Estrogenic and anti-estrogenic activity of 23 commercial textile dyes.

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Estrogenic and anti-estrogenic activity of 23 commercial textile dyes

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Abstract

The presence of dyes in wastewater effluent of textile industry is well documented. In contrast, the endocrine disrupting effects of these dyes and wastewater effluent have been poorly investigated. Herein, we studied twenty-three commercial dyes, usually used in the textile industry, and extracts of blue jean textile wastewater samples were evaluated for their agonistic and antagonistic estrogen activity. Total estrogenic and anti-estrogenic activities were measured using the Yeast Estrogen Screen bioassay (YES) that evaluates estrogen receptor binding-dependent transcriptional and translational activities. The estrogenic potencies of the dyes and wastewater samples were evaluated by dose-response curves and compared to the dose-response curve of 17β-estradiol (E2), the reference compound. The dose-dependent anti-estrogenic activities of the dyes and wastewater samples were normalized to the known antagonistic effect of 4-hydroxytamoxifen (4-OHT) on the induction of the lac Z reporter gene by E2. About half azo textile dyes have anti-estrogenic activity with the most active being Blue HFRL. Most azo dyes however have no or weak estrogenic activity. E2/dye or E2/waste water ER competitive binding assays show activity of Blue
HFRL, benzopurpurine 4B, Everzol Navy Blue FBN, direct red 89 BNL 200% and wastewater samples indicating a mechanism of action common to E2. Our results indicate that several textile dyes are potential endocrine disrupting agents. The presence of some of these dyes in textile industry wastewater may thus impact the aquatic ecosystem.

*Keywords:* textile dyes, estrogenic activity, anti-estrogenic activity, industrial textile effluent.

