

1 **COMPARISON OF COAL/SOLID RECOVERED FUEL (SRF) WITH COAL/REFUSE**
2 **DERIVED FUEL (RDF) IN A FLUIDISED BED REACTOR**

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4 **S.T. Wagland^a, P. Kilgallon^a, R. Coveney^a, A. Garg^b, R. Smith^a, P.J. Longhurst^a, S.J.T.**
5 **Pollard^a and N. Simms^{a*}**

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7 ^aSchool of Applied Sciences, Sustainable Systems Department, Cranfield University, Cranfield,
8 Bedfordshire, UK, MK43 0AL

9

10 ^bCentre for Environmental Science and Engineering (CESE), Indian Institute of Technology
11 Bombay, Powai, Mumbai, INDIA, 400076

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1 **Abstract**

2 An experimental study was undertaken to compare the differences between municipal solid
3 waste (MSW) derived solid recovered fuel (SRF) (complying with CEN standards) and refuse
4 derived fuel (RDF). Both fuels were co-combusted with coal in a 50 kW fluidised bed combustor
5 and the metal emissions were compared. Synthetic SRF was prepared in the laboratory by grinding
6 major constituents of MSW such as paper, plastic, textile and wood. RDF was obtained from a local
7 mechanical treatment plant. Heavy metal emissions in flue gas and ash samples from the (coal +
8 10% SRF) fuel mixture were found to be within the acceptable range and were generally lower than
9 that obtained for coal + 10% RDF fuel mixture. The relative distribution of heavy metals in ash
10 components and the flue gas stream shows the presence of a large fraction (up to 98%) of most of
11 the metals in the ash (except Hg and As). Thermo-gravimetric (TG) analysis of SRF constituents
12 was performed to understand the behaviour of fuel mixtures in the absence and presence of air. The
13 results obtained from the experimental study will enhance the confidence of fuel users towards
14 using MSW-derived SRF as an alternative fuel.

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16 **Key Words:** Fluidized bed combustion; coal; municipal solid waste; solid recovered fuel; refuse
17 derived fuel; metal analysis; thermo-gravimetric analysis

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19 * Corresponding author. Tel.: +44 1234 750111 extn 2954; Fax: +44 1234 754755,

20 E-mail address: n.j.simms@cranfield.ac.uk (N. Simms)

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1 **1. Introduction**

2 Combustion of biomass rich waste derived fuels is currently being promoted worldwide, as
3 the use of such fuels in energy intensive industries can mitigate greenhouse gas (GHG) emissions
4 and can be economical for energy users. The biomass fraction of such fuels is considered 'carbon
5 neutral' and thus these may also bring incentives in the form of renewable energy for end users. The
6 EU Landfill Directive (Council of the European Union, 1999) offers further incentive by prescribing
7 targets to member states for diverting biodegradable municipal solid waste (MSW) from landfill.
8 Failure to achieve these targets may incur a heavy financial penalty on the local authorities (LAs)
9 who fail to meet their own targets set by national government. In the context of the UK, the situation
10 is particularly challenging due to the heavy reliance on landfilling in the past. The most recent data
11 suggest that England still landfilled 54.4% of the total MSW in 2007-08 (Defra, 2008), which is
12 15.5 million tonnes (Defra, 2008).

13 In recent years, several combustion studies have been performed using unsorted or pre-
14 treated MSW (Chang et al., 1998; Ferrer et al., 2005; Garg et al., 2007b; Guo et al., 2001; Kobayashi
15 et al., 2005; Kouvo and Backman, 2003; Piao et al., 2000; Piao et al., 1998). However, there are still
16 concerns surrounding the heterogeneity of waste material, calorific value (CV) and heavy metal
17 emissions. To overcome this ambiguous situation regarding fuel quality, the European Commission
18 has given mandate to CEN Technical Committee CEN/TC 343 "Solid Recovered Fuels" to prepare
19 a document classifying solid recovered fuels (SRF) (European Committee for Standardisation,
20 2006). Recently, SRF co-combustion studies have been performed in a pulverised fuel-fired
21 combustion reactor to observe the fate of mercury and chlorine (Hilber et al., 2007b). Two different
22 qualities of SRF complying with CEN standards were being used as co-fuel. One of these was
23 derived from the high calorific fractions of MSW and the resulting SRF was mixed with commercial
24 waste (25%) to prepare the other SRF. Addition of commercial waste reduces the quantity of

1 chlorine in the resulting product. It was found that chlorine concentrations in the flue gas stream
2 rises with increase in SRF share. However, use of SRF did not change total mercury content
3 significantly. Gaseous elemental mercury was found to be reduced behind the filter, whereas the
4 HgCl₂ concentration increased behind the filter. In another study, Hilber *et al.* (Hilber et al., 2007a)
5 also developed a method to characterise the de-volatilisation process of SRF meeting CEN
6 requirements.

7 In order to prepare a fuel complying with CEN standards, residual or mixed MSW can be
8 treated using mechanical biological treatment (MBT) or extensive mechanical treatment (MT)
9 process. MSW derived fuel is known as solid recovered fuel (SRF) or refuse derived fuel (RDF)
10 depending upon the composition and characteristics of fuel. SRF is considered more homogeneous
11 and less contaminated fuel, and is market-driven due to tighter quality specifications, whereas RDF
12 has an input-driven specification (Juniper Consultancy Services Ltd, 2005). The present study was
13 undertaken to compare the heavy metal emissions resulting from the co-combustion of synthetic
14 SRF and real RDF (derived from MSW) with coal.

15 MBT is a method for treating residual or mixed MSW and in some parts of Europe it is
16 relatively widespread, e.g. Germany, Italy and Austria (Steiner, 2005). In MBT processes, a series
17 of biological and mechanical treatment unit operations are employed to separate out recyclables,
18 compost, biogas and fuel. MSW derived SRF or RDF has paper, plastic, textile and wood as their
19 main constituents. The Ecodeco process (Ecodeco, 2010) is an example of a commercially available
20 MBT process producing SRF, which is currently operating in the UK in East London, England and
21 Dumfries and Galloway, Scotland. This process employs a bio-drying method for partial
22 stabilisation of untreated MSW (Friends of the Earth, 2008). The bio-drying process reduces the
23 moisture in the waste, thus making mechanical separation more efficient. SRF has the potential use
24 as co-fuel in cement kilns, power plants and dedicated incineration facilities (Garg et al., 2009; Garg

1 et al., 2007a). However, it is necessary to address the emissions from the co-combustion of this
2 material prior to adoption in the UK, as limited information is available in the open literature.

3 The aim of this study was to produce synthetic SRF in the laboratory by grinding paper,
4 plastic, wood and textile and perform co-combustion studies with coal in a fluidized bed combustor.
5 The heavy metal emissions obtained from SRF co-combustion studies were compared with those
6 from coal and coal + RDF mixture combustion studies. The MSW derived SRF and RDF proportion
7 in the final fuel mixture was kept at 10% (w/w) as, in practice, the supply of SRF or RDF could not
8 easily exceed 10% of total fuel for coal-fired power stations. The distribution of trace elements in
9 the ash components and flue gas stream was also calculated. The thermal behaviour of the paper,
10 plastic and coal was investigated in the presence and absence of oxygen using thermo-gravimetric
11 analysis (TGA).

12 13 **2. Experimental**

14 15 ***2.1. Fluidized bed combustor***

16 The combustion of the fuel or fuel blends was carried out in a 50 kW capacity pilot scale
17 fluidised bed combustor test facility located in the Centre for Energy and Resource Technology
18 (CERT) at Cranfield University, UK (Figure 1).

