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A SHORT REVIEW ON REGULATION OF STABILITY OF AQUEOUS SUSPENSIONS OF CARBON NANOTUBES

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This review analyses the authors' recent and related works on electrokinetic properties and colloidal stability of aqueous suspension of multi-walled carbon nanotubes (CNTs) in the presence of surfactants and nanoparticles. Selected adsorptive properties of carbon nanotubes are also considered.

The applicability of classical theories of electrophoresis for description of electrophoretic mobility of carbon nanotubes is discussed. Examples on the ζ -potential of CNTs in aqueous suspensions as a function of pH and concentration of electrolytes (KCl, CaCl₂ and AlCl₃) are given. Additions of cetyltrimethylammonium bromide (CTAB) cationic surfactant below the critical micelle formation concentration (CMC) values give a reduction and then an overcharging the CNT surface because of accumulation of the surfactant ions in the Stern layer. At concentrations above CMC the substantial drop in ζ -potential is observed. It is due to the shift of the shear plane toward solution as a result of formation of hemi-micelles on the surface. An increase in the mass ratio of artificial mineral Laponite (Lap) in suspension (X) from 0 to 0.4 results in a monotonic decrease of the ζ -potential of CNT + Lap hybrids with reaching its plateau value ≈ -32 mV at $X \ge 0.4$ that corresponds to the ζ -potential of "pure" Lap platelets. This evidences the high surface coverage of nanotubes surface with Lap at $X \ge 0.4$.

The major methods to improve the dispersibility of carbon nanotubes, their colloidal stability and adsorptive properties are briefly discussed, namely: stabilization in mixtures of water and organic solvents; functionalization by chemical treatment; stabilization by additives of surfactants or polymers and hybridization by addition of supplementary stabilizing nanoparticles. CNTs can be significantly debundled in "good" solvents, such as 1-cyclohexyl-2-pyrrolidone, N,N-dimethylformamide, or N-methyl-2-pyrrolidone. It is demonstrated that the dispersibility of CNTs is a complex function of the type of surfactant, its concentration and the surfactant/CNTs ratio. The optimum concentration of CTAB to achieve homogeneous aqueous dispersion of carbon nanotubes was identified to be near the CMC. Additions of CTAB significantly modify the size distribution of CNTs and a sharp transition from small primary aggregates to large secondary aggregates at concentrations above CMC is observed.

We have shown also that CNT + Lap hybrids can be effectively used for removal of methylene blue dye from aqueous systems. The kinetics and mechanisms of adsorption are elucidated.

Keywords: carbon nanotubes, aqueous suspension, colloidal stability, zeta potential, surfactants, hybrids

INTRODUCTION

The composite materials doped with carbon nanotubes (CNTs) have demonstrated many attractive physico-chemical properties (electrical, magnetic, mechanical, thermal etc.) [1]. On the base of these materials different types of electrochemical biosensors, nanosensors for chemical and biological applications, piezoresistive strain sensors, electrical energy, memory storage devices, and super-capacitors, nanofluids for heat transfer devices, sorbents, materials for nanomedicine and promising biomedical applications, cancer therapy, biopharmaceutics

drug delivery and tissue engineering have been proposed.

However, the poor dispersibility of CNTs in most of the solvents and tendency of their aggregation present a serious obstacle for preparation of materials with good functionality. Preparation of CNT products are often realized through use of their stable aqueous suspensions [2]. Commonly, these suspensions possess low colloidal and sedimentation stability. The strong aggregation of CNTs is due to the high surface energy and aspect ratio, presence of strong van der Waals and hydrophobic attractive forces. Moreover, because of high hydrophobicity of CNT surface, the water is a poor solvent for preparation of stable aqueous CNT suspensions. The electrical, mechanical and thermal properties of composite materials can be significantly deteriorated by aggregation of CNTs. Therefore, the prevention aggregation and regulation of stability of suspensions of the CNTs is important scientific and technical task. In addition, the colloidal stability of the CNTs may be crucial with respect to the environmental impact of these materials.

In this review we briefly discuss our resent results on electrokinetic properties and colloidal stability of aqueous suspension of carbon nanotubes in the presence of surfactants and nanoparticles. Selected adsorptive properties of carbon nanotubes are also considered.

ELECTROPHORETIC MOBILITY AND ZETA POTENTIAL OF CARBON NANOTUBES

The electrophoretic mobility of charged particles dispersed in a solution is defined by interaction of their electrical double layers (EDL) with an external field. At relatively low fields (typically several or several dozen V/cm) the velocity of motion V is proportional to the electric field strength E, *i.e.*, $V = \mu E$, where μ is a electrophoretic mobility [3]. Electrophoresis is widely used in colloid and material science, medical and biological applications for characterization of the particles surface, separation of biocolloids, macromolecules, in electrodeposition/electrocoating technologies for fabricating of advanced materials, ion batteries and ceramic coatings [4].

