Waste ashes as catalysts for the pyrolysis–catalytic steam reforming of biomass for hydrogen-rich gas production

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Abstract
Combustion ashes from coal, refuse-derived fuel (RDF) and waste tyres have been investigated as potential catalysts for the production of a hydrogen-rich gas from waste biomass. The process used a two-stage reactor involving pyrolysis of the biomass followed by catalytic steam reforming of the evolved pyrolysis gases using the ash catalysts. The ashes were also impregnated with 10 wt% nickel to determine the influence on hydrogen production. In the presence of the ash samples, the total gas yield and hydrogen yield significantly increased, particularly for the refuse-derived ash. The ash samples contained a high metal content, including Al, Ca, Mg, Cu and Fe, K, Na and Zn. All such metals have been reported to act as catalysts for hydrogen production. In the absence of catalyst, the total gas yield from the biomass was 39.9 wt% which increased to 52.7 wt% with the tyre rubber ash, to 50.3 wt% with coal ash and 59.5 wt% with RDF ash. The highest hydrogen yield of 7.90 mmol g−1 biomass was produced in the presence of the RDF-derived ash, representing 29.73 vol% H2. Addition of nickel to the combustion ash samples showed a further significant increase of ~ 20% in the yield of hydrogen.

Keywords Biomass · Pyrolysis · Reforming · Waste · Ash

Introduction
The range of fuels used in combustion systems for the generation of power has expanded from the combustion of coal in power plants, to include the use of a variety of waste materials [1]. For example, municipal solid waste and refuse-derived fuels combusted for energy recovery in waste to energy plants, biomass forestry residues for large and smaller scale power production and waste tyres to provide the energy in cement kilns. However, the combustion process generates a residual ash which may be produced in large quantities. For example, it is estimated that 750 Mt of coal ash are produced world-wide each year [2] and ~ 100 million tonnes per year in the EU [3]. Incineration of municipal solid waste in waste to energy plants generates approximately 25 wt% of bottom ash consisting of more than 90% bottom ash and ~ 10% fly ash and air pollution control residues. Bottom ash is regarded as inert, but fly ash and pollution control residues are hazardous. It has been estimated that approximately 18 million tonnes of bottom ash is produced in the EU each year [4].

Often the residual ash from coal combustion or municipal solid waste incineration is deposited in waste landfill sites. However, there is current interest in the sustainable management of waste ashes to maximise the recovery of resources. Residual combustion ashes have been investigated for use as alternative products [1]. For example, coal ash has been processed for use in the production of bricks [5], for use in concrete [6] and for cement production [7]. The ash residue from the incineration of municipal solid waste has been used in the production of light weight aggregate material [8] and also incorporated in asphalt mixtures for paving material [9] and in cement composites [10].

The ash produced from the combustion of the solid hydrocarbon fuels, coal, municipal solid waste and waste tyres contains a range of metal-based components. The main constituents in coal ash from power plants are SiO2, Al2O3, Fe2O3, CaO and MgO [2]. Goodarzi [11] reported a wide range of oxides identified in coal-fired power plant fly ash including, SiO2, Al2O3, Fe2O3, TiO2, P2O5, CaO, MgO, SO3, Na2O, K2O, BaO, SrO, V2O5, NiO, MnO and Cr2O3. Lynn et al. [12] reported that the composition of municipal solid waste incinerator bottom ash comprised largely, SiO2, CaO
and Al₂O₃, with others oxides at lower concentrations identified as Fe₂O₃, Na₂O, MgO, SO₃, P₂O₅, ZnO and CuO. A detailed compositional analysis of incinerator bottom ash showed the presence of various silicates, oxides and carbonates, such as, Ca₃Al₂Si₂O₇, Ca₃Mg₂Si₂O₇, SiO₂, Fe₂O₃, Fe₂O₅, Fe₂O₇, CaCO₃, MgCO₃, Ca(OH)₂, CaSO₄, NaCl and KCl and elemental Fe, Al and Cu [13]. Species identified in waste incinerator fly ash include, Pb₃SiO₅, Pb₃O₂SO₄, Pb₃Sb₂O₇, PbSiO₄, Cd₃(AsO₄)₃Cl₂, CdSO₄, K₂ZnCl₄, ZnCl₂, ZnSO₄, Fe₂O₅, Fe₂O₇, SiO₂, CaSiO₃, Al₂SiO₅, Ca₂Si₃O₉, Ca₃Al₂SiO₆, Ca₃Al₂Si₃O₁₆, NaAlSiO₅, and KAlSiO₅ [14]. Singh et al. [15] reported that the main constituents of the ash produced from waste tyre rubber combustion comprises, SiO₂, Al₂O₃ and ZnO with much lower (< 2.5 wt%) concentrations of Fe₂O₃, K₂O, CaO, TiO₂, MgO, Na₂O and P₂O₅.

