Electrooptical properties of liquid crystal n-pentil-n’-cyanobifenil with J-aggregates of astrofloxine

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Abstract. It is shown that organic nanocryallites – J-aggregates – can be formed and oriented in a liquid crystalline matrix. External electric field can reorient and destroy these nanocryallites. Transmission and photoluminescence spectra of astrofloxine solutions in the nematic liquid crystal are investigated in d.c. and a.c. electric fields with the frequencies of 1 Hz–1 kHz and amplitudes $F = 1.2\times10^{4}$ V·cm$^{-1}$. Intense photoluminescence was observed at two-photon excitation. Efficiency of IR laser radiation transformation into visible photoluminescence was as high as 10%.

Keywords: liquid crystal, nanocryallite, photoluminescence, two-photon excitation.

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1. Introduction

Nematic liquid crystals (LC) are often used as matrices for parallel orientation of elongated dye molecules [1]. In an external electric field these LC exhibit pronounced electrooptical properties, i.e. the changes of absorption spectra (electrochromic effect) and photoluminescence (PL) spectra. Electrochromic effect is caused by two different physical processes [2]:

i) external electric field changes the orientation of molecules and, hence, the absorption spectrum intensity,

ii) the field changes the distances between energy levels of the molecules, shifts the absorption bands and varies the transition moments.

Changes in the absorption spectra observed in optical experiments are not related to the shift due to Stark effect and separate lines splitting. They are caused by the extinction coefficient variation [2,3]. The spectral features of a dye are essentially influenced by the medium where it is dissolved or dispersed. The medium also determines the phase state of the dye and evolution of its spectrum in the electric field. It was shown recently [4] that organic nanocryallites – J-aggregates – can be formed and oriented in a LC matrix.

At present the above-mentioned organic and inorganic semiconductor particles of sub-micron size are widely investigated. E.g., in [5] it was shown that J-aggregates of pseudoisocyanines have very large nonlinear cubic susceptibility [5]. Elongated (rod-like) shape and large electric dipole along the aggregate axis are the key properties for the creation of «macroanisotropic» media on the base of aligned J-aggregates. One of the possible methods to create «microanisotropy» is to align the aggregates in the flow of solutions [6] and polymers. But this «microanisotropy» is static and uncontrollable. The present paper deals with the investigation of J-aggregates behavior in a nematic LC that has a long-range order, on the one hand, and can be reoriented in the electric field, on the other hand.

In the present study we continue the investigations of electrooptical properties of the system of J-aggregates of the dye in the LC that have been started in [4]. The orienting medium for J-aggregates is the LC n-pentil-n’-cyanobifenil(5CB). To obtain J-aggregates we used the dye of the pseudoisocyanine family – astrofloxine (AF). Nanocrystals of AF molecules in LC matrix have the spectra and PL lifetime that differs from those of the AF molecules. External electric field can reorient and destroy J-aggregates. The influence of the electric fields of various frequencies and amplitudes is investigated. The role of the surface in
these phenomena is analyzed. High efficiency of two phonon transformation of IR laser radiation into visible light was found for J-aggregates of very long AF in the 5CB LC matrix.

2. Experimental

For the preparation of AF solutions in LC matrix we used n-pentyl-n'-cyanobiphenyl produced by the firm Merk (Germany). 5CB has the temperatures of the phase transitions crystalline LC and LC-isotropic fluid equal to 22.5 and 36.5°C, respectively. To obtain necessary solutions, AF has been at first dissolved in toluene, then mixed with LC and then heated to 70–80°C. During the heating the toluene evaporated and the AF remained in the lquid 5CB where it formed J-aggregates at high concentrations. To obtain molecular solutions of AF toluene and chloroform were used.

To study the influence of the external electric field on the absorption spectra and photoluminescence we used the cells with transparent In2O3 electrodes deposited on the quartz wafers. Internal surfaces of the electrodes in the 10-micrometer thick LC cell were covered with orientant. The orientation of the nematic phase of the mixture was controlled by optical polarizing microscope. The AF dye concentration in LC was 0.5 weight % or less.

