Commercial LC ZLI 1840 (Merck, Germany) was used as a host. It exhibits a stable nematic phase over a broad temperature range (−15° to 90°C) and high positive dielectric anisotropy [11]. The dyes were initially screened for solubility in LC and then dissolved in the medium at a concentration of 0.5 w/v %, which is suitable for the intended application [12]. The measurements on the dyes in LC were carried out in "sandwich" cells of 20 μm thickness. Planar homogeneous molecular orientation of the dye/LC systems in the cells was achieved by coating the cell surfaces from inside with polyimide layers orientated by rubbing [13, 14].

Coloured thin films of polystyrene (0.5% dye) were prepared from a 10% solution of polystyrene in benzene with the dye dissolved there.

**Results and Discussion**

*Influence of the Substituents at C-3 Position on the Dyes Absorption*

The electron donor-acceptor interaction in the molecule of 3-substituted benzanthrone occurs between the electron acceptor carbonyl group and the electron donating groups at C-3 position of the chromophoric system [9]

\[
\text{Scheme 1}
\]

The molecules are polarised in dependence on the electron donating power of the substituent R. Therefore their absorption properties are different.

The 3-substituted benzanthrone derivatives exhibit the following order of the energetic levels [15, 16]

\[
T^* < S^* < Tn^* < Sn^*.
\]

In the absorption spectra of these compounds, the longwave band in the visible region has charge transfer, CT, character, due to \( \pi \rightarrow \pi^* \) electron transfer during the \( S_0 \rightarrow S_1 \) transition. The absorption and emission bands largely depend both on the electron donating power of the substituents at C-3 position and the nature of solvents. The absorption properties of the dyes determined in a polar proton donating solvent (ethanol) and in a nonpolar solvent (benzene) are presented in Table 1. The exchange of the polar solvent with the nonpolar one brings in most cases a substantial hypsochromic shift, which is well known for CT bands [17].

Let us consider the influence of both groups of substituents, azomethine and oxy groups, upon the absorption properties of the dyes. In the case of dye 1, the CT band exhibits a bathochromic effect because of the strong donor-acceptor interaction. The substitution of the primary amino group with the azomethine group (dyes 2–5) leads to decrease of the donor-acceptor interaction and, correspondingly, to hypsochromic shift of the absorption maximum, with the exception of dye 4. The different nature and position of the substituents in the benzene ring of the azomethine fragment in dyes 2–5 cause also themselves a substantial effect upon the electronic spectra. Dye 2, with no substituents in the benzene ring, exhibits a yellow-green colour in ethanol and benzene solutions. In the case of dye 3, the electron donating OH- group at o-position with regard to the azomethine group changes only weakly the position of the CT band compared to that of dye 2. Most probably this can be related to the formation of an intramolecular hydrogen bond according to Scheme 2:

\[
\text{Scheme 2}
\]

The electron donating OH and \(-N(CH_3)_2\) groups at p-position in the benzene ring (dyes 4 and 5) cause stronger
polarisation of the benzanthrone molecule due to enhanced donor-acceptor interaction. Consequently, their absorption maxima are bathochromically shifted with regard to those of dye 2.

The substitution of the primary amino group in dye 1 with alkoxy and phenoxy groups (dye 6–8) leads to even larger hypsochromic shifts of the absorption maxima than for dyes 3–5. This is related to the weaker electron donating nature of the OR groups in comparison with the primary amino group. In the case of the oxy derivative dyes 6–8, there is no substantial influence of the substituents connected to the O atom at 3r position on the positions of the CT bands.

Influence of the Environment on Dyes Absorption

The spectral properties of benzanthrone dyes depend also on the nature of the environment (polarity, viscosity, formation of hydrogen bonds or other intermolecular interactions, etc.) [18–20]. The dependence of the absorption maximum of the CT bands in the case of dyes 1, 2 and 6, as example, upon the empirical parameter of solvent polarity $E_{r(30)}$ [21] is presented in Figure 1. It is seen from Fig. 1 that the dyes 1, 2 and 6 exhibit positive solvatochromism. The linear relations observed indicate that dipol-dipol interactions prevail in the dye solution. In acetone, a specific interaction between the free electron pair of the O atom of the solvent and H atoms of the primary amino group in dye 1 is possible. In this case, the polarisation of the molecule is enhanced and the absorption maxima are bathochromically shifted with respect to those in toluene, benzene and chloroform, where such intermolecular H-bond complex formation cannot occur. The absorption maxima of dye 1 in the proton donating solvents ethanol and methanol are bathochromically shifted to 508 and 512 nm, respectively. In that case, the polarisation of the chromophoric system is enhanced by the formation of intermolecular H-bonds between the free electron pair of the carbonyl group in benzanthrone and the hydroxy H-atoms of the solvents, which causes the bathochromic shift. Dyes 2 and 6 have similar absorption behaviour in organic solvents. For both dyes the absorption maxima in alcoholic solvents are bathochromically shifted in comparison with those in non polar aprotic solvents. From Fig. 1 it is seen too that the absorption maxima of dyes 2 and 6 are less influenced by the nature of the solvents.

