Towards a Novel Optical Trace Oxygen Sensor for Commercial Use

G.R. McDowell, F.G. Farrow, M. Uttamal, A.S. Holmes-Smith
School of Engineering & Built Environment
Glasgow Caledonian University
Glasgow, UK
gary.mcdowell@gcu.ac.uk

C. Mitchell, P. H. Shannon
SST Sensing Ltd
Coatbridge, UK
www.sstsensing.com

Abstract—This paper presents the development of a novel luminescence-based trace oxygen sensor that will eventually lead to the production of a commercial sensor for use in gaseous sensing applications. The proprietary sensing formulation provides the means of a robust and stable sensor using a metalloporphyrin complex as the luminescent indicator that is efficiently quenched by oxygen. The spectroscopic properties of the oxygen sensitive layer have been carried out and appropriate optical components and electronics sourced and built into a low cost, compact, miniaturized device. This novel oxygen sensor operates in the 0–10,000 parts per million (ppm) range and offers an alternative to traditional trace oxygen sensors, such as galvanic electrochemical, zirconia and other luminescence-based ppm sensors.

Keywords—Gas sensor, luminescence quenching, trace oxygen

I. INTRODUCTION

Oxygen (O2) is one of the most important chemical species and is vital for the survival of most living organisms [1]. However, in applications such as nitrogen (N2) generation, industrial gas manufacturing and food packaging there is a need to ensure O2 is absent, even at the lowest concentration levels. Therefore, trace O2 parts per million (ppm) sensors have been developed with the most common technologies summarised in Table 1.

<table>
<thead>
<tr>
<th>Property</th>
<th>Electrochemical</th>
<th>Zirconia</th>
<th>Paramagnetic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cost</td>
<td>Low</td>
<td>High</td>
<td>High</td>
</tr>
<tr>
<td>Operating temperature (°C)</td>
<td>-20 to +55</td>
<td>-100 to &gt;500</td>
<td>0 to +50</td>
</tr>
<tr>
<td>Calibration intervals</td>
<td>6–12 months</td>
<td>~1 year</td>
<td>&gt;1 year</td>
</tr>
<tr>
<td>Life</td>
<td>2–3 years</td>
<td>&gt;5 years</td>
<td>&gt;5 years</td>
</tr>
</tbody>
</table>

TABLE I. SUMMARY OF TECHNOLOGIES USED FOR TRACE OXYGEN SENSING.

Electrochemical sensors are the most widely used as they are low cost, small, accurate and have a reasonable operational lifetime. However, they suffer from long recovery times and deplete when exposed to ambient O2 (air) thus calibration and/or replacement is required periodically. Zirconia and Paramagnetic sensors offer longer operational lifetimes however, they are expensive and Zirconia sensors require a high temperature for operation. The conventional method is the Clark Electrode [2] but O2 is consumed, prone to electrical interferences and has a limited operational lifetime. Clark-type microelectrodes have been developed but they are expensive to fabricate, fragile and their usefulness is limited. [3]. For applications where these considerations are important, optical methods, in particular, luminescence-based trace O2 sensors have been developed commercially by Aanderaa, Centec, PreSens and Pyro Science to name a few.

It is well known that O2 is an efficient quencher of luminescence [4]. Luminescence-based sensing is now regarded as one of the best techniques for O2 sensing [5] due to high detection sensitivity, no analyte consumption and there are a vast array of luminescent indicators and sensing platforms that can be selected to suit the desired application [6]. The luminescence emission intensity and/or excited state lifetime (decay time) of a luminescent indicator is related to O2 concentration through the Stern-Volmer equation:

\[
\frac{I_0}{I} = \frac{\tau_0}{\tau} = 1 + K_{SV} \cdot ppO_2 = 1 + k_q \tau_0 \cdot ppO_2
\]

where \(I_0, I, \tau_0\) and \(\tau\) are the luminescence intensity and decay time in the absence and presence of the quencher respectively. \(k_q\) is the biomolecular quenching constant, ppmO2 is the partial pressure of oxygen and \(K_{SV}\) is the Stern-Volmer quenching constant. Plots of \(I_0/I\) and \(\tau_0/\tau\) versus O2 concentration should yield a straight line. In many cases the line deviates from linearity which indicates that the luminophore exists in more than one environment and/or dynamic and static quenching are occurring concurrently. In these cases, nonlinear forms of the Stern-Volmer equation have been proposed [7], [8].