In aqueous systems the electrophoretic mobility is greatly affected by the sort and concentration of electrolytes, and character of ions hydration. Effect of ions can be represented as the Hofmeister or lyotropic series which orders them with respect to macroscopic properties like surface tension, solubility of hydrocarbons, protein denaturation, *etc.* The Hofmeister series [5] reflect the orders of the ion effect on hydration of anions and cations:

	Increasing hydration of anions	
$CO_3^{2-} < F^- < CH_3COO^- < Cl^- < Br^- < NO_3^- < ClO_4^- < SCN^-$,		
	Increasing hydration of cations	
·	$Cs^+ < Rb^+ < K^+ < Na^+ < Li^+$	•
\uparrow	Surface tension	\downarrow
\downarrow	Solubility of hydrocarbons	\uparrow
\downarrow	Protein denaturation	↑.

Generally, anions show more specific and more pronounced effects than cations. This distinction can be explained by the fact that anions are commonly bigger and more polarisable than cations. The simple explanation of the Hofmeister series is based on the hydration. Generally, there is no unique Hofmeister series for the electrophoretic mobility, it can depend on the nature of the surface involved (sign of charge, polarity, hydrophilic or hydrophobic nature, *etc.*), type of electrolyte, pH, temperature and other conditions of the experiment [6]. Depending on parameters, the ordering of the ions within the Hofmeister series can be direct, indirect and partially altered.

The measured electrophoretic mobility μ may be converted into ζ -potential (zeta potential) using the Smoluchowski's approximation [3]

$$\zeta = \mu \eta / \varepsilon \varepsilon_o = \rho / (\varepsilon \varepsilon_o \kappa), \tag{1}$$

where ε and η are the dielectric constant and dynamic viscosity of the solution, respectively, ε_0 is the permittivity of the free space, ρ is the charge density at the slip plane in the EDL, $\kappa = 1/\lambda_D$ is the inverse Debye screening length. The ζ -potential corresponds to the potential drop between the shear plane (*i.e.* the plane where the abrupt change in the viscosity of liquid adjacent to the surface takes place) and the bulk of solution.

The validity of Smoluchowski's equation assumes that the ζ -potential is distributed uniformly, the local radii of curvature of the particles are much larger than the thickness of the electric double layer λ and the electric double layer remains in equilibrium. For heterogeneous charge distribution and for arbitrary shaped particles, the relevant corrections to the Smoluchowski's equation are necessary. The theory of electrophoretic mobility was mainly developed for cylindrical, spherical and disk-like particles [7]. However, the proposed models are rather complicated for mathematical analysis of the experimental systems.

In general, the electrophoretic mobility of CNT aggregates cannot be treated in the same manner as for spherical particles due to the large size and possible complexity of the aggregates. For thin double layers, the particles of any shape and size mostly behave very closely to that as expected for spheres of equivalent volume and charge. In this case, the classical Smoluchowski's equation can be applied for estimation of the "effective" ζ -potential (which is equal to the electrophoretic mobility multiplied by the ratio of the viscosity of the medium to its permittivity) [8], and it becomes even more appropriated for the large aggregates of CNTs. Such a procedure is accepted in the literature, and it can serve for comparison purposes, *i.e.* to follow the effects of different variables on the electrophoretic mobility and ζ -potential values. The Smoluchowski's equation was widely used for estimation of the effective ζ -potential of CNTs [9].



Fig. 1. ζ -potential (the electrophoretic mobility μ at right axes) of MWCNTs (0.01 % wt.) in water versus pH (*a*) and versus concentration of electrolyte, *C*, for KCl, CaCl₂ and AlCl₃(*b*) (adapted from [10])

The effects of pH and the validity of the Hofmeister series for suspensions of multiwalled carbon nanotubes (MWCNTs) in water has been checked [10]. Fig. 1 a shows an example of the pH dependence of ζ -potential (and electrophoretic mobility μ) of 0.1 % wt. MWCNTs in distilled water. The nanotubes were charged negatively in the whole interval of pH values (pH=2-12), and an increase in pH resulted in a noticeable increase of ζ -potential from \approx -19 mV at pH=2 to \approx -44 mV at pH=12. The step-like behavior with near-constant ζ -potential $(\zeta \approx -30 \text{ mV})$ in the interval of pH 4-8 was observed. Such behavior can be explained by the impact of pH on dissociation of weak acidic functional groups on the surface of nanotubes, presumably hydroxyl and carbonyl groups (the presence of such groups was justified by the IR spectroscopy data) [10]. These functional groups (e.g., -OH and -COOH) can arise on the surface of CNTs as the result of the applied synthesis and purification procedures. The step-like behavior was explained by presence of carbonyl functional groups and phenolic hydroxyls with pK ~ 2-3 and 10, respectively. Therefore, the observed dependence ζ (pH) evidently reflected the contribution of functional groups on the surface of MWCNTs, and changes in the synthesis and purification procedures can result in changes of the shape of ζ (pH).

Figure 2 *b* shows the effects of electrolytes KCl, CaCl₂ and AlCl₃ on the ζ -potential of MWCNTs (0.01 % wt) in water [10]. Addition of a KCl solution resulted in an initial increase of the ζ -potential and at high KCl concentrations (above 6.6·10⁻³ mol/dm³) a decrease of the ζ -potential was observed. Such a dependence with a maximum at low ionic strength values can be attributed to surface conductivity effects [11]. Polarization of the EDL can induce the dipole moment oriented oppositely to the external electric field. The dipole moment magnitude increases with increasing the electrolyte content.

