# SIMULATION OF OXY-FUEL COMBUSTION PROCESSES IN INDUSTRIAL FURNACES

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#### Abstract.

For a variety of reasons, oxy-fuel combustion, i.e. the combustion of a fuel (most often natural gas) with pure oxygen instead of air, has become firmly established in thermal processing applications in recent years. Increased thermal efficiencies and heat transfer rates allow for a more productive operation of industrial furnaces, even despite the increased operational cost for the oxygen supply. In glass manufacturing, oxy-fuel combustion is considered to be one of the foremost solutions to operate glass melting furnaces at very high furnace temperatures without producing excessive amounts of pollutants such as nitrous oxides ( $NO_X$ ).

The use of computational fluid dynamics to design and optimize industrial furnaces has become more and more widespread. Advances in computer hardware and the development of ever more sophisticated models and numerical solver algorithms allow for the numerical description of complex physical and chemical processes such as turbulent flows, combustion and heat transfer in realistic geometries; the analysis and optimization of industrial furnaces by means of numerical simulation has become routine. However, these models for turbulence, combustion and other phenomena are based on simplifying assumptions which may pose limits to their applicability in certain situations.

In a recent research project, Gas- und Wärme- Institut Essen e. V. (GWI) investigated the applicability of a number of combustion models for industrial oxy-fuel combustion processes. The results of OH chemoluminiscence imaging techniques and detailed 2D field measurements of characteristic chemical species distributions in a semi-industrial burner test rig were compared with CFD simulations based on a variety of commonly used combustion models. The comparisons showed that many popular combustion models found in commercial CFD codes fail to capture the characteristics of oxy-fuel combustion. Models using a mixture fraction approach in combination with a non-adiabatic data table generated by either an equilibrium or a flamelet model seemed to have problems with the impact of strong heat losses on the chemistry, while models based on the Eddy Dissipation Model suffered from the drastically simplified chemistry description found in the 2- and 4-step reaction mechanisms that were used. Only the more sophisticated Eddy Dissipation Concept in combination with a rather more comprehensive reaction mechanism (17 species, 50 elementary reactions) was able to describe the oxy-fuel combustion process in good agreement with the measured data, although at prohibitive computational cost. A more detailed chemical reaction pathway

analysis showed that the severely reduced 2- and 4-step reaction mechanisms were not applicable for oxy-fuel combustion. This result was corroborated by spectral radiometric measurements. It was found that at the moment, the capability to simulate oxy-fuel industrial furnaces is severely limited.

In the course of a follow-up research project, GWI, in cooperation with its partners, is currently working on improving reaction models for oxy-fuel combustion processes in industrial applications, combining sufficient accuracy with manageable numerical cost.

# **1 INTRODUCTION**

Glass production on an industrial scale requires furnace temperatures of more than 1600 °C, depending on glass quality. The flame temperatures are, of course, even higher. In conventional furnaces, these temperature levels can only be achieved by intensive recuperative or regenerative air pre-heating with pre-heat temperatures rising to about 800 °C for recuperators and even 1400 °C for regenerative air pre-heaters [1]. However, these high temperatures in combination with long residence times and an inexhaustible supply of nitrogen in form of the combustion air can lead to high emissions of thermal nitrous oxides (NO<sub>x</sub>). The glass industry is therefore very much interested in modifications of the combustion treatment of the flue gas. One such approach is the so-called oxy-fuel combustion where pre-heated air is substituted by almost pure oxygen as oxidizer.

There is occasionally some confusion about the term "oxy-fuel combustion" as it describes a different process in the power plant sector. There, "oxy-fuel combustion" refers to the substitution of combustion air with a mixture of oxygen and recirculated flue gas in order to facilitate carbon capture and storage techniques (CCS) [2]. In the glass industry as well as in other manufacturing processes, however, "oxy-fuel combustion" is considered to be the combustion of fuel with (almost) pure oxygen in order to achieve high combustion temperatures without resorting to air-preheating.