SST currently market a luminescence-based O2 sensor (LuminOx) which measures O2 in the 0–25% range or 0–300 mBar ppmO2 and can measure O2 in environments where the temperature range is –30 °C to +60 °C. This is achieved through an extensive calibration process. Here, a new development of the LuminOx sensor for trace O2 measurement is described. The aim is to manufacture a low cost, small and robust gaseous oxygen sensor for this market place which is capable of operating for more than 5 years. The luminescent indicator within a proprietary polymer formulation will be
changed from a ruthenium complex to a metalloporphyrin complex to ensure good sensitivity to low concentrations of O2. Also, the sensor will be developed to have a working temperature range of –30 to +60 °C which is an improvement on other commercially available luminescence-based trace O2 sensors which tend to operate from –5 to 50 °C.

II. EXPERIMENTAL

All chemicals, reagents and polymers were purchased from commercial suppliers and were used as received. The new O2 sensitive layer is based on the luminescent metalloporphyrin complex Platinum (II) 5,10,15,20-Tetrakis(2,3,4,5,6-pentafluorophenyl)-porphyrin (PtTFPP) immobilised within a proprietary sensing formulation.

A. LuminOx Sensor Design for Trace Oxygen

The O2 sensitive layer is embedded within a miniaturised (18 x 20 mm), compact, optical housing shown in Fig. 1. The sensing layer is deposited onto a 600 nm cut-on dichroic mirror and fixed within an optical carrier (Fig. 1(a)) at an angle of incidence (AOI) of 45°. In the same carrier, a UV blocking 535 nm bandpass filter is inserted at AOI 0° that sits directly in front of a 535 nm green LED (Fig. 1(b)) connected to a customised printed circuit board (PCB) containing a photodiode that collects the luminescence signal passing through the dichroic mirror. The PCB also contains temperature and pressure sensors so that measurements can be corrected for changes in the luminescence response due to these environmental variables. The optical carrier is fixed on top of the PCB (Fig. 1(c)). For the current LuminOx sensor a housing is added containing an oleophobic and hydrophobic PTFE membrane (Fig. 1(d)) and for trace O2 (ppm) sensing a flow through housing is fixed into place (Fig. 1(e)).

B. Spectroscopic and Sensor Characterisation

Spectroscopic characterization of the sensing polymer was carried out on commercial spectrometers prior to insertion into the LuminOx sensor. Luminescence excitation and emission spectra were collected using a HORIBA Jobin Yvon Ltd FluoroMax-3 spectrophotometer. Luminescence lifetime (decay time) measurements were carried out on a HORIBA Jobin Yvon 5000U lifetime instrument using either Multi-channel scaling (MCS) and/or Time-correlated single photon counting (TCSPC) depending on ppO2. O2 and O2 free nitrogen (OFN) gases were purchased from BOC gases. An Environics Series 2020 computerized gas dilution system was used to control the concentration of O2. A factory calibrated SST LuminOx (0–300 mBar ppO2) sensor was used as a reference. Trace oxygen measurements were carried out by characterizing the luminescence decay time of the sensing matrix to changes with O2 concentration from 0–10,000 ppm using the gas blender to mix a cylinder of 98.7% N2/1.3% O2 or >99.98% N2/130 ppm O2 with OFN.

III. RESULTS

A. Oxygen Sensor Characterisation

The metalloporphyrin complex PtTFPP has an intensely absorbing Soret-band (~390 nm) corresponding to an S0 → S2 transition and weaker absorbing Q-bands (~504 nm and ~538 nm) corresponding to S0 → S1 transitions in the visible spectrum [6]. When illuminating PtTFPP continuously for 1 hour at λex = 390 nm and λem = 535 nm in air (207 mBar ppO2) the luminescence intensity dropped by 0.036 ± 0.010% and 0.003 ± 0.002% at each wavelength respectively. Therefore, the rate of photodegradation of PtTFPP was reduced 10-fold when λex = 535 nm. Importantly, bright LEDs emitting at this wavelength are readily available. The % photodegradation is a marked improvement on that observed for the same luminescent indicator in other sensing matrices where λex = 540 nm [9]. The longer excitation wavelength ensures that PtTFPP is less reactive in the excited state due to the S0 → S1 transition. In addition, it has also been suggested that PtTFPP exhibits excellent photostability due to the presence of the strongly electron-withdrawing character of the pentafluorophenyl substituents. The sensing formulation in this work contains a fluoropolymer which may further enhance the photostability. This renders PtTFPP less reactive toward oxidation by singlet oxygen and prolongs the life of the luminescent indicator [9].

