Effect of long-time heating for elements from flame retardants in acrylonitrile butadiene styrene and polycarbonate resin disks

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Abstract

The effect of long-time heating for elements from flame retardants (FRs) such as dechlorane plus (DP), tetrabromobisphenol A (TBBPA), triphenyl phosphate (TPhP) and antimony trioxide (Sb$_2$O$_3$) in both acrylonitrile-butadiene-styrene (ABS) and polycarbonate (PC) resin disks was examined in the present study. The heating temperature by drying oven was 80 $^\circ$C, which was expected as harder environment than that of usual dump site to discard the electric and electronic equipment, and the long-time heating was carried out for up to 722 days. As a result, the increases for both ether (C−O−C) and ketone (C=O) groups and the decrease of C=C double bond as well as methylene group (=CH$_2$) were found for ABS resin disk by fourier transform infrared (FTIR) spectrometry, which was expected to result in an increase of mass of the disk up to ca. 4 % by oxidation during the long-time heating. The changes in concentrations for bromine (Br), chlorine (Cl), phosphorous (P) and Sb, which were originated elements from FRs contained, were also evaluated by energy dispersive X-ray fluorescence (ED-XRF) spectrometry and the decreases in concentrations of ca. 5 % in ABS resin disk were observed which could be almost explained by increasing ca. 4 % mass of the disk. On the contrary, the mass and chemical form of PC resin disk as well as the concentrations of containing elements did not change during the long-time heating. From these results, it could be considered that the most of the elements from DP, TBBPA, TPhP and Sb$_2$O$_3$ contained as FRs in the ABS and PC resin disks examined in the present study were not lost during the long-time heating, even though the color, shape, mass and chemical form of ABS resin disk were changed.

Keywords
Flame retardant, Acrylonitrile-butadiene-styrene, Polycarbonate, Long-time heating, Mass, Chemical form, Loss of element, Oxidation
1. Introduction

