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GAS TEMPERATURE AND RADIATIVE HEAT TRANSFER IN OXY-FUEL FLAMES

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ABSTRACT

This work presents measurements of the gas temperature, including fluctuations, and its influence on the radiative heat transfer in oxy-fuel flames. The measurements were carried out in the Chalmers 100 kW oxy-fuel test unit. The in-furnace gas temperature was measured by a suction pyrometer and by an optical system based on FTIR-spectroscopy. The radiation intensity was measured by a Narrow Angle Radiometer and the gas radiation was calculated with a Statistical Narrow Band model. The overall agreement between the two temperature measurement techniques was good. The optical system showed a lower temperature than the suction pyrometer in the low velocity regions of the furnace, a difference which is likely to be an effect of the purge gas added in the optical probe. The measured temperature fluctuations were evaluated by modeling of the gas radiation. The influence from the measured fluctuations on the radiative heat transfer shows no effect of turbulence-radiation interaction. However, by comparing with temperature fluctuations in other flames it can be seen that the fluctuations measured here are relatively small. Further research is needed to clarify to which extent the applied methods can account for the turbulence-radiation interaction in the investigated flame.

INTRODUCTION

One of the main options proposed for capture of CO₂ from fossil based power production is oxy-fuel combustion [1]. In this technology the combustion air is replaced by oxygen and recirculated flue gases thereby replacing the N₂ in the furnace with CO₂ and H₂O. This has an impact on the radiative heat transfer since both CO₂ and H₂O absorb and emit radiation in contrast to N₂, which is an inactive gas with respect to radiation. Radiation is the main heat transfer mechanism in combustion chambers and has accordingly been identified as a key research question for development of the oxy-fuel technology. The emitted radiation is proportional to the temperature to the power of four, and it is therefore crucial to accurately measure the in-furnace temperature for understanding of the radiative heat transfer. The temperature in flames is constantly changing. Applying a time average of these fluctuations in modeling of radiation can therefore not be done without introducing errors due to the strongly

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non-linear relation between temperature and radiation. The interaction between these fluctuations and radiation is called Turbulence-Radiation Interaction (TRI). In a review on the subject [2], it was concluded that it is important to include TRI in the modeling of heat transfer in turbulent reacting flows for accurate heat transfer predictions.

Therefore, it is important for the understanding of radiative heat transfer to be able to measure temperature fluctuations in furnace flames.

There are several ways to measure the flame temperature, ranging from simple techniques based on a thermocouple to advanced laser techniques. When bare thermocouples are used, the radiation loss from the thermocouple to a colder surrounding is a problem. To overcome the problem, a suction pyrometer can be used. In this technique flue gases are sucked with a high velocity over a shielded thermocouple, thus increasing the convective heat transfer and suppressing the radiative heat loss. A problem with the suction pyrometer is that the flame is disturbed by the removal of gas and by the cold probe itself. The technique has also a poor time resolution due to thermal inertia. Therefore, optical measurement techniques are appealing since they allow non-intrusive measurements with high temporal resolution. The optical techniques can be divided into line-of-sight techniques and point measurement techniques. Examples of line-of-sight techniques used in combustion diagnostics are pyrometers or spectroscopic techniques. Point measurement techniques in combustion are often advanced laser based techniques. The laser technique most frequently used for temperature measurements in large scale combustion applications is Coherent Anti-Stokes Raman Spectroscopy (CARS). In this work an optical technique based on FTIR-spectroscopy [3] is applied as it is relatively simple compared to laser techniques, at the same time it gives more information on the flame characteristic than a pyrometer since the entire spectrum instead of single wavenumbers is analyzed.

Measurements of radiation intensity and gas temperature have been carried out in the Chalmers 100 kW oxy-fuel test rig since 2003 [4-7]. The gas temperature has so far been measured with a suction pyrometer and no information on temperature fluctuations in the flames is therefore available. The aim with the experiments carried out in this work was to also obtain information of temperature fluctuations in the Chalmers oxy-fuel test rig. This was done by measuring the gas temperature with the optical FTIR technique [7]. The optical measurements are compared with suction pyrometer measurements and their performance is evaluated for the flames investigated. The influence of the measured fluctuations on the radiative heat transfer is examined by modeling of gas radiation, and the modeled gas radiation is compared with measurements of the total radiation intensity.
METHODOLOGY

Experiments
The measurements were carried out in the Chalmers 100 kW oxy-fuel test rig, a down-fired and refractory lined furnace presented in more detail by Andersson et al. [5]. The cylindrical furnace has an inner height of 2.4 m and an inner diameter of 0.8 m. The burner is a swirl burner with two air registers and a central fuel lance for either propane or coal. In the experiments of this work propane was used as a fuel in order to establish well-defined combustion conditions and dry recycling was applied in the oxy-fuel flames. The test rig allows in-furnace measurements in seven measurements ports distributed along the height of the reactor. Measurements were done by traversing water cooled probes radially through the furnace.