The high metal content in such wastes indicates that the ash has the potential to be used as catalytic materials for a variety of applications. For example, Wang et al. [16] investigated the use of coal ash catalysts for the catalytic steam reforming of acetic acid and phenol as representative model bio-oil compounds for the production of hydrogen. They reported that the presence of metals in the ash catalysed the steam-reforming reaction, resulting enhanced carbon conversion and production of hydrogen. Shahbaz et al. [17] investigated the catalytic steam gasification of palm shell as a biomass feedstock with coal ash as a catalyst for the production of syngas and hydrogen. The optimised process conditions with the coal ash catalyst produced a hydrogen yield of 36.9 vol% and syngas yield of 61.9 vol%. They attributed the increased yield of syngas and hydrogen to the presence of Fe₂O₃, MgO, Al₂O₃ and CaO in the ash. Loy et al. [18] used coal ash as a catalyst for the catalytic pyrolysis of rice husks for the production of syngas. They reported that coal ash catalysed the production of hydrogen and syngas compared to non-catalytic pyrolysis, producing 68.3 vol% syngas and also enhancing the hydrogen content of the syngas by 8.4 vol%.

Incinerator bottom ash has been used as a catalyst for the cracking of tars produced from the gasification of municipal solid waste [19]. The tar was represented as toluene model compound and using the incinerator bottom ash catalyst, the toluene cracking conversion was 40.1% at 750 °C, which was reported to be significantly higher than that of thermal cracking. At a higher ash catalyst temperature of 950 °C the toluene conversion was 94.2%.

The analysis of the literature shows that ash derived from combustion of solid fuels possesses catalytic properties for several applications. The motivation for the work presented here was to expand this knowledge to investigate the potential of ash residues derived from the combustion of coal, refuse-derived fuel (RDF) and waste tyre rubber to act as catalysts for the catalytic steam reforming of biomass pyrolysis gases. A two-stage reactor has been used, whereby the first stage involves pyrolysis of biomass and the product pyrolysis gases are then passed directly to a catalytic steam-reforming reactor containing the combustion ashes. The gases from the process have been analysed in detail with the aim of maximising the production of a hydrogen-rich gas.

**Materials and methods**

**Materials**

Waste wood pellets with a particle size of 1 mm were used as the biomass feedstock for pyrolysis reforming/steam gasification experiments. The wood pellets were produced as compressed saw dust pellets from waste wood processing by Liverpool Wood Pellets Ltd, Liverpool, UK. The proximate analysis of the wood pellets gave 75.0 wt% volatiles, 7.0 wt% moisture, 2.0 wt% ash, and 15.0 wt% fixed carbon. Elemental analysis gave 46.0 wt% carbon, 5.6 wt% hydrogen, 0.7 wt% nitrogen and 45.7 wt% oxygen (by difference).

Ash samples derived from coal, waste tyre and RDF were used as a catalyst for gas and hydrogen production during the biomass pyrolysis–catalytic steam-reforming process. The coal, waste tyre and RDF were ashed in a furnace at 700 °C for 4 h. The produced ash samples were further examined as a catalyst support using an impregnation method to produce Ni-ash catalysts. For this purpose, the coal, waste tyre, and RDF ash samples were impregnated with nickel using an aqueous solution of Ni(NO₃)₂·6H₂O (corresponding to Ni loading of 10 wt%). The mixture of ash and Ni(NO₃)₂·6H₂O was stirred using a magnetic stirring apparatus at 100 °C to produce a slurry, dried overnight and calcined followed by calcination at 750 °C for 3 h in an air atmosphere. Finally, the obtained catalysts were reduced at 800 °C under hydrogen.

