DEGRADATION OF STANDARD DYSES AND TEXTILE WASTEWATER AS A POLLUTANT MODEL USING GAMMA RADIATION

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ABSTRACT

DEGRADATION OF STANDARD DYSES AND TEXTILE WASTEWATER AS A POLLUTANT MODEL USING GAMMA RADIATION. A degradation and a decoloration of textile wastewater using ionizing radiation are already applied which some factors influence i.e. original wastewater, pH, pollutant concentration, solubility and variation of pollutant. A dyes standard had been treated using ionizing radiation and the mechanism degradation will be discussed. The percentage of the dye degradation about 90-99% at doses of 2-10 kGy. The degradation of textile wastewater using gamma radiation is was carried out at doses of 25 kGy. The combination of irradiation and the addition of coagulant caused the decoloration and the degradation of the textile wastewater.

Keywords: Dispers dyes, Basic dyes, Reactive dyes, Textile wastewater, Ionizing radiation

INTRODUCTION

The growth of the world population, the development of various industries, and the use of fertilizers and pesticides in modern agriculture have overloaded not only the water resources but also the atmosphere and the soil with pollutants [1]. In the last few decades the handling of wastewater appeared to be one of the most important. Textile industry which is one of the largest water consumer in the world, produces the wastewater comprising of various recalcitrant agents such as dye, sizing agents and dying aid. Therefore it has to be really concerned in releasing these types of wastewater to the environment. In the disposal of textile wastewater, color is of very important due to the aesthetic deterioration as well as the obstruction of penetration of dissolved oxygen and sun light into natural water bodies [2]. The sunlight penetration and reduction of dissolved of oxygen into natural water sources affects to aquatic life. The dye precursors and the degradation products of the dye are proven to be carcinogenic and mutagenic in nature.

In general, different types of dyes are used for colorization of textile in the dying process. The dyes (soluble or insoluble) that used in textile industry are synthetic and have a complex chemical structure and persist in nature. Therefore, they can not be degraded easily by ordinary treatments. Conventional method for eliminating dyes from the waste water include floculation with lime, activated charcoal adsorption and bio treatment. Lime treatment and charcoal adsorption generate solid wastes which require
costly disposal methods. Biotreatment processes rely on indigenous soil microorganisms to degrade dye compounds. Since the synthetic dyes are usually resistant to aerobic biodegradation, this process is likely to be inefficient [2-3]. It was reported that ionizing radiation has the ability to reduce biological oxygen demand (BOD) and chemical oxygen demand (COD) to convert refractory compounds to a more oxidizable by biological or chemical agents, to destruct bacteria and to make the improvement in handling disposal of solids [4].

The objective of the study is to degrade the standard (anthraquinone and azo) dyes and textile wastewater as a model of pollutant using gamma radiation. Variation parameters i.e. the change of the spectrum, total organic carbon (TOC), the addition of coagulant, absorbed dose, variation of pH and radiation, and combination of radiation with the addition of coagulant will be discussed in this paper.

Previous researchers have reported that radiation induced decolouration and degradation of disperse, acid and reactive dyes, so that radiation technology could be an alternative method to solve the textile industries problem [6-12]. The radiation technique helps to solve environmental problems very efficiently, especially in the degradation of water pollutant.

Ionizing radiation can be used for the treatment of dye standards and textile wastewater effluents treatment. Radiation-induced decolorization and degradation of water soluble azo and anthraquinone dyes especially in the presence of oxygen during irradiation.

In this report some determining factors which can strongly influence the efficiency of the radiation-induced pollutant degradation are kind and energy of radiation, absorbed dose and dose rate, effect of pollutant concentration, influence of pH, temperature influence, effect of oxygen and ozone, and effect of the molecular structure.

The research was carried out to degrade and decolour the dyes standard and textile wastewater as a model of pollutant using gamma radiation. The addition of coagulant, radiation, variation of pH and radiation, and the combination of radiation with the addition of coagulant were used to degrade and decolour the dyes standard and textile wastewater will be discussed in this paper.