1. Introduction

Dyes are widely used in most industries such as those manufacturing papers, plastics, food, cosmetics, textiles or leathers. These dyes are useful to colour the final products. Dyes are classified depending on their colors, their chemical structures and/or their origin (natural or synthetic). Natural dyes most frequently originate from plants (such as riboflavin or β-carotene). Determination of the chemical structures of natural dyes and the accomplishment of their synthesis allow the gradual replacement of these natural dyes by their synthetic counterparts. Approximately 10,000 commercial dyes are used in the coloring industry. More than 10% of dyestuff used during the coloring processes does not bind to the fibers and therefore these excess dyes are released into the environment producing serious environmental pollution (Pearce et al., 2003; Rajeswari et al., 2011). The presence of these dyes in wastewater and subsequently in water resources, even at very low concentrations, is easy to observe visually as the result of textile industry activities. They may increase effluent toxicity and lead to environmental damage (Robinson et al., 2002). In addition, many synthetic dyes are poorly biodegradable. In some dyehouse effluents, dye concentration can reach up to 400 mg/l (O’Neill et al., 1999). A specific study has even demonstrated that these concentrations can exceed 600 mg/L in Nigeria (Yusuff and Sonibare, 2004). Their presence in water reduces light penetration and has a negative impact on photosynthesis. Moreover, the dyeing process itself generally also contributes to the water body contamination by chromium, zinc and copper which are all toxic to aquatic plants and fish below 1.0 mg/L (Eremektar et al., 2007; Sharma et al., 2007; Verma, 2008). The research on textile wastewater toxicity, which has been carried out so far, shows how the action of toxic dyes occurs at different levels of the food chain, from food supply (i.e. algae and plants) to consumers (i.e. crustaceans and fishes) (Manu et al., 2003; Robinson et al., 2002; Sharma et al., 2007; Soni et al., 2006; Tigini et al., 2011). Most research effort has been devoted towards elucidating or
improving the degradation mechanisms of textile dyes with the hope to reduce their toxicity levels (Phugare et al., 2011a,b). Environmental regulations in most countries (i.e. EU directive 91/271) have prioritized in wastewater dye decontamination in order to minimize environmental damage (Robinson et al., 2001). Although the Environmental Protection Agency (EPA) has emitted new practice guidelines for environmental management, the presence of pollution dyes remains a serious environmental issue specifically for small textile industries in various countries (e.g. China, India, Taiwan) where working conditions and low economic status does not allow an efficient wastewater treatment before disposal into water sources (Mathur et al., 2005; Gregory et al., 2007; You et al., 2009). Since the mid nineties, the links between dyes, environmental impact and cancer emergence have been the subject of considerable interest both from researchers and from the general public. However, the causal relationship between dyes and certain types of cancer is difficult to establish. Several studies have been conducted on the toxicity, mutagenicity and genotoxicity of textile dyes (Bakshi et al., 2003; Ben Mansour et al., 2007; Dogan et al., 2005; Durnev et al., 1995; Mathur et al., 2007; Schneider et al., 2004). In the case of azo dyes, the increase in bladder cancer incidence, observed among textile industry workers, has been linked to prolonged exposure to these dyes. A report of experts from the “Cosmetic Ingredient Review Committee” confirmed that some anthraquinone dyes, like the disperse Blue 7 dye, used in the cosmetic industry as hair colorant and in textile, induce genotoxicity in bacteria (Cosmetic ingredient, 2007). Since then, research on these dyes confirmed their carcinogenic effects for humans and animals (Tsuda et al., 2001). This toxic effect has been linked to their presence in the environment (Dogan et al., 2005; Chou et al., 2007; Tigini et al., 2011). From an environmental point of view, toxicity, genotoxicity and mutagenicity of industrial effluents have been demonstrated (Alves de Lima et al., 2007; Grinevicius et al., 2009; Tigini et al., 2011). However, these effects have generally not been linked to the presence of dyes. Chou and collaborators associated the dioxin-like activity of some dyeing wastewater with the presence of specific anthraquinone dyes (Chou et al., 2006, 2007). In the same studies, these authors demonstrated that disperse blue 56 can bind the aryl hydrocarbon receptor (AhR), which is involved in many physiological functions such as cell regulation and reproduction. Two Brazilian studies have identified that the mutagenic activity of the Cristais River, a drinking water source of São Paulo, is caused by the presence of three blue dyes; C.I. Disperse Blue 37 contributing to 55% of this effect (De Aragão et al., 2005).
Up to now, the estrogenic activity of dyes has seldom been studied. Rare studies mainly report the effects of food colors, such as tartazine (E102) and erythrosin B (E127). Both compounds affect chromosome structure and increase Estrogen Receptor (ER) site-specific DNA binding to Estrogen Response Element (ERE) in HTB 133 cells (Roychoudhury et al., 1989) and in the E-screen test (Datta et al., 2008). The present manuscript investigates both the estrogenic and anti-estrogen activities of twenty three pure commercial textile dyes using the Yeast Estrogen Screen (YES). This in vitro assay has been developed for the detection of endocrine disrupting compounds (EDCs). In order to identify estrogen and anti-estrogen activity of textile dyes, the YES assay has been used to probe two different modes of action. First, a reporter gene assay measures the impact of dye binding on ER on its ability to promote binding-dependent transcriptional and translational activity. Second, an ER competitive binding assay measures how a dye competes with 17β-estradiol (E2)-dependent ER activation. In addition, the endocrine activity (estrogenic and anti-estrogenic activity) has been assessed on a textile effluent coming from a blue jean manufacturing Tunisian industry in an attempt to correlate the endocrine activity observed for commercial compounds and that obtained for dyeing wastewater.