This leads to an increasing retardation of the particle movement in the electric field. However, at high electrolyte concentrations, this effect is overcompensated by the EDL compression and this results in a decrease in mobility. Typically, appearance of unusual changes in the $\zeta(C)$ dependence corresponds to electrolyte concentrations at which contribution of the surface conductivity to the conductivity of dispersions is quite large. For bivalent (CaCl₂) and trivalent (AlCl₃) counter-ions increase in concentration, C, resulted in considerable decrease of the ζ -potential of MWCNTs, moreover with adding of AlCl₃ the overcharging of nanotubes above some critical concentration $(\approx 2 \times 10^{-4} \text{ M})$ was observed (Fig. 1 b). Such behavior is rather typical for hydrophobic colloidal particles [11].

REGULATION OF COLLOIDAL STABILITY

To improve the dispersibility of carbon nanotubes, their colloidal stability and adsorptive properties (increase of their surface activity and specific surface area) several approaches have been proposed:

• stabilization in mixtures of water and organic solvents [12];

• functionalization by chemical treatment of the MWCNT surface [13];

• stabilization by additives of surfactants or polymers de-bundling the aggregated MWCNTs [14, 15];

• hybridization by addition of supplementary stabilizing nanoparticles (formation of hybrid MWCNT-based particles) [16].

Mixed solvents. The dispersibility of carbon nanotubes can be noticeably improved in nonaqueous solvents or mixtures of water with suitable organic solvent [12]. MWCNTs can be significantly debundled in "good" solvents, such as 1-cyclohexyl-2-pyrrolidone (CHP), N,N-dimethylformamide (DMF), and N-methyl-2pyrrolidone (NMP) [12]. For example the aprotic CHP can disperse MWCNTs up to 3.5 mg/mL with very large populations of individual particles [12]. The dispersibility of carbon nanotubes in the given solvent is defined by the energetic costs of nanotube exfoliation. The "good" solvents have surface tension close to $\approx 40 \text{ mJ/m}^2$ [12], moreover, it was speculated that the dispersibility can be predicted from estimation of Hansen solubility parameter [17].

The dispersibility of MWCNTs in the mixture of CHP ("good") and water ("poor") solvents has been studied [18]. The CHP and water have significantly different total Hansen $\delta_T = 47.8 \text{ MPa}^{1/2}$ parameters: and $\delta_T = 20.5 \text{ MPa}^{1/2},$ respectively [17]. This difference is related to the differences in dispersion and hydrogen bonding components of the Hansen parameters for CHP and water. Nevertheless, the CHP is fully miscible with water below the lower consolute temperature $T \approx 320$ K [19].

Fig. 2 shows examples of photos of nanotubes suspensions (for mass concentration of nanotubes $C_n = 0.1$ % wt.) at two small and large mass fraction of water, x_w , in mixed solvent ($=m_w/(m_w+m_{CHP})$, where m_w and m_{CHP} are the masses of water and CHP, respectively) [18].



Fig. 2. Examples of optical microscopy images of 0.1 % wt. MWCNT suspensions at different mass fractions of water, x_m , in mixed water + CHP solvents: $x_w = 0.15$ (*a*) and $x_w = 0.75$ (*b*). Different colors correspond to the different clusters. The temperature was 294 K, and $x_w = m_w/(m_w + m_{CHP})$, where m_w and m_{CHP} are the masses of water and CHP, respectively (adapted from [18])

The nanotubes clusters were not detected visually in pure CHP, but even at a low concentration of water ($x_w = 0.05$), they became visible and their size increased with an increase of the water content. For example, the mean radius of two-dimensional projections of clusters was about 25 µm at $x_w = 0.05$ and it was about 850 µm at $x_w = 1.0$ (pure water). Observed clusters were ramified, their surface was tortuous and fractal. The estimated fractal dimensionality of the aggregate surface increased with x_w .

The changes in the dispersibility of MWCNTs greatly affected the electrical conductivity of suspensions, σ , in mixed water + CHP solvents. Fig. 2 shows σ versus the concentration of MWCNTs, C_n , at different mass fractions of water, x_m . [18]. Electrical conductivity significantly increased (by several

orders of magnitude) even at rather small concentrations of nanotubes (below 1 %). This clearly evidenced the presence of percolation transition related to the bridging of different clusters. In absence of water $(x_w = 0)$ the electrical conductivity was low and the percolation transition was rather smooth. The inset to Fig. 3 shows the value of σ_s determined at fixed $C_n = 1.0$ % wt. versus the value of x_w . The value of σ_s significantly increased with x_w , and it was estimated that in the poor solvent (water) it is approximately 25 times higher than that in the good solvent (CHP). Such behavior in water can be explained by easier constant ramified between large clusters and improvement of electrical contacts between different nanotubes because of the appearance of hydrophobic attraction.



Fig. 3. Electrical conductivity σ versus concentration of MWCNTs, C_n , in mixed water+CHP solvents at different mass fraction of water, x_m . Inset shows the value of electrical conductivity σ_s at fixed $C_n = 1.0$ % versus x_w . The temperature was 294 K (adapted from [18])

Functionalization of CNT surface. Colloid stability of aqueous suspensions can be improved by oxidation or functionalization of the MWCNTs [13]. The surface functional groups determine the wetting properties, surface charge density, and ζ -potential of nanotubes. For example, oxidation of CNTs by acids, hydrogen peroxide, ozone, etc. can lead to formation of carboxyl, carbonyl and hydroxyl groups on the CNTs surface [20]. Correlation between ζ potential, dispersion in solvents and oxygen content of multi-walled CNTs, modified by functionalization, have been discussed [21]. The observed electrokinetic behavior of the CNTs was similar to that typical for hydrophobic colloids. MWCNTs modified with application of stabilization or chemical functionalization techniques were used for production of filtering sheets (so called, bucky-papers), and absorbing additives [22] for purification of water [23].

