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# Addition of X-ray fluorescent tracers into polymers, new technology for automatic sorting of plastics: Proposal for selecting some relevant tracers

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## ABSTRACT

A description of a new technology for automatic sorting of plastics, based on X-ray fluorescence detection of tracers, added in such materials is presented. This study describes the criteria for the selection of tracers, and concluded that the most adapted for XRF are some rare earth oxides. The plastics chosen for tracing and identification are the ones contained in ELV and WEEE from which discrimination is difficult for the existing sorting techniques due to their black colour.

### Keywords:

Sorting  
Recycling  
Polyolefin  
Rare earth oxides  
Tracer  
X-ray fluorescence

## 1. Introduction

The substantial benefits of plastics in terms of low weight, durability, and low cost together with their properties to be used at a wide range of temperatures, be chemical and light resistant as well as to be easily workable as a hot melt helps explains that since 1950, their production has increased an average of almost 10% per year on a global basis and that the annual worldwide demand has grown to 225 million tonnes in 2004 (Andrady and Neal, 2009; APME, 2006).

In the European context, according to PlasticsEurope (APME, 2006) the thermoplastic and thermosetting demand by converters was 47 million tonnes in 2004. The major countries of plastic production are Germany, Italy, France, UK (United Kingdom) and Spain, which together account for around 70% of all European conversion.

Table 1 presents a breakdown of plastic demand and post-consumer waste in Germany, Italy, France, UK and Spain during the year 2004 (APME, 2005). Of the plastics consumed, 37% was used for packaging, 15% for durable consumer applications such as electronics goods and vehicles and the remainder for building/construction applications, domestic/household use, agriculture, etc. Post-consumer plastic waste generation across these five countries was 14.6 million tonnes in 2004, for a demand of

33.1 million tonnes. Table 1 confirms that the main source of plastic wastes has its origin in the field of packaging. However, it is clear that post-consumer waste from electronic and electrical equipment (WEEE) and end-of-life vehicles (ELV) are becoming significant sources with approximately 10% of plastic wastes.

Fig. 1 summarizes the recovery and disposal for post-consumer plastic waste by sector in France, Spain, Italy, UK and Germany in 2004 (APME, 2005). More than 50% of post-consumer plastic waste is sent to landfill for disposal. The automotive and electrical industries are the worst performers with less than 10% of plastic wastes recycled. Moreover, the energetic recovery of plastic wastes in these sectors is achieved through easy options such as incineration, which may cause the emission of harmful gases together with generation of toxic fly and bottom ash that contain lead and cadmium (Dodbiba and Fujita, 2004; Patel et al., 1998; Curlee and Das, 1991).

For giving a real policy of raw resource management based on the recycling of end-of-life products, the European Commission has set up two Directives, 2000/53/CE and 2002/96/CE, dealing with materials of ELV and WEEE, respectively. The objective for the ELV is that from January 1st 2015, the reuse and recovery rate shall be increased to a minimum of 95% of average weight per vehicle and year, whereas the recovery quotas for 10 WEEE categories were fixed to a range of 70–80% for the end of 2006.

The recycling of polymer materials coming from ELV and WEEE is difficult because large numbers of polymer types, grades and blends are used. In addition, the presence of additives can produce significant changes to the mechanical and thermal properties of the polymers. Consequently, the separation and automatic sorting

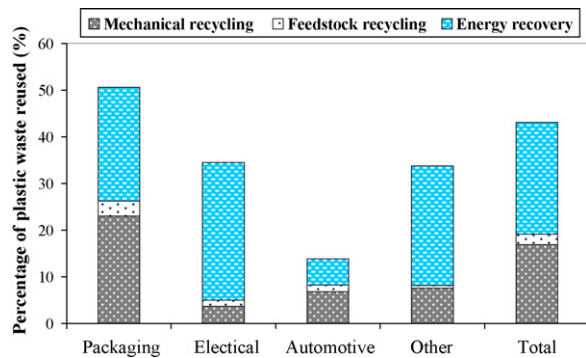
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**Table 1**

Plastic demand and post-consumer waste by sector in France, Spain, Italy, UK and Germany in 2004.

	Demand		Waste	
	ktonnes	%	ktonnes	%
Packaging	12,250	37.0	8970	61.3
Electrical and electronics	2480	7.5	600	4.1
Automotive	2480	7.5	820	5.6
Other	15,890	48.0	4240	29.0
Total	33,100		14,630	



**Fig. 1.** Recovery and disposal of post-consumer plastic waste by sector in France, Spain, Italy, UK and Germany in 2004 (APME, 2005).

of polymers as part of a recycling scheme is a major issue. Moreover, for today's optical sorting technologies it is impossible to discriminate the black plastic parts in cars and electronic equipments.

