Magnetic Field Effects on Aqueous Anionic and Cationic Surfactant Solutions Part I: Water Evaporation

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Abstract
Static magnetic field (0.5 T) effects on water evaporation rate from anionic Sodium Dodecyl Sulfate (SDS) and cationic Dodecyl Trimethyl ammonium Bromide (DoTAB) 1 mM solutions were studied at room temperature and humidity for up to several hours. Keeping in mind possible practical application of the effects the experiments were intentionally carried out in a common laboratory environment and not in any sophisticated conditions. The evaporation of water from Magnetic Field (MF) treated and untreated samples were carried out simultaneously in the same environment. Although the quantitative differences in the evaporated amounts of water between MF treated and untreated samples changed from run to run, the qualitative MF effects were always reproducible. Therefore, it is believed that the observed changes are significant. It was found that the MF affects evaporation rate of water from solutions of both surfactants causing increase in the evaporated water amount in comparison to that of MF untreated sample. Prior to MF experiments first the water evaporation rate from the untreated surfactants solutions was studied. From the MF-untreated anionic surfactant solution water evaporated slower than from pure water, while from the cationic one water evaporated faster than from pure water. This difference was explained taking into account the properties of the polar (ionic) head of the surfactants, i.e. their size, ability to hydrogen bonding formation with water molecules, and the reduction of water surface tension. The MF treatment caused an increase in the evaporated water amount from both surfactants. However, a greater effect was observed for cationic DoTAB. Because the hydrocarbon tail in both surfactants is the same (C12) the observed differences were assigned to the differences in their ionic heads. Gibbs adsorption equation and Lorentz force in the gradient MF were applied to explain the differences.

Keywords: Magnetic field effects, Anionic and cationic surfactant, Water evaporation

Introduction
Investigation of Magnetic Field (MF) effects on properties of water and aqueous solutions are still of interest although they have been studied for at least 50 years. Hundreds of papers have been published where magnetic fields effects and application of MF in industry, agriculture, medicine, and others are described. Nevertheless, some of the results are debatable or even incompatible. Initially MF studies were focused to eliminate the hard scale formation at elevated temperatures in industrial pipes or house heating installations. If MF would successfully protect against deposition of the carbonates this could be beneficial elimination of chemicals used for water softening which are expensive and harmful for the environment. Later studies of MF effects in many systems and applications were carried out. Generally, using the classical magnetic field theory it is hardly to explain the observed effects which, however, often are well documented and statistically validated. The latest theories claim that to obtain an MF effect more important is the field gradient than its strength [1-3]. Also, the non-classical theory of nucleation mechanism and formation of dynamically ordered, so called liquid like oxyanion polymers, are used to explain the magnetic field action [4,5].

However, to our knowledge only in few papers the investigations of MF effects in different systems where a surfactant was present are reported [6-10]. However, no paper describes any MF effects on pure surfactant solutions. On the other hand, surfactants are present in the surface and waste waters, soil, and many industrial waters, sewage treatment plants, laundry, etc. [11,12]. Therefore, it seemed us interesting to carry out study in a natural room environment to learn whether some MF effects will appear in pure anionic or cationic surfactant aqueous solutions, and if so, whether the effects are reproducible qualitatively and/or quantitatively. In this paper first the MF effects on the rate of water evaporation from a surfactant solution was studied. The enhanced water evaporation from pure water was already reported in some papers [13-20].

For this purpose static MF (max. 0.5 T) originating from ring Nd magnets (MP 86 x 58 x 35 mm) was applied in which the sample of 10 cm diameter was placed ca.1.5 m apart from that with the MF and the samples we separated by a wooden board.

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weighed with the accuracy 0.1 mg, and the evaporated amount of water was calculated. For the reference purposes evaporation of water from the surfactant solution and pure Milli-Q water without MF presence were investigated too. In the previous paper similar studies have been conducted on MF effects on evaporation rate of pure water and water surface tension [21]. These are preliminary studies and depending on the obtained results further systematic investigations in aspect of possible practical applications will be continued. Application of MF for enhancing or hindering water evaporation, depending on the need of the process, would be beneficial.

Experimental

Materials

Sodium dodecyl sulfate >99.0% was purchased from Fluka and dodecyl trimethylammonium bromide approx. 99% was from Sigma. Both were used without further purification. For preparation of their 10 M aqueous solutions water from Milli-Q Plus system was used and the solutions were prepared a day before their first usage.

