salt was necessary to remove the dye from water with a reverse effect about the nature of used salts. The efficiency of the adsorption process occurred improved by using NaCl. Indeed, by comparing 0.1 as salt concentration, excellent results were obtained quickly when the adsorption process was studied in presence of NaCl (Fig. 6B and Fig. 6C) than with Na₂SO₄. The differences observed between NaCl and Na₂SO₄ could be ascribed to different effects in screening the charges of DB78 dye and adsorbent. However by comparing Fig. 6A, Fig. 6B and Fig. 6C, the efficiency of the adsorption process appeared overall improved when the electrolytes were added in DB solution at pH 6. So, along with the differences obtained changing the salt and pH, these results suggested that the ionic strength alone cannot be considered to obtain a comprehensive description of the adsorption process. The presence of more important hydrophobic interactions should be considered under these experimental conditions. The thickness of the electric double layer surrounding the dye molecule decreased as the concentration of electrolytes increased allowing the dye molecules to move closer to each other favoring hydrophobic interactions.

Not surprisingly, when the release of dye was studied under these conditions, at pH 12, its recovery was not obtained suggesting that the nature of interactions was changed. The results discussed so far suggested as the best work condition was obtained at pH 2 with also the dye recover. Therefore, from now on, the attention was focused on the adsorption process performed at pH 2 evaluating the effect of the adsorbent dosage and dye concentration on the adsorption process.

3.1.3 Adsorbent dosage and dye concentration. The amount of OP was changed from 0.10 g to 1.00 g fixing the concentration of DB78 at 5×10⁻⁵ M in 10 mL. The q values arisen from these experiments were calculated using (1) and are reported in Figure 7. At first glance, observing the time necessary to reach the plateau region, point in which theoretically the dye could be completely adsorbed, increasing the amount of OP increased the removed percentage of dye from water. This behavior was imputed to the presence of more free active sites, able to host dye molecules. On the other hand, when an high amount of OP was used the active sites were not saturated. These results were confirmed when the amount of dye was decreased. Adopting 0.10 g as fixed amount of OP and 1×10⁻⁵ M as dye concentration, 15 minutes were enough to completely adsorb DB78 from water (Fig. S2), against 90 minutes necessary to remove it from a concentrated solution, under the same experimental condition. Lowering the concentrations, all dye molecules interacted with the binding sites of the biosorbent. On the other hand, the adsorbent is characterized by a limited number of binding sites, which become saturated at a certain concentration. At higher concentrations, the great part of dye molecules are left in the solution due to the saturation of binding sites. Lowering the concentrations, all dye molecules interacted with the binding sites of the biosorbent. On the other hand, the adsorbent is characterized by a limited number of binding sites, which become saturated at a certain concentration. At higher concentrations, the great part of dye molecules are left in the solution due to the saturation of binding sites, observing a decreased dye removal percentage.

3.1.4 Effect of Temperature. As reported by several papers, studying the adsorption processes, temperature values could be an important factor in affecting the removal of dyes from water. Indeed, in accordance with the thermodynamic nature of process, that can be endothermic or exothermic, changing the temperature can positively or negatively influence the kinetic of adsorption. For such purpose, the temperature was changed from 25°C to 70°C by using the lowest amount of pomace, i.e. 0.10 g, fixing the concentration of dye at 5×10⁻⁵ M.
10^3 M. The obtained results are reported in Figure S3 and are related to the q values obtained under these conditions. The temperature played an important key role during the adsorption process. Passing from 25°C to 70°C the contact time necessary to remove DB78 from water changed from 90 minutes to 15 minutes, obtained an improvement in the process efficiency similar to that observed using 0.25 g of pomace at room temperature.

Several factor could considered to justify this behavior: (i) increasing the temperature values, the mobility of dye molecules increased; (ii) or as well described by Akkaya et al.[8], increasing the temperature there is also an increase of the number of active sites available for adsorption on the material surface along with the desolvatation of the adsorbing species and the decrease in the thickness of the boundary layer surrounding the adsorbent, so that the mass transfer resistance of adsorbate in the boundary layer decreased; (iii) last but not least, the swelling effect of the internal structure of the pomace, enabling large dye molecules to penetrate into the structure in higher quantities, that could not be excluded.[9] Indeed, as documented in our recent paper related to disperse dye removal, this effect was very important. This observation is extremely important. Interestingly, during the dyeing process of textiles, hot water is used and the generated wastewater is discharged at considerable high temperatures. These results with the extraordinary effect of salts in dye solution, auxiliary agents used to dyeing the textiles, offer the concrete possibility to use bleached olive pomace for the removal and recovery of dye from wastewater. For these purpose, experiments were performed using the desorbed dye from olive pomace to color the fibers as suggested by Colorprint Fashion. The effect of salt is clear evident in the camera pictures reported in the Fig. 8. As expected, the adsorption of dyestuff increased increasing the sodium sulfate concentration and it could be ascribed to processes affecting the dyeing method, i.e. the neutralization of negative charge of cotton fibers by sodium ions during the dyeing.

