FIBER-OPTIC SENSOR FOR THE EXPRESS CONTROL OF THE CHEMICAL COMPOSITION

Y. P. Sharkan¹, N. B. Jytov², I. I. Sakalosh¹, J. J. Ramsden³, M. Y. Sichka¹, I. I. Popovich¹, S. O. Korposh¹

¹Institute of Solid-State Physics & Chemistry, Uzhgorod National University, Voloshina St 54, 88000 Uzhgorod, Ukraine, e-mail: shark@univ.uzhgorod.ua
²“Technomedica”, Moscow, Russia
³Department of Advanced Materials, Cranfield University, MK 43 0AL, UK

Abstract. On the example of the aqueous-ethanol solutions we proposed fiber-optic sensor system of the evaluation of the concentration of the liquid solutions with the known qualitative composition. The deposition of the thin film chalcogenide layer with the high refractive index permitted to improve in one order the precision of the refractive index measurements of the aqueous solutions for the quartz Y-shaped splitter. We proposed and tested the method of the definition of the aqueous solutions concentration thanks to the measurement of the time of the total drying of the film of the solution on the fiber end, and thanks to the measurements of the changes of the interference signal which appears on the film in course of the drying process.

Keywords: Fiber-optic sensor, Fabry-Perot interferometer, aqueous solutions concentration, the process of drying of the film of the solution

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**Introduction**

Determination of the chemical composition of liquid solutions continues to be a task of current importance, covering a very wide spectrum of applications ranging from the control of physico-chemical technological processes and ecological monitoring to the analysis of medical and biological processes. The development and creation of sensors for rapid analysis is especially topical. The sensors should also be cheap, small in size, requiring a small amount of the investigated medium and able to work under the conditions of intensive electromagnetic fields and ionizing radiation.

Fiber-optic sensors meet the demands for precise, rapid and reliable monitoring of media of different kinds [1]. The small size of the mono-fiber permits its use for the analysis of a small amount of the investigated medium. This is especially important for medical purposes. Sensors with sensitive elements on the fiber end are suitable for the determination of chemical compositions [2–5]. The sensitive element can either be the fiber end itself or a film or layer from a material sensitive to the influence of the investigated medium [6].

The principle of operation of such sensors can be the amplitude measurement [7–9], in which the change of a signal is due to change of absorption or refractive index of the investigated environment, with which the fiber end contacts. More sensitive are interference fiber-optic sensors, in which the film from the investigated material [10, 11] or created at the fiber end beforehand the Fabry–Perot interferometer, the parameters of which vary during the interaction with the investigated medium [3], are used in the role of interferometer [12].

In this work, a fiber-optic sensor for the determination of quantitative changes in chemical composition of solutions is presented. The operating principle of the sensor is a method of determination of the reflection coefficient at the boundary between the fiber end and the investigated medium and the determination of the parameters of the film drying process, which occurs at the fiber end during withdrawal of the fiber from solution.

**Direct measurements of the solutions concentration**

The most convenient method for the determination of the refractive index of the solutions is the measurement of the amplitude of the inverse light reflection due to reflection coefficient at the boundary between fiber end and the investigated medium. In this case, using the single-mode quartz optic fiber the reflection coefficient is defined as follows:

\[ R = \left( \frac{n_f - n_m}{n_f + n_m} \right)^2, \]  

where \( n_f \) — the refraction coefficient of quartz core of the single-mode fiber; \( n_m \) — refraction coefficient of the investigated medium.

The sensor (see, please, Fig.1) contains a Y-shaped splitter that divides the power equally between the input and output channels. From the input channel the signal propagates into the common channel, which is placed in contact with the sample. The optical signal is reflected from the fiber-sample interface, returns to the common channel, and then propagates into the output channel. The output sig-
nal is amplified and sent through an ADC to reach a computer for information processing.

![Diagram of the mono-fiber optical sensor](image)

Fig. 1. Scheme of the mono-fiber optical sensor: 1 — light source ($\lambda$ = 0.95 μm in the present work); 2 — investigated medium (sample); 3 — fiber optic Y-shaped splitter; 4 — liquid film on the fiber end; 5 — photodiode and amplifier; 6 — ADC.

Fiber-optic Y-shaped splitter contains the quartz fiber with core diameter and covering 8 and 150 μm, correspondingly, as far as exactly the quartz fibers, comparatively with polymer, due to the chemical stability provide the possibility to investigate the parameters of the physiological liquids in the “in vivo” mode.

