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Consideration of recommendations to the Conference of the Parties

Guidance on flame-retardant alternatives to commercial pentabromodiphenyl ether (<u>C-</u>PentaBDE)

Note by the Secretariat

- At its third meeting, the Committee suggested that the risk management evaluation of pentabromodiphenyl ether should include guidance on alternatives and Ms. Liselott Säll (Norway) and Mr. Bo Wahlström (Sweden) offered to make an initial effort to prepare guidance for commercial pentabromodiphenyl ether.1
- Accordingly, the Secretariat entrusted Ms. Säll and Mr. Wahlström with the preparation of the document providing guidance on flame-retardant alternatives to pentabromodiphenyl ether (PentaBDE). The document is contained in the annex to the present note as submitted and has not been formally edited by the Secretariat.

Comment [spencerv1]:

This guidance addresses the alternatives to use of Commercial Pentabromodiphenyl ether the main components of which were: 2,2', 4,4'- tetrabromodiphenyl ether (BDE-47 CAS No. 40088-47-9);

-2,2',4,4',5 pentabromodiphenyl ether (BDE-99 CAS No. 32534-81-9).

Penta BDE is a congener family of PBDEs. To be consistent references throughout this guidance should always be to C-PentaBDE.

UNEP/POPS/POPRC.3/20, para 47.

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UNEP/POPS/POPRC.4/1.

Annex

Guidance on flame-retardant alternatives to pentabromodiphenyl ether (PentaBDE)

Preface

In 2005 Norway nominated the brominated flame retardant pentabromodiphenylether (PentaBDE) as a persistant organic pollutant (POP) to be evaluated for inclusion in the Stockholm convention. Based on the Risk Profile developed in 2006 and the Risk Management Evaluation Report developed in 2007 the POP Review Committee (POPRC) concluded that global action on PentaBDE is warranted. At the POPRC meeting in November 2007 Norway was commissioned to issue a guide of alternative flame retardants to PentaBDE. The Norwegian Pollution Control Authority (SFT) has therefore commissioned Swerea IVF (Sweden), to perform this guide that will be presented to the POPRC-meeting in Geneva in October 2008.

SFT, Oslo, June 2008

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Summary

Flame retardants represent a large group of chemicals that mainly consist of inorganic and organic compounds based on bromine, chlorine, phosphorus, nitrogen, boron, and metallic oxides and hydroxides. They are either additive or reactive.

Reactive flame-retardants are added during the polymerisation process and become an integral part of the polymer and forms a co-polymer. The result is a modified polymer with flame retardant properties and different molecule structure compared to the original polymer molecule.

Additive flame-retardants are incorporated into the polymer prior to, during, or more frequently after polymerisation. Additive flame-retardants are monomer molecules that are not chemically bounded to the polymer. They may therefore, in contrast to reactive flame retardants, be released from the polymer and thereby also discharged to the environment.

In contrast to most additives, flame-retardants can appreciably impair the properties of polymers. The basic problem is the trade-off between the decrease in performance of the polymer caused by the flame-retardant and the fire requirements. In addition to fulfilling the appropriate mandatory fire requirements and rules, a feasible flame-retardant shall, at most, fulfil the whole range of physical, mechanical, health and environmental properties and simultaneously be cost efficient for the market.

Halogenated flame-retardants are primarily based on chlorine and bromine. A large group of additive flame-retardant is the polybrominated diphenylethers (PBDEs), which include all congeners of commercial pentaBDE (C-PentaBDE). PBDEs are used in many different applications worldwide, and have the second highest production volume of brominated flame retardants currently used (today mainly represented by decabromodiphenylether).

C-PentaBDE has been produced in Israel, Japan, US and the EU, but production in these regions ceased in the beginning of this millenium. There are indicative reports of an expanding production of brominated flame retardants in China. No official information is available for production of C-PentaBDE in China, this is also the case for Israel and Eastern European countries outside EU.

PBDEs are used in different resins, polymers, and substrates at levels ranging from 5 up to 30% by weight. The main historic use of C-PentaBDE was in flexible polyurethane foam (PUR), but it has also been used in epoxy resins, PVC, unsaturated thermoset polyesters (UPE), rubber, paints and lacquers, textiles and hydraulic oils. The quantities used for each specific application are not publicly available.

Like any other additives, a flame retardant will be selected for the particular properties it imparts to make the final product satisfy the specifications established by the customer. New flame retardant solutions are constantly introduced and some disappear from the market for a number of reasons. However, there is a variety of optional chemical systems available on the market that actually work as alternatives to C-PentaBDE. Their use in commercial applications are illustrated in table 4, and their environmental and health properties are described in table 7 in this report. However, it needs to be clearly understood that each flame retardant application is specific and unique and there are no single universal solutions for fire protection of materials and applications.

Even though there are toxicological and ecotoxicological data gaps for the potential alternatives to C-PentaBDE, the data available clearly show that there are commercially available alternative flame retardants that are less hazardous than C-PentaBDE. It is important to search for further health and environmental data on a sound scientific basis for potential alternative flame retardants and avoid those flame retardants that may pose any risk to health and the environment.

1. Introduction

1.1 Flame retardants

With the increasing use of thermoplastics and thermo sets on a large scale for applications in buildings, transportation, electrical engineering and electronics, a variety of flame-retardant systems have been developed over the past 40 years. They mainly consist of inorganic and organic compounds based on bromine, chlorine, phosphorus, nitrogen, boron, and metallic oxides and hydroxides. Today, these flame-retardant systems fulfil the multiple flammability requirements developed for the above-mentioned applications (EHC 1921997).

