

Spectroscopic (Polarization, Excited State, FT-IR, UV/Vis and ¹H NMR) and Thermophysical Investigations of New Synthesized Azo Dye and Its Application in Polarizing Film

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Abstract: Quantum-chemical calculations using the Density Functional Theory (DFT) approach for structural analysis of a new azo benzene derivative: *Sodium 6-amino-5-hydroxy-3-((E)-(2-methoxy-4-((E)-(4-sulfonatophenyl)diazanyl)phenyl)diazanyl)naphthalene-2-sulfonate* (A) (trans isomer) was designed using B3LYP method with 6-311++G** basis set. The electronic absorption spectrum of the A in water solution was calculated using TDB3LYP/6-311++G** method. The excitation energies, electronic transitions and oscillator strengths for studied structure have also been calculated. On the basis of calculations a new dichroic azobenzene dye A was synthesized. The experimental UV-Vis, FT-IR and ¹H NMR spectra of the title molecule were presented. On the basis of polyvinyl alcohol (PVA) and the dichroic synthesized azobenzene dye A thermostable polarizing film absorbing in the seen region of the spectrum ($\lambda_{\max} = 550\text{nm}$) with Polarization Efficiency (PE) in the absorption maximum ($\lambda_{\max} = 92\%$) and Stretching Degree (R_s) = 3.0 was developed. The experimental spectral-polarization parameters of stretched colored PVA-films were measured. During the work it was established that oriented PVA-films is phenomenon of *anisotropy of thermal conductivity* ($\lambda_{\parallel}/\lambda_{\perp}$). Thermal conductivity in a direction of orientation (λ_{\parallel}) is higher than in a direction perpendicular orientations (λ_{\perp}).

Keywords: Polarizer Film, Dichroic Dye, Thermal Conductivity, Polarization Efficiency

1. Introduction

The polarizing thin film presents uniaxial oriented polymeric film containing dichroic agent. Light-polarizing films are the basis of sheet polarizers which are traditionally used in processes of manufacturing of liquid crystal indicators for displaying devices. They are constituted usually from polymer matrix, e.g., polyethylene or polyvinyl alcohol containing uniaxial oriented dichroic dyes or iodine

[1, 2]. The most used polarizers are the H-type polarizers which contain molecular iodine as dichroic agent. They are excellent in the initial polarization performance but are weak against water or heat. L-type polarizers contain a dichroic dye or mixture of the dyes as a polarizing agent [3, 4]. Examples of dichroic molecules include dye compounds, such as azo dyes or as well as iodine have a high polarization degree (90-99%) and can be maintained in a wide range of temperatures. Dichroic dyes when employed as polarizing

films are generally used in combination with other dyes, having absorption in a particular wavelength range to provide the polarizing films with a neutral color and wide range spectra [5, 6]. PVA-films are easily exposed to uniaxial orientation, then it becomes optically anisotropic and partially polarize passing light beams. These properties of PVA have defined theoretical and practical interest in it from the point of view of their use for the manufacturing of film polarizers. The majority of films with dichroic dyes finding practical application polarize light in a visible area (400-700 nm) electromagnetic spectrum where the basic strips of absorption of molecules with stretched chains of the interfaced double bond conjugation are located. Theoretical quantum chemistry methods based on HF and DFT are used for the calculation of optimized geometry, absorption spectrum, FT-IR and ^1H NMR spectra of the organic molecules [5-17]. In this work on the basis of PVA and new synthesized dichroic azobenzene dye A thermostable polarizing film absorbing invisible spectral range of spectrum ($\lambda_{\text{max}} = 550$ nm) for application in Display Technologies and Optoelectronic applications was developed.

2. Experimental

2.1. Reagent and Apparatus

All chemicals used were of analytical reagent grade. PVA used in this work «Mowiol28-99» was purchased from (Hoëchst Akiengesllschaft Co., Germany). The experimental UV/Vis absorption spectrum of the molecule was recorded on UV-Visible Spectrophotometer Cary 300 (Varian, USA). The optical transmission spectra were measured in polarized light with UV-NIR Spectrophotometer HR 4000 (Ocean optics, USA). FT-IR spectrum of films was measured by a spectrophotometer of Protégé 460 (Nicolet, US). The experimental FT-IR spectrum was recorded by KBr pellet method with spectral resolution 2cm^{-1} . ^1H NMR spectrum was recorded in DMSO- d_6 by NMR AVANCE-500 spectrometer (Bruker, Germany) with a working frequency of 600 Hz (1H). During calibration of the chemical shifts as the internal standard the residual signal of the solvent was used. Thermal conductivity of films was measured on the complex equipment LC-201 (Alfa Laval Group, Sweden) using indicator method for the determination of heat conductivity of polymer materials and films.