Methods

For the experiments of evaporation rate 65 mL (in some cases 50 mL, see Figure 8) samples of the surfactant solution or water from Milli-Q Plus system (resistivity 18.2 MΩ cm) were used. The magnetic field originated from a neodymium ring magnet 86 mm (outer diameter) x 58 (inner diameter) x 35 mm (height) directed with its north N pole upward. The solution or water surface in the vessel was exactly on the level of sample center. In the surfactant solution and pure water published earlier [21]. As it is seen in Figure 2 during the evaporation experiments the MF-treated and MF-untreated samples evaporated simultaneously at the room temperature (23 ± 1°C) and the relative humidity was 32-38%. The surface area of the sample on which the MF acted was 21.24 cm² and the circular surface radius was 26 mm. Therefore the meniscus curvature effect on the vapor pressure (Kelvin equation) did not play any role at this sample diameter. From Figures 1b and 1c it is seen that the strongest MF effect in vertical direction appears at the sample vessel walls and it vanishes toward the sample center. In the surface radial direction it decreases from 0.35 to 0.05 T. After every 30 min the closed samples were weighed and then their location together with the magnet was replaced. To avoid any possible influence of air fluctuations in the room the samples were placed in plastic tubes (Figure 2). The samples were weighed using a high precision balance Sartorius with an accuracy of 0.1 mg. The amounts of evaporated water were calculated by subtracting from the initial weight of the MF-treated or MF-untreated sample its weight after given time of the evaporating experiment, respectively. Then the difference between these two amounts was calculated and plotted versus time.

Results and Discussion

**Anionic sodium dodecylsulfate, SDS, solution**

First to learn whether water evaporation rate is different from aqueous surfactant solutions than that from pure water experiments without MF were performed using 10 M SDS solution and pure water. In Figure 3 are shown the results where it can be seen that more water evaporates from pure water than from 10 M SDS solution (the negative differences in mg). However, the differences obtained in these three separate experiments differ between themselves. The room temperature was the same 22°C and the relative humidity did not differ much. The plateau on the curve indicates that the rate of water evaporation is the same from MF treated and untreated samples and an extremum show the greatest difference in the evaporation rate. The difference in the evaporated amounts can be also presented as the relative percentage taking the evaporated amount from the pure water (or MF-untreated sample) as the reference 100%. In Figure 3 are shown these relative negative percentages calculated after 150 min of the experiment duration. Thus, the reduced evaporation of water from the SDS solution amounts to 2.0%, 10.8% and 14.3%, respectively which gives mean reduction percentage 9%. In experiment 1 (Figure 3, curve 1) the
2.0% relative decrease after 150 min results from 511.2 mg and 521.4 mg evaporated water from the SDS solution and pure water. However, after 30 min of the MF treatment in this experiment the percentage reduction amounted to 18.7% (Figure 3). It is because during this time 94.1 mg and 115.8 mg of water evaporated from the SDS solution and pure water, respectively. Therefore, to better depict the changes in the next figures, presenting the differences in amounts of evaporated water in mg, also the relative percentage values are given for 150 min of the experiment duration. As can be seen in Figure 3 in next two experiments much larger difference in the evaporated amounts of water has been obtained. Hence, the mean relative smaller amount of the evaporated water from the SDS solution after 2h is 9.0%. These results show that even without MF presence it is difficult to reproduce exactly the evaporation rate of water in a typical room environment using the same experimental setup, the surfactant lot and water. Nevertheless, an important finding is that water evaporates faster from pure water than 10^{-3} M SDS solution. However, one would expect an opposite relationship, i.e. faster and enhanced evaporation of water from the SDS solution whose surface tension is lower than pure water, 49.01 ± 0.26 mN/m (10^{-3} M SDS) and 72.30 ± 0.22 mN/m, respectively. This indicates that the cohesion forces between water molecules are stronger on the surface in pure water (2 x 72.3 = 145.6 mN/m) than in the SDS solution. Despite that 10^{-3} M SDS solution concentration is less than its critical micelle concentration, CMC=8.2 x10^{-3} M at 25 °C, there is already significant amount of SDS molecules adsorbed on this solution surface (compare the surface tensions). The molecules are oriented with their hydrocarbon chains toward the air and the ionic heads are located between the water molecules. The -SO4 head of SDS possesses oxygen atoms which interact with water molecules by hydrogen bonds. They are much stonger than the London dispersion and Keesom dipole-dipole forces, i.e. the strength of hydrogen bonds in water is ca. 20 kJ/mol while that of London and Keesom is 0.4-4 kJ/mol [22]. Van Oss and Constanzo [23] reported surface tension for SDS molecules immersed in water to be 23.8 mN/m for the hydrocarbon tail and 34.6 mN/m for the electron-donor parameter -r due to the presence of -SO4 head responsible for the hydrogen bonds formation. The average distance between the -SO4 groups in water was evaluated to be 0.907 nm. While between two alkyl chains the attraction amounts to 0.21 mJ/m² so strong repulsion of the electrostatic and polar nature exists between the sulfate heads. Therefore, they have to diverge with at least one -CH2 to which -SO4 is attached. From the surface tension values of SDS solution cited above it results that the -SO4 interactions are responsible for SDS surface tension to a great degree.