2. Materials and methods

2.1. Selected dyes

All the dyes selected (23) in this study are used in the textile industry (Table 1 and 2). Only, 17 on 23 dyes have a known or communicated structure. The structure of the 6 other dyes were confidential and not available in the chemical abstracts service. However, the safety and chemical information certificate of analysis usually indicates the nature of the dye (azoic or anthraquinone-type chemical structure). For each dye, a stock solution (10 g/L) was prepared by dissolving in distilled water, followed by filtration through Whatmann No. 5 filter paper. All these dyes appeared perfectly soluble in water at this concentration as assessed by the absence of precipitate. For each dye, the endocrine activity was assessed at four concentrations in the range of 1×10⁻³ g/L to 1 g/L. The natural fluorescence of the dyes and their interference on the fluorescence emitted by yeast was determined beforehand in order to avoid spurious signals generated by the YES test in our experimental conditions. The fluorescence of each dye has been measured in the absence of yeast cells and subtracted from
the data obtained for estrogenic tests. The quenching potential of the dyes was also assessed on the yeast fluorescence emission during the antiestrogenic test as follows. The fluorescence of yeast cell has been measured after 6 h of incubation of the reference compound E2 (from Sigma-Aldrich, St Quentin-Fallavier, France). Then, each dye was added to the medium at various concentrations and the fluorescence quenching was measured immediately. Compounds inducing more than 10% of fluorescence quenching were not investigated further for anti-estrogenic activity (noted by an asterisk in Table 2). The cytotoxicity of each dye was evaluated by measuring yeast growth at an O.D. of 600 nm.

2.2. Wastewater sampling and extraction

Wastewater samples (1 L) were collected from the textile effluent of an industrial blue jeans factory in Tunisia. Raw water samples were centrifuged (2000 g, 15 min) to eliminate solids in suspension. Supernatants were then extracted by solid-phase extraction as described by Pillon et al. (2005). Briefly, aqueous samples were concentrated on reverse-phase C18 (5 g, 20 mL) cartridges (Sigma-Aldrich, St Quentin-Fallavier, France) preconditioned with methanol. Compounds elution from the column was triggered using methanol followed by hexane. Eluates were dried at 37°C in a rotary evaporator and residues were taken up in 2 mL methanol (concentration factor: 500).

2.3. Yeast estrogen screen assay (YES)

2.3.1. Assay of estrogenic and anti-estrogenic activity

Both agonist and antagonist activities of chemical dyes were examined using the Yeast strain BY4741 (Euroscarf, Frankfurt, Germany). This recombinant yeast strain carries the β-galactosidase reporter gene under the control of the ERE and contains the human ER cloned into the constitutive yeast expression vector pAAH5 (Garcia-Reyero et al., 2001). The test measures β-galactosidase activity (fluorescence at 460 nm and excitation at 355 nm) with a fluorimeter (Fluoroskan Twinkle LB 970, BERTHOLD Technologies) after 6 h of exposure to the compounds to evaluate. Tests were performed in 96-well plates. To determine the estrogen agonist activity of dyes and effluent samples, E2 was used as a positive control and distilled water was used as a negative control. Four dye concentrations were tested in the range of
1×10^{-5} \text{g/L} \text{ to } 1 \text{ g/L}. \text{ For the effluent samples, a series of dilutions (1 to 1000 of concentrated}
extract) were tested \text{ for their estrogenic activity. To determine the estrogen antagonist activity}
of \text{ the dyes, 4-OHT was used as a positive control and E}_2 (1 \text{ nM}) was used as a negative
control. \text{ In the antagonist test, the ability of the dye to compete with E}_2 \text{ for binding to ER}
and/or inhibit the receptor functions was evaluated. The tested dyes or 4-OHT, the positive
control, were combined with 1 \text{nM E}_2 \text{ before the start of the assay. All experiments were}
performed in triplicate. For estrogen agonist activity, the half maximal effective concentration
(EC_{50}) \text{ was calculated based on the sigmoidal dose–effect curve of E}_2. \text{ For estrogen antagonist}
activity, the half-maximal effective concentration (AC_{50}) \text{ was calculated based on the}
sigmoidal dose–effect curve of 4-OHT.