Flame-retardants are either additive or reactive. Reactive flame-retardants are added during the polymerisation process and become an integral part of the polymer and forms a co-polymer. The result is a modified polymer with flame retardant properties and different molecule structure compared to the original polymer molecule. This prevents them from leaving the polymer and keeps the flame retardant properties intact over time with very low emissions to the environment (Danish EPA 1999). Reactive flame-retardants are mainly used in thermosets, especially polyester, epoxy resins and polyurethane's (PUR) in which they can be easily incorporated (Posner 2006).

Additive flame-retardants are incorporated into the polymer prior to, during, or more frequently after polymerisation. They are used especially in thermoplastics. If they are compatible with the plastic they act as plasticizers, otherwise they are considered as fillers. Additive flame-retardants are monomer molecules that are not chemically bound to the polymer. They may therefore be released from the polymer and thereby also discharged to the environment.

1.2 Categories of flame retardants

Flame retardants are added to various kinds of polymers, both synthetic and natural, to enhance the flame retardant properties of the polymers. Around 350 different chemical flame retardant substances are described in literature, with no specific reference to national or international fire regulations. Such a reference would strengthen the case for the use of the particular substance, for any specific market.

The Index of Flame retardants 1997, an international guide, contains more than 1000 flame retardant products (preparations and substances) listed by trade name, chemical name, application and manufacturer. This index describes around 200 flame retardant substances used in commercial flame retardant products.

There are four main families of flame retardant chemicals.

- Inorganic
- Organophosphorous
- Nitrogen based
- · Halogenated flame retardants

Inorganic flame-retardants are metal hydroxides (such as aluminium hydroxide and magnesium hydroxide), ammonium polyphosphate, boron salts, inorganic antimony, tin, zinc and molybdenum compounds, and elemental red phosphorous. Both aluminium hydroxide, also sometimes called aluminium trihydrate (ATH), and magnesium hydroxide are used as halogen free alternatives to brominated flame retardants and they also function as smoke suppressants. Inorganic phosphorus compounds are widely used as substitutes to brominated flame retardants. Inorganic flame-retardants are added as fillers into the polymer and are considered immobile in contrast to the organic additive flame-retardants. Antimony trioxide and zinc borate are primarily used as synergists in combination with halogenated flame-retardants. Alternative synergists include zinc hydroxystannate (ZHS), zinc stannate (ZS), and certain molybdenum compounds. The whole group of inorganic flame-retardants represents around 50% by volume of the global flame retardant production, mainly as aluminium trihydrate, which is in volume the biggest flame retardant category in use on the market.

Organophosphorous flame-retardants are primarily phosphate esters and represent around 20% by volume of the total global production. This category is widely used both in polymers and textile cellulose fibres. Of the halogen-free organophosphorous flame-retardants in particular, triaryl phosphates (with three benzene rings attached to a phosphorous group) are used as alternatives to brominated flame-retardants. Organophosphorous flame-retardants may in some cases also contain bromine or chlorine.

Nitrogen based organic flame-retardants inhibit the formation of flammable gases and are primarily used in polymers containing nitrogen such as polyurethane and polyamide. The most important nitrogen-based flame-retardants are melamine's and melamine derivatives and these act as intumescent (swelling) systems.

Halogenated flame-retardants are primarily based on chlorine and bromine. These flame retardants react with flammable gases to slow or prevent the burning process. The polybrominated diphenylethers (PBDEs) are included in this group, where all the isomers of PentaBDE are represented. The group of halogenated flame-retardants represent around 30% by volume of the global production, where the brominated flame-retardants dominate the international market (SRI Consulting 2005).

Halogenated flame-retardants can be divided into three classes:

- Aromatic, including PBDEs in general and PentaBDE in particular.
- Cycloaliphatic, including hexabromocyclododecane (HBCDD).
- Aliphatic, globally representing a minor group of substances.

2. Requirements for feasible flame retardants

2.1 Fire requirements

Tightened legislation and tougher fire requirements are the major forces that have driven forward development towards functionally better and more effective flame retardants. In the light of this trend, a large number of specific fire standards with unique fire requirements have been developed internationally for various widely differing situations. Customer's requirements are absolute, whether they are public institutions, international organisations or businesses on the market. If the fire requirements are not met, there is no market for the individual supplier and the manufacturer. On the other hand, there are no prescriptive fire requirements at all stipulating that particular flame retardants have to be used to meet the requirements. The choice of flame retardants is left entirely to the manufacturer of the final product.

In some cases the requirements are so strict that the alternatives are not economically feasible or the environmental requirements or regulations in that part of the world do not make the manufacturers choice of flame retardants possible to apply. Worse quality characteristics may also be limiting factors in the manufacturer's choice of flame retardants (Posner 2005).

2.2 Quality properties on fire retarded materials

In contrast to most additives, flame-retardants can appreciably impair the properties of polymers. The basic problem is the trade-off between the decrease of performance of the polymer caused by the flame-retardant and the fire requirements. In addition to fulfilling the appropriate mandatory fire requirements and rules, a feasible flame-retardant shall, at most, fulfil all of the qualities mentioned below.

Fire retardant properties

- Commence thermal activity before and during the thermal decomposition of the polymer
- Not generate any toxic gases beyond those produced by the degrading polymer itself
- Not increase the smoke density of the burning polymer

Mechanical properties

- Not significantly alter the mechanical properties of the polymer
- Be easy to incorporate into the host polymer
- Be compatible with the host polymer

Physical properties

- Be colourless or at least non-discolouring
- · Have good light stability
- · Be resistant towards ageing and hydrolysis
- Not cause corrosion

Health and envrionmental properties

- · Not have harmful health effects
- · Not have harmful environmental properties

Commercial viability

· Be commercially available and cost efficient

3. Characteristics of C-PentaBDE

Brominated diphenylethers (PBDEs) are a large group of additive brominated flame retardants with a versatile use in many applications worldwide. PBDEs are the second highest production group of brominated flame retardants currently used, mainly represented today by the fully brominated decabromodiphenylether.