Table 2 lists the main types of polymers and their consumption in Europe in 2004 for the sectors of automotive and electronic industry. PP together with ABS represents the largest volume of commodity plastics and could be excellent secondary sources of materials.

In this paper, we will review the current technologies for the sorting of plastics and point out their limits, in order to propose a new technology for automatic sorting of plastic wastes through the use of a tracer system, mainly orientated for the ELV and WEEE.

## 2. Sorting and separation technologies for plastics

For an economically efficient recycling of polymer materials, waste plastics need to be sorted cheaply and automatically into individual types and grades due to the various characteristics that each of the different resin types hold. Bruno has separated the automated sorting of plastic wastes into two categories: macrosorting and microsorting. The macrosorting section deals with the sorting of whole bottles or containers whereas the microsorting section follows the sorting of plastics after it has been chopped into pieces. A comparison of the applications of macrosorting and microsorting and their limits are given in Table 3.

**Table 2**

Plastics consumption in automotive industry and electronic and electrical equipment by resin type in Europe in 2004 (APME, 1999, 2001; Maudet-Charbuillet, 2009).

Abbreviation	Name	Automotive		Electrical	
		ktonnes	%	ktonnes	%
PP	Polypropylene	1516	43	635	18
ABS	Acrylonitrile-butadiene-styrene	247	7	1163	33
PE	Polyethylene	282	8	35	1
PS	Polystyrene	-	-	670	19
PVC	Polyvinyl chloride	106	3	141	4
PA	Polyamide	423	12	106	3
PC	Polycarbonate	106	3	141	4
Other	-	846	24	635	18

As it can be seen in Table 3, for the macrosorting systems, optical sorting is limited for colour separation of plastics only; the near infra-red is unsuitable for dark objects whereas middle infra-red can identify them but cannot provide a high-speed identification. The X-ray technology, transmission or fluorescence, is limited to the separation of PVC from PET and the laser induced breakdown spectroscopy is unsuitable for high speed automatic sorting.

Regarding the microsorting techniques, density separation and froth flotation are slow processes which require at least one separation step for each material, and do not provide polymers of high purity. The triboelectric/electrostatic separation could be used for plastics of significantly different dielectric constant, but this technique requires dry and clean plastic surfaces. Moreover, all the sorting techniques mentioned above could not identify different grades of the same polymer.

## 3. Tracing: a new concept for plastic identification and automatic sorting

As shown previously, the existing technologies of sorting do not provide the versatility or flexibility needed for separating the dark plastics into monopolymeric fractions which are the essential prerequisite materials for any efficient recycling process. In the end of 1990s, Simmons et al. (1998) and Ahmad (2000) proposed a new concept on identification of plastics by marking them with a binary combination of fluorescent tracers detectable by UV (ultra-violet) spectroscopy. The use of a tracer system could provide high purity of the sorted materials, separation by polymer grade as well as polymer type, separation by additive system, high speed positive identification and high speed sorting. This project, founded by the European Economic Community, was focused on the sorting of rigid plastics from packaging household waste for demonstrating the concept. They concluded that the speed and purity of sorting were limited by the mechanical singulation inadequacy of the conveyor system at high speed for the clear plastics and that the presence of pigments reduced the fluorescence yield. In the case of black pigments, the reduction was too drastic to allow identification. Aside from these limitations, UV spectroscopy is a surface detection method and this may imply a "clean" surface for tracer identification. The use of "tags" for plastic identification by UV/Vis spectroscopy has also been studied by Corbett et al. (1994). They showed that the addition of phosphor luminescent "tags" to different sort of polymers is viable. However, the detection can be disturbed by the stability of the organic "tags" during the reprocessing of polymers and contaminants, which may be luminescent.

The specialized companies in magnetic sorting, as Eriez (Mankosa and Luttrell, 2005), also proposed a magnetic sorting process of polymers in which a magnetic substance was dispersed. The main advantage of the magnetic detection was the lack of sensitivity with respect to the additives contained in polymers. However, the magnetic tracer system provided only binary separation and required high tracer amount.

**Table 3**

Comparison of the applications of macrosorting and microsorting technologies and their limits.