19 The fuel was fed continuously into the combustor through a hopper via a drop tube above the
20 surface of fluidized silica sand bed. The combustion was controlled by adjustments to the fuel feed
21 rate and the preheated air flow rate. The flue gas passed into a second (refractory lined) chamber
22 then through a water-cooled heat exchanger assembly. The resulting gas stream was then entered
23 into the cyclone separator, where fly ash was separated from flue gas. After the cyclone, flue gas
24 was sent to an extraction fan before release to atmosphere via a stack. The combustor was operated

1 at below atmospheric pressure to prevent the release of gases into the combustion hall. The
2 combustor was equipped with bottom and fly ash removal devices and thermocouples were installed
3 to measure the fluidised bed temperature at the top of the combustion zone and pre-heater. Fly ash
4 samples were collected from the fly ash silo located under the cyclone assembly. The CO
5 concentration was kept as low as possible (near to zero) to achieve optimum combustion.

6 The fluidised bed combustor was used for relatively short periods with solid fuels allowing a
7 sufficient steady state period; therefore no extraction of bed material was necessary during its
8 operation. The bed was emptied only when cool after an operational run and replaced with fresh
9 sand before the next experimental run.

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11 ***2.2. Sampling, measurement and analytical methods***

12 Gas sampler ports were provided in the combustor for sampling. One port was used for
13 online CO and CO₂ monitoring of the flue gases, while the other one was used for trace metal
14 sampling. The air flow rate was measured by means of rotameters and this was kept at *ca.* 1400
15 l/min. The flue gas samples for metal analysis were collected by three bubbling traps immersed in a
16 cold bath. The first two traps contained 10% nitric acid to dissolve the metals into solution and the
17 third one was a guard of water to protect the pump from the acid. The metals measured in the flue
18 gas samples include Cr, Zn, Cu, Ni, Pb, Fe, Ca, Mg, Na, K, Al, Mn, Cd and Co. Measurements of
19 metal concentrations were made using an Atomic Absorption Spectrophotometer (AAS) (Hitachi
20 Z8100 Polarized Zeeman Spectrophotometer).

21 Bottom ash and fly ash samples were sent to a commercial testing agency (TES Breiby,
22 Staffordshire, UK) for the determination of heavy metals (including Hg).

23 TGA was carried out using a thermogravimetric analyser manufactured by CI Instruments
24 Ltd. (ROBAL with a 5 gram head). Experimental runs were performed at a heating rate of 10 °C/

1 min with a target temperature of 800 °C. Two types of experimental runs were undertaken; N₂ was
2 used for pyrolysis volatilization and air used for the combustion runs. Gas flow rate in both was kept
3 at 20 ml/min. A sample size of 200 mg was used for the experimental runs.

5 **2.3. Fuel samples**

6 The coal used for the combustion experiments was a bituminous coal from Daw Mill mine,
7 UK. For experimental purposes, the coal was sieved to provide a particle size between 5 and 13 mm.
8 Synthetic SRF was prepared by shredding and grinding paper, saw dust, polypropylene,
9 polyethylene, PVC, and textiles. The composition of synthetic SRF sample was similar to that
10 obtained from the Ecodeco process (Cozens, 2004) as this is a commercially operational facility .
11 The synthetic fuel comprised 58% paper, 22% plastic, 15% textile and 5% wood by weight (w/w).
12 Polypropylene (8% w/w), polyethylene (12% w/w) and PVC (2% w/w) were used as representative
13 of plastics, whereas conifer saw dust was used for wood. Polypropylene and PVC were supplied by
14 Aldrich, UK. Polyethylene was in the form of unused carrier bags obtained from a local
15 supermarket. Junk mail and household sheets were used to represent paper and textile, respectively.
16 In addition, 8% (w/w) water was added to the dry mixture.
17 MSW derived RDF was obtained from a local mechanical treatment plant situated in Leicester, UK.
18 The RDF comprised 82% paper and card, 13% plastic, 5% others (wood, textiles and miscellaneous
19 combustibles). RDF was also shredded and homogenised in a blender prior to pelletization.

20 SRF and RDF were used in the form of pellets having a diameter of 18 mm and a length of
21 30 mm in the combustion runs. This was a constraint of the combustion rig.

22 Samples of coal, synthetic SRF components, SRF and RDF were sent to TES Bretby,
23 Staffordshire, UK for proximate, ultimate and ash analyses. Clean silica sand of grade 16/30 (size
24 0.50 mm- 1.00 mm) (supplied by Garside sands, Leighton Buzzard, Bedfordshire, UK) was used as

1 bed material during the combustion trials and was also sent to the same laboratory for the metal and
2 oxide determination. The metals, including Silicon, were determined by solubilising the ash samples
3 by acid dissolution and analysis of the solution by inductively coupled plasma (ICP) spectroscopy.
4 The measured content of metals is then reported in the form of their highest oxide. Results obtained
5 from the analyses are presented in Tables 1 and 2.

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7 **3. Results and discussion**

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9 ***3.1. Synthetic SRF characterisation***

10 The results from proximate and ultimate analyses of the fuels revealed that the net CV of the
11 synthetic SRF was found to be 21.4 MJ/kg (as received basis) in comparison to 13.15 MJ/kg (as
12 received) for RDF. The net CV of SRF was comparable to that of coal (26.75 MJ/kg, as received
13 basis). The moisture and ash contents for RDF were found to be much higher in comparison to the
14 other two fuels. The chemical analyses of the SRF constituents suggested that paper contributed
15 mainly in the ash content of SRF. However, the volatile matter is much higher in SRF (79%) with
16 RDF and coal only containing 46% and 33% volatile matter, respectively. Other major components
17 such as, Cl and Hg in SRF were also found to be much lower quantity than RDF and coal. Higher
18 concentrations of Hg, Zn, Cd, Pb, Cr metals in RDF shows the presence of materials containing
19 hazardous components like batteries, cosmetics and paint (Sharma et al., 1997). Paper and plastics
20 are also sources of Pb, Cu, Zn and Cd. Synthetic SRF contains smaller concentrations of these
21 metals due to the absence of hazardous materials. RDF has the much higher quantities of Na and K
22 in comparison to synthetic SRF and coal. This indicates the presence of salt in the RDF originating
23 from food waste (Ward and Litterick, 2004). According to the CEN classification, the class code of

1 synthetic SRF used in the present study will be designated as (European Committee for
2 Standardisation, 2006):

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4 Class code Net CV 2; Cl 1; Hg 1.

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6 SRF of this specification is suitable for cement kilns, power plants and fluidized bed
7 combustion plants (van Tubergen et al., 2005). However, power plant operators may be more
8 interested in the biomass content (Juniper Consultancy Services Ltd, 2005) as this can increase
9 revenues in the form of Renewable Obligation Certificates (ROCs) (Ofgem, 2009). A material will
10 qualify as biomass if 90% of energy could be derived from the biogenic fraction (Department of
11 Trade and Industry (DTI), 2006). It is anticipated that MSW derived fuel can be made rich in
12 biomass fraction by introducing partially stabilised biodegradable organic material in SRF, as the
13 biogenic content of the SRF used in this study is likely to be significantly lower than 90% due to the
14 plastic content (15%).