For the investigation of the photoluminescence spectra and kinetics of AF in different phases, including liquid crystalline and crystalline states of 5CB matrix, we used the technique with nanosecond resolution [8]. PL was excited by nitrogen laser (λ = 337.1 nm, pulse duration τ1 = 8 ns, pulse power N = 5 kW, frequency ν = 100 Hz). Time-resolved PL signal was registered by stroboscope system that provided the measurements of PL kinetics and PL spectra at different values of delay time τd as to maximum of the laser pulse. Time resolution of the installation for the PL lifetime measurements was about 0.1 ns. We used SPM-2 monochromator for the spectra analysis. The spectral width of the slit varied in the limits of 0.2–0.4 nm. The same monochromator was used for the study of absorption spectra of the solutions LC-AF.

3. From the molecule to the crystal: photoluminescence and absorption of the solutions and crystalline films of astrosefoxine

The formation of J-aggregates in the solutions can be proved using the PL spectra and lifetime measurements. Usually the radiation of the dyes in various solvents at low temperatures exhibits a narrow PL band with a short lifetime. In the PL spectra the band of J-aggregates is shifted to the long wavelengths as compared to the molecular solutions and the lifetime is essentially shorter. Fig. 1 shows the PL spectra of AF in different phases at T = 4.2 K under nitrogen laser excitation. Curves 1, 2 are the PL spectra of AF solution in chloroform with the concentrations C = 10^6 and 10^5 molecule/l, respectively. When the concentration is C = 10^5 molecule/l the PL band at 557 nm with the lifetime 3.5 ns is observed. This spectrum corresponds to the molecular radiation of AF. At C = 10^4 molecule/l the PL spectrum shifts towards the long wavelengths by 10^3 cm^-1. In the spectrum, the bands with the maxima at 590 and 637 nm can be seen, which can be attributed to the formation of the aggregates of different lengths. The lifetime in the PL band at 590 nm is close to 1 ns. Investigations of the PL spectra of J-aggregates with different time delays τd showed that these spectra are rather similar to the PL spectra of the crystalline AF films. At small values of τd the 590 nm band predominates, while with the increase of τd its intensity decreases as compared to the 637 nm band.

Curves 3 and 4 correspond to the PL spectra of the crystalline AF film measured at different time delays. At small τd (curve 3, τd = −3 ns) three bands 592, 612 and 635 nm can be seen in the PL spectrum. The lifetime of the bands 592 and 612 nm is 0.4 ns. With the increase of τd (curve 4, τd = 15 ns) the intensity of 592 and 612 nm bands decreases and the band at 635 nm with the lifetime 2.5 ns predominates. One can suggest that 592 and 612 nm bands correspond to the radiation from the microcrystalline regions with one-dimensional ordering of the molecules while the 635 band corresponds to the radiation of the defects inside these crystalline regions. This suggestion was confirmed by the investigation of the PL of the crystalline AF films under two-photon excitation. In [9] the highly efficient two-photon transformation of IR laser radiation into visible radiation was observed at low excitation levels in several organic crystals. Further investigations demonstrated that the observed high efficiency of the two-photon absorption of the crystals under investigation is caused by high cubic nonlinearity that is peculiar for the highly-conjugated systems of π-electrons with one-dimensional ordering. Investigations of the PL with two-photon excitation at low levels of IR laser radiation can be used as a testing technique for the selection of one-dimensional highly-conjugated nanocrystalline structures with large cubic nonlinearity.
PL of AF crystalline films was excited with two photons of Nd:YAG laser radiation (λe = 1064 nm, N=0.1–0.5 MW). At low excitation levels (unfocused laser beam) the spectra of the luminescence of AF films exhibit three bands with maxima at 620, 650 and 700 nm, corresponding to the crystallites of different sizes.

Investigations of the PL spectra of the AF solutions in LC 5CB demonstrated that they have the properties of both J-aggregates in concentrated AF solutions in chloroform and J-aggregates formed in crystalline AF films. At high concentrations of AF molecules in LC the isolated J-aggregates of certain length and concentration are formed. Extra AF is gathered in the ordered regions of J-aggregates as colloidal particles. These colloidal particles are not uniformly distributed in the bulk of the LC but form dendrimers. With the temperature increase dendrimers are easily destroyed. Colloidal particles of J-aggregates of AF in LC 5CB can be observed in two-photon luminescence spectra at low levels of IR laser excitation. Probably, it is caused by larger length of J-aggregates in colloidal particles as compared to the one in the bulk of LC.

