Gasification of poultry litter in a lab-scale bubbling fluidised bed reactor: Impact of process parameters on gasifier performance and special focus on tar evolution

Giannis Katsaros a,⇑, Daya Shankar Pandey a,b,⇑, Alen Horvat c, Guadalupe Aranda Almansa e, Lydia E. Fryda e, James J. Leahy d, Savvas A. Tassou a

a RCUK Centre for Sustainable Energy Use in Food Chains (CSEF), Brunel University London, Uxbridge UB8 3PH, UK
b School of Engineering and the Built Environment, Anglia Ruskin University, Chelmsford CM1 1SQ, UK
c Carlos III University of Madrid, Energy Systems Engineering Group, Thermal and Fluids Engineering Department, Avda. de la Universidad 30, 28911 Leganés, Madrid, Spain
d Carbolea Research Group, Department of Chemical Sciences, Bernal Institute, University of Limerick, Limerick V94 T9PX, Ireland
e Energy Research Centre of the Netherlands, Biomass and Energy Efficiency, Petten, the Netherlands

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Abstract

Poultry litter (PL) gasification was experimentally investigated using a lab-scale bubbling fluidised bed reactor. Characterisation of the gasification process was performed in terms of yields and compositions of both gas and tar, lower calorific value (LCV) of the product gas, cold gas efficiency (CGE) and carbon conversion efficiency (CCE). Experiments were carried out at different temperatures (700–750 °C) and equivalence ratios (ERs). The effect of gasifier temperature at a constant ER of 0.21 shows that an increase in temperature improved the gasification process performance whilst the total tar content decreased, implying that higher temperature enhances the conversion of biomass to product gas. The total gas yield increased from 0.93 to 1.24 N2-free m3/kgfeedstock-daf, LCV increased from 3.38 MJ/m3 to 4.2 MJ/m3, while the tar content was reduced by 24% (5.6–4.25 gtar/kgfeedstock-daf). The detailed analyses of tar compositions reveal that styrene and xylenes were the most abundant compounds in the secondary tar group. Moreover, naphthalene and 1, 2-methyl naphthalene were the dominant compounds found in tertiary polycyclic aromatic hydrocarbons (PAH) and alkyl tertiary groups, respectively. Furthermore, at the highest tested temperature of 750 °C and ER of 0.25, bed agglomeration took place causing the shutdown of the gasifier. The defluidisation of the bed occurred due to the high ash content of PL comprising of low melting temperature alkali compounds. The results obtained from this study showed the performance and potential challenges associated with gasifying PL in a fluidised bed reactor for the combined heat and power production at farm level.

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1. Introduction

The production of poultry meat is expanding throughout the world, imposing significant challenges related to the efficient management of associated litter generation. In 2018, on a monthly basis approximately 80 million birds, mostly chickens were slaughtered in the UK alone (Rumsey, 2018). A recent study estimated that the amount of PL produced falls between 1.75 and 5.7 kg of litter/bird over a 42 days production cycle (Dalólio et al., 2017). In the UK, the total amount of PL generated annually ranges between 140,000 and 456,000 tonnes, while its LCV on an as received basis ranges between 8.75 GJ/tonne and 14.27 GJ/tonne (Lynch et al., 2013). Therefore, the estimated potential energy from PL varies between 1.22 PJ and 6.5 PJ. Considering its energy potential, PL can be utilised as a renewable feedstock for bioenergy production.

The growing demand for poultry meat has led to intensive livestock farming which outperformed the prevalent traditional farming based on small installations. Although the intensive farming is
more efficient and cost effective, it creates serious environmental concerns due to the large amount of waste accumulated within a confined area where the available arable land for manure application as a nutrient source is limited (Bernal et al., 2015). Excessive soil fertilisation with nutrient rich animal manure can lead to eutrophication, nitrate leaching, crop toxicity, odours and emissions of greenhouse gases (NH₃, NOx, N₂O) to the atmosphere (Billen et al., 2015; Joseph et al., 2012; Lynch et al., 2013; Taupe et al., 2016).

