Gas sensing technologies in combustion: A comprehensive review

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Abstract
Irrespective of the change in kind of energy to renewables there are many processes in which combustion of conventional fuels and biofuel are necessary. For cement and glass production, paper fabrication and air traffic it is absolutely essential to control the combustion processes by intelligent sensors in order to maximize the efficiency and to minimize the emission of harmful substances such as NOₓ and CO. Mainly oxygen sensors based on solid electrolytes and calorimetric sensors using the heat formation by catalytic combustion of CO and hydrocarbons with two resistance temperature detectors (RTD) are utilized. Recently, tunable diode laser spectroscopy (TDLS) became attractive in chemical plants. Several analytical companies are offering in-situ or extractive laser analyzers for combustion gases.

Keywords
Combustion control; in situ sensors for oxygen and hydrocarbons; solid electrolyte sensors; calorimetric sensors; tunable diode laser spectroscopy.

Combustion process
The combustion of fuel is a complex process which has to be carried out in a small operation window concerning the air/fuel ratio. By reasons of economy as well as restriction of law the main components as oxygen and unburnt species (CO and hydrocarbons) have to be measured in situ to gain real-time information. NOₓ as the main pollutant must be measured at the end of exhaust to ensure the regulations by law. An overview on the concentrations of combustion flue gases and the small operation window is given in Figures 1 and 2.

Due to the high viscosity of flue gases the concentration of oxygen and CO is local-dependent (stratification) so that several sensors have to be installed to obtain an average value of oxygen and combustibles (Figure 3).
There are several technologies for combustion process gases monitoring (Figure 4). Only few of them, such as electrochemical high temperature sensors, calorimetric sensors and laser techniques, are suitable to measure in situ across stack application. The last one is expensive as compared with solid electrolyte sensors and it needs some effort for maintenance.

**Solid electrolyte sensors**

Electrochemical oxygen sensors based on zirconia solid electrolytes, which are widely used today in different industrial applications and for controlling fuel/air ratio in internal combustion engines (lambda sensor), were first introduced ~60 years ago by Peters and Möbius (Germany) [4] and Weissbart and Ruka (Westinghouse, USA) [5]. An overview on the basic principles of solid electrolyte sensors is given in Figure 5 [6]. Common to all these sensors is the high operation temperature which is necessary to ensure high electrochemical kinetics for the oxygen exchange within the electrodes and sufficient conductivity of the solid electrolyte. On the other hand, it allows a high operation temperature to be measured in hot gases such as in or near the combustion chamber. Most industrial zirconia oxygen sensors are based on electrochemical cells with stabilized zirconia solid electrolytes operating in potentiometric mode [7-12]. This electrochemical cell is made of zirconia solid electrolyte (mostly yttrium stabilized zirconia, YSZ) and two platinum electrodes printed and sintered on the opposite sides of a dense zirconia ceramics and exposed to the process and reference gases, respectively. Gas-tight sintered ceramics is formed as tubes, discs, planar substrates or thick films. Zirconia sensors are supplied as steel or ceramic probes up to 4 m in length.
The service life depends on application temperature and loads of pollutants and can be up to 510 years. Several producers from US (Emerson, Ametec), Japan (Yokogawa) and Germany (ENOTEC, Zirox, STG) are on the market. YSZ based oxygen sensors can be applied not only for optimization of combustion, but also to measure in processes in which reducing or oxidizing gases are necessary in order to ensure a certain oxidation state. That is important for the production of ceramics, porcelain, bricks.

| potentiometric | mixed potential | amperometric | coulometric |
|---------------|-----------------|--------------|-------------|
| $U_m = U_0 - A \ln p_{\text{HC}}$ | $I_d \equiv p(O_2)$ | $I_F = n(O_2)zF$ | |
| measuring range | 10$^{-30}$ ... 10 bar | 1 ... 20 000 vol.-ppm | 10$^{-4}$ ... 1 bar | 10$^{-5}$ ... 1 bar |

**Figure 5.** Basic working principles of solid electrolyte sensors for oxygen containing gases. The equations refer to the measured signals [6].

