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Surfactant adsorption to soil components and soils

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ABSTRACT

Soils are complex and widely varying mixtures of organic matter and inorganic materials; adsorption of surfactants to soils is therefore related to the soil composition. We first discuss the properties of surfactants, including the critical micelle concentration (CMC) and surfactant adsorption on water/air interfaces, the latter gives an impression of surfactant adsorption to a hydrophobic surface and illustrates the importance of the CMC for the adsorption process. Then attention is paid to the most important types of soil particles: humic and fulvic acids, silica, metal oxides and layered aluminosilicates. Information is provided on their structure, surface properties and primary (proton) charge characteristics, which are all important for surfactant binding. Subsequently, the adsorption of different types of surfactants on these individual soil components is discussed in detail, based on mainly experimental results and considering the specific (chemical) and electrostatic interactions, with hydrophobic attraction as an important component of the specific interactions. Adsorption models that can describe the features semi-quantitatively are briefly discussed. In the last part of the paper some trends of surfactant adsorption on soils are briefly discussed together with some complications that may occur and finally the consequences of surfactant adsorption for soil colloidal stability and permeability are considered. When we seek to understand the fate of surfactants in soil and aqueous environments, the hydrophobicity and charge density of the soil or soil particles, must be considered together with the structure, hydrophobicity and charge of the surfactants, because these factors affect the adsorption. The pH and ionic strength are important parameters with respect to the charge density of the particles. As surfactant adsorption influences soil structure and permeability, insight in surfactant adsorption to soil particles is useful for good soil management.
Keywords: humic substances, silica, metal (hydr)oxides, kaolinite, montmorillonite, soil components

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References
1. Introduction

Surfactants have an amphiphilic character, which means that they are composed of a polar (hydrophilic) and an apolar (hydrophobic) part. A fundamental property of surfactants in solution is their ability to form micelles (colloidal-sized surfactant aggregates) at a characteristic surfactant concentration, the critical micelle concentration or CMC. At the CMC the surfactant monomers aggregate into micelles with the hydrophobic tail groups located in the core of the micelle and the hydrophilic head groups in the aqueous interface. In this way the unfavourable contacts of water with the apolar tails are minimized. Micellization gives surfactants their excellent detergency and solubilization properties; due to solubilization (partially) hydrophobic organic compounds dissolve in a micellar solution much better than in water alone. Another characteristic feature of surfactants is their tendency to adsorb at interfaces, mostly in an oriented fashion. The name surfactant is derived from this property; it is a contraction of surface-active agent. By surfactant adsorption the particle surface characteristics change; for instance, by adsorbing with their hydrophobic part to a hydrophobic surface the hydrophilic part protrudes in solution and makes the particle surface hydrophilic, which facilitates wetting and dispersion of the particles in aqueous solutions [1]. Due to their specific character surfactants are widely used chemicals. In household applications they are used in detergents, soaps and shampoos. In industrial applications surfactants are used for emulsification, dispersion, flocculation, wettability, flotation, foaming and so on [2]. In agricultural applications, surfactants are used for the formulation of hydrophobic agricultural chemicals to make these chemicals easily dispersible in water and as anticaking agents for chemical fertilizers [3]. In some cases surfactants are used in soil and groundwater remediation [4, 5]. The disadvantage of the application of large amounts of surfactants worldwide (>15 Mton/year) [6], is that their
Discharge is a cause of water pollution [7] and from the aqueous environment surfactants can easily bind to soil components. Dissolved surfactants are considered contaminants in aqueous environments and, in principle, they should be removed from wastewater before entering receiving waters. The ‘Handbook of Detergents, Part B, Environmental impact’ [8] considers the environmental aspects of most surfactant types. Recent information on the aquatic toxicity, biodegradability, and bioaccumulation, which are relevant for the assessment of surfactant ecotoxicity, can be found in [9-13]. Rebello et al. [10] conclude in their review: “…regarding surfactants as non-pollutants is a mistake. Visible manifestations of surfactant toxicity are available in the case of microbes, plants and animals”.