Growing environmental concerns associated with excessive fertilisation demands the development of alternative viable options for treating animal waste. Due to its relatively high energy content and fixed carbon, PL in particular, has gained attention lately in energy conversion processes. Currently two different pathways, biochemical and thermochemical conversion are being exploited. The choice of technology depends on the feedstock properties, the desired end product, economic feasibility and environmental regulations (Pandey et al., 2016). The slow production rate of the anaerobic digestion process, the need for feedstock with high moisture content (moisture content of PL varies significantly from batch to batch and has relatively high solid content) and related high capital costs make this method less suitable for PL treatment (Burra et al., 2016; Joseph et al., 2012). Composting, although producing a fertiliser with a value in the market, poses some serious drawbacks such as the odours generated during the process, the need for land availability and high equipment costs (Joseph et al., 2012). Thermochemical conversion seems a promising option for PL treatment, since it can reduce the volume of the waste by up to 80–95%, upgrade PL to higher value products (e.g. bio-oil, synthetic natural gas), destroy pathogens due to high operating temperatures, while it also offers the possibility of electricity, heat generation and biofuel production (Arena, 2012).

Thermochemical conversion routes are divided into three core technologies, namely combustion, gasification and pyrolysis. Amongst these technologies, combustion is already proven and mature whereas gasification and pyrolysis technologies are still in their early commercialisation stage and pose different challenges that need to be addressed prior to their deployment at full commercial scale. In a gasification process, a carbon-based feedstock is partially oxidised at high temperatures (700–1200 °C) in the presence of an oxidant (air, steam, oxygen, or mixtures thereof) under sub-stoichiometric conditions. The result of this process is the production of a combustible gas known under the different names “producer gas”, “product gas” or “syngas” consisting mainly of CO, H₂, CO₂, CH₄ and small amount of C₂, compounds, along with impurities, such as fine particulates, tars and alkali metals (Arena, 2012). To maintain consistency, the term “product gas” is used throughout the paper, since “syngas” refers to gas consisting only of H₂ and CO utilised as basis for the production of chemicals and fuels (Arena et al., 2010). The calorific value of the product gas depends on the oxidant supplied, for instance if air is used as an oxidant it results in a low calorific value (4–7 MJ/Nm³) gas as the product gas is diluted by atmospheric nitrogen (up to 60%) (Arena, 2012).

Gasification of biomass has emerged as a cleaner technology compared to combustion offering higher efficiency and lower gaseous pollutants such as SOₓ, NOₓ, heavy metals and particulate emissions that are in compliance with emission standards (Arena, 2012; Pan and Pandey, 2016). However, gasifying PL poses significant challenges due to the presence of compounds with low eutectic temperature in its high ash concentration, which that can lead to fluidised bed agglomeration as well as slagging and fouling issues on the heat transfer surfaces and subsequent equipment train (Bartocci et al., 2017).

Different type of reactors can be employed in a gasification process, namely fixed bed, fluidised bed and entrained flow. Among the various reactors considered, fluidised bed reactors have gained attention recently as they offer greater fuel flexibility and are able to maintain the temperature below the ash melting point, secure high heat transfer rates, achieve a higher production capacity and are suitable in wider range of applications (Belgorno et al., 2003).

High interest in PL valorisation as a gasification feedstock together with the specific challenges of this fuel is reflected in the large number of publications in the recent years. The effect of limestone addition to prevent agglomeration while gasifying PL has been studied. The authors reported that by adding limestone, agglomeration did not occur below 800 °C compared to the case without limestone where agglomeration was observed at 750 °C (Pandey et al., 2016). The effect of ash composition on PL gasification has been investigated in a pre-pilot reactor by comparing two batches of manure taken from an industrial chicken farm. The experiments were carried out at different ERs (0.27–0.4) and temperatures (700–800 °C). The findings revealed the role of ash composition, since all the process parameters were significantly reduced in the batch with the higher ash content (Di Gregorio et al., 2014). A recent study focused on producing energy from an innovative 300 kW thermal power gasification plant installed on a poultry farm located in central Italy. Aspen Plus v8.0 model was developed by the authors to predict the outlet gas composition and its LCV which was found in the range 3–5 MJ/m³ for an ER of 0.2 (Cavalaglio et al., 2018). Six different model based energy integration schemes were applied to a small-scale gasification process for onsite power generation. The findings revealed CE and energetic efficiency ranging between 58.4–79.5% and 46.8–65.7% respectively (Font Palma and Martin, 2013). The techno-economic feasibility of generating biochar, electricity and heat production from PL was investigated using a model developed on ECLIPSE software. The authors concluded that gate fees, carbon credits and renewable energy certificates greatly influence the breakeven selling price of produced biochar (Huang et al., 2015).