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16 ***3.2. Fluidized bed combustor results***

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18 ***3.2.1. Combustion performance***

19 In this study, coal, coal + 10%w/w synthetic SRF and coal + 10%w/w RDF fuels were
20 tested. The feed rate of the mixtures was varied from 6-9 kg/h to maintain the temperature of the
21 combustion unit. The fuel mixture having lower CV and higher moisture content required a higher
22 feed rate. During combustion experiments, the fluidized bed temperature varied between 750 and
23 950 °C. Temperature profiles with time for three fuel/ fuel mixtures revealed that temperature would
24 rarely reach 900 °C for coal and RDF mixture during the whole run (2.5 h), whereas for the coal and

1 SRF mixture, the combustor attained a 950 °C temperature. This can be attributed to the higher
2 moisture content of RDF (~30%) in comparison to synthetic SRF (~3%). Another reason could have
3 been the presence of more volatile matter in SRF (~79%) that leads to a sharp increase in
4 temperature. The fuel having lower moisture content and high volatile matter exhibited a faster
5 increase in temperature, consistent with previous literature (Ferrer et al., 2005). A summary of the
6 process conditions during the combustion trials is provided in Table 3.

7 Bottom ash samples obtained after coal and coal + 10% SRF fuels combustion were tested
8 for carbon content. Samples were heated in a furnace at a temperature of 550 °C for 2 h. No loss in
9 weight of coal bottom ash sample was observed, whereas a slight reduction (around 1%) was
10 obtained in the latter case. These results indicate zero or low levels of carbon containing compounds
11 in the ash.

12 ***3.2.2. Metal concentrations in flue gas samples***

13 Figure 2 shows the concentrations of different metals in flue gases as a result of the
14 combustion of different fuels. Metal emissions from coal combustion were found to be much lower
15 in comparison to the other fuel mixtures except for Cr, Zn and Ni. However, little difference was
16 observed in metal emissions for the (coal +10% RDF) and (coal + 10% SRF) mixtures.

17 The presence of Cu and Zn in the flue gas is indicative of the chloride species of the metals,
18 as reported earlier by Kouvo and Backman (Kouvo and Backman, 2003). Pb concentrations were
19 similar (~ 120 µg/m³) in the flue gas samples obtained from coal and coal + 10% SRF. However,
20 the flue gas sample from coal + 10% RDF combustion exhibited high Pb release (more than twice
21 that obtained with other fuels). This may be due to the higher Pb concentration (~ 70 mg/kg) in
22 RDF. According to a study reported by the Environment Agency in 2001 (Environment Agency,
23 2001), other metal emissions (such as Cr, Cu, Ni, Pb, Co, Mn etc.) must be below 1000 µg/m³ for
24 co-combustion in cement kilns. In the present study, the sum of these metals was 660 µg/m³ for coal

1 and coal + 10% SRF fuels and $830 \mu\text{g}/\text{m}^3$ for coal + 10% RDF. Nevertheless, Cd was higher in the
2 flue gas sample obtained from coal + 10% SRF sample than samples from the other two fuels,
3 though this was well below the prescribed limits [max. $(\text{Cd} + \text{Tl}) < 100 \mu\text{g}/\text{m}^3$] (Environment
4 Agency, 2001), as is evident from Figure 2. Tl was found to be in non-detectable quantities in the
5 fuel samples. In comparison to unsorted MSW combustion (Chang et al., 1998), the coal + SRF
6 mixture showed much lower amounts of Pb and Cd.

7

8 **3.2.3. Metal analysis in ash samples**

9 Metal concentrations were measured in bottom ash and fly ash samples obtained after the
10 combustion of coal, coal + 10% SRF and coal + 10% RDF fuels. Results are shown in Figures 3 and
11 4. The release of different trace elements depends upon the volatility of metals and the fuel feed
12 rate. The more volatile metals will be present in the fly ash samples, and a higher fuel feed rate will
13 result in more fuel being passed through the system, therefore introducing higher quantities of the
14 metals.

15 Among the tested elements, As, Cd, Hg and Pb are considered of principal concern. These metals
16 are the most volatile between 800 and 900 °C temperatures (Miller et al., 2002). From Figure 4, it is
17 indicated that the major fraction of these metals was found in fly ash. Cd and Pb releases were lower
18 in bottom and fly ash samples obtained from (coal + 10% SRF) fuel combustion than the other two
19 fuel/fuel blends. From Figure 3, it is observed that in general the use of RDF as co-fuel increases
20 heavy metals accumulation in bottom ash except Zn, Mn and Hg. A higher metal release with the
21 coal and RDF mixture was expected due the metals content being the highest in this fuel. However,
22 no significant difference in the metal concentrations was observed in fly ash samples obtained as a
23 result of the combustion studies (Figure 4).

24

1 **3.2.4. Relative distribution of metal contents in ash and flue gas stream**

2 The distribution of different metals among bottom ash, fly ash and flue gas was also seen for
3 three combustion scenarios (Figures 5a, b and c). It is evident from the figures that a major part of
4 the trace elements is retained in the ash. Only the significant quantities of Hg and As are released
5 with flue gas into the atmosphere. Combustion studies with sewage sludge have revealed that
6 typically between 75-98% of Cd, Cr, Cu, Pb and Zn are retained in the ash (Cenni et al., 1998;
7 George et al., 2010; Lopes et al., 2003; Werther and Ogada, 1999). It has also been found that most
8 of the mercury is released to the atmosphere via flue gas due to high volatility (Germani, 1988; Lee
9 et al., 2006; Otero-Rey et al., 2003; Reddy et al., 2005; Sandelin and Backman, 2001; Werther and
10 Ogada, 1999). During combustion, generally mercury is found in three forms, namely, particle-
11 bound, gaseous elemental mercury and in oxidized form (Lee et al., 2006). At high temperature in
12 the combustion zone, Hg vaporises into the elemental form. With a temperature reduction, it is
13 oxidized by flue gas components forming HgCl_2 and HgO . A small fraction of Hg can also be
14 condensed on ash particles due to temperature reduction. Otero-Rey *et al.* have reported that Cd, Mn
15 and Pb are readily incorporated in the bottom ash, whereas the larger fractions of As, Cr, Cu and Zn
16 are found in fly ash (Otero-Rey et al., 2003). Our results are consistent with Cu, Zn, As, Mn and Cr
17 in the samples obtained from coal only and (coal + 10% SRF) combustion studies. Cd was found in
18 the fly ash rather than in bottom ash in all three combustion runs; this result is similar to that
19 obtained for coal combustion by Reddy *et al.* Significant quantities of arsenic were found in flue gas
20 stream (ca. 50%, 30% and 30% for coal, (coal + 10% SRF) and (coal + 10% RDF) combustion,
21 respectively). Variation in the behaviour of metals can be attributed to the different fuel properties.
22 The mass of a particular element can also affect the mobility of the metal (Sushil and Batra, 2006).
23 For example, the lighter element such as Cu and Zn can be carried with fly ash. It has also been
24 found that trace metals composition can be varied even from a single power plant. In addition to

1 these aspects, the particle size of feedstock may have an impact on metal fractionation in different
2 streams. The presence of heavy metals in the ash samples is also influenced by the composition of
3 waste and its handling (Chang et al., 1998).

5 **3.3. Thermogravimetric experiments**

6 Thermogravimetric experiments were carried out for the two major components of SRF
7 (paper and various plastics) in an air and nitrogen atmosphere to investigate the thermal behaviour
8 of these components. The results from this analysis are shown in Figures 6a and 6b.