Sorting technology	Application	Limits
<i>Macrosorting</i>		
Optical (Pascoe, 2003)	Sorting of polymers by colour, removing coloured impurities	Does not identify the polymer, limited for only colour separation
Near infra-red (Alam et al., 1994; Scott and Waterland, 1995)	Bottle sorting	Unsuitable for dark objects
Mid infra-red (Pascoe, 2003)	Technology which can identify dark plastics	Cannot be used for high-speed identification and requires relatively smooth, clean surface
X-ray (Dinger, 1992; Kenny and Bruner, 1994)	Proven and established technology for the identification of PVC	Cannot identify the polymer families since they are composed of the same elements. Only used for the separation of PVC from PET
Laser induced breakdown spectroscopy (Gondal and Siddiqui, 2007; Cuesta et al., 2009)	Identification of PE, PP, ABS.	Unsuitable for high speed automatic sorting, time of analysis: 1-5 s.
<i>Microsorting</i>		
Density separation (Bruno; Altland et al., 1995)	A low cost separation technique used for any material mixture of different densities	Similar densities for some plastics (PE vs. PP, PVC vs. PET), which leads to a low purity of sorted fractions. Slow process, requires at least one separation step for each material
Froth flotation (Shent et al., 1999; Fraunholcz, 2004)	Used for any kinds of polymers. The material is treated with a surfactant for changing the wettability	Selectivity may be difficult to achieve. Requirement of a surfactant to modify the plastics. Separation of only one component at a time
Triboelectric/electrostatic separation (Hearn and Ballard, 2005; Iuga et al., 2005)	Used for plastics of significantly different dielectric constant	Requires a dry and clean plastic surface

**Table 4** summarizes the advantages and drawbacks of some spectroscopic techniques which can be used for detecting a tracer system together with the magnetic detection as reported by Froelich et al. (2007).

From the comparison between the detection techniques of **Table 4**, it comes out that the XRF spectrometry shows potential for detecting a tracer system dispersed into dark polymer materials coming from ELV and WEEE.

XRF spectrometry is a volume, non-destructive elemental analysis. Compared to ultra violet fluorescence, the detection process is not affected by black pigments, and a "clean" surface is not required due to a volume detection of around 1 mm depth. However, as XRF is a spectroscopic method based on elemental analysis, thus the number of tracers is limited by the Periodic Table.

#### 4. The selection of tracers

##### 4.1. The detection system in XRF

For having an efficient sorting, the detection system must be able to reliably identify the tracers, singly or in any combination, at very high speeds and analyse the data generated. To achieve the above requirements, the identification system is based on four components, as shown in Fig. 2:

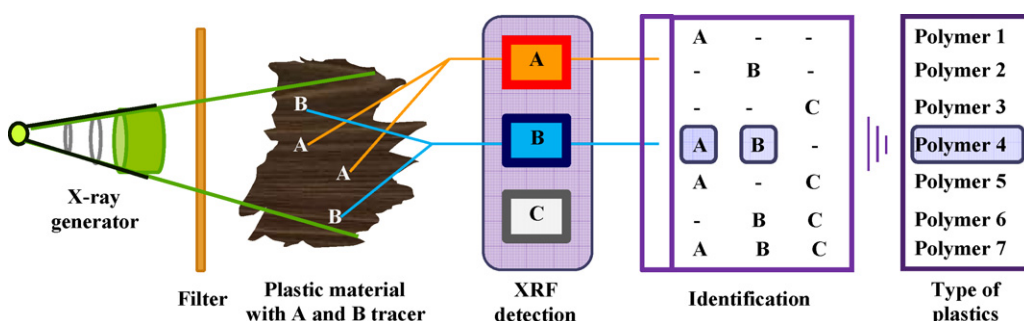
- *Illumination source*: an X-ray generator.
- *Filter system*: a copper filter, for reducing the noise of measurements.

**Table 4**

Comparison of the detection techniques which can be used for detecting a tracer system (Froelich et al., 2007).

Detection technique	Advantages	Drawbacks
UV	High speed identification. Tracer concentration: 1–10 ppm	High quantities of tracer for dark plastics. Fluorescence of the polymer matrix.
Near infra-red	High speed identification	Surface detection
Mid infra-red	Can identify the tracers in dark plastics	Unsuitable for the identification of tracers in dark plastics
XRF (X-ray fluorescence)	Can identify the tracers in dark plastics. High speed identification. Volume detection	Unsuitable for high-speed identification The number of tracers limited by the Periodic Table
Neutron activation	Can identify the tracers in dark plastics	Case study for biological applications
Magnetic detection	Lack of sensitivity with the additives contained in plastics. Proven and established technology	Only binary discrimination. Requires an elevated quantity of tracers

- *XRF detection*: silicon, high purity germanium or cadmium tellurium detectors, proceeding to 200,000–1,000,000 counts/s.
- *Data processing electronics*: processed the output from the detectors for identifying the tracers.