2.2. Synthesis of Sodium 6-amino-5-hydroxy-3-((E)-(2-methoxy-4-((E)-(4 sulfonatophenyl) diazenyl) phenyl) diazenyl) naphthalene-2-sulfonate (A)

Azo coupling diazotized sulfanilic acid with *o*-anisidine was carried out in formic acid (I). Thus was obtained 4-(4-amino-3-methoxyphenylazo)-benzolsulfonic acid as a chromatographically pure product (58 % yield), which was diazotized and treated with γ -acid in an alkaline medium by the classical method (II). The diazo compound was isolated by treating the reaction mixture with sodium chloride. The yield after recrystallization from 70 % ethanol was 79.4 %. The dye was dried in vacuum at 10 mm Hg at 150°C . The

final product is dark violet crystals (III) (Fig.1). This used method avoids the formation of triazenes and products of their transformation.

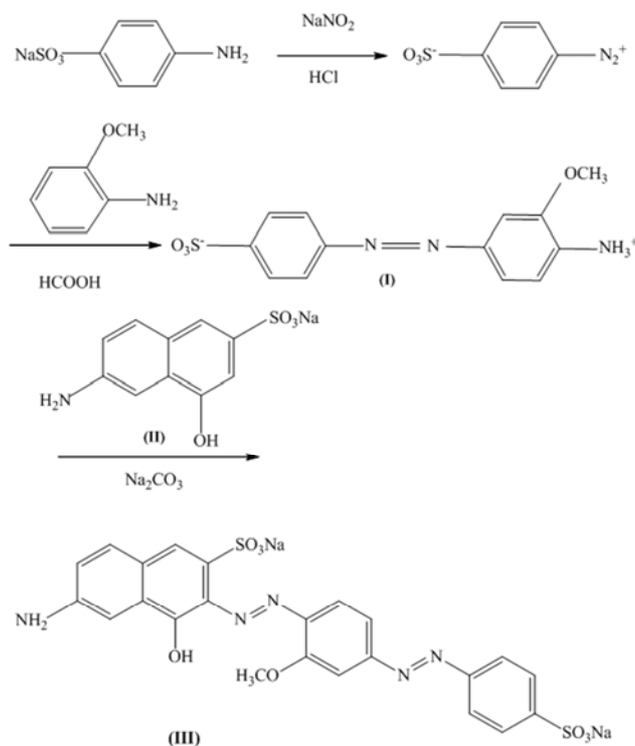
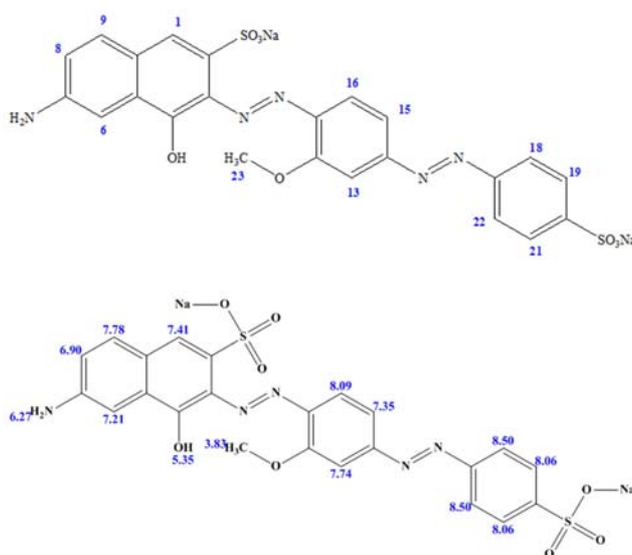


Figure 1. Synthesis of the dye A.

Experimental ^1H NMR Spectrum of the dye A δ : (DMSO- d_6 , ppm) at 6.27 MHz (S, 2H, NH_2), 3.83 MHz (S, 3H, CH_3), 7.74 MHz (d, 1H, $J = 6.4$ Hz, H-13), 7.78 MHz (d, 1H, $J = 6.4$ Hz, H-9), 7.21 MHz (d, 1H, $J = 1.2$ Hz, H-6), 6.90 MHz (S, 1H, H-8), 8.06 MHz (S, 2H, H-19,21), 7.35 MHz (d, 2H, $J = 6.4$ Hz, H-15), 8.09 MHz (d, 2H, $J = 6.4$ Hz, H-16), 8.50 MHz (d, 2H, $J = 6.8$ Hz, H-18,22), 7.41 MHz (S, 1H, H-1), 5.35 MHz (S, 1H, O-H) (Fig. 2).



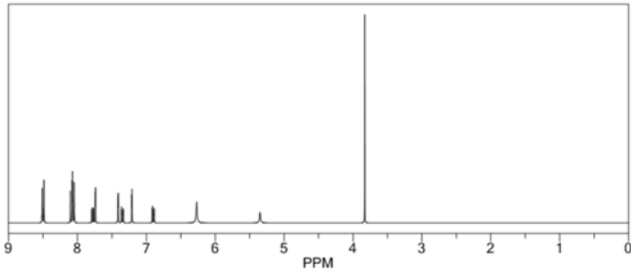


Figure 2. Experimental ^1H NMR Spectrum of the dye A.