10 **3.3.1. In the presence of air**

11 Figure 6a illustrates the thermal behaviour of paper and plastic fractions (polyethylene,
12 polypropylene and PVC) in the presence of air. In all materials, initial weight loss with an increase
13 in temperature takes place due to moisture loss and volatilization. This stage is followed by gradual
14 or no further weight reduction before volatilisation. Polyethylene and polypropylene profiles
15 exhibited the sharp decrease in weight in the temperature range of 400-500 °C. However, almost no
16 ash was found for polypropylene. An ash content of *ca.* 10% was obtained for polyethylene. For
17 paper, a weight loss of 10% was observed up to 300 °C heating due to the loss of moisture. This was
18 followed by two zones of sharp weight reduction (First zone: 300-375 °C and second zone: 375-500
19 °C). Beyond 500 °C, there was a continuous gradual reduction in weight to *ca.* 30% of the original,
20 which may go further with an increase in temperature due to the presence of inorganic fillers.
21 Results for paper are consistent with those found by other researchers (Guo et al., 2001; Lu et al.,
22 1996; Lu et al., 1999). However, the ash content was found to be much higher in comparison with
23 previous studies. This may be a function of different types of paper. For PVC, weight loss due to
24 moisture was negligible (as for other plastic components). Three zones for weight loss were

1 observed; (a) 300-350 °C, (b) 400-525 °C, and (c) 525-800 °C. It was found that the weight losses
2 in each temperature zone decreased with increasing temperature. A major fraction of weight
3 reduction (*ca.* 65%) was obtained during the first heating zone followed by *ca.* 15% and 8% weight
4 reduction in the second and third zones, respectively. The first weight loss for PVC may be due to
5 the loss of HCl as reported by Lu *et al.* (Lu et al., 1996). The second and third weight loss might be
6 due to the decomposition of the materials and combustion of carbon residue (Lu et al., 1996; Lu et
7 al., 1999).

8

9 **3.3.2. In the presence of nitrogen**

10 The thermal behaviour of paper, polyethylene, polypropylene and PVC was also observed in
11 an inert atmosphere (N₂). TGA curves for different components are shown in Figure 6b. Similar
12 curves were obtained for polypropylene and polyethylene. A single degradation zone was observed
13 for each of these plastics. No char yield was observed for polypropylene, whereas, *ca.* 8% char was
14 formed for polyethylene. The TG curve for paper shows the loss of moisture in the first degradation
15 zone followed by faster weight loss over a temperature range of 300-350 °C in the second thermal
16 degradation zone. Reduction in weight up to 350 °C was 50%. Beyond 350 °C, the gradual decrease
17 in weight loss might be attributed to the thermal degradation of calcium carbonate, which is used as
18 filler in the paper manufacturing process (Cozzani et al., 1995). The TG pattern for PVC reveals a
19 weight reduction of *ca.* 60% in the first thermal degradation zone (in a temperature range of ~ 300-
20 350 °C). This might be due to the elimination of HCl and a small amount of benzene (Ma et al.,
21 2002). In the second thermal degradation zone, *ca.* 20% weight loss was observed, this might be
22 attributed to the pyrolysis of the polyethylene sequences that formed during the first stage (Ma et al.,
23 2002).

24

1 **4. Conclusions**

2 The synthetic SRF (58% paper, 22% plastic, 15% textile and 5% wood) contained less
3 chlorine and mercury than Daw Mill coal and RDF. However, as for RDF, the composition of real
4 SRF is dependent on MSW composition and the type and performance of specific unit operations
5 employed for preparing fuel from waste. The SRF prepared in the present study was suitable for
6 cement works and power sectors, as indicated by its characteristics. The biomass content of the SRF
7 was *ca.* 70%. The pure coal and coal + 10% SRF mixture showed a comparable combustion
8 performance for heavy metals release from flue gas and ash (bottom and fly ash) samples. The coal
9 and 10% RDF mixture produced slightly higher emissions.

10 The relative distribution study confirms the findings of previous studies that most of the Hg
11 releases into the atmosphere with flue gas and other heavy metals remain in ash. Volatility and mass
12 also affect the mobility of trace elements in fly ash and bottom ash samples. A major portion of Cu,
13 Zn, Cd, Pb, Cr and As were found in fly ash, whereas a large fraction of Mn remained in the bottom
14 ash.

15 TG analysis was carried out to study the combustion behaviour of the main components of
16 waste derived fuel. This provides the information on temperature ranges for various reactions taking
17 place during the combustion or pyrolysis of the fuel.

18 The results obtained from this study highlight the potential use of waste derived fuels, and
19 support the findings of an options appraisal of SRF by Garg *et al* (2009). A high calorific value, low
20 chlorine and mercury content, lower metal emissions and the comparatively inexpensive availability
21 of fuel could be attractive to cement kiln operators, power plants and industrial boiler facilities.

22 Future studies should be targeted at measuring gaseous emissions. The share of SRF in the
23 fuel mixture should be increased (up to 20%) and the resulting emissions should be determined. The
24 kinetics of the TG analysis can be studied by performing differential thermal analysis.

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References

Cenni, R., Frandsen, F., Gerhardt, T., Spliethoff, H., Hein, K.R.G., 1998. Study on trace metal partitioning in pulverized combustion of bituminous coal and dry sewage sludge. *Waste Management* 18, 433-444.

Chang, Y.H., Chen, W.C., Chang, N.B., 1998. Comparative evaluation of RDF and MSW incineration. *J. Hazard. Mater.* 58, 33-45.

Council of the European Union, 1999. Directive 1999/31/EC on the Landfill of Waste. *Official Journal of the European Communities* L 182, 1-19.

Cozens, P., 2004. EfW - an alternative vision, in: E.K. Papadimitriou and E.I. Stentiford (Ed.), *Biodegradable and residual waste management: 1st UK Conference and exhibition*. Cal Recovery Europe Ltd, Harrogate, UK, pp. 464-472.

Cozzani, V., Petarca, L., Tognotti, L., 1995. Devolatilization and pyrolysis of refuse derived fuels: characterization and kinetic modelling by a thermogravimetric and calorimetric approach. *Fuel* 74, 903-912.

Defra. 2008. Key Facts about: Waste and Recycling, Municipal waste management, 2007/08, Retrieved 01.05.10, from <http://www.defra.gov.uk/evidence/statistics/environment/waste/kf/wrkf06.htm>.

Department of Trade and Industry (DTI), 2006. Renewable Obligation Order 2006 – Final Decisions.

Ecodeco. 2010. Intelligent transfer stations, Retrieved 20.09.2010, from <http://www.ecodeco.it/gruppo/cms/ecodeco/en/municipal/its/>.

Environment Agency, 2001. Solid waste derived fuels for use in cement & lime kilns - an international perspective. Entec, Bristol, UK, p. 151pp.

European Committee for Standardisation, 2006. CEN/TC 343/WG 2, Solid recovered fuels - specifications and classes.

Ferrer, E., Aho, M., Silvennoinen, J., Nurminen, R.V., 2005. Fluidized bed combustion of refuse-derived fuel in presence of protective coal ash. *Fuel Process. Technol.* 87, 33-44.

Friends of the Earth, 2008. Mechanical biological treatment (MBT) briefing note.

Garg, A., Smith, R., Hill, D., Longhurst, P.J., Pollard, S.J.T., Simms, N.J., 2009. An integrated appraisal of energy recovery options in the United Kingdom using solid recovered fuel derived from municipal solid waste. *Waste Management* 29, 2289-2297.

1 Garg, A., Smith, R., Hill, D., Simms, N., Pollard, S., 2007a. Wastes as Co-Fuels: □ The Policy
2 Framework for Solid Recovered Fuel (SRF) in Europe, with UK Implications. *Environ. Sci.*
3 *Technol.* 41, 4868-4874.