\[
\Gamma_2^{(1)} = -\frac{a_2}{RT} \left( \frac{\partial \gamma}{\partial a_2} \right)_A
\]

Where:
- \(\Gamma_2^{(1)}\) is the surface excess concentration of component 2 (surfactant) relative to its concentration in the bulk solution at zero excess concentration of main component 1(water).
- \(a_2\) is the surfactant activity in the bulk solution (concentration in the case of diluted solutions).
- \(\gamma\) is the surface tension of solution.
- \(A\) is the surface area (should be constant).

Therefore, the reason of increased water evaporation from MF treated solution might be due to weakening of Van der Waals interactions and hydrogen bonds in water intra-clusters [24,25] and formation of hydrogen bonds of water with the oxygen atoms from -SO3 groups. The average increase in the evaporated water after 150 min MF treatment amounts to 4.5% but it changes between 1.0% and 8.5% depending on the experiment run (Figure 4). To better depict the MF effects and compare water evaporation from the SDS solution and from pure water the differences are plotted in Figure 5 where curve 1 is that one from Figure 4 (these values are about the mean ones) and the curves 2 and 3 were calculated using the values obtained for pure water on the same day as those of curve 1. As can be seen in Figure 5 during 1h less water evaporated from the MF-ununtreated 10^{-3} M SDS solution than from pure water (1.7%, curve 2). From the MF-treated SDS solution more water evaporated from this solution than the MF-ununtreated one (1.6%, curve 1). After timespan 150 min the differences increased to 4.8% (curves 2) and 4.2% (curves 1), respectively. However, within 2h the evaporation rates of water from MF-treated SDS and pure water appeared to be practically the same and decreased by 0.8% only after 150 min (curve 3).

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Cationic dodecyltrimethylammonium bromide, DoTAB, solution

Analogous experiments to those with the anionic SDS surfactant were carried out with 10^-3 M cationic dodecyltrimethyl ammonium bromide, DoTAB, solution. Similarly as in the case of SDS solution first it was interesting to compare water evaporation rate from 10^-2 M DoTAB aqueous solution and pure water without MF presence. Two series of experiments have been carried out and the results are plotted in Figure 6. While from the anionic SDS solution water evaporated faster from pure water samples, so in the case of cationic DoTAB solution water evaporated faster from its solution than from pure water. In other words, at the same experiment duration more water evaporated from MF-untreated DoTAB sample than pure water, hence the differences in Figure 6 are positive. Also the percentage changes of evaporated water amounts are calculated for 150 min experiment duration and also at the end of particular experiment. The mean value from 7 experiments amounts 5.1% of the increased amount of evaporated water from DoTAB while in the case of SDS solution 9% less of water evaporated from the solution than from pure water (Figure 3). In the room environment at a slightly changing humidity and temperature obtained differences vary from run to run of the experiment but no doubt each time the evaporated amount of water from the solution is larger than from pure water.