Soils are complex materials; they contain both inorganic particles and organic matter, see e.g.[14]. Roughly speaking, inorganic particles can be divided into primary minerals (physically weathered minerals from rocks), silicas, metal (hydr)oxides and layered aluminosilicates (clays) and within these three groups the particle sizes can vary largely. In general the primary minerals and silica particles make-up the sand and silt fraction, metal (hydr)oxides and aluminosilicates contain the smaller particle size fractions with large specific surface areas. All these soil particles contain in the normal pH range a surface charge, depending on the nature of the particles, which is largely or partly variable by changing the pH. This primary charge is due to the adsorption of protons and/or hydroxyl ions to functional groups (sites) present at the surface of the particles. Soil organic matter, which potentially acts as sorbent, originates from living organisms. It consists of structurally randomized macromolecular residues due to partial degradation, rearrangement and recombination of the original organic structures [15]. Mostly a distinction is made between non-humic and humic substances. Non-humic substances are chemically recognizable in biochemistry and are transitory in soil. Humic substances are not easily degradable and carry specific functional groups that can dissociate depending on the
structure of the group. As most groups are acidic the humic substances also carry a charge that is variable with pH.

The nature of the surface sites of the inorganic particles and the functional groups of the organic particles, determine the particle charge and chemistry and charge govern to a large extent the sorption of other components, including surfactants, to the particles. Important for surfactant binding are the precise sites present at the surface as they determine the chemical nature of the surface and the primary particle charge. The chemical nature determines the “specific” or “chemical” interactions with the polar and apolar parts of the surfactant molecules and the charges determine the electrostatic interaction, which can be either attractive or repulsive, depending on the charge signs of surfactant and particles. Together these interactions determine the mode of the surfactant adsorption to the particles (with their hydrophilic or their hydrophobic part directed to the surface). As far as we are aware, there is no comprehensive review on surfactant sorption to soils, most likely this is due to the complicated and diverse nature of the soils and the many different types of surfactants.

In the present paper first the properties of surfactants are considered in some detail, including surfactant adsorption on the air/water interface, which can illustrate the importance of the CMC for the adsorption process and gives an impression of surfactant adsorption to a hydrophobic surface. Surfactant–soil interactions are determined by the soil components; therefore attention is given to the different types of soil particles: humic substances, silica and metal oxides and aluminosilicates. Information is provided on their structure, surface properties and primary charge characteristics, which are all important for surfactant binding. Subsequently, the adsorption of different types of surfactants on these individual soil components is discussed in detail, based on mainly experimental results and considering the various specific (chemical) and electrostatic interactions. Adsorption models that can describe the features semi-quantitatively are explained and discussed. In the last
part of the paper some trends of surfactant adsorption on soils are discussed together with some complications that may occur and finally the consequences of surfactant adsorption for soil colloidal stability and permeability are considered.

2. Surfactants and their characteristics

2.1. Surfactant types

The chemistry of the polar and apolar part of surfactants can be quite different for different surfactants and based on the nature of the polar head group surfactants are classified into anionics, cationics, nonionics, and zwitterionics. The principal surfactant demand is composed of four types of surfactants: linear alkylbenzene sulfonates (LAS), fatty alcohol ethoxylates (AE), fatty alcohol ether sulfates (FES) and fatty alcohol sulfates (FAS). Both Western Europe and the US rely on the alcohol derivatives for 60-70% of the major surfactants but in the rest of the world it is 15-30%, globally LAS is the most used surfactant [16]. Next to synthetic surfactants also natural surfactants and biosurfactants exist. Some examples of synthetic and natural surfactants are depicted in Fig. 1.

Biosurfactants are surfactants that are produced extracellularly or as part of the cell membrane by bacteria, yeasts and fungi. Mulligan [17] and Bustamante et al. [18] give examples of biosurfactants and their structures and review the environmental applications for soil remediation and water treatment. Biosurfactant applications in the environmental industries are promising due to their biodegradability, low toxicity and effectiveness in enhancing biodegradation and solubilization of low solubility compounds.