There are several advantages to oxy-fuel combustion in glass melting furnaces. First of all, the main source of nitrogen in the combustion system is removed, thus drastically reducing the potential  $NO_x$  emissions. The firing efficiency of the furnace is increased and costly airpreheating systems are no longer required. Conventional furnaces can easily be retrofitted to oxy-fuel combustion by swapping the burners and dismantling (or simply closing off) the air pre-heating equipment. On the other hand, the economic feasibility of oxy-fuel furnaces is largely dependent on the cost of oxygen production on-site. Also,  $NO_x$  formation may actually increase if impure oxygen is used or if there are major air leakages or the furnace process is not properly controlled.

Already, there are many oxy-fuel glass melting furnaces in operation worldwide, but most of these are converted conventional furnaces [3, 4, 5]. This means that the furnace space was not optimized to take into account the changes in heat transfer in such a furnace due to the changes in atmospheric composition. In an oxy-fuel system, the flue gas consists almost exclusively of species such as  $CO_2$  and  $H_2O$  which participate actively in radiative heat transfer.

In the course of a research project carried out by Gas- und Wärme- Institut Essen e. V. (GWI), the oxy-fuel combustion process was investigated both by experimental and numerical means in order to determine design criteria for new, purpose-built oxy-fuel glass melting furnaces. As the modeling of such industrial furnaces via CFD simulations is increasingly gaining importance for industrial applications, one aim of the project was to evaluate the performance of common reaction models in CFD codes for oxy-fuel combustion systems.

#### **2 VALIDATION MEASUREMENTS IN A SEMI-INDUSTRIAL TEST RIG**

The experimental investigations of this research project were carried out at one of GWI's semi-industrial burner test rigs. Detailed 2D field measurements of relevant species (CO, CO<sub>2</sub>, O<sub>2</sub>, NO and NO<sub>2</sub>) were performed as well as flame visualization by means of OH chemoluminiscence. The experiments were accompanied by spectralradiometric measurements by the "Hüttentechnische Vereinigung der Deutschen Glasindustrie" (HVG). The aim of this measurement campaign was to provide insight into the basic processes of oxyfuel combustion on a semi-industrial scale and to observe the impact of different burner configurations and fuels on the combustion process as a whole. Also, the experimental data can be used for CFD validation purposes.

Five different burners were investigated. Sketches of these burners are shown in Fig.1. Among the burners, there are high and low momentum burners as well as a flat flame burner. Four of the burners use natural gas as fuel, while Burner 2 uses liquid fuel. All burners have a thermal load of 400 kW.

Fig. 2 shows OH chemoluminiscence images of the various burners, all taken at identical operating conditions (P = 400 kW,  $\lambda$  = 1.05). As can be expected, the geometry of the burner has a significant impact on the flame shape and structure.

This is especially obvious in the comparison of Burners 4 and 5 which differ only in the diameters of their nozzle orifices. The jets of the high velocity burners entrain much hotter, but chemically inert exhaust gas from the surroundings of the jet and force them into the reaction zone, thus diluting it. The low velocity Burner 5 on the other hand shows significantly higher local OH concentrations in the reaction zone as the momentum of the jets is insufficient to entrain large quantities of flue gas. The image of Burner 2 shows that of course the choice of fuel plays a major role in the OH distribution in the reaction zone. Even in a high velocity burner system, large concentrations of OH will be found in the reaction zone when using complex fuels such as oil.

As mentioned above, detailed two-dimensional field measurements were carried out for a variety of species. Fig. 3 shows the measuring plan of GWI's 1.3 MW semi-industrial test rig. Each of the fourteen axial positions can be accessed by a port through which a water-cooled probe can be inserted into the furnace. This probe can also be moved in the transversal direction to obtain a two-dimensional grid of measuring positions. Flue gas samples are taken at each measuring point and then analyzed.



Figure 1:Sketches of the investigated oxy-fuel burners



Figure 2: OH chemoluminiscence images of the investigated burners

Fig 4 shows a comparison of the measured CO concentration distributions for the five burners. The differences between the various burner systems are obvious. Similar distributions were obtained for species such as  $O_2$ ,  $CO_2$  and  $NO_x$ . These detailed data sets are well suited to evaluate the performance of reaction models in CFD codes. In addition to the flame visualization and field measurements, this campaign was accompanied by spectralradiometric measurements carried out by HVG in order to provide data for spectral radiation modeling for CFD codes.