One of the largest technical obstacles that hinders further development and commercialisation of gasification technology is the presence of tar in the product gas. Tar is a mixture of complex hydrocarbons which may condense in the process installation if the temperature drops below the tar dew point. Condensation of tar leads to the formation of a black and sticky material which causes system malfunctioning due to clogging and fouling. Multiple definitions of tar can be found across literature. One of the most representative definition given by Basu (2010) is the one derived from IEA’s gasification task force which defines tar as “the organics produced under thermal or partial-oxidation regimes (gasification) of any organic material, are called tar and are generally assumed to be largely aromatic”.

There are two different tar classifications considered, either based on the temperature regime under which tar compounds are formed (Milne et al., 1997) or based on water solubility, dew point temperature, and aromatic ring number (Kiel et al., 2004). According to Milne et al. (1997), tar is classified in primary, secondary, alkyl tertiary and tertiary tar groups. Primary tar derives from pyrolysis reactions of ligno-cellulosic materials at temperatures between 200 and 500 °C. Primary tar consists of highly oxygenated compounds such as acids, sugars, alcohols, and ketones (Horvat et al., 2016c). As the temperature increases and with the presence of the gasification agent, primary tar releases functional groups and reforms into light non-condensable gases and heavier compounds called secondary tar such as phenols and olefins which remain stable up to the temperature of 750 °C. Above 750 °C the secondary tar undergo rearrangement into tertiary tar by completing the condensation pathway resulting in purely aromatic species (Rios et al., 2018). Tertiary tar consists mainly of polycyclic aromatic compounds (PAHs) such as naphthalene, acenaphthylene, and pyrene. PAHs increase exponentially with temperature due to
to polymerisation. Alkyl tertiary tar such as methyl naphthalene and biphenyl are intermediates between secondary and PAH tar. The yield of PAHs appears to peak at 850 °C followed by a gradual decrease (Rios et al., 2018; Van Paasen and Kiel, 2004). Tertiary tar is not present in the initial biomass but rather as a product of decomposition and rearrangement of secondary tar. Typically tertiary and primary tar do not co-exist in the reactor (Molino et al., 2016; Rios et al., 2018) nevertheless, in some reactor arrangements this scenario is possible.

Fluidised bed gasifiers generate a blend of secondary and tertiary tar groups in the order of 10 g/Nm³ (Rabou et al., 2009). Furthermore depending on the end-use application, different tar concentration limits are in place. As an example if the product gas is supplied in internal combustion engines tar limits are in the range of 50–100 mg/Nm². Different product gas applications and the respective limits in tar are given in Rios et al. (2018). When the amount of tar is excessive gas cleaning is imperative, which increases the process complexity and costs. In addition to the total amount of tar, the nature of the individual tar compounds is a crucial parameter. The nature of these compounds determines the tar dew point, a decisive factor for downstream applications. In general, the presence of tar compounds with higher molecular weight tends to increase tar dew point and vice versa.

In the recent past PL gasification has been studied extensively but most of the studies have focused on the composition and calorific value of the product gas. The present work apart from studying the effect of temperature on various process parameters (LCV, CCE, CCE etc.), also aims to give a useful insight into the amount and composition of tar derived from PL gasification in a fluidised bed reactor. To the best of authors’ knowledge, detailed analysis of the tar generated from PL does not exist in the literature. The temperature range used in this work was chosen based on the findings of previous researchers taking into the consideration of the challenges associated with de-fluidisation and agglomeration of the bed during the PL gasification (Pandey et al., 2016; Taupe et al., 2016; Di Gregorio et al., 2014). Overall, the findings of this study reveal an effective method of managing the unavoidable waste generated and additionally they can be helpful in examining a potential application of the product gas for heat and power generation at a farm level.