Polymers are well known to be widespread used in electrical and electronic equipment and several hazardous substances contained in polymers are regulated in EU RoHS (restriction of the use of hazardous substances in electrical and electronic equipment) directive which became effective since July in 2006. The polymers such as acrylonitrile-butadiene-styrene (ABS) and polycarbonate (PC) are found as major ones in office machinery such as personal computers, printers, and data CDs, and many of them contain flame retardants (FRs) which are utilized in polymers since the 1960s as one of useful chemical additives for preventing combustion and/or delaying of the spread of fire after ignition of materials. Generally, FRs consist of chlorine (Cl), bromine (Br), phosphorus (P), and compounds such as aluminum hydroxide [Al(OH)₃] and antimony trioxide (Sb₂O₃). Particularly, organochlorine, organobromine and organophosphorus are major ones of organic FRs and are widely used in commercial electric and electronic equipment. Recently, Sb₂O₃ has also been widely used to enhance the effect of FRs containing halogens. On the other hand, commercial products containing FRs may cause environmental contaminations due to the release of FRs. Elevated concentrations of FRs such as polybromodiphenyl ethers (PBDEs) and 1,3-phenylene bis(diphenyl phosphate) have been found in leaves and are believed to affect biosphere from the air and the soils near dump sites for discarded electric and electronic equipment. In addition, commercial products containing FRs are not readily managed throughout the supply chain. Therefore, it is considered that properly elucidation of the changes in concentrations of FRs in these products is very important. The ageing of additives and polymers such as combinations of FRs and resins as well as antioxidants and rubbers was evaluated in earlier studies; however, it was difficult to catch slight changes in concentrations of additives since their homogeneities in polymers were not evaluated. From this point of view, we prepared plastic disks for both ABS and PC resins containing FRs and homogeneities of FRs were evaluated in our previous study. The FRs contained in these plastic disks were dechlorane plus (DP), tetrabromobisphenol A (TBBPA) and triphenyl phosphate (TPhP) which were selected as
organochlorine, organobromine and organophosphorus FRs, respectively. The Sb$_2$O$_3$ was also added to the ABS and PC resin disks as an inorganic FR. The concentrations of FRs prepared in these plastic disks were ca. 0.1 % in compliance with the RoHS directive on organobromine FRs and their homogeneities evaluated in compliance with ISO Guide 98-3:2008$^{15}$ were sufficiently small (maximum inhomogeneity 4.3%) to examine the change in concentrations of FRs by the ageing or weathering tests. In addition, the plastic disks could be used to evaluate the behavioral differences among FRs in polymers under environmental changes since three or four FRs were contained in the same disks and their concentrations were properly determined.$^{13,14}$ Consequently, the simulated weathering tests, which were independently set as outdoor or indoor exposure in compliance with the ISO 4892-2:2006,$^{16}$ were also performed with respect to the plastic disks.$^{13,14}$ As results, FRs in the plastic disks showed small changes in their concentrations due to the partly decomposition of their chemical forms under the simulated weathering tests which assumed as scrap-yard or office environments for ca. 6 months.$^{13,14,17-19}$ On the other hand, another question came out if the plastic disks were exposed to harder environment such as higher temperature. It is well known that X-ray fluorescence (XRF) spectrometry is expected for direct and speedy analysis of the hazardous substances in electrical and electronic equipment$^{20-23}$ and also described in the IEC 62321 as a screening method with respect to RoHS directive.$^{24}$ However, the loss of elements in plastics during XRF measurement was also reported for an organometallic compound of Hg in plastic certified reference material (CRM) due to its volatility.$^{25}$ In our previous study, we also observed the loss of Cr from the PP resin disk CRM (NMIJ CRM 8136-a) for long-time measurements by XRF spectrometry.$^{22}$ The temperature observed at the surface of the PP resin disk CRM was ca. 90 $^\circ$C after 1 hour measurement by wavelength dispersive XRF (WD-XRF) spectrometer and the loss of ca. 10 % Cr was observed during the long-time measurement for 120 hours from the side of the surface irradiated by X-ray.$^{22}$ Since the chemical forms of Cr for both inorganic and organometallic compounds were mixed in the PP resin disk CRM, the organo one was considered to be lost by the heating from the long-time X-ray irradiation. The effect of long-time
heating at 80 °C was also examined for both PVC and PP resin pellet CRMs to explore the source of the loss of elements in our previous study and the loss of organometallic compound of Cr was observed.

From these points of view, the long-time heating as harder environment compared to usual dump site to discard the electric and electronic equipment were examined with respect to the elements from FRs contained in plastic disks in the present study. The plastic disks were exposed at 80 °C in a drying oven for long-time up to 722 days and the concentrations of elements originated from FRs as well as the change of the masses and chemical forms of the plastic disks were evaluated in the present study.

2. Experimental

2-1. Samples

The plastic disks of ABS or PC resin containing known concentrations of FRs of DP, TBBPA, TPhP, and Sb$_2$O$_3$ were used in the present study. The concentrations of DP, TBBPA, TPhP, Cl, Br, P, and Sb in ABS and PC resin disks are listed in Table 1 and details of the preparation procedure are described in our previous study. The original ABS and PC resin disks (without long-time heating) used in the present study can be seen in photos 1 and 2, respectively. The diameter and the thickness of the original disk were 30 mm and 2 mm, respectively; and the original masses of ABS and PC resin disks were ca. 1.5 g and ca. 1.7 g, respectively. Though the detailed thermal properties were unknown for both plastic disks, the heat resisting temperatures for ABS and PC resins were generally known to be 70-100 °C and 120-130 °C, respectively.