Commercial pentabromodiphenylether (C-PentaBDE) is a mixture of two major congeners i.e. 2,2',4,4'´tetrabromodiphenylether (BDE-47), and 2,2',4,4'´,5-pentabromodiphenylether (BDE-99). Trace amounts of 2,2',4-tri-bromodiphenylether (BDE-17) and 2,4,4'´-trisbromodiphenylether (BDE-28) are also present in C-PentaBDE. Both BDE-17 and BDE-28 are precursors in the formation of major congeners in C-PentaBDE such as BDE-47.

Continued bromination of BDE-47 yields mainly BDE-99 and 2,2′,4,4′,6-pentabromodiphenylether (BDE-100). Percentages of BDE-99 and BDE-100 are 35% and around 7% respectively. Further bromination yields 2,2′,4,4′,5,5′-hexabromodiphenylether (BDE-153) and 2,2′,3,4,4′,5′,6 – heptabromodiphenylether (BDE-154), that are also present in C-PentaBDE (Alaee et. al 2003).

Table 1 Composition of C-PentaBDE.

Categories	Tridiphenyl		Tetradiphenyl	Pentadiphenyl		Hexadiphenyl	Heptadiphenyl
of PBDEs	Es ethers eth		ether	ethers		ether	ether
Congeners	BDE-17	BDE-28	BDE-47	BDE-99	BDE-	BDE-153	BDE-154
					100		
Content	Traces	Traces	Major	Major	Minor	Minor	Traces

PentaBDE is widespread in the global environment. Levels of components of C-PentaBDE have been found in humans in all UN regions. Most trend analyses show a rapid increase in concentrations of PentaBDE in the environment and in humans from the early 1970s to the middle or end of the 1990s, reaching plateau levels in some regions in the late 1990s, but continuing to increase in others. The levels in North America and the Arctic are still rising. Vulnerable ecosystems and species are affected, among them several endangered

species. Some individuals of endangered species show levels high enough to be of concern. Toxicological studies have demonstrated reproductive toxicity, neurodevelopmental toxicity and effects on thyroid hormones in aquatic organisms and in mammals. The potential for the toxic effects in wildlife, including mammals, is evident.

Based on the information in the risk profile, PentaBDE, due to the characteristics of its components, is likely, as a result of long-range environmental transport and demonstrated toxicity in a range of non-human species, to cause significant adverse effects on human health or the environment, such that global action is warranted.

4. Commercial use and production of PentaBDE

4.1 Historic production of PentaBDE

Based on the latest available information from Bromine Science and Environmental Forum (BSEF), the total market demand of C-PentaBDE has decreased from 8,500 tons in 1999 to 7,500 tons in 2001. The estimated cumulative use of C-PentaBDE since 1970 was in 2001 estimated to 100 000 t (BSEF 2001),(UNEP/POPS/POPRC.3/20/Add1 2007).

Table 2 C-PentaBDE volume estimates: Total market demand by region in 2001 in metric tons (and by percent) (BSEF 2001).

	Americas	Europe	Asia	Rest of the world	Total	% of total world usage of BFR
Penta-mix PBDE formulation	7100	150	150	100	7500	4

C-PentaBDE has been produced in Israel, Japan, US and the EU. China may have produced for their market as well. Since 2004 C-PentaBDE is no longer produced at least by any BSEF member company. Today there is no production in Japan and C-PentaBDE was voluntarily withdrawn from the Japanese market in 1990 (UNECE 2007) There is no official information available from the Israeli government of any present production or use of PentaBDE, however as BSEF members do not make or supply PentaBDE and Israel Chemicals Ltd is a BSEF member, it is unlikely to be made or used there.

Since the late 1990's it is believed to have been produced primarily in the US although there are some indications. The sole producer of C-PentaBDE in the US, the Great Lakes Chemical Corporation (now Chemtura), voluntary ended their production of C-PentaBDE in 2004². Before the phase-out in US the majority of C-PentaBDE formulation produced globally was used in North America (>97%). At the end of 2004, in the US, approximately 7.5% of the more than 1 million tonnes of flexible polyurethane foam produced each year in the US contained the C-PentaBDE formulation (UNECE 2007).

Production in the EU ceased in 1997. Usage in EU has been declining during the second half of the 1990's and was estimated to be 300 metric tonnes in year 2000, used solely for PUR production (EU 2000). The use of C-PentaBDE was banned in the EU in 2004 through the restrictions on marketing and the use of dangerous substances in the Council directive 2003/11/EC. From 1st of July 2006 PentaBDE was restricted in electrical and electronic appliances through the RoHS –directive [2002/95/EC].

Results from a survey conducted in Canada in 2000 indicated that approximately 1300 tonnes of PBDE commercial products were imported into Canada. Based on quantities reported, C-PentaBDE was imported in the greatest volume. Now PentaBDE is on the list of toxic substances in the Canadian Environmental Protection Act (CEPA 1999).

On total market, demand for C-PentaBDE was: 8,500 metric tons 1999; and 7,500 metric tons in 2001

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² Landry S Albermarle, personal communication (2008)

4.2 Historic use of PentaBDE

PBDEs are used in different resins, polymers, and substrates at levels ranging from 5 up to 30% by weight. The quantities used for each specific application are not publicly available (USEPA (Dfe) 2004).

Table 3 Historic use of PentaBDE in various materials and applications (EHC 162, (Danish EPA 1999), (Renner 2000), (UNEP/POPS/POPRC.3/20/Add1).

