APPLICATION OF SUB-CRITICAL WATER FOR RECOVERY OF TIN AND GLASS SUBSTRATES FROM LCD PANEL E-WASTE

Hiroyuki Yoshida 1,2, Shamsul Izhar 1,2*, Eiichiro Nishio 3, Yasuhiko Utsumi 3, Nobuaki Kakimori 3 and Salak Asghari Feridoun 2

1 Department of Chemical and Environmental Engineering, Faculty of Engineering, Universiti Putra Malaysia, 43400 UPM Serdang, Selangor, Malaysia
2 Ecology Research Center, Research Organization for the 21st Century, Osaka Prefecture University, 1-1 Gakuen-Cho, Naka-ku, Sakai, Osaka 599-8570, Japan
3 Environment Research and Development Center, Environmental Protection Group, Sharp Corporation, 1 Takumi-Cho, Sakai-ku, Sakai, Osaka 590-8522, Japan

ABSTRACT
Tin and transparent glass substrate were efficiently recovered from color filter (CF) and thin-film transistor (TFT) glasses in LCD panel wastes using sub-critical water (sub-CW) at various treatment temperatures and reaction time. Treatment for 5 min using sub-CW added with NaOH resulted in a 95% recovery of tin from CF and TFT glasses. The tin oxide did not liquify in the liquid phase instead stayed in the organic multi-layers, which exfoliated together from the CF glass and TFT glass. This is a huge advantage because the organic multi-layers were readily separated by sub-CW and tin was recovered by filtration with ease. Transparent and clean glass was also recovered from LCD panel waste. The amount of tin oxide recovered depended on the sub-CW reaction temperature, reaction time and NaOH concentration. Treatment of smaller CF glass improved the recovery of tin. With this advance, we have showed that sub-CW method is technically feasible for tin oxide recovery in LCD waste.

1. INTRODUCTION

Liquid crystal display (LCD) panel consists of layers of polymer, thin-film transistor (TFT), liquid crystal, and color filter (CF) substrates. These layers contain complex circuits, electrodes, multiple organic layers and glass substrates as shown in Figure 1. The electrode used in LCDs comprises of indium tin oxide (ITO), a mixture of 90 wt% indium (III) oxide (In₂O₃) and 10 wt% tin (IV) oxide (SnO₂) (Alfatanzani et al., 2003). Thus, many studies have been carried out to recover valuable material from scrap LCD screens. Most studies have concentrated on hydrometallurgical methods (Hsieh et al., 2009) such as acid dissolution or acid leaching (Li et al., 2009; Li et al., 2011; Gabriel et al., 2017a; Gabriel et al., 2017b), solvent extraction (Fortes et al., 2003; Honma and Muratani, 2005) and chlorination (Park et al., 2009) methods. However, those studies require the need for wastewater treatment after recovering the metals. Furthermore, clean glass cannot be recovered because only a part of the metal is extracted while multilayers of polymer films, TFT, CF and liquid crystals are still stuck on the glass surface.

In the present work, sub-critical water (sub-CW) was employed. Sub-critical water has been developed for solid waste resources recovery and is gaining interest due to its potential as solvent and catalyst for organic reaction. This practice is based on the use of water as medium, at temperatures between its boiling point (100°C) and critical point (374°C) and at pressures higher than or equal to the pressure of saturated vapor pressure. At sub-critical conditions, the dielectric constant of water decreases, thereby lowering its polarity as depicted in Figure 2. Secondly, the magnitude of ionic product of water increases three orders of magnitude around 250°C compared to room temperature. This will facilitate the dissociation of water making sub-CW advantageous for hydrolysis and decomposition of organic compounds including polymeric materials. Only a few papers have been published for metal recovery from multilayer films using sub-CW. Pure aluminum foil from waste composite laminates by decomposing plastic films by sub-CW hydrolysis reaction (Kulkarni et al., 2011). The plastics were also recovered as monomers.

Previously, a study revealed that 90% of indium was recovered from CF glass but only less than 7% indium from TFT glass (Yoshida et al., 2014). However, adding alkali improved the process where indium was totally removed and recovered from CF and TFT glass (Yoshida et al., 2014).
et al., 2015). However, the recoveries of other crucial elements, such as SnO$_2$ and high-quality LCD glass substrate was not considered. Thus, in the present work, we investigated tin and glass substrate recovery using sub-CW. In addition, the recovery of clean and transparent glass was carried out by changing the sample sizes before sub-CW treatment.