Three different flames were investigated: one air fired and two oxy-fuel flames with 25 respectively 30 vol-% O₂ in the oxidizer. The oxy-fuel flames are referred to as OF25 and OF30. The stoichiometric ratio was kept constant at 1.15 in all experiments, and the concentration of O₂ in the oxidizer was controlled by changing the recycling rate of the flue gas. Table 1 lists the test conditions. The wall temperature in the furnace was monitored by a number of thermocouples mounted inside the wall. During the experiments, temperature, gas concentrations and radiation intensity were measured in one measurement port in the high temperature flame zone (384 mm from the burner).

<table>
<thead>
<tr>
<th>Test case</th>
<th>O₂ in oxidizer [vol-%]</th>
<th>Stoichiometric ratio</th>
<th>O₂ in stack [vol-%]</th>
<th>Fuel input [kW]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>21</td>
<td>1.15</td>
<td>3.0</td>
<td>80</td>
</tr>
<tr>
<td>OF25</td>
<td>25</td>
<td>1.15</td>
<td>3.5</td>
<td>80</td>
</tr>
<tr>
<td>OF30</td>
<td>30</td>
<td>1.15</td>
<td>4.3</td>
<td>80</td>
</tr>
</tbody>
</table>

The gas temperature was measured with a water cooled suction pyrometer and with the optical technique based on FTIR. The suction pyrometer is equipped with a B-type thermocouple allowing temperature measurements up to 1700°C.

The system used in the optical measurement have previously been described by Bak and Clausen [8]. In principle, the technique measure emitted radiation from a gas column between a cooled background and the probe tip, at a distance g in Fig. 1. The optical measurement system gives an estimation of temperature fluctuations since the time resolution is in the order of 10⁻⁴ s. The time resolution is limited by the scanning velocity of the spectrometer and the width of the spectral region used for estimation of the temperature. The time to scan the entire spectrum was approximately 1 s in these measurements, giving a measurement frequency of 1 Hz. To be able to carry out in-situ measurements of temperature, the setup is integrated in a water cooled probe (Fig. 1). The optics is protected by an uncooled ceramic tube, and by a small purge gas flow of argon.
Figure 1. Cross-section of the tip of the optical probe used: a) water cooled steel probe, b) fibre optic cable, c) cooling water tube, d) focusing lens, e) ceramic protection tube, f) water cooled background. The distance (g) between the ceramics and the beam stop was 200 mm in the air flame and 51 mm in the oxy-fuel flames.

To determine the gas temperature from the measured spectra, the strongly absorbing CO$_2$ band around 2349 cm$^{-1}$ is analyzed. The emissivity at these wavelengths is close to unity for a short distance, and the gas does therefore act as a blackbody in this spectral region. The gas temperature can accordingly be found by fitting a Planck curve to the measured spectra at the strongly absorbing region 2250-2350 cm$^{-1}$, as shown in Fig. 2. Due to the non-uniform path, the shape of the measured spectra is not a perfect Planck curve, Fig. 2b). Therefore one temperature for each data point in the spectral region is determined, and an average of these temperatures is calculated.

The temperature given by fitting the Planck curve is an estimate of the instantaneous temperature of the gas column near the probe tip. By combining a series of measured temperatures, information of the fluctuations inside the flame is obtained. The spatial resolution of the measurements, the length of the gas column needed to reach an emissivity close to unity, depends on the concentration of CO$_2$. A higher CO$_2$ concentration will give higher absorption and a shorter path length is needed, the spatial resolution can therefore be increased in the oxy-fuel flames. For pure gas temperature measurements the water cooled beam stop is not required, but it was used here since additional information on gas concentrations can be obtained if the optical path length is known. The equipment was calibrated with a two point method [9].