THEORY

Radiolysis of Water

Radiation interaction between gamma rays with water will produce some species as described in the equation 1.

\[
H_2O \quad \rightarrow \quad H^+, \cdot OH, e_{aq}, H_2O_2, H_2 \quad \text{and} \quad H^+ \quad (1)
\]
Mechanisms interaction of ionizing radiation with water will produce species (H\(^+\), H\(^\cdot\), \(^{\cdot}\)OH, \(\text{e}_{aq}^{-}\), \(\text{H}_2\text{O}_2\), and \(\text{H}_2\)) in the spur [5]. This species are the primer product that produced from radiolysis of water. These species will react with pollutant in a solution and they will degrade them. Those species are hydrogen atom (H\(^+\)), hydroxyl radical (\(^{\cdot}\)OH), hydrated electron (\(\text{e}_{aq}^{-}\)), hydrogen peroxide (\(\text{H}_2\text{O}_2\)), \(\text{H}_2\) and hydrogen ion (H\(^+\)). From these products, the most reactive are hydroxyl radical, hydrated electron and hydrogen atom. In general, wastewaters contain dissolved oxygen, therefore H\(^+\) and \(\text{e}_{aq}^{-}\) can react with O\(_2\) to form an additional oxidizing species, perhydroxyl radical (\(\text{HO}_2\)\(^{\cdot}\)). The reactions are as followed:

\[
\begin{align*}
\text{H}^{\cdot} + \text{O}_2 & \rightarrow \text{HO}_2^{\cdot} \quad (2) \\
\text{e}_{aq}^{-} + \text{O}_2 & \rightarrow \text{O}_2^{-} \quad (3) \\
\text{O}_2^{-} + \text{H}^{\cdot} & \rightarrow \text{HO}_2^{\cdot} \quad (4) \\
\text{HO}_2^{\cdot} + \text{HO}_2^{\cdot} & \rightarrow \text{H}_2\text{O}_2 \quad (5)
\end{align*}
\]

**Oxidative Destruction of Organic Compounds in Aqueous Solutions**

The most hydroxyl radical produced from radiolysis of water solution subsequently react with organic pollutant (RH) in wastewater with the mechanisms as follows [4]:

\[
\begin{align*}
\text{RH} + \text{^{\cdot}OH} & \rightarrow \text{R}^{\cdot} + \text{H}_2\text{O} \quad (6) \\
\text{R}^{\cdot} + \text{O}_2 & \rightarrow \text{ROO}^{\cdot} \quad (7) \\
\text{ROO}^{\cdot} + \text{RH} & \rightarrow \text{ROOH} + \text{R}^{\cdot} \quad (8) \\
\text{ROOH} & \rightarrow \text{RO} + \text{OH} \quad (9) \\
\text{RO}^{\cdot} + \text{RH} & \rightarrow \text{ROH} + \text{R}^{\cdot} \quad (10) \\
\text{ROH} + 2 \text{^{\cdot}OH} & \rightarrow \text{R}^{\cdot\cdot}\text{C(OH)H} + \text{H}_2\text{O} \quad (11) \\
\text{R}^{\cdot\cdot}\text{C(OH)H} + \text{O}_2 & \rightarrow \text{R}^{\cdot\cdot\cdot} + \text{CO}_2 + \text{H}_2\text{O} \quad (12)
\end{align*}
\]

The wastewater contain some complex substrate which is unknown and variable, therefore evaluation of any oxidation process can be calculate by measuring the total amount of organics and total oxidation equivalents in the wastewater.

There are two important general measurements on the wastewater are chemical oxygen demand (COD) and total organic carbon (TOC). The COD measures essentially all of the organic and inorganic oxygen demand except
that of ammonia. The TOC measures the total amount of organic in the wastewater. Therefore the decrease in the TOC represents oxidation some of the organic material to $\text{CO}_2$ and indicates the amount of completely destructive oxidation occurring in the system.

The degradation product such as acetic acid, oxalic acid, succinic acid etc described in equation 11 and 12 can be measured by high performance liquid chromatography (HPLC) system.