Adding of surfactants. Effect of different surfactants on stabilization of aqueous dispersion of CNTs has been discussed in the literature in detail [14, 15, 24]. It has been demonstrated that the dispersibility of carbon nanotubes is a complex function of the type of surfactant (cetyltrimethylammonium bromide (CTAB), sodium dodecyl sulfate, Triton X-100, *etc.*), its concentration and the surfactant/CNTs ratio [25].

Cationic surfactant CTAB was widely used for stabilization of carbon nanotubes in aqueous suspensions [26–30]. The CTAB molecule has length of 2.33 nm, and its critical micelle concentration (CMC) in water is $\approx 1 \text{ mM}$ (0.0364 % wt).

Isotherm of adsorption of CTAB on the MWCNT surface and behavior of ζ -potential in the presence of CTAB were investigated in a number of works [27–30]. The aggregation stability was quantified using photometric dispersion analyzer (PDA) in a flow system by the PDA aggregation ratio R_{PDA} [31]. Fig. 2 presents examples of ζ -potential and PDA aggregation ratio, R_{PDA} , versus concentration of CTAB, C_s , in aqueous suspension of MWCNTs ($C_n = 0.003 \%$ wt.) [29, 30]. A change in the

 ζ -potential from negative to positive values with increasing the CTAB concentration, C_s , was observed. The position of the isoelectric point (IEP) was found at about $C_s(\text{IEP}) \approx 3.7 \cdot 10^{-5}$ M. It corresponds to 0.51 mg of CTAB/m² of the CNT surface ($x_s = m_s/m_n = 0.13$ g/g). The decrease of ζ -potential at larger concentrations of CTAB, above $C_s(\text{IEP})$, was observed. This reflected the formation of CTAB micelles and their adsorption onto the MWCNT surfaces. The adsorption of surfactant micelles shifts the slip plane toward solution and decreases the ζ -potential value.



Fig. 4. ζ -potential (*a*), and PDA aggregation ratio, R_{PDA} , (*b*) versus concentration of CTAB, C_s , for aqueous suspensions of MWCNT ($C_n = 0.003 \%$ wt.) Insert to (*a*) shows the sediment suspension photographs at 24 h after its preparation. The temperature was 298 K (adapted from [29, 30])

The similar overcharging behaviour was also observed in other works [27, 28]. For example, for 0.175 % wt. aqueous suspension of MWCNTs the position of the isoelectric point (IEP) was about $C_s \approx 1$ mM at pH 6.3 and the zeta potential reached the saturation level of about 50 mV for $C_s \ge 2$ mM [28].

Also the CTAB/MWCNTs ratio affects the character of adsorption and stabilization of nanotubes in aqueous suspensions [26]. For example, at $C_s = 0.27$ mM (below CMC) the ζ -potentials of nanotubes were negative ($\zeta = -46$ mV at $C_n = 10^{-3}$ % wt., and $\zeta = -25$ mV at $C_n = 10^{-2}$ % wt.), whereas at $C_s = 2.7$ mM (above CMC) they turned out to be positive ($\zeta = 66$ mV at $C_n = 10^{-3}$ % wt., and $\zeta = 40$ mV at $C_n = 10^{-2}$ % wt.). For low concentration of MWCNTs, $C_n = 10^{-3}$ % wt., the effective stabilization over a 15-day period was observed

for both $C_s = 0.27 \text{ mM}$ (below CMC) and $C_s = 2.7 \text{ mM}$ (above CMC). For high concentration of MWCNTs, $C_n = 2.5 \times 10^{-2} \%$, the stabilization was more efficient at $C_s = 0.27 \text{ mM}$ than that at $C_s = 2.7 \text{ mM}$. The observed phenomena were explained by differences in adsorption of individual CTAB molecules and worm-like CTAB micelles on the surface of nanotubes [26].

The dispersibility of carbon nanotubes in the presence of CTAB and aggregation stability of MWCNTs were studied in different works [26, 29, 30, 32]. The optimum concentration of CTAB to achieve homogeneous dispersibility of carbon nanotubes in aqueous suspensions was identified to be near the CMC [26, 32]. Remarkable that, in spite of the high negative charge of the carbon nanotubes, they were unstable in aqueous suspensions in the absence

of CTAB [29, 30]. At $C_s = 0$ M the nanotubes at $C_n = 0.003$ % formed large visible aggregates and the measured values of PDA ratio were rather large ($R \approx 8$) (Fig. 4 b). However, addition of CTAB in the concentration interval between $5 \cdot 10^{-6}$ and $3.7 \cdot 10^{-5}$ M resulted in stabilization of CNT suspension and the R values diminished to 1.5-3.0. It is unusual that stabilization was observed even in the vicinity of the isoelectric point, *i.e.* at $C_s(\text{IEP}) = 3.7 \cdot 10^{-5}$ M. At high CTAB concentrations, above the IEP (*e.g.*, at CMC, $C_s \approx 1$ mM), destabilisation and aggregation were again observed (Fig. 4). However, the character of this aggregation and aggregation at $C_s = 0$ M were significantly different.