2-2. Instruments

A drying oven with maximum temperature of 250 °C was used for long-time heating at 80 °C in the present study. The heating temperature of 80 °C is the drying temperature generally used for plastic CRMs for chemical analysis to remove any moisture before analysis. The heating
temperature was also close to the temperature of ca. 90 °C observed at the surface of the plastic disk after 1 hour measurement by WD-XRF spectrometer in our previous study. Approximately 1.5 g or 1.7 g of ABS and PC resin disks, respectively, were weighed accurately by electric valance (AX 205, Mettler-Toledo International Inc.) and heated in the drying oven at 80 °C during the heating periods of 0 (without heating), 10, 25, 152, 385 and 722 days. After the long-time heating, the mass of plastic disks and the change in concentrations of elements from FRs were evaluated. In case of mass measurement, the electric valance was used to weigh the mass of both plastic disks. The change in concentrations of elements from FRs were evaluated by energy dispersive X-ray fluorescence (ED-XRF) spectrometer (Shimadzu XRF Rayny EDX-720, Kyoto, Japan) and the operating condition was listed in Table 2. In the present study, fourier transform infrared (FTIR) spectrometry was also carried out to identify the chemical forms of ABS and PC disks, and the instrument used was Nicolet 6700 Continuum FTIR (Thermo Fisher Scientific). FTIR with attenuated total reflection (ATR) method was used. The internal reflection element, incidence angle and number of reflections for ATR-FTIR were diamond, 45 degrees and one time, respectively; and the spectral range measured was 650−4000 cm$^{-1}$ in the present study. All measurements were carried out after all plastic samples shown in photos 1 and 2 were ready, that is, after the long-time heating of 722 days.

3. Results and discussion

3-1. Plastic disks after long-time heating

The long-time heating was carried out with respect to ABS and PC resin disks with FRs and the obtained disks were shown in photos 1 and 2, respectively. The original colors of ABS resin disks (without heating) containing 3 FRs and 4 FRs (without and with Sb, respectively) were milky-white (see photos 1). On the other hands, the original PC resin disk containing 3 FRs (without Sb) was transparent even though the color changed to light-brown when Sb was added in the PC resin disk (see photos 2). As can be seen in photos 1, the shapes and colors of ABS resin disks changed
compared to the original one without heating (0 day) due to the long-time heating at 80 °C for more than 10 days even though FRs were contained. The colors observed between ABS resin disks containing 3 FRs and 4 FRs for 10 and 25 days heating were also found to be slightly different, even though others were seemed to be similar. Since the heat resisting temperature of ABS resin were generally known to be 70-100 °C, the change of shape and color were not surprised. The slightly different colors observed between ABS resin disks containing 3 FRs and 4 FRs for 10-25 days heating might be attributed to the effect of Sb$_2$O$_3$ as FR. On the other hand, no obvious changes were observed for both PC resin disks with and without long-time heating. Since the heat resisting temperature of PC resin was generally known to be 120-130 °C, it was considered that the shape and color of the PC resin disks did not change under the heating at 80 °C. From these observations, though the PC resin disks seemed to be strong and stable enough with respect to the long-time heating at 80 °C, it could be considered that the ABS resin disks did not have enough strength even though FRs were contained.