**Fig. 2.** X-ray fluorescent tracer detection system.

The excitation of the tracers is achieved through the use of an X-ray generator and the detection by X-ray fluorescence spectrometry. Each tracer emits a unique radiation in XRF, which depends on the atomic number of the element. The detection system, coupled with a data processing system, detects the emitted radiation and identifies the signature of the tracer, and thus the nature of the polymer matrix. The tracer concentration must be in the range of [100–1000 ppm] in order to not affect the properties of the polymer matrix and to be cost effectiveness. To achieve an automatic sorting of plastics by grade and type, the tracers might be used in a matrix, such that each combination corresponds to a specific type of polymer. For example, in Fig. 2, by using only 3 tracers, it is possible to identify  $7(2^3 - 1)$  variations of different plastics. The detection of tracers will be achieved by EDXRF (energy dispersive X-ray fluorescence).

#### 4.2. Selection of elements composing the tracers

The selection of tracers adapted for the XRF spectrometry detection system is based on a two steps process. As XRF is an elemental analysis, the first step is to choose the more adapted elements from the Periodic Table. Once these selected, the second step is to find the chemical form of the elements which will be conformed to a number of key criteria based on the material properties and tracer identification, summarized as follows:

- *High X-ray fluorescent yield*: for an intense signal and distinguished peaks.
- *Toxicity and radioactivity*: elimination of toxic and radioactive elements.
- *Element availability and physical state*: elimination of elements having poor reserves and of elements which cannot be added as solids in the polymer matrix.
- *Singularity of tracer signal*: elimination of elements contained in polymer additives, such as “natural tracers”.

##### 4.2.1. High X-ray fluorescent yield

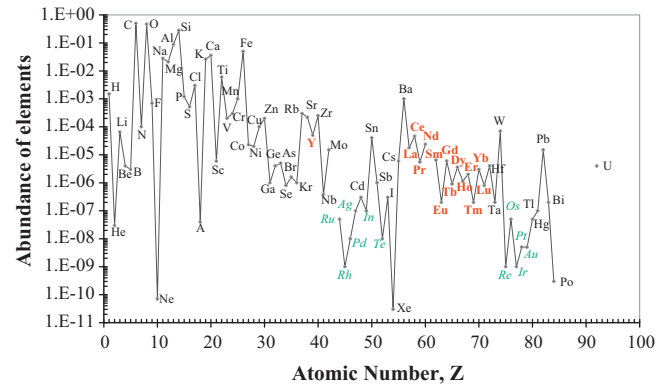
With the aim of having visible and well distinguishable peaks from the background sample, the intensity of tracers has to be maximalist and their fluorescent transmissions should lie in the energy range of [10–60 keV], where the detectors performed the best. The fluorescent peak should preferably be narrow, for reducing the background noise, and, furthermore, they should differ from other selected tracers by at least 0.5 keV, (resolution of commercial detectors) for ensuring adequate and reliable discrimination between tracers.

The tracer identification will be carried out by detecting their K energy lines in order to operate at energies in which the detectors have high efficiency. Therefore, elements emitting at energies lower than 10 keV will be dismiss, since these energies are more likely absorbed by air, plastic dust coverings, thin metal foils, which may be present in the construction of an “industrial” sorting environment. Consequently, by applying these criteria, the elements having an atomic number lower than 30 ( $ZnK_{\alpha 1} = 9.67$  keV) could not be chosen as tracers.

##### 4.2.2. Toxicity and radioactivity

The elements which can be selected as tracers must display very low toxicity and radioactivity due to regulations in processing and use. Toxic elements such as Gallium (Ga), Arsenic (As), Selenium (Se), Cadmium (Cd), Tin (Sn), Antimony (Sb), Mercury (Hg), Thallium (Tl), Lead (Pb) and Bismuth (Bi) will be rejected for a potential XRF tracer application (Lauwers et al., 2007).

Furthermore, radioactive elements such as Curium (Cm), Americium (Am), Plutonium (Pu), Uranium (U), Thorium (Th), Radium (Ra), Polonium (Po) and Actinium (Ac) or synthetic ones such as



**Fig. 3.** Abundance in weight percentage of elements in Earth's upper continental crust as a function of atomic number. The rare earth elements are labelled in red bold whereas the 11 rarest “metals” are labelled in green italic (Pannetier, 1985).

Americium (Am), Berkelium (Bk), Bohrium (Bh), Curium (Cm), Californium (Cf), Darmstadtium (Ds), Dubnium (Db), Einsteinium (Es), Fermium (Fm), Hassium (Hs), Lawrencium (Lr), Mendelevium (Md), Meitnerium (Mt), Neptunium (Np), Nobelium (No), Rutherfordium (Rf), Seaborgium (Sg) and Technetium (Tc) will also be eliminated.