Fig. 2 shows the absorption spectra of AF solutions in LC 5CB for small concentrations of AF (curve 1) and for saturated solutions (curve 2). Curve 3 corresponds to the absorption of the thin crystalline film of AF. It is seen that relative intensity of the long wave length band 560 nm increases and somewhat shifts to longer wavelengths with the increase of AF concentration, thus this band can be ascribed to the aggregates formation. Moreover, the shoulder in the spectral range 570–600 nm appears in the spectrum of the concentrated AF solution. In this spectral region two bands 575 and 600 nm are observed in the spectra of crystalline AF films. One can suggest that both the increase of the long wavelength band and the shoulder appearance at 570-600 nm in absorption spectra of concentrated AF solutions in LC matrix are related to the formation of J-aggregates of different lengths. Longer J-aggregates absorb the light with the wavelengths larger than 575 nm.

4. Electrooptical properties of the solutions of J-aggregates of astrofoxine in liquid crystalline matrix 5CB

Experimentally observed variation of the optical density of AF solution in LC 5CB under external electric field is caused mostly by LC reorientation. AF molecules, that are organized into J-aggregates and oriented along the long axes of 5CB molecules, follow the change of 5CB molecules reorientation. When the LC orientation is planar and the molecules of 5CB together with J-aggregates of AF are oriented parallel to the electrode plane, the absorption of the normal incident light in the AF absorption region will be the largest. The reason is that the transition dipole moment in J-aggregate is oriented along the aggregate axis and is almost equal to zero in the perpendicular direction. Maximum absorption takes place when the E-vector is parallel to long axis of the aggregate. The nematic LC 5CB that is used as a matrix has the positive anisotropy and changes the orientation from planar to homeotropic in the electric field; thus, the dye molecules reorient perpendicular to the E-vector of the light and the sample becomes almost totally transparent. Variation of the optical density of AF solutions in LC 5CB in the external field in the visible range is mostly observed in the region of AF absorption. The reason is that 5CB LC absorbs in the UV region and at λ ≥ 350 nm the LC is transparent.

On the other hand, when the wavelengths of excitation, λ = 365 and 337 nm, correspond to the transparency region of 5CB, the intense luminescence of both LC and dye is observed. The spectral region of LC 5CB radiation is 390–500 nm. The intensity of PL is maximum when E-vector of incident light is parallel to the long axes of 5CB molecules. The PL of LC 5CB was analyzed in [8]. When LC 5CB is excited in the region of molecular absorption (λ < 300 nm) monomer and excimer radiation takes place. The band of the excimer absorption is observed at 410 nm. Taking into account that 5CB molecules in LC phase are not dimerized yet one may suggest that some part of the molecular pairs of 5CB is close to the dimer configuration and gives weak absorption at 300–400 nm. Under the excitation of LC 5CB in this spectral range both excimer and dimer PL is observed. In the external electric field the PL anisotropy of both LC and the dissolved dye molecules changes. Therefore investigation of the PL of the solutions of AF J-aggregates in LC matrix in external fields gives additional information about their electro-optical properties.

In external field the cells with AF solution in LC 5CB exhibit pronounced electrochromic properties. Investigations of the transmission of AF solutions in LC 5CB in d.c. and a.c. electrical fields with the frequencies 1 Hz–1 kHz and amplitude F = 1.2 104 V cm⁻¹ showed that the best reproducibility of the results is observed in the fields with sinusoidal modulation at the frequencies exceeding 100 Hz. The decrease of the electric field frequency from 10³ to 10 Hz at
$F = 1.2 \times 10^4 \text{ V cm}^{-1}$ caused the decrease of the intensity and shift of the absorption band from 564 to 556 nm (Fig. 3). It can be ascribed to higher inertia of J-aggregates in electric field. In the d.c. electric field the AF solution in LC becomes transparent; the process of its initial properties restoration after the field switch-off is rather long. The behaviour observed can be explained by the destruction of J-aggregates in the bulk and on the electrodes surface that causes rather long restoration after field switch-off.