**Characterization of the produced ash catalysts**

The composition of the ash samples was determined using a Thermo Advant XP X-ray fluorescence (XRF) spectrometer. The samples were prepared as fused beads to provide a homogenous sample. A lithium borate flux was added to the ash samples and heated to 900 °C in a platinum crucible and then cast into a mould. The resultant fused bead was analysed using OXSAS programme. The surface area of the produced coal, waste tyre and RDF ash samples was determined according to the Brunauer–Emmet–Teller (BET) method using a Quantachrome NOVA 2200e series apparatus. Samples were degassed at 250 °C for 3 h under vacuum prior to measurement. A Bruker D-8 diffractometer with Cu Kα radiation operated at 40 kV and 40 mA was used to obtain the X-ray diffraction (XRD) patterns of the original
The XRD pattern identification was obtained using High-Score Plus software. A Hitachi SU8230 scanning electron microscope (SEM) coupled to an energy-dispersive X-ray spectrometer (EDXS) system was used to characterise and examine the surface of the tested catalysts.

Reactor system

It is a two-stage fixed bed reactor system consisting of a first stage involving pyrolysis of the wood pellets and a second-stage catalytic reactor containing the produced residual coal, tyre and RDF ashes. A schematic diagram of the system is shown in Fig. 1. The reactors were constructed of stainless steel with an inner diameter of 22 mm and a length of 160 mm with monitoring and control of gas flow and temperature.

The experiments consisted of the initial heating of the second-stage catalytic reactor containing 1.0 g of ash catalyst to a temperature of 800 °C and maintained at that temperature throughout the experiment. In addition, water was injected into the second-stage catalytic reactor using a syringe pump at a rate of 4.0 g h⁻¹ (steam/biomass (S/B) ratio of 2). After the catalytic reactor temperature was stabilised at 800 °C, the first-stage pyrolysis reactor containing the biomass (1.0 g) was heated at a rate of 40 °C min⁻¹ to a final temperature of 600 °C. The evolved pyrolysis gases from biomass thermal degradation mixed with the steam and underwent catalytic steam reforming over the coal, tyre and RDF ash catalysts and Ni-ash catalysts. For comparison with the ash-based catalysts, clean and washed quartz sand was used in place of the catalyst in the second-stage reactor. The sand would have none of the catalytic metals found in the ash-based catalysts. Any condensable liquid products, including condensed steam, were collected in a condenser system cooled with dry ice. Gases passing through the condenser system were collected in a Tedlar™ gas sample bag.

The product gases in the gas sample bag were analysed immediately after each experiment using three Varian CP-3380 gas chromatographs (GC). N₂, H₂, and CO gases were analysed using a 60–80-mesh molecular sieve packed column, Ar carrier gas and thermal conductivity detector (TCD). CO₂ gas was analysed on a second Varian GC with a 80–100-mesh HayeSep packed column, Ar carrier gas and TCD. CH₄, C₂H₆, C₂H₄, C₃H₆, C₃H₈, C₄H₁₀, C₅H₁₀ and C₆H₁₂ hydrocarbon gases were analysed on a third Varian GC with an HayeSep 80–100-mesh molecular sieve column, N₂ as carrier gas and flame ionisation detector (FID).

Results and discussion

Characterization of the ash catalysts

The ash samples that were produced from the coal, waste tyre rubber and RDF and used for the pyrolysis–catalytic steam-reforming experiments were characterised using XRF, surface area analysis, XRD, and SEM. Table 1 shows the metal-oxide compositional analysis of the derived combustion ashes produced by XRF. The waste tyre rubber combustion ash is dominated by the presence of zinc which is added to the tyre vulcanisation process to improve the physical properties of the rubber. In addition, silica (SiO₂) is present in high concentration, and the silica is often added as filler material for the tyre formulation. The RDF ash contains typical metal compounds that are found in waste incinerator combustion ash. RDF is prepared from municipal solid

| Sample    | Tyre ash (wt%) | RDF ash (wt%) | Coal ash (wt%) |
|-----------|----------------|---------------|----------------|
| SiO₂      | 24.2           | 3.31          | 58.2           |
| ZnO       | 41.5           | 0.6           | –              |
| Al₂O₃     | 2.3            | 21.0          | 20.8           |
| CaO       | 3.3            | 16.9          | 2.9            |
| MgO       | 0.7            | 4.6           | 1.4            |
| CuO       | 0.9            | 0.1           | –              |
| K₂O       | 0.9            | 2.5           | 1.7            |
| Fe₂O₃     | 5.7            | 3.3           | 9.3            |
| Na₂O      | 3.0            | 1.9           | 2.3            |
| SO₃       | 15.5           | 3.03          | 0.9            |

Table 1 Chemical composition of the ash samples (wt%)
waste with the aim of raising the calorific value of the product fuel by removing metals, glass and ceramic materials. The RDF ash prepared in this work contained mainly Al2O3 and CaO with minor concentrations of other alkali metals. The coal ash sample consisted of mainly SiO2, Al2O3 and Fe2O3.