2.2. Surfactant adsorption at water/air surfaces

The attraction of surfactants for aqueous solution/air surfaces is due to hydrophobic attraction. Hydrophobic attraction is the thermodynamically favorable tendency of the
surfactant hydrocarbon moiety to escape from the aqueous environment to an apolar environment (air). With adsorption the hydrophobic attraction is the difference between the Gibbs energy (energetic and entropic contributions) of the surfactant tails in aqueous solution and that in the adsorbed state. The adsorption of surfactants at liquid interfaces is generally studied by measuring the surface tension, \( \gamma \), and using the Gibbs equation to obtain the adsorption [19, 20].

\[
d\gamma = -s_a^{(1)} dT - \sum_{i=2}^{\infty} \Gamma_i^{(1)} d\mu_i
\]

where \( s_a^{(1)} \) is the relative surface entropy per unit area, \( T \) the absolute temperature, \( \Gamma_i^{(1)} \) is the relative adsorption of component \( i \) and \( \mu_i \) is the chemical potential of \( i \). The quantities \( s_a^{(1)} \) and \( \Gamma_i^{(1)} \) are conveniently determined by assuming that the solvent (component 1) adsorption is zero (\( \Gamma_1 = 0 \)). In the case of surfactant adsorption from ideally dilute aqueous solutions the Gibbs equation can be approximated as [21-23]:

\[
d\gamma = -pRT \Gamma_{\text{surface}} d\ln c_{\text{surface}}
\]

where \( p = 1 \) for nonionic surfactants (they are only affected by the presence of electrolytes when the electrolyte changes the solvent quality), but for ionic surfactants

\[
p = 1 + \frac{c_{\text{surfactant}}}{c_{\text{surfactant}} + c_{\text{salt}}}
\]

This notation of the Gibbs equation has used the equality \( d\mu_i = RT d\ln c_i \) for all components present and the implicit assumptions are made that the ion concentrations from the background electrolyte are constant, that \( T \) is constant, that the adsorption, \( \Gamma_{\text{surface}} \) is a simplified notation for \( \Gamma_i^{(1)} \) and that for ionic surfactants the surfactant co-ion adsorption can be neglected. The latter assumption holds best for highly charged surfaces, i.e. at sufficient surfactant ion adsorption. Schematically the effect of surfactants on the surface tension of aqueous solutions is shown in Fig. 2.
In Fig. 2a $\gamma(\ln c)$ curve is shown, firstly the surface tension progressively decreases with increasing surfactant concentration (region I). In region II the decrease becomes approximately linear and at the start of region III the critical micelle concentration or CMC is reached. Typically, micelles are clusters of 50–200 surfactant molecules. For surfactant concentrations somewhat above the CMC (region III) the monomer concentration stays about constant and so does the chemical potential of the monomeric surfactant. With the Gibbs equation the adsorption, $\Gamma$, can be calculated from the derivative of the $\gamma(\ln c)$ curve. A typical result is depicted in Fig. 2b. The adsorption first increases but in region II, where the slope of the $\gamma(\ln c)$ curve becomes constant (within experimental error), the adsorption reaches a plateau value. In general, the maximum adsorption depends on the effective size of the hydrated head group. In region III the monomer concentration remains constant.

Application of the Gibbs equation in region III is not possible because the monomer surfactant concentration stays equal to the CMC, but there is general consensus that the adsorption stays at the level reached at the CMC. For ionic surfactants the $\gamma(\ln c)$ curve depends on the ionic strength, see Fig. 3. For the calculation of the adsorption with the Gibbs equation the factor $p$ comes into play, the extreme situations are $p=2$ in the absence of salt and $p=1$ when the salt concentration is much larger than the surfactant concentration. For the intermediate situations $p$ has to be calculated and can gradually increase with increasing surfactant concentration. The maximum attainable adsorption, $\Gamma^m$, depends on the salt concentration and increases with increasing ionic strength. This is due to a decrease in electrostatic repulsion between the surfactant head groups, which reduces the effective head group size. The order of magnitude of adsorption of a univalent surfactant is 1-4 $\mu$mol/m$^2$ [24], which corresponds to about 0.1-0.4 C/m$^2$ and indicates that the adsorption occurs (due to hydrophobic attraction) even at a considerable electrostatic repulsion between the surfactant ions.
2.3. Micelles and their shape