Fig. 5 shows examples of size distribution functions V(d) in aqueous suspension of MWCNTs in the absence ($C_s = 0$ M) and presence ($C_s = 2.64$ mM) of CTAB [30]. The distribution functions were measured in gently stirred suspensions with applied hydrodynamic perturbations to prevent sedimentation of large aggregates. The distributions were monomodal and logarithmically symmetric without appearance of several fractions of nanotubes. In the absence of CTAB, the median size of aggregates was rather small, $d_{50} \approx 22.5 \,\mu\text{m}$, and this size was roughly twofold higher than the maximum length of individual carbon nanotubes (5–10 μ m). These primary aggregates in the absence of CTAB are stable and resistant to the applied shear.

Addition of CTAB significantly modified the size distribution and a rather sharp transition from small primary aggregates to large secondary aggregates at concentrations above $C_s^{\rm CMC}$ was observed [30]. For example, the median size was $d_{50} \approx 154 \,\mu\text{m}$ at $C_s \approx 2.64 \,\text{mM}$ (Fig. 5). A one-week aging of nanotube suspensions containing different amounts of CTAB (with C_s between 0.001 and 5.2 mM), or supplementary ultrasonic treatment in the measuring unit after formation of initial aggregates (for 10 sec $-2 \,\text{min}$) did not significantly affected the stability of the systems neither in diluted (below CMC) nor in concentrated (above CMC) systems.



Fig. 5. Size distribution functions F(d) in aqueous suspensions of MWCNTs in the absence and presence of CTAB. The aging time after ultrasonic pre-treatment time for 5 min was 2 h. The temperature was 298 K (adapted from [30])

Different mechanisms responsible for stabilization of MWCNT suspensions in the presence of CTAB were identified (Fig. 6). At small values of C_s (below IEP) the decorated adsorption of CTAB can be realized in the form of patches with different organization of CTAB molecules on the nanotube surface (Fig. 6 *a*):

• the positively-charged groups of CTAB $[CH_3(CH_2)_{15}(CH_3)_3N^+]$ can neutralize the negatively charged surface groups. This results

in the screening of nanotube charge and formation of hydrophobic patches around nanotubes.

• the long alkyl chain (length of $\sim 2 \text{ nm}$) can be attracted toward the "pure" carbon skeleton by hydrophobic forces. This charges the nanotubes positively and results in formation of hydrophilic shells around nanotubes.

The observed effect of CTAB induced stabilization of aqueous suspension of CNTs can

reflect competition between these two adsorption mechanisms. Note that structural repulsion forces can originate between the layers of adsorbed strongly hydrated CTAB molecules. In the vicinity of CMC the differences in adsorption of individual CTAB molecules and worm-like CTAB micelles on the CNT surface may be also important [27, 28]. At concentrations of CTAB above IEP more complex mechanisms of adsorption of individual CTAB molecules and micelles/semi-micelles on the hydrophobic surface might be realized [33]. The different types of encapsulation of nanotubes in a cylindrical surfactant micelle, hemi-micellar adsorption and random adsorption can be also considered (Fig. 6 b).



Fig. 6. Adsorption mechanisms of CTAB on the surface of carbon nanotubes at low (*a*) and large (*b*) concentration of CTAB (with permission from [29])

Hybridization with other nanoparticles. Hybridization with other nanoparticles has attracted a great attention during the last decades [34–36]. In this technique the MWCNTs are used as a base for preparation of hybrid particles with superior chemical activity, surface area and adsorptive properties as compared to intact MWCNTs. Different methods were proposed for preparation of hybrids: covalent and noncovalent interaction, *in situ* sol-gel process, chemical vapor deposition, electrochemical deposition, *etc.* [37].

Recently, a new type of hybrid based on MWCNTs covered by platelets of Laponite RD® (Lap) was described [16]. Lap represents a synthetic clay with the formula of Na_{0.7}[(Si₈Mg_{5.5}Li_{0.4})O₂₀(OH)₄], and it is composed of charged platelets with thickness of about 1 nm and average diameter of about 25–30 nm. The negative surface charge of Lap faces is counterbalanced by the positive charge of exchangeable sodium ions (for other details, see the recent review [38]). Adhesion of charged Lap nanoparticles on the surface of nanotubes improved their dispersion [10, 29, 39–44]. Addition of Lap resulted in efficient dispersion of MWCNT aggregates and separation of individual nanotubes (Fig. 7). In the presence of optimal concentration of Laponite ($x_l \approx 0.3$), all nanotube particles got stabilized, and a step-wise increase of sedimentation stability was observed [16]. Stabilization of nanotubes by Lap nanoparticles was explained by formation of the hydrophilic charged shells on the nanotube surface. The presence of Lap shells around the MWCNT surface was recently visualized using high resolution TEM technique [43].