2.3.2. Competitive binding assay

\text{ Competition between dyes and/or sample effluent and E}_2 \text{ was measured at various}
concentrations of E}_2. \text{ Textile dye concentration used is 0.01g/L and the textile dyeing effluent}
sample \text{ at a dilution factor 5 of the concentrated extract. Thus any decrease of \beta-galactosidase}
activity after 6 h of exposure indicates that the dye induces a decrease in E}_2 \text{ binding.}

2.4. UV/Visible spectrum deconvolution

\text{ The absorbency spectrum of a water sample can be decomposed into a few numbers of}
spectra (reference spectra). \text{ The shape of the UV spectrum can be considered as a linear}
combination of defined spectra (REF_1, ..., REF_p) related to potential compounds present in
studied water sample (Thomas et al., 1996). Sw = \Sigma a_i \times REF_i +/- r, \text{ where Sw is the final}
spectra, a_i and r are the coefficient of the i^{th} reference spectra and the admitted error,
respectively. The Secomam company (Alès, France) has developed the UVPro software based
on advanced UV spectral deconvolution (UV PRO, 2000) which allows creating dedicated
models and determination of reference spectra from a set of studied wastewater UV spectra.
The UVPro software has been applied to the textile effluent of an industrial blue jeans factory
of Tunisia. The reference spectra used for the deconvolution are obtained from the 23 selected
textiles dyes (Table 1). Using the deconvolution model, it is possible to assess the
contribution of studied dyes in the wastewater spectrum.
3. Results / Discussion

3.1. Estrogenicity / anti-estrogenicity studies of individual dyes

YE5 allows a fast determination of both endocrine agonist and antagonist actions. The relative estrogenicity of each dye has been estimated using YES by comparison to the estrogenicity of E2 as reference compound. Typical dose-response curves for E2 have been established (data not shown). In the present study, we have essentially analyzed azo dyes. These dyes are extensively used for dyeing cotton in textile industries. By using YES assay in the present work, we have analyzed the degree of interference of textile dyes with the endocrine system considering both the potential agonist and antagonist actions. A great variability between dyes is observed. The YES assay data illustrate a dose-dependent estrogen agonist activity from 1×10⁻⁵ g/L to 1 g/L for three dyes (Yellow Flavina CXL, Reactive dye red 3BS and Solvent yellow 56) (Figure 1). All these dyes have weak estrogenic effects since the maximum activity is obtained at 1 g/L. In addition, the estrogenic effect of the dyes did not saturate precluding the determination of the EC₅₀ value, contrary to the positive control E2 (EC₅₀ value at 1 ng/L) (Figure 1). Higher dye concentrations were not investigated, since water body dye concentration in the environment never exceeds 1 g/l. Table 2 also summarizes the fact that the twenty other dyes had no estrogenic effect by themselves. These results indicate that these dyes are not xenoestrogens that should highly impact the environment, as compared to E2 or other known xenoestrogens (e.g. bisphenol A, paraben...) (Routledge et al., 1998).