Materials/polymers/resins	Applications	Commercial commodities for	
		the applications	
Epoxy resins	Circuit boards, protective coatings	Computers, ship interiors,	
		electronic parts.	
Polyvinylchloride (PVC)	Cable sheets	Wires, cables, floor mats,	,
		industrial sheets.	
Polyurethane (PUR)	Cushioning materials, packaging,	Furniture, sound insulation	Ų,
	padding	packaging, padding panels,	ì
		wood imitations,	
		transportation.	
Unsaturated (Thermoset)	Circuit boards, coatings	Electrical equipment, coatings	
polyesters (UPE)		coatings for chemical	
		processing plants mouldings,	
		military	
		and marine applications:	
		construction panels.	l
Rubber	Transportation	Conveyor belts, foamed pipes for	
		insulation.	
Paints/lacquers	Coatings	Marine and industry lacquers for	
		protection of containers	
Textiles	Coatings	Back coatings and impregnation	`
		for carpets,	
		automotive seating,	
		furniture in homes and official	
		buildings, aircraft, underground.	
Hydraulic oils	Drilling oils, hydraulic fluids	Off shore, coal mining	

Greater than 95% of the US production of PentaBDE in the years leading up to the voluntary withdrawal of production and sales in 2004 was consumed in the US for the production of flexible polyurethane foam for the production of cushioning and other uses. Although there is no data available on the proportions of use of C-PentaBDE for the different applications in table 3, this indicates that these were minor or non-existent applications by the time production stopped.

4.3 Present use and trends in production of PentaBDE

C-PentaBDE is no longer produced anywhere worldwide by BSEF member companies. There is no current production of C-PentaBDE in Europe, Japan, Canada, Australia and the US. In addition, through direct personal contacts with BSEF, it has been confirmed that C-PentaBDE is not produced in Africa, any other Asian countries (including China) and South and Latin America, 4.

China's flame retardant market has seen rapid growth in the last five years aided by favorable economic growth, increasing demand from the end user market, increasing awareness on fire safety issues and numerous other factors. China's flame retardant market is one of the most dymanic flame retardant markets around the world, and establishment and implementation of government regulations and standards are playing an important role in the growth of the market. The current growth levels are expected to continue with more environment-friendly non-halogenated types driving the growth of the market.

Comment [spencerv2]:

BSEF confirms that there is no production of C-PentaBDE in Eastern European countries outside the EU, by our member companies.

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Investigations through direct contacts with industry and studies of relevant sources information on any historic or present production or use of C-PentaBDE in Eastern European countries outside the EU have been sought, but no information of such activities was found.

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Comment [b3]: c-PentaBDE was a minor use flame retardant in "potting compounds" and encapsulants around components attached to the board and in early versions of low grade printed wiring boards.

Comment [b4]: Another very minor use for c-PentaBDE use in plasticized PVC

Comment [b5]: Another very minor use for c-PentaBDE use UPE

Comment [b6]: Another very minor use for c-PentaBDE use in rubber

Comment [b7]: Another very minor use for c-PentaBDE use in paints/laquers

Comment [b8]: Another very minor use for c-PentaBDE use in textile coatings

Comment [b9]: c-PentaBDE might have been used in hydraulic oils but not in the last 20 years

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³ Dr. Didier, M Trimbos, Eurobrom, personal communication (2008)

⁴ Baker A, Dead Sea bromine Ltd, personal communication (2008)

Xu Dan, Industry Analyst for the Chemicals, Materials and Food Group of Frost & Sullivan's Asia Pacific reveals that, "China has increasingly become the global production base for electronic products and the main region for consumption of plastics." She declares, "The increase in demand for E&E equipments, as well as the development of Building & Construction, and Automotive industries is driving the growth of the flame-retardant market in China." (Frost & Sullivan 2007).

In China there are two major global suppliers, and possible producers, of C-PentaBDE that have been identified through the internet. There is no official data available of any possible production of C-PentaBDE in China⁵. However, China has its own RoHS since 2006, where the use of PBDEs in electronics is banned (SJ/T 11363 2006).

Comment [b10]: This information is not relevant for BSEF. BSEF member companies do not produce C-PentaBDE in China.

5. Alternative flame retardants and alternative technical solutions to PentaBDE

Like any other additives, a flame retardant will be selected for the particular properties it imparts to make it satisfy the specifications for the final compound established by the customer. As mentioned earlier, different flame retardants may be chosen to give different levels of fire protection depending on the specific levels defined by the customer and that particular market. New flame retardant solutions are constantly introduced and some disappear from the market for a number of reasons. Therefore table 4 is a on-the-spot account and cannot be complete, but only act as a guide that illustrates the variety and optional chemical systems that are available and actually work as viable alternatives to C-PentaBDE. However, it needs to be clearly understood that each flame retardant application is specific and unique, and there are no single universal solutions for fire protection of materials and applications.

⁵ DiGangi J, personal communication (2008)

Table 4 Overview of use of alternative flame retardants to PentaBDE in several materials and applications. (EHC 162 1994), (UNEP/POPS/POPRC.3/INF/23 2007), (KemI 2006), (Timpe 2007), (Haglund 2000), (Troitzsch 2007), (Supresta 2008).

