NOVEL CYANINE DYES AS POTENTIAL AMYLOID PROBES: A FLUORESCENCE STUDY

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The applicability of the novel heptamethine cyanine dyes AK7-5 and AK7-6 to the detection and characterization of one-dimensional protein aggregates (amyloid fibrils) associated with numerous pathologies has been evaluated using the method of fluorescence spectroscopy. It was found that both the monomeric and aggregated forms of these dyes can bind to amyloidogenic protein lysozyme, but the concomitant changes in the electronic structure of H-aggregates render them capable of fluorescing. The growth of the hypsochromic bands with negligible changes of the monomeric peaks induced by the native protein and the opposite effects induced by the lysozyme fibrils suggest that the native lysozyme has more binding sites for the dye aggregates than fibrillar protein, while the fibril grooves represent specific binding site for the dyes monomers. The observed spectral behavior of the cyanine dyes, viz. significant distinctions in the fluorescence responses produced by the monomeric and fibrillar forms of lysozyme, suggest the possibility of recruiting these compounds as fluorescent amyloid markers along with the classical amyloid marker Thioflavin T.

KEYWORDS: Heptamethine cyanine dyes, amyloid marker, H-aggregates, fluorescence, lysozyme, amyloid fibrils

НОВІ ЦІАНІНОВІ БАРВНИКИ ЯК ПОТЕНЦІЙНІ АМІЛОІДНІ ЗОНДИ: ФЛУОРЕСЦЕНТНЕ ДОСЛІДЖЕННЯ

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За допомогою методу флуоресцентної спектроскопії проведено оцінку можливості використання нових гептаметинових ціанінових барвників AK7-5 та AK7-6 для детекції і характеристики одномерних білкових агрегатів (амілоїдних фібрил), пов’язаних із розвитком численних патологій. Показано, що мономерна та агрегована форми цих барвників можуть зв’язуватися з амілоїдогенним білком лізозимом, а супутні зміни в електронній структурі H-агрегатів надають їм здатність флуорескувати. Зростання гіпсохромної смуги, поряд з незначними змінами мономерного піка, індукувани нативним білком, і протилежні ефекти, викликані фібрилами лізозиму, дозволяють припустити, що нативний лізозим має більшу кількість сайтів зв’язування для барвника, ніж фібрилярна форма білка, в той час як жолобки фібрил являють собою специфічні сайти зв’язування для мономерів барвника. Спостережувана спектральна поведінка ціанінів, а саме значні відмінності флуоресцентних відповідей, що викликані мономерною та фібрилярною формами лізозиму, свідчать про можливість застосування цих сполук в якості флуоресцентних амілоїдних маркерів, поряд з класичним маркером Тіофлавіном Т.

КЛЮЧОВІ СЛОВА: Гептаметинові ціанінові барвники, амілоїдний маркер, H-агрегати, флуоресценція, лізозим, амілоїдні фібрили