Figure 3: Measuring plan of GWI's semi-industrial test rig



Figure 4: Measured CO distributions for the investigated burners

# **3 NUMERICAL SIMULATION OF OXY-FUEL COMBUSTION**

One of the main objectives of this research project was to evaluate the performance of commonly used reaction models in standard CFD applications with regards to oxy-fuel

combustion. Most of the models available today in commercial codes were developed and validated for conventional fuel-air combustion for which they yield reasonable results. However, the situation in an oxy-fuel-fired furnace is rather different from a conventional furnace. Therefore, the applicability of these reaction models, which need to rely on some quite drastic simplifying assumptions to reduce computational cost, to oxy-fuel furnaces has to be verified.

This is why a series of RANS simulations of the experiments presented above was carried out in which various combustion models were applied. The software used was the commercially available CFD tool FLUENT (versions 6.3 and 13). Burner 5 was chosen as a reference burner, with an air ratio of 1.05 at a thermal load of 400 kW (cf. Figure 5).





In all cases, the realizable k- $\epsilon$  turbulence model and the Discrete Ordinates model (DOM) were used, while a variety of common reaction models were chosen to describe the chemistry of oxy-fuel combustion: PDF-Equilibrium, PDF-Flamelet, Eddy Dissipation Models (EDM) with 2- and 4-step mechanisms (Westbrook-Dryer [6] and Jones-Lindstedt [7] respectively) and the Eddy Dissipation Concept (EDC) [8] with a more detailed reduced mechanism containing 17 species and 50 reaction equations [9]. In the following, the results for simulations of Burner 5, a low velocity pipe-in-pipe burner, will be presented.

Fig. 6 shows a comparison of measured and simulated  $CO_2$  concentrations in the burner plane. It is obvious that the PDF-Equilibrium model, at least in the implementation used in FLUENT, is unable to predict the species distribution adequately, the  $CO_2$  concentrations are much too small. The Eddy Dissipation Model based on the 2-step Westbrook-Dryer reaction mechanism yields better results, the flame shape in the simulation is quite similar to that found in the measured data, but overall, the calculated flame is too short by about 50 %. The EDM simulation using the 4-step Jones-Lindstedt mechanism yields results similar to that of the 2-step mechanism. The results of the PDF-flamelet simulation were close to that of the PDF equilibrium simulation which is why they are not presented here.

Only the EDC simulation with a more comprehensive reaction mechanism can describe the flame structure with a reasonable degree of accuracy.



Figure 7: Comparison of measured and simulated CO concentration distributions

The simulated CO concentrations show similar results, as can be seen in Figure 7. The PDF-Equilibrium model obtains a completely unphysical solution. Since the predicted  $CO_2$  concentrations are too low, the predicted CO concentrations are much too high. The EDM with a 2-step-mechanism again yields better results, but only the more sophisticated EDC approach with a rather detailed reaction mechanism can achieve a good agreement with the measured distribution, especially with regards to the size and shape of the reaction zone. Cuoci et al. [10] report similar findings in their investigations of oxy-fuel combustion. However, the EDC approach requires significantly more computational resources and CPU time which make it ill-suited for numerical parameter studies in order to improve furnace design. Nevertheless, a reliable prediction of flue gas composition in the furnace space is crucial since the distribution of species also influences radiative heat transfer modeling, the main heat transfer process in glass melting furnaces and other high temperature processes.

The reason for the failure of the PDF-based models (the PDF-Flamelet model yields similar results which is why it is not presented here) seems to be that the models are unable to deal with the strong non-adiabaticity of the system. In the experimental setup, water-cooled tubes were used to emulate the heat loss of the furnace space into the glass melt. These heat losses were of course also part of the boundary conditions of the simulations. The calculated distributions of CO and CO<sub>2</sub> would agree better with an adiabatic oxy-fuel combustion system at these high temperature levels, but are evidently wrong for a strongly non-adiabatic system. In the course of this project it was unfortunately not possible to clarify if this behavior is a problem of these models in general or merely an issue with the specific implementations of the models in the used CFD codes.