Materials /polymers /resins	Inorganic alternatives to PentaBDE	Phosphorous/ nitrogen organic alternatives to PentaBDE	Halogen organic alternatives to PentaBDE	Alternative flame inherent materials	Applications	Commercial commodities for the applications	
Epoxy resins	Aluminium hydroxide (ATH) Magnesium hydroxide Ammonium poly phosphate Red phosphorous Zinc hydroxystannate (ZHS), Zinc stannate (ZS) & ZHS/ZS-coated ATH	Metallic phosphinates Reactive nitrogen and phosphorous constituents (unspecified) DOPO ⁶	Tetrabromobis phenol A (reactive) Etylenebis (tetrabromo) phtalimid	Polyethylene sulphide	Circuit boards, protective coatings	Computers, ship interiors, electronic parts.	
Polyvinylchloride (PVC)	Aluminium hydroxide (ATH) Zinc borate Zinc- molybdenum compounds (together with phosphate esters) Zinc hydroxystannate (ZHS), Zinc stannate (ZS) & ZHS/ZS-coated ATH	Tricresyl phosphate (also plasticizer)	Tris (dichloropropyl) phosphate Vinylbromide	Rigid PVC is flame inherent itself	Cable sheets	Wire end cables, floor mats, industrial sheets.	
Polyurethane (PUR)	Ammonium poly phosphate Red phosphorous Reofos (non-halogen flame retardant)	Melamine (nitrogen based) Dimethyl propane phosphonate (DMPP)	Bromoalkyl phosphates Tetrabromophtalic anhydride Tris(chloroethyl) phosphate (TCPP) (together with brominated polyols or red phosphorous)	Intumescent systems	Cushioning materials, packaging, padding	packaging padding panels, we imitations transporta	ment [RCC11]: Reofos is ename belonging to tutra Corporation. Chemtura tutra Corporation. Chemtura tember of BSEF. They offer aber of products under the os brand. Some of these are n legacy c-PentaBDE rations (e.g. PUR foam). But of the Reofos brand products organic compounds.

 $^{^6}$ DOPO=Dihydrooxaphosphaphenanthrene oxide

Comment [b12]: Please see comments above. In most of the applications mentioned in this table c-PentaBDE was never used or uses were very minor.

Materials /polymers /resins	Inorganic alternatives to PentaBDE	Phosphorous/ nitrogen organic alternatives to PentaBDE	Halogen organic alternatives to PentaBDE	Alternative flame inherent materials	Applications	Commercial commodities for the applications	
Unsaturated (Thermoset) polyesters (UPE)	Ammonium polyphosphate Aluminium hydroxide (ATH) Magnesium hydroxide Zinc hydroxystannate (ZHS), Zinc stannate (ZS) & ZHS/ZS-coated ATH	Triethyl Phosphate Dimethyl propane phosphonate (DMPP)	Dibromostyrene Tetrabromophtalic anhydride based diol, Tetrabromophtalic anhydride Bis (tribromophenoxy) ethane	Intumescent systems	Circuit boards, coatings	Electrical equipment, coatings coatings for chemical processing plants mouldings, military and marine applications: construction panels.	
Rubber	N/A	Alkyl diaryl phosphates (nitril rubber)	N/A	Intumescent systems	Transportation	Conveyor belts, foamed pipes for insulation.	
Paints/lacquers	N/A	Triaryl phosphates (unspecified)	Tetrabromo phthalate diol Tetrabromophtalatic anhydride based diol Bis (tribromophenoxy) ethane	Intumescent systems Silicone rubber	Coatings	Marine and industry lacquers for protection of containers	
Textiles	Aluminium hydroxide Magnesium hydroxide Ammonium compounds (unspecified) Borax	Tetrakis hydroximethyl phosphonium salts such as chloride (THCP) or ammonium (THPX) Dimethyl phosphono (N-methylol) propionamide Diguanidine hydrogen phosphate Aromatic phosphates (unspecified)	Trichloropropyl phosphate	Intumescent systems Aramide fibres (certain protective applications) Wool Modacrylic	Coatings	comm	ment [b13]: Please see nents above. In most of the actions mentioned in this c-PentaBDE was never used

Materials	Inorganic	Phosphorous/	Halogen organic	Alternative	Applications	Commercial	
/polymers	alternatives to	nitrogen	alternatives to	flame		commodities	
/resins	PentaBDE	organic	PentaBDE	inherent		for the	
		alternatives		materials		applications	
		to PentaBDE					
Textiles cont.		Dimethyl					
		hydrogen					
		phosphite					
		(DMHP)					
		Melamine					
		(nitrogen					
		based)					
		Phospho					
		nitrilic					
		chloride					
		(PNC)					
Hydraulic oils	N/A	Butylated	N/A	N/A	Drilling oils,	Off shore, coa	al
		triphenyl			hydraulic	mining - Co	mment [b14]: Please see
		phosphate			fluids		nments above. In most of the
		esters					lications mentioned in this le c-PentaBDE was never used

N/A : not available or not applicable

6. Present manufacture and use of alternative flame retardants to PentaBDE

6.1 Inorganic flame retardants and synergists

6.1.1 Aluminium hydroxide (ATH)

ATH has been used as a flame retardant and smoke suppressant since the 1960's and it is available in a variety of particle sizes as commercial products. Flame retardation by ATH has been shown to be partly due to the heat sink effect and partly due to the dilution of combustible gases by the water formed as a result of dehydroxylation. Alumina which is formed as a result of thermal degradation of ATH slightly above 200 °C has been shown to form a heat- insulating barrier on the surface that prevents further fire propagation of the matrix material.

The major concern with ATH is the required high loading levels in order to obtain equivalent flame retardant properties as by other additives. These loads can be reduced with a correct choice of particle size, surface modification and proper dispersion in the matrix material (Swaraj 2001). Furthermore, recently developed coated filler products (e.g. ZHS-coated ATH) offer the possibility of equivalent or better flame retardancy and smoke suppression at significantly reduced incorporation levels.

6.1.2 Magnesium hydroxide

Magnesium hydroxide acts in general the same way as ATH, but it thermally decomposes at slightly higher temperatures around 325 °C. Combinations of ATH and magnesium hydroxide function as very efficient smoke suppressants in PVC.

6.1.3 Red phosphorous

Red phosphorous has been reported to be most efficient as a flame retardant in oxygen containing polymers such as polycarbonate, polyethylene terephatalate (PET), polyamide and phenolic resins. Flame retardancy takes place due to formation of phosphorous-oxygen bonds that reduces the ester cleavage into cross linking

aromatic structures with lesser volatility. In addition the red phosphorous creates a heat shield on the polymer surface that result in flame retardant properties. Some drawbacks with the use of red phosphorus are the red colour that could lead to discoloration of polymers and the formation of toxic phosphine gas during combustion.