Fig. 8 *a* illustrates the effect of Lap additions on the ζ -potential of MWCNT + Lap hybrids. An increase in $x_l = m_l/m_n$ ratio (here m_l and m_n are the masses of Lap and MWCNTs, respectively) from 0 to 0.4 resulted in a monotonic decrease of -40 to ζ -potential from -32 mV. The ζ -potential reached plateau value $\approx -32 \text{ mV}$ at $x_l \ge 0.4$ that corresponds to the ζ -potential of "pure" Lap platelets, $\zeta \approx -30 \text{ mV}$ [10], i.e. the MWCNT + Lap hybrids acquired the ζ -potential of the "pure" Lap platelets. Such behaviour evidenced the high surface coverage of nanotubes surface with Lap at $x_l \ge 0.4$.

The observed heterocoagulation and formation of MWCNT + Lap hybrids reflect the presence of attractive interactions between similarly charged species. Such attractive forces can exist between similarly and highly charged colloidal particles (see [45] for a review). The charge non-uniformity on the surface of MWCNTs (see *e.g.*, Fig. 6) may significantly contribute to the attractive interactions. The Lap disks can be effectively immobilized on the neutral parts of the nanotube surface due to strong attractive electrostatic image forces between the charged particles and neutral graphite-like surface. The examination of size distribution function of MWCNT + Lap hybrids revealed the presence of species with different sizes (Fig. 8 *b*). At small content of Lap $(x_l = 0.1)$, the relatively large aggregates of nonstabilized nanotubes (several µm), intermediate aggregates of partially stabilized nanotubes (300–1000 nm) and small species of stabilized hybrids (100–200 nm) were observed. Note that even at small content of Lap the size of particles was significantly smaller than that in absence of Lap (Fig. 5). At higher concentrations of Lap $(x_l = 0.6 \text{ and } x_l = 0.7)$ the content of the partially stabilized and stabilized hybrids was noticeably higher.



Fig. 7. Examples of optical microscopy images of 0.01 % MWCNT aqueous suspensions at different ratios Lap and MWCNT: $x_l = 0.0$ (a) and $x_l = 0.2$ (b). Here $x_l = m_l/m_n$, where m_l and m_n are the masses of Lap and MWCNTs, respectively. The temperature was 298 K (adapted from [16])



Fig. 8. ζ -potential (the electrophoretic mobility μ at right axes) of MWCNTs+Lap hybrids in water *versus* ratio Lap and MWCNT, x_l , (a), and size distribution functions F(d) at different values of x_l . Here $x_l = m_l/m_n$, where m_l and m_n are the masses of Lap and MWCNTs, respectively. The aging time after ultrasonic pre-treatment time for 5 min was 2 h, the temperature was 298 K (adapted from [10])

ADSORPTIVE PROPERTIES OF CNTS

Adsorptive properties of carbon nanotubes are determined by their high specific surface area and functional groups on the surface. Typically they have cylindrical shape and consist of graphene sheets wrapped around themselves. The multi-wall carbon nanotubes (MWCNTs) contain two or more concentric shells with intershell distance $d_{ss} = 0.34$ nm. For known outer diameter, d_o , and number of walls, n, of MWCNTs (of order 10–20 nm) the inner diameter d_i can be evaluated as $d_i = d_o - 2nd_{ss}$. The mass of one nanotube with length of L (of order 5–10 µm) can be evaluated as $m_n = \pi L d_0 n (1 - (n+1)d_{ss}/d_0)/S_c$ where $S_c ~(\approx 1315 \text{ m}^2/\text{g})$ is the surface area of the graphene sheet [46, 47]. Then the specific surface area can be evaluated as [10]:

$$S = s_n / m_n = S_c / [n(1 - (n+1)d_{ss} / d_0],$$
(2)

where $s_n = \pi L d_0$ is the surface of one nanotube.

Fig. 9 presents evaluated specific surface area, S, versus the external diameter of MWCNTs, d_0 , at different values of n. Note that increase of the number of walls, n, or the outer diameter, d_0 , results in decrease of the value of S.



Fig. 9. Calculated specific surface area of MWCNTs, S, (Eq. (2) versus its outer diameter, d_0 , at different numbers of walls, n. Inset shows example of TEM image of MWCNTs (adapted from [10])

Recent studies have shown the good potential of MWCNTs for adsorptive purification of water contaminated by toxic metals [48], organic matters [49], and synthetic dyes [50].

The specific surface area of CNTs can be experimentally estimated using the gas adsorption isotherm (e.g., N₂), or dye adsorption from the liquid phase. The methylene blue (MB) frequently used as a model dye in is investigations of adsorption on carbon-based substances. The kinetic and equilibrium data for adsorption of MB on MWCNTs at different temperatures have been analyzed [51]. Adsorption equilibrium was attained within $\approx 2 \text{ h}$ and the estimated thermodynamic parameters suggested that adsorption was spontaneous and endothermic. Adsorption was rapid in the initial stage and then followed by a slower process to reach the plateau value [52]. Based on the studies of adsorption of MB on SWCNTs, it was

concluded that adsorption is controlled by charge-transfer and hydrophobic interactions [53]. From studies of adsorption of MB on MWCNTs at different pH it was speculated that the adsorption mechanism may be due to π - π electron donor acceptor interactions and electrostatic attraction between positively charged dye ions and nanotubes [54].