The reason why the EDM simulations in combination with the Westbrook-Dryer mechanism fail (the Jones-Lindstedt mechanism with four species behaves similarly) lies in the nature of the reduced reaction mechanisms used. Both the Westbrook-Dryer and the Jones-Lindstedt mechanism were developed to describe near-stoichiometric methane-air combustion in such a reduced manner to be used in CFD simulations with reasonable numerical effort. However, in an oxy-fuel system, the kinetics of the radical species behave differently while the radical species occur in bigger concentrations as well due to the lack of diluting nitrogen. The reduced mechanisms are unable to describe the different kinetics as they were never developed or validated for these conditions in the first place.

Using kinetic simulations of one-dimensional freely propagating premixed flames at stoichiometric conditions, elemental flux analyses of a non-preheated and a strongly preheated air-fuel flame as well as an oxy-fuel flame were carried out. They show drastic differences between reaction pathways for the different flames (cf. Fig. 8 and 9). In all cases, the GRI 3.0 reaction mechanism [11] was used.

Fig. 8 shows the elemental flux analysis for carbon atoms for the non-preheated fuel-air case. It can be seen that there is only a relatively small number of relevant pathways in this combustion process. If the combustion air is pre-heated to temperature levels common in the glass industry, the reaction pathways change somewhat, but are still relatively small in number. Experience shows that even at elevated temperature levels, common drastically reduced reaction mechanisms yield reasonable results.

However, the situation changes completely when the combustion air is substituted by pure oxygen, as can be seen in Fig. 9. The reaction pathways become more numerous and complex,

as a larger number of radical species is involved. Also, the concentrations of these radical species are higher since the diluting nitrogen is no longer present in the system.



Figure 8: Elemental flux analysis for carbon atoms in a non-preheated stoichiometric premixed methaneair flame



Figure 9: Elemental flux analysis for carbon atoms in a non-preheated stoichiometric premixed methaneoxygen flame

This greater impact of radical species on the combustion process is corroborated by spectralradiometric measurements performed by HVG [12]. In the case of oxy-fuel combustion, significant concentrations of  $C_2$ -species were detected in the flame region which are not commonly found in fuel-air-flames. The pathway analysis confirms that these species take part in the oxy-fuel combustion process to a much greater extent than in air-fired combustion.

The results of the CFD simulations as well as those of the kinetic investigations indicate that a detailed description of chemical kinetics is important in order to describe oxy-fuel processes adequately, since higher temperatures und local radical concentrations have a profound impact on the combustion process as a whole. This is why only the EDC approach with a relatively detailed reduced reaction mechanism of Kee et al. [9] was able to obtain reasonable approximations for crucial flame characteristics, though at increased numerical cost.

In order to describe oxy-fuel furnaces in a more time-efficient manner, smaller reduced mechanisms are necessary which were specifically developed and validated for such systems. This is especially important since the predicted local flue gas composition also serves as a boundary condition for radiation models.

#### 4 CONCLUSION

In the glass industry as well as in many other thermal processing industries, the use of oxyfuel burners becomes more and more widespread. In this context, oxy-fuel combustion means the combustion of fuel (usually natural gas) with almost pure oxygen as oxidizer in order to obtain high process temperatures without resorting to intensive air pre-heating and hence strong thermal  $NO_x$  formation in the furnace. Since most currently operating oxy-fuel glass melting furnaces are based on conventional air-fired furnace designs, design criteria are needed in order to better exploit the specific characteristics of oxy-fuel combustion. In the course of a recent German research project, Gas- und Wärme-Institut (GWI) investigated oxyfuel combustion by both numerical and experimental means in order to gain a deeper understanding on the processes involved and to propose design criteria for purpose-built oxyfuel furnaces.

As computer-assisted design of industrial furnaces becomes ever more important, the validity of the models applied has to be assured. The comparison of detailed measurements with the corresponding simulations shows that many popular combustion models found in modern CFD codes fail to accurately describe oxy-fuel combustion. Only a rather sophisticated and numerically expensive model, the Eddy Dissipation Concept, in combination with a relatively detailed reduced reaction mechanism, yields reliable results. However, this approach requires too many resources to be used for large scale parameter studies.

In order to be able to perform robust and reliable simulations of oxy-fuel combustion processes in furnaces, reduced reaction mechanisms are required which are specifically developed for oxy-fuel conditions.

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