Studies on degradation and decoloration of disperse dyes (insoluble dyes) in water using irradiation was carried out using disperse dyes standard as a model pollutant i.e. Terasil Black CMS (TB CMS), Terasil Red 4G (TR 4G), Terasil Golden Yellow (TGY) and Terasil Blue BGE (TB BGE). The dyes were insoluble in water but completely soluble in methanol and chloroform, indicating that they are non-ionic substances. The TB CMS, TR 4G, and TGY are commercial azo dyes and TB BGE is a commercial anthraquinone dye.

EXPERIMENTAL METHODS

Reagents

$\text{Al}_3(\text{SO}_4)_2$, $\text{NH}_4\text{Fe(SO}_4)_2$ as coagulant, standard azo dyes as commercial name i.e : Terasil Black CMS (TB CMS), Terasil Red 4G (TR 4G), Terasil Golden Yellow (TGY), and anthraquinone dye: Terasil Blue BGE (TB BGE). Standard basic, direct, and reactive dyes are Maxillon Blue (MB), Direct Red (DR) and Cibacron Violet (CV), respectively. Samples of dyes waste water were taken from the effluent of textile industry of PT.Unitex, Bogor Indonesia in four different times.

Experimental Design

The samples from PT Unitex Bogor were taken four times from the effluent at different time, namely A, B, C and D, and one sample from outlet, E. Irradiation was done using cobalt-60 gamma rays at room temperature with various dose 0-30 kGy. The dose rate of 5 kGy/h was determined by Fricke dosimeter. During irradiation oxygen was bubbled through a porous plate at the bottom of the reaction vessel. The absorbance and the pH of the solution were measured by HP 8453 spectrophotometer uv-vis and Methrohm 620 pH-meter, respectively. The degradation products were determined by a High Pressure Liquid Chromatography (HPLC LC 5A0) using uv detector and Shodex Pax column. The experiment was performed at the Center For Application of Isotopes and Radiation Technology, BATAN.
RESULTS AND DISCUSSION

Degradation of Anthraquinone and Azo Dyes Using Ionizing Radiation as a Pollutant Model

Disperse Dyes (Insoluble Dyes)

The TB CMS, TR 4G and TGY are commercial azo dyes and TB BGE is a commercial anthraquinone dye. Figures 1, 3, 5 show the changes of TB CMS, TR 4G and TGY after irradiation in oxygen-saturated condition. After irradiation, the intensity of the main absorption decreased gradually as the dose increased. These intensity decreases are obviously due to the destruction of conjugated system of the dyes molecules by the attack of OH radicals. After irradiation, when concentrated nitric acid was added to reduce the pH to around 1 and the solution was kept for two days, the upper layer of the irradiated sample became clear and precipitated was occurred.

Figure 1. Absorption spectra of irradiated TB CMS in O₂ bubbling solutions (0, 2, 4, 6, 8, 10 and 12 represent the dose in kGy).

Degree of precipitation was determined by measuring the spectra of the upper layer as shown in Figures 2, 4 and 6. The main absorption decreased markedly, especially at doses above 6, 12 and 8 kGy for TB CMS, TR 4G and TGY, respectively, while another peak appeared at 295 nm sharply for all doses (See also Figure 9 for TB BGE solutions).
Figure 2. Absorption spectra of irradiated TB CMS in O₂ bubbling solutions + HNO₃ (0, 2, 4, 6, 8, 10 and 12 represent the dose in kGy).

Figure 3. Absorption spectra of irradiated TR 4G in O₂ bubbling solutions (0, 2, 4, 6, 8, 10 and 12 represent the dose in kGy).
The changes of total organic carbon (TOC) of TB CMS, TR 4G and TGY solution irradiated in oxygen-saturated condition are shown in Figure 4. In oxygen saturated solution, the TOC content decreased slowly at doses below 6 kGy for TB CMS, then decreased rapidly at doses above 6 kGy. But for TR 4G the TOC content decreased steadily with dose especially in nitric acid added solution. These results correspond to the change of absorption spectra of TR 4G as illustrated in Figure 4. The TOC content of TGY did not change much either after irradiation or after the addition of nitric acid. This fact does not correspond to the results of its absorption spectra measurements (Figure 6) which may be caused by the increasing of insoluble components from colloid particles of the disperse dye due to irradiation [6-7].