3-2. Change in concentrations of Br, Cl, P and Sb from FRs in ABS and PC disks

The elements originated from FRs in ABS and PC resin disks such as Br, Cl, P and Sb were measured by ED-XRF spectrometer. Figures 1 and 2 show typical ED-XRF spectra obtained by ED-XRF spectrometry with the operating conditions listed in Table 2. The Cl (K$_\alpha$), Cl (K$_\beta$) and Sb (L$_\alpha$) shown in Figs 1, which were observed at lower energy, showed small signals even though Br (K$_\alpha$) and Sb (K$_\alpha$) shown in Figs 2 revealed larger ones. Figures 3 show the relative changes of Br (K$_\alpha$) signals observed for both ABS and PC resin disks. In the present study, a signal correction method by Compton scattering was carried out to compensate possible signal fluctuations due to the different shapes of plastic resin disks, especially for ABS ones observed as shown in photos 1. It could be seen in Figs 3 (a) and (b), the Br (K$_\alpha$) signals without the correction by Compton scattering showed more than 100 % as well as larger variations which were attributed to the changing shapes of the ABS resin disks obtained after the long-time heating. On the other hand, the maximum
decreasing rate of ca. 5 % was observed for ABS resin disks when the correction method by
Compton scattering was carried out. Since the Br \( (K_\alpha) \) signal was observed at higher energy range
of 11.6-12.2 keV as listed in Table 2, the decreasing rate could be related to the concentration
changes of Br on bulk. On the other hand, the decreasing of Br \( (K_\alpha) \) signals was not observed for
PC resin disks by ED-XRF spectrometry in the present study. Since PC resin showed higher thermal
tolerance than that of ABS,\(^{26}\) these elements could also be considered to be stable in the PC disk.
Figures 4 show the relative changes of Cl \( (K_\alpha) \) and P \( (K_\alpha) \) signals observed for ABS resin disks. No
relative changes were observed for PC resin disks (data not shown). As can be seen in Figs 4, Cl
and P in ABS resin disks with long-time heating decreased to 20-30 % compared to the one without
heating. It was also found that the decreasing rate of these elements in ABS resin disk containing
4 FRs (including Sb\(_2\)O\(_3\)) under the heating within 10-25 days was 10-20 % which was smaller than
the one containing 3 FRs (without Sb\(_2\)O\(_3\)). From these results, it was considered that the ABS disk
containing 4 FRs was seemed to show enhanced thermal tolerance compared to the one containing
3 FRs even though the effect was not enough for long-time heating for more than 25 days. Figures
5 show the relative changes of Sb \( (K_\alpha) \) and Sb \( (L_\alpha) \) signals observed for ABS resin disks. It was
found that the maximum decreasing rates of ca. 5 % and ca. 25 % were observed for Sb \( (K_\alpha) \) and Sb
\( (L_\alpha) \) signals, respectively. Since the Sb \( (K_\alpha) \) signal was observed at higher energy range of 25.9-26.7
keV as listed in Table 2, the relative changes could be related to the concentration changes of Sb on
bulk. On the other hand, it was considered that Sb \( (L_\alpha) \) signal reflected Sb from surface of the disk
since the signal was observed at lower energy range of 3.4-3.8 keV. From this point of view, the
relative changes of Cl and P shown in Figs 4 were also considered from the surface of the disk since
these signals were observed at lower energy ranges of 2.4-2.8 and 1.8-2.2 keV, respectively. The
analytical depths for the measurements of Cl, P and Sb \( (L_\alpha) \) could be 0.1-0.4 mm from the surface of
ABS disk.\(^{30,31}\) From these results, decreasing rate of Cl, P and Sb including Br could be considered
to be maximum ca. 5 % for the ABS resin disk.
3-3. Changing mass and chemical form of ABS and PC resin disks by long-time heating