The presence of micelles at surfactant concentrations above the CMC is a very important property of surfactants. Micelles are governed by energetic and geometric considerations. A comprehensive review on the models that describe micellization can be found in [25]. In a micelle the hydrophobic attraction between the hydrophobic moieties is just balanced by the repulsion between the hydrated head groups. The hydrophobic attraction is in this case the difference between the Gibbs energy of the surfactant tails in the aqueous solution and that in the core of the micelles. The shape of the micelles depends on the geometric characteristics of the surfactant molecules. In general, the concentration at which the CMC occurs depends on the surfactant structure and solution conditions. With respect to the surfactant structure the variation in the length of the hydrophobic tail of the surfactant is the most important factor and, in general, the CMC decreases as the hydrophobic character of the surfactant increases, see Fig. 2c. For nonionics also the head group size is important; in general the CMC and the surface tension at the CMC increases with the head group size, see Fig. 2d.

The solution conditions affect the CMC of nonionic surfactants when the solvent quality for the head groups is affected. For ionic surfactants the ionic strength and type of surfactant counterion are important. By way of illustration, Fig. 3 shows how the NaCl concentration affects the surface tension of an ionic surfactant: sodium dodecyl pyridinium chloride (C12PC). It is obvious that the salt concentration changes both the CMC and the slope of the curves. The presence of electrolytes in solution reduces the CMC by shielding the electrostatic repulsion between the ionic head groups. In soils, the CMC of ionic surfactants is likely to be reduced due to the salts present in the soil-solution.

The above illustrates that the CMC of surfactants in aqueous solution varies with surfactant structure and presence of electrolyte in the case of ionic surfactants; the presence of various organic compounds and the temperature also affect the CMC. Although the CMC of a surfactant may vary under different conditions, the CMC is a characteristic property of a
surfactant. Tabulated values of many surfactants (for ionic surfactants with specification of
the salt concentration or ionic strength) can be found in [24, 26, 27].

The shape of the micelles varies from surfactant to surfactant and for ionic surfactants
the salt concentration is also affecting the micelle shape. In general, the micelle shape
depends on the area of the apolar tail, $V_o/l_o$, where $V_o$ is the tail volume in the micelle and $l_o$
the extended tail length, and the equilibrium area, $a_e$, of the polar/ionic head group at the
micelle surface. The parameter $V_o$ varies with the number of hydrophobic groups, chain
unsaturation, chain branching and chain penetration by other compatible hydrophobic
groups, while $a_e$ is mainly governed by electrostatic interactions and head group hydration.
For ionic surfactants $a_e$ depends on the ionic strength and for large tails the tail packing in
the micelle is non-uniform and affects also the equilibrium head group area [28, 29]. The
ratio tail area over head group area, $V_o/a_e l_o$, is called the “critical packing parameter” or
CPP and the CPP value provides a good indication of the shape of the micelles [30-32]. In
aqueous media and $0<\text{CPP}<0.3$ the micelles are spherical (single chained surfactants with
large head group areas), for $0.3<\text{CPP}<0.5$ the micelles are cylindrical (single chained
surfactants with small head group areas) and for $0.5<\text{CPP}<1$ curved bilayers (lamellar) are
formed (double chained surfactant with large head group areas) and for CPP=1 flat bilayers
(double chained surfactant with small head group areas). For CPP>1 only inverted micelles
can exist. When the surfactant concentration is raised above the CMC, linear growth of
micelles is the dominating type of growth. Some knowledge about the shape of the micelles
is also important because in the adsorbed state surfactants already tend to form micellar type
aggregates before the CMC is reached.