Experimental FT-IR Spectrum of the dye A: 451.06 cm^{-1} bending vibrations of $=\text{C}-\text{H}$ aromatic, 530.08 cm^{-1} stretching

vibrations $-\text{NH}$, 570.58 cm^{-1} out of plane aromatic ring vibrations, 630.43 cm^{-1} stretching vibrations $\text{S}-\text{O}$, 705.89 cm^{-1} $-\text{C}-\text{H}$ bending vibrations $-\text{NH}$, 1007.40 cm^{-1} stretching vibrations $\text{C}=\text{C}$, 1035.38 cm^{-1} stretching vibrations $\text{C}-\text{N}$, 1052.12 cm^{-1} stretching vibrations $\text{S}=\text{O}$, 1122.17 cm^{-1} stretching vibration of $\text{C}-\text{O}$, 1155.04 cm^{-1} stretching vibration of $\text{C}-\text{N}$, 1402.08 cm^{-1} stretching vibration of CH_3 , 1496.50 cm^{-1} stretching vibration of $\text{N}=\text{N}$, 1631.07 cm^{-1} stretching vibration of $\text{C}=\text{C}$ aromatic, 1657.73 cm^{-1} stretching vibrations of $\text{C}=\text{C}$ aromatic (Fig. 3).

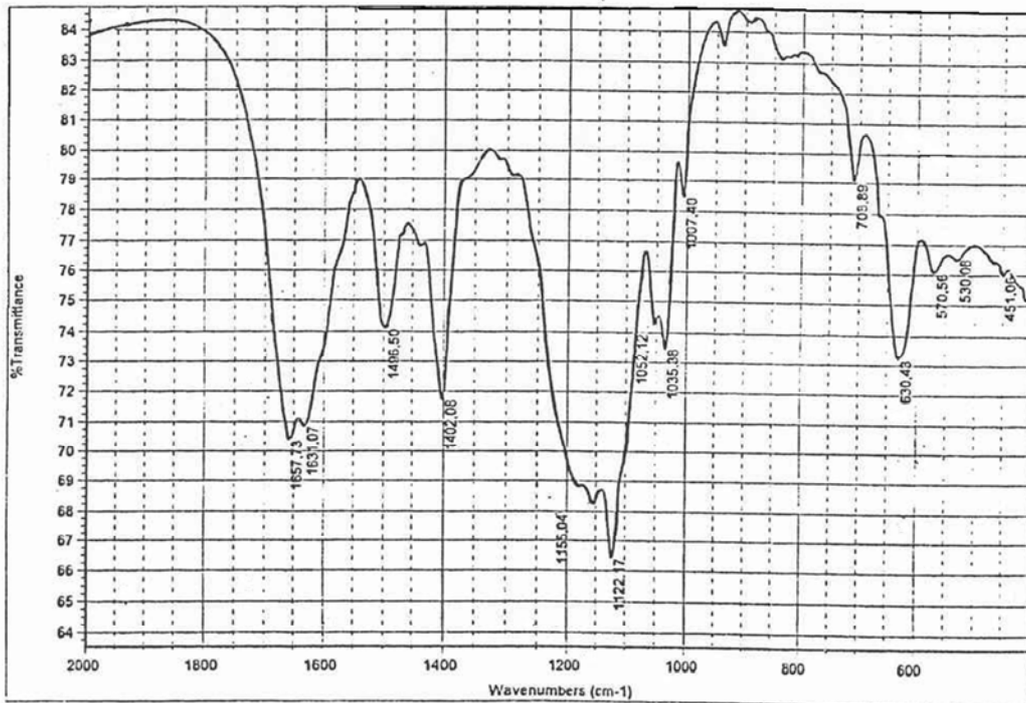


Figure 3. FT-IR spectrum of the dye A.

2.3. Preparation of PVA-Films Containing the Dye A

The PVA-films were prepared from 10 % PVA solution, containing the 0.01-0.05 wt. % the dye A, 0.01 wt. % boric acid (H_3BO_3), 6.0 wt. % ethyl alcohol ($\text{C}_2\text{H}_5\text{OH}$) and water. An initial composition was prepared by dissolving PVA in distilled water and ethyl alcohol. The composition was mixed at temperature $85-90\text{ }^\circ\text{C}$. The dye A and additives were added after 3 hours after starting of heating PVA solution at intervals 20 minutes. The mixture was heated for 3 hours. The hot solution was filtered through two layers of technical nylon. Deaeration was occurred during 12-15 hours. The composition was cast on the polished glasses and dried in the closed box at temperature $20-22\text{ }^\circ\text{C}$. Uniaxial orientation was done in the 4 % boric acid solution at temperature $42-45\text{ }^\circ\text{C}$. The washed film was dried for 30 minutes at temperature $60-63\text{ }^\circ\text{C}$. The value of R_s was determined as the ratio between length of the films after and before ($l_{\text{aft}} / l_{\text{bef}}$) uniaxial orientation. The thickness of the resulting films was between

50 to $55\text{ }\mu\text{m}$ [1-4]. The film thickness was measured with a micrometer with an accuracy of $\pm 5\text{ }\mu\text{m}$ (GS SSSR 6507-90). Accuracy of measurements in this research limits the accuracy of the results [1]:

- Sample of the starting composition for preparation of solutions and films for casting taken on an analytical balance accurate to 0.0005 g ;
- Temperature measurement error was:
 - $\pm 2.0\text{ }^\circ\text{C}$ in the preparation of polymer solutions with additives,
 - $\pm 1.0\text{ }^\circ\text{C}$ in the chemical treatment of the films,
 - $\pm 3.0\text{ }^\circ\text{C}$ at thermal fixation and drying films;
- Relative error of spectrophotometric analysis did not exceed 1.0% and thermal conductivity 0.1% .

