Савицкая Т.А., Шахно Е.А., Босько И.П., Матулис Вадим Э., Мелеховец Н.А., Гриншпан Д.Д., Ивашкевич О.А. Комплексообразование канамицина с сульфатом ацетатом целлюлозы: перспективный путь от инъекционной лекарственной формы к пероральной

https://doi.org/10.33581/2520-257X-2021-1-3-20

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Water-insoluble polyelectrolyte complexes of cellulose acetate sulphate in the form of sodium salt (Na-CAS) and aminoglycoside antibiotic (AB) kanamycin (KAN) were obtained by mixing of the components aqueous solutions. The composition of the complexes was determined in accordance with the medium pH and mixing order. The increase of Na-CAS cellobiose units per mole of AB has been shown to correlate with the decrease of pH value. The complex formation was studied by Fourier transform infrared spectroscopy, thermal analysis, X-ray analysis, laser diffraction, motion trajectory of nanoparticles analysis and scanning electron microscopy. Quantum-chemical study of the relative stability of the protonated forms of KAN in aqueous solution was performed to determine the preferred protonation sites of KAN molecule. The pK\textsubscript{a} values of KAN were calculated by means of isodesmic reactions method. The structures and binding energy for the KAN dimer and the KAN – CAS complex were also investigated by quantum-chemical methods. Na-CAS – KAN complex itself and immobilised on the activated carbon was shown to demonstrate \textit{in vitro} two times antibacterial activity of the standard (injectable) form of KAN against \textit{Mycobacterium tuberculosis}. It can be recommended for \textit{in vivo} clinical trials as a new form of aminoglycoside AB for oral administration.

\textbf{Keywords:} tuberculosis; kanamycin; cellulose acetate sulphate; complex; activated carbon; oral dosage form.

\section*{Introduction}

Taking a heavy toll on the world, tuberculosis (TB) as an infectious disease was named among the 2016’s top 10 causes of death worldwide. In 2018, the first-ever UN General Assembly high-level meeting on tuberculosis endorsed a global strategy to accelerate efforts in ending TB [1]. Along with social-economic determinants, the main risk factors for TB include prolonged treatment and difficulties associated with chemotherapy. TB treatment usually involves the repeated continuous administration of large doses of different drugs, which leads, especially in the case of parenteral use, to various side effects. Addressing these issues requires improvement of already existing drugs formulation as well as the development of new anti-TB drugs with preference oral versus injectable forms.

Recently, many strategies based on new delivery materials have been developed to enhance the antibacterial efficacy and comfort taking of commonly used TB antibiotics. For example, dry powder aminoglycoside
antibiotic (AB) aerosol products were proposed for inhaled therapy for TB [2], but there is a discussion in literature about safety and long-term effects of this therapy. The hydrophobic ion-pairing approach has been recently suggested as a chemical strategy to reversibly modify the properties of AB. Using the hydrophobic ion-pairing in drug molecules containing ionisable groups, polar counter-ions are stoichiometrically replaced with lipophilic ones. The modification by lipoamino acid was described for aminoglycoside AB kanamycin [3]. The resulting ion-pairs improve drug permeability, allowing a better systemic (e.g., intestinal) adsorption and enhancing cellular uptake. At the same time, the modification by polymers, which is considered as a method creating new easy-to-use medications and ameliorating drug-induced allergic and toxicological reactions, is recently a subject of numerous research but still not implemented in anti-TB antibiotic design at the industrial level [1; 4; 5].

Aminoglycosides are one of the first AB. Produced by Streptomyces kanamyceticus kanamycin (KAN) was discovered over 60 years ago but in spite of such a long history it remains the successful broad-spectrum antibacterial agent, in particular the important second-line AB in the standard treatment regimen for multi-drug resistant TB. It exhibits the bactericidal action towards the majority of gram-positive and gram-negative microorganisms, and also the acid-resistant bacteria (including Mycobacterium tuberculosis) [3; 6]. Its irreversible binding to mRNA decodifying region of the bacterial ribosome 30S subunit allows the inhibition of the protein synthesis [7].

KAN is used in the form of mono- or disulphates intravenously and intramuscularly, since it is practically not absorbed from the gastrointestinal tract (GIT); absorption capacity is less than 1 % in the absence of mucosa damage [2]. The oral application route of KAN in capsules, tablets or syrup is used pre-eminently for local sterilisation of the gastric cavity before operative treatment [6]. Therefore, the replacement of the injections with solid oral dosage forms will make it possible to improve and substantially simplify the treatment regimen for tuberculosis patients, making it less traumatic and invasive.

The focus of the present study was to modify KAN sulphate into new oral drug form through its converting into the polymer salt with cellulose acetate sulphate in the form of sodium salt (Na-CAS) as counter-ion and by further immobilization of this ionic-pair on the activated carbon (AC).