2. Experimental

2.1. Materials

PL is a blend of poultry excreta, waste feed, feathers and bedding material such as straw, peat or sand. PL was collected from a local poultry farm in Finland and it was partially dried and sieved to a particle size range of 0.5–0.98 mm before being fed into the reactor. Table 1 reports the ultimate and proximate analysis along with the chemical composition of PL ash. Fixed carbon was calculated by subtracting the percentages of volatile matter and ash from 100% on a dry basis (d.b.). Similarly, the oxygen content was determined by the difference from the elements presented in ultimate analysis. For the determination of the chemical composition of PL ash (generated at 550 °C according to BS EN 14775:2009), the generated ash was digested and analysed using an Agilent Inductively Coupled Plasma-optical emission spectrometry. The ash contained high concentrations of alkali metals such as K and Na that promote agglomeration and consequently can cause disruption of continuous fluidised bed gasification. These alkali metals along with the high concentration of Cl in PL contribute significantly to the potential challenges associated with fouling, agglomeration and corrosion.

2.2. Experimental facility

The experimental setup illustrated in Fig. 1 is located at the Energy Research Centre of the Netherlands (ECN part of TNO). The experimental campaign was carried out under the framework

| Table 1 Ultimate, proximate analyses of PL and chemical composition of PL ash. |
|-----------------------------------------------|
| Proximate analysis (wt. %)                      |
| Moisture (a.r.)                                | 9.70 |
| Ash content (d.b.)                             | 14.30 |
| Volatile matter (d.b.)                         | 69.60 |
| Fixed carbon* (d.b.)                           | 16.10 |
| Ultimate analysis (wt.% d.b.)                  |
| Carbon                                         | 42.8 |
| Hydrogen                                       | 5.5 |
| Nitrogen                                       | 3.9 |
| Sulphur                                        | 0.60 |
| Chlorine                                       | 0.25 |
| Oxygen*                                        | 32.65 |
| LHV (in MJ/kg)                                 | 16.78 |

| Chemical composition of PL ash                  |
|-----------------------------------------------|
| Major elements                                 |
| Elements Amount (mg/kg, d.b.)                  |
| Alumnum (Al)                                   | 1200 |
| Calcium (Ca)                                   | 15,500 |
| Iron (Fe)                                      | 1600 |
| Magnesium (Mg)                                 | 8200 |
| Manganese (Mn)                                 | 600 |
| Phosphorous (P)                                | 10,200 |
| Potassium (K)                                  | 27,700 |
| Silicon (Si)                                   | 7300 |
| Sodium (Na)                                    | 4200 |
| Sulphur (S)                                    | 6100 |
| Titanium (Ti)                                  | 95 |
| Zinc (Zn)                                      | 450 |
| Minor elements                                 |
| Elements Amount (mg/kg, d.b.)                  |
| Arsenic (As)                                   | <0.50 |
| Barium (Ba)                                    | 29.0 |
| Cadmium (Cd)                                   | 0.14 |
| Cobalt (Co)                                    | 1.90 |
| Chromium (Cr)                                  | 16.0 |
| Copper (Cu)                                    | 84.0 |
| Mercury (Hg)                                   | <0.02 |
| Molybdenum (Mo)                                | 4.80 |
| Nickel (Ni)                                    | 16.0 |
| Lead (Pb)                                      | 1.50 |
| Antimony (Sb)                                  | <0.50 |
| Thallium (Tl)                                  | <0.50 |
| Vanadium (V)                                   | 4.20 |

*Calculated by difference; Oxygen = 100-(C + H + N + S + Cl + Ash content); Fixed carbon = 100 – (Volatile matter + Ash content).
of the European project BRISK2. Fuel is fed into an atmospheric bubbling fluidised bed reactor by two mechanical screw feeders under 1 Nl/min flow of N2 (flush gas) in order to avoid backflow of gases. The reactor consists of two different zones: (i) bed section with an internal diameter of 74 mm and 500 mm height and (ii) the freeboard section with an internal diameter of 108 mm and height of 600 mm. The lab-scale reactor operates in allothermal mode, implying that the desired temperature cannot be achieved by controlling the ER alone. Therefore external heat source is needed, which was realised by electrical means under inert conditions. The fluidising medium (a mixture of N2 and air calculated to achieve a particular ER value while maintaining a constant fluidisation velocity) is adjusted and introduced from the bottom of the reactor through the perforated distributor plate. The fluidising medium is preheated to 160°C before being introduced into the reactor. The product gas exits the freeboard section passing through a cyclone where entrained particles of char and ash were removed. After the cyclone, part of the raw product gas is sampled for chemical analysis, while the rest is combusted in a flare. Product gas for chemical analysis flows through the hot filter to remove the finest particles that escape from the cyclone. The downstream section of the reactor including a hot filter is maintained at 400°C, preventing tar condensation inside the pipes. Tar and moisture samples were collected via a sampling port located after the hot filter. Successive cold filter removes tar prior to an online micro-gas chromatography (GC) analyser.