The light emitting diode was used as a light source with the illumination wavelength 950 nm and possesses the width of the emission band at the level of 0.5 of about 30 nm. Unfortunately use of quartz fibers with the core refractive index of 1.45, leads to a low reflection when measuring the aqueous solutions, in which the refractive index doesn’t differ too much from the $n_f$. The use of the fibers with the high refractive index, for example chalcogenides, will considerably increase the reflection coefficient at the fiber end—investigated medium interface. Fig. 2 shows the calculated dependences of the fiber end reflection value from the investigated medium for quartz and chalcogenides fibers with a refraction index of 2.05 in the range of refractive indexes inherent to the aqueous solutions.

![Graph of fiber end reflection](image)

Fig. 2. Calculated dependences of fiber end reflection from the investigated medium for a quartz fiber $n_f$ = 1.45 (curve 1), and for a chalcogenide fiber $n_f$ = 2.05 (curve 2); the inset expands the plot for refractive indexes typical for aqueous solutions.

Changing the concentration of the investigated medium, with which the common channel end of the chalcogenide monofiber sensor is in contact, the reflection coefficient $R$ also changes; and change of signal amplitude is recorded:

$$U = kR = k \left( \frac{r_{12} + r_{23}}{1 + r_{12} r_{23}} \right) - 4 r_{12} r_{23} \sin^2 \left( \frac{\delta}{2} \right),$$  (2)

$$r_{12} = \frac{n_f - n_{lm}}{n_f + n_{lm}}, \quad r_{23} = \frac{n_f - n_m}{n_f + n_m}, \quad \delta = \frac{2 \pi n_f d}{\lambda},$$  (3)

where $k$ is a proportionality coefficient; $U$ — the signal amplitude output of the photodiode; $r_{12}$ and $r_{23}$ — reflection coefficients on the boundary quartz single-mode fiber — thin film chalcogenide layer — the investigating medium correspondingly; $\delta$ — phase shift at the light passing at wavelength $\lambda$ through the thin film chalcogenide layer with thickness $d$; $n_f$ — the refractive index of the core of the optic single-mode fiber; $n_{lm}$ — the refractive index of the thin film chalcogenide layer; $n_m$ — the refractive index of the investigated solution. Hence by simply making optical contact of the mono-fiber end of the sensor with the investigated medium it becomes possible to determine the index of refraction of the given medium. Upon a change of the chemical composition of the solution its refractive index varies.

Fig. 3 shows the reflection coefficient of the fiber end—air interface and how the reflection coef-
ficient changes during immersion of the fiber end into solutions containing different concentrations of ethanol.

Fig. 3. Experimental determination of the reflectivities of aqueous ethanol solutions and pure water.

From these data the calibration curve of the dependence of the optical signal on the ethanol concentration was plotted (Fig. 4).

The measured value of the refractive indexes of the ethanol solutions are in a good agreement with tabular data [13] within an accuracy 10^{-3}. It is worth noting that the repeatability of these results was very good. Thus the suggested fiber-optic sensor (Fig. 1) of the amplitude type with single-mode quartz Y-shaped splitter permits directly to measure the refractive index of the investigated solution and due to the change of the refractive index to determine the concentration change.

Also, monitoring of the solutions concentration of the known chemical composition may be carried out using the sensor calibration on the standard samples, in this case there is no necessity to measure directly the refractive index of the investigated solution, that permits to use the multi-mode optic fiber and by this to decrease the loses of the optic signal on the optic connections and to increase the reliability of the sensor as a whole.

The process of solution drying on the fiber end

The second procedure reported here is the study of the dynamics of optical signal changes during the withdrawal of the fiber end from solution and during the drying processes of the film which is formed on the fiber end; during withdrawal of the fiber from a wetting solution, a drop forms on the fiber end, which then remains due to surface tension; its size depends on the composition of the solution from which it is formed (we neglect here any possible changes of the drop composition compared with the bulk composition due to the proximity of the air water interface in the drop).

The drying process was studied with the help of the sensor depicted on figure 1. It was held at the same conditions of 25 °C and 55 % relative humidity. Aqueous ethanol solutions were chosen for the experiment.

In order to explain the processes which occur on the fiber end of the sensor after breaking optical contact of the fiber with the investigated medium, a CCD camera was used.

Dependence of the change of the signal value in the process of measurement is shown at Fig.5.

Fig. 4. Calibration curve, linking the optical output signal with ethanol concentration.

Fig. 5. The change of the signal value in the process of measurement: 1 — the monofiber end-air interface; 2 — fiber end-investigated medium interface; 3 — withdrawal of the fiber end from solution and formation of a drop-shaped film on the monofiber end; 4 — reduction of film thickness as a result of drying.
The 1st region corresponds to the fiber end-air interface; the 2nd region to the fiber end-investigated medium interface; the 3rd region to the formation of the drop-shaped film on the fiber end.