According to [8–10] the dimerisation and higher clustering of aminoglycoside AB is known. However, literature on aggregation of KAN or related aminoglycosides is extremely sparse, both on the experimental and on the theoretical side. It is reported that KAN aggregates into bare dimers and dimers with certain physiological cations, preferentially sodium one [8]. Covalent linked dimers have been obtained by using a «tether» and studied [9]. Formation of long fibers as a linear noncovalent assembly of KAN molecules on negatively charged surfaces was determined as well [10].

The experimental study included the synthesis of KAN and Na-CAS complexes, the establishing of the components concentration and pH medium impact on the complexes composition, complexes solubility and dispersion, bacteriological activity in parallel with theoretical research such as quantum-chemical calculations, molecular dynamics simulations of KAN protonation, dimerisation and complexation with Na-CAS.

Materials and methods

Materials. KAN sulphate (2-(aminomethyl)-6-[4,6-diamino-3-[4-amino-3,5-dihydroxy-6-(hydroxymethyl) oxan-2-yl]oxy-2-hydroxyclohexyl]oxyxane-3,4,5-triol; sulphuric acid) in an injectable powder form was purchased from Kyivmedpreparat (Ukraine) in the form of acid sulphate, KAN base – from North China Pharmaceutical (China). We took into account that as commonly occurs with natural products, this antibiotic turned out to be a complex of several closely related substances: a major component, kanamycin A (usually designated as kanamycin), and two minor variants (less than 5 %), kanamycin B and C [11]. However, we considered that AB behaviour in the complexation process determines mainly by kanamycin A and this form was used in quantum-chemical calculations.

Na-CAS (poly-((1β→4)→(2-O-acetyl-6-sulpho-D-glucopyranose)) was homogeneously synthesised according to [12] and had viscosity average molecular weight ($M_v$) 36000, (14.4 ± 0.1) % of combined acetic acid and (32.0 ± 0.1) wt. % of combined sulphuric acid. Cellobiose unit was chosen as a configuration repeated unit of Na-CAS (fig. 1) in spite of the fact that some authors consider glucose unit but as chemical composition repeated unit [13] (the molecular weight of cellobiose mole-link was 574) and polymer concentration was expressed in the moles of these units.

AC in the form of a fine powder of birch activated charcoal from Sorbent JSC (Russia) was used for immobilisation of a complex when tablets prepared. AC was considered as a carrier for AB – polymer complex in the design of the solid dosage form.
Formation of Na-CAS and KAN complex. AB and polymer solutions of appropriate concentrations were prepared and mixed in molar ratio between AB and polymer from 1 : 8 to 8 : 1, changing mixing order and pH medium. Distilled water, buffer solutions with pH 2.0, 6.0 and 7.5 were used. Before mixing, the solutions were filtered through 0.45 μm pore size membrane. The composition of obtained complexes was determined using colloid titration technique on CPC-3-01 photoelectric colorimeter (ZOMZ, Russia) at 540 nm wavelength. This technique is based on the system ability to scatter light when a dispersion of insoluble particles is formed. Under the direct complex synthesis, polymer solutions with a concentration of 10^{-3} mol/dm^3 were titrated by the solutions of AB with a concentration of 10^{-2} mol/dm^3 by sequentially adding 5 μL portions of the solution under stirring until the growth of the solid ceased, which was indicated by reaching constant optical density. Optical density D was measured 1 min after the addition of each portion of titrant. Using the inverse mixing order, solution of AB with a concentration of 10^{-3} mol/dm^3 was titrated by the solution of polymer with a concentration of 10^{-2} mol/dm^3 by sequentially adding 5 μL portions of the solution. The content of AB in the complex was determined by chemical analysis (Kjeldahl technique). All experiments were performed at ambient temperature.

Fourier transform infrared spectroscopy (FTIR) and scanning electron microscopy (SEM). Samples for FTIR and SEM were prepared in the form of coevaporates obtained by applying Na-CAS – KAN complex particles suspensions. The above-mentioned suspensions were obtained by mixing 10 wt. % solutions of the components in a ratio corresponding to the complex composition, then pouring on a glass substrate, followed by evaporating water in air at the ambient temperature. The FTIR spectra were recorded on a Nicolet iS10 FTIR spectrometer (Thermo Fisher Scientific, USA) through 46 scanning at a 0.5 cm^{-1} resolution. The morphology of the films surface after deposition of gold (a Emitech K550 instrument was employed) was examined on a LEO-1420 scanning electron microscope (Carl Zeiss, Germany) at magnifications from 500× to 20 000×.