The standard deviation of parallel measurements determined by the formula [3]:

$$S(d) = \sqrt{\sum_i (d_i - d)^2 / n - 1} \quad (1)$$

Where d_i - the result of a single measurement,
 d - Arithmetic mean
 n - Number of measurements.

3. Results and Discussion

3.1. Optical Properties: Polarization of PVA-Films Containing the Dye A

The main optical properties of polarizing films such as Transmittance (T_{\perp} , T_{\parallel}) and Polarizing Efficiency (PE) were evaluated at the absorption maximum of the polarizing films according to Eqs. (2) [1-3]:

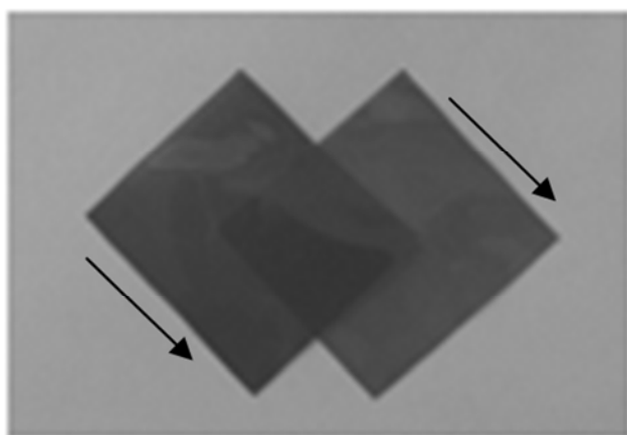
$$PE = (T_{\perp} - T_{\parallel}) / (T_{\perp} + T_{\parallel}) * 100 \quad (2)$$

Where, T_{\parallel} , T_{\perp} - Transmittance for linearly polarized light parallel (\parallel) and perpendicular (\perp) to direction of stretching of colored film. Polarizing Efficiency of colored oriented PVA-films depends on the concentration of injected dye and Stretching Degree of the film [2-4], therefore the optimum concentration of the dye A in PVA-film was obtained (Table 1). Changes in concentration of the dye A from 0.10 to 0.50 wt. % in the colored oriented PVA-film show that with increasing concentration of the dichroic agent, maximum light transmission in parallel and perpendicular direction are reduced. And at $[A] = 0.18$ wt. %, $PE = 91$ %. With increasing concentration of the dichroic agent (0.50 wt. %) PE decreases. It is due to the increasing intensity of the absorption of light by the dye.

Table 1. The optical characteristics of PVA-films at different concentration of the dye A, $R_s = 3.5$.

| [A] in film, wt. % | Transmittance at $\lambda = 550$ nm, % | | PE, % |
|--------------------|--|-----------------|-------|
| | T_{\perp} | T_{\parallel} | |
| 0 | 91.9 | 90.6 | 0 |
| 0.10 | 66.9 | 18.8 | 56 |
| 0.14 | 56.6 | 10.1 | 70 |
| 0.18 | 34.8 | 1.6 | 91 |
| 0.50 | 15.3 | 1.0 | 88 |

Dichroism (anisotropy of absorption) (Fig. 4) appears under uniaxial orientation of the colored PVA-films (Fig. 4(b)).



(a)

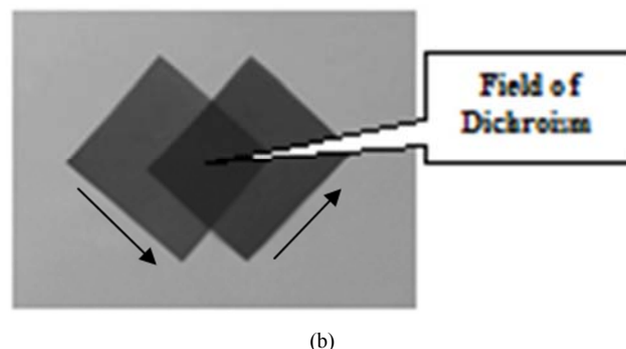


Figure 4. PVA-films containing the dye A in parallel (a) and perpendicular (b) directions of stretching.

We have studied effect of R_s on PE of the PVA-films (Table 2). Stretching (uniaxial orientation) of film containing 0.18 wt. % of the dye A from 2.5 to 4.5 times leads to increase in its T_{\perp} from 31.6 % to 42.0 % and decrease PE from 92 to 85 %.