The sharp increase of the signal amplitude initially observed is due to the drop-shaped form of the film; all the light is reflected into the mono-fiber; the subsequent decrease of the signal is due to a change of the drop geometry, and correspondingly the radius of curvature of the spherical mirror, which exists at the investigated medium-air interface is continually changing. Further drying leads to an increase of the radius of curvature, and eventually the solution film on the fiber end becomes plane-parallel and light interference arises, due to the formation of a Fabry-Perot interferometer on the fiber end.

The 4th region shows a changing in the reflected signal as the film thickness on the fiber end decreases due to evaporation of the plane-parallel solution film; when the film thickness reaches the coherence length for the sensor, the interference occurs.

Additional confirmation of such division of the dependence of the amplitude on the film form was received in the course of the investigation of evaporation with the help of the video recording of the process on the CCD camera. On the end of the mono-fiber the drop of the investigated solution is formed. In time the evaporation of the liquid from the end of the mono-fiber occur, the volume decreases and the radius of the drop curvature increases, as a result on the end of the mono-fiber the plane parallel film is formed in which the multiple-beam interference is taking place.

Fig. 6 shows the optical signal of the drying dynamics for aqueous ethanol solutions of different concentrations. The correlation between the drying time and the solution concentration is evident, i.e. with increasing ethanol concentration film drying is faster. The calibration graph of the drying time dependence on ethanol concentration $C=C(t)$ is plotted at Fig. 7.

![Fig. 7. Dependence of the drying time of the film on the concentration of the aqueous solution of the ethanol.](image)

In such way by the measurement of the time of the process of the total drying of the aqueous solution on the end of the fiber it is possible to determine its concentration.

**Interference in the drying solution film**

The interference that appears in the drying film on the fiber end enables the development of interference fiber optic sensors for the determination of solution concentrations, which are more sensitive in comparison with amplitude sensors.

For studying the interference of the solution drying on the fiber end a fiber-optic spectrophotometer (Ocean Optics HR2000) and a white light source (Ocean Optics HL-2000 tungsten halogen lamp) were used. Measurements of the reflection spectrum from the end of the fiber-optic Y-shaped splitter with a diameter of 400 $\mu m$ was carried out; the diameter of the connector of the common channel, which was immersed into the investigated medium, was 3 mm. Aqueous ethanol solutions were also chosen for this experiment.

At Fig. 8, the reflection spectra of the drop formed on the end of the optical connector for different drying time of 50% aqueous ethanol solution are shown.

![Fig. 8. Reflection spectra of the drop formed on the end of the optical connector for different drying time.](image)
Curve 1 corresponds to the reflection spectrum from the investigated solution when the fiber end is in solu-
tion, curve 2 to the spectrum of a solution drop formed on the fiber end after withdrawing it from the 
investigated liquid. During the drying process a change of reflection occurs due to change of the drop 
form (curve 3). Gradually the radius of curvature of the drop is so increased, that interference in the 
solution film appear (curve 4), characteristic for a thin-film Fabry-Perot interferometer.

Further decreasing of the solution film thickness due to drying leads to a shift of the interference 
maxima in the reflection spectrum (curve 5).

Using the well-known equation for interference, it is possible to assess only the optical thickness $n_md$, 
but it is impossible to define the absolute value of the refractive index and thickness, because both values 
vary simultaneously in time for the solutions:

$$\frac{\lambda_2 - \lambda_1}{\lambda_2 \lambda_1} = 2n_md$$

where $\lambda_1$ and $\lambda_2$ are the wavelengths corresponding to the positions of two neighbor interference peaks; 
$n_m$ is refractive index of the solution film at the time of measurement; and $d$ is the geometric film thick-
ness at the time of measurement.

The decrease of the geometrical film thickness takes place due to evaporation; and the change of the index of refraction takes place due to the change of the film chemical composition during the dry-
ing, which occurs because of the differential rate of evaporation of the various solution components.

The index of refraction will be constant in time only for pure liquids. Thus, measuring the reflection 
spectrum, in other words the change of the interference peak positions in time, and calibrating the sig-
als for given solution compositions, application of the given technique for the quantitative determina-
tion of solution compositions whose composition is qualitatively known is possible.

However, the processing of the spectral interference curves requires powerful software, because of the 
presence of large data arrays, and needs also quite expensive and complex hardware, which consider-
ably complicates the application of this method.

Therefore, a system of monitoring the interference in the drying liquid film without spectral de-
termination is offered. For this purpose the scheme depicted in figure 1 was used, single mode quartz 
fiber was used as the fiber-optic Y-shaped splitter, and a semiconductor laser diode with an illumina-
tion wavelength of 1320 nm and width of emission band at the level of 0.5 of about 5 nm, mean emitted 
power of 1 mW and frequency of direct modulation 10 kHz, was used as the emitter. The connector of a 
single mode optical fiber with a diameter of 2.5 mm was used as a fiber end of the common channel, 
which was immersed into the investigated solution and on which the drying drop was formed.