The changing mass for ABS and PC resin disks was investigated as shown in Figures 6 (a) and (b), respectively; since their masses could be changing due to the loss of water from disk surface or the changing their chemical forms by the long-time heating. As can be seen in Figure 6 (a), the masses of ABS resin disks containing 3 FRs and 4 FRs increased up to ca. 4% during the long-time heating for 722 days and the changes of their masses were similar to each other except for the ones observed for 10-25 days heating. The observed phenomena could fit the changes of the colors between ABS resin disks containing 3 FRs and 4 FRs observed for 10-25 days (see photos 1). From these results, the concentrations of FRs and related elements should be changed since the masses of ABS resin disks increased. If the FRs and related elements are not lost, their concentrations become ca. 4% lower than those of original ABS resin disks during the long-time heating for 722 days. As mentioned in previous section, the decreasing rates of Cl, P and Sb including Br could be considered to be maximum ca. 5% and the rates were similar to the increasing rate of mass (ca. 4%) for the ABS resin disk, even though both rates were not completely matched. On the other hand, PC resin disks showed almost no mass changes from original one to the ones with long-time heating for 722 days as shown in Fig. 6 (b). The changes of masses between the PC resin disks containing 3 FRs and 4 FRs were also similar. Since the PC resin disks showed no changes of colors and masses, it could be considered that the disk seemed to be enough strong and stable with respect to the long-time heating at 80 °C for 722 days, which might be attributed to the thermal tolerance of PC resin. In order to examine the changing mass of ABS resin disk during the long-time heating, FTIR measurement was carried out to evaluate the chemical form. Figures 7 show ATR-FTIR spectra observed for both ABS and PC resin disks. The negative spectra with some noises observed were attributed to the effects of CO$_2$ (2300-2400 cm$^{-1}$) and H$_2$O (ca. 1600 cm$^{-1}$ and 3600-3800 cm$^{-1}$) in air from blank measurement. It could be seen from Figs 7 (a) and (b), the differences during long-time heating were found for ABS resin disks especially in 910 cm$^{-1}$, 1000-1250 cm$^{-1}$, 1600-1700 cm$^{-1}$ and 2800-2900 cm$^{-1}$ which were corresponded to C=C double
bound, ether (C-O-C), ketone (C=O) and methylene (=CH₂) groups, respectively.³² It could be seen that the C=C double bond as well as methylene group decreased and both the ether and ketone groups increased in ABS resin disk. Since the C=C double bond could be found in butadiene group in ABS, the double bond was considered to be oxidized by the oxygen from ambient air and changed to ether and ketone groups during long-time heating which resulted in the increasing mass of ABS resin disk. Similar phenomenon was observed in our previous study.³³ On the contrary, no differences were found for PC resin disks as shown in Figs 7 (c) and (d). Since the PC resin does not have C=C double bond like a butadiene group in ABS, the oxidation did not occur which sustained its chemical form stable without mass changing. From these results, though the PC resin disk seemed to be strong and stable enough with respect to the long-time heating at 80 °C, the ABS resin disk was considered to be oxidized which resulted in increasing its mass as well as decreasing concentrations of FRs contained. Though both increasing and decreasing rates were not completely matched, most of the changes in concentrations of elements from FRs could be due to the changes in mass of the ABS resin disk observed during the long-time heating. Therefore, it could be also considered that the most of the elements from FRs in ABS and PC resin disks was not lost even if these plastics were discarded at harder environment such as higher temperature examined in the present study.

4. Conclusion

The effect of long-time heating for elements from FRs such as DP, TBBPA, TPhP and Sb₂O₃ in both ABS and PC resin disks was examined in the present study. The decreases in concentrations of ca. 5 % for Br, Cl, P and Sb originated from FRs were found for ABS disk; however, no changes were observed for PC disk. The increases in mass of ABS resin disk was found up to ca. 4 % during the long-time heating, even though the PC resin disk did not reveal any large changing. The increases for both ether (C-O-C) and ketone (C=O) groups and the decreases of C=C double bond as well as methylene group (=CH₂) for ABS resin disk were found and it was considered to result in an increase...
of ca. 4% mass by oxidation. Though both decreasing and increasing rates were not completely matched, most of the changes in concentrations of elements could be due to the changes in mass of the ABS resin disk observed during the long-time heating. Though the long-time heating could be considered to be harder environment than that of usual dump site for discarding electric and electronic equipment, it was concluded that the most of elements from DP, TBBPA, TPhP and Sb$_2$O$_3$ contained as FRs in the ABS and PC resin plastics examined in the present study were not lost, even though the color, shape, mass and chemical form of ABS resin disk were changed.

Acknowledgements

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References


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Table 1  Concentrations (mg kg\(^{-1}\)) of FRs and related elements in ABS and PC disks

<table>
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<th>Element</th>
<th>ABS disk</th>
<th>PC disk</th>
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<tr>
<td>DP</td>
<td>449 ± 6</td>
<td>450 ± 11</td>
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<tr>
<td>TBBPA</td>
<td>909 ± 26</td>
<td>924 ± 42</td>
</tr>
<tr>
<td>TPhP</td>
<td>1011 ± 27</td>
<td>1023 ± 31</td>
</tr>
<tr>
<td>Cl</td>
<td>329 ± 3</td>
<td>338 ± 5</td>
</tr>
<tr>
<td>Br</td>
<td>561 ± 10</td>
<td>568 ± 10</td>
</tr>
<tr>
<td>P</td>
<td>90 ± 1</td>
<td>91 ± 1</td>
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<tr>
<td>Sb</td>
<td>426 ± 6</td>
<td>395 ± 1</td>
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Table 2  Operating conditions of ED-XRF spectrometer (Rayny EDX-720, Shimadzu)