##### 4.2.3. Availability and material behaviour

Elements having poor reserves (Fig. 3), and which are expensive such as Ruthenium (Ru), Rhodium (Rh), Palladium (Pd), Silver (Ag), Indium (In), Tellurium (Te), Osmium (Os), Iridium (Ir), Platinum (Pt), Gold (Au), Protactinium (Pa), Rhenium (Re) and Astatine (At) will be rejected as potential tracers for XRF application.

With respect to the material behaviour the elements of the noble gases group such as Helium (He), Neon (Ne), Argon (Ar), Krypton (Kr), Xenon (Xe) and Radon (Rn) will be eliminated since their standard state is gas.

##### 4.2.4. Singularity of tracer signal

In order to avoid confusion caused by the emissions from the elements composing the additives incorporated within the polymer matrix, it is necessary to identify these elements, by considering them as “natural tracers”, for ensuring an adequate and reliable discrimination between tracers. Table 5 summarizes the main additives found in plastic materials.

The elements composing the additives will be eliminated as “natural tracers” and can probably be used for identifying the prop-

**Table 5**

The main additives used in plastics according to Murphy's ‘Additives for Plastics Handbook’ (Murphy, 1996).

Additives	Specifications of additives
Lubricants	Ca, Zn, Li, Ba, Al, Pb stearates, organic compounds
Plasticizers	Chlorinated, phosphor and amino organic polymers
Release agents	Lubricants, alcohols, metallic stearates (Ca, Zn, Li, Ba, Al, Pb, Mg, Cd, Na)
Fillers and reinforcements	Glass fibers, carbon fibers, talc, mica, wollastonite, barite, calcium carbonate, Pb, Mn, Ba, Ti, Fe, Co, Ni oxides
Colourants	(75% inorganic compounds): TiO <sub>2</sub> (white), Pb (coloured or white), iron oxides (yellow, red, black-brown), Cr (yellow), Mo (orange), Cd (red), Co (blue), Zn (white), aluminium (green), carbon black and organic pigments
Anti-foaming agents	Insoluble oils, silicones, alcohols, stearates and glycols
Stabilizers	Salts of Ba, Ca, Sr, Cd, Zn, ZnO
Antioxidants	S-organic, P-organic, carbon black, organic compounds
Flame retardants	Chlorinated paraffin, Sb <sub>2</sub> O <sub>3</sub> , P/Br-copolymers

erties that they add in the polymer matrix. These elements are the following one: Fluor (F), Magnesium (Mg), Aluminium (Al), Phosphor (P), Chlorine (Cl), Calcium (Ca), Titanium (Ti), Chrome (Cr), Manganese (Mn), Iron (Fe), Cobalt (Co), Nickel (Ni), Zinc (Zn), Strontium (Sr), Bromine (Br), Cadmium (Cd), Tin (Sn), Antimony (Sb), Barium (Ba), Mercury (Hg) and Lead (Pb).

#### 4.2.5. The potential elements for XRF tracer application

By applying the key criteria defined above the list of potential elements is the following one:

- **Rare earth elements:** Yttrium (Y), Lanthanum (La), Cerium (Ce), Praseodymium (Pr), Neodymium (Nd), Samarium (Sm), Europium (Eu), Gadolinium (Gd), Terbium (Tb), Dysprosium (Dy), Holmium (Ho), Erbium (Er), Thulium (Tm), Ytterbium (Yb).
- **Solid metals:** Germanium (Ge), Zirconium (Zr), Niobium (Nb), Molybdenum (Mo), Iodine (I), Lutetium (Lu), Hafnium (Hf), Tantalum (Ta) and Tungsten (W).

#### 4.3. Selection of tracers – rare earth oxides

The selection process developed above has shown that rare earth elements are the most adapted to be used as tracers in XRF. For these elements, it is convenient to use as tracers their most stable chemical form, the oxide ([Hussein, 1996; Sastry et al., 1966](#)). In order to confirm their selection, we have first overviewed their application and availability. Secondly, we have considered their use phase as tracers by analysing their toxicity and stability. Finally, we have evaluated their end-of-life phase by studying what it will be the danger of such particles in the plastic wastes during the incineration or disposal.

##### 4.3.1. Rare earth elements: application and availability

In the contrary of what their name indicates rare earth elements are not so rare elements. Their content in the earth's upper crust is estimated to 0.02% and they have more important reserves than the copper and the lead ([Fig. 3](#)). The three main minerals from which rare earth oxides are extracted are the monazite, bastnaesite and xenotime ([Leveque and Maestro, 2005; Gupta and Bose, 1989](#)).