2. MATERIALS AND EXPERIMENTAL PROCEDURE

2.1 Material preparation and sub-critical water treatment

Preparation of raw material and sub-CW treatment were referred to Yoshida et al (2015). The waste LCD panel was a 40-inch TFT type (Sharp Corp.), composed of a polarizer, liquid crystals and arrangement of TFT and CF glasses. The polarizing films on the outer sides of the LCD were stripped off and separated into CF glass and TFT array by cutting the panel edge using a glass cutter. Liquid crystals on both glass surfaces were washed away by acetone. Each glass was crushed into 5-10 mm size, small enough to fit the reactor. The reactor as shown in Figure 3 was a stainless-steel tube with end-caps (Swagelok) were fitted to both ends of the tube (16 mm internal diameter, 150 mm length). Pure water or NaOH aqueous solution with volume approximately 20 mL together with 6 g of either CF glass or TFT glass was filled into the reactor. The air in the reactor was replaced with argon gas before the reactor was sealed and weighed.

For the sub-CW treatment, the reactor was immersed quickly in a preheated molten salt bath (Tomasu Kagaku, Celsius 600) at a desired temperature. After the desired time was reached, the tube was taken out and immediately cooled at room temperature. The pressure inside the reactor was the saturated vapor pressure of water at the subsequent reaction temperature. The reaction time was defined as the time from immersing the reactor into the salt bath until before immediate cooling. The sample was then transferred from the tube to a beaker. The solid residues (substrate glass) were separated from the aqueous phase by means of vacuum filtration through a 1 µm pore-sized membrane filter (Advantec, cellulose acetate). The filters were dried at room temperature and weighed.

2.2 Measurement and analysis

After sub-CW treatment, concentration of metal in each medium was determined. Tin was extracted using 7% hydrochloric acid solution from the remaining glass and membrane filter according to equation (1).

$$\text{SnO}_2 + 2\text{HCl} \rightarrow \text{SnCl}_4 + \text{H}_2\text{O}$$

(1)

![FIGURE 1: Detail cross section layout of a single pixel in a common liquid crystal display panel.](image1)

![FIGURE 2: Dielectric constant and ionic products of water at sub-critical and supercritical state (Wolfgang and Kretzschmar, 2008).](image2)
The tin concentrations, defined as mass of tin for every kg of LCD glass [mg/kg-LCD] in the solution extracted from the remaining glass substrate ($W_g$), filter ($W_f$) and liquid-phase ($W_l$) were measured using a plasma atomic emission spectroscopy ICP (Shimadzu, ICPE-9000). The mass balance of fresh tin ($W_{init}$) before the reaction and after the reaction ($W_g$, $W_f$ and $W_l$) was verified. The recovery of tin ($\eta_{Sn}$) was defined by the amount of tin that was filtered ($W_f$) to the amount of tin in the fresh LCD ($W_{init}$):

$$\eta_{Sn} = \frac{W_f}{W_{init}} \times 100$$  \hspace{1cm} (2)

3. RESULTS AND DISCUSSION

3.1 The concentrations of tin on CF and TFT glasses before sub-CW reaction

To evaluate the amount of tin recovery, the total amount of tin on each fresh CF and TFT glasses were determined by extracting tin using 7% hydrochloric acid aqueous solution. Tin quantity in fresh CF glass and TFT glass ($W_{init}$) were 36 and 28 mg/kg-LCD, respectively. The tin in TFT glass is 80% of the CF glass due to the sensitivity of the TFTs towards backlight illumination during application (Lee et al., 2008).

3.2 Sub-CW reaction in CF and TFT glasses (without NaOH)

TFT and CF glasses were treated by sub-CW without NaOH for 5 min reaction time at different reaction temperatures. The amounts of tin dissolved in the liquid-phase ($W_l$), in the organic multi-layers on the membrane filter ($W_f$) and in the residues on the glass surface ($W_g$) after the reaction was detected by the same method as the untreated fresh TFT and CF glasses as mentioned in section 3.1. Figure 4 shows the results from CF glass treated with sub-critical water for 5 min reaction time at various temperatures. Tin was not detected in the liquid-phase ($W_l$) when reaction temperature was lower than 360°C. $W_g$ reduced steadily with increasing reaction temperature beginning from 220°C to 360°C. At supercritical state (400°C), $W_g$ reduced to 4 mg/kg-LCD. Identically, $W_f$ showed the opposite trend to $W_g$ where $W_f$ increased with increasing treatment temperature. At supercritical state (400°C), $W_f$ was 31 mg/kg-LCD. Higher treatment temperature resulted in a more tin exfoliation from the LCD surface, although small amount of tin was detected in the liquid phase ($W_l$) when temperature was higher than 360°C. This is because the exfoliated multilayer organics, in which tin oxide was sandwiched, decomposed at very high temperature and tin oxide exposed to sub-CW.