Thermal analysis. Thermal analysis was carried out through thermogravimetric (TG), differential thermogravimetric analysis (DTG), differential scanning calorimetry (DSC) on a thermal analyser STA 449 Jupiter (Netzsch, Germany). Samples of the complexes in the form of powders were obtained by mixing components solutions, precipitates filtration through the paper filter and under-vacuum drying. Analysis was performed at a heating rate of 10 K/min in the temperature range 303–600 K with nitrogen gas flow of 25 cm^3/min and 8–10 mg samples.

Laser diffraction method. The particles hydrodynamic size in the solutions of Na-CAS and in the sols of its complexes with KAN, obtained by mixing of solutions with Na-CAS concentration of 2 · 10^{-2} mol/dm^3 and AB concentration corresponding to the compositions of the complexes, were estimated using laser diffraction method on a Zetasizer nano ZS device (Malvern, UK) at the temperature 298 K. The kinetics of the complex particles growth was evaluated by a Mastersizer 3000 laser particle size analyser (Malvern) using a wet dispersion unit with a measuring vessel volume of 120 cm^3 at a rotation speed of 1500 min⁻¹. The formation of complex particles dispersions resulted directly in the vessel by adding of the polymer solution (0.02 mol/dm^3) to the AB solution with the concentration of 0.002 mol/dm^3 and by different mixing order.
The particles size distribution was also estimated by the analysis of motion trajectory of nanoparticles on a Nanosight LM0 multiparametric analyser of nanoparticles (Nanosight Ltd., UK) in the HS – BF configuration (Andor Luca high-sensitivity video camera; a semiconductor laser with a wavelength of 405 nm and a power of 45 mW).

**X-ray diffraction analysis (XRD).** Powders of KAN, Na-CAS and their complex XRD analysis was carried out on a Drone 3.0 X-ray diffractometer (Bourevestnik, Russia), CuKα radiation, at an accelerating voltage of 50 kV. Data were systematically collected from 23° to 3° in 2θ with a scan speed of 2 °/min in 2θ and a step size of 0.05°.

**Bacteriological tests.** The content of the bacteriologically active AB form was determined with the bioassay technique on agar plates. Tablet forms of the water-insoluble complexes immobilised on the activated carbon sorbent were obtained by triturate method with the laboratory setup, followed by drying at $T = (303 \pm 1) \ K$ in an air flow to a constant weight. Tests to determine the drug susceptibility of *M. tuberculosis* to the action of this new KAN pharmaceutical form were carried out by the absolute concentration method using Löwenstein – Jensen medium [15].

**Computational details.** Protonation of KAN A in aqueous solution. The starting geometry of KAN A was taken from the Cambridge Crystallographic Data Centre\(^1\). The geometrical parameters of KAN A molecule and corresponding protonated structures were fully optimised using CAM-B3LYP/6-31G(d) level of theory [16]. Our previous investigations showed that this computational level provides good agreement of the calculated geometries of heterocyclic compounds with the experimental data [17]. The optimised geometries were used for single point energy calculation using CAM-B3LYP/6-31+G(d, p) level of theory. The solvent effects were evaluated using the polarised continuum model [18] with the default parameters for water.

The pK\(_a\) values were calculated by means of isodesmic reactions method. We considered the following isodesmic reaction:

$$\text{BH'} + \text{OH} \rightarrow \text{B} + \text{OH},$$

where B is kanamycin A molecule.

The isodesmic reaction Gibbs energy in an aqueous solution was calculated using the following equation:

$$\Delta_{rs} G_s = \sum_{\text{products}} G_s - \sum_{\text{reactants}} G_s.$$

Further, it can be proved that

$$\text{pK}_a(\text{B}) = \text{pK}_a(\text{ethanolamine}) + \frac{\Delta G_s}{RT} \ln 10,$$

where pK\(_a\) (ethanolamine) is equal 9.5 [19].

Our previous studies have showed that this computational method gave the calculated pK\(_a\) values for heterocyclic compounds which were consistent with the experimental data [20]. Obviously, the KAN A molecule and the corresponding protonated structures has many different conformations. The data below refer to the most stable ones.

**Binding energy of KAN dimer and KAN – CAS complex.** By virtue of the fact that complex composition corresponds to one mole of KAN per two CAS glucose mole-links we used cellobiose unit as a polymer chain fragment. The semi-empirical GFN-xTB method [21] has been used for the conformational search to obtain the initial structures for global minimum searching. The resulting structures were ranked by energy, and the geometry of the ten lowest energy structures have been optimised using PBEh-3c/def2-mSVP method [22], which is computationally cheap and shows good performance for non-covalent interaction energies in complexes. The most stable structures were then optimised on the wB97XD/6-31G(d) level of theory.