2.3. Test procedure

Considering the high ash content in PL and possible agglomeration issues, experiments were conducted at lower temperature starting from 700°C. Air and N2 were continuously supplied from the bottom of the reactor at a total flow rate of 12 Nl/min in order to maintain an adequate fluidisation regime while ensuring the set gasification conditions. The minimum theoretical fluidisation velocity at the specified operating conditions was calculated using correlation proposed by Wen and Yu (1966). Experiments were conducted at different ERs (adjusting the flow rates of air and N2) and temperatures whilst keeping the same fluidisation condition (fluidisation velocity 4.2 times the minimum one). To adjust for lower ER, the flow rate of air was reduced while the N2 flow was increased and vice versa. Sieved silica sand (0.25–0.5 mm) was used as the bed material with bulk and absolute densities of 1422 kg/m³ and 2620 kg/m³, respectively. To avoid accumulation of ash in the bed (which would distort the results, due to the potential catalytic activity of certain compounds in the fuel ash), 1.06 kg of fresh silica sand was introduced into the reactor on each experimental day.

In order to ensure stable, representative conditions for the calculation of the performance parameters, each test lasted at least 1.5 h. Three different ER levels were tested at each temperature (tests 1–3 at 700 °C and tests 5–7 at 750 °C). Due to time limitations, only one ER was tested at 725 °C (test 4). No sign of agglomeration was observed at this point. The tests at 750 °C and ERs of 0.17 and 0.21 were successfully completed (5, 6) however, during the final test at an ER of 0.25 (test 7), fluctuations of bed temperature and pressure indicated bed agglomeration. A summary of the experimental tests are illustrated in Table 2.

2.4. Measurement methods

Continuous online measurement of product gas composition was carried out by an ABB gas analyser (CO, CO2, CH4, H2, O2) and Varian micro-GC analysis (Ar/O2, Ne, CO, CO2, CH4, C2H2, C2H4, C2H6, C6H6, C7H8, H2S, and COS). The micro-GC measurements took place continuously at 4 min intervals. Neon (10 ml/min) was added as a tracer gas to measure the flow rate of dry product gas enabling the calculation of carbon conversion, gas yield and cold gas efficiency. The product gas flow rate was calculated according to the formula proposed by Pandey et al. (2016).

The solid phase adsorption (SPA) method was employed for the tar sampling. The SPA protocol coupled with GC detection offers reliable measurement of phenolic and 2–5 rings PAH tar compounds (Horvat et al., 2016b; Rabou et al., 2009). However, SPA is not ideally suitable for the detection of hydrocarbons too heavy to pass through GC instrument. Notable deviations were also observed during the measurement of light hydrocarbons such as benzene and toluene. This may be attributed to their high volatility making these compounds difficult to trap on the solid sorbent (Padban et al., 2000). Three SPA samples were taken for each test.
condition. 100 ml of dry product gas was withdrawn from the SPA sampling port with an automatic syringe pump. The amount of total GC-detectable tar as well as the amount of each individual tar compound is expressed as an average of the three repetitive measurements. The SPA tar samples were taken in 2 min intervals where the tar vapours either adsorbed or condensed on 500 mg of amino propyl silica sorbent. These tars were subsequently desorbed from the amino phase by the addition of 3 × 600 ml of dichloromethane before being analysed by gas chromatography.

An Agilent 7890A GC coupled with a triple-axis MSD 5975C was used for identification of the most abundant tar compounds. A Thermo Scientific Trace 1310 GC with a flame ionisation detector (GC-FID) was used for tar quantification. Tert-butylcyclohexane as a function of temperature at a constant ER (0.21). The concentration of the COS is very small and showed almost a

Table 2
Process conditions of the experimental tests.