Figure 2. Example of a measured spectrum without the beam stop and a Planck curve, a) the entire scanned spectrum b) the strongly absorbing region used to determine the temperature.
The line-of-sight radiation intensity was measured with a narrow angle radiometer of IFRF-type, as shown in Fig. 3. The measured radiation is the total intensity including both gas and particle radiation. A small purge gas flow of argon was used to prevent flue gas and particles to enter the tube. The sensor used is a thermopile, and to ensure a stable signal, the sensor housing is maintained at a constant temperature by a separate water cooling circuit. The measured intensity corresponds to radiation emitted by the gas column between the probe tip and the background, a window at the opposite side of the furnace. The probe, and the procedure of the measurements are similar to what is described by Andersson et. al. [4].

![Figure 3. Schematic picture of the narrow angle radiometer used to measure the radiation intensity: a) thermopile, b) focusing lens, c) shutter, d) PT-100, e) water cooled sensor housing, f) collimating tube and g) water cooled probe.](image)

The time average in-furnace gas concentrations were measured by extracting flue gas with a water cooled extraction probe. The inner tube of the probe is electrically heated to ensure a temperature above the dew point of the flue gas. The extracted flue gas is analyzed by both a conventional gas analyzing system and a system based on FTIR- and UV spectroscopy. The gas species of interest in this work are H$_2$O, CO$_2$, CO and O$_2$. Due to the different gas composition in air- and oxy-fuel flames, several analyzers calibrated under conditions corresponding to both air- and oxy-fuel combustion were applied. The instruments were calibrated each morning before the measurements.

**Modeling of gas radiation**

The measured temperature and gas concentrations were used as input for modeling of the gas radiation intensity along the measured profiles in the furnace. The dominating species contributing to gas radiation are CO$_2$ and H$_2$O. The radiative properties of these gases are well known and are here calculated with the Malkmus Statistical Narrow-Band (SNB) model [10]. The transmissivity, $\bar{t}_k$, of each narrow band is calculated according to

$$\bar{t}_k = \exp \left[ -\frac{2\gamma}{d_k} \left( \frac{YPLk_kd_k}{\gamma} - 1 \right) \right]$$

where $Y$ is the mole fraction of the absorbing species, $P$ is the total pressure and $L$ is a characteristic length. The parameters $k_k$ and $d_k$ are species specific temperature dependent parameters, and are taken from the work of Soufiani and Taine [11]. The half-widths, $\gamma$, of the absorbing species are calculated with $P_s = 1$ bar and $T_s = 296$ K as
\[
Y_{\text{CO}_2} = \frac{P}{P_s} \left( \frac{T_s}{T} \right)^{0.7} \left[ 0.07Y_{\text{H}_2\text{O}} + 0.058(1 - Y_{\text{CO}_2} - Y_{\text{H}_2\text{O}}) + 0.1Y_{\text{H}_2\text{O}} \right] \tag{2}
\]
\[
Y_{\text{H}_2\text{O}} = \frac{P}{P_s} \left\{ 0.462Y_{\text{H}_2\text{O}} \left( \frac{T_s}{T} \right)^{0.5} + \left( \frac{T_s}{T} \right)^{0.5} \left[ 0.079(1 - Y_{\text{CO}_2} - Y_{\text{O}_2}) + 0.106Y_{\text{CO}_2} + 0.036Y_{\text{O}_2} \right] \right\} \tag{3}
\]

To account for the non-uniform path the Curtis–Godson approximation is applied [12] and the transmissivity (\(\bar{\tau}_{k,g}\)) of the gas mixture is calculated as the product of the absorbing gases:

\[
\bar{\tau}_{k,g} = \tau_{k,\text{H}_2\text{O}} \tau_{k,\text{CO}_2} \tag{4}
\]

The path between the probe and the quartz window is discretized into a number of equally sized cells. The intensity at each cell face is calculated by using the correlated formulation of the Radiative Transfer Equation (RTE) for each narrow band, Eq. (5).

\[
I_{k,n} = I_{0k} \tau_{k,0-n} + \sum_{k}^{n-1} \left( \tau_{k,i+1-n} - \tau_{k,i-n} \right) I_{b,k,i+1/2} \tag{5}
\]

The total gas radiation, \(I_{\text{tot},n}\), can then be calculated as a sum of all band intensities, \(I_{k,n}\), multiplied by the band width, Eq. 6.

\[
I_{\text{tot},n} = \sum_{k} I_{k,n} \Delta \nu \tag{6}
\]

As been mentioned above, the influence of temperature fluctuations on the radiative heat transfer can be significant. The influence of the fluctuations is evaluated by using two temperature profiles when calculating the black body radiation, \(I_{b}\). The first temperature profile is based on the time average of the measured temperatures. It is calculated as an arithmetic mean value of the instantaneous temperatures, \(T_i\), measured with the optical system, \(j\) refers to the number of measured temperatures as

\[
T_{\text{avg}} = \frac{1}{j} \sum_{i=1}^{j} T_i \tag{7}
\]