For both dimer and complex structures basis set superposition error (BSSE) was estimated and subtracted from energies of these structures. All the calculations have been carried out for isolated molecules (gas phase).

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\(^1\)The Cambridge Crystallographic Data Centre (CCDC) [Electronic resource]. URL: https://www.ccdc.cam.ac.uk/ (data of access: 20.06.2019).
The binding energy of KAN dimer and KAN – CAS complex has been calculated using the following equations:

\[
\Delta E_d = E_d - 2E_m,
\]

\[
\Delta E_c = E_c - E_m - E_{\text{CAS}},
\]

where \( E_d, E_m, E_c, E_{\text{CAS}} \) are the total energies of dimer, monomer, complex and CAS fragment respectively.

**Results and discussion**

Na-CAS and KAN aqueous solutions mixing results in the formation of insoluble hydrophobic product due to the co-operative interaction between functional groups of polyelectrolyte chain and AB molecules. The terms «complex of an antibiotic with a polymer» or «polymeric salt of an antibiotic» are used in literature to refer to similar substances [18].

Depending on the components solutions concentrations, the complex dispersion can be obtained as either a sol \((c < 10^{-5} \text{ mol/dm}^3)\) or suspension \((c > 10^{-5} \text{ mol/dm}^3)\), the optical densities of which are proportional to the size and concentration of the complex particles.

As follows from fig. 2, the composition of the complex corresponds to \([\text{Na-CAS}] : [\text{KAN}] = 2.4 : 1.0\) and \([\text{Na-CAS}] : [\text{KAN}] = 2.0 : 1.0\) at direct and inverse titration respectively.

There are four possible protonation sites in the KAN molecule (nitrogen atoms of four amino groups). The protonation of AB molecules affects their conformation and interaction with target biomolecules. For this reason, data on the thermodynamics of aminoglycoside protonation together with exact assignment of the protonation sites are necessary for the interpretation of their activity. According to the data [23] the stepwise protonation constants \(pK_a\) of KAN are 9.16 (C6′ (ring A)), 8.27 (C1 (ring B)), 7.52 (C3″ (ring C)) and 6.28 (C3 (ring B)). In this work, we performed a quantum-chemical study of the relative stability of the protonated forms of KAN in aqueous solution to determine the preferred protonation sites. Figure 3 represents the optimised structures of four KAN A ions with different protonated amino groups.

Table 1 lists the relative (the smallest value has been taken as zero) Gibbs free energies of the ions. The results of quantum chemical calculations show that the stability of protonated forms of KAN A decreases as follows

\[ \text{C6}' \ (\text{ring A}) \sim \text{C1} \ (\text{ring B}) > \text{C3} \ (\text{ring B}) > \text{C3}'' \ (\text{ring C}). \]

The results obtained agree with the experimental data [23] on the protonation sequence of the amino groups of KAN A molecule in aqueous solution:

\[ \text{C6}' \ (\text{ring A}) \rightarrow \text{C1} \ (\text{ring B}) \rightarrow \text{C3}'' \ (\text{ring C}) \rightarrow \text{C3} \ (\text{ring B}). \]

The \(pK_a\) values of the deprotonation for each amino group of KAN A ions were calculated and compared with experimental data [23] (table 2). The calculation results agree with experimental data.
Thus, the preferred protonation sites of KAN in aqueous solution were determined: amino groups at the C6’ (ring A) and C1 (ring B). Further, this will be taken into account when calculating the structure of the Na-CAS – KAN complex.

Gradual protonation of KAN base caused by adding calculated amounts of hydrochloric acid to KAN aqueous solution followed by the rearrangement of AB hydrogen bonds system. Table 3 shows that there is a tendency to shift bands in FTIR spectra of KAN solutions with different concentrations of HCl but to a certain limit. The position of the bands ceases to change after the tetra protonated KAN is formed. For example, OH group of one of the terminal ring involved in the hydrogen bond with NH₂ group is responsible according
to [9] for the peak at 3250 cm$^{-1}$ in a dimer, that essentially corresponds to the monomer peak at 3338 cm$^{-1}$. In our case there was 3258.9 cm$^{-1}$ peak in the initial solution. After the protonation of all NH$_2$ group it places at 3100.0 cm$^{-1}$.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Bands position, cm$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>KAN acid sulphate dry powder</td>
<td>3142.1 1614.7 1521.1 1022.6 –</td>
</tr>
<tr>
<td>Aqueous solution of KAN base without HCl</td>
<td>3258.9 1634.8 1531.1 1057.6 –</td>
</tr>
<tr>
<td>Aqueous solution of KAN base with HCl equivalent for 2 amino groups</td>
<td>3171.0 1624.9 1523.3 1023.6 –</td>
</tr>
<tr>
<td>Aqueous solution of KAN base with HCl equivalent for 3 amino groups</td>
<td>3170.0 1613.6 1509.9 1023.3 863.7</td>
</tr>
<tr>
<td>Aqueous solution of KAN base with HCl equivalent for 4 amino groups</td>
<td>3100.0 1609.6 1506.5 1022.0 861.5</td>
</tr>
<tr>
<td>Aqueous solution of KAN base with HCl equivalent for 6 amino groups</td>
<td>3100.0 1609.1 1505.9 1022.7 860.6</td>
</tr>
<tr>
<td>0.1 mol/L HCl (pH 1.0)</td>
<td>3100.0 1605.7 1505.1 1040.1 858.7</td>
</tr>
</tbody>
</table>