In addition, some textile dyes have an anti-estrogenic activity. The anti-estrogenic reference compound 4-OHT was used to demonstrate efficient anti-estrogenic activity when incubated in the presence of 1 nM E2. The AC₅₀ value of 4-OHT was 0.5 µM or 0.02 mg/L (Table 2). Similar experiments were conducted with the dyes. Only some dyes have been tested (Figure 2), the other ones, marked by an asterisk, could not be tested because they quenched the β-galactosidase fluorescence. At a concentration of 1 g/L, antagonistic activities of these dyes vary from 100% to 10% inhibition according to their colour (Table 2 and Figure 2A). Blue and red dyes show the highest inhibition potential. Everzol navy blue FBN, blue HFRL, Direct Red 89 BNL 200% and Benzopurpurine 4B (Red 4B) are the most potent inhibitors tested. Their inhibitory effects remain however inferior to that of 4-OHT. At 1 g/L, these dyes are the only ones that induce a complete inhibition of the reporter gene in yeast.
cells. Among these four dyes, Blue HFRL is the strongest antagonist; its anti-estrogenic potency being approximately 750-fold less than that of 4-OHT. A reduction of 50% of β-galactosidase activity is observed for concentrations as low as 15 mg/L. These concentration values are thus considerably lower than the total color level found in the environment (400 mg/L). The antagonist effect of three other dyes (Benzopurpurine 4B (Red 4B), Everzol Navy Blue FBN, and Direct Red 89 BNL) were, respectively, 3500-, 2500-, and 1250-fold less than that of 4-OHT. In contrast, all other dyes were either weak ER inhibitors or non-ER inhibitors. For instance, Everzol Blue ED, Red alpacide 3BL, Direct Blue 71, Blue ED 250, Direct Black VSF and Blue DERF were weak ER inhibitors. The anti-estrogenic activity for Blue DERF and Direct Blue 71 at 1 g/l, are 10% and 22%, respectively. Among these dyes, most are of the azo class with sulfonated aniline (Benzopurpurine 4B, Direct Red 89 BNL 200%) and one is an anthraquinone dye, Everzol Blue ED. The Direct Black VSF AZO-FREE, Blue ED 250, both being polyazo dyes, Everzo Yellow ED and Red ED (for which no structures can be disclosed) induce inhibition of β-galactosidase expression by E2 with \( \text{AC}_{50} > 1 \text{ g/L} \).

Competition between the textile dyes and E2 was also measured by varying the concentration of E2 and maintaining a constant dye concentration (0.01 g/L, a concentration that does not fully inhibit E2 effect, Figure 2A). Benzopurpurine 4B, Blue HFRL and Everzol Navy Blue FBN can compete with E2 for binding to ER (Figure 2B). These dyes induces a slight inhibition of E2 estrogenic activity when E2 is used at concentrations equal or lower than \( 1.10^{-8} \text{ M} \). No effect is observed for higher concentrations of E2. For example, this activity decreased from 200000 RLU without dye to 100000 RLU in presence of the Blue HFRL at \( 10^{-8} \text{ M E2} \). This study also shows that the Red Alpacide 3BL induces a 44% inhibition of estrogenic activity and the metal-complexed azo dye yellow 4G induces a 13% inhibition (Table 2). The competitive binding assay in the YES assay showed that these dyes can bind specifically to ER in order to induce these antagonist effects.

3.2. Estrogenicity / anti-estrogenicity studies of textile effluent

The textile effluent sample analyzed presents a weak estrogenic effect at the maximum concentration tested (5-fold dilution). At higher concentrations of the sample, the sample compounds reduce cell viability (data not show) precluding the use of the sample for the estrogenic assay. For the dilution range (5- to 10000-fold), results from the YES bioassay...
indicate that the industrial textile effluent presents a low (below 15%) but significant estrogenic activity (Figure 3A). Also, the effluent sample inhibits 60% of estrogenic activity (Figure 3B). In competition experiments, in which E2 concentration is varied and a single dilution dose of the effluent sample is used (5-fold dilution), the anti-estrogenic effect is observable until $10^{-5}$ M of E2 (Figure 3C). The competition effect of the sample is most significant at $10^{-8}$ M.

The composition of the effluent sample in dyes has not been fully determined. Spectral analyses indicate that most of its components are blue dyes. Indeed, UV/Visible deconvolution of the spectrum of a blue jeans manufacturer’s effluent of Tunisia indicates that the sample contains mainly Blue DERF, Direct Blue 71, Everzol navy Blue FBN, and Grey GGL dye according to the spectrum of each individual dye (Figure 4). The deconvolution of the sample spectrum did not predict the presence of other dyes. Interestingly, our studies have demonstrated that at least three of these dyes presented anti-estrogenic activity with the YES assay. Most blue and navy blue dyes are derived from the Reactive Black 5 azo dye, which is classified Xn (harmful). It is also suspected to be mutagenic and to be associated to bladder cancer development (You et al., 2009). Nevertheless, without a complete characterization of the nature of the dyes being released in the local environment of this factory, the anti-estrogenic effect of the effluent sample can’t be unequivocally be attributed to these blue dyes. Since this effluent is essentially blue, and that some blue dyes are among those presenting the highest anti-estrogenic effect, suspicion remains however high that this Tunisian factory releases harmful components in the environment. In addition, it is worth noting that the most active dyes have anti-estrogenic AC$_{50}$ values (between 15 mg/L and 70 mg/L) that are well below the dye concentration that can be detected in some water bodies (400 mg/L). If these same dyes are indeed present in the environment, they would present a real anti-estrogenic activity to local water consumers.