Next the DoTAB sample was MF treated during water evaporation and simultaneously water from a reference untreated sample evaporated too (Figure 2). The results of differences in the amounts of evaporated water from MF treated and untreated solutions are presented in Figure 7 for four individual experiments. It can be seen in the figure that the MF causes increase in the evaporated amount of water in comparison to the MF-untreated solution. The relative percentage from 150 min experiment duration lies between 1.9-9.3%, giving mean value 4.6% which is practically the same as the mean value for SDS MF-treated solutions (4.5%, Figure 4). Similarly as in the case of SDS solution the MF field direction (north or south pole upward) does not make any visible difference. Also it looks that the small changes in relative humidity (32% and 36%) do not influence significantly the evaporation. The surface tension of 10^-2 M DoTAB solution after 60 min MF treatment decreased by 11.4 mN/m from 60.7 mN/m to 49.3 mN/m, i.e. more than in SDS solution, by 5.7 mN/m (to be published in the Part II of the paper). Taking again into account the Gibbs’ adsorption equation (Equation 1) for binary solutions a decreasing surface tension with the increasing bulk concentration (activity) of a surfactant means an increase in the surface excess concentration of this surfactant \( \Gamma_1 \). Therefore, it can be concluded that MF causes an increased adsorption of these two surfactant molecules on the surface, especially that of cationic DoTAB. In other words, it can be concluded that MF affects the structure of surfactants surface layer in a similar way as it occurs during the increasing bulk concentration of surfactant.

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Comparison of the MF effects on the surfactant solutions

To compare the observed MF effects on the two kinds of surfactants, first the differences in their ionic heads should be discussed. This is because both surfactants have the same hydrocarbon chain length, C12, therefore the different behaviour can be ascribed to drastic differences in the head group properties. They possess completely different ionic heads, i.e.

\[ \text{OSO}_3 \text{Na}^+ \] and \[ \text{-N}^+ \text{(CH}_3\text{)}_3\text{Br}^- \].

First of all, the size of these groups is different which is 0.17 nm\(^3\) for SDS and 0.54 nm\(^3\) for the hard-core area of DoTAB. These areas were “estimated from knowledge of the bond lengths, bond angles and atomic volumes using a molecular model of the headgroup” [26]. Also the distance from the hydrophobic core surface to the centre of the counterion location is 0.545 nm and 0.345 nm, respectively [26]. Therefore the Na\(^+\) counterions are at a larger distance from the \(-\text{O-SO}_3^-\) headgroup than Br\(^-\) from the \(-\text{N}^+ \text{(CH}_3\text{)}_3\text{Br}^-\). The Critical Micelle Concentration (CMC) of these surfactants at 25°C amounts to 8.2 mM and 11.0 mM, respectively. Moreover, the SDS molecule’s \(-\text{O-SO}_3^-\) group can form hydrogen bonds with water molecule oxygen atom but this is not the case for DoTAB whose head is more hydrophobic because of the presence of three methyl -CH\(_3\) groups and only a weak N–H hydrogen bond the \(-\text{N}^+ \text{(CH}_3\text{)}_3\text{Br}^-\) group can form [27].

The determined surface tension of 10\(^1\) M DoTAB is 60.7 mN/m while that of SDS was 49.0 mN/m (will be publish in Part II of the paper). Because the DoTAB head group is much larger than the SDS therefore it can be expected that the same surface area occupies less DoTAB head groups -\(-\text{N}^+ \text{(CH}_3\text{)}_3\) than \(-\text{O-SO}_3^-\) of SDS. Moreover, the Br\(^-\) counterions are located closer to the head [26] and hence at the solution the interactions between water molecules and the head group are much weaker than those at the SDS solution surface. Therefore this cationic surfactant can also reduce some of the hydrogen water-water molecules bonding. In consequence evaporation of water from the 10\(^1\) M DoTAB solution is easier than from pure water (Figure 8), contrary to the SDS solution (Figure 4).

The surface activity of surfactants can be described by Sprow and Prausnitz equation [20].

\[ \gamma_{LV} = \gamma_s + \frac{RT}{omega} \ln \frac{a_s^t}{a_s^b} \]  
(2)

Where:

- \( \gamma_{LV} \) - is the surface tension of aqueous solution of surfactant.
- \( \gamma_s \) - is the surface tension (surface free energy) of surfactant.
- \( omega \) - is the molar area of surfactant at the water–air interface.
- \( a_s^t \) and \( a_s^b \) - are the activity of surfactant in the surface layer and bulk phase, respectively.