This temperature does not account for the effect of the strongly non-linear relation between temperature and radiation. In the second approach the fluctuations are accounted for by a simplified approach since it is very complex to fully account for TRI. The gas concentration is assumed to be constant at each position in the flame, and the only fluctuating term included in the modeling is the temperature. An average radiation temperature, \(T_{\text{rad}}\), is calculated as the mean of the instantaneous temperatures to the power of four, Eq. (8). This average reflects the influence of a fluctuating temperature on the non-linear relation between temperature and radiation.
\[ T_{\text{rad}} = \sqrt[4]{\frac{1}{j} \sum_{i=1}^{j} T_i^4} \]  

Eq. (8) is a simple way of accounting for TRI in modeling of radiative heat transfer, as it only accounts for temperature fluctuations and the temperature fluctuations in each position in the flame are assumed to be independent.

RESULTS AND DISCUSSION

Figure 4 gives the measured temperature profiles in the flames investigated. The temperature measurements were not performed on the same day with the optical system and the suction pyrometer, but the wall temperatures and combustion conditions were the same in each case i.e. similar flame temperatures are expected. The peak temperature in the air- and the OF30 flame is above 1500°C, while the peak temperature in the OF25 flame is lower, around 1400°C. The temperature difference between the two oxy-fuel flames is due to a higher recycle rate in the OF25 flame, i.e. more cold CO\textsubscript{2} is added to the flame. The increased emission and absorption from CO\textsubscript{2} and H\textsubscript{2}O is a likely reason for the slight broadening of the temperature profile in the oxy fuel flames. Cold CO\textsubscript{2} is absorbing radiation emitted by the flame, thus increasing the gas temperature close to the walls.

In general, the agreement is good between the two measurement techniques. Yet, the optical measurement gives a slightly higher temperature in the center of the flame and a lower temperature in the low temperature regions. The region outside the flame where the temperature and temperature gradients are relatively low, the suction pyrometer should give a good estimation of the average temperature. The difference of up to 200°C between the optical technique and the suction pyrometer may be explained by cooling and dilution by the purge gas (argon) that is added to protect the optics. In this region the velocity is low and it is therefore likely that the flue gas flow is not able to entirely remove the argon, thus decreasing the concentration of CO\textsubscript{2} as well as cooling the CO\textsubscript{2} which makes the calculated temperature from the spectra too low. To avoid this effect, the flow of purge gas should be kept at a minimum, especially in low-velocity regions. The problem of self-absorption from cold CO\textsubscript{2} is further discussed in the work by Clausen [3].

The spatial resolution of the optical measurements in the air fired flame is lower than in the oxy-fuel flames due to a lower CO\textsubscript{2} concentration. The distance between the beam stop and optics can therefore be decreased in the oxy-fuel flames. The influence from the purge gas, discussed above, seems to increase when the measurements are performed on a shorter gas column and the distance between the beam stop and the optics is decreased. This can be seen by looking at the temperature difference between the suction pyrometer and the optical system close to the walls in the air flame compared to the oxy-fuel flames.
Figure 4. Temperature profiles for the three different flames; a) air, b) OF25 and c) OF30. The probe is inserted at 0 m and traversed through the flame.

For optimum performance of the optical temperature measurements, the temperature and gas concentration should be constant in the measured line-of-sight path. However, when measuring in a turbulent diffusion flame the temperature and gas concentration in the measurement volume will vary. With a non-uniform temperature in the measurement volume, the measured temperature will be biased towards higher temperatures [13]. The influence of the lower spatial resolution, and thus larger temperature differences in the measurement volume, can be seen as a broadening of the temperature profile measured with the optical system compared to the profile measured with the suction pyrometer. This broadening is more pronounced in the air flame, in which the optical distance is larger, Fig. 4 a), compared to the oxy-fuel flames, Fig. 4 b) and c). The temperature profiles given in Fig. 4 are measured with the water cooled beam stop. Measurements were also performed without the beam stop, and the result was similar.