Simultaneously, the increasing of Na-CAS mole-links per KAN mole takes place as shown in fig. 4. It demonstrates that the more the acidity of the medium, the more colloidal titration curves shift to the right along the abscissa axis. This means that NH$_2$ group protonation causes an increase in the amount of Na-CAS mole-links connected with KAN mole.

![Fig. 4. Colloid titration curves of Na-CAS aqueous solution by the KAN base solutions (1) with different amount of HCl added for the protonation of 1, 2, 3 and 4 amino groups respectively 2, 3, 4 and 5.](attachment:figure4.png)

Turning back to fig. 2, one can see that the composition of the complex in the medium of acetate buffer (pH 6.0) corresponds to a molar ratio of [KAN] : [Na-CAS] = 1.0 : 2.4 at direct titration. Figure 2, a, shows that adding of AB excess leads to a slight decrease in turbidity due to the hydrophilisation of the complex particles via binding of the extra amount of ionogenic KAN molecules, that are relatively small in size compared to polymer macromolecules. In the case of inverse titration, the curve maximum corresponds to a ratio KAN : Na-CAS = 1.0 : 2.0 with the observed decrease in optical density following by the addition of polymer solution to the resulting dispersion of the complex particles. In this case, the system becomes transparent, which can be attributed to the transition of the complex into the dissolved state due to hydrophilisation asso-
associated with the decrease in the number of contacts of the Na-CAS sulphate groups with the protonated KAN amino groups. The decrease in turbidity in such a circumstance is not caused by dilution, since, as follows from fig. 2, b (curve 1), when diluted with water, the optical density of the system does not change.

Table 4 data confirms that the decrease in turbidity is caused by the complex dissolution, rather than by its degradation. As can be seen from table 4 the size of the complex particles in hydrosol exceeds that of the particles in a solution of polymer itself with a concentration corresponding to its partial concentration in hydrosol. The excess amount of the polymer followed by the growth of particles size. At the same time, the size of the particles in the solutions of individual polymer falls down with a concentration that is typical for polyelectrolyte solutions behaviour. Attention is drawn to the fact that, when the [Na-CAS] : [KAN] molar ratio of 3.0 : 1.0 is achieved, the size of the complex particles practically stops changing, due to the saturation of the protonated amino groups, that are capable of interacting with the polymer. Figure 5 illustrates the particle size distribution for all these cases.

<table>
<thead>
<tr>
<th>Na-CAS : KAN molar ratio</th>
<th>Average hydrodynamic diameter of the particles, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hydrosol of Na-CAS – KAN complex</td>
</tr>
<tr>
<td>1.0 : 1.0</td>
<td>350</td>
</tr>
<tr>
<td>2.0 : 1.0</td>
<td>380</td>
</tr>
<tr>
<td>3.0 : 1.0</td>
<td>395</td>
</tr>
<tr>
<td>4.0 : 1.0</td>
<td>390</td>
</tr>
</tbody>
</table>

Fig. 5. Differential curves of particles hydrodynamic size distribution estimated via laser diffraction technique for hydrosols of Na-CAS – KAN complexes for different amount of Na-CAS moles per KAN mole: 1 – 1; 2 – 2; 3 – 3 and for aqueous solutions of Na-CAS with corresponding partial concentration 4, 5, 6 respectively.

Determination of Na-CAS – KAN complexes particle size in the dispersions, obtained through the blending of solutions with different concentrations, revealed that the higher the concentration of the solutions, the larger the particles size. For instance, with an increase in the concentration of the polymer solution from 0.02 to 0.16 mol/dm³, the particle size increases from 20 to 45 μm. At the same time, direct titration produces lar-
ger particles than the inverse titration, since the amino groups of the same AB molecule can, in the first case, interact with the sulphate groups belonging to different Na-CAS macromolecules, stitching them together. In the second case, the number of contacts of these complementary groups decreases. Indeed, with the reverse method of components solutions blending, the average particle size in the Na-CAS – KAN suspensions only reached 0.5 μm. As follows from fig. 6, a, the kinetics of particle growth also depends on the method of the components solutions mixing. With the inverse method, particles of a finite size are formed immediately, whereas with the direct method, it takes at least 15 min to achieve a sizable growth. Figure 6, b, illustrates the distribution of the particles for Na-CAS – KAN complexes.