4. Conclusions

Our results on the endocrine effects of textile dyes come in complement of those already published on the carcinogenic and mutagenic effects of dyes. They point to the fact that some dyes, mainly blue and red dyes, may be endocrine disruptor compounds. These observations raise two issues. Concerning environmental pollution, some of these dyes may be present at concentrations high enough that they may indeed affect life quality by promoting
tumor generation. Textile sewage is therefore a potential health hazard that should require a better communication on the dyes used by the industry and its method of disposal. In that respect, it appears as particularly important to efficiently treat industrial effluent containing azo dyes before they get discharged into the environment. The second issue concerns the pharmacological effect of these textile dyes. Besides 4-OHT, few compounds have anti-estrogenic activity. It is therefore of interest to note that some of the dyes that we characterized have similar anti-estrogenic activity, suggesting that they may be lead compounds for the development of new 4-OHT like anti-estrogenic compounds.

References


Figure legends

**Figure 1**: Induction of estrogenic activities of some textile dyes. Average ± standard deviation (n=3).

**Figure 2**: Anti-estrogenic activity of textile (A) dyes as measured by the yeast estrogen screen (B). Competitive binding essay of 3 textile dyes (0.01 g/l) against various concentrations of E2. Average ± standard deviation (n=3).

**Figure 3**: Estrogenic (A) and anti-estrogenic (B) activity of textile effluent, (C) competitive binding assay of textile effluent (dilution factor 5-fold) against various concentrations of E2. Average ± standard deviation (n=3).

**Figure 4**: UV/Visible deconvolution of textile effluent spectrum.
Table 1: Textile dyes analysed in this study. *Dyes and informations obtained by Everlight Chemical SA.

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<th>Compound</th>
<th>CAS #</th>
<th>Family</th>
<th>Formula</th>
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*Dyes and informations obtained by Everlight Chemical SA. *Reactive azo dye (copper complex, 70% pure for Blue HFRL) kindly provided by a textile industry unfortunately there is no available information on its structure (Baêta et al. 2011)

Table 2: Relative estrogen agonistic and antagonistic activities of industrial dyes. Estrogenic activity of textile dyes (1g/l). The anti-estrogenic activity of textile dyes has been determinate at 1g/l and expressed in 4-OHT%. The half maximal effective concentration (AC50) was calculated based on the sigmoidal dose–effect curve of 4-OHT.

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<th>Test Estrogen Screen antagonist activity (4-OHT %)</th>
<th>Test Estrogen Screen antagonist AC 50 (g/l)</th>
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<td>-</td>
<td>100</td>
<td>2*10^{-5}</td>
</tr>
<tr>
<td>Dye Name</td>
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<td>Value 2</td>
<td>Value 3</td>
</tr>
<tr>
<td>-----------------------------------------------</td>
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<td>---------</td>
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<tr>
<td>Blue DERF</td>
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<td>10</td>
<td>&gt;1</td>
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<tr>
<td>*Everzol Yellow ED</td>
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<tr>
<td>Reactive dyes red 3BS (Red 195)</td>
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<tr>
<td>Everzol Blue ED</td>
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<tr>
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<tr>
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<tr>
<td>Blue ED 250</td>
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