The drying of aqueous ethanol solution drops of various concentrations formed on the end of the 
single mode optical fiber was investigated.

Fig. 9 shows the graph of the reflection change for the complete process including immersion into the 
solution, withdrawal from the solution and drying. 1 — pure water; 2 — 50 % ethanol solution; 3 — 80 % ethanol solution.

Fig. 9. Reflection changes for the complete process comprising immersion into solution, withdrawal from solu-
tion and drying. 1 — pure water; 2 — 50 % ethanol solution; 3 — 80 % ethanol solution.

The drying of aqueous ethanol solution drops
described in the first part of this paper, the times of drying are also much bigger.

The course of the complete drying process, as shown in figure 9, correlates with the data presented in the first part of this work, i.e. with increasing ethanol concentration the time for complete drying is decreased.

Fig. 10. The interference picture for a 50% aqueous ethanol solution.

At the same time, for the given measurement system, the region of evaporation of the formed planar film appears more clearly. The reason is connected with the different film geometry at the fiber end (the bigger dimensions in comparison with the first part of this work) and coherence length of the emitter.

The study of the interference picture for the investigated solutions has shown that as well as the spectral changes of reflection during film drying, the rate of the change of the film optical thickness \((n_d)\) varies in time for solutions. This is evident from the change of period of the interference maxima (Fig. 10 and Table 1). During the film drying of their pure components individually (ethanol or water), the period remains at a constant value.

Thus, by measuring the speed of the period change of the interference maxima the determination of the concentration of the drying liquid is possible. In other words, it is necessary to solve the task of measurement of the distance between the peaks on the interference picture. One possible hardware implementation of a solution to this task is introduction of a differential chain after the signal amplifier (Fig. 11, a), which will allow the creation of impulse fronts from the incoming signal (Fig. 11, b) with a much higher amplitude than the oscillations of the average signal level (Fig. 11, c). After the differential chain, a comparator (threshold device) is placed, which will evaluate the moments of intersection of the impulse with the value of the threshold level (Fig. 11, d).

Fig. 11. Schematic illustration of the determination method of the distance between the peaks on the diagram of the interference pattern.

Table 1

| Number of period | \(H_2O\) | \(C_{2}H_{5}OH\) 50% | \(C_{2}H_{5}OH\) 80% | \(C_{2}H_{5}OH\) 96% |
|-----------------|---------|---------------------|---------------------|---------------------|
| 1.6809          | 1.0588  | 0.6856              | 0.4874              |
| 1.6816          | 1.0630  | 0.6901              | 0.4867              |
| 1.6808          | 1.0658  | 0.6930              | 0.4853              |
| 1.6813          | 1.0697  | 0.6975              | 0.4873              |
| 1.6803          | 1.0731  | 0.7012              | 0.4863              |
| 1.6812          | 1.0776  | 0.7035              | 0.4867              |
| 1.6813          | 1.0816  | 0.7078              | 0.4853              |
| 1.6805          | 1.0847  | 0.7109              | 0.4873              |
| 1.6812          | 1.0877  | 0.7157              | 0.4874              |
| 1.6808          | 1.0921  | 0.7189              | 0.4863              |
| 1.6816          | 1.0961  | 0.7222              | 0.4853              |

Hence, the periods of the interference maxima will be estimated. Further measurement of the temporal distance between impulses can be accomplished by filling the measured interval by time impulses and counting them, which can be easily carried out with the help of a personal computer.

On the base of the received results the plot of the dependence of the middle value of the period of the
interference maxima on the concentration of the aqueous ethanol solution was constructed (Fig. 12), which permits to measure the ethanol concentration in aqueous solutions.

![Graph showing dependence of the middle value of the period of the interference maxima on the concentration of the aqueous ethanol solution.](image)

Fig. 12. Dependence of the middle value of the period of the interference maxima on the concentration of the aqueous ethanol solution.

The method of determination of the components concentration for the qualitatively known solution composition was developed that permits to observe the quantitative changes in the composition of the film of the investigated solution and also according to the value of the change of the period of the interference maxima in the process of the measurement, to determine whether the investigated solution is ideal or not ideal [14].

Conclusions

The work carried out and discussion have shown that:

1. By attaching a chalcogenide glass fiber with a high refractive index to the common channel of a fiber-optic Y-shaped splitter, the sensitivity of the refractometer is increased by almost one order of magnitude compared with conventional silica glass fibers.

2. The measurement of the changes in the reflection coefficient during drying of a solution film on the end of the optical fiber allows the sensitive determination of the concentration of aqueous solutions whose composition is known qualitatively.

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