6.1.4 Ammonium polyphosphate (APP)

APP is mainly used as an acid source in intumescent systems, which are described in more detail in chapter 6.4. APP alone as a flame retardant has been found effective in polyamides and similar polymers.

6.1.5 Antimony trioxide

Antimony trioxide does not function as a flame retardant, but in combination with halogenated flame retardants it functions as a synergist. This term means that the desired effect of two or more components working together is greater than the effect of each of the components separately. As a synergist, the main advantage by the addition of antimony trioxide is to reduce the amount of halogenated flame retardants applied to the polymer.

6.1.6 Zinc borate

Zinc borate (used mainly in PVC) cannot be used alone to achieve desired flame retardant properties in polymers, since it is used as synergist together with other flame retardants, often brominated compounds.

6.1.7 Zinc hydroxystannate (ZHS) and Zinc stannate (ZS)

ZHS and ZS have primarily found use as alternative non-toxic synergists to antimony trioxide in PVC and other halogen-containing polymer systems. However, they have recently found growing applications in halogen-free formulations, and are particularly effective as partial replacements for hydrated fillers such as ATH and magnesium hydroxide, either in the form of powdered mixtures or as coated fillers (Cusack 2005).

6.2 Organophosphorous flame-retardants

6.2.1 Triethyl phosphate

Triethyl phosphate is either used alone or together with a bromine synergist, such as antimony trioxide, for unsaturated polyester resins. Less volatile types of flame retardants include trialkyl phosphates with longer alkyl chains such as tributyl, trioctyl or tris-butoxyethylphoshates. Several high alkylated phosphorous products are commercially available, which is required in lower added concentrations in the polymer.

6.2.2 Aryl phosphates

This large group of organophosphorous flame retardants include triphenyl, isopropyl – and t-butylsubstituted triaryl and cresyl phosphates. Phosphates with larger substitution carbon chains (therefore less volatile) are commercially available beside those mentioned above.

Aryl phosphates are used as flame retardants for phthalate plasticized PVC. It has been shown that although PVC does not require any flame retardancy as a polymer, the addition of phthalate plasticizers makes PVC flammable. Triaryl phosphates are more efficient flame retardants than the alkylated triaryl phosphates. However, the alkylated triaryl phosphates is shown to be more efficient plasticizers than triaryl phosphates.

6.2.3 Halogen containing phosphorous flame retardants

Several halogen containing phosphates, such as chloro and bromophosphates, are commercially available as shown in table 5 below.

Table 5 Commercial phosphorous organic flame-retardant chemical formulations (UNEP/POPS/POPRC.3/INF/23),(Supresta 2008).

Albemarle Corporation	Ameribrom, Inc. (ICL Industrial Products)	Great Lakes Chemical Corporation (now Chemtura)	Supresta (Akzo Nobel)
SAYTEX® RX-8500	FR 513	Firemaster® 550	Fyrol® FR-2
Proprietary reactive	Tribromoneopentyl	Proprietary halogenated aryl	Trs(1,3-dichloro-2-
brominated flame	alcohol	esters, proprietary triaryl	propyl) phosphate
retardant, proprietary aryl	CAS 36483-57-5	phosphate isopropylated,	CAS 13674-87-8
phosphate, triphenyl		triphenyl phosphate	
phosphate			
CAS 115-86-6			
SAYTEX® RZ-243		Firemaster® 552	AB053
Proprietary		Proprietary halogenated aryl	Tris(1,3-dichloro-2-
tetrabromophthalate,		esters, proprietary triaryl	propyl) phosphate
proprietary aryl		phosphate isopropylated,	
phosphate, triphenyl		triphenyl phosphate	
phosphate			
ANTIBLAZE® 195			AC003
Tris(1,3-dichloro-2-			Proprietary organic
propyl) phosphate			phosphate ester,
CAS 13674-87-8			triphenyl phosphate
ANTIBLAZE ® 205			AC073
Proprietary chloroalkyl			Proprietary aryl
phosphate, aryl phosphate			phosphates, triphenyl
and triphenyl phosphate			phosphate
ANTIBLAZE® 180			Fyrquel 150, Fyrquel
<i>Tris</i> (1,3-dichloro-propyl)			220, and Fyrquel 300
phosphate			Butylated triphenyl
CAS 13674-87-8			phosphate esters
ANTIBLAZE® V-500			
Proprietary chloroalkyl			
phosphate, aryl phosphate			
and triphenyl phosphate			
ANTIBLAZE® 182			
Proprietary chloroalkyl			
phosphate, aryl phosphate			
and triphenyl phosphate			
ANTIBLAZE®TL10ST			
(proprietary			
chlorophosphate)			
CAS # propietary mixture			

Chloro alkyl phosphates have been found effective in flexible polyurethane (PUR) foams, but since they are not stable during curing reactions of PUR, which is a strong exotermic reaction with heat generated, they render discolouring problems. Therefore a blend of PentaBDE and triaryl phosphates was, and may still be, used in flexible PUR to avoid this discoloration problem.

Brominated phosphates have been reported to be effective flame retardants without the use of antimony trioxide as synergist, for polyesters and other polymers, such as HIPS and other polystyrenes that may not be relevant for this report.

6.2.4 Reactive phosphorous flame retardants

Reactive phosphorous polyols, including phosphine oxide diol and triol, have also been reported as useful flame retardants in PUR, PET and epoxy resins. No specified information of their flame retardant efficiency in these polymers has been found in this study.

6.3 Nitrogen based organic flame-retardants

Nitrogen containing polymers have been found to be synergetic with phosphorous compounds. For example, polymers containing amine and amide groups were synergistic whereas polymers containing nitrile have been found to be antagonistic. One common example of nitrogen based flame retardants is melamine, which is also a common constituent in intumescent systems.