Non-identical composition of the complex particles in the dependence of mixing order was confirmed by the agar diffusion test. It was held to determine the percentage of the bacteriologically active AB form in a complex and showed (table 5) that an inverse order of the components mixing results in almost 10 wt. % higher KAN content in the complex than a direct one. The elemental chemical analysis confirmed this result.

The visualisation of Na-CAS – KAN complex particles was carried out by the SEM technique (fig. 7). The evidence presented suggests that, after removing of the dispersion medium, the complex particles tend to form spherical aggregates with sizes ranging from 2 to 8 μm for the direct mixing order, and less than 1 μm for the inverse one. This morphological picture is similar to that one for Na-CAS film that we published earlier [12].

Fig. 6. Distribution of the particles for Na-CAS – KAN complexes:

a – kinetic curves of the particles size growth for Na-CAS – KAN suspensions under direct titration at the concentration of Na-CAS solutions 0.02 mol/dm³ (1), 0.04 mol/dm³ (2), 0.08 mol/dm³ (3), 0.016 mol/dm³ (4);

b – inverse titration curve;

c – histogram of particle size distribution for hydrosol of Na-CAS – KAN complex (b) obtained by the analysis nanoparticles motion trajectory respectively

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Table 5

The content of the bacteriologically active AB form in Na-CAS – KAN complexes determined with the bioassay technique on agar plates and chemical analysis

<table>
<thead>
<tr>
<th>Complex preparation method</th>
<th>KAN content, wt. %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Bioassay technique</td>
</tr>
<tr>
<td>Direct</td>
<td>43,8 ± 0,9</td>
</tr>
<tr>
<td>Inverse</td>
<td>52,3 ± 0,5</td>
</tr>
</tbody>
</table>

Table 6

Bands position in FTIR spectra for individual components and their complex

<table>
<thead>
<tr>
<th>Composition</th>
<th>Band position, cm⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>KAN</td>
<td>3150 1618 1529 – –</td>
</tr>
<tr>
<td>Na-CAS</td>
<td>3440 1637 – 1225 799</td>
</tr>
<tr>
<td>Complex Na-CAS – KAN</td>
<td>3380 1631 1536 1220 798</td>
</tr>
</tbody>
</table>

Fig. 7. SEM photos of the Na-CAS – KAN complex particles in the case of direct (a) and inverse (b) order of the components solutions mixing

Fig. 8. XRD curves of Na-CAS (1); Na-CAS – KAN complex (2); KAN (3)
Initial salt form of KAN destroys as a result of complex formation and AB becomes connected strongly with polymer in the insoluble complex. The complexion is accompanied by the changing in the individual components’ bands position in FTIR spectra. As follows from table 6, significant changes take place in the field of hydrogen bonds. According to fig. 8, the structural features of KAN preserves in the complex (curves 2 and 3). To imagine the structure of the complex quantum chemical calculations have been done. Computational studies on KAN are rarer still [9]. Molecular dynamics and docking studies were performed for aminoglycoside monomers and RNA [24; 25], using standard force fields and scoring functions. Apparently only in two cases electronic structure theory methods were applied to KAN: one of them again in the context of KAN binding to RNA, the other one for the isolated monomer in the gas phase [26]. Both studies, however, only employed HF calculations with small basis sets. Theoretical studies on KAN A complexion with cellulose derivative appear to be non-existent. In this work, we performed a quantum-chemical study of the structure and binding energy of the KAN dimer and the Na-CAS – KAN complex. The structures of the lowest energy conformations of the KAN A dimer and the Na-CAS – KAN complex are shown on fig. 9 and 10. The calculated BSSE corrected binding energies of KAN A dimer and KAN – CAS complex are –199 and –299 kJ/mol respectively. This shows that complex formation is an energetically preferable process compared to dimerisation.

Figure 10 shows that in KAN-CAS complex KAN molecule is protonated on amino groups at С6’ (ring A) and С1 (ring B). This is consistent with the results of our calculation of pK\textsubscript{a} value of KAN A [24], showing that amino group at С6’ (ring A) is the most basic.

The involving of polymer and drug functional groups into different kinds of interaction influences on the properties of their complex as opposed to the initial components.