The luminescence decay time of PtTFPP in the sensing matrix in the absence of O2 (t0) and at ambient temperature was 83.5 µs and dropped to 1.1 µs in air (201 mBar ppO2). The Stern-Volmer plots (SVPs) of the PtTFPP sensor and the LuminOx sensor are shown in Fig. 2. Comparing the sensitivity shows that the PtTFPP sensor is 9 times more sensitive as the Stern-Volmer quenching constants (KSV) were 0.3893 ± 0.0066 mBar⁻¹ (R² = 0.9997) and 0.0433 ± 0.0015 mBar⁻¹ (R² = 0.9986) for the PtTFPP sensor and the LuminOx sensor respectively.

![Fig. 2. Stern-Volmer plots of t0/t versus ppO2 for the SST LuminOx sensor (λex = 440 nm, λem = 590 nm) and the new trace O2 sensor (λex = 375 nm, λem = 645 nm) at ambient temperature.](image)
Fig. 3. Luminescence intensity response of the PtTFPP O₂ sensor when switching between nitrogen (N₂) and air (207 mBar ppO₂).

\[ \lambda_{ex} = 535 \text{ nm, } \lambda_{em} = 645 \text{ nm.} \]

Fig. 3 shows the luminescence intensity response of PtTFPP in the polymer matrix in going from N₂ to air (207 mBar ppO₂) and back for several cycles on the commercial luminescence spectrophotometer. Critically, the luminescence intensity was reversible and highly repeatable when switching between N₂ and air, indicating that the sensor is unaffected by exposure to ambient O₂, an advantage over electrochemical ppm sensors as they deplete with time when exposed to ambient O₂. Within the LuminOx system, the T₉₀ (increase or decrease of luminescence signal to 90% of its original value) response and recovery times were 13.9 ± 0.5 s and 23.9 ± 0.8 s respectively. It should be noted that the response and recovery time of a luminescent indicator is not only affected by the permeability of the polymer matrix to O₂ but is also dependent on the thickness of the sensing layer. In this work, the sensing layer was ~500 µm. Decreasing the thickness of the sensing layer will result in a faster response and recovery time due to faster equilibration of O₂ through the polymer matrix.

B. Trace Oxygen Analysis

Fig. 4. Decay time of the sensor with increasing O₂. Note that ppm can also be expressed as µBar. Inset is a Stern-Volmer plot of τ₀/τ versus O₂ concentration.

The decay time of PtTFPP in the absence of O₂ (τ₀) was 85.6 µs and decreased with increasing O₂ shown in Fig. 4 to 18.2 µs at 10,000 ppm thus τ₀/τ = 4.71 (Fig. 4 inset). A linear correlation (R² = 0.9999) between τ₀/τ and ppm was observed over the wide O₂ ppm concentration range, (Fig. 4 inset) beneficial for calibration of the sensor. Going from 0 ppm to 25 ppm O₂ caused the decay time of PtTFPP to drop ~0.9 µs demonstrating the high sensitivity of the sensor to changes in O₂. Preliminary tests were also conducted in an O₂ free environment at −30 °C and +60 °C with decay times of PtTFPP being 100.5 µs and 62.9 µs respectively to determine the sensitivity and usefulness of the sensor to changes with temperature.

IV. CONCLUSIONS & FURTHER WORK

A novel luminescence-based trace O₂ sensor has been described that will lead to the production of a commercial product for SST. The sensor will be able to compete within the market (Table 1) and will offer an attractive alternative to these commercial trace O₂ sensors as it will be low cost, robust, compact and does not deplete when exposed to ambient O₂.

Most commercial luminescence-based trace O₂ sensors are used in the detection of dissolved oxygen (DO) thus have been developed to detect parts per billion (ppb) as well as ppm. One drawback is their operating temperature range which tends to be −5 to +50 °C. In addition, the life of most luminescence-based trace O₂ sensors is ~3 years if stored in the dark and under ambient conditions. The aim is to improve on these aspects with the new trace O₂ sensor i.e. −30 to +60 °C and >5 years life under continuous operation. Also, the majority of trace O₂ sensors are opt(r)ode devices. The new LuminOx is unique as it is a compact, complete, miniaturized device thus no additional electronics or accessories are required.

Further work is underway to determine the repeatability of the PtTFPP based trace O₂ sensor over the full temperature (−30 to +60 °C) and O₂ concentration range (0–10,000 ppm). This will include rigorous photo- and long-term stability tests to determine the longevity of the sensor.

REFERENCES