Table 2. Influence of R_s on the optical characteristics of films containing $[A] = 0.18$ wt. %.

| R_s | Transmittance at $\lambda = 550$ nm, % | | PE, % |
|-------|--|-----------------|-------|
| | T_{\perp} | T_{\parallel} | |
| 2.5 | 31.6 | 1.3 | 92 |
| 3.0 | 32.8 | 1.4 | 92 |
| 3.5 | 34.8 | 1.6 | 91 |
| 4.0 | 38.1 | 2.5 | 88 |
| 4.5 | 42.0 | 3.3 | 85 |

Thus we have found that a PVA-films containing 0.18 wt. % of the dye A at $R_s = 2.5$ and 3 times have the best optical properties ($PE = 92$ %, $T_{\perp} = 31.6$ %, $T_{\parallel} = 1.3$ % and $PE = 92$ %, $T_{\perp} = 32.8$ %, $T_{\parallel} = 1.4$ %) (Tables 1,2).

3.2. Absorption Spectrum and Electronic Structure of the Dye A

Theoretical absorption spectrum of the molecule A optimized in a solvent (Water) was calculated using TDB3LYP/6-311++G** method at 300-700 nm by the IEFPCM solvent model with Water as the solvent. All the DFT computational calculations were carried out with the Gaussian 09W package [18]. For visualization of UV/Vis spectrum of the dye A Chemcraft software was used [19]. The equations were solved for 10 excited states, where the computational studies were performed in presence Water as solvent, using the IEFPCM (integral equation formalism PCM) method coupled to UAKS radii. The IEF method originally developed by Cancès and Mennucci introduces one function of permittivity dependent of the position, with this change [20]. The TDDFT method is able to detect accurate absorption wavelength at a relatively small computing time, which corresponds to electronic transitions computed on the ground state geometry (Table 3). The experimental absorption spectrum of the dye A (Fig. 5 (a)) has a wide absorption at about 300-620 nm which characterized by maxima at 440-460 and 550 nm and calculated spectrum (Fig. 5 (b)) shows oscillations in 456 and 555 nm that the strongest is at 555 nm ($f = 2.00$). The strong

peak at 550 nm is due to the Charge-Transfer (CT) excited state. The other peaks are local excited states corresponding to electrons going into anti-bonding orbitals associated with the benzene rings. Excitation of one electron at 550 nm belonged to the transition into the excited singlet state S_1 and describes by a wave function corresponding to a superposition of two configurations for one-electron excitations (134→138, and 134→147). Excitation of an electron from 134 to 138 molecular orbital (MO) gives the main contribution to the formation of the absorption band at 555 nm. Excitation of one electron at 456 nm belonged to the transition into the excited singlet state S_2 and describes by a wave function corresponding to a superposition of twelve configurations for one-electron excitations (112→136, 112→144, 118→135, 118→136, 118→144, 127→135, 127→136, 127→140, 127→144, 129→135, 129→136 and 129→144). Excitation of an electron from 127 to 144 molecular orbital (MO) gives the main contribution to the formation of the absorption band at 456 nm (Table 3). The next strong absorption at 453 nm belonged to the transition into the excited singlet state S_3 and describes by a wave function corresponding to a superposition of six configurations for one-electron excitations (128→135, 132→135, 133→135, 133→136, 133→140 and 134→135). Excitation of an electron from 132 to 135 and 133 to 135 MO gives the main contribution to the formation of the absorption band at 453 nm (Table 3, Fig. 5). The next absorption peak at 440 nm belonged to the transition into the excited singlet state S_4 and describes by a wave function corresponding to a superposition of five configurations for one-electron excitations (131→138, 131→147, 134→136, 134→139 and 134→140). Excitation of an electron from 134 to 139 MO gives the main contribution to the formation of the absorption band at 440 nm. Excited state S_5 at 416 nm describes by a wave function corresponding to a superposition of three configurations for one-electron excitations (134→138,

134→142 and 134→143). Excitation of an electron from 134 to 142 MO gives the main contribution to the formation of the absorption band at 416 nm (Table 3, Fig. 5). Excited states in UV region of spectrum (S_6 + S_{10}) are presented in Table 3 but not discussed.

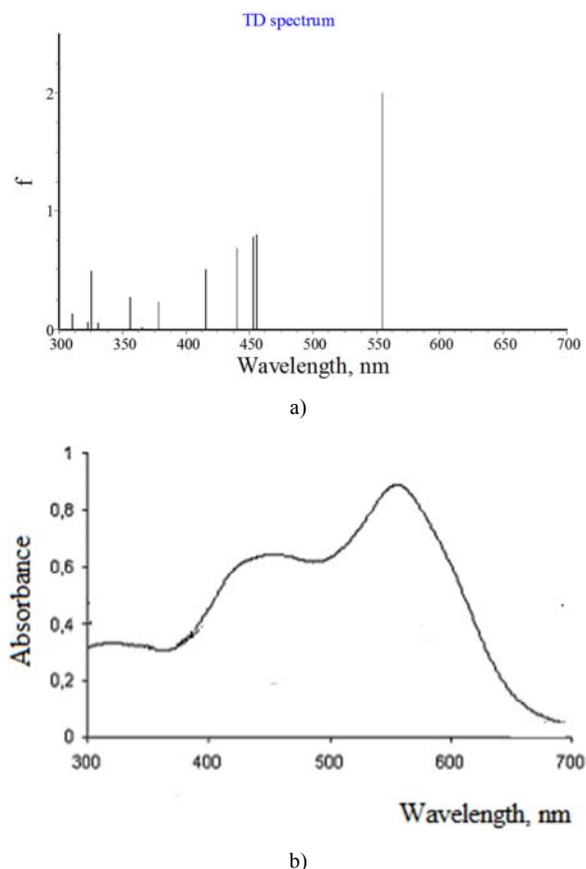


Figure 5. UV-Vis Spectrum of the dye A in Water: (a) experimental at dye concentration $3.6 \cdot 10^{-4}$ M/L and (b) calculated by TDB3LYP/6-311++G** method.