Figure 5 demonstrates the results in TFT glass treated with sub-critical water for 5 min reaction time at various temperatures. Tin was not discovered at all in $W_f$. Treatment below 260°C and above 340°C showed no drop of $W_g$. 

![FIGURE 3: Layout of the reactor used for subcritical water treatment.](image1)

![FIGURE 4: Tin distribution in filter ($W_f$), liquid phase ($W_l$) and glass ($W_g$) after sub-CW treatment of CF glass for 5 min reaction time at various temperatures.](image2)

![FIGURE 5: Tin distribution in filter ($W_f$), liquid phase ($W_l$) and glass ($W_g$) after sub-CW treatment of TFT glass without NaOH for 5 min reaction time at various temperatures.](image3)
suggesting no tin was removed from TFT glass. At around 280°C, \( W_g \) was 26 mg/kg-LCD, demonstrating small amount of tin was removed. This could be attributed to hydrolysis, because at this temperature the ionic product is high as indicated in Figure 2, suggesting hydrolysis is active. At 400°C, \( W_g \) was 28 mg/kg-LCD indicating greater part of the tin remained on TFT glass. These results suggest that multilayer organics which sandwiched tin oxide were difficult to be exfoliated by sub-CW and even by super critical water. TFT layers are usually covered with a thin alignment layer (~80-100 nm thickness), with a function to insulate the optomer layer and to protect the passivation layers (Lee et al., 2008). The optomer film shown in Figure 1 is an organic mixture consisting of mostly diethylene glycol methyl ethyl ether and acrylic resin. The passivation layer wrapping the drain and source terminals are made of silicon nitride (Si\(_N_x\)). This implied that the optomer and passivation layers on the TFT glass were insufficiently decomposed by treatment with only sub-CW and supercritical water. However, it is worth noting that \( W_g \) was near zero in every circumstance. Consequently, the results in Figures 4 and 5 showed that tin did not dissolve in the liquid-phase but was appended with the organic multilayers and were removed together from the CF and TFT glasses.

### 3.3 Tin recovery from CF and TFT glasses with Sub-CW added with NaOH

To overcome the insufficient decomposition, NaOH was added in sub-CW. Figure 6 illustrates the effect of NaOH concentration in Sub-CW on tin recovery \( \eta_{Sn} \) from TFT and CF glasses. The reaction time was 5 min. For TFT glass, at 220°C, almost no tin was recovered when treatment was performed using less than 0.03 M NaOH. However, when treated with concentrations above 0.1 M, the amount of tin recovered was more than 80% for TFT glass. This suggests that concentration of NaOH in sub-CW is important for decomposing the organic multilayers in CF and TFT glasses.

In CF glass at 160°C, a similar result with TFT glass at 220°C was observed. But maximum recovery was more than 90% when NaOH concentration was higher than 0.2 M. Then a slightly higher reaction temperature (180°C) for CF glass was examined. The recovery of tin increased substantially, and 90% tin recovery were obtained in 0.1 M NaOH concentration.

Figure 7 demonstrates the effect of reaction temperature on the recovery of tin \( \eta_{Sn} \) by sub-CW treatment for 5 min with water-only and with 0.1M NaOH added. For CF glass (Figure 7A), \( \eta_{Sn} \) with water-only treatment was found to be inactive below 240°C. The \( \eta_{Sn} \) increased from none at 240°C to 80% at 360°C. \( \eta_{Sn} \) remained 80% until 400°C. These results exhibited that treatment temperature directly affects tin exfoliation from CF glass. However, when 0.1 M NaOH was present, the tin recovery was 45% even at 100°C. This suggests that hydrolysis process was probably active and had already occurred at water boiling temperature. The tin recovery increased to 95% at 160°C and remained about 90% at 360°C. With the presence of NaOH, Si\(_N_x\) from the passivation layer has dissolved into the liquid-phase as a result of the exfoliation of the organic multi-layers. Addition of NaOH probably caused corrosion of the substrate glass in NaOH environment causing the formation of sodium silicate. This showed the significant effect of the presence of NaOH to the tin recovery from CF glass.