All benzanthrone dyes under study have similar absorption properties in proton donating solvents and, correspondingly, their long-wave CT bands are strongly bathochromically shifted with regard to those in benzene and toluene. In acetone, intermolecular hydrogen bonds are formed only in case of the dyes 1 and 4, by the labile H atoms of amino and hydroxy groups.

The absorption properties of dyes 1–8 in solid polystyrene (PS) film (Table 1) are similar to those in non-polar solvents as a result of comparable polarity constants: $\varepsilon$ (benzene) = 2.28, $\varepsilon$ (PS) = 2.40 – 2.65 [22]. An exception is dye 4 where the absorption maximum in PS is largely hypsochromically shifted ($\Delta \lambda = 34$ nm) with regard to that in benzene, probably because of aggregate formation and lack of conformational changes of the dye in the solid matrix.
Figure 2 presents the maxima of the CT band of the dyes in LC. As shown, dyes 1–8 cover the colour interval from yellow-green to red depending on their structural specificity. Dye 1 has an intense red colour while dyes 2, 3, 6–8 have yellow-green colour, enhanced in the case of dyes 6–8 by their strong green fluorescence; dyes 4 and 5 exhibit yellow colour. The absorption and fluorescence maxima of the dyes in LC are bathochromically shifted with regard to $\lambda_A$ and $\lambda_F$ in the nonpolar solvents in polystyrene film, and hypsochromically shifted in the polar solvents, indicating that the polarity of LC is in-between.

Fluorescence and other Optical Characteristics of the Dyes in Isotropic and Anisotropic Media

It is known that the substituents at C-3 position of the benzanthrone molecule have strong influence on all the basic photophysical characteristics [18, 19, 23]. This is established also here not only for the absorption maxima but also for the other characteristics of the dyes under study (Table 1).

The fluorescence maxima of these dyes are in the region 525–653 nm, depending on the structure of the chromophore. In the series of N-substituted derivatives (dyes 1–5), the nature of the substituent has a strong influence on the fluorescence maxima $\lambda_F$. The substitution of the primary amino group in dye 1 by the azomethine group (dye 2) leads to a large blue shift of the fluorescence maximum, of about 40 nm. Substantial influence on the fluorescence maxima show also the substituents in the azomethine group. The electron donating dimethyamino group in the benzene ring in p-position (dye 5) causes a large red shift of about 70 nm. The hydroxy group in o-position (dye 3) causes also a red shift of the fluorescence maximum due to the intramolecular hydrogen bond formation (Scheme 2), suggesting that the hydrogen bond is more stable in the excited $S_1$ state than in the ground state $S_0$. The largest red shift of the fluorescence maximum $\lambda_F$, of 128 nm (dye 4 with hydroxy group in p-position) is probably connected with the prototropic state of the dye molecule presented in Scheme 3 [9]:

![Scheme 3](image)

Table 2. Optical characteristics of substituted 3-benzanthrone dyes in liquid crystal ($ld$-molecular aspect ratio, $D(l)$: dichroism measured by absorption A or fluorescence F).

<table>
<thead>
<tr>
<th>Dyes</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\lambda_F$ nm</td>
<td>496</td>
<td>434</td>
<td>432</td>
<td>464</td>
<td>462</td>
<td>430</td>
<td>418</td>
<td>420</td>
</tr>
<tr>
<td>$D(\lambda_F)$</td>
<td>2.76</td>
<td>4.54</td>
<td>5.03</td>
<td>4.65</td>
<td>4.62</td>
<td>3.64</td>
<td>3.04</td>
<td>3.76</td>
</tr>
<tr>
<td>$\lambda_A$ nm</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>502</td>
<td>504</td>
</tr>
<tr>
<td>$D(\lambda_A)$</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.84</td>
<td>2.06</td>
<td>2.82</td>
<td></td>
</tr>
</tbody>
</table>

In the case of alkoxy and phenoxy substituted benzanthrone dyes, the substituents have almost no influence on the fluorescence maxima.

The energy of the first excited state $E_{S1}$ of dyes 1–8 in ethanol (Table 1) is determined from the cross point of the absorption and fluorescence spectra. Dyes 4 and 5 show the lowest values, i.e. they need lower energy supply for the $S_0 \rightarrow S_1$ transition.

The substituents at C-3 have definitely a strong influence on the quantum yields of fluorescence $\Phi_F$ (Table 1). In the case of the N-substituted derivatives, the quantum yield of fluorescence is very low, indicating a substantial nonradiating deactivation during the transition from the excited to the ground state. Much higher values of the quantum yield have the oxy substituted benzanthrone dyes, where $\Phi_F = 0.42 - 0.55$, their molecules being subjected to less conformational changes.

An important characteristic of the dyes is the oscillator strength $f$ determined by [24]

$$ f = 4.32 \times 10^{-9} \Delta \nu_{1/2} \epsilon_{\text{max}}, \quad (4) $$

where $\Delta \nu_{1/2}$ is the width of the absorption band (in cm$^{-1}$) at 1/2 $\epsilon_{\text{max}}$. The $f$ values are highest for azomethine derivatives (dyes 2–5), showing the tendency to increase with the higher molar absorption (Table 1).