Electrochromic effect can be qualitatively characterized using electrochromic coefficient $K_e$, that can be approximately calculated using the experimental data [11]

$$K_e = \frac{[I(0) - (I(F) / I(0))]}{F^2 D \ln 10}$$  \hspace{1cm} (1)

where $D$ is the optical density of the solution at zero electric field, $I(F)$ and $I(0)$ are the intensities of the transmission light with and without field, respectively. Relative variation of light transmittance $\Delta I / I(0) = [I(0) - I(F)] / I(0)$ of the cells in the region of dye transparency at $10^{-3}$ Hz and $F = 1.2 \times 10^4 \text{ V cm}^{-1}$ is shown in Fig.4. It is seen that main changes of transmission take place in the range of J-aggregates absorption at low electric field frequencies. Depending on the electric field modulation frequency this value varies from $-0.2$ to $-2.2$.

When the electric field changes from 0 to $1.2 \times 10^4 \text{ V cm}^{-1}$ the experimentally observed variations of the optical density of AF solution in LC 5CB are mostly caused by the reorientation of the liquid crystal that is verified by anisotropy investigations. Electrochromic effect for J-aggregates can be seen by eye. At the absence of electric field the cell is pink while in the field it becomes almost transparent.

Fig. 5 shows the spectra of polarized PL of the planar nematic phase of the LC with J-aggregates at Hg line 363 nm excitation: curve 1 – polarization of the incident light is parallel to the direction of the 5CB molecules and to the polarization of the measured PL signal, external electric field is absent; curve 2 – polarizations are the same as for the curve 1 but the electric field with the frequency $f = 1$ kHz and amplitude $U_0 = 12 \text{ V}$ is applied to the electrodes of the cell; curve 3 – incident light polarization is parallel to the direction of 5CB molecules and perpendicular to the polarization of the measured PL signal, external field is absent;

\begin{equation} \hspace{1cm} \begin{array}{c}
\text{Fig. 4. Spectral dependence of } \frac{(I(0) - I(F))}{I(0)} \text{ for the astrofoxine solution in liquid crystalline 5CB matrix with planar orientation at 300 K for } C = 0.5 \text{ w.%. } 1) F = 1.2 \times 10^4 \text{ V cm}^{-1}, \\
\text{frequency } f = 10 \text{ Hz}; \hspace{0.5cm} 2) F = 1.2 \times 10^4 \text{ V cm}^{-1}, \hspace{0.5cm} f = 100 - 1000 \text{ Hz.} \end{array} \end{equation}

\begin{equation} \hspace{1cm} \begin{array}{c}
\text{Fig. 5. Polarized photoluminescence spectra of astrofoxine J-aggregates in liquid crystalline 5CB matrix with planar orientation. } \\
T = 300 \text{ K. } C = 0.5 \text{ w.%. Other experimental conditions are described in the text.} \\
\end{array} \end{equation}
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curve 4 – polarizations are the same as for the curve 3 but the electric field with the frequency $f = 1$ kHz and amplitude $U_0 = 12$ V is applied to the electrodes of the cell.

If the radiative dipole moment of the 5CB cell is parallel to the molecule axis the anisotropy of the uniaxial LC can be written as [4]:

$$A = (I_{pl} - I_{pp})/(I_{pl} + 2I_{pp}),$$

(2)

where $I_{pl}$ and $I_{pp}$ are the intensities of the PL components polarized parallel and perpendicular to the polarization of the incident light, respectively.

At low AF concentrations in LC 5CB the PL spectrum of the solution (Fig. 6, curve 1) is similar to the PL spectrum of the pure LC 5CB [8]. For the planar cell the anisotropy parameter calculated in accordance with (1) is equal to 0.5 in the spectral region 390–500 nm. In the a.c. electric field $(F = 1.2 \times 10^4 \text{ V cm}^{-1}, f = 1 \text{ kHz})$ the PL intensity decreases in the whole spectral range (curve 2) in the limits $\Delta I_{pl}/I_{pl}(0) = (I_{pl}(0) - I_{pl}(F))/I_{pl}(0) = 0.2 - 0.3$ (curve 3). Moreover the spectrum shifts to the long wavelengths by 5 nm which is, probably, due to the influence of the electric field on the electronic subsystem of 5CB molecules.