The alkali-activated nanotubes demonstrated excellent adsorption capacity of MB [55]. The effects were attributed to the multiple adsorption interaction mechanisms on the functionalized surface (hydrogen bonding, $\pi-\pi$ and electrostatic interactions, *etc.*). The adsorption capacity of MB on modified nanotubes increased after heat treatment, and decreased after acid modification [56]. The method of removal of MB from aqueous solution using magnetic MWCNTs was proposed [57, 58]. Magnetic nanotubes can be easily separated in magnetic field after they have been loaded with dye. The MWCNT-based

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hybrid sorbents can be used for water purification from different inorganic, organic and biological compounds, and nanoparticles [34–36].

MWCNT + Lap hybrids can be also effectively used for removal of MB from aqueous systems. Adsorption of MB on MWCNT + Lap and dead-end membrane have been studied [42, 44]. The size and the shape of hybrid particles were determined by nanotubes, whereas the adsorption properties were determined by Lap particles attached to the surface of nanotubes. Due to the presence of Lap, the adsorbing capacity of hybrid was much higher as compared to the adsorbing capacity of pure nanotubes.



Fig. 10. Adsorption curve of MB on MWCNT + Lap hybrid ($x_l = 0.5$) represented as the relative quantity of the MB adsorbed by hybrid, *a*, *versus* the MB/MWCNTs ratio in suspension, C_d . Inset shows the saturation value a_m versus the ratio $x_l = m_l/m_n$, where m_l and m_n are the masses of Lap and MWCNTs, respectively. Duration of adsorption time was 3 h, the temperature was 298 K (adapted from [42])

Fig. 10 presents example of adsorption curve of MB on MWCNT + Lap hybrid ($x_l = 0.5$) represented as the relative quantity of the MB adsorbed by hybrid, a, versus the MB/MWCNTs ratio in suspension, C_d . For $C_d < 0.2$ (low MB/MWCNTs ratio) the purified solution was practically colourless, which is the evidence of complete adsorption of the MB. The adsorption reached its saturation level, a_m , at a certain critical MB/MWCNTs ratio ($C_d \ge 0.2$), and remained constant with the further increase of the MB concentration (Fig. 10). Inset to Fig. 10 presents the saturation adsorption of the MB, a_m , at different values of x_l . The maximal adsorption of the MB in suspensions of pure MWCNTs (at $x_l = 0$) was relatively low, $a_m = 0.053 \pm 0.007$ g MB/g MWCNTs. The increase of x_l resulted in increase of a_m and at $x_l = 0.2-1$, the direct proportionality between a_m and x_l was approximately observed, $a_m \approx 0.46 x_l$.

The MWCNT + Lap hybrids may be used either as a purifying additive or as a filtering layer if it is deposited on the surface of a supporting membrane. Due to relatively large size of hybrid particles, they can be easily separated from the purified solution by filtration or centrifugation. Figure 11 shows the hybrid membrane, obtained by filtration of hybrid MWCNT + Lap suspension ($x_l = 0.5$) through the nylon membrane with nominal pore size 0.2 µm (a), and ESEM image of the dynamic layer of MWCNT + Lap hybrids [42].

The dynamic homogeneous layers of hybrids have a micro-porous felt-like structure with a carcass composed of disaggregated MWCNTs. The average pore size of the layer, formed by hybrids, was rather small (< 1 µm). The estimated specific filtration resistance of hybrids increased from 2.0 10¹² m/kg to 7.1 10¹³ m/kg with increase of x_i from 0 to 0.5. For comparison, the specific filtration resistance of pure Lap was much high 1.5 10¹⁵ m/kg.

The kinetics of adsorption and parameters of equilibrium adsorption of MB on MWCNT + Lap hybrids have been also investigated in detail [44]. For elucidation of the mechanism of MB adsorption, the electrical conductivity and ζ -potential were measured [42, 44]. Three different stages in the kinetics of adsorption were observed: a fast initial stage I (t = 0-10 min), a

slower intermediate stage II (up to $t \approx 120$ min) and a long-lasting final stage III (up to $t \approx 24$ h). The presence of these stages was explained accounting for different types of interactions between MB and hybrids, as well as for the changes in the structure of aggregates and the long-range restructuring of Lap platelets on the surface of MWCNTs. The analysis of the adsorption data of MB by MWCNT + Lap hybrid particles evidenced in favor of the model with linear contacts between rigid Lap platelets and MWCNTs [44].



Fig. 11. Hybrid membrane, obtained by filtration of hybrid MWCNT + Lap suspension ($x_l = 0.5$) through the nylon membrane with nominal pore size 0.2 μ m (*a*), and ESEM image of the dynamic layer of MWCNT + Lap hybrids (adapted from [42, 59])

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Короткий огляд про регулювання стабільності водних суспензій вуглецевих нанотрубок

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У цьому огляді аналізуються нещодавні роботи авторів та споріднені роботи щодо електрокінетичних властивостей та колоїдної стійкості водних суспензій багатошарових вуглецевих нанотрубок (CNTs) у присутності поверхнево-активних речовин (ПАР) та наночастинок. Дано розгляд також адсорбційних властивостей вуглецевих нанотрубок.