Table 3. Electronic absorption spectrum of the compound A calculated by TDB3LYP/6-311++G** method.

| Excited state | Wavelength (nm) | Excitation energy (eV) | Configurations composition (corresponding transition orbitals) | Oscillator strength (f) |
|---------------|-----------------|------------------------|--|-------------------------|
| S_1 | 555 | 2.23 | 0.59(134→138)-0.28(134→147) 0.11(112→136)+0.11(112→144)- | 2.00 |
| S_2 | 456 | 2.72 | 0.19(118→135)+0.22(118→136)+0.24(118→144)+0.19(127→135)- 0.19(127→136)+0.11(127→140)-0.27(127→144)+0.15(129→135)-0.14(129→136)- 0.22(129→144) | 0.80 |
| S_3 | 453 | 2.74 | 0.16(128→135)+0.35(132→135)+0.35(133→135)+0.17(133→136)-0.12(133→140)- 0.15(134→135) | 0.78 |
| S_4 | 440 | 2.82 | 0.17(131→138)-0.16(131→147)+0.13(134→136)+0.45(134→139)+0.44(134→140) | 0.68 |
| S_5 | 416 | 2.98 | 0.14(134→138)+0.58(134→142)-0.30(134→143) | 0.50 |
| S_6 | 379 | 3.27 | -0.17(128→135)-0.25(130→135)-0.20(132→135)+0.12(132→136)+0.13(132→139)- 0.13(132→140)+0.21(133→135)+0.18(133→136)-0.16(133→140)-0.21(134→145) 0.18(127→137)-0.11(127→141)-0.27(127→146)+0.25(128→135)-0.18(128→136)- 0.10(128→144)- | 0.24 |
| S_7 | 356 | 3.48 | 0.18(130→135)+0.26(130→136)+0.19(130→144)+0.14(132→136)+0.14(132→144) | 0.27 |
| S_8 | 331 | 3.75 | -0.11(128→135)-0.11(133→135)-0.42(134→135)+0.24(134→136)+0.20(134→139)- 0.27(134→140)-0.15(134→144)+0.13(134→150) | 0.06 |
| S_9 | 326 | 3.80 | -0.14(123→135)+0.11(127→137)-0.11(128→136)-0.12(129→146)+0.32(131→138)- 0.14(131→142)-0.26(131→147)-0.13(133→135)-0.12(134→139)-0.11(134→140)- 0.27(134→145) | 0.49 |
| S_{10} | 311 | 3.99 | -0.12(119→143)-0.10(122→135)+0.10(125→135)+0.10(125→136)- 0.11(130→135)+0.11(132→135)+0.11(132→139)+0.46(134→145) | 0.14 |

3.3. Thermo-Physical Properties

To clarify the nature of the effect of dyes on thermal stability of polyvinyl alcohol matrix, thermal conductivity of colored films has been studied. Experimental data on the thermal properties of an anisotropic medium allow estimating the role of structural parameters and the strength of bond between molecules and their aggregates in the process of heat transfer. In polymeric substances covalent bond generally predominates, and the motion of atoms in a polymer substance appears as collective oscillations of the atoms of the entire system [2]. Heating polymer increases the amplitude of the oscillations of the atoms. Heat is transferred through the interatomic forces from the hotter region to less heated one until all parts of the body acquire one common temperature. Besides the thermal motion of molecule segmental areas which appears at the glass transition temperature of the polymer, movement of individual fragments or monomeric units is possible [2, 3]. One should also distinguish heat transfers inside a macromolecule and between molecules. It is established [3] that the thermal fluctuations in the macromolecule are transmitted much faster on the main valences than on (auxiliary ones) the side. Intermolecular heat transfer in certain PVA is characterized by the existence of a potential barrier (or thermal resistance) in the heat pulse transfer caused by Vander Waals forces of interaction. As a result, the thermal conductivity (λ) in this direction is characterized by smaller values. It is known [2] that fillers and modifiers of plastics material modify the thermal conductivity of polymers. The degree of influence depends on the size and shape of the particles, their distribution in the polymer matrix and interaction of filler or a modifier with a polymer [3]. Consequently, coloring materials as modifiers of the film properties influence their heat conductivity. There are various types of polymer polarizers, but the ones that are mostly used in the practice are obtained on the basis of PVA, colored with iodine or organic dyes such as azo benzene derivatives, which are more resistant to high temperatures and humidity of the environment. The search and synthesis of new effective dichroic dyes remains an urgent task of research in the field of thermostable polarizing films [2, 3, 5].