For the potential use of the dyes in the “guest-host” LC displays it is of interest to evaluate the fluorescence properties of dyes 1–8 in LC matrix. In the case of dyes 1–5, however, one could observe on irradiation with UV-light only a faint fluorescence emission of orange-red colour. In the case of dyes 6–8, bearing oxy groups, the emission is very intensive in the range 450–650 nm, where the maximum of fluorescence is observed at 502–504 nm (Table 2).

It is known that some dyes, when oriented selectively to absorb polarised light, exhibit dichroism [25]. Figure 3 presents as an example the polarised absorption spectra of dye 3 in LC. From such spectra, the dichroic
ratio of the dyes in LC is determined according to the equation [26]

\[ D(\lambda_A) = A_{||}/A_{\perp}, \]

where \( \lambda_A \) is the absorption maximum, and \( A_{||} \) and \( A_{\perp} \) are the corresponding absorbances in polarised light at parallel and perpendicular orientation of the polarizer with respect to the LC director, respectively. The data obtained (Table 2) show that the dichroic ratios of the benzanthrone dyes under study are not proportional to the length to width ratio, \( l/d \), of the molecules and obviously depend not only on the linearity of the molecules but also on other factors. The calculation of \( l/d \) values was made by using the bond lengths in the molecules including the end groups [27].

It is evident that both substituent R and the flexibility of the C-N-C link at C-3 of the benzanthrone molecule influence the dichroic properties of dyes 1–8. Azomethine derivatives (dyes 2–5) show higher values of \( D(\lambda_A) \) than the oxy derivatives 6–8, suggesting better orientational order of their solutions in LC [28]. The azomethine fragment there is involved in a system of conjugated double bonds of the chromophoric system, in this way increasing its linearity, while the oxy substituents at C-3 of dyes 6–8 exhibit only their electron donating power. In Table 2, the respective values of the dichroic ratio determined from the fluorescence maxima, \( D(\lambda_F) \), are also presented. The values are lower than those determined from the absorption maxima, \( D(\lambda_A) \), which is probably due to the existence of non zero intramolecular angles between the absorption and emission oscillators [29, 30].

**Vibronic Transitions in Benzanthrone Dyes**

The width of an absorption band in the electronic spectra of conjugated substances is mainly a result of both vibronic and intermolecular interactions.

The observed vibronic modes of all benzanthrone dyes studied in different media, ethanol, benzene, PS film and in LC cell, are determined from the dyes second derivative spectra [31] and are presented in Table 3. Vibronic transitions within the \( S_0 \rightarrow S_1 \) transition are less excited in the polar proton donating ethanol. When comparing the vibronic structure of azomethine and oxy derivative dyes, it is seen that the oxy substituted benzanthrone dyes (6–8) are characterised by a more pronounced vibronic structure of the spectra, probably connected with the greater rigidity of their molecules. This is also valid for the azomethine dye 2.

The vibronic structure is more pronounced in benzene due to the lower solvent polarity and to the lack of hydrogen bond formation between dye and solvent. It is noteworthy that the vibronic structure of the dyes in benzene and PS film is similar, indicating that it depends substantially on the polarity of the medium but not on its viscosity.
For the dyes oriented in LC, the vibronic structure of dyes 1–5 (especially of dye 1 and 4) is richer than that of the oxy derivatives 6–8. Obviously, the oriented state of LC prevents the possible conformational changes connected with a rotation of the azomethine fragment. Within the series of azomethine derivative dyes 2–5, dyes 2 and 3 have the less pronounced vibronic structure, probably due to molecular rigidity which in the case of dye 3 is due to the intramolecular H bond formation (Scheme 2).

**Concluding Remarks**

The spectral properties of a number of recently synthesised 3-benzanthrone derivatives bearing azomethine and oxy groups have been determined in isotropic and anisotropic (LC) media. The substituents at 3-C position of the benzanthrone molecule have strong influence on the spectral properties of the dyes under study, which confirms the prevailing role of the electron donor-acceptor interaction. The influence is stronger in case of azomethine substitutes (dyes 2–5), where the polarization is large, and smaller for the oxy derivatives with their lower polarization (dyes 6–8).

There is a well pronounced solvatochromic effect established for dyes having groups capable to solute-solvent interaction (dyes 1, 4 and 5). When such groups are not present (dyes 2, 6–8) or are screened (dye 3) or undergo transformation (dye 4), the effect is small.

The vibronic interactions of the dyes depend both on the nature of the substituents and the polarity of environment in a complex way. The dye molecules are more stable in the excited state $S_2$ in the nonpolar media benzene and polystyrene film and in the oriented state in the liquid crystal matrix.

The functional properties of the dyes for use in electro-optic displays can be resumed as follows. The dyes cover a wide colour interval, from yellow-green to red-orange. The oxy substituted benzanthrone dyes are expected to enhance the image contrast of the displays because of their good fluorescence properties. The azomethine substituted dyes are also of interest because of their dichroic properties which suggest a better orientation order of the dye/LC system in the display and a well developed vibronic structure supposing enhanced contrast.