3. Soil components

3.1. Humic substances
The soil organic fraction contains the non-humic substances (e.g., proteins and natural macromolecules) and the humic substances. The humic substances are divided into fulvic acid, humic acid and humin by their aqueous solubility; humic and fulvic acids are soluble in the normal pH range and humins are not. Fulvic and humic acids are distinguished by their solubility at low pH: fulvic acids are soluble in all pH, whereas humic acids precipitate at low pH. For the isolation and separation of the humic substances the procedure described by Swift et al. [33] is mostly used as the International Humic Substances Society recommends it.

Humic particles found in soil, sediment and water have an extreme structural complexity. In an overview Leenheer [34] discusses the successive approximations of model structures to molecular structures that have appeared in literature for dissolved humic particles. Present days insight is that the old “polyelectrolyte model” that represented the dissolved humic particles as randomly coiled macromolecules resulting from various chemical condensation and coupling reactions of smaller biomolecular constituents is no longer adequate. New structural models view dissolved humic particles as dynamic associations of relatively low molar mass components stabilized by hydrophobic interactions and hydrogen bonds [35] or as complex supramolecular assemblies of small molecules, oligomers, and polymers aggregated into supramolecular assemblies by non-covalent forces [36].

From physicochemical point of view the dynamic character of the aggregation [37], the presence of a diversity of functional groups that protrude in the aqueous phase [38] and the fact that the aggregates have an open structure [39] that allows the aqueous solution to partly permeate through the particles has to be taken into account. The functional groups can dissociate or associate with protons and this causes the primary charge of the particles that governs the electrostatic interactions. With physicochemical treatments humic particles should therefore be considered as amorphous, polydisperse, chemically complex and internally structured supramolecular particles that are held together by hydrophobic
attraction and hydrogen bonding and that are partly permeable by the aqueous solution. Through their structure and dissociation or association of the functional groups with protons, the particles have in aqueous solutions a polyelectrolyte nature that can be classified as soft-colloidal matter and both their charging and electrokinetic behavior [40] are different from that of rigid solid particles. Due to their ‘open structure’ humic and fulvic acid have very large sorption capacities per kg of material. The main differences between humic and fulvic acids are their functional group density, average molar mass and hydrophobicity: fulvic acids have a relatively low molar mass, a relatively high functional group density and a relatively low hydrophobicity. Humin particles are only poorly investigated, but based on the isolation procedure it may be expected that they are solid like (much less permeable than humic acid) with a relatively low functional group density, have a large particle mass and are hydrophobic in nature.

For a further physicochemical characterization of humic and fulvic acids the proton binding characteristics of humic and fulvic acids are relevant. The heterogeneity of the proton binding groups of humic substances is considerable with, in general, two main classes of groups each with a wide distribution [41, 42]. The low proton affinity groups are mostly referred to as ‘carboxylic’ groups (pK_H around 3-4) and the high proton affinity groups as ‘phenolic groups’ (pK_H around 8-9). For the pH conditions in natural waters humic and fulvic acid are negatively charged and pH and ionic strength affect the proton charge density [43], the hydrodynamic radius of the humic particles [44] and their electrokinetic behavior [40]. Fig. 4 gives a typical example of proton binding to a humic acid at four different salt concentrations [45].

Much attention has been paid to modeling proton (and heavy metal ion) binding to humic and fulvic acids and the literature is reviewed in [46-50]. In general, models should (i) take explicitly into account the heterogeneity of the functional groups of humic and fulvic acids, (ii) distinguish between electrostatic and ion-specific binding and (iii) account for
competition for binding among ions. Models that confirm to these conditions are based on intrinsic parameters that are independent of the environmental conditions (e.g., pH, ionic strength). The two most successful models that suit the above conditions are Tipping’s ion-binding Model VI/VII [51-53] and the NICA-Donnan model [38, 43, 54-56]. These models can be applied across a wide range of environmental conditions and be used to make predictions outside the calibration range. The heterogeneity of the humic substances is described in Model VI/VII with a wide discrete distribution of (intrinsic) proton affinity constants [47, 51] and in the NICA-Donnan model with a bi-modal continuous distribution [57, 58]. For the electrostatic interactions a simple Donnan model is used in the NICA-Donnan model [54] and recently also in Tipping’s model [52, 53]. The Donnan models describe the electrostatic interaction with only one average smeared-out electrostatic Donnan potential at given environmental conditions [59]. The Donnan model leads to calculated Donnan potentials that are highly non-Nernstian, see [54]. This means that the Donnan potential is non-linear with a change in pH and that its value is also strongly ionic strength dependent. By considering a great deal of the existing data up to 2003, Milne et al. [55, 56] have been able to establish two sets of parameters that can describe the proton and other ion binding to, respectively, generic-HA and generic-FA. These generic-HS compounds can be used as substitutes when no detailed information is present on a specific HS to calculate the ion binding for many ions.