Figure 4. Absorption spectra of irradiated TR 4G in O₂ bubbling solutions + HNO₃ (0.2, 4, 6, 8, 10 and 12 represent the dose in kGy).
Figure 5. Absorption spectra of irradiated TGY in O₂ bubbling solutions (0,2,4,6,8,10 and 12 represent the dose in kGy).

Figure 6. Absorption spectra of irradiated TGY in O₂ bubbling solutions + HNO₃ (0,2,4,6,8,10 and 12 represent the dose in kGy).
Figure 7. The Measurement of total organic carbon (TOC) and the effect of nitric acid on TOC changes in TB CMS, TR 4G and TGY.

Figure 8 shows the spectra change of irradiated TB BGE in oxygen bubbling solutions. After irradiation, the intensity of the main absorption decreased slightly with dose. However, after irradiation when nitric acid was added and settled for two days, precipitation occurred even at low dose as shown in Figure 9. These results suggest that chemical state of the surface of dye particles was changed by irradiation [6-7].

Figure 8. Absorption spectra of irradiated TB BGE in O₂ bubbling solutions (0,2,4,6,8, and 10 represent the dose in kGy).
In general, two important factors in measurements waste water are chemical oxygen demand (COD) and total organic carbon (TOC). The COD measures essentially all of the organic and inorganic oxygen demand except that of ammonia. On the other hand, the TOC measures the total amount of organic carbon in the wastewater. Therefore the decrease in the TOC represents oxidation of some of the organic material to CO₂ and thus indicates the amount of completely destructive oxidation occurring in the system. Figure 7 shows the change in TOC content in oxygen-saturated of TB BGE solutions due to irradiation and the addition of nitric acid after irradiation. The TOC content did not change much after either irradiation or the addition of nitric acid. The fact does not correspond to the results of its absorption spectra measurements (Figure. 5). This is probably due to increase of soluble components from colloidal particles of the disperse dye after irradiation.

The structures of the two disperse dyes are different as TB BGE is anthraquinone with an aromatic ring that is difficult to degrade using gamma radiation. On the other hand TB CMS has N=N bond that is weak and easy to degrade. The same results have been also reported by Suzuki et.al [15] where the decoloration degree of soluble azo dye at 1 kGy was almost 100%, while that of soluble anthraquinone dye at 1 kGy was only 50%. After the addition of nitric acid, the phenomena of both disperse dye solutions were slightly different. Sedimentation occurred at higher dose for azo dyes but for anthraquinone dyes occurred at lower dose. This phenomena is also influenced...
by the shape and the length of molecules. Anthraquinone dyes has thin shape, so neutralization and aggregation occurred easily at a low dose, while azo dye which is rather thick in shape, required a higher dose for degradation [6-8].

Water Soluble Dyes

Studies on degradation and decoloration of soluble dyes in water using irradiation was carried out using soluble dyes standard as a model pollutant i.e. basic, direct and reactive dyes. The basic dye is maxillon blue (MB), the direct dye is direct red (DR), and the reactive dye is cibacron violet (CV)[9-11,13].

Figure 11 shows the spectra change of irradiated maxillon blue in oxygen bubbling solutions. At dose of 4 kGy the percentage of degradation was achieved 100%, in this condition the color change from blue into a decolored solution. maxillon blue could be degraded using gamma radiation at low dose (4 kGy), because this dye is soluble in water. Most of the soluble dyes could be degraded easily at low dose [15].
Figure 11. Absorption spectra of irradiated MB in O$_2$ bubbling solutions (1,2,3, and 4 represent the dose in kGy).