Figure 11 demonstrates DSC, DTG and TG curves for KAN, Na-CAS and their complex in the temperature range up to 400 °C. As can be seen, KAN shows three endothermic peaks. The first broad one corresponds to the water loss, the rest ones can be due to decomposition phenomena [27]. Several peaks appear on the Na-CAS DSC curve: the first endothermic one at 71 °C corresponds to the evaporation of water molecules. Exothermic processes occurring at the temperature 189; 203 and 216 °C can be caused, most probably, due to the removal of the sulfuric acid, formed as a result of interaction between two closely located sulphate groups and their hydration [12]. Endothermic peak at 253.1 °C very likely corresponds to the thermal destruction of sulphate groups meanwhile it shifts to 238.2 °C at the DSC curve for complex because of these groups participation in the ionic interaction with AB. The shift of 216.0 °C peak to 228.5 °C has the same reason. The appearance of 283.4 °C peak in the DSC profile for the complex provided it’s absence for the components confirms their interaction as well. DTG curve for complex has a peak of maximum decomposition rate at 228.5 °C in the comparison with 216.0 °C peak for Na-CAS and 256.4 °C – for KAN.

Taking into account the complex formation between Na-CAS and KAN, the preservation of AB activity has become the subject of the study. Herein, we propose and detail a simple drug delivery design approach,
exploiting the complexation and adsorption of the complex on the AC, to achieve a sustained and complete release of KAN from GIT to blood. Na-CAS – KAN complexes themselves and immobilised on the AC were tested on the archival strain of *M. tuberculosis*. As a result, it was established that the KAN complex exhibits twice as many activities of the standard (injection) KAN form (with a minimum inhibiting concentration of 30 μg/cm³), while the active substance concentration was only 15 μg/cm³ (table 7). Figure 12 demonstrates the visual similar effect of tableted form of Na-CAS – KAN complex immobilised on the AC and injection form on the bacterial growth.

*Fig. 11.* DSC (*a*) and DTG (*b*) curves for Na-CAS (*1*), KAN (*2*), Na-CAS – KAN complex (*3*)
Table 7

Table 7

<table>
<thead>
<tr>
<th>Antibiotic concentration per 1 cm³ of nutrient medium, μg/cm³</th>
<th>Standard KAN injection form</th>
<th>KAN in a complex with Na-CAS</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>No growth</td>
<td>No growth</td>
</tr>
<tr>
<td>30</td>
<td>»</td>
<td>»</td>
</tr>
<tr>
<td>15</td>
<td>Moderate growth</td>
<td>»</td>
</tr>
</tbody>
</table>

It should be stressed that there is a fresh opinion [28] on the AC may prevent negative effects of AB on the microbiome. AC could solve this problem by allowing antibiotics to kill pathogenic microbes while sparing those that are important to gut health. It does not impact on the amount of drug entering the bloodstream. But it releases slowly the absorbed drug at the large intestine.

An in vivo study of the effectiveness of the combined dosage form of an AB and AC showed that the concentration of the antibiotic entering the blood from the tablet dosage form corresponds to its concentration entering the blood from the injection form (fig. 13). Figure 13 confirms almost equal bioavailability of injection and new dosage form of KAN. One can see that an adequate plasma level of KAN is achieved in both cases, when using a similar dose orally or intramuscularly. So, Na-CAS – KAN complex immobilised on the AC is a good carrier for KAN delivery. The concentration of KAN in the blood was determined by agar diffusion test with respect to Bacillus subtilis. A distinctive feature of the tablet form is the prolongation of action.

The present study enables us to suggest the explanation for the role of polymer in the creation of bioactive oral dosage form of KAN. Oral usage drugs, before they enter the bloodstream and target the bacterial RNA, must penetrate the numerous biological membranes of epithelial and endothelial cells of the GIT and then the bacterial membranes [29]. Molecular properties that are important for the penetration include molecular weight, size, conformation, degree of ionisation, polar surface area, non-polar surface area, lipophilicity and number of H-bonding acceptors and donors [30].

It is a matter of general experience that hydrophilic ionised drugs transport across biological membranes is poor. They have low oral bioavailability. For KAN it is reported about the increase of permeability by the ion-pair transport. But this concept still not extensively used because of the difficult search for specific lipophilic counter ions with a convenient physiological compatibility. It has been shown that at a physiological pH 7.4, KAN molecules are dimerised [8].

Dimerisation provides the formation of hydrogen bonds between hydroxyl groups of AB molecules. In this case, the amino groups of AB in the dimer remain free, which does not allow such an ionised aggregate to
penetrate through the lipid membranes (see fig. 9 and 10). Thus, the ionised AB dimer cannot penetrate through the phospholipid bilayer of the membranes on the one hand, and, on the other hand, it does not have enough free hydroxyl groups for binding with the transporter protein molecule, which could ensure AB transfer by the mechanism of active diffusion through transmembrane protein channels with its subsequent release into the bloodstream from the opposite side of the membrane [31]. Complexion of KAN with Na-CAS increases the lipophilicity of the drug that can change its behaviour in GIT.