Analogical equation can be written for water molecules in the surface layer.

\[ \gamma_{LV} = \gamma_w + \frac{RT}{omega} \ln \frac{a_w^t}{a_w^b} \]  
(3)

Where:

- \( \gamma_{LV} \) - is the surface tension of the surfactant solution.
- \( \gamma_w \) - is the surface tension of water.
- \( R \) - is the gas constant.
- \( T \) - is the absolute temperature.
- \( omega \) - is the molar area of water at the water–air interface.
- \( a_w^t \) and \( a_w^b \) - are the activity of water in the surface layer and bulk phase, respectively.

In Equations 2 and 3 the activity \( a \) of the components are defined in symmetrical system, i.e. \( a_s \rightarrow 1 \) if \( x_s \rightarrow 1 \). On the basis of the above equations Żdzenicka et al. [28] found for many surfactant solutions the maximum reduction of water surface tension to be ca. 41 mN/m, what indicated that the chains are oriented parallel toward the surface. This value is smaller than the values of our SDS and DoTAB solutions measured after MF treatment. The bigger decrease in surface tension of the cationic surfactant than the anionic after MF treatment can be ascribed to the presence of three \(-\text{CH}_3\) groups in the DoTAB head group whose surface tension is lower than \(-\text{CH}_2^-\) group present in the hydrocarbon tail [29,30].

Possible mechanisms of MF action

It is important to recognize possible mechanisms of the MF force action. Some approaches were discussed in the previous paper dealing with water evaporation from pure water surface [21]. Nakagawa et al. [13] and others [2-5,18] found that for water evaporation in the MF field more important is the field gradient \( B \partial dB/dx \) than the field itself. Moreover, oxygen present in the air can cause a susceptibility gradient in the direction normal to evaporating water surface which can enhance magnetic convection and in consequence a decrease in the water vapor density. This is because volume susceptibility \( \chi \) of oxygen is much greater than that of water and nitrogen. The bulk magnetic force was calculated by the authors [13] from Equation (4).

\[ \Delta F_m = \left( \frac{2a \mu_0}{\mu w} \right) B \frac{dB}{dx} \]  
(4)

Where:

- \( \Delta F_m \) - is the bulk magnetic force.
- \( \Delta \chi \) - is the susceptibility difference between the dry \( \chi_{dry} \) and wet air \( \chi_{wet} \) (which is near the water surface).
- \( \mu_0 \) - is the vacuum permeability (permeability of free space).
- \( B \partial dB/dx \) - is the magnetic field gradient.

In the magnetic field \( B = 8 \text{T} \) and the field gradient \( B \partial dB/dx = 320 \text{T/m} \) the force corresponded to as much as 17% of the gravitational force acting on the air which can be compared to the thermal convection effect that would be caused by 50K temperature increase from 293K [13]. In the case of our experiment at the water surface close to the inner magnet wall the gradient \( B \partial dB/dx \) amounted to 15 T/m but only 0.42 T/m at the magnet centre (Figures 1b and 1c) [21]. Hence the maximum force difference \( \Delta F_m \) (Equation 4) amounted to 0.089 N/kg which is only 0.91% of the gravitational force and its contribution in the magnetic convection is rather minimal. Then the Lorentz force acting on the ionic surfactant solutions can be analyzed.

\[ F = qE + q(\nu \times B) \]  
(5)

Where:

- \( F \) - is the Lorentz force.
- \( q \) - is the charge.
- \( E \) - is the electric field density.
- \( \nu \) - is the charge velocity.
- \( B \) - is the magnetic field.

In the case of electrolyte solution first term in Equation (5) equals zero because the electric field density \( E = 0 \). The second term expresses the magnetic force whose direction is perpendicular both to velocity \( \nu \) of