Table 2 shows the surface areas and pore diameter for the ash samples and the nickel-ash catalysts. The BET surface areas of the ash samples produced from RDF, tyre rubber and coal were found to be 21.8, 13.7 and 2.5 m2 g−1, respectively. However, nickel catalysts supported on RDF ash showed a slightly reduced surface area than the original ash samples.

The morphologies of the produced ash catalysts and Ni-ash catalysts were characterised by scanning electron microscopy (SEM) and are presented in Fig. 2. The coal ash shows the characteristic cenospheres produced during coal combustion. The tyre rubber-derived ash and RDF-derived ash show less structure. The addition of 10 wt% nickel to the ash appears to show surface deposits of the metal.

Figure 3 shows XRD profiles of the produced ash catalysts and Ni-ash catalysts. The diffraction peaks observed around 2-Theta 44° and 52° with the ash-supported nickel catalysts were related to the presence of metallic Ni and no peaks corresponding to the NiO phase were identified. This was expected as the catalysts were reduced before the catalytic steam-reforming process. The XRD diffraction peaks observed with Ni/tyre ash were smaller compared to the XRD peaks obtained with either Ni/coal or Ni/RDF catalysts, which might indicate that the particle size of Ni is smaller for the Ni/tyre ash sample.

Pyrolysis–catalytic steam reforming of biomass with ash catalysts

Table 3 shows the product yield and gas composition for the pyrolysis–catalytic steam reforming of biomass with the ash-based catalysts. In addition, experiments were carried out with clean quartz sand in place of the catalyst and the results shown in Table 3 indicate that catalytic cracking and steam reforming are taking place, even in the presence of sand. The particles of sand in the second-stage reactor is maintained at 800 °C, producing a hot surface for thermal cracking and steam reforming of the evolved pyrolysis gases from the biomass thermal degradation. In the presence of the sand, the total gas yield in relation to the feedstock biomass was 39.9 wt% which increased to 52.7 wt% when tyre rubber ash was introduced and with coal ash and RDF ash, increased to 50.3 wt% and 59.5 wt%, respectively. The metals in the ash sample produce a catalytic effect producing more product gas, particularly for the RDF ash sample. The residual char yield derived from the biomass pyrolysis in the first-stage reactor was constant at ~ 19.0 wt% for all the experiments since it was unaffected by the second-stage reactions. Consequently, the liquid yield from biomass (by difference) represented 41.1 wt% for the sand, 28.3 wt% for the tyre ash, 30.7 wt% for the coal ash and 21.5 wt% for the RDF ash. Suggesting that cracking of the higher molecular weight biomass volatile compounds to gas occurred in the presence of the ash samples compared with the non-catalytic sand.

The gas composition shown in Table 3 shows that the product gases consist of mainly CO, CO2, H2, CH4 and C2H4 hydrocarbons. The influence of ash addition to the catalytic reactor is to increase the level of catalytic steam reforming of the hydrocarbons (CH4 and C2H4) which show a decrease in concentration compared to the non-catalytic experiment (sand) and a consequent increase in H2 production. In addition, the decrease in CO and increase in CO2 and H2, compared to the non-catalytic (sand) results indicates that the presence of the ash enhances the water gas shift reaction. Overall, the presence of the ash with their inherent metal content acts as a catalyst for the production of hydrogen from the biomass pyrolysis gases. The highest hydrogen gas yield of 29.73 vol% representing 7.90 mmol H2 g−1 biomass was produced in the presence of the RDF-derived ash.