| Type of feedstock | PL | Test number | 1 | 2 | 3 | 4 | 5 | 6 | 7 |
|-------------------|----|-------------|---|---|---|---|---|---|---|
| Feedstock flow rate (kg/h, a.r) | | 0.548 | 0.21 | 0.25 | 0.21 | 0.17 | 0.21 | 0.25 |
| Equivalence ratio, ER (-) | 0.17 | 0.21 | 0.25 | 0.21 | 0.17 | 0.21 | 0.25 |
| Air flow rate (NI/min) | 6.05 | 7.6 | 9.08 | 7.6 | 6.05 | 7.6 | 9.08 |
| Nitrogen flow rate (NI/min) | 5.95 | 4.4 | 2.92 | 4.4 | 5.95 | 4.4 | 2.92 |
| Minimum fluidisation velocity Umf (m/s) | 0.033 | 0.033 | 0.033 | 0.033 | 0.033 | 0.033 | 0.033 |
| Superficial fluidisation velocity U (m/s) | 0.138 | 0.138 | 0.138 | 0.138 | 0.138 | 0.138 | 0.138 |
| Gasifier temperature (°C) | 700 | 700 | 700 | 725 | 750 | 750 | 750 |

3.1. Composition of the product gas

Fig. 2(a) presents the composition of the major gas components as a function of temperature at a constant ER (0.21). The concentrations of H2, CO and CH4 increase with temperature, while the CO2 content shows the opposite trend. These tendencies stem from the fact that higher temperatures favour char gasification reactions (C + H2O <=> CO + H2, C + CO2 <=> 2 CO). CH4 is mainly evolved during the devolatilisation process. Therefore an increase in CH4 concentration at higher temperatures might indicate a larger extent of devolatilisation and tar decomposition into lighter molecules such as CH4. Taupe et al. (2016) reported that at higher temperatures the hydrogen content rises because oxygen reacts preferably with carbon forming CO2 rather than water. On the other hand the decrease in CO2 concentration can be attributed to the Boudouard equilibrium (C + CO2 <=> 2CO). The results obtained are in line with the relevant literature (Nilsson et al., 2016; Pandey et al., 2016; Taupe et al., 2016). The evolution of minor gas compounds is presented in Fig. 2(b). C2H4 shows an increasing trend with rising temperature. The decline in ethane (C2H6) concentration may be the result of thermal reforming into C2H2 and CH4 at elevated temperatures (Pandey et al., 2016; Taupe et al., 2016). C2H6 increases slightly with temperature, whilst the concentration of C2H2 shows the opposite trend. The increase in C2H6 concentration may be attributed to the conversion of phenols and toluene via demethylation (Dufour et al., 2011; Horvat et al., 2016c). C3H8 is a thermally stable compound. For its decomposition, an adequate gas residence time and temperatures above 1100°C are required (Van Paasen and Kiel, 2004). The results of minor gas compounds are in agreement with Xue et al. (2014) where the authors investigated the gasification of raw and torrefied miscanthus × giganteus at temperatures between 660 and 850°C and an ER of 0.18–0.32. Sulphur is present in the gas phase mainly in the form of H2S and COS. There are likely traces of other S compounds, such as thiophenes and mercaptans present in the gas, but those were not measured during the tests. The H2S increases with temperature while the concentration of the COS is very small and showed almost a negligible change with temperature, hence it is not reported in the graph.

3.2. Gas yield, CCE, CGE, and LCV

Fig. 3(a) shows gas yield and CCE as a function of temperature at a constant ER (0.21). Gas yield is reported on a N2 and dry ash free basis in order to ascertain the actual gas production without any
dilution effects. A gas yield increment of 33% (from 0.93 to 1.24 N₂ free m³/kg feedstock-daf) correlates with elevated temperature. Higher temperature favours the breakdown of molecular bonds (i.e. char conversion and release of volatiles). CCE also rises with temperature from 67% to 85% (an increase of 27%). It should be mentioned that for the experiment conducted at 750 °C and an ER of 0.21, some extraction of bed material took place prior to the test due to ash accumulation in the bed, which may underestimate the carbon conversion. In industrial gasifiers bed extraction usually takes place in order to prevent agglomeration phenomenon due to the build-up of alkaline metals contained in the ash (Nilsson et al., 2016).