At high AF concentrations $(C > 0.5 \text{ w} \text{. %})$ in the LC 5CB solutions the PL spectra exhibit intense bands at 577 nm (Fig. 5, curves 1 and 2). Characteristic time and concentration dependencies of this band evidence that the long wavelengths bands in the Fig.5 correspond to the radiation of AF J-aggregates. Due to high AF aggregation degree the radiation of dye monomers is almost absent. The bands in the short wavelengths range in Fig.5 correspond to the PL of the LC matrix molecules. Application of the electric field $(F = 1.2 \times 10^4 \text{ V cm}^{-1}, f = 1 \text{ kHz})$ to the LC cell with the planar orientation of the molecules causes the essential decrease of both LC and J-aggregates PL intensity. Relative change of the PL intensity in the whole spectral range (400–600 nm) is equal to $\Delta I_{pl}/I_{pl}(0) = 0.35$.

Using (2) one can estimate (using the data in the Fig. 5) the spectral dependence of the PL anisotropy of the planar LC and J-aggregates without field (Fig. 7, curve 1) and in the field (Fig. 7, curve 2). At zero field the anisotropy in the PL maximum for LC is $A_{LC} = 0.61$ and for J-aggregates $A_J = 0.47$.

In the electric field the values of the PL anisotropy decrease. It is seen that LC matrix causes the orientation of J-aggregates parallel to the LC molecules, however, the degree of orientation order for the aggregates is lower than for 5CB molecules. Analyzing curves 2 and 4, Fig. 5 one can suppose that the obtained values of A are not the saturated ones. Nevertheless, under the influence of the high frequency electric field LC efficiently reorients J-aggregates along the field. With the increase of the dye concentration in LC up to about 1 w. % the microcrystals (colloidal particles) of the dye «grow» in the matrix; they can be seen in the optical microscope. One can observe the reorientation of these microcrystals when the low frequency electric field is applied to the electrodes of the cell. In this case the characteristic value of the electrochromic effect is $\Delta I_{pl}/I_{pl}(0) = 0.35$, but this is not the limiting value.

Let us analyze the case of the LC with the homeotropic orientation of 5CB molecules in the cell. For the case of ideal homeotropic orientation when 5CB molecules are strictly perpendicular to the cell surface the result is obvious. For all electric field frequencies and amplitudes and all wavelengths of the PL excitation neither 5CB nor J-aggregates in LC matrix exhibit electrochromic effect. The case of the tilted orientation when LC molecules are not exactly perpendicular to the substrate reminds the case of the planar orientation. But the value $\Delta I_{pl}/I_{pl}(0)$ in this case

![Fig.6. PL spectra (curves 1, 2, 3) of the astroloxine J-aggregates in liquid crystalline 5CB matrix with planar orientation and $(I_{pl}(0) - I_{pl}(F))/I_{pl}(0)$ (curve 3) at $T = 300$ K and $C = 0.1$ w.%. 1) $F = 0 \text{ V cm}^{-1}$; 2) $F = 1.2 \times 10^4 \text{ V cm}^{-1}$, $f = 100–1000 \text{ Hz}$.

![Fig.7. Spectral dependence of the photoluminescence anisotropy of the astroloxine solution in the liquid crystalline 5CB matrix with the planar orientation at $T = 300$ K, $C = 0.5$ w.%. 1) $F = 0 \text{ V cm}^{-1}$; 2) $F = 1.2 \times 10^4 \text{ V cm}^{-1}$, $f = 100–1000 \text{ Hz}$. Other experimental conditions are described in the text.

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is much smaller (Fig.8). Depending on the cell (i.e. on the tilt angle of 5CB molecules) the value $\Delta I_{PL}/I_{PL}(0)$ can vary but it does not exceed 0.1–0.15.