Table 1 showed a high metal content in the ash, including Al2O3, CaO, MgO, CuO and Fe2O3, depending on the waste ash used. During the catalytic steam-reforming process, the metal oxides would undergo reduction to the metal via the reducing gases such as hydrogen and carbon monoxide produced during the process. The metal oxides, Al2O3, CaO, MgO, and Fe2O3, have been used as catalysts for the catalytic steam reforming of biomass [20, 21]. Magnesium, acting as a promoter in nickel-based catalysts, has also been investigated by Garcia et al. [22] where it was reported to produce a higher hydrogen yield in the steam reforming of an aqueous fraction of bio-oil. In a later report, Garcia et al. [23] also showed that magnesium containing NiMgAl2O5 catalyst produced high total gas and hydrogen yields for the hydrogen production from biomass catalytic steam gasification for biomass in the form of pine sawdust. The high yields of H2 and CO2 coupled with low yields to CH4, C2 hydrocarbons and CO suggesting high catalytic activity for
the steam-reforming reactions of hydrocarbons and the water gas shift reaction. The presence of copper as a Ni-catalyst promoter metal for the production of hydrogen has been shown to enhance the yield of hydrogen and decomposition of methane [24]. The addition of zinc to catalytically steam reform wood sawdust biomass pyrolysis gases using a Ni/Zn/Al₂O₃ catalyst showed that the total gas yield and hydrogen yields were increased producing a maximum gas yield of 74.8 wt% and H₂ yield of 20.1 mmol H₂ g⁻¹ biomass [25]. In the absence of the Ni/Zn/Al₂O₃ catalyst, the total gas yield was only 33.0 wt% and hydrogen yield only 2.4 mmol H₂ g⁻¹ biomass. On the one hand, zinc modifies the surface structure and the surface chemistry of the catalysts by formation of zinc aluminates, and on the other hand, zinc oxide can be reduced to metallic zinc under reaction conditions, thus modifying the catalytic properties of the active phase. The presence of Zn increases the ethanol conversion to gaseous compounds as compared with the catalyst supported on the

Fig. 2 Scanning electron micrographs of the ash and Ni-ash catalysts

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Zn-free commercial alumina. Alkali metals (K₂O, Na₂O) were also identified in the different waste ashes (Table 1) used in this study. Sodium and potassium compounds have also been reported as catalysts for hydrocarbon decomposition and improved gas quality [21]. It has been suggested that the presence of alkali metal compounds act as catalysts to increase hydrocarbon cracking and reforming and thereby increase hydrogen production from biomass [20, 26].

The wide range of metal and alkali metal catalysts present in the combustion ash samples will clearly have a catalytic effect on the catalytic steam reforming, dry reforming, thermal cracking and gasification reactions involving the evolved biomass pyrolysis gases. Other researchers have investigated the use of combustion ashes and reported a catalytic effect. Wang et al. [16] used a combustion ash produced from coal as a catalyst for the catalytic steam reforming of acetic acid and phenol and compared their results with several commonly used steam-reforming catalysts. They showed that the coal ash sample produced similar carbon conversion efficiencies and hydrogen for acetic acid to that of an Fe–Al₂O₃ catalyst. For phenol, the results were lower in terms of conversion and H₂ yield.

Pyrolysis–catalytic steam reforming of biomass with nickel-ash catalysts

Table 4 shows the influence of the addition of 10 wt% nickel to the combustion ash catalysts in terms of gas yield and gas composition derived from the pyrolysis–catalytic steam reforming of biomass. In addition, comparison is again made with the yields in relation to no-catalyst (in the presence of quartz sand). There was only a small influence of nickel addition to the combustion ashes in terms of any increase in total gas yield compared to the non-nickel-impregnated combustion ash samples shown in Table 3. For example, the total gas yield increased by 6.6 wt% for the Ni-tyre ash, 4.6 wt% for the Ni-coal ash sample, whilst the Ni-RDF ash sample showed a similar total gas yield compared to the ash-only catalysts. However, the hydrogen yield in relation to the mass of biomass feedstock showed a significant increase for all the nickel-impregnated ash catalysts. For example the

![XRD analysis of the ash and Ni-ash catalysts](image)

**Table 3** Gas yield and gas composition from the pyrolysis–catalytic steam reforming of biomass with the ash-based catalysts

| Sample       | Sand | Tyre ash | Coal ash | RDF ash |
|--------------|------|----------|----------|---------|
| Gas yield (wt%) | 39.9 | 52.7     | 50.3     | 59.5    |
| Gas composition (vol%) |
| CO         | 43.50 | 35.29    | 42.16    | 34.51   |
| CO₂        | 14.10 | 19.22    | 15.49    | 20.63   |
| H₂         | 19.92 | 27.98    | 22.08    | 29.73   |
| CH₄        | 15.33 | 12.10    | 14.38    | 10.74   |
| C₂H₄       | 7.15  | 5.42     | 5.88     | 4.39    |
| H₂ yield (mmol g⁻¹ biomass) | 3.39 | 6.54     | 4.79     | 7.90    |