НОВЬІ ЦІАНІНОВІ БАРВНИКИ КАК ПОТЕНЦИАЛЬНЫЕ АМІЛОІДНЫЕ ЗОНДЫ: ФЛУОРЕСЦЕНТНОЕ ИССЛЕДОВАНИЕ

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С помощью метода флуоресцентной спектроскопии проведена оценка возможности использования новых гептаметиновых цинаниевых красителей AK7-5 и AK7-6 для детектирования и характеристики одномерных белковых агрегатов (амиллоидных фибрилл), связанных с развитием множества патологий. Показано, что мономерная и агрегированная формы этих красителей могут связываться с амиллоидогенным белком лизоцимом, а сопутствующие изменения в электронной структуре H-агрегатов делают их способными флуоресцировать. Рост гипсохромной полосы, арду с незначительным изменениями мономерного пика, индуцируемым нативным белком, и противоположные эффекты, вызванные фибриллами лизоцимом, позволяют предположить, что нативный лизоцим обладает большим количеством сайтов связывания для агрегатов красителя, чем фибрилярная форма белка, в то время как жолобки фибрил представляют собой специфические сайты связывания для мономеров красителя. Наблюдаемое спектральное поведение цинанинов, а именно значительные различия флуоресцентных ответов, вызванных мономерной и фибрилярной формами лизоцимами, свидетельствует о возможности применения этих соединений в качестве флуоресцентных амиллоидных маркеров, наряду с классическим маркером Тиофлавином T.
The accumulation of the highly ordered protein aggregates, amyloid fibrils, in various tissues and organs is associated with a variety of human diseases, including Alzheimer’s, Parkinson’s, systemic amyloidosis, type II diabetes, etc. [1]. Amyloid fibrils represent highly ordered one-dimensional protein aggregates up to several micrometers in length and several nanometers in width, consisting of cross-β-structure core, in which β-sheets are oriented along the long axis of fibril [2-4]. One of the most powerful approaches to the detection of amyloid fibrils is based on the use of fluorescent dyes [5]. The classical amyloid markers are Thioflavin T and Congo Red [6-10], whose association with β-sheet structure of amyloid fibrils is accompanied by a substantial increase of fluorescence intensity and significant shifts in the absorption, fluorescent and excitation spectra [11-13]. Nevertheless, these well-known dyes have some drawbacks, among which are low result reproducibility and poor specificity to amyloid fibrils, the propensity for binding to α-helical polypeptide fragments, etc. [14,15]. In view of this, extensive research efforts are currently focused on looking for the more effective amyloid-specific fluorophores.

Near-infrared (NIR) cyanine dyes, a wide class of organic molecules with unique photophysical properties, such as high extinction coefficients, high fluorescence quantum yield and long-wavelength absorption and fluorescence maxima [16-18]. They have found numerous applications in different areas including optoelectronics, laser technologies, bioanalytics, photoelectrochemistry, etc. [19-22]. Due to their advantageous spectral characteristics cyanines have been employed in biomedical research and diagnostics as non-covalent labels for detection of proteins, nucleic acids, lipids and strong oxidizing agents [23-27]. Remarkably, cyanine dyes have been also successfully applied to the identification of amyloid fibrils and inhibition of their growth [28-32]. The advantages of these compounds over the classical amyloid marker Thioflavin T are fluorescence and absorption in the NIR region, high reproducibility of the results and more pronounced changes in fluorescence and absorption spectra upon fibril binding [33].

Most cyanine dyes are known to aggregate in an aqueous solution and show the changes in the emission and absorption spectra with respect to the monomeric species. These assemblies are stabilized by van der Waals, H-bonding, hydrophobic, electrostatic, steric and stacking intermolecular interactions [34,35]. The aggregates that exhibit hypsochromically shifted band (H-band) in their absorption/emission spectrum are called H-aggregates, while J-aggregates display bathochromic shift relative to the monomer band [36]. This behavior was explained by the exciton theory, that considers the individual dye molecule as a point dipole [37]. In the first approximation, taking up only two dye molecules, the interaction between their transition dipoles splits the exciton state of the aggregate into two levels [38]. H-dimers with sandwich-type arrangement (or plane-to-plane stacking) during self-assembly transit to an upper energy level, resulting in the short-wavelength shift of the absorption maximum. The head-to-tail arrangement (end-to-end stacking) of J-dimers is coupled with the transition into lower state with respect to the monomer species, thus exhibiting the long-wavelength shifted absorption band. The structural packing of the dye molecules in the aggregate is described by the slippage angle α defined as the angle between the direction of any one of the parallel molecules and long axis passing through the centers of the aggregated molecules. In particular, when α = 90° the molecules are in a parallel orientation, while when α = 0° a linear molecular orientation is observed [39]. In a general case, the slippage angles ranging from 0° to 32° correspond to J-aggregates, while the angles {32°–90°} are observed for H-aggregates [17].