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the charge \( q \) and to magnetic field \( \mathbf{B} \). The force action depends on the charge and the magnitude of so called cross product of \( \mathbf{v} \times \mathbf{B} \), i.e. the velocity and flux density vectors, where the relative directions of these two vectors are taken into account. The magnitude of the force equals \( q \mathbf{v} \times \mathbf{B} \sin \phi \), where \( \phi \) is the angle between \( \mathbf{v} \) and \( \mathbf{B} \). If the angle \( \phi = 90^\circ \), i.e. \( \mathbf{v} \) is perpendicular to \( \mathbf{B} \), the particle trajectory is circular with a radius of \( r = m/\mu \). For angles \( \phi \) smaller than \( 90^\circ \) the charge moves along a helix with the axis parallel to the field lines. Obviously, if \( \phi = 0^\circ \) no action of magnetic force is observed. Silva et al. [31] taking \( v \equiv 0.992 \text{ m/s} \) (determined experimentally) and \( q = 3.2 \cdot 10^{-19} \text{ C} \) (divalent cation) in the field \( B = 1 \text{ T} \) calculated the Lorentz force to be \( 3.17 \cdot 10^{-19} \text{ N} \). Because the ion mass is equal to \( 10^{-21} \cdot 10^{-25} \text{ kg} \), the acceleration (\( F/m \)) can be as large as \( 10^{-10} \text{ m/s}^2 \) which can cause the ion polarization.

In our experiments the MF in the ring magnet changes radially from the top inner edge to its center from 0.347 T to 0.053 T, which occurs on the distance of 19 mm. Hence \( B/\Delta x \) on the sample surface level equals to 43.2 T/m and 7.9 T/m, respectively. Then the MF gradient changes from 14.96 T/m to 0.42 T/m, respectively [13]. Because of the field gradient and some mixing during the samples weighing every 30 min, the ions moves in the solution. Let us assume a \( v \) value 0.5 m/s and if some of the ions cross perpendicularly the field lines, the Lorentz force \( F = qvB \) for a monovalent ion amounts to \( (1.6 \cdot 10^{-18} \text{ C} \times 0.5 \text{ m/s} \times 0.347 \text{ T}) = 0.278 \cdot 10^{-20} \text{ N} \). Hence the acceleration force \( F/m \) acting on the dodecylsulfate ion \( \text{C}_{12}\text{O}_4\text{SO}_4^- \) (4.406 \cdot 10^{-6} \text{ kg/ion}) would be \( 6.3 \cdot 10^{-10} \text{ m/s}^2 \) and that acting on \( \text{C}_{12}\text{N}(-\text{CH}_3)_2^- \) (3.79 \cdot 10^{-6} \text{ kg/ion}) would amount to \( 7.3 \cdot 10^{-10} \text{ m/s}^2 \). The force at the magnet center is ca. 6.5 times lower than those at the edge. Although above calculations are very rough ones they show possible way to understand the observed MF effects.

Conclusions

These preliminary experiments showed for the first time that in a common laboratory environment preserving comparable conditions of temperature and humidity the static MF effects on the evaporation of water from both cationic and anionic surfactant solutions are present. Although they are not quantitatively reproducible, they are reproducible qualitatively. Nevertheless the amount of experiments is too small to be evaluated statistically the obtained changes can be assumed as significant because in each experiment simultaneously with MF- treated sample the reference MF-un-treated sample was present. Thus in most carried out experiments the MF affects evaporation of water from cationic and anionic surfactant solutions. Water from the MF-treated samples evaporates faster than that from the untreated ones thus leading to a larger evaporated amount of water during the same time. Larger MF effect observed in the experiments for the cationic than anionic surfactant solutions can be understood by taking into account the different properties of the two head groups, the anionic –O-SO_3^-Na^+ and cationic –N(CH_3)_2Br^- . The cationic group is over 3 times larger than the anionic and possesses three hydrophilic methyl groups –CH_3. Also the Na^+ counterions are located at a larger distance from the head group than Br^- . The sulfate group can form relatively strong hydrogen bonds with water molecules while the hydrogen bond with the ammonium group is weak, if ever. These differences reflect in the observed differences of the MF effects on these surfactant solutions. Generally, MF increases evaporated amount of water from both surfactant solutions and the mean relative values from several experiments are comparable for up to 150 min of their duration. The rough calculations indicate that MF can interact both perpendicularly to the liquid surface as the bulk magnetic force \( \Delta F_{\text{eff}} \), as well as horizontally as the Lorentz force. In the Part II of this paper the MF effects on the surface tension of these two surfactants solutions will be described. To our knowledge such MF effects on the surfactant solutions are published for the first time in the literature although a number of papers have been published on the MF effects on water evaporation from pure water. These results suggest that more systematic study at well-defined conditions are needed to better recognize these effects and the MF mechanisms causing their appearance. Such study will be conducted next. Potentially these MF effects may have a practical meaning in the processes where water is evaporated from surfactant solutions.

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