3.4. Anisotropy of Thermal Conductivity of PVA-Films

We have studied thermal conductivity of the systems: PVA-A for development of thermostable polarizing films. In the researches were used special paints able to sharply change the initial color at a critical temperature (T_{cr}). The isothermal surface moves at a certain speed in the proportion to a local gradient of the temperature field. Thermo-physical characteristics of material can be judged by the speed of isotherm movement and the form of flowed surface. Thermal indicators of Ciba Com. (Switzerland) with small sizes of T_{cr} are used in order to avoid thermal destruction of polymer materials. Thermal indicator is put on a film by a thin layer and at regular intervals by a draw plate. Then, after some

drying period its unpainted surface is resulted on some time (30 sec.) in dense contact to a dot source of heat (it is heated up approximately to 55 °C metal needles). For estimation of thermo-physical properties of films, heat conductivity of samples was determined in parallel ($\lambda_{||}$) and perpendicular (λ_{\perp}) directions of stretching axis. During this work it was established that the oriented PVA-films with the dye A is the phenomenon of *anisotropy of thermal conductivity* ($\lambda_{||} / \lambda_{\perp}$). Anisotropy of thermal conductivity in unstretched PVA-films is observed not appreciably ($\lambda_{||} = 0.875$ W/m.°C; $\lambda_{\perp} = 0.869$ W/m.°C) as comparison to stretched PVA-films that is observed very clearly (Table 4).

Table 4. Dependence of thermal conductivity on stretching degree in pure PVA-films.

| R_s | λ | | |
|-------|-------------------------|----------------------------|----------------------------------|
| | $\lambda_{ }$, W/m.°C | λ_{\perp} , W/m.°C | $\lambda_{ } / \lambda_{\perp}$ |
| 1.5 | 0.876 | 0.764 | 1.15 |
| 2.0 | 0.878 | 0.636 | 1.38 |
| 2.5 | 0.880 | 0.549 | 1.60 |
| 3.0 | 0.881 | 0.503 | 1.75 |
| 4.0 | 0.882 | 0.475 | 1.86 |

Results of thermal conductivity measurements of PVA-films containing the dye A depending on stretching degree are given in Table 5. It is clear from the results that the thermal conductivity in a direction ($\lambda_{||}$) orientation is higher than perpendicular orientations (λ_{\perp}). On resulting anisotropy at a known degree of extension it is possible to judge anisotropy of chain structure. It is necessary to notice that during thermal expansion and thermal conductivity, geometric parameters of molecule and intermolecular forces play a significant role.

Table 5. Thermal conductivity of PVA-films containing the dye A at concentration 0.18 wt. %.

| R_s | λ | | |
|-------|-------------------------|----------------------------|----------------------------------|
| | $\lambda_{ }$, W/m.°C | λ_{\perp} , W/m.°C | $\lambda_{ } / \lambda_{\perp}$ |
| 2.0 | 0.865 | 0.261 | 3.31 |
| 3.0 | 0.851 | 0.247 | 3.44 |
| 4.0 | 0.844 | 0.186 | 4.54 |
| 5.0 | 0.840 | 0.110 | 7.36 |

The occurrence of anisotropy of thermal conductivity is connected that at orientation of PVA occurs orientation of the amorphous part of polymer and also formation of additional number of intermolecular connections. Thermal conductivity of PVA films changes after the injection of dye (Table 5) and along an axis of orientation and in perpendicular axis decreases.

4. Conclusions

In the present work, we have calculated the electronic structure and UV/Vis spectrum of the molecule: *Sodium 6-amino-5-hydroxy-3-((E)-(2-methoxy-4-((E)-(4-sulfonatophenyl)diazanyl)phenyl)diazanyl)naphthalene-2-sulfonate* (A) by using B3LYP/6-311++G** method. Based

on quantum-chemical calculations have been synthesized new azo benzene dichroic dye for seen region of spectrum (at 440-460 and 550 nm). At the first time on the basis of polyvinyl alcohol (PVA) and synthesized compound: *Sodium 6-amino-5-hydroxy-3-((E)-(2-methoxy-4-((E)-(4-sulfonatophenyl)diazenyl)phenyl)diazenyl)naphthalene-2-sulfonate (A)* thermostable polarizing film absorbing at $\lambda_{\max} = 550$ nm was developed. Stretched colored PVA-films have phenomenon of *anisotropy of thermal conductivity*. Thermal conductivity in a direction of orientation (λ_{\parallel}) is higher than in a direction perpendicular orientations (λ_{\perp}). In stretched dyed PVA-films anisotropy of thermal conductivity is observed very clearly (at $R_s = 5.0$, $\lambda_{\parallel} / \lambda_{\perp} = 7.36$).