Cibacron violet solution shows a clear absorption at 281 nm and 550 nm in the UV-VIS spectra as shown in Figure 12. The former absorption is considered to be the substituted aromatic rings and the latter can be assigned to conjugated system of the dye molecules. After irradiation at dose of 2 kGy the intensity of the 550 nm decreased sharply. The degradation could be achieved at dose of 10 kGy at this condition the percentage of degradation almost 100%.

Figure 12. Absorption spectra of irradiated CV in O$_2$ bubbling solutions (0,2,4,6,8, and 10 represent the dose in kGy).
Direct red shows a clear absorption at 528 nm and weak absorption at 288 and 372 nm (Figure 13). The 528 and 372 nm are assigned to be azo conjugated system and auxochromic effect of (-NH-CO-NH-) respectively. The 288 nm is substituted aromatic ring. At dose of 1 kGy the red color has change into decolored solution. The decreased in intensity are obviously due to the attack of OH radical into azo group. At dose of 3 kGy the percentage degradation is 99% in the UV area. This results indicated that the structure of the dye molecules already degraded. The degradation mechanisms of the dye can be explain as follow [15]:

\[
Z + \cdot OH \rightarrow ZOH \quad \text{(decolored product)} \quad (13)
\]
\[
Z + \cdot OH \rightarrow ZOH' \quad \text{(colored product)} \quad (14)
\]

Figure 13. Absorption spectra of irradiated Direct Red in O\textsubscript{2} bubbling solutions ( 1= 0 kGy; 2= 0.5 kGy; 3= 1 kGy; 4=1.5 kGy; 5= 2 kGy; 6 = 3 kGy and 7= 4 kGy)

Degradation of Textile Wastewater Using Ionizing Radiation

Sample of dyes wastewater were taken from the effluent of textile industry of PT Unitex Bogor at a certain time and labelled as A, B, C and D. One sample was taken from the outlet of the wastewater process (E). Preparation of these samples were carried out by the addition of coagulant, irradiation without the addition of coagulant, combination irradiation and coagulant and variation of pH before irradiation. From the results it is suggested that some determining factors can strongly influence the efficiency
of the radiation-induced pollutant degradation as illustrated in Table.1 [1].

Figure 14. shows the textile wastewater spectra that easily degraded at dose of 4 kGy and pH 3. From the results it is assumed that the sample containing some soluble dyes. SUZUKI et.al reported that soluble dyes could be degraded at low dose. This is indicated that the sample may containing some dyes that soluble in water.

Figure 14. Spectra of textile wastewater after irradiation at pH 3 in O₂ bubbling solutions.

Table 1. Important experimental factors determining the efficiency of radiation-induced pollutant degradation [1].

<table>
<thead>
<tr>
<th>No</th>
<th>Factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Kind and energy of radiation, absorbed dose and dose rate</td>
</tr>
<tr>
<td>2</td>
<td>Effect of pollutant concentration</td>
</tr>
<tr>
<td>3</td>
<td>Influence of pH</td>
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<tr>
<td>4</td>
<td>Temperature influence</td>
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<tr>
<td>5</td>
<td>Effect of oxygen and ozone</td>
</tr>
<tr>
<td>6</td>
<td>Synergistic effect of radiation, O₃ and TiO₂</td>
</tr>
<tr>
<td>7</td>
<td>Effect of the molecular structure</td>
</tr>
<tr>
<td>8</td>
<td>Application of pulse radiolysis technique as an aid for better understanding of the reaction mechanisms and hence for optimizing the degradation efficiency</td>
</tr>
</tbody>
</table>

Figure 15 shows the spectra of textile wastewater that has strong absorption at 291 nm and weak absorption at 530 nm. These waste could be degraded after the pH was adjusted into pH 3 with dose up to 30 kGy. The maximum degradation of this sample was obtained to be 85%. From these results it is assumed that this sample may contain some insoluble dyes i.e disperse dyes, certain surfactant and inorganic additive due to the insolubility of the dye in water [7].