Oxidizing and reducing states differ in Nernst’s Voltage (Figure 6). At the lambda-leap ($\lambda = 1$) the equilibrium potential changes abruptly. In the 1960s and 1970s the limits of measurement in terms of temperature and oxygen partial pressure have been determined [7]. Sometimes the load of the flue gas with sulfur cause problems regarding the poisoning of electrode material [13]. Hence, nowadays there are only few research activities in this field [14].

For automotive applications, lambda probes (oxygen sensors) are based on the potentiometric or combined potentiometric and amperometric dual sensing technologies [8,9]. Such sensors are produced in millions of pieces by thick film techniques in a co-firing process. Recently, mixed potential sensors for CO and hydrocarbons were developed [15,16] and are commercially available in Germany by Lamtec and Enotec. The components in flue gas like CO, hydrocarbons ($H_nC_m$), nitric oxides ($NO_x$) and residual oxygen ($O_2$) to be measured do not equilibrate at temperatures < 700 °C in the vicinity of the hot electrode. Thus, gas components which are not thermodynamically stable are electrochemically active. In hydrocarbons and oxygen containing atmosphere at least two electrode reactions can take place: oxygen electrochemical reduction and electrochemical oxidation of hydrogen or other combustibles (Figure 7). The measured open circuit voltage, also called mixed potential, does not obey the Nernst equation but depends logarithmically on the concentration of CO, methane or hydrocarbons. Low catalytic activity materials such as gold or gold platinum alloys are usually employed for these sensors combined with catalytically active Pt-electrode.

In many cases composite materials consisting of gold and oxides show a high sensitivity to CO as can be seen in Figure 8. Furthermore, complex perovskite-type oxides are very promising sensitive materials (Figure 9).
It was established that beside the morphology, the electronic and ionic defects in such substituted oxides play an important role for sensitivity and selectivity optimization and trimming [17-19]. The main drawback of all mixed potential sensors is their quite high cross-sensitivity to sulfur dioxide as well as long term stability in severe combustion flue gas environment [1]. Often carbon monoxide equivalent expression “COe” is applied to these sensors meaning CO concentration response used for the calibration.

Calorimetric catalytic sensors for combustibles (CO, CH₄)

In sensors using calorimetric effects by catalytic turnover of combustibles such as CO or methane, two resistance temperature detectors (RTD) are compared. On of this is covered with catalytically active layer, e.g., platinum. The catalyst film is merged with RTD using highly thermally conductive packaging oxide materials to ensure more efficiently heat transfer of the combustion reaction on catalyst film to RTD. The surface of the other one is catalytically inactive. The reaction heat of the active resistor leads to an increase of temperature linearly proportional to the concentration of combustibles [20,21].
Oxidation reaction occurs between combustible species and oxygen with defined heat released is depending highly on the combustible concentration. Calorimetric catalytic CO-sensor can measure CO-concentration down to ppm level with detection limit \( \sim 5\ldots10 \) ppm CO. This technology is permitting fast and precise (Figure 10) \( \pm 25 \) ppm CO or 5 \% of the reading measurements in flue gas up to \( \sim 1750 \) °C temperature.

Figure 10. Calorimetric catalytic CO-sensor response at low CO concentration [1]

Figure 11. Calorimetric CO-sensor linearity [1]

Calorimetric CO-sensor has good >99.9 \% linearity (Figure 11) and reproducibility (Figure 12) in a wide CO concentration range up to 2.5 \% CO but to improve accuracy a calibration would be always recommended using upper limit of CO concentration measurement range. Calorimetric catalytic CO-sensor is highly flow-sensitive and gas flow stabilization and a flow sensor should be implemented to improve CO-sensor reliability in extractive combustion analyzer [1].

Figure 12. Calorimetric catalytic CO-sensor response at high CO concentration

Conventional catalysts for calorimetric methane sensors and combustion reaction are based on noble metals, \textit{i.e.}, palladium (Pd) or platinum (Pt) deposited on oxides such as cerium oxide (CeO\(_2\)) or metals. A calorimetric catalytic CH\(_4\) sensor has a good linearity (Figure 13) and \( \sim 30 \) s response for 90 \% signal (Figure 14). Flue gas flow control extractive environment would contribute to CH\(_4\)
detection reliability with better accuracy and reproducibility in the very challenging temperature, pressure, flow and chemicals flue gas combustion environment [20].