3.2 Silica and metal (hydr)oxides

In many parts of the world, silica (SiO₂) is the major constituent of sand and silt. The fine silica particles (<2 μm) are classified as clay. Oxides of iron and aluminum are most prominent in soils of the tropics but are widespread elsewhere. Silica has a number of distinct crystalline forms in addition to amorphous forms. Amongst the various crystallographic forms of silica, α-quartz is the most abundant, as it is the most stable
crystalline silica phase. Thus, $\alpha$-quartz has been widely used as a model for the investigations of water/silica interface systems in environmental processes. Due to their coarse size, quartz particles have a low specific surface area and do not have colloidal properties, yet they exhibit adsorption. In contrast, amorphous silicas can be very fine and mostly have a large specific surface area. Also because their industrial uses, the adsorption properties of amorphous silicas (mostly synthetic) have attracted much attention.

The surface structure of silica is composed of siloxane sites (-Si-O-Si-) and silanol (-SiOH) sites. Siloxane sites are hydrophobic and silanol sites are hydrophilic; therefore, the higher the silanol content is, the larger is the hydrophilicity of the surface. The silanol groups can be subdivided into isolated (or single), geminal (-Si(OH)$_2$) and vicinal or bridged silanols. The vicinal OH groups are two nearest neighbors that are bound together by a hydrogen bond. The silanol sites are formed by hydroxylation of siloxane bridges. Based on dehydration studies, Zhuravlev [60] has shown that amorphous silica surfaces will contain predominantly single and geminal silanol groups and siloxane bridges, the relative content of each type of group depending on the temperature of dehydration and the subsequent rehydration conditions. Direct information on the distribution of silanol groups over isolated, geminal and vicinal groups can be obtained from NMR studies. Chuang and Maciel [61] have reported that the surface of amorphous silica powders have a roughly equal distribution of isolated and geminal silanols. Yang and Wang [62], who reviewed computational simulations, conclude that the normal siloxane bond is hard to be hydrolyzed because of the high reaction barrier, but siloxane surface structures with a large strain stress and surface defects are rather readily broken by water attack, especially with the aid of extra water molecules. The presence of silanols next to siloxane groups thus also follows from the simulation studies.

From hydration-dehydration and tritium and deuterium exchange studies it can be concluded that when an amorphous silica surface is hydroxylated to the maximum state the
silanol density is around 4.9 groups per nm$^2$ independent on the history of the sample [60]. This number is about two times smaller than the classical value of number of silanols of about 8 groups per nm$^2$ derived by Iler using the assumption that for each surface Si there is one OH group [63]. The difference can be explained by the fact that only half of the Si atoms are capable of holding OH groups [64]. Tamura et al. [65] give an overview of hydroxyl site densities of silicas and a range of metal oxides (Al, Fe, Mn, Ti, Sn, Zn, Mg).

From the collected densities it follows that the hydroxyl content of silica is substantially lower than that of the metal oxide surfaces. The high hydroxyl content of the metal oxides explains that these surfaces are more hydrophilic than the silica surface.

The sites responsible for the charging of the aqueous interface of silica and metal oxides can be derived from the mineral structure. For crystalline minerals the structure is well known through X-ray diffraction measurements and in combination with electron microscopy also the dominant crystal faces that make up the surface can be detected. Using this information in combination with Pauling’s bond valence concept [66] and its refinement [67-69] the protonation properties of the oxygen containing surface groups can be derived.

In the bond valence concept, