Ignition behaviour of coal and biomass blends under oxy-firing conditions with steam additions

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Abstract

The ignition behaviour of coal and biomass blends was assessed in air and oxy-firing conditions in an entrained flow reactor. Four coals of different rank, an anthracite, a semianthracite and two high-volatile bituminous coals, were tested in air and O₂/CO₂ (21-35% O₂) environments. For all the coals, a deterioration in ignition properties was observed in the 21%O₂/79%CO₂ atmosphere in comparison with air. However, the ignition properties were enhanced when the oxygen concentration in the O₂/CO₂ mixture was increased. Coal and biomass blends of a semi-anthracite and a high-volatile bituminous coal with 10 and 20 wt% of olive residue were also used in the ignition experiments under air and oxy-firing conditions. The ignition behaviour of the coals improved as the additions of biomass increased both in air and oxy-firing conditions. In particular, the effect of biomass blending was more noticeable for the ignition of the high rank coal. Since industrial oxy-coal combustion with a wet recycle would result in higher concentrations of H₂O(ν), the effect of steam addition on ignition behaviour was also studied. A worsening in ignition behaviour was observed when steam was added to

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the oxy-fuel combustion atmospheres, although an increase in the steam concentration from 10 to 20% did not produce any significant difference in the ignition characteristics of the fuels.

**Keywords:** biomass; coal; ignition behaviour; oxy-combustion, steam

### 1. Introduction

The use of coal in power plants generates a large amount of CO\(_2\) which is the chief cause of global climate change. A diverse power generation portfolio including Carbon Capture and Storage (CCS) technologies and renewable energies is needed to reduce atmospheric CO\(_2\) to below 1990 levels\(^{1,2}\). During oxy-coal combustion, coal is burnt in a mixture of oxygen and recycled flue gas (RFG), mainly CO\(_2\) and water vapour, to yield a rich CO\(_2\) stream, which after purification is ready for sequestration\(^3\). In addition, biomass is a source of energy which is considered carbon neutral as the carbon dioxide released during its combustion is recycled as an integral part of the carbon cycle. The combination of oxy-coal combustion with biomass co-firing can help to increase CO\(_2\) capture efficiency\(^4\). However, the successful implementation of the oxyfuel technology in pulverised coal boilers depends on fully understanding the differences that may result from replacing N\(_2\) with a mixture of CO\(_2\) and water vapour in the oxidiser stream. In oxy-firing conditions, due to the higher concentrations of CO\(_2\) and H\(_2\)O\(_{\text{v}}\), compared to conventional air combustion, several aspects such as heat transfer, flame ignition, pollutant formation and volatiles and char combustion are affected\(^5\)\(^-\)\(^9\). The use of biomass in existing coal power plants requires only minor modifications compared to
the construction of new biomass-only fired power plants, making the co-firing of coal and biomass an easy and cheap way to obtain biomass energy\textsuperscript{10}.

The application of the oxy-combustion technology has been implemented at a higher scale in Vattenfall’s 30 MW\textsubscript{th} oxyfuel pilot plant in Schwarze Pumpe, Germany\textsuperscript{11}, which was constructed in order to investigate the oxyfuel firing process. The combustion investigations have focused on the radiation heat flux behaviour, NO\textsubscript{x} formation, as well as combustion performance and reaction rates between air and oxyfuel operation modes. Details on the operation with three different burners (one combined jet-swirl burner and one swirl burner both from Alstom, and one swirl DST-burner delivered by Hitachi) have been provided and it was found that the operation of the oxyfuel boiler with the three burners tested so far has proven to be very reliable and a good flame ignition and high stability over the entire load range has been achieved\textsuperscript{12}.

The recycling of flue gas and the injection of the oxygen add more complexity to the design and operation of the oxyfuel burners and boilers\textsuperscript{13}. Under oxyfuel conditions, the burner geometry has to be modified in order to achieve a stable flame attached to the burner quarl at oxygen contents in the O\textsubscript{2}/RFG mixture close to that in air. In this regard a series of test runs was performed at the oxycoal test facility at RWTH Aachen University with the aim to achieve an experimentally confirmed database needed for development of a swirl burner able to operate at a wide range of O\textsubscript{2} concentrations (from 18 to 34\% vol. O\textsubscript{2}) under oxy-firing conditions. Thus, a new burner concept based on aerodynamic stabilization of an oxyfuel swirl flame has been developed\textsuperscript{14}.

The ignition of solid fuel particles is an important preliminary step in the overall combustion process due to its influence on the stability, shape and length of the flame, and on the formation of pollutant. In practice, the ignition behaviour of solid fuels may
be decisive for identifying the optimal location for injecting them into industrial pulverised fuel burners. The ignition and combustion behaviour of pulverised coal particles are not inherent properties of the coals, as they are dependent on the operating conditions\textsuperscript{15}. In the oxy-fuel combustion of pulverised coal, poor ignition quality has often been observed during pilot-scale burning trials when operating with substantial flue gas recirculation\textsuperscript{16}. Several efforts have been made recently to understand the fundamentals of ignition (i.e., particle ignition, flame propagation and flammability) in oxy-firing conditions when designing combustion systems\textsuperscript{17,18}.

The present work studies the influence of fuel type, CO\textsubscript{2} dilution and oxygen concentration on the temperature and ignition mechanisms for a wide number of coals and coal/biomass blends (up to 20%wt biomass). It needs to be appreciated that, when blending different fuels, certain aspects such as ignition behaviour, burnout or NO emissions cannot always be estimated from the behaviour of the individual fuels\textsuperscript{19}. As Smart et al.\textsuperscript{20} have pointed out, another important aspect to consider is the effect of wet recycling in oxy-firing conditions on coal ignition and flame stability. In the present work, the effect of adding 10 and 20% of steam on the ignition characteristics, under air and oxy-firing conditions, was also studied.

2. Experimental

2.1. Materials

Four coals of different rank were used in this work: an anthracite from Cangas del Narcea, in Asturias, Spain (AC), a semi-anthracite from the Hullera Vasco-Leonesa in León, Spain (HVN), a South African high-volatile bituminous coal from the Aboño power plant in Asturias, Spain (SAB), and a washed coal supplied by the Batán coal
preparation plant in Asturias, Spain (BA). A biomass, olive residue (OR) was also employed. This biomass is the wet solid residue that remains after the process of pressing and extraction of the olive oil. The coal and biomass samples were ground and sieved to obtain a particle size fraction of 75-150 µm. The proximate and ultimate analyses together with the high heating values of the samples are presented in Table 1.

2.2. Experimental device and procedure

The ignition characteristics of the coals and coal/biomass blends were studied in an entrained flow reactor (EFR), which has been described in detail elsewhere. Briefly, the reactor has a reaction zone 140 cm in length and an internal diameter of 40 mm. It is electrically heated and is capable of reaching a maximum temperature of 1100 ºC. Fuel samples were introduced through a cooled injector before entering the EFR reaction zone. The gases were preheated up to the reactor temperature before being introduced into the EFR, where they passed through two flow straighteners. The reaction products were quenched by aspiration in a stream of nitrogen using a water-cooled probe. The probe was inserted into the reaction chamber from below. Particles were removed by means of a cyclone and a filter. The exhaust gases were monitored using a battery of analysers (O₂, CO, CO₂, SO₂, and NO).

During the ignition tests, the reactor was heated at 15 ºC min⁻¹ from 400 to 900 ºC. The gas flow used in these tests ensured a particle residence time of 2.5 s. Air (21%O₂/79%N₂) and three binary mixtures of O₂/CO₂ (21%O₂/79%CO₂, 30%O₂/70%CO₂ and 35%O₂/65%CO₂) were employed to study the ignition characteristics of the coals and coal/biomass blends. Also, several ternary mixtures of O₂/N₂/H₂O(ν) and O₂/CO₂/H₂O(ν) were employed. The addition of 10 and 20% of steam
was evaluated for all the air and oxy-fuel combustion atmospheres as a substitute for N₂ or CO₂ in order to study the effect of the wet recirculation of flue gas on coal ignition properties.

3. Results and discussion

3.1. Effect of the coal rank

Faúndez et al. 21 have stated that ignition is characterised by a rapid decrease in CO production, a significant consumption of O₂, and an increase in the production of CO₂ and NO. Prior to ignition, at low temperatures, the production of CO increases due to the release of volatiles and incomplete char combustion. The production of CO₂ and NO shows only a slight increase, and there is some O₂ consumption due to the evolution and subsequent combustion of coal volatiles at low temperatures. The criterion for determining the ignition temperature was based on the first derivative temperature curves of the gases produced. The ignition temperature was taken as the temperature at which the first derivative curve, normalised from the maximum derivative value, reached an absolute value of 10% 21. The ignition temperatures, derived from derivative curves of the gases, for coals AC, HVN, SAB and BA are shown in Table 2. As can be seen, coals SAB and BA tend to ignite at lower temperatures than coals AC and HVN. This may be due to their higher volatiles content (which enhances subsequent char combustion), and to their higher reactivity 22.

Wall et al. 23 tracked the changes in gas composition during the ignition of pulverized coal in air in a laboratory scale drop-tube reactor, and associated these changes with the homogeneous ignition of volatiles and heterogeneous char combustion. In general, two types of mechanisms have been observed for coal particle ignition 24: gas mode ignition
(ignition of the volatiles in an enveloping flame that surrounds a devolatilising char particle), heterogeneous mode ignition (which often signifies char ignition) or a combination of both. The ignition mechanism is closely related with the particle combustion mode. In a recent paper by Khatami et al.\textsuperscript{25}, the authors employed the term \textit{two-mode} combustion to signify events where the gas-phase (homogeneous) combustion of volatiles in an enveloping flame that surrounds a char particle, is distinct from the ensuing heterogeneous combustion of the solid char, as it occurs in homogeneous ignition. On the other hand, the term \textit{one-mode} combustion was used to refer to events where either (i) the combustion of devolatilised char takes place or (ii) the combustion of the volatiles in the proximity of the char surface occurs with, presumably, simultaneous burning of the char occurs, as in heterogeneous ignition. In the present paper, the ignition mode was elucidated from the gas evolution profiles. Examples in the air atmosphere are provided for a better comparison between coal samples.

The evolution of the gases during the ignition tests in air conditions for anthracite coal AC is shown in Fig. 1. The CO concentration increases up to a value of \textasciitilde750 ppm. At around 760 °C there is a reduction in CO concentration, which is accompanied by a drastic reduction in O\textsubscript{2} and a sudden increase in NO and CO\textsubscript{2} corresponding to the ignition of the char. This is confirmed by the continued decrease in CO after the ignition event. The coal ignites heterogeneously as a result of the direct attack of oxygen on the surface of the char. For the ignition mode of various coal ranks, see also Fig. 2. As can be seen in the cinematographic observations, most of the AC coal particles burn heterogeneously, there being no evidence of the burning of the volatiles.
The gas emissions during semi-anthracite coal HVN ignition in air can be seen in Fig. 3. Above 400 °C there is an increase in the CO concentration due to coal devolatilisation (the major devolatilisation products are CH₄, CO and CO₂). Two changes in the CO profile and its derivative value can be observed: an initial decrease in CO concentration takes place at around 625 °C, which corresponds to the combustion of volatiles. The second change involves another decrease in CO concentration which takes place at around 670 °C. This is accompanied by a sudden decrease in O₂ concentration and an increase in NO and CO₂. These latter events correspond to char ignition, which is confirmed by the constant increase in the CO₂ produced after the ignition event. This coal presents the most difficult mechanism to be elucidated, i.e., it seems to be homogenous since the ignition of volatiles and char took place sequentially. Coal HVN is a physical blend of approx. 90% anthracitic and 10% low volatile bituminous coal from the same mine. For the ignition mode of semi-anthracite HVN coal, see also Figure 2. As observed in the cinematographic records, some of the semi-anthracite HVN particles burn heterogeneously, but for other particles a small surrounding flame is observed corresponding to the combustion of volatiles. This enveloping flame burns up prior to the heterogeneous combustion of the char. However, in the EFR it is not possible to appreciate the ignition of single particles, it can only be seen that, in global, the stream of HVN particles burned homogenously (although much less volatile matter was released in comparison with hvb coals, see Fig 4).

The gas evolution profiles for the SAB ignition tests in air are shown in Fig 4. Above 400 °C there is a significant increase in CO concentration (due to the higher volatile matter content of SAB, the amount of CO released is much higher than for anthracitic coals). Also a continuous decrease in O₂ concentration is observed, which suggests that
part of the volatiles released are oxidised. At a temperature of 530 °C the CO concentration starts to decrease, which suggests that coal devolatilisation has ended, and that more CO is being consumed than formed. From Fig. 4 it can be inferred that the ignition of the char takes place at around 545 °C (i.e., there is a sudden decrease in the concentration of CO and O2, and an increase in NO and CO2). This suggests that the ignition of high-volatile coals takes place via a homogeneous mechanism, with the sequential ignition of volatiles and char. However, the ignition delay between the extinction of the volatiles and char combustion is much shorter than in the case of semi-anthracite coal HVN. For the ignition mode of bituminous coal SAB, see also Figure 2. As can be seen from the cinematographic records, the combustion of coal SAB includes particle devolatilisation with ignition and combustion of the volatiles in a flame that surrounds the particle, followed by the ignition and combustion of the resulting char.

Coal BA is a high-volatile bituminous coal whose ignition mechanism is also homogeneous. Since its gas evolution profiles are very similar to those of coal SAB, they are not shown in this paper.

3.2. Effect of the O2/CO2 atmosphere

In order to evaluate the effect of the presence of CO2 in large concentrations, ignition tests were conducted in both O2/N2 and O2/CO2 environments. As can be seen in Table 2 higher ignition temperatures are required when N2 (21%O2/79%N2) is replaced by CO2 (21%O2/79%CO2). Stivers et al.26 and Khatami et al.27 observed a delay in ignition in an O2/CO2 environment compared to an O2/N2 environment with identical O2 concentration. They attributed the longer ignition delay partly to the effect of the volumetric heat capacity of the gas mixtures. The temperature rise during ignition is
inversely proportional to the heat capacity of the surrounding gas and, since the heat
capacity of CO₂ is higher than that of N₂, a reduction in gas temperature occurs.
However, the heat capacity and temperature of the surrounding gas are not the only
factors that affect ignition properties; the oxygen concentration, the heating rate of the
gas and devolatilisation rates of particles, and the coal volatiles content also have a
considerable influence. It should be noted that in this study the highest increase in ignition temperature was
observed for coal BA, the coal with the highest volatile matter content, due to the lower
mass diffusivity of the volatiles in the CO₂ mixture. It can also be observed that for
semi-anthracite and bituminous coals HVN, SAB and BA and for oxygen
concentrations up to 30% or 35%, the ignition temperature is lower than in air, even
though the heat capacity of the gas atmospheres with O₂ concentrations up to 30-35%
are still higher than the heat capacity of the air. In the case of anthracite coal AC—which
has the lowest volatile matter content—increasing the oxygen concentration to 35% was
not enough to compensate for the negative effects of CO₂ on ignition temperature.
From the values of the ignition temperatures it is difficult to determine whether the
worsening of the ignition properties under oxy-fuel conditions is due to the reactions
affecting the char, the reactions involving the volatiles, changes in heat transfer or a
combination of all three factors. For this reason the ignition mechanism of the different
coals was inferred from the evolution curves of the gases.
The gas evolution during the ignition of anthracite coal AC in two of the oxy-firing
conditions studied (21%O₂/79%CO₂ and 35%O₂/65%CO₂) is shown in Fig. 5. The
ignition mechanism is the same as that observed under air-firing conditions; i.e., the
char ignites heterogeneously due to the direct attack of oxygen. The only difference
between the air and oxy-fuel atmospheres is the larger amount of CO formed, which may be attributed to char-CO₂ reactions. Pyrolysis experiments for the coals studied in the N₂ and CO₂ atmospheres at 1000 °C have been carried out previously in the EFR²². The results for volatile yields are presented in Table 3, and they show that the volatile yield is enhanced in a CO₂ atmosphere, as the CO₂ reacts with the resulting chars. The high amounts of CO which persist as a thick protective sheath, even with oxygen contents of 35%, prevent particle ignition.

The gas evolution profiles for semi-anthracite coal HVN in oxy-fuel conditions are shown in Fig. 6. An increase in CO concentration occurs due to the devolatilisation of HVN at temperatures above 400 °C. In the 21%O₂/79%CO₂ atmosphere the combustion of part of these volatiles takes place above 625 °C, as in air-firing conditions. The ignition of the char (i.e., a marked reduction in O₂ concentration and a marked increase in CO₂ and NO) occurs at around 725 °C. The ignition mechanism is the same as that of air ignition. However, higher CO concentrations are obtained. These higher CO concentrations contribute to the deterioration in ignition properties, via the formation of a persistent cloud around the particle that prevents the oxygen from gaining access to the surface of the particle. As other authors have observed²⁵,²⁹ in a 21%O₂/79%CO₂ atmosphere, the volatiles released remain partially unburnt and form a thick cloud of volatiles. Also, higher concentrations of CO are formed, partly due to the incomplete combustion of the volatiles, and partly as a result of char gasification by CO₂. To compare the intensity and brightness of the burning coal particles in air and CO₂ atmospheres see also Figure 2. From the cinematographic records, when N₂ is replaced by CO₂ for the same oxygen concentration, the burning particles appeared dim and blurry, which is indicative of slow oxidation. The brightness and intensity of the coal
combustion increases drastically with oxygen in the O₂/CO₂ environments. However, the combustion images of the 30%O₂/70%CO₂ atmosphere resemble those of air. Since the gas evolution profiles corresponding to 30%O₂/70%CO₂ and 35%O₂/65%CO₂ are similar, only those for the 35%O₂/65%CO₂ atmosphere are shown in Fig. 6. At temperatures above 400 °C the process of coal devolatilisation starts with the consequent increase in CO concentration. As in the case of the N₂ and CO₂ atmospheres with a 21% oxygen content, the CO starts to oxidize above 625 °C. The ignition of the char occurs at 642 °C. Thus, in these cases ignition also takes place with the sequential ignition of volatiles and char. However, the time delay between volatiles and char combustion decreases as the oxygen concentration increases and ignition occurs at a lower temperature than in air. Khatami et al.²⁵,²⁷ have also observed that the ignition delay in O₂/CO₂ atmospheres becomes smaller as O₂ increases. The ignition and combustion of volatiles provide extra heat that enhances the ignition of the char. However, this effect is not observed for coal AC due to its low volatile matter content. Also, when there is sufficient oxygen, the combustion of CO to form CO₂ provides extra heat.

The gas evolution of bituminous coal SAB during its ignition in oxy-firing conditions is shown in Fig. 7. High amounts of CO are released during coal devolatilisation, which are later oxidised at temperatures of around 545 °C. Subsequently, char ignition takes place. The ignition mechanism is therefore homogeneous with the sequential ignition of volatiles and chars. However, the time delay between them is much shorter than in the case of semi-anthracite coal HVN, and becomes even shorter with increasing oxygen concentrations. Also larger amounts of CO are produced under oxy-firing conditions. It can be observed in Table 3 that the volatile yield in CO₂ for coal SAB is much higher.
than in N\textsubscript{2} in comparison with coals AC and HVN. These findings are in accordance with those reported by Zhang et al.\textsuperscript{29} who have found that the replacement of N\textsubscript{2} by CO\textsubscript{2} enhanced coal particle pyrolysis prior to ignition, as CO\textsubscript{2} reacted with the resulting char to form additional combustible gases, i.e., CO, in the vicinity of the particle. The cinematographic records showed an increase in both char and volatile burning times in the 21\%O\textsubscript{2}/79\%CO\textsubscript{2} atmosphere in comparison with air-firing conditions. Also, there is a decrease on burning times in the O\textsubscript{2}/CO\textsubscript{2} environments with increasing oxygen concentrations, and a decrease in the time delay between the extinction of the volatiles and char ignition.

Bituminous coal BA experiences the highest ignition delay, when N\textsubscript{2} is replaced by CO\textsubscript{2} for the same oxygen concentration, of all the coals under study. As for coal SAB, during its ignition in 21\%O\textsubscript{2}/79\%CO\textsubscript{2}, the CO concentration remains very high over a wide range of temperatures, preventing the oxygen from gaining access to the surface of the particle and causing a big delay in ignition. In a recent paper by Khatami et al.\textsuperscript{27} the authors observed coal and char particle ignition in O\textsubscript{2}/N\textsubscript{2} and O\textsubscript{2}/CO\textsubscript{2} atmospheres and found that in a N\textsubscript{2} atmosphere the presence of volatiles accelerated the ignition process, as the coals ignited faster than the chars, whereas in CO\textsubscript{2} the chars ignited faster than the coals, because the presence of a thick cloud of volatiles appeared to have impeded the ignition process. In the present study, coal BA has the highest content of volatiles, and therefore it will release more CO than the other coals. Also, as can be seen from Table 3, it shows the highest volatile yield in a CO\textsubscript{2} atmosphere (not shown). When the oxygen concentration is increased to 30\% or 35\% the oxidation of CO to CO\textsubscript{2} is favoured and, as a consequence, ignition takes place at lower temperatures.
3.3. Effect of the addition of biomass

The effect of blending coals and biomass on ignition behaviour was studied under air and O₂/CO₂ (21-35% O₂) conditions. Two coals of different rank, the semianthracite HVN and the high volatile bituminous coal SAB, were blended with the olive residue OR. The ignition temperatures are shown in Table 4. It can be observed that the addition of olive residue, OR, causes a significant reduction in the ignition temperature of both coals in all the atmospheres studied. This decrease is proportional to the amount of biomass in the blend and is more pronounced for the HVN-OR blends.

As can be seen in Fig 8, the ignition mechanism of the HVN-OR blends in air is homogeneous (i.e., with the sequential ignition of char and volatiles). For the case of oxy-fuel combustion, the addition of biomass does not affect the ignition mechanism because it remains homogeneous, so their gas evolution profiles are not shown in this paper.

In any case, a significant reduction in ignition temperature is observed in both the air and oxy-fuel atmospheres when coal is blended with biomass. Biomass is a highly reactive fuel and has a much higher volatile matter content than coal. The biomass OR will release far more volatiles when it is devolatilised than coal HVN, and this is reflected in the higher CO concentrations observed in the coal/biomass blends. As mentioned before, the ignition of the volatiles and char takes place sequentially. For coal HVN the ignition of the volatiles occurs at around 600 °C (see Fig. 3), whereas for blends 90HVN-10OR and 80HVN-20OR it takes place at around 530 °C and 510 °C, respectively; whereas the combustion of char occurs at 636 °C and 575 °C. In summary, the addition of increasing quantities of biomass leads to a reduction in the ignition temperatures and in the delay between the ignition of the volatiles and char.
From Fig. 9 it can be seen that the ignition mechanism of the SAB-OR blends in air is homogeneous, as in the case of the individual coal SAB. The ignition mechanism for oxy-fuel conditions is also homogeneous (figures not shown). Furthermore ignition occurs at lower temperatures as the biomass content in the blends increases. For coal SAB the volatiles ignite at around 530 °C and the char at 543 °C (see Fig. 4); whereas for blend 90SAB-10OR the ignition temperatures are of 500 °C and 510 °C, respectively. In the case of blend 80SAB-20OR there is a marked reduction in ignition delay between the volatiles and char ignition almost to the point where they seem to happen simultaneously at around 461 °C.

Although there is a decrease in ignition temperatures for both char and volatiles when increasing the biomass percentage in the SAB-OR blends, these decreases are less influenced by the addition of biomass than in the case of HVN-OR blends. This suggests that the effect of the addition of biomass on the ignition temperature of coal is more marked for high rank coals. When two fuels are fired as a blend, the ignition properties of the blend may be different to those exhibited when each component is ignited individually. The ignition properties of high rank coals, which have far fewer volatiles and are less reactive than low rank coals, will be more easily enhanced by the addition of biomass. Faúndez et al. have observed that, when blending fuels with different volatile matter contents, the ignition of the higher volatile component of the blend enhances the ignition of the lower volatile component. However, when both fuels have similar volatile contents, they compete for the oxygen available. As was shown by Khatami et al. when biomass particles are burned, large volatile flames are formed. This also happens when burning low rank coals. The simultaneous burning of biomass and low rank coals leads to a competition for oxygen and in some zones oxygen
depletion will result. Consequently, the enhancement of ignition properties will be less marked than when biomass is blended with high rank coals.

In summary, the ignition mechanism of high rank and low rank coals does not change when they are blended with biomass (up to 20% by mass). However, the addition of biomass improves their ignition properties, i.e., the coal and biomass blends ignite at lower temperatures than the individual coals.

3.4. Effect of the addition of steam

In order to study the effect of wet recirculation of flue gas, ignition tests were performed under air and oxy-firing conditions with the addition of steam as a substitute for N₂ or CO₂, respectively. Two coals of different rank, a semianthracite (HVN) and a high volatile bituminous coal (BA), were chosen for the ignition experiments. The ignition temperatures are shown in Table 5. The partial replacement of N₂ or CO₂ by steam causes a slight increase in the ignition temperatures, but no significant differences are observed between the results for 10 and 20% of steam. It should be noted that in the case of coal BA no significant differences were observed when steam was added to the oxy-fuel atmosphere with 30 or 35% oxygen content.

Fig. 10 shows the gas evolution curves during the ignition of coal HVN under air and oxy-firing conditions with 20% steam addition. Similar gas evolution curves were obtained for 10% steam addition (not shown). The addition of steam does not affect the ignition mechanism of coal HVN. However, higher CO concentrations are observed with the addition of water vapour. In the atmospheres with a lower oxygen content (21%), the CO concentrations are higher and they remain higher over a wider range of temperatures. The reasons for these high CO concentrations with the addition of steam
are not yet clear. They may be partly due to unburnt volatiles. Binner et al.\textsuperscript{31,32} observed that the ignition of a volatile flame was delayed during wet coal combustion, as well as a decrease in particle temperature. The same authors also observed that steam gasification of the char could take place to some extent. As was observed for oxy-firing conditions without the addition of steam, the CO preferentially remains in the vicinity of the particle surface, forming a thick protective sheath. If the CO remains on the char surface for a long time, this will result in O\textsubscript{2} depletion on the char surface, delaying the ignition process. When the oxygen concentration is increased, the combustion of CO to form CO\textsubscript{2} is favoured and, the ignition of the coal particles is enhanced.

A similar conclusion can be drawn from the evolution of gases during the ignition of coal BA in air and oxy-firing conditions with steam addition. The ignition mechanism remains homogeneous for air and oxy-fuel conditions, both in wet and dry conditions. Also higher CO concentrations are observed with increasing steam addition.

4. Conclusions

The aim of this work was to study the ignition characteristics of coal and biomass blends in oxy-firing conditions with and without steam addition. The most important conclusions of this work are as follows:

(a) A significant increase in ignition temperature was observed when N\textsubscript{2} was replaced by CO\textsubscript{2}, for the same oxygen concentration for the four coals studied (an anthracite, a semi-anthracite and two high-volatile bituminous coals). This increase is not only due to the higher heat capacity of the background gas, but also due to the persistence of a thick cloud of volatiles around each particle which prevents its ignition. Not all the coals are affected in the same way by the
background gases. In the O₂/CO₂ with low oxygen content (i.e., 21%) coals with a higher volatile matter content experience higher ignition delays due to the larger concentrations of volatiles and CO formed around the particle. The anthracite coal ignites in a heterogeneous mode in both air and oxy-firing conditions, semi-anthracite coal partially ignite heterogeneously whereas the bituminous coals ignite in a homogeneous mode.

(b) Co-firing coal (a semi-anthracite and a high-volatile bituminous coal) and biomass results in an improvement in the ignition properties of the blend in both air and oxy-firing conditions. However, this improvement is more significant in the case of the blends with the semi-anthracite. The ignition properties of the bituminous coal seem to be less affected by the addition of biomass.

(c) A worsening of ignition properties is observed when N₂ or CO₂ is partially replaced by H₂O(v) for both semi-anthracite and high-volatile bituminous coals. Higher CO concentrations are observed when the H₂O(v) concentrations are increased. The effect of steam addition is less noticeable in atmospheres with a high oxygen content.

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References


Table 1. Proximate and ultimate analyses and high heating value of the fuel samples

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<td>Ash</td>
<td>14.2</td>
<td>10.7</td>
<td>15.0</td>
<td>6.9</td>
<td>7.6</td>
</tr>
<tr>
<td>V.M.</td>
<td>3.6</td>
<td>9.2</td>
<td>29.9</td>
<td>33.9</td>
<td>71.9</td>
</tr>
<tr>
<td>F.C. a</td>
<td>82.2</td>
<td>80.1</td>
<td>55.1</td>
<td>59.2</td>
<td>20.5</td>
</tr>
<tr>
<td><strong>Ultimate Analysis (wt.%, daf)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>94.7</td>
<td>91.7</td>
<td>81.5</td>
<td>88.5</td>
<td>54.3</td>
</tr>
<tr>
<td>H</td>
<td>1.6</td>
<td>3.5</td>
<td>5.0</td>
<td>5.5</td>
<td>6.6</td>
</tr>
<tr>
<td>N</td>
<td>1.0</td>
<td>1.9</td>
<td>2.1</td>
<td>1.9</td>
<td>1.9</td>
</tr>
<tr>
<td>S</td>
<td>0.7</td>
<td>1.6</td>
<td>0.9</td>
<td>1.1</td>
<td>0.2</td>
</tr>
<tr>
<td>O a</td>
<td>2.0</td>
<td>1.3</td>
<td>10.5</td>
<td>3.0</td>
<td>37.0</td>
</tr>
<tr>
<td><strong>High heating value (MJ kg⁻¹, db)</strong></td>
<td>29.2</td>
<td>31.8</td>
<td>27.8</td>
<td>33.1</td>
<td>19.9</td>
</tr>
</tbody>
</table>

an: anthracite; sa: semi-anthracite; hvb: high-volatile bituminous coal
db: dry basis; daf: dry and ash free bases

* Calculated by difference
Table 2. Ignition temperatures (°C) of coals AC, HVN, SAB and BA under air and O₂/CO₂ (21-35 vol.% O₂)

<table>
<thead>
<tr>
<th>Coal</th>
<th>21%O₂/79%N₂</th>
<th>21%O₂/79%CO₂</th>
<th>30%O₂/70%CO₂</th>
<th>35%O₂/65%CO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>757</td>
<td>782</td>
<td>767</td>
<td>761</td>
</tr>
<tr>
<td>HVN</td>
<td>700</td>
<td>723</td>
<td>669</td>
<td>642</td>
</tr>
<tr>
<td>SAB</td>
<td>543</td>
<td>565</td>
<td>524</td>
<td>498</td>
</tr>
<tr>
<td>BA</td>
<td>509</td>
<td>554</td>
<td>498</td>
<td>490</td>
</tr>
</tbody>
</table>
Table 3. Experimental devolatilisation yields at 1000 ºC in the EFR under N\textsubscript{2} and CO\textsubscript{2} atmospheres

<table>
<thead>
<tr>
<th>Coal</th>
<th>Volatile yield-N\textsubscript{2}</th>
<th>Volatile yield-CO\textsubscript{2}</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>3.6</td>
<td>4.2</td>
</tr>
<tr>
<td>HVN.</td>
<td>6.0</td>
<td>7.2</td>
</tr>
<tr>
<td>SAB</td>
<td>44.9</td>
<td>53.1</td>
</tr>
<tr>
<td>BA</td>
<td>49.9</td>
<td>62.2</td>
</tr>
</tbody>
</table>
Table 4. Ignition temperatures (ºC) for blends HVN-OR and SAB-OR in air and O$_2$/CO$_2$ (21-35 vol.% O$_2$)

<table>
<thead>
<tr>
<th></th>
<th>21%O$_2$/79%N$_2$</th>
<th>21%O$_2$/CO$_2$</th>
<th>30%O$_2$/CO$_2$</th>
<th>35%O$_2$/CO$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>HVN</td>
<td>700</td>
<td>723</td>
<td>669</td>
<td>642</td>
</tr>
<tr>
<td>90HVN-10OR</td>
<td>636</td>
<td>662</td>
<td>615</td>
<td>567</td>
</tr>
<tr>
<td>80HVN-20OR</td>
<td>574</td>
<td>612</td>
<td>551</td>
<td>503</td>
</tr>
<tr>
<td>SAB</td>
<td>543</td>
<td>565</td>
<td>524</td>
<td>498</td>
</tr>
<tr>
<td>90SAB-10OR</td>
<td>510</td>
<td>532</td>
<td>491</td>
<td>455</td>
</tr>
<tr>
<td>90SAB-20OR</td>
<td>461</td>
<td>478</td>
<td>444</td>
<td>425</td>
</tr>
</tbody>
</table>
Table 5. Ignition temperatures (°C) for coals HVN and BA in air and O₂/CO₂ (21-35 vol.% O₂) with steam addition (the H₂O (v) is added as a substitute of N₂ or CO₂)

<table>
<thead>
<tr>
<th></th>
<th>21%O₂/N₂</th>
<th>21%O₂/CO₂</th>
<th>30%O₂/CO₂</th>
<th>35%O₂/CO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>HVN</td>
<td>700</td>
<td>723</td>
<td>669</td>
<td>642</td>
</tr>
<tr>
<td>HVN+10%H₂O (v)</td>
<td>713</td>
<td>733</td>
<td>682</td>
<td>657</td>
</tr>
<tr>
<td>HVN+20%H₂O (v)</td>
<td>703</td>
<td>730</td>
<td>678</td>
<td>656</td>
</tr>
<tr>
<td>BA</td>
<td>509</td>
<td>554</td>
<td>498</td>
<td>490</td>
</tr>
<tr>
<td>BA+10%H₂O (v)</td>
<td>529</td>
<td>560</td>
<td>499</td>
<td>491</td>
</tr>
<tr>
<td>BA+20%H₂O (v)</td>
<td>548</td>
<td>566</td>
<td>498</td>
<td>492</td>
</tr>
</tbody>
</table>
Figure captions

Fig 1. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air for anthracite coal AC.

Fig 2. High-speed, high-magnification cinematographic images of single particles (75-150µm) of various coals (anthracite AC, semi-anthracite HVN, and two bituminous SAB and BA) and a biomass (olive residue OR) in air and in two different simulated oxy-fuel conditions (21%O₂-79CO₂ and 30%O₂-70%CO₂). In each case, a particle is shown prior and after ignition takes place. Different coal ranks experience different ignition modes.

Fig 3. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air for semi-anthracite coal HVN.

Fig 4. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air for bituminous coal SAB.

Fig 5. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for anthracite coal AC.

Fig 6. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for semi-anthracite coal HVN.

Fig 7. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for bituminous coal SAB.

Fig 8. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air-firing conditions for blends HVN-OR.
Fig 9. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air-firing conditions for blends SAB-OR.

Fig 10. Gas emissions and normalised derivative curves of gas concentration during ignition tests for semi-anthracite coal HVN in air and O₂/CO₂ (21-35%O₂) with steam addition (the H₂O(v) is added as a substitute of N₂ or CO₂).
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Fig 3. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air for semi-anthracite coal HVN.
Fig 4. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air for bituminous coal SAB.
Fig 5. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for anthracite coal AC.
Fig 6. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for semi-anthracite coal HVN.
Fig 7. Gas emissions and normalised derivative curves of gas concentration during ignition tests in oxy-firing conditions for bituminous coal SAB.
Fig 8. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air-firing conditions for blends HVN-OR.
Fig 9. Gas emissions and normalised derivative curves of gas concentration during ignition tests in air-firing conditions for blends SAB-OR.
Fig 10. Gas emissions and normalised derivative curves of gas concentration during ignition tests for semi-anthracite coal HVN in air and O₂/CO₂ (21-35%O₂) with 20% steam addition (the H₂O(v) is added as a substitute of N₂ or CO₂).