Figure 5 a-c) shows both measured and modeled radiation intensity profiles in the radial cross-section 384 mm from the burner. Modeling is based on the profile of average temperature measured with the suction pyrometer as well as the profile measured with the optical system. The radiation intensity in the air-fired flame is lower than the intensity in the two oxy-fuel flames, due to the increased concentration of CO₂ and H₂O in the oxy-fuel flames. The higher and broader temperature profiles measured with the optical system results in a higher modeled gas radiation in all three flames (dashed curve). The difference between the modeled gas radiation and the measured total radiation can be caused by radiation from particles [4], and possibly by the TRI effects which are not accounted for in the modeled profiles in Fig. 5. In the air fired flame it should be noted that the purge gas flow added to prevent gas from entering into the collimating tube, was higher than in the oxy-fuel flames. This has probably affected the measured intensity in the region 0.4-0.8 m in Fig 5 a), where the modeled gas radiation is higher than the measured intensity. It is believed that the purge gas has diluted and cooled the flue gas in front of the probe, thus decreasing the measured intensity. The difference in temperature measured with the suction pyrometer and the optical system in the low temperature region of the furnace can also be seen in the modeled intensity profile, as a larger decrease in intensity close to the measurement port in the profile based on the optical temperature in the oxy-fuel flames, Fig 5 b) and c).
Figure 5. Measured total intensity and modeled gas radiation intensity profiles in a cross-section of the investigated flames; a) air, b) OF25 and c) OF30.

Figure 6 shows the measured instantaneous gas temperature at a position 300 mm from the measurement port in the OF30 flame. Similar fluctuations as in Fig. 6 can be seen in all positions in the flame, but the fluctuations are largest in the flame boundaries where the temperature gradients are highest. The measured instantaneous temperatures are used to calculate an arithmetic average temperature and an average of the instantaneous temperature to the power of four, according to Eqs (7) and (8), also shown in Fig. 6. The difference between the two calculated average temperatures is small in all positions indicating that the choice between Eqs (7) and (8) should have a small influence on the radiative heat transfer in this flame.

Figure 6. Instantaneous gas temperature measured with the optical system at a position 300 mm from the wall in the OF30 flame.

To investigate the influence of the measured temperature fluctuations on the radiative heat transfer the two calculated average temperatures are applied in the modeling of the gas radiation. The result is shown in Fig. 7, and as seen the influence of the fluctuations measured is negligible. The difference between the highest and lowest temperature in Fig. 6 is 470°C, which indicates that significant temperature differences can be captured with this measurement technique. However, the influence on the average value from these fluctuations is small. The optical measurement technique has previously been compared to CARS measurements in a coal flame [3]. From that comparison it was seen that the Probability Density Function (PDF) was wider in the CARS measurements, which indicates that the fluctuations measured with CARS can be expected to be higher than with the FTIR-based setup. A comparison with the commonly used reference flame Sandia Flame D [14] shows
that the temperature fluctuations measured with Raman/Rayleigh scattering in the Sandia flame is larger than what is measured in the Chalmers flame. Another study by Hughes et al. [15] in which CARS was used to measure temperature in a swirling coal and oil flame with a thermal input of 280 kW, showed a difference between the highest and lowest temperature in a position downstream of the oil flame of 600°C. The standard deviation was also higher than what is observed in the flame investigated here, indicating that the influence from the fluctuations measured by Hughes et. al. [15] would be larger than from the ones observed in this work. The fact that the optical temperature measurement technique is a line-of-sight measurement can mean that not all temperature fluctuations in the flame are captured, which could explain the differences in measured fluctuations. Further research is needed to clarify these issues as well as the influence of other fluctuating variables such as concentration of gas species and particles.

Figure 7. Radiation intensity with modeled profiles based on temperatures calculated with both Eq. 6 and 7 a) air, b) OF25 and c) OF30.

CONCLUSIONS

Gas temperature, radiation intensity and gas composition have been measured in Chalmers oxy-fuel test unit. Two techniques for measuring gas temperature have been used and evaluated: a suction pyrometer and an optical technique based on FTIR. The FTIR-based technique was used since it provides information of temperature fluctuations. The influence of the measured temperature fluctuations are evaluated by modeling the gas radiation in the flames. In general the agreement between the temperature profiles measured with the suction pyrometer and the optical system is good. However, the FTIR-based system showed a somewhat lower temperature in the low velocity regions compared to the suction pyrometer measurements, the difference likely being due to the influence of the purge gas on the FTIR measurements. The measured temperature fluctuations do not show any significant effect on the radiative heat transfer. However, studies in other flames have indicated larger temperature fluctuations and more work is needed to clarify if the optical technique is capable of capturing all fluctuations in the investigated flames before conclusions can be drawn to what extent, if at all, the temperature fluctuations are of importance for modeling the conditions studied in this work.
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REFERENCES