4 Garg, A., Smith, R., Longhurst, P.J., Pollard, S.J.T., Simms, N.J., Hill, D., 2007b. Comparative
5 evaluation of SRF and RDF co-combustion with coal in a fluidised bed reactor, *Proceedings*
6 *Sardinia 2007, Eleventh International Waste Management and Landfill Symposium.* CISA, S.
7 Margherita di Pula, Cagliari, Italy, p. 411.

8 George, A., Larrion, M., Dugwell, D., Fennell, P.S., Kandiyoti, R., 2010. Co-firing of single, binary,
9 and ternary fuel blends: Comparing synergies within trace element partitioning arrived at by
10 thermodynamic equilibrium modeling and experimental measurements. *Energy and Fuels* 24, 2918-
11 2923.

12 Germani, M.S., 1988. Vapor-phase concentrations of arsenic, selenium, bromine, iodine, and
13 mercury in the stack of a coal-fired power plant. *Environ. Sci. Technol.* 22, 1079-1085.

14 Guo, X.F., Yang, X.L., Li, H., Wu, C.Z., Chen, Y., Li, F., Xie, K.C., 2001. Release of hydrogen
15 chloride from combustibles in municipal solid waste. *Environ. Sci. Technol.* 35, 2001-2005.

16 Hilber, T., Martensen, M., Maier, J., Scheffknecht, G., 2007a. A method to characterise the volatile
17 release of solid recovered fuels (SRF). *Fuel* 86, 303-308.

18 Hilber, T., Thorwarth, H., Stack-Lara, V., Schneider, M., Maier, J., Scheffknecht, G., 2007b. Fate of
19 mercury and chlorine during SRF co-combustion. *Fuel* 86, 1935-1946.

20 Juniper Consultancy Services Ltd, 2005. Mechanical biological treatment- a guide for decision
21 makers

22 Kobayashi, N., Itaya, Y., Piao, G., Mori, S., Kondo, M., Hamai, M., Yamaguchi, M., 2005. The
23 behavior of flue gas from RDF combustion in a fluidized bed. *Powder Technol.* 151, 87-95.

24 Kouvo, P., Backman, R., 2003. Estimation of trace element release and accumulation in the sand
25 bed during bubbling fluidised bed co-combustion of biomass, peat, and refuse-derived fuels. *Fuel*
26 82, 741-753.

27 Lee, S.J., Seo, Y.C., Jang, H.N., Park, K.S., Baek, J.I., An, H.S., Song, K.C., 2006. Speciation and
28 mass distribution of mercury in a bituminous coal-fired power plant. *Atmos. Environ.* 40, 2215-
29 2224.

30 Lopes, M.H., Abelha, P., Lapa, N., Oliveira, J.S., Cabrita, I., Gulyurtlu, I., 2003. The behaviour of
31 ashes and heavy metals during the co-combustion of sewage sludges in a fluidised bed. *Waste*
32 *Management* 23, 859-870.

33 Lu, H., Purushothama, S., Hyatt, J., Pan, W.P., Riley, J.T., Lloyd, W.G., Flynn, J., Gill, P., 1996.
34 Co-firing high-sulfur coals with refuse-derived fuel. *Thermochim. Acta* 284, 161-177.

35 Lu, R., Purushothama, S., Yang, X., Hyatt, J., Pan, W.P., Riley, J.T., Lloyd, W.G., 1999.
36 TG/FTIR/MS study of organic compounds evolved during the co-firing of coal and refuse-derived
37 fuels. *Fuel Process. Technol.* 59, 35-50.

38 Ma, S., Lu, J., Gao, J., 2002. Study of the low temperature pyrolysis of PVC. *Energy and Fuels* 16,
39 338-342.

40 Miller, B.B., Kandiyoti, R., Dugwell, D.R., 2002. Trace element emissions from co-combustion of
41 secondary fuels with coal: A comparison of bench-scale experimental data with predictions of a
42 thermodynamic equilibrium model. *Energy and Fuels* 16, 956-963.

43 Ofgem, 2009. Renewables Obligation: fuel measurement and sampling guidance. Ofgem, London.

44 Otero-Rey, J.R., López-Vilariño, J.M., Moreda-Piñeiro, J., Alonso-Rodríguez, E., Muniategui-
45 Lorenzo, S., López-Mahía, P., Prada-Rodríguez, D., 2003. As, Hg, and Se Flue Gas Sampling in a
46 Coal-Fired Power Plant and Their Fate during Coal Combustion. *Environ. Sci. Technol.* 37, 5262-
47 5267.

1 Piao, G., Aono, S., Kondoh, M., Yamazaki, R., Mori, S., 2000. Combustion test of refuse derived
2 fuel in a fluidized bed. *Waste Management* 20, 443-447.

3 Piao, G., Aono, S., Mori, S., Deguchi, S., Fujima, Y., Kondoh, M., Yamaguchi, M., 1998.
4 Combustion of refuse derived fuel in a fluidized bed. *Waste Management* 18, 509-512.

5 Reddy, M.S., Basha, S., Joshi, H.V., Jha, B., 2005. Evaluation of the emission characteristics of
6 trace metals from coal and fuel oil fired power plants and their fate during combustion. *J. Hazard.*
7 *Mater.* 123, 242-249.

8 Sandelin, K., Backman, R., 2001. Trace elements in two pulverized coal-fired power stations.
9 *Environ. Sci. Technol.* 35, 826-834.

10 Sharma, V.K., Caudatelli, M., Fortuna, F., Cornacchia, G., 1997. Processing of urban and agro-
11 industrial residues by aerobic composting: Review. *Energy Convers. Management* 38, 453-478.

12 Steiner, M., 2005. Status of mechanical-biological treatment of residual waste and utilisation of
13 refuse-derived fuels in Europe, Proceedings, The future of residual waste management in Europe.
14 Orbit Association, Luxembourg, p. 7.

15 Sushil, S., Batra, V.S., 2006. Analysis of fly ash heavy metal content and disposal in three thermal
16 power plants in India. *Fuel* 85, 2676-2679.

17 van Tubergen, J., Glorius, T., Waeyenbergh, E., 2005. Classification of solid recovered fuels.
18 Ward, C., Litterick, A., 2004. Assessment of the potential variation of composted materials across
19 the UK, Banbury, UK.

20 Werther, J., Ogada, T., 1999. Sewage sludge combustion. *Prog. Energy and Combustion Sci.* 25, 55-
21 116.

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1 **Table 1 – Composition of simulated SRF Components**

Fuel	Textile	Paper	Polyethylene	Sawdust	PVC	Polypropylene
Proximate and ultimate analyses (as received)						
Moisture (%)	1.5	3.3	0.1	8.2	0.1	0.1
Ash (%)	1.0	15.9	8.7	2.6	0.2	0.2
Volatile Matter (%)	89.1	72.2	91.3	70.6	95.8	99.8
HHV/ Gross CV	20.3	16.7	40.8	19.5	21.8	46.3
LHV/ Net CV (MJ/kg)	19.2	15.8	38.0	18.2	20.7	43.2
C (%)	52.5	33.2	76.2	46.5	38.6	86.6
H (%)	4.9	4.0	12.9	5.3	4.9	14.7
N (%)	0.60	0.04	0.06	0.42	0.17	0.18
S (%)	0.15	0.08	0.10	0.05	0.08	0.21
Cl (%)	0.01	0.02	0.01	0.01	5.97	0.01
Metal analysis (mg/kg_{fuel}, dry basis)						
Pb	1.2	0.4		6.5		
Cu	2.2	10.7		3.4		
Zn	7.1	18.6		40.4		
As	<0.1	<0.1		0.2		
Cd	0.02	0.04		0.21		
Mn	1.6	49.3		640		
Cr	6.5	4.9		3.7		
Na	245	472		193		
K	91	1056		1489		
Tl	<0.1	<0.1		<0.1		
Hg	0.05	0.05		0.04		
Elemental oxide analysis (% , in ash)						
SiO ₂	13.4	9.8		40.4		
Al ₂ O ₃	4.6	5.8		7.5		
Fe ₂ O ₃	1.2	0.4		6.1		
TiO ₂	7.5	0.1		0.4		
CaO	27.5	74		19.8		
MgO	2.2	2.1		3.2		
Na ₂ O	3.3	0.4		1		
K ₂ O	1.1	0.8		6.9		
P ₂ O ₅	22.5	0.1		2.3		