Fig.9 demonstrates the influence of the electric fields of various frequencies on J-aggregates. The field with $f > 100$ Hz only reorients J-aggregates along the field; the value $\Delta I_{PL}/I_{PL}(0)$ is constant in the whole frequency range and is close to 0.15. Low frequency electric field with $f < 10$ Hz efficiently destroys J-aggregates. The lower the frequency of the electric field the higher the rate of the conversion of J-aggregates into monomers. Fig. 9 shows that low frequency electric field suppresses J-band and enhances the intensity of the monomolecular PL; note, that the intensity of 5CB radiation is not essentially changed, the intensity of the aggregates PL ($\lambda > 550$ nm) decreases and monomolecular radiation at $\lambda < 550$ nm increases. Destruction of aggregates is caused by electrohydrodynamic flows between the electrodes that occur in the cell in the d.c. or low frequency electric field.

5. Influence of the electrode surface on the AF aggregation

Investigations of the PL spectra of AF J-aggregates solutions in 5CB LC matrix showed that aggregation processes in the bulk and near the electrode are different. These differences can influence the electrooptical properties of the cell as a whole. Fig.10 shows the PL spectra of the cell with AF solution in 5CB LC at 300 K. The measurements were carried out for two cases of the cell filling. In the first case the cell is completely filled with LC solution of AF (curve 1). In the second case the LC with AF is pressed out of the cell (curve 2) and the layer on the electrode surface only partially fills the cell. The thickness of the subsurface layer of LC is smaller than the laser radiation penetration depth in LC. Relative radiation of AF molecules situated just on the surface is rather large in this case. Curve 2 in Fig. 10 is normalized using the short wavelengths part of the curve 1. In the PL spectrum of the surface layer one can distinguish the radiation of AF J-aggregates with the maximum at 585 nm that is characteristic for the radiation in the bulk (curve 1). Moreover, in the differential spectrum (curve 3) which was obtained by the subtraction of the curve 1 from curve 2, two bands, 600 and 615 nm, are seen; they can be as-

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Fig.9. Spectral dependence of the $(I_{PL}(0) - I_{PL}(F))/I_{PL}(0)$ for the astrophiloxine J-aggregates solution in the liquid crystalline 5CB matrix with planar orientation at $T = 300$ K and $C = 0.5$ w.%, and $F = 1.2 \times 10^4$ V·cm$^{-1}$. 1) $f = 100–1000$ Hz; 2) $f = 10$ Hz; 3) $F = 1$ Hz.

Fig.8. PL spectra of the astrophiloxine J-aggregates solution in the liquid crystalline 5CB matrix at the tilted orientation of 5CB molecules: 1) electric field is absent; 2-4) the amplitude of the applied electric field is $U_0 = 12$ V and the frequency is 100–1000 Hz, 10 Hz and 1 Hz for 2, 3 and 4 curves, respectively.

Fig.10. Photoluminescence of the astrophiloxine J-aggregates solution in the liquid crystalline 5CB matrix at the excitation of the thick (1) and adsorbed on the electrodes surfaces (2) layers at 300 K. 3) Differential spectrum for the curves 2 and 3. Curve 2 is normalized by the front part of the curve 1.
6. Photoluminescence of the astrofloxine solution in the liquid crystalline matrix of 5CB at two-photon excitation

J-aggregates are the low-dimensional organic nanocrystals and the collective interaction of the molecules in them causes their pronounced nonlinear properties. First of all the nonlinear properties of J-aggregates are caused by the high degree of \( \pi \) electrons delocalization [13] that leads to anomalously high values of cubic nonlinearity. E.g., nonlinear cubic susceptibility of the solution of pseudoisocyanine J-aggregates solution was measured at room temperature by the method of four-wave light scattering and the value of \( 10^{-3} \text{cm}^2/\text{erg} \) was obtained [5].

Two-photon absorption is related to the cubic susceptibility of the third order. Two-photon absorption can be described by microscopic polarization \( P(\omega) \) that is proportional to the imaginary part of the cubic susceptibility \( \chi^{(3)} \) and the square of the exciting light intensity. One can verify the above statement by calculating the generalized absorption coefficient \( \alpha \) [14] that is equal to the energy absorbed in the unit volume during the unit time, averaged over the time and divided by the energy flux. If \( P(\omega) = \chi^{(3)} EE * E(\omega) \exp(-i\omega t) \) then \( \alpha = -\text{Re}(JE) \rangle / 2I \) where \( J = \partial \rho / \partial t \) is the current density. Taking the differential and dividing by \( I = \langle c / 8\pi \rangle |E| \) one obtains

\[
\alpha_2 = \left\{ \frac{32\pi^2 \omega / c^2}{c} \right\} \chi^{(3)}
\]

(4)

where \( \chi^{(3)} \) is the imaginary part of the third order susceptibility and \( \alpha_2 \) is the coefficient in the formula \( \alpha = \alpha_0 + \alpha_2 I^2 \).