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<td>X-ray irradiated diameter (circle)</td>
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<td>Atmosphere</td>
<td>Atmospheric pressure</td>
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<tr>
<td>Measurand</td>
<td>Br (K(_\alpha))</td>
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<td>Filter</td>
<td>Ag</td>
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<td>Current ((\mu)A)</td>
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<tr>
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<td>Upper limit of detector dead time (%)</td>
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<td>Detector</td>
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</table>
Figure captions

Photos 1  ABS resin disks containing (a) 3 FRs and (b) 4 FRs observed with long-time heating at 80 °C for 722 days.

Photos 2  PC resin disks containing (a) 3 FRs and (b) 4 FRs observed with long-time heating at 80 °C for 722 days.

Figure 1  Typical ED-XRF spectra observed for Cl (Kα), P (Kα) and Sb (Lα) from ABS and PC resin disks. The ED-XRF spectra are shown in the range of (a) 0-20 keV and (b) 0-5 keV. The actual values of voltage, current and detector dead time for ED-XRF spectrometer were 15 kV, 756 μA and 39 %, respectively.

Figure 2  Typical ED-XRF spectra observed for (a) Br (Kα) and (b) Sb (Kα) from ABS and PC resin disks. The actual values of voltage, current and detector dead time for ED-XRF spectrometer were (a) 50 kV, 350 μA and 39 %, and (b) 50 kV, 36 μA and 40 %, respectively.

Figure 3  Relative changes of Br (Kα) originated from FRs of TBBPA in ABS and PC resin disks containing 3 FRs and 4 FRs. (a) and (b) or (c) and (d) show the changes for ABS or PC resin disks containing 3 FRs and 4 FRs, respectively. The symbols of ○ and × show the data with and without corrections by Compton scattering, respectively.

Figure 4  Relative changes of Cl (Kα) and P (Kα) originated from FRs of DP and TPhP, respectively, in ABS resin disks containing 3 FRs and 4 FRs. (a) and (b) or (c) and (d) show the changes for Cl or P in ABS resin disks containing 3 FRs and 4 FRs, respectively. The symbols of ○ and × show the data with and without corrections by Compton scattering, respectively.
Figure 5  Relative changes of (a) Sb (L\textsubscript{\alpha}) and (b) Sb (K\textsubscript{\alpha}) originated from FRs of Sb\textsubscript{2}O\textsubscript{3} in ABS resin disks containing 4 FRs. The symbols of ○ and × show the data with and without corrections by Compton scattering, respectively.

Figures 6  Changes in the masses of (a) ABS and (b) PC resin disks as a function of the heating time. The symbols of ● and × show the data for plastic disks containing 3 FRs and 4 FRs, respectively.

Figures 7  ATR-FTIR spectra observed for ABS and PC resin disks containing 3 FRs or 4 FRs. (a) and (b) or (c) and (d) show the spectra for ABS or PC resin disks containing 3 FRs and 4 FRs, respectively.
(a) ABS resin disks containing 3 FRs of DP, TBBA and TPhP

(b) ABS resin disks containing 4 FRs of DP, TBBA, TPhP and Sb$_2$O$_3$

Photos 1 Ohata et al.

(a) PC resin disks containing 3 FRs of DP, TBBA and TPhP

(b) PC resin disks containing 4 FRs of DP, TBBA, TPhP and Sb$_2$O$_3$

Photos 2 Ohata et al.
Figures 1 Ohata et al.

Figures 2 Ohata et al.
Figures 3  Ohata et.al.

Figures 4  Ohata et.al.
Figures 5 Ohata et al.

Figures 6 Ohata et al.
Figures 7  Ohata  et.al.