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1 **Table 2 – Composition of fuel samples and silica sand**

Fuel	Coal	SRF	RDF	Silica sand
Proximate and ultimate analyses (as received)				
Moisture (%)	6.2	3.0	30.4	-
Ash (%)	11.4	11.1	16.2	-
Volatile Matter (%)	33	79.6	46.1	-
HHV/ Gross Calorific Value (MJ/kg)	27.6	13.0	14.8	-
LHV/ Net Calorific Value (MJ/kg)	26.5	11.8	13.2	-
C (%)	66.3	40.5	28.1	-
H (%)	4.2	5.3	3.4	-
N (%)	1.2	0.03	0.98	-
S (%)	1.7	0.07	0.32	-
Cl (%)	0.28	0.02	0.25	-
Metal analysis (mg/kg_{fuel}, dry basis)				
Pb	13.8	1.0	71.2	-
Cu	13.6	6.4	66.3	-
Zn	5.6	10.8	1970	-
As	12	0.01	2.1	-
Cd	0.07	0.05	0.30	-
Mn	207	39.6	164	-
Cr	14	4.5	25.2	-
Na	868	247	5362	-
K	1845	92	3454	-
Tl	0.3	<0.1	<0.1	-
Hg	0.3	0.07	0.19	-
Elemental oxide analysis (% , in ash)				
SiO ₂	49	7.5	48.1	95.4
Al ₂ O ₃	22.8	4.3	9.5	0.1
Fe ₂ O ₃	11.5	4.5	2.7	1.0
TiO ₂	1	8.1	1.8	<0.1
CaO	4	60.4	18.5	<0.1
MgO	2.2	1.2	2	<0.1
Na ₂ O	1	0.3	3.3	<0.1
K ₂ O	1.9	0.1	1.9	<0.1
P ₂ O ₅	0.3	0.8	1.5	<0.1

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 2 **Table 3 – Summary of process conditions during combustion trials**

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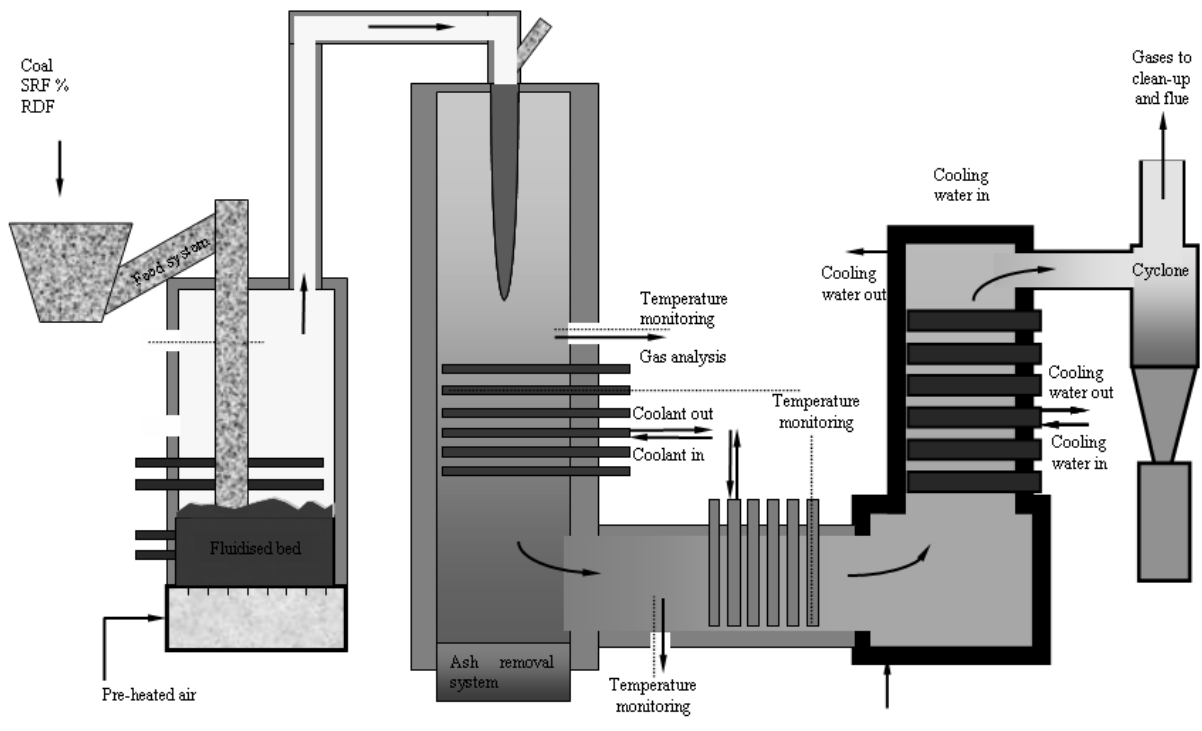
Process parameters	Fuel		
	Coal only	Coal + 10% SRF	Coal + 10% RDF
Fuel feed rate (kg/h)	6.0	7.0	9.0
Duration of the run (h)	2.5	4.0	2.5
Fluidised bed	800 - 900	800 - 950	750 - 900
Temperature (°C)			
Secondary chamber	550 - 600	600 - 700	500 - 600
temperature (°C)			

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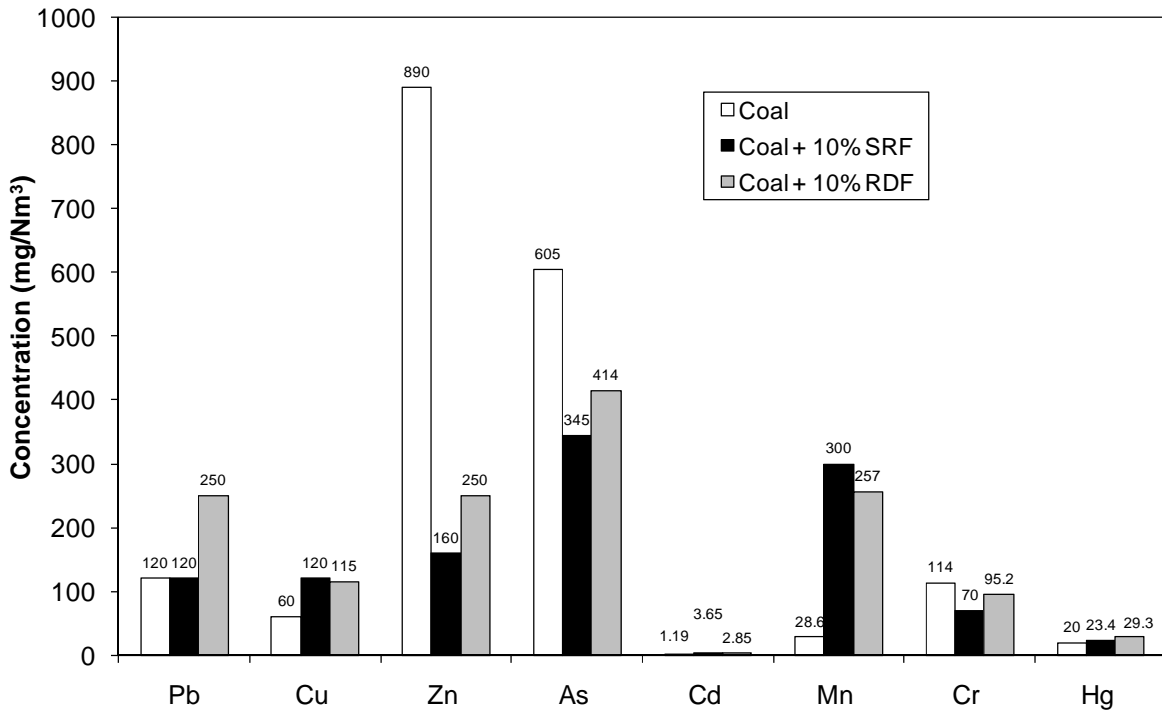