Since their discovery, rare earth elements have found many applications in various fields of materials science, thanks to their

**Table 6**  
World reserves of rare earth ores ([USGS, 2010a,b](#)).

Country	Mine production in 2009		Reserves	
	tonnes	wt%	ktonnes	wt%
China	129,000	97	36,000	36.4
Commonwealth of Independent States	Not available	Not available	19,000	19.2
United States	–	–	13,000	13.1
Australia	–	–	5400	5.5
India	2755	2	3100	3.1
Brazil	665	0.5	48	0.05
Others	580	0.5	22,500	22.7
World total (rounded)	133,000		100,000	

in United States, 5.5% in Australia and 3.1% in India; the remaining reserves can be found in places such as Canada, Brazil, Malaysia and Africa.

Because of its relative abundance of rare earth containing minerals, China has been very active in 2009 and has developed a major rare earth element industry, by eclipsing all other countries in production of both ore and refined products, by producing almost the totality (97 wt%) of rare earth minerals. It can also be seen in [Table 6](#) that countries with important mine ore reserves such as United States and Australia, have no production since 2009; for the reason that the prices proposed by the Chinese market are very low, due to much lower labour and regulatory costs ([USGS, 2005](#)). This fact shows also the importance of integrating other countries than China in the rare earth mine production, mostly due to the total dependence on a unique international supplier.

As can be seen in [Fig. 3](#) and [Table 7](#), there are differences in the abundances of individual rare earth oxides. First, the even atomic number rare earth elements ( $^{58}\text{Ce}$ ,  $^{60}\text{Nd}$ ,  $^{62}\text{Sm}$ , ...) have greater abundances than the odd ones ( $^{57}\text{La}$ ,  $^{59}\text{Pr}$ ,  $^{63}\text{Eu}$ , ...). Second, the lighter rare earth elements are more incompatible and therefore more strongly concentrated in the continental crust than the heavier ones ([USGS, 2005](#)).

If it is considered that the objective is to trace 1000 ktonnes of plastics per year in Europe for WEEE and ELV, at a tracer concentration of about 100 ppm, this means that the quantity of each rare earth oxide used as tracer in binary combination, will be approximately 400 tonnes.

$$\frac{1000 \text{ (ktonnes\_plastics)} \times 3 \text{ (tracers)} \times 4 \text{ (repeatability)}}{3 \text{ (number\_tracers)}} \times 0.1 \text{ g(tracer)/kg(plastics)} = 400 \text{ tonnes (tracers/year)}$$

For reserve and production data presented in [Table 7](#), it was considered that the distribution of rare earth elements (except Yttrium) is similar to the one of the Earth's crust. For the yttrium oxide, the values were obtained according to the U.S. Geological Survey data.

Based on [Table 7](#), the 400 tonnes tracer demand of rare earth oxides does not seem to be a problem with respect to reserves and production of the light rare earth elements such as Y, La, Ce, Pr, Nd and Sm, since they will present 1–6% of the total production of 2009. For the even atomic number rare earth elements, such as Gd, Dy, Er and Yb, the availability does not seem to be a problem. However, regarding the tracer demand compared to the production of 2009 (6–19%), this could be a potential issue causing a bottleneck for the market distribution.

The use of rare earth elements in automotive pollution control catalysts, permanent magnets, and rechargeable batteries are expected to continue to increase as future demand for conventional and hybrid automobiles, computers, electronics and portable equipment grows. This means that markets are expected to require greater amounts of higher purity mixed and separated products for meeting the demand ([Hedrick, 2007; Maestro and Huguenin, 1995](#)). For these reasons, the availability and the production of rare earth

specific electronic structure, by offering high specificity and unit value. For example, cerium oxide plays an important role in the domain of multifunctional catalysts by reducing the emission level of pollutants such as nitrogen oxide and monocarboxylic oxide ([Nunan et al., 1992; Diwell et al., 1991](#)). Cerium oxide together with erbium oxide is also known to be one of the best polishing agents for glass ([Niinisto, 1987](#)). Thanks to their small, lightweight and high-strength, rare earth element magnets, such as alloys containing Nd, Sm, Gd, Dy, or Pr, have allowed miniaturization of numerous electrical and electronic components used in appliances, computers, automobiles, and military gear. Some of rare earth elements, such as Y, La, Ce, Eu, Gd and Tb, have found applications to the new energy-efficient fluorescent lamps and batteries containing La and Ce are gradually replacing Ni–Cd batteries in computer as well as communications applications and could probably replace lead-acid batteries in automobiles ([USGS, 2005; Falconnet, 1993](#)).