Fig. 3(b) depicts the effect of temperature on LCV and CGE at a constant ER of 0.21. The LCV of the product gas rises by 24% ranging from 3.4 MJ/Nm³ to 4.2 MJ/Nm³ as the temperature increases. It is noteworthy to mention that the highest LCV at 750 °C (4.2 MJ/m³) doesn’t exceed the limit of 4.71 MJ/m³ reported to be suitable for internal combustion engine applications (Kim et al., 2013). However, the sum of all tar content represented solely as naphthalene gives a LCV of 5.85 MJ/Nm³. Similarly, Arena and Di Gregorio (2014) in their study on gasification of industrial plastic wastes reported a significant increase in the LCV of the product gas when adding up the energy stored in the tar (i.e. naphthalene). Therefore it is evident that, when tar is removed from product gas, its calorific value reduces significantly. CGE rises considerably with temperature, reaching the value of approximately 60% at the highest tested temperature. The explanation stems from the fact that both gas yield and LCV increase with temperature as described above.
3.3. Tar content and composition

The identified tar compounds in this work together with their retention times are given in Table S2 (supplementary file). Compound classification is based on the system proposed by Milne et al. (1997). Compared to typical lignocellulosic biomass, PL is expected to give lower tar yields due to the lower lignin content in PL with respect to wood. Lignin is considered as a tar precursor leading to the formation of higher amount of GC-detectable tar and PAHs (Horvat et al., 2016c; Yu et al., 2014). Furthermore, high alkali and alkali earth metal content (e.g. calcium, magnesium, sodium) in PL ash should catalyse tar cracking reactions. The tar composition of PL feedstock also varies with respect to lignocellulosic feedstock. In particular the high nitrogen content found in waste feed, excreta, and feathers, leads to the formation of nitrogen-containing hydrocarbons (pyridine, pyrrole, methyl pyridine).

The evolution of total GC-detectable tar and associated tar groups as a function of temperature is presented in Fig. 4 (see Tables S3–S5 in supplementary file). Total GC-detectable tar accounts for ~1 wt% of the initial dry and ash free feedstock. For the temperature range tested, the total GC-detectable tar decreased by 24% (from 5.6 to 4.25 gtar/kgfeedstock-daf). Detected but not identified tar compounds account for 20–30% of total GC-detectable tar. The yield of secondary tar dominates the tar groups while alkyl tertiary tar is the least abundant category over the entire range of tested temperatures. The alkyl tertiary tar group evolves at 750–850 °C as an intermediate between secondary and PAH tertiary tar. At temperatures between 850 and 950 °C, alkyl tertiary tar reforms into unsubstituted PAHs (Horvat et al., 2016c; Yu et al., 2014). Furthermore, high alkali and alkali earth metal content (e.g. calcium, magnesium, sodium) in PL ash should catalyse tar cracking reactions. The tar composition of PL feedstock also varies with respect to lignocellulosic feedstock. In particular the high nitrogen content found in waste feed, excreta, and feathers, leads to the formation of nitrogen-containing hydrocarbons (pyridine, pyrrole, methyl pyridine).

The dew points were calculated using an online tool developed by the ECN (“Tar dew point”) to be between 101 and 105 °C and show minor effects of tested temperatures on its values. However, the high tar dew points confirm the need for gas cleaning if the gas is to be used in e.g. internal combustion engines, gas turbines or synthesis processes. The dominant factor determining tar dew point is the yield of the PAH compounds in the product gas. The dew point of PAHs correlates with their molecular mass and concentrations in the product gas. Thus, PAH growth amplifies the risk of tar condensation on the cold surfaces of the gasifier.

Nitrogen-containing hydrocarbons show different thermal behaviour. Pyridine increases steadily while the concentration of pyrrole reduces significantly. Methyl pyridine shows a very small decrease, ranging approximately to 0.12 g/kgfeedstock-daf. The opposite trends for pyridine and pyrrole may be attributed to the higher thermal stability of the former compound. Zhao et al. (2010) investigated the transformation of nitrogen during pyrolysis and combustion of coal in a flow reactor. Pyridine and pyrrole were considered as model compounds while measuring the amount of HCN and HCN in order to identify their thermal stability. The findings revealed that pyridine appeared to be more stable generating high amounts of HCN at 825 °C, while the respective temperature for pyrrole was at 775 °C.

Methyl naphthalene is the most abundant of the alkyl tertiary tar compounds, indicating decreased yields in the tested temperature range. Biphenyl yield remains constant, while 2-ethenyl naph-
thalene reduces slightly. Dufour et al. (2011) reported that reforming of methyl naphthalenes into naphthalene and acenaphthylene occurs at 800–900 °C. Horvat et al. (2016c) observed a peak in the yield at 800 °C for alkylated naphthalenes. Steady increase of biphenyl yield was observed at the temperature range 660–850 °C using torrefied feedstock. Biphenyl may act as an intermediate in the polymerisation pathway promoted by higher temperatures (Berrueco et al., 2014).

Tertiary PAH tar evolution shows an upward trend with rising temperature. The findings are in line with other researchers who observed that the production of PAH is driven by increased temperature (Horvat et al., 2016c; Berrueco et al., 2014). Yu et al. (2014) argued that at 850 °C the composition of tar consists mainly of PAHs. Naphthalene is the most abundant PAH compound ranging between 0.43 and 0.55 g tar/kg feedstock-daf. The low reforming rate of naphthalene is explained by its thermal stability. Naphthalene formation initiates either by the decomposition of heavier PAHs or by polymerisation reactions (Devi et al., 2005a; Nilsson et al., 2016). In this study the relatively low operating temperature resulted in low production of PAHs and the dominance of secondary tar.

### 3.4. Agglomeration issues

Agglomeration is a crucial phenomenon as regards the operational stability of fluidised bed gasifiers. The occurrence of bed agglomeration results in de-fluidisation conditions leading to local temperature and pressure deviations and consequent shutdown of the gasifier. The reason behind this phenomenon is the presence of inorganic compounds (P, K, Na, etc.) in the feedstock ash characterised by low melting temperatures. Agglomeration is exacerbated when silica sand is used as bed material as the reaction between silica and potassium may form low melting potassium silicate. Prevention or mitigation of such formation may be realised with the addition of calcium forming calcium phosphate instead with higher melting temperature (Pandey et al., 2016). Agglomeration in the first minutes of test 7 resulted in the interruption of the fluidisation conditions and feeding was stopped. After 10 min the feeding started again in order to investigate if the fluctuations appear again and although for 10 min (3–12 min in the Fig. S1 in supplementary file) the gasifier seemed to operate smoothly, deviations of pressure and temperature occurred again leading to the termination of the experiment.
4. Conclusions
The effect of temperature on the gasification behaviour of PL was experimentally studied in a lab-scale fluidised bed reactor. Gas yield, LCV and CGE showed an upward trend with increasing temperature from 700 °C to 750 °C. Although the LCV of 4.2 MJ/m³ is low, if the presence of tar in the gas stream is taken into account represented as naphthalene, the value rises to 5.8 MJ/Nm³, a fact that is useful if the product gas is destined directly for combustion without prior cleaning. Due to the high ash content of PL comprising of inorganic components characterised by low melting temperatures, agglomeration occurred at a temperature as low as 750 °C and ER 0.25. Total GC-detectable tar yield is expected to be lower compared to lignocellulosic biomass, due to both the low content of lignin in PL and the presence of inorganic compounds which act as tar reducing catalysts. Total GC-detectable tar yield decreased with temperature (from 5.6 g/tar/kgfeedstock-daf at 700 °C to 4.25 g/tar/kgfeedstock-daf at 750 °C). For the temperature range tested, secondary tar was the dominant category among the tar groups consisting of oxygen-containing phenolic compounds, substituted one-ring aromatics and nitrogen-containing hydrocarbons.

Despite the fact that the PL is considered a low-quality fuel, the present study reveals its potential as alternative energy resource for onsite (farm) energy generation. Nevertheless, increasing the operating temperature to improve the gasification performance while avoiding agglomeration conditions is a crucial aspect that needs to be further addressed. Addition of minerals in the fuel intake (use of additives) or mixing PL with conventional woody biomass that could change the ash composition are two viable options that can be further investigated as potential measures. On contrary to the lab-scale reactors that operate under auto-thermal conditions, industrial scale gasifiers operate in an auto-thermal mode implying that the reactor temperature is regulated by the ER. Therefore, modelling of the gasification process is proposed for future work in order to investigate if any of the conducted tests are close to auto-thermal conditions, which will provide the possibility to scale-up the experimental findings to an industrial gasifier level.

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Appendix A. Supplementary material
Supplementary data to this article can be found online at https://doi.org/10.1016/j.wasman.2019.09.014.

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