Обговорюється застосування класичних теорій електрофорезу для опису електрофоретичної рухливості вуглецевих нанотрубок. Наведено приклади ζ-потенціалу CNTs у водних суспензіях як функції рН та концентрації електролітів (KCl, CaCl₂ та AlCl₃). Додавання катіонної поверхнево-активної речовини цетилтриметиламмонію броміду (CTAB) нижче значень критичної концентрації міцелоутворення (CMC) призводить до зменшення, а потім до перезарядження поверхні CNTs шляхом накопичення іонів CTAB в шарі Штерна. При концентраціях вище CMC спостерігається значне спадання ζ-потенціалу. Це пов'язано зі змішенням площини зсуву в бік розчину в результаті утворення на поверхні напівміцел. Збільшення масового співвідношення штучного мінералу Лапоніт (Lap) у суспензії (X) від 0 до 0.4 призводить до монотонного зниження ζ -потенціалу гібридів CNT + Lap із досягненням його плато \approx -32 мВ при $X \ge 0.4$, що відповідає ζ -потенціалу «чистих» пластинок Lap. Це свідчить про високе покриття поверхні нанотрубок за допомогою Lap при $X \ge 0.4$.

Коротко обговорюються основні методи поліпшення диспергованості вуглецевих нанотрубок, їхня колоїдна стійкість та адсорбційні властивості, а саме: стабілізація в сумішах води та органічних розчинників; функціоналізація хімічною обробкою; стабілізація добавками поверхнево-активної речовини або полімерів та гібридизація шляхом додавання додаткових стабілізуючих наночастинок. CNTs можуть бути значно відокремлені в «гарних» розчинниках, таких як 1-циклогексил-2-піролідон, N,N-диметил-формамід або N-метил-2-піролідон. Продемонстровано, що диспергованість CNTs є складною функцією типу ПАР, його концентрації та співвідношення ПАР/CNTs. Оптимальна концентрація CTAB для досягнення однорідної водної дисперсії вуглецевих нанотрубок була визначена поблизу СМС. Додавання CTAB значно модифікують розподіл за розмірами CNTs і спостерігається різкий перехід від малих первинних агрегатів до великих вторинних агрегатів при концентраціях вище СМС. При оптимальній концентрації Lap частинки нанотрубок добре стабілізуються і спостерігається ступінчасте підвищення стабільності осадження. Mu також показали, що гібриди CNTs + Lap можуть ефективно використовуватися для видалення барвника метиленового синього з водних систем. Висвітлено кінетику та механізми адсорбції.

Ключові слова: вуглецеві нанотрубки, водяна суспензія, колоїдна стабільність, поверхнево-активні речовини, гібриди

Краткий обзор по регулированию стабильности водных суспензий углеродных нанотрубок

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В этом обзоре анализируются недавние работы авторов и близкие работы по электрокинетических свойствам и коллоидной утойчивости водных суспензий многослойных углеродных нанотрубок (CNTs) в присутствии поверхностно-активные веществ (ПАВ) и наночастиц. Дано рассмотрение также адсорбционных свойств углеродных нанотрубок.

Обсуждается применение классических теорий электрофореза для описания электрофоретической подвижности углеродных нанотрубок. Приведены примеры ζ -потенциала CNTs в водных суспензиях как функция pH и концентрации электролитов (KCl, CaCl₂ и AlCl₃). Добавление катионной поверхностноактивного вещества цетилтриметиламмония бромида (CTAB) ниже значений критической концентрации мицеллообразования (CMC) приводит к уменьшению, а затем к перезарядке поверхности CNTs путем накопления ионов CTAB в слое Штерна. При концентрациях выше CMC наблюдается значительное падение ζ -потенциала. Это связано с смещением плоскости сдвига в сторону раствора в результате образования на поверхности полумицелл. Увеличение массового соотношения искусственного минерала Лапонит (Lap) в суспензии (X) от 0 до 0.4 приводит к монотонному снижению ζ -потенциала гибридов УНТ + Лап с достижением плато ≈ -32 мВ при $X \ge 0.4$, что соответствует ζ «чистых» пластинок Lap. Это

Коротко обсуждаются основные методы улучшения диспергованости углеродных нанотрубок, их коллоидная устойчивость и адсорбционные свойства, а именно: стабилизация в смесях воды и органических растворителей; функционализация химической обработкой; стабилизация добавками поверхностноактивного вещества или полимеров и гибридизация путем добавления дополнительных стабилизирующих наночастиц. CNTs могут быть значительно отделены в «хороших» растворителях, таких как 1-циклогексил-2-пирролидон, N, N-диметилформамид или N-метил-2-пирролидон. Продемонстрировано, что диспергируемые CNTs является сложной функцией типа ЮАР, его концентрации и соотношения ЮАР/СNTs. Оптимальная концентрация CTAB для достижения однородной водной дисперсии углеродных нанотрубок была определена вблизи СМС. Добавление СТАВ значительно модифицируют распределение по размерам CNTs и наблюдается резкий переход от малых первичных агрегатов к большим вторичным агрегатам при концентрациях выше СМС. При оптимальной концентрации Lap частицы нанотрубок хорошо стабилизируются и наблюдается ступенчатое повышение стабильности осаждения.

Мы также показали, что гибриды CNTs + Lap могут эффективно использоваться для удаления красителя метиленового синего из водных систем. Освещены кинетика и механизмы адсорбции.

Ключевые слова: углеродные нанотрубки, водные суспензии, коллоидная стойкость, зета-потенциал, поверхностно-активные вещества, гибриды

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