In Figure 7B, for TFT glass treated by sub-CW without NaOH, \( \eta_{Sn} \) gradually increased from none at 240°C to 6% peak at 280°C. However, when 0.1 M NaOH existed in sub-CW, \( \eta_{Sn} \) increased drastically with reaction temperature. When the temperature was higher than 160°C, \( \eta_{Sn} \) increased to 95% at 220°C. The \( \eta \) was around 93% between 220 and 340°C. These results implied that the decomposition of the passivation and optomer layers on the TFT glass was enhanced significantly by the presence of NaOH in sub-CW. However, at 360°C the \( \eta_{Sn} \) slightly dropped to 70%. A previous study by Yoshida et al., has shown that high content of Si ion in \( W_I \) caused of the formation of sodium silicate at...
very high temperature. During ICP analysis of \( W \), due to the adjacent wavelength of tin and silica, the high intensity of silica interfered with tin causing the appearance of tin in \( W \). Thus, this resulted in the small reduction of \( \eta_{Sn} \).

Figure 8 illustrates the time course of the recovery of tin \( \eta_{Sn} \) from CF glass during sub-CW treatment with (a) 0.1 M NaOH at 160°C, (b) 0.05 M NaOH at 180°C, and 0.1 M NaOH at 360°C. The three curves were close to each other until 30 min. They showed the reactions were rapid because after only 5 min, \( \eta_{Sn} \) immediately became 80-90%. The reaction time of 5-15 min showed maximum recovery at 91-95%. After 30 min, curve (a) was close to curve (b) and \( \eta_{Sn} \) in both (a) and (b) slightly decreased with time. However, \( \eta_{Sn} \) in (c) decreased significantly with time. When reaction time was extended, the exfoliated multilayer organics that sandwiched \( SnO_2 \) were decomposed in small amount for (a) 160°C and (b) 180°C, but in large portion for (c) 360°C. The drop of \( \eta_{Sn} \) at 360°C is probably due to the difficulty to discriminate the intensity of silica and tin when analyzing the ICP results.

Figure 8 illustrates also the tin recovery from TFT glass by treatment with 0.1 M NaOH at 220°C. The \( \eta_{Sn} \) was 75% when treatment was performed for 2 min, but was maximum at 80% when treated for 5 min. The \( \eta_{Sn} \) did not go above 80% when treated above 5 min. Thus, when the concentration of NaOH was 0.1 M and reaction time was 5-15 min, maximum recovery of tin from CF and TFT glasses was attained. Most leaching using acid solution was able recover 89% to 99% of tin in the form of ITO as reviewed in the literature (Ueberschaar et al., 2017). The result in the present study using subcritical water was a 80 to 95% recovery from CF and TFT glasses, which is comparable to the hydrometallurgical methods. Furthermore, only a short time of 5 min is required thus making subcritical water method a promising technology and excellent alternative to the hydrometallurgical method.

3.4 The effect of sizes of CF glass on sub-CW reaction

Figure 9 illustrates the effect of glass size on the remaining color of CF glass after sub-CW treatment with 0.1 M NaOH for 5 min at 180°C. The CF glass was cut using a glass cutter into about 20 mm and crushed to about 5 mm length. After the sub-CW treatment, the larger size glass still has color remained from color filter pigment. While the smaller sized glass became completely clear of any pigment color. This showed that smaller sized glass which consisted of large amount of surface area effected the removal of CF pigments. As for the large sized glass, the color remained at the middle of the glass surface, indicating that as the surfaces of the glasses adhered each other, sub-CW contacted slightly with the surface but probably penetrated through the side of the glasses adhered.

Table 1 indicates the recovery of tin from CF glass cut into 5-10 mm, 20 mm and crushed to 5 mm length. Treating the glass with sub-CW method resulted in a maximum recovery of 80% to 95%. The result is comparable to the hydrometallurgical methods. Furthermore, only a short time of 5 min is required. This makes subcritical water method a promising technology and excellent alternative to the hydrometallurgical method.
TABLE 1: Effect of treated CF glass size on tin recoveries when they were treated with sub-CW added 0.1 M NaOH at 180°C for 5 min.

<table>
<thead>
<tr>
<th>Treated CF glass size</th>
<th>Tin Recovery [%]</th>
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<tbody>
<tr>
<td>5 mm</td>
<td>92</td>
</tr>
<tr>
<td>5-10 mm</td>
<td>92</td>
</tr>
<tr>
<td>20 mm</td>
<td>90</td>
</tr>
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2. The reaction temperature 160-180°C in CF glass and 220°C in TFT glass, NaOH concentration 0.05-0.1N, and reaction time 5-15 min were enough to get maximum recovery of tin (η = 91-95%). More tin was recovered, and the glass substrate became completely clean and transparent when sub-CW treatment was performed on smaller glass size (5 mm).

REFERENCES


H. Yoshida et al. / DETRITUS / Volume 04 - 2018 / pages 98-103