The aim of the present study is to assess the amyloid-sensing propensity of the two newly synthesized near-infrared cyanine heptamethine dyes, AK7-5 and AK7-6. To this end, we compared the fluorescence responses of the examined dyes in the presence of the native and fibrillar lysozyme. Lysozyme is a well-characterized multifunctional protein with bactericidal, antitumor and immunomodulatory activities. The mutants of human lysozyme (I56T, F57I, W64R, D67H) are prone to pathological fibrillization implicated in the molecular etiology of familial nonneuropathic systemic amyloidosis, a disease affecting kidney, liver and spleen [40].

**EXPERIMENTAL SECTION**

**Materials.** Hen egg white lysozyme (Lz) was obtained from Sigma (St. Louis, MO, USA). Cyanine probes AK7-5 and AK7-6 (Fig. 1) were synthesized in the University of Sofia, Bulgaria, as described previously [41]. Stock solutions of AK7-5 and AK7-6 were prepared by dissolving the dyes in dimethylsulfoxide (DMSO), then diluted by 5 mM sodium phosphate buffer (pH 7.4) and used for spectroscopic measurements. The concentration of the dyes was determined spectrophotometrically, using the extinction coefficients $\varepsilon_{DMSO}^{\text{LzN}} = 0.197 \mu\text{M}^{-1}\text{cm}^{-1}$ and $\varepsilon_{914}^{\text{DMSO}} = 0.209 \mu\text{M}^{-1}\text{cm}^{-1}$ for AK7-5 and AK7-6, respectively.

**Preparation of amyloid fibrils.** Amyloid fibrils were obtained from the egg yolk lysozyme by the protein incubation in 10 mM glycine buffer at pH 2 and 60 °C for 14 days [41]. Protein concentration in the stock solution was 10 mg/mL. The working solutions of the native and fibrillar lysozyme were prepared in 5 mM sodium phosphate buffer (pH 7.4). Hereafter, the native and fibrillar protein forms are designated as LzN and LzF, respectively.

**Fluorescence measurements.** Fluorescence spectra of AK7-5 and AK7-6 were recorded with the spectrofluorimeter Shimadzu RF-6000 (Japan) at 25 °C using 5 nm excitation and emission slit widths. The excitation and emission wavelengths are indicated in the legends to figures.
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RESULTS AND DISCUSSION

At the first step of study we analysed the emission and excitation spectra of AK7-5 and AK7-6 in an organic solvent and an aqueous solution. Recently, using the absorption spectroscopy technique we have found that both the above dyes can form non-fluorescing H-aggregates in an aqueous solution, but tend to disaggregate in the presence of protein [42]. In an organic solvent, DMSO, AK7-5 and AK7-6 monomers display fluorescence emission maxima at 835 and 839 nm, respectively, while fluorescence excitation maxima are observed at 816 nm for both dyes (Fig. 2). In the sodium-phosphate buffer solution the examined dyes have the emission maxima similar to DMSO, but there were no bands corresponding to H-dimers or H-aggregates, suggesting that such kinds of aggregates are non-fluorescent. Notably, this observation is in concert with the literature data [43].

The monomeric peaks of AK7-5 and AK7-6 gradually increased upon the dye titration with the native lysozyme, but above a certain protein concentration the fluorescence intensity reduced to a nearly initial level. This process was accompanied by the appearance of a hypsochromic band at 700 nm whose intensity increased with the protein concentration (Figs. 3a, 4a).

The dye concentration was 14.4 µM. The excitation wavelength was 557 nm.

Fig. 1. Structures of the AK7-5 (a) and AK7-6 (b) dyes.

Fig. 2. Emission and excitation spectra of AK7-5 and AK7-6
a) in DMSO, b) in sodium-phosphate buffer. The concentrations of AK7-5 and AK7-6 were 14.4 and 9.8 µM, respectively.

Fig. 3. Fluorescence spectra of AK7-5
a) at the increasing concentration of the native lysozyme, b) at the increasing concentration of fibrillar lysozyme
The dye concentration was 14.4 µM. The excitation wavelength was 557 nm.
In the presence of lysozyme fibrils, both heptamethine dyes demonstrated significant rise of monomeric bands (by a factor of 4.5/3 for AK7-5/AK7-6, respectively) without any decrease of the hypsochromic bands and their slight enhancement compared to the native protein (Figs 3b, 4b).