![Camera pictures of dyeing experiments related to desorbed DB78 (5 × 10-5 M in 10 mL) on pieces of a cotton textiles with increasing concentrations of sodium sulphate (10, 15 and 20 g/L). The experiments were performed at 95°C for 60 minutes.](image)

Fig. 8: Camera pictures of dyeing experiments related to desorbed DB78 (5 × 10^{-5} M in 10 mL) on pieces of a cotton textiles with increasing concentrations of sodium sulphate (10, 15 and 20 g/L). The experiments were performed at 95°C for 60 minutes.

IV. CONCLUSIONS

In this paper, the operational parameters affecting the adsorption of a commercial direct dye (Direct Blue 78) from wastewater were evaluated, proposing also an alternative use of oil mill waste (named olive pomace, OP) and obtaining excellent results both in the removal and recycle of dye and adsorbent. The biosorbent was modified through previous treatments with NaOH and NaClO for removing impurities and lignin from the surface of the substrate. A cellulosic material was thus obtained. These findings were confirmed using FTIR-ATR spectroscopy, evidencing the main bands of cellulose and cellulose-like structures, arisen after the treatment. A bleached adsorbent was obtained. Experiments were performed both using OP as grains and in powder (OPp). The latter exhibited extraordinary performance removing in 15 minutes the DB78 dye from water using only 0.25 g of the biosorbent. However, such a behavior was obtained settling the pH of dye solution at 2 units. The presence of electrostatic interactions inducing the adsorption process was thus evidenced, since at pH>2 there was no dye adsorption. The use of electrolytes as NaCl and Na2SO4 further confirm these findings. More specifically the latter salt increased the percentage of dye removal if compared with experiments performed in the presence of NaCl. Overall the influence of ionic strength was taken in account, showing as the process was affected by appropriate amount of salts. These results were better emphasized when the recycle of the dye was studied. Indeed, adopting pH 12 as medium of work, DB78 molecules were desorbed from OPp enabling the recycle of both the dye and biosorbent. As an example three cycles of adsorption/desorption were performed showing as the material exhibited the same efficiencies in dye removal with the not complete desorption of dye molecules. A mean value of 80% was obtained for each cycle. Interestingly, when experiments were performed at pH 6 in presence of the listed salts, the adsorption occurred, suggesting a change in the type of adsorbate/adsorbent interactions, favoring hydrophobic interactions instead of electrostatic ones. In that condition the adsorption occurred with a greater efficiency if compared with results obtained at pH 2, however the desorption of dye was not obtained. Starting from these considerations, the attention was focused on experiments performed at pH 2 studying the effect of the biosorbent amount and dye concentration. Both the experiments

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involving the change of OP<sub>p</sub> weight and dye concentration evidenced the role of free active sites hosting DB78 molecules. Indeed, increasing the amount of OP<sub>p</sub> the efficiency in the adsorption process increased. Accordingly, the same results were obtained decreasing the amount of dye. The adsorption process was also influenced by temperature exhibiting an endothermic character. The increase of temperature values induced an increase of the process efficiency.

Thus, the comprehensive investigation about the parameters affecting the adsorption process involving DB78 and OP<sub>p</sub>, carefully studied in this paper, evidenced as without salt and in acid conditions, the recycle of dye is possible. Whereas, the presence of salt in neutral medium, although improves the efficiency of the dye removal process, prevents the dye recover. In conclusion, the reuse of dye adsorbed results possible at pH 2 using different amount of Na<sub>2</sub>SO<sub>4</sub>, auxiliary agent generally used during the industrial dying process.

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Operational parameters affecting the removal and recycling of direct blue industrial dye from wastewater proposing bleached oil mill waste as alternative adsorbent material.

Fig. S1: The effect of salts on the adsorption capacity $q_t$ (mg g$^{-1}$) for DB removal from aqueous solutions ($5 \times 10^{-5}$ M DB, 0.10 g OP$_P$ in 10 mL) at pH 2 comparing the efficiency between NaCl and Na$_2$SO$_4$ at $1 \times 10^{-2}$ M and 1 M.

Fig. S2: Effect of DB dye concentration ($5 \times 10^{-5}$ M and $1 \times 10^{-5}$ M, in 10 mL) on the adsorption capacity $q_t$ (mg g$^{-1}$) using 0.10 g of OP$_P$ at pH 2.
Fig.S3: Effect of temperature values ranging from 2°C to 70°C on the adsorption capacity $q_t$ (mg g$^{-1}$) of DB dye removal ($5 \times 10^{-5}$M) from aqueous solution at pH 2; 0.10 g of OP$_p$ were used.