**Table 4** Gas yield and gas composition from the pyrolysis–catalytic steam reforming of biomass with the nickel-ash-based catalysts

| Sample       | Sand | Nickel-tyre ash | Nickel-coal ash | Nickel-RDF ash |
|--------------|------|----------------|-----------------|---------------|
| Gas yield (wt%) | 39.9 | 59.3           | 54.9            | 58.03         |
| Gas composition (vol%) |
| CO         | 43.50 | 34.64         | 39.40           | 33.60         |
| CO₂        | 14.10 | 15.19         | 14.07           | 17.30         |
| H₂         | 19.92 | 35.79         | 28.51           | 34.40         |
| CH₄        | 15.33 | 10.00         | 12.78           | 10.60         |
| C₂H₄       | 7.15  | 4.37          | 5.25            | 4.20          |
| H₂ yield (mmol g⁻¹ biomass) | 3.39 | 10.50         | 7.29            | 9.66          |
Ni-tyre ash catalyst increased the hydrogen yield to produce the highest yield 10.5 mmol^{-1} \text{H}_2$ from the biomass. The presence of the nickel in the ash catalysts showed that nickel metal catalytic steam reforming of the hydrocarbons (CH$_4$ and C$_2$H$_2$) occurred, producing increased hydrogen yield in terms of H$_2$ yield as mmol g$^{-1}$ biomass and also a volumetric increase in hydrogen content in the gas product. The results suggest that the high metal content of the combustion ashes acts as the main catalytic effect and that addition of a further active nickel content to the catalyst produces an approximate 20% increase in hydrogen yield.

The addition of nickel to combustion ash to enhance the catalytic effect of the metals and compounds in the ash has been examined by other researchers. For example, Wang et al. [16] investigated the catalytic steam reforming of bio-oil model compounds in the form of acetic acid and phenol using a nickel-coal ash catalyst with a 15 wt% nickel loading. The conversion of acetic acid was 57.5% and for phenol conversion was 26.3% with the combustion ash, but increased to 98.4% and 83.5%, respectively, with the nickel-ash catalyst. In addition, they reported that the yield of H$_2$ from the catalytic steam reforming of acetic acid using the combustion ash was 50.2% and for phenol was 19.6%. However, with the addition of nickel to the ash, the yield of H$_2$ improved to 85.6% for acetic acid and 79.1% for phenol at a catalyst temperature of 700 °C and steam/carbon ratio of 9.2. The influence of temperature was to increase the conversion of the bio-oil model compounds and increase hydrogen yield. In a later report, Wang et al. [2] added both nickel and iron to a coal combustion ash to produce a Ni–Fe-ash catalyst for the catalytic steam reforming of acetic acid. They reported 100% conversion of the acetic acid and a hydrogen yield of 89.6% at a catalyst temperature of 700 °C.

**Conclusions**

The work reported here has shown the potential of combustion ash produced from coal, municipal solid waste and tyres to the combustion ash samples showed a further significant increase of ~20% in the yield of hydrogen.

The ashes contained Al$_2$O$_3$, CaO, MgO, and Fe$_2$O$_3$ which have been used in the catalytic steam reforming of biomass. In addition, the ashes contain metals such as Mg, Zn and Cu which have been used as catalyst promoters in steam-reforming catalysts and K$_2$O, Na$_2$O which have been used as catalysts for hydrocarbon cracking and reforming. However, it is difficult to assign particular metals present in the ashes for the catalytic effect on hydrogen production from biomass. For example, the ash samples contained mixtures of metals and metal species which would contribute to the catalytic activity of the ashes and also would have had a synergistic or promotional effect on enhancing hydrogen production from the biomass. Combustion ashes are regarded as problematic wastes with low commercial value and often are disposed of in waste landfill sites, representing a waste of resource. This work has identified combustion ashes as a potentially useful and low cost catalyst for catalytic steam reforming of biomass pyrolysis gases.

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