Only two available major methane gas sensing technologies – catalytic calorimetric and TDLS/QCLS – have so far found practical application in instrumentation on market because they produce quite reliable and accurate (about 2...5 % error) values in the extractive or across the stack mode.

Tunable diode laser spectroscopy (TDLS) for CO, CH₄ and oxygen measurements

Tunable diode laser (TDL) and quantum cascade laser (QCL) spectroscopies are innovative optical measurement techniques utilizing lasers with different wavelength (Figure 15), e.g., tunable diode and quantum cascade [22-27].

TDL and QCL spectroscopies are highly distinguished from the conventional process photometry with the laser’s ability to be scanned across the narrow combustion species absorption peaks many times per second by trimming the current through the laser. With a typical scan in the range of 0.2 to 0.3 nm, the laser would provide much higher selectivity in applications. The IR absorption spectrum of oxygen (O₂), carbon monoxide (CO) and methane (CH₄) is like a fingerprint, providing gas identification at different detection wavelengths (see Figure 16 for O₂).

The detection wavelengths and limits of some combustion gases are summarized in Table 1. A series of process adaption application tools were developed for in-situ, extractive or in cross-pipe installations. The probe design would not require a special optical path alignment but would deliver single point measurement. In the probe design the laser source and detector are located in the probe housing with the laser beam being reflected at the probe end, back to the detector by
quartz prism, with max operation temperature of <250 °C or by gold mirror with max operation temperature of <430 °C [20].

Table 1. Combustion gases absorption lines and detection limits [20]

| Target gas                      | Wavelength (µ) | Detection limit (ppm) |
|---------------------------------|----------------|-----------------------|
| Methane (CH₄)                   | 1.65 (TDL)     | 0.15                  |
|                                 | 7.9 (QCL)      | 0.003                 |
| Carbon monoxide (CO)           | 1.57           | 30                    |
|                                 | 2.33           | 0.5                   |
|                                 | 4.80           | 0.01                  |
| Oxygen                          | 0.76           | 8.956                 |
| Carbon dioxide (CO₂)           | 1.96           | 3                     |
| Nitric oxide (NO)              | 1.8            | 60                    |
|                                 | 2.65           | 1                     |
| Nitrogen dioxide (NO₂)         | 0.68           | 0.3                   |
| Water (H₂O)                    | 1.39           | 0.06                  |

With a unique folded-path design, the probes can be installed in almost all pipes and stacks with no alignment needed. The TDL or QCL analyzer can serve as a probe using deflection mirror or quartz prism (Figure 17) for single point near real-time measurement or would provide extractive measurement across the duct or pipe (Figure 18). Temperature and pressure variation in the process has to be compensated and might cause an additional error in the measurements. IR light reflection at high temperatures, combined with wide background radiation from the fire box and process windows fouling, might bring additional challenges in many applications. There are quite accurate TDL analyzers with ~±1-2 % error and ppm or even sub-ppm detection limits in some applications.

![Figure 17. In-situ probe, 250-430°C limit (Au-mirror or quartz prism)](image)

![Figure 18. Across the stack installation [29]](image)

However, laser analyzers are much more expensive with additional expensive and very time-consuming installation compared to the zirconia O₂-analyzers. TDL O₂, CO and CH₄ analyzers are mostly covering special chemicals production and very high >1000°C temperature applications.

Conclusions

After more than 50 years, zirconia potentiometric oxygen gas sensing technology is still dominating combustion market with the high reliability, accuracy and low price. State of the art is the simultaneous measurement of oxygen and COₑ by means of one and the same probe. Calorimetric catalytic and mixes potential sensors are implemented for COₑ measurement. TDLS/QCLS are also applied for CO and for oxygen measurements. They are about 5 times more expensive compared to the zirconia O₂-analyzers and additional require very expensive installation. TDLS O₂ analyzers mainly cover the production of specialty chemicals and very high >1000 °C
temperature applications. TDLS/QCLS and calorimetric catalytic sensors are used in combustion process for methane measurement. Only two methane gas sensing technologies – catalytic calorimetric and TDLS/QCLS – have so far found practical application in instrumentation in the combustion market as in extractive devise or across the stack for in situ measurements.

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