References

- [1] Agabekov V., Muravsky A., Ariko N., Shahab S., Tolstic A. J.: New dyes for «gust-host» mode, *Semicond. Phys. Quantum. Electro. Optoelectro.* 13 (2010) 32-35.
- [2] Shahab S., Agabekov V., Ariko N., Filippovich L.: Anisotropy of polarizing polyvinyl alcohol films, *Advanced Display Technologies: proc. Of the 15 Intern. Symp., Moscow, Russia, Oct. 3-5, 2006/Soc. for Inform. Display.* 163-166.
- [3] Agabekov V., Shahab S., Ariko N., Filippovich L.: Anisotropy of polarizing films with azo dyes, *XV Intern. Symp. On Advanced Dis. Tech. SPIE. San Diego, USA, August 27-31, 2007. V.19/4. -P. 174-179.*
- [4] Agabekov V., Ivanova N., Shahab S., Kulevskaya I., Ariko N.: Film transfect polarizer, *Advanced Display Technologies: Basic Studies of Problems in Inform. Display (FLOWERS 2000), Moscow, Russia, Oct. 10-12, 2000./ed. V.V. Belyaev, I.N. Kompanets; Soc. for Inform. Display.-Bellingham, 2000. -P. 141-143.*
- [5] Almodarresiyeh H. A., Shahab S. N., Zelenkovsky V. M., Agabekov V. E.: Electronic structure and absorption spectra of sodium 2-hydroxy-5-((2-methoxy-4[4-sulfophenyl]diazenyl]phenyl]diazenyl)benzoate, *J. App. Spect.* 81 (2014) 181-183.
- [6] Almodarresiyeh H. A., Shahab S. N., Zelenkovsky V. M., Ariko N. G., Filippovich L. N., Agabekov V. E.: Calculation of UV, IR and NMR spectra of diethyl 2,2'-[(1,1'-biphenyl)-4,4'-diylbis(azanediyl)]diacetate, *J. App. Spect.* 81 (2014) 42-47.
- [7] Yaroschuk O., Sergan T., Lindau J., Lee S., Kelly J., Chien L.C.: Light induced structures in liquid crystalline side-chain polymers with azobenzene functional groups. *J. Chem. Phys.* 114 (2001) 5330-5337.
- [8] Kim J. J., Funabiki K., Muramatsu H., Shibata K., Kim S. H., Shiozaki H., Hartmann H., Matsui M.: Synthesis, structure, and UV-Vis absorption spectra of azo dyes derived from (dialkylamino) thiazole dimmers, *J. Chem. Soci. Perkin Transactions 2* (2001) 379-387.
- [9] Oprea C., Moscalu F., Dumbrava A., Ioannou S., Nicolaidis A., Girtu M.: DFT study of the optical and vibration spectra of a series of platinum-olefin complexes. *Roman. J. phys.* 56 (2011) 125-133.
- [10] Bezerra E. M., Bezerra-Neto J. R., Sales F. A., dosSantos R. P., Martins A., deLima-Neto P., Caetano E. W. S., Albuquerque E. L., Freire V. N.: Optical Absorption of the Antitrypanocidal Drug Benznidazole in Water, *Molecules* 19 (2014) 4145-4156.
- [11] Masoud M. S., Ali A. E., Shaker M. A., Ghani M. A.: Solvatochromic behavior of the electronic absorption spectra of some azo derivatives of amino pyridines, *Spectrochimica Acta Part A* 60 (2004) 3155-3159.
- [12] Shahab S., Filippovich L., Kumar R., Darroudi M., Yousefzadeh Borzehandani M., Gomar M., Haji Hajikolae F.: Photochromic properties of the molecule Azure A chloride in polyvinyl alcohol matrix, *J. Mol. Struct.* 1101 (2015) 109-115.
- [13] Demukhamedova S., Alieva I., Gadzhiev Z., Godzhaev N.: AB initio calculation structure and spectral analysis of the vibrational spectra of the carnosine molecule, *J. Qafqaz Univ. Chem.* 27 (2009) 15-25.
- [14] Fleming S., Mills A., Tutte T.: Predicting the UV-Vis spectra of oxazine dyes, *J. Org. Chem.* 7 (2011) 432-441.
- [15] Anbarasan P. M., Kumar P. S., Vasudevan K., Govindan R., Prakasam A., Geetha M.: Geometrical, electric structure, nonlinear spectroscopic investigations of 4-(phenylthio) phthalonitrile dye sensitizer for solar cell using quantum chemical calculations, *Eur. J. Chem.* 2 (2011) 206-213.
- [16] Shahab S., Alhosseini Almodarresiyeh H., Kumar R., Darroudi M.: A study of molecular structure, UV, IR and ¹H NMR spectra of a new dichroic dye on the basis of quinoline derivative, *J. Mol. Struct.* 1088 (2015) 105-110.
- [17] Shahab S., Kumar R., Darroudi M., Yousefzadeh Borzehandani M.: Molecular structure and spectroscopic investigation of sodium(E)-2-hydroxy-5-((4-sulfonatophenyl)diazenyl)benzoate: A DFT study, *J. Mol. Struct.* 1083 (2015) 198-203.
- [18] Frisch M. J., Gaussian 09, Revision A.1 Gaussian Inc., Wallingford CT (2009).
- [19] <http://www.chemcraftprog.com/index.html>
- [20] Mennucci B., Cancés E., Tomasi, J.: Evaluation of Solvent Effects in Isotropic and Anisotropic Dielectrics and in Ionic Solutions with a Unified Integral Equation Method: Theoretical Bases, Computational Implementation, and Numerical Applications, *J. Phys. Chem. B* 101 (1997) 10506-10517.