Spectrally Resolved White Light Interferometry for NaCl Aqueous Solutions Refractive Index Measurement

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In this work, a technique for comparing refractive indices for solutions of low concentrations has been developed. The method of direct spectral detection developed by the authors has been using. The solutions of low and ultra-low NaCl concentrations were compared with each other, as well as with the respective water controls. The solutions were prepared using the method of serial dilution with different types of mixing at each step: conventional stirring or intensive shaking. Measurements were made for water solutions of NaCl subjected to 6–12 and 30 hundredfold dilution, as well as for water controls, i.e. water subjected to the same technological procedure as the tested salt solutions.

Keywords: interferometry, spectral coding, aqueous solution, refractive index, NaCl

INTRODUCTION

It is known that the properties of low-concentration aqueous solutions have attracted great interest in recent years [1–3]. These studies require the use of modern spectroscopic or mass spectroscopic equipment. This is a rather laborious and costly procedure. There are no such disadvantages in an interferometric fiber-optic sensor, which combines a simple design, small dimensions and high sensitivity. The measured value of the interferometer optical path difference is proportional to the refractive index of the medium. One of the necessary conditions for the use of such systems is the development of a precise and reliable algorithm for processing signals. In this work, we develop the algorithm proposed before [4–6 and others] for the interferometer length measurement to measure the small variations of aqueous solutions refractive index.

In this work the highly diluted NaCl solutions were investigated. It was shown that their refractive index varies by values of $(1...5) \times 10^{-5}$ depending on the method of their preparation (using stirring or shaking). Differences were established between solutions of low and ultra-low NaCl concentrations prepared using shaking, and the respective water controls. The differences were observed between the refractive indices of water and water controls. Moreover, differences were shown between different samples of water controls prepared using shaking, i.e., water subjected to different numbers of serial dilutions.

METHODS

The experimental setup is shown in Figure 1. Broadband radiation from a source (superluminescent diode from www.superlumdiodes.com) with a central wavelength of ~800 nm and a spectral width of...
~35 nm is introduced into a single-mode fiber optic coupler (made of Corning HI-780 fiber) and enters the probe, reflecting from which it is fed to the spectrometer through the other arm of the coupler. The probe is a Fabry–Perot interferometer formed by two ends of the same fiber waveguide. The fiber ends were Ag sputtered to increase the interferometry visibility. The frequency dependence of the interferometer reflection coefficient was sinusoidal, and this fact is used in the signal processing algorithm. The measured spectrum (Figure 2) is an oscillating function with a smooth envelope, which repeats the shape of the spectrum of the radiation source; the period of oscillations and their position determine the optical thickness of the interferometer (the product of the refractive index of the medium and the interferometer length).

For highly accurate determination of the refractive index of liquid media, a specific measuring cell was developed. It is presented at Figure 3.

The cell is a quartz cube with an edge length of 9 mm with two holes—one is horizontal with a diameter of 2.5 mm, the other is vertical with a diameter of 3 mm. The horizontal hole is intended for gluing the ends of optical fibers in ceramic ferrules (2.5 mm diameter), the gap between which formed a Fabry–Perot interferometer. The typical interferometer length was 50 µm. Vertical holes are designed to supply and drain the investigated liquid. To minimize the effect of temperature drift on the gap between the ends of the fibers, quartz was chosen as a cube material having a small thermal expansion.

**PROCESSING OF THE MEASURED SPECTRA**

The processing algorithm for the measured spectra is mostly based on [4, 6] and illustrated in Figure 4 (for clarity, discrete signals are shown as continuous).

The processing algorithm for the measured spectrum includes the Fourier transform, adaptive filtering (rectangular windowing is used for necessary correlation peak selection) in the time domain and the next Fourier transform, which gives the so-called “analytical signal”, the modulus of which coincides with the spectrum of the radiation source, and its phase gives the dependence of the phase difference of interfering waves on the frequency [4].

But in this work, we measured the increment of the refractive index of the medium filling the Fabry–Perot interferometer with unchanged interferometer length. In this case, we do not need the “phase unwrapping” procedure usually used for determining the optical thickness of the interferometer. As is known in interferometry, this procedure results in the problem of uncontrolled phase jumps caused by errors in the measurement of phase readings and the discrete nature of calculations. Thus, in our case, this problem is absent. The measurement results is a change (Δn) in the refractive index of NaCl aqueous solutions.
the medium caused by the replacement of the "reference" solution with the "sample" investigated.

The phase difference $\phi(k)$ of interfering waves at the frequency $\nu(k)$ ($k = k_{\min}, \ldots, k_{\max}$) is determined by the relation:

$$\phi(k) = \frac{4\pi L}{c} n(k) \nu(k) + \alpha$$  \hspace{1cm} (1)

where $k = k_{\min}, \ldots, k_{\max}$—optical frequency number, $L$—the interferometer length, $n(k)$—the refractive index at the frequency $\nu(k)$, $c$—velocity of light in vacuum, $\alpha$—the initial phase difference. As in [4, 6], the spectrum processing algorithm allows calculating an array of phase factors

$$e^{i\phi(k)} = \mu(k)$$  \hspace{1cm} (2)

where $k = k_{\min}, \ldots, k_{\max}$, which we denoted by $\mu(k)$, here $i$ is the imaginary unit.