In view of the importance of polar and non-polar area distribution on the surface of the drugs influence on their GI absorption, the molecular electrostatic potential (MEP) surfaces were calculated using the B3LYP/6-31G(d, p) basis set [32]. Figures 14–17 illustrate the distribution of the electrostatic potential (EP) values at the surface for KAN monomer, dimer, CAS (cellobiose fragment) and their complex. They are shown by different colours. EP increases in the order red < orange < yellow < green < blue. The negative regions of MEP, which are related the ability interact with electrophilic reagents, are seen as red and yellow colours, the positive regions that are related to the ability to interact with nucleophilic reagents, are seen as blue colour.

As can be seen from the diagram (see fig. 14), the regions with the largest negative charge are located near to the lone electron pairs of the nitrogen atoms of the amino groups. And oppositely the regions with the largest positive charge are located near to the hydrogens of hydroxyl groups.

There is no principle changing of MEP diagram in the case of KAN A dimer (see fig. 15) but it has more extended regions of intermediary potential.

As can be seen from fig. 17, there are two wide area of the negative EP that looks like an entirely red blob, and the positive EP area as an entirely blue colour blob on the complex KAN – CAS surface. This means a great electronegativity difference and indicates the presence of ionic bonds in the complex along with hydrogen bonds. It may be assumed that this specific localisation of the surface EP will enhances the transporter-dependent permeability of KAN complexed with Na-CAS. Taking into account that in the Na-CAS – KAN complex positively charged —NH\textsubscript{3}+ groups of AB molecule are associated with negatively charged —SO\textsubscript{4}2– groups of polymer (see fig. 10) one can conclude that at the periphery of the complex the number of charged —NH\textsubscript{3}+ groups decreases compared to the KAN dimer or ion pairs of the KAN cation with the sulphate anion. Thus, the binding of KAN molecules to the complex with Na-CAS makes the surface of the KAN molecules less polar to allow their transport through GIT membranes.
Further investigation will be necessary for the clarification of drug GIT absorption mechanism. For now, we have only the experimental results of the efficacy of ionic drug combination with the electrostatically complementary polyelectrolyte.

**Conclusion**

Aminoglycoside antituberculosis AB KAN has been shown to complex with Na-CAS. The composition of resulting hydrophobic product that is generated at pH 6.0 and 7.4 corresponds to molar ratio Na-CAS : KAN that is close to 1.0 : 1.0. Acidification of the solvent leading to the protonation of amino groups increases the amount of Na-CAS mole-links per mole of KAN in the complex till 2.9. The mixing order and the concentration of the components solutions allows to adjust particle size and distribution. Quantum-chemical calculations of the binding energies of KAN – CAS complex and KAN dimer shows that complex formation is an energetically preferable process compared to dimerisation. As a result of the complexation, the rearrangement of intra- and intermolecular bonds both in AB dimer and Na-CAS cellobiose unit takes place. According to the spectroscopic and thermal analysis data, ionised NH$_2$ groups, which impede the penetration of AB through the membranes of the epithelial layer, are connected in the complex with sulphate groups by ionic electrostatic interaction along with hydrogen bonds. Additionally, the great electronegativity difference over the complex surface may cause its special interaction with lipid membrane of the GIT walls, making it slightly loose. The hydroxyl groups released from intermolecular hydrogen bonds are able to bind to membrane proteins, which ensures the transport of AB through the membrane with its subsequent release into the bloodstream from the opposite side of the membrane. Both the hydroxyl and amino functional groups are involved in membrane interaction with the hydroxyl groups being deeper inserted into the membranes. After the disintegration of the complex, the carrier proteins ensure the transfer of AB monomer through the membrane protein channels.
A further mechanism of action of KAN is associated with irreversible inhibition of protein synthesis at the level of ribosomes in microorganisms sensitive to them. The monomeric form of AB is required for binding to the 16S site of rRNA. Using of the Na-CAS–KAN complex ensures the delivery of the monomer to the bloodstream, while when using the injection form, bacterium is directly attacked by the KAN dimer, with required preliminary destruction before ribosome attack. It is not yet clear as to what role the absolute configuration of KAN plays apart in this codon–anticodon interaction.

Unquestionably, the mechanism of increasing the activity of the complexes, as compared to individual antibiotics in injections, requires further research. But the results obtained allow us to recommend the polymer salt of KAN with Na-CAS as a new pharmaceutical form for oral administration to the clinical trials with the aim of current injection form replacement.

References


Received by editorial board 30.11.2020.