6.4 Intumescent systems

Intumescent (or swelling) systems have existed since the 1940s, principally in paints. Several intumescent systems linked to textile applications have been on the market for about 20 years, and have successfully shown their great potential. Intumescent systems include use of expandable graphite impregnated foams, surface treatments and barrier technologies of polymer materials.

Almost all intumescent systems comprise, in general, of three basic components

- · a dehydrating component, such as APP
- a charring component, such as pentaerythritol (PER)
- a gas source, often a nitrogen component such as melamine

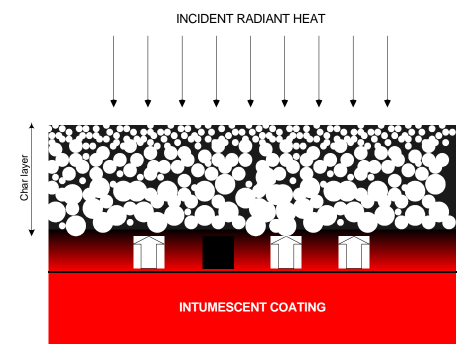


Figure 1 Mechanisms for intumescent systems.

The main function of APP is to catalyse the dehydration reaction of other components in the intumescent system. It has been shown that in spite of the fact that APP functions as a catalyst it has been used in rather large concentrations partly due to its participation in the formation of a char structure. In polyolefin polymers it has been shown that melamine and PER act as synergists to APP.

Other intumescent systems have been applied in polymers such as expandable graphite and silica based and metal hydroxide compounds, some of them incorporated as nanocomposites. Recent research describes extended nanoparticles of clay as promising char-forming fillers for good fire protection. These applications are however still on a research level and wait to become commercial. (Kashiwagi et al. 2005).

Metal complexes such as zinc-molybdenum compounds together with phosphate esters and ATH have been used to effectively increase the char formation for PVC.

Whatever the detailed mechanisms for intumescents systems are, the formation of a thick char layer, high carbon concentration, high viscosity of pyrolyzing melt and low penetration capability for propagation of heat, makes intumescent systems efficient to reduce flammability and the exposure of fume gases (Swaraj 2001), (Posner 2004).

6.5 Halogenated flame retardants

Several types of halogenated flame retardants, mainly brominated flame retardants, are described in the literature which includes compounds belonging to families of polybrominated diphenylethers (PBDEs), where congeners of PentaBDE are a part, tetrabromobisphenol-A (TBBPA), tribromophenol (TBP) and brominated phthalic anhydride. Such use of flame retardant additives depends mainly on the type of polymer to be applied for flame retardancy. Within this broad class of PBDE chemicals, the use of commercial Octabromodiphenyl ether and commercial Pentabromodiphenyl ether is restricted in all applications in Europe. The use of Decabromodiphenyl ether has limited restrictions in Europe and in two States in the United States.

7. Historic, present and future consumption of alternative flame retardants to PentaBDE

In general it is very hard to forecast the international market for flame retardants since there are so many market driving forces involved, such as environmental, health and safety regulations, consumer awareness etc., that has a tendency to change rapidly over a limited period of time.

This means that the assumptions made in this chapter are either conservative, meaning that there is a linear approach to the development of the present flame retardants markets, or an innovative approach, meaning that there are incentives to introduce less hazardous flame retardants systems as a result of more stringent regulations and awareness from the public world wide.

Around 90% of the world's production of flame retardants ends up in electronics and plastics, while the remaining 10% ends up in coated fabrics and upholstery furniture and bedding products. In 2004, the production of halogenated flame retardants was equivalent to around 27% of total global production. By then the brominated flame retardants (BFR) constituted around 21% of the total production and use of flame retardants world wide. A further consideration is that the market for plastic in electronic enclosures is growing at around 5% per year (SRI Consulting 2005). With a conservative approach the use of BFRs would then grow by around 63% over a 10 year period.

Table 6 Global consumption of flame retardants and their geographical distribution (SRI Consulting 2005).

Category	United States	Europe	Japan	Other Asia	Total volume [1000 metric tonnes]	Value [million USD]
Aluminium hydroxide	315	235	47	48	645	424
Organo phosphorous FRs	65	95	30	14	205	645
Brominated FRs	66	56	50	139	311	930
Antimony trioxide	33	22	17	44	115	523
Chlorinated FRs	33	35	5	10	82	146
Other FRs	51	47	11	14	123	197
TOTAL	564	489	160	269	1481	2865

Comment [RCC15]: This statement questioning the use of all brominated flame retardants on the basis of their "potentially harmful environmental and health characterisitics" should be removed for two reasons: 1) This statement is a prejudicial and is inconsistent with conclusions of Risk Assessments and other independent scientific evaluations of several major Brominated Flame Retardants in use today. And 2) It is of no value to the reader. This same statement could be said of any group of chemical compounds. Virtually every chemical substance has the capacity to cause harm to health or the environment if uses and exposures are not properly managed.

Deleted: Although the use of brominated flame retardants is still growing by around 5% per year, their use is strongly questioned due to their potentially harmful environmental and health characteristics. A number of brominated flame retardants are already restricted in several countries worldwide. Due to further restrictions and public concern against health and environmentally hazardous chemicals, brominated flame retardants have no future.

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The conservative estimated growth of around 63% in demand of BFRs till 2017 will lead to a total demand for BFRs round 500-600.000 metric tonnes per year, primilary as the predominant brominated flame retardant substances on the international market today, namely decabromodiphenylether and TBBP-A⁷.