For small (compared to unity) phase increments $\delta\phi(k)$ from (2), we obtain:

$$\delta\mu(k) = (k) 2i\delta\phi(k)$$  \hspace{1cm} (3)

this implies

$$\delta\phi(k) = \frac{\delta\mu(k)}{2i\mu(k)}$$  \hspace{1cm} (4)

In turn, for the phase increments $\delta\phi(k)$ formula (1) gives

$$\delta\phi(k) = \frac{4\pi L}{c} \nu(k) \delta n(k)$$  \hspace{1cm} (5)

where $\delta n(k)$ is the increment of in the refractive index caused by the medium replacement at the frequency $\nu(k)$ and $k = k_{\min}, \ldots, k_{\max}$.

By eliminating the values of $\delta\phi(k)$ from formulas (4) and (5), we obtain the change in the refractive index at the frequency with the number $k$:

$$\delta n(k) = \frac{c}{8\pi L \nu(k)} \frac{\delta\mu(k)}{i\mu(k)}$$  \hspace{1cm} (6)

where $k = k_{\min}, \ldots, k_{\max}$.

And, finally, we find the required value ($\delta n$)—the change in the refractive index caused by the medium change averaged over the considered optical frequency range:

$$\delta n = \frac{\sum_{k_{\min}}^{k_{\max}} \rho(k) \delta n(k)}{\sum_{k_{\min}}^{k_{\max}} \rho(k)}$$  \hspace{1cm} (7)

where the values $\rho(k)$ of the analytical signal modulus are taken as the most natural coefficients for spectral data weighting.

Each measurement of the difference between the refractive indices of the two media ($\delta n_{\text{exp}}$) was obtained by averaging $N = 100$ readings:

$$\delta n_{\text{mean}} = \frac{1}{N} \sum_{j=1}^{N} (\delta n)_j$$  \hspace{1cm} (8)

where $(\delta n)_j$ is the result of $j$ measurement ($j = 1, 2, \ldots, N$).

The root-mean-square error $\sigma(n)$ of measuring the difference in refractive indices was determined using the standard formula:

$$\sigma(n) = \sqrt{\frac{1}{N-1} \sum_{j=1}^{N} \left[(\delta n)_j - (\delta n)_{\text{mean}}\right]^2}$$  \hspace{1cm} (9)

Typical example of the measured ($\delta n$) time series under static conditions for some sample is shown in Figure 5. It allows us to estimate the accuracy of the method. The result is

$$\sigma(n) = 4.9 \times 10^{-6}$$

In the experiment, we used aqueous NaCl solutions subjected to hundredfold dilution from 0 to 30 times. Note that the preparation of the solutions included vigorous shaking at each stage of dilution.
RESULTS

In the experiment, the difference between the refractive index of the corresponding sample and the refractive index of the initial water was measured. The results of measurements for samples with pure water and aqueous NaCl solutions are presented for various degrees of their hundredfold dilution. The results were obtained by averaging over six measurements. Each measurement is a 100-point average.

The figures show the results of measuring the difference between the refractive index of a sample of the corresponding dilution and the refractive index of deionized water. As one can see from the figures, the refractive index of the solution changes nonlinearly and increase with the number of dilutions. In the case of pure water (Figures 6, 7), at the beginning, we see the sameness of the original water and water without dilutions. And it is natural to assume that water diluted and just water should have the same properties. However, we see the difference appears after 6 hundredfold dilution between water with shaking and without. That is, there is some feature at 6-fold dilution that leads to a sharp difference between the properties of the solution and those of solutions with a higher or lower concentration.

We observe a similar picture for an aqueous solution of NaCl (Figures 8, 9), but here the characteristic dip corresponds to a 12-fold dilution.

It should be remembered that these points were obtained by averaging 100 measurements and are highly reliable. Thus, our results show that the amount of dilution affects the physical properties of the solution.

Certainly, the obtained results cannot be interpreted as a measurement of NaCl concentration. With such dilutions,
there should not be any amount of a substance left in the solution. However, we see changes in the signal. This may indicate the effect of the solution preparation procedure on the properties of the medium. As shown in [7], mechanical action on an aqueous solution can change its molecular composition and, consequently, the refractive index. However, it is not clear why the maximum change occurs at any particular amount of dilution. The purpose of further research should be to study the properties of the environment itself by various methods and under various external influences. In this case, one of the most essential requirements is increasing the sensitivity. In this technique, an increase in the measurement accuracy can be achieved by optimizing the reflection coefficients of the interferometer mirrors and its length, by increasing the spectrum width of the radiation source, and also by improving the resolution of the spectrometer used.

CONCLUSION

A high-precision technique has been developed for comparing the refractive indices of solutions, one of which is considered to be a "reference". The results of the performed measurements show that the procedure for preparing aqueous solutions affects their physical properties. The experiments have shown that the proposed technique can be used to assess changes in the optical properties of aqueous solutions of ultra-low concentrations. To increase the reliability of the results, it is necessary to conduct further research with an increase in the accuracy of measurements and an increase in the amount of statistical data.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

IL and BU conducted experiments. IL and VP participated in the processing of the results and their discussion. IL and VP participated in writing the text of the manuscript.

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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