By and large, the rise in the fluorescence intensity of monomer band can be caused by the two processes: i) the growth of the monomer concentration due to a shift of equilibrium between the different dyes forms in the buffer after addition of the protein, as was previously demonstrated using absorption spectroscopy [42]; ii) increase of the dye quantum yield arising from its immobilization on the protein binding site. The interference between these processes complicates the quantitative analysis of the obtained results. Taking into account the absorption spectroscopy data [42] it can be assumed that the emission band of 600-800 nm corresponds to the H-dimers and H-aggregates of the examined dyes. This finding is especially interesting because, as mentioned above, in an aqueous media H-dimers and H-aggregates are commonly uncapable of fluorescing. Allowing for this fact, it can be supposed that the binding of H-aggregates to the native or fibrillar protein is followed by the changes in the geometric characteristics and electronic structure of H-dimers and H-aggregates of cyanine dyes, thereby causing an increase in the fluorescence quantum yield.

Remarkably, such an unexpected fluorescence was previously observed for the H-aggregates of thiazole orange and merocyanine dyes [44, 45]. It is also noteworthy that the intensity of this hypsochromic band in the presence of the native protein was 1.5 - 2.5-fold higher compared to the lysozyme fibrils.

![Fig. 4. Fluorescence spectra of AK7-6](image)

**a)** at the increasing concentration of the native lysozyme, **b)** at the increasing concentration of fibrillar lysozyme

The dye concentration was 4.9 µM. The excitation wavelength was 561 nm.

Qualitatively, this finding can be rationalized by: i) the formation of the complexes “monomer-protein”, “dimer-protein”, “H-aggregate-protein” shifting the monomer-aggregate equilibrium; ii) the binding of the dye monomers, H-dimers and H-aggregates to the native and fibrillar lysozyme manifesting itself in the changes in the fluorescence spectra. The fact that the more significant increase of the hypsochromic band 600-800 nm was found for LzN, suggest that the native protein has more binding sites for cyanine aggregates than LzF. Presumably, the hydrophobic, electrostatic, and van der Waals interactions between the dye monomers and the native protein promote the formation of the dye dimers and H-aggregates. The electrostatic interactions between the positively charged molecules of AK7-5/AK7-6 and negatively charged lysozyme cavities increase the dye-protein affinity. At the same time, despite the LzF has more exposed hydrophobic residues compared to LzN, the contribution of hydrophobic interactions to the formation of the complexes “H-dimer-protein” and “H-aggregate-protein” is less than that of electrostatic forces. It can also be noted that LzF produced a significant increase in the monomeric peak and much less pronounced rise in the peak at 700 nm compared to LzN. The observed substantial fluorescence increase at 840 nm can be regarded as an unambiguous proof of specific binding of the dye monomers to lysozyme fibrils.

**CONCLUSIONS**

- Both the monomeric and aggregated forms of the novel heptamethine cyanines, AK7-5 and AK7-6, can bind to lysozyme, but the concomitant changes in the electronic structure of H-aggregates render them capable of fluorescing. The growth of the hypsochromic bands with negligible changes of the monomeric peaks induced by the native protein and the opposite effects brought about by the lysozyme fibrils suggest that the native lysozyme has more binding sites for the dye aggregates than fibrillar protein, while the fibril grooves represent specific binding site for the dyes monomers.

- The changes of the monomer band observed in the presence of the native lysozyme is indicative of the disassembly of the H-aggregates in solution followed by their assembly with presumably modified properties on the protein surface. At the same time, the fluorescence intensity of monomeric dyes gradually increases with the concentration of lysozyme fibrils without any re-assembly of the H-aggregates within the fibril structure.
• Cumulatively, the revealed spectral behavior of the cyanine dyes, viz. significant distinctions in the fluorescence spectra in the presence of monomeric and fibrillar forms of lysozyme, point to the possibility of using these compounds as fluorescent amyloid markers along with the classical amyloid marker Thioflavin T.

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REFERENCES