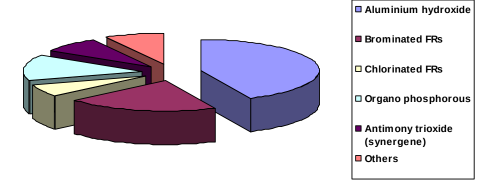


Figure 2 The global market share of groups of flame retardants (SRI Consulting 2005).

With the innovative approach, there will be an introduction of new and innovative specific formulations of less hazardous flame retardant systems that are (or will be) feasible from a commercial and technical point of view, i.e. intumescent systems. These and other less harmful commercial flame retardant systems will balance a conservative market growth of halogenated flame retardants consumption over the coming years.

8. Health and environmental properties of alternative flame retardants to PentaBDE

Since there is a lack of data on health and environmental properties, it is not always possible to perform a comprehensive comparison of all known flame retardant systems described in literature in general and in this report in particular. However, in order to evaluate the toxicity and ecotoxicity of potential alternatives to PentaBDE, the ranking system in table 7 below can be applied. The intention in this report is to present an overall picture of the hazard characteristics as comparable as possible as done in table 7. The system has been used by some US authorities in their recent surveys on alternatives to PBDEs (Illinois EPA 2007), (US EPA 2004).

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⁷ Kirschner M, personal communication (2008)

Table 7 Health and environmental properties of a range of alternative flame retardants to C-PentaBDE.

Chemical	Toxicological properties	Ecotoxicological properties	Comments
Inorganic flame retar	dants and synergists	1.	
Aluminium hydroxide	Low concern	Low concern	Ecotox: Few data available
Magnesium hydroxide	Low concern	Low concern	Tox and ecotox: few data available
Red phosphorous	Non toxic in pure form Low concern	Highly flammable and very toxic to aquatic organisms	May form toxic phosphine gas during combustion in combination with moisture
Ammonium poly phosphate	Insufficient data for assessments	Insufficient data for assessments	May be slightly irritating to skin
Zinc borate	High concern on zinc toxicity	High acute aquatic toxicity	Limited tox and ecotox data available
Boron compounds other than zinc borate, (Borax and disodium tetra borate)	Moderate concern due to 2- generation reproductive/developmental effects	Limited data available	Limited tox and ecotox data available
Antimony trioxide	Ranked as possible carcinogen by IARC ⁸ and EU	Low concern	May produce toxic or irritating vapours during combustion conditions
Zinc hydroxystannate & Zinc stannate	Low concern	Low concern	Very low acute toxicity. Very low aqueous solubility
Organophosphorous	flame-retardants	•	
Triethylphosphate	No data available	No data available	
Aryl phosphates	Low concern	A few compounds show high acute aquatic toxicity	
Halogen containing phosphorous compounds	A few compounds show moderate reproductive toxic properties	A few compounds show moderate or high persistence and acute aquatic toxicity	
Tris (2-chloro-1- methylethyl) phosphate (TCPP or TMCP)	Concern	Low concern	Subject to risk assessment in the EU under the 4 th Priority List Will be transferred to REACH
Reactive phosphorous	No data available	No data available	
Nitrogen based organ	ic Hame-retardants		
Melamine	Low concern	Low concern	Allergic dermatitis has been reported among workers

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 $^{^{8}}$ IARC – International Agency for Research on Cancer

There are toxicological and ecotoxicological data gaps for the potential alternatives to PentaBDE, but the data available clearly show that there are commercially available alternative flame retardants that are less hazardous than C-PentaBDE.

9. Example of costs related to substitution of C-PentaBCD in flexible PUR foam

As mentioned earlier, chloro alkyl phosphates are effective and frequently used as flame retardants in flexible polyurethane (PUR) foams as alternatives to C-PentaBDE in combination with organophosphorous substances. Not only the technical and environmental properties are important for feasibility of flame retardant systems, but also that they are commercially available and cost efficient. Table 8 illustrates an example of a market cost comparison for flame retarded flexible PUR-foam that contain C-PentaBDE in combination with organophosphorous substances and another flexible PUR foam that contain tris (2-chloro1-methylethyl) phosphate (TCPP).

Table 8 Comparison of flame retarded PUR- flexible foam⁹

Application	Content of FRs	Cost of flame	Comments
		retarded PUR	
		per kg	
Flexible	10% PentaBDE in addition to approx	Approx	Price for PentaBDE was set to 6
PUR foam	2% inexpensive organophosphorous	0,70 € per	€ per kg, by 2005 when it was
	substances	kg PUR	phased out in EU
Flexible	20% TCPP	Approx	Present price of TCPP is
PUR foam		0,35 € per	1,80 € per kg
		kg PUR	

This example in table 8 show that flexible PUR foam that contain TCPP is more cost efficient than the use of C-PentaBDE together with inexpensive organophosphorous substances.

10. Conclusion

The objective of this report has been to review possible alternatives to PentaBDE. The available data illustrate that there are alternative flame retardants commercially available which at present appear to be less hazardous than C-PentaBDE. It should be the overall target to replace harmful substances with safer options, but it is also important to point out that the alternative flame retardants presented need to be evaluated based on their range of application. A case by case assessment will be necessary to find the best suitable alternative for a specific use. The data presented in this report are just suggestive and not conclusive, and it is crucial to search for further health and environmental data to get a better understanding of toxicological and ecotoxicological effects of the alternatives presented.

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Comment [spencerv16]:

As the author notes in the conclusion, and demonstrates in table 7, there is little available data on the toxicity and ecotoxicity of the alternative flame retardants. With this in mind it is inconsistent in the context of these guidelines to conclude that the alternatives are definitively less hazardous than C-PentaBDE.

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⁹ Beard A, Clariant, Personal communication (2008)

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Since there should be no current production of C-PentaBDE in Europe, Japan, Canada, Australia and the US, remaining production would be located in other parts of the world. The bromine industry has representation worldwide beside Europe, Japan and North America

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no information was provided of any production or use of C-PentaBDE