According to the 2010 edition of the U.S. Geological Survey ([USGS](#)) Mineral Commodity Summaries ([USGS, 2010a,b](#)), the world's rare earth reserves are estimated to 100 Mtonnes. As shown in [Table 6](#), 36.4% of the world reserves are located in China, 13.1%

**Table 7**

Rare earth oxides production in 2009, reserves, tracing demand and prices in 2010 given by AMPERE for a purity of 99.9%.

Tracers	Production in 2009 (tonnes)	Reserves (ktonnes)	Tracing demand based on 2009 production (%)	Prices for 1 kg (€)
Y <sub>2</sub> O <sub>3</sub>	8900	540	4	12
La <sub>2</sub> O <sub>3</sub>	18,900	15,300	2	14
CeO <sub>2</sub>	48,500	39,100	1	6
Pr <sub>2</sub> O <sub>3</sub>	5800	4700	7	31
Nd <sub>2</sub> O <sub>3</sub>	25,200	20,400	2	28
Sm <sub>2</sub> O <sub>3</sub>	6800	5500	6	50
Eu <sub>2</sub> O <sub>3</sub>	200	170	200	560
Gd <sub>2</sub> O <sub>3</sub>	6300	5100	6	85
Tb <sub>2</sub> O <sub>3</sub>	1000	770	40	800
Dy <sub>2</sub> O <sub>3</sub>	3800	3100	11	340
Ho <sub>2</sub> O <sub>3</sub>	1200	1000	33	500
Er <sub>2</sub> O <sub>3</sub>	2100	1700	19	30
Tm <sub>2</sub> O <sub>3</sub>	200	170	200	300
Yb <sub>2</sub> O <sub>3</sub>	3200	2500	13	250

elements are very important parameters which have to be taken in consideration for the selection of tracers.

#### 4.3.2. Use phase: toxicity and stability of rare earth oxides

The tracers must have very low toxicity and need to meet the regulations associated with their processing and use. **Table 8** presents the hazard classification of some rare earth oxides. With respect to Annexe I of European Directive 67/548/EC, the majority of rare earth oxides are classified as irritant and their risk code is R36, which means that they may be irritating to eyes. Regarding the median lethal dose (LD<sub>50</sub>), the results show that these substances have a very low toxicity. Moreover, a study done by Hirano and Suzuki (1996) on the exposure, metabolism, and toxicity of rare earths and related compounds showed that these elements are not highly toxic, and their cytotoxicity to macrophages is comparable to that of Cd or silica in vitro. The rare earth oxides, used as tracers are intended for plastics used in automotive and electronic industry, thus their property for being irritating to eyes will not be a problem in this approved applications.

The substances used as tracers must not chemically react with the host materials and cause changes to any of their thermal, mechanical or physical properties beyond acceptable limits. The characteristics of some rare earth oxides are given in **Table 9**. Rare earth oxides are inert particles and thus cannot chemically react with the host polymer material. For most of them, they crystallise to a cubic close packing arrangement with some exceptions such as lanthanum and neodymium oxides which crystallise to a hexagonal one. They can be provided to a micrometric size in order not to act as probable nucleator agents by modifying the thermal properties of the polymer matrix (Xiaomin et al., 1996; Ye et al., 1996; Liu et al., 1993).

**Table 8**

Hazard classification of some rare earth oxides.

Rare earth oxides	CAS number	Hazard symbols <sup>a</sup>	Risk code	LD <sub>50</sub> <sup>d</sup> (mg/kg)
Y <sub>2</sub> O <sub>3</sub>	1314-36-9	Xi <sup>b</sup>	R36 <sup>c</sup>	>5000
La <sub>2</sub> O <sub>3</sub>	1312-81-8	Xi	R36	>5000
CeO <sub>2</sub>	1306-38-3	Xi	R36	>5000
Pr <sub>2</sub> O <sub>3</sub>	12037-29-5	Xi	R36	>5000
Nd <sub>2</sub> O <sub>3</sub>	1313-97-9	Xi	R36	>5000
Gd <sub>2</sub> O <sub>3</sub>	12064-62-9	Xi	R36	>5000
Dy <sub>2</sub> O <sub>3</sub>	1308-87-8	Xi	R36	>5000
Er <sub>2</sub> O <sub>3</sub>	12061-16-4	Xi	R36	>5000
Yb <sub>2</sub> O <sub>3</sub>	1314-37-0	Xi	R36	>5000

<sup>a</sup> Classification with respect to Annexe I of European Directive 67/548/EC.

<sup>b</sup> Irritant.