Our investigations showed that the AF solutions in LC exhibit intense PL under two-photon excitation, and this PL is first of all observed for colloidal particles (microcrystals). The efficiency of the IR laser radiation transformation into the visible light was as high as 10%. Thus, investigations of the PL of AF J-aggregates in LC can also be used as a method of these solutions uniformity investigation. To excite the J-aggregates in colloidal particles one needs much lower intensity of the exciting light than for the excitation of isolated J-aggregates in the bulk of LC. The reason is that colloidal particles (microcrystals) consist of rather large J-aggregates with high degree of \( \pi \)-electronic system conjugation. With the increase of the number of molecules in J-aggregate the optical susceptibility of the aggregate should increase.

Fig.11 shows the PL spectra of the AF solution in 5CB LC at 300 K at one- and two-photon excitation. For the two-photon and one-photon excitation the first harmonic \( (\lambda_0 = 1060 \text{ nm}, N = 0.1 - 0.5 \text{ MW}) \) and the second harmonic \( (\lambda_0 = 532 \text{ nm}, N = 0.1 - 0.5 \text{ MW}) \) of Nd:YAG laser were used, respectively. Second order dependence of the PL on the intensity of excitation was measured using the technique of the comparison with the intensity of the second harmonic of organic nonlinear crystal [9]. At the same level of excitation one- and two-photon excited PL spectra are different. In the spectra of one-photon excited PL two bands – 595 and 633 – nm are seen; they can be ascribed to the radiation of the J-aggregates of different lengths. Apparently, in this case we observe the radiation of both isolated and incorporated into the colloidal particles J-aggregates. The spectrum of the two-photon excited PL is shifted to the long wavelengths and is observed in the range 590-800 nm. It consists of two bands with maxima at 650 and 700 nm. One can suppose that the difference of the sizes of short and long J-aggregates in solutions is essential because the radiation at 595 nm is noticeably less intense than in the long wave lengths range. Comparison of the intensities of the two-photon excited PL for AF J-aggregates in LC and for polydiacetylene crystals showed that in both cases it occurs at almost the same excitation levels. This evidences that the coefficients of the cubic susceptibility of these substances are comparable. For the polydiacetylene \( \chi^3 = 1.4 \times 10^{-11} \text{ CGSE units} \) [13].

Difference of the one-and two-photon excited PL spectra of AF J-aggregates in 5CB LC can also be caused by the influence of different states. Two-photon transitions are symmetry-forbidden in the dipole approximation for the states of different parity and are allowed for the states of the same parity. And vice versa for the one-photon transitions – the allowed are the transitions between the states of different parity.

**Fig.11.** Photoluminescence spectra of the astrofloxine J-aggregates solution in the liquid crystalline 5CB matrix at the one-photon (curve 1, \( \lambda_0 = 532 \text{ nm} \)) and two-photon (curve 2, \( \lambda_0 = 1064 \text{ nm} \)) excitation.
Conclusions

The transmission and photoluminescence spectra of astrophloxine solutions in nematic liquid crystal are investigated in the d.c. and a.c. electric fields. AF molecules incorporated into J-aggregates and oriented along the long axes of 5CB molecules follow the changes of the 5CB molecules orientation. When the orientation of the liquid crystal is planar and 5CB molecules with AF J-aggregates are oriented parallel to the electrode the absorption of the normal incident light in the region of the AF absorption has the largest value. Maximum absorption by an AF J-aggregate occurs when the E-vector of the light is parallel to the long axis of J-aggregate. In the external electric field the long axes of AF J-aggregates together with 5CB molecules orient perpendicular to the electrode and E-vector is perpendicular to the long axes of the J-aggregates. In this case the light absorption is minimum. Intense photoluminescence was observed under the two-photon excitation. Efficiency of the IR laser radiation transformation into the visible photoluminescence was as high as 10%.

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