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Figure 15. Spectra of textile wastewater after irradiation at pH 3 in O₂ bubbling solutions.

Experiment of sample D was carried out by combining radiation and addition of coagulant. Sample D has strong absorption at 282 nm in the UV region and slight shoulder at 345 and 454 nm. First, [Al₂(SO₄)₃] was added into sample D as a coagulant, furthermore the sample was settled for four days. Subsequently, the upper layer was taken out and irradiated at doses of 0-25 kGy. Figure 16(a) shows the spectrum of original sample D before the addition of coagulant. Figure 16(b) shows the spectrum of sample D after the addition of coagulant and irradiated at dose of 25 kGy. The intensity in the visible region decreased clearly after addition of coagulant but the intensity of the spectrum in the UV region hardly to decrease as shown in Figure 16(b) (curve 0 kGy). After irradiation at dose of 25 kGy the intensity in the UV region decreased markedly Figure 16(b) (curve 25 kGy). The intensity in the uv region is due to the aromatic rings from the dye molecules which is hard to degrade. These decreases in intensity are obviously due to the destruction of the aromatic rings of the dye molecules by attack of OH radicals that formed from radiolysis of water. The percentage degradation after treatment with combination of addition coagulant and radiation could be achieved 84.3%.

This results then compare with the sample that taken from outlet as shown in Figure 16(b) (curve outlet). By comparing the absorbance of sample outlet and sample D, it is seen that the intensity of the absorbance closed to each other. From the results above it is suggested that combination of radiation and coagulation could be used as an alternative method to treat textile wastewater to get better achievement.
Figure 16. Spectra of sample D, (a) before the addition of coagulant and non-irradiated, (b) after the addition coagulant and irradiated at dose of 25 kGy.

**Mechanism Azo Dyes**

Degradation of the dye molecules is induced by the reaction with oxidative species from water radiolysis[3,14]. The oxalic acid might be formed by the oxidation of benzene which is generated by the attack of oxidative species such as hydroxyl radicals on the dye molecules (Figures 17 and 18). The OH radicals attack the azo linkage-bearing carbon (C-4) of a hydroxy or amine-substituted ring. The resulting OH adduct breaks down to produce phenildiazene and phenoxy radical. Phenildiazene is extremely unstable, OH or molecule oxygen can readily oxidize it by one electron to yield a phenildiazene radical, (Figure 11). The latter intermediate is also unstable and cleaves homolytically to generate a phenyl radical and molecular nitrogen. The phenyl radical might abstract a hydrogen radical to produce benzene. Nickelsen proposed the oxidation of benzene into carboxylic acid, such as oxalic acid through the formation of muondialdehyde (Figure 18).
Figure 17. The formation of benzene from azo dye [3].

Figure 18. The formation of oxalic acid from benzene [14].
The degradation product of reactive dye cibacron golden yellow was analysed using high performance liquid chromatography (HPLC). The retention time of the sample was compared to that of organic standard. Figure 19 shows the chromatogram of irradiated of cibacron golden yellow at 10 kGy, it has the retention time at 4.292 and 7.413 minute. Chromatogram organic acid standards have retention time at 4.257 (oxalic acid), 7.383 minute (succinic acid), 8.223 minute (formic acid), 8.862 (acetic acid), and 10.598 (propionic acid) (Figure 20). By matching these two chromatogram it is assumed that degradation product of cibacron golden yellow are oxalic acid and succinic acid.

Figure 19. Chromatogram of irradiated cibacron golden yellow 10 kGy.
CONCLUSION

Decoloration and degradation of dye standards and textile wastewater can be done by irradiation, but the required dose varies with the type of the dye and the structure of the molecules. The addition of coagulants in textile wastewater treatment using gamma radiation, namely Al₂(SO₄)₃, NH₄Fe(SO₄)₂ induced degradation. Degradation of textile wastewater could be done but certain experimental factors i.e molecular structure, effect of oxygen, and pH can strongly influence the degradation. The degradation product that could be detected by HPLC were oxalic acid and succinic acid which less toxic.

REFERENCES