<sup>c</sup> Irritating to eyes.

<sup>d</sup> Median lethal dose of a toxic substance (LD<sub>50</sub>): dose required to kill half the members of a tested population.

Regarding the effect of tracers in the thermal and mechanical properties as well as in their stability under weathering, the authors have already reported in previous works (Bezati et al., 2011, 2010a,b) that the addition of 1000 ppm of rare earth oxides of micrometric particle size into PP matrix has a minor effect on the mechanical and thermal properties of the traced materials as well as in the photo-degradation of the polymer after UV irradiation exposure.

#### 4.3.3. End-of-life phase: rare earth element risks during incineration or disposal

Even if the best scenario for the traced plastics is to consider that hundred percent will be recovered, although, one possibility could be that part of the post-consumer traced plastic wastes are sent to landfill for disposal or to incineration for energy recovery. Therefore it is important to analyse the life cycle of the traced plastics and obtain more knowledge in the thermal decomposition of rare earth oxides and their potential risk to soils.

Diatloff et al. (1995) as well as Wahid et al. (2000) have reported the positive effects of rare earth elements on agricultural production. In particular, La, Ce, Pr and Nd can promote the root growth of coconut, corn and mungbean at a low rate of application. On the contrary, high application of rare earth elements may lead to their scattering and bioaccumulation in the environment and cause environmental pollution (Haley, 1965; Sabbioni et al., 1982).

Zhang et al. (2001) have studied the rare earth element content in various waste ashes and the potential risk to Japanese soils. The results showed that Y, La, Ce, Pr, Nd, Dy, Yb, Ho, Er, Tm and Lu in the waste ash samples were normally distributed, whereas Sc, Sm, Eu, Gd and Tb were not. Ce, with an average of 26 mg/kg, was the most abundant of the whole rare earth elements, followed by La, with an average of 14 mg/kg, whereas the least abundant was Tm with an average of 0.2 mg/kg. They also found that the content in waste ashes followed the sequence of Ce > La = Y > Nd > Sm > Pr > Gd > Dy > Eu > Tb > Er > Yb > Ho > Lu > Tm.

**Table 9**

Characteristics of some rare earth oxides.

Tracers	Mean particle size (μm)	Density (g/cm <sup>3</sup> )	Crystalline form of the oxide	Melting point (°C)
Y <sub>2</sub> O <sub>3</sub>	1–10	2989	Cubic	2415
La <sub>2</sub> O <sub>3</sub>	1–10	5872	Hexagonal	2250
CeO <sub>2</sub>	1–10	6110	Cubic	2600
Nd <sub>2</sub> O <sub>3</sub>	1–10	6453	Hexagonal	2272
Gd <sub>2</sub> O <sub>3</sub>	1–10	7600	Cubic	2340
Dy <sub>2</sub> O <sub>3</sub>	1–10	8161	Cubic	2340
Er <sub>2</sub> O <sub>3</sub>	1–10	8660	Cubic	2355
Yb <sub>2</sub> O <sub>3</sub>	1–10	9200	Cubic	2346

Based on the studies described above, the selection of rare earth oxide as tracers will probably not affect the life cycle of the plastic material during the incineration and disposal. However, additional studies have to be done in order to confirm this conclusion.

## 5. Conclusion

In this paper we have shown that the existing sorting techniques cannot discriminate the polymers in different families and grades and cannot identify the black plastics, which are the main polymer materials used in automotive and electronic industry. Based on this research work, it appears that X-ray fluorescence, used with a tracer system, could be a solution for the high speed identification and automatic sorting of black plastics into different families and grades.

The addition of X-ray fluorescent tracers to polymer materials provides positive and specific versatility, high purity of sorted fractions as well as efficient and effective sorting. The major advantage of XRF compared to other optical techniques, such as UV spectroscopy, is its ability to detect the tracers even if the polymer matrix is black. However, for XRF the number of tracers is limited to the elements of Periodic Table.

This study has concluded that the tracers the more adapted for the XRF detection process are some rare earth oxides such as  $Y_2O_3$ ,  $CeO_2$ ,  $Nd_2O_3$ ,  $Gd_2O_3$ ,  $Dy_2O_3$  and  $Yb_2O_3$ . The selected components are non toxic, non radioactive, with important reserves, and in concentration levels below 1000 ppm do not affect the properties of the polymer matrix.

The results of this work together with recent publications of the authors (Bezati et al., 2011, 2010a,b; Bezati, 2010c) has lead to a four-years project, mainly funded by the French Agency of Research, with the objective of demonstrating the technical feasibility and commercial viability of sorting of plastics containing X-ray fluorescent tracer.

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