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2 **Figure 1. Schematic diagram of fluidized bed combustor**

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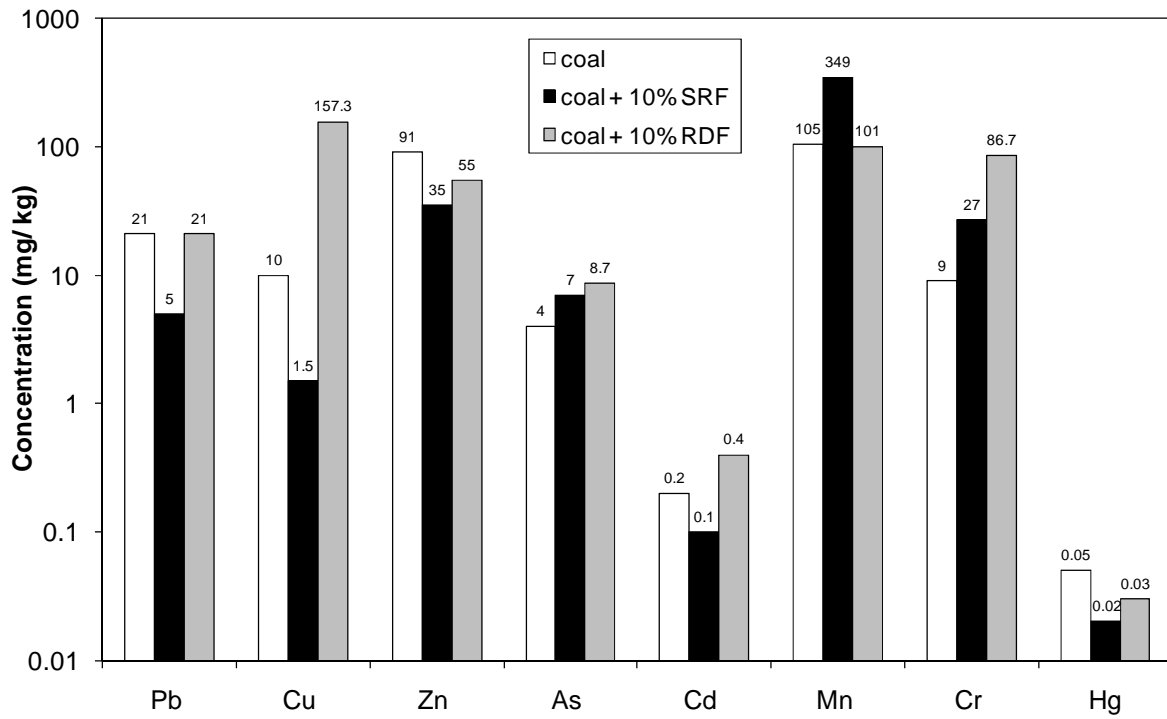


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2 Figure 2. Metals concentration in flue gas samples obtained during the combustion of coal, (coal + 10%

3 SRF) and (coal + 10% RDF) fuels

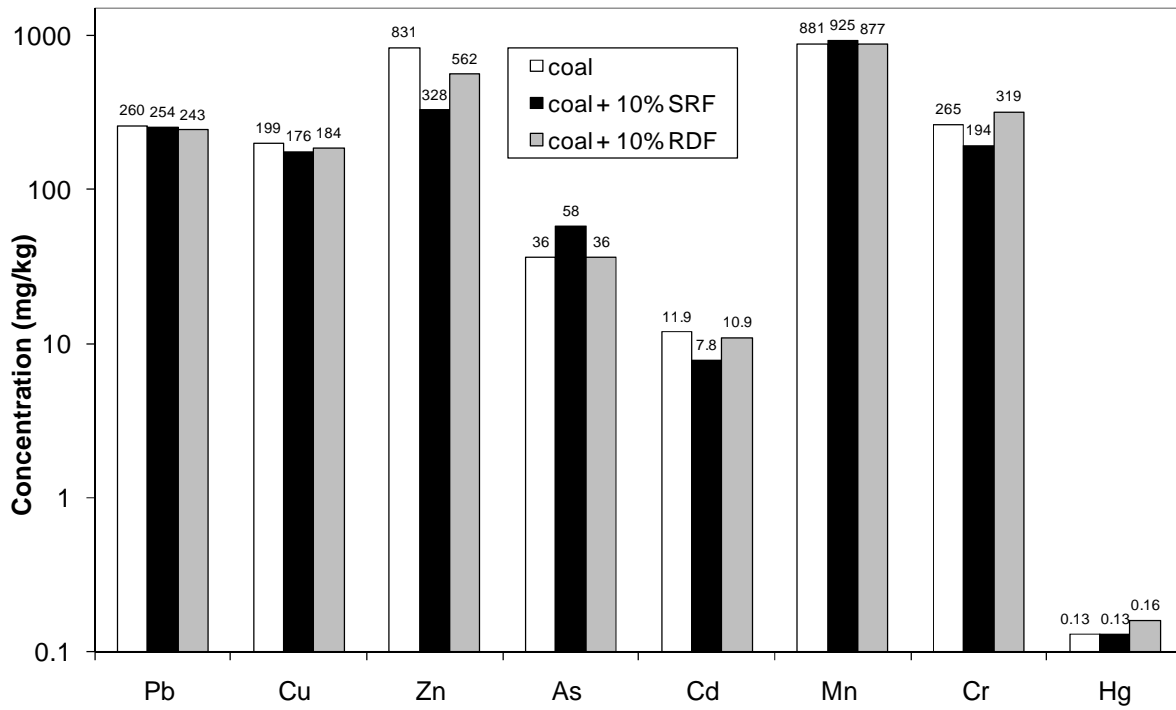
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2 **Figure 3. Metal concentrations in the bottom ash samples obtained after combustion of fuels**

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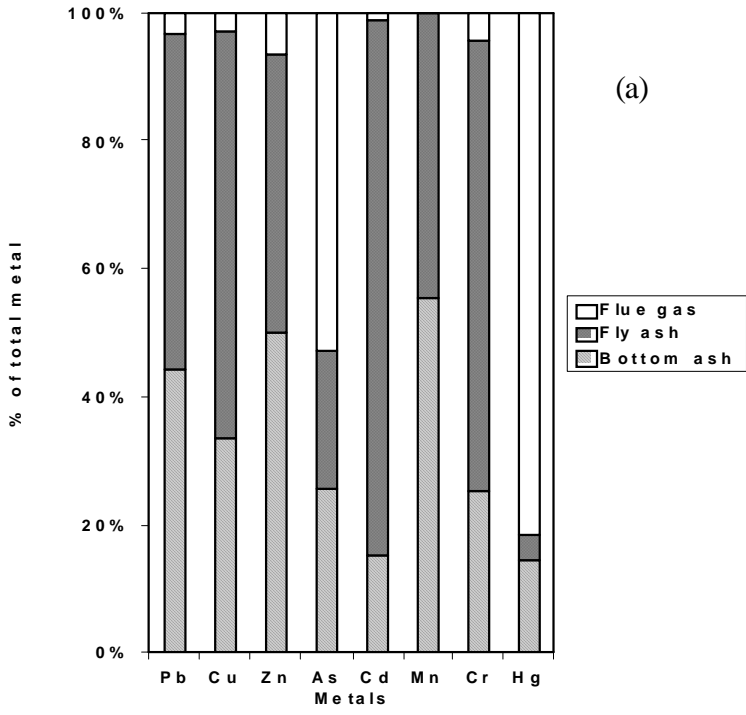


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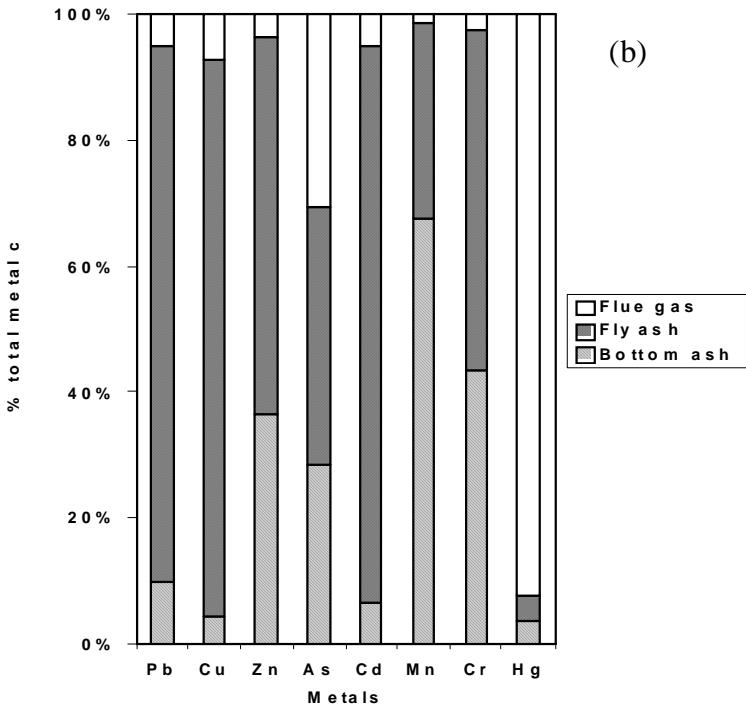
2 **Figure 4. Metal concentrations in the fly ash samples obtained after combustion of fuels**

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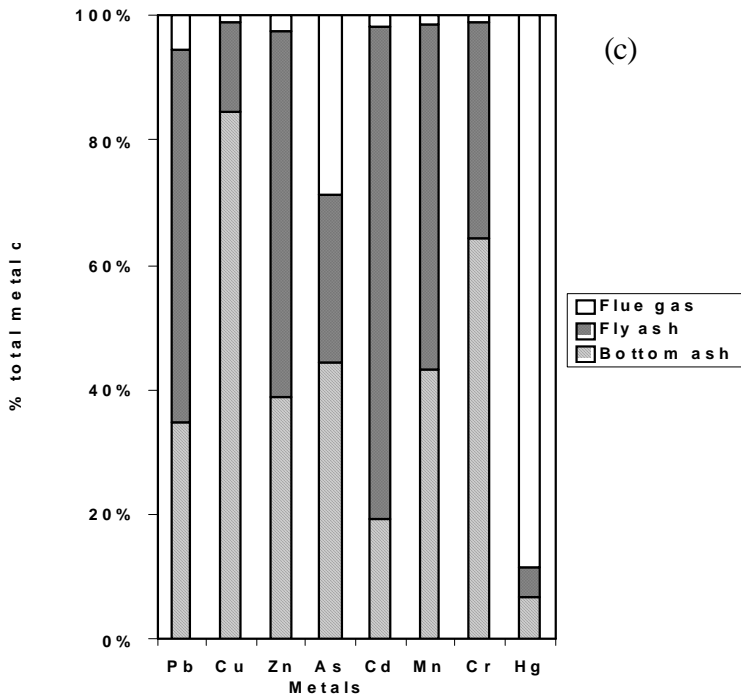
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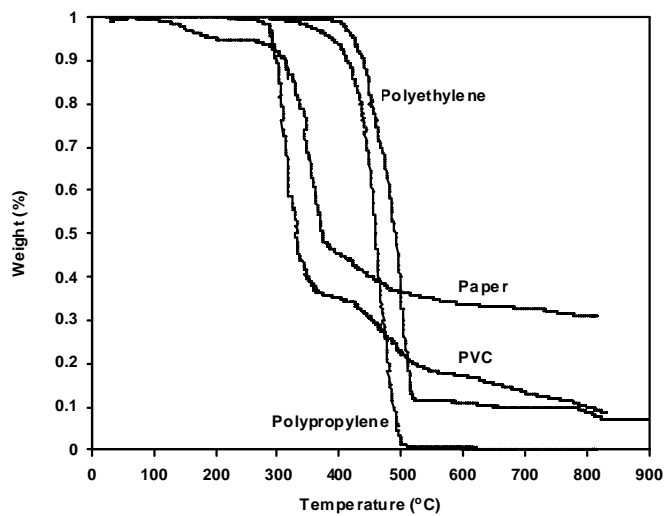
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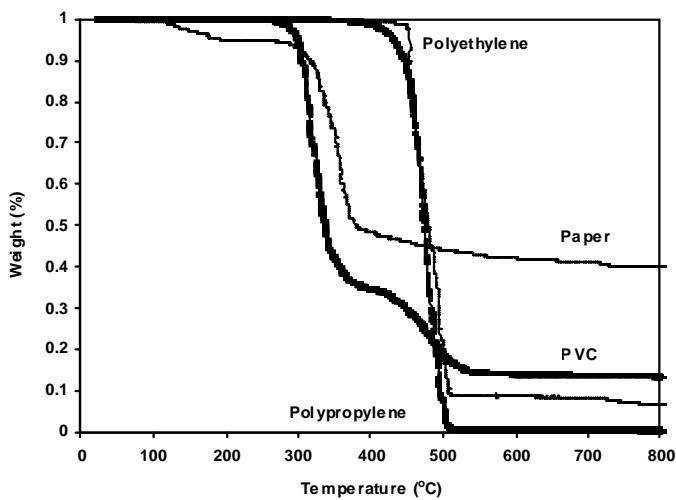
2 **Figure 5. Metal distribution in ash and flue gas as a result of (a) coal combustion, (b) coal + 10%**
 3 **SRF, and (c) coal + 10% RDF**

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2 **Figure 6a. Comparison of TGA curves for paper and plastics (Heating rate = 10 °C/min, Air flow**
 3 **rate = 20 ml/min)**



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5 **Figure 6b. Comparison of TGA curves for paper and plastics (Heating rate = 10 °C/min, N₂ flow**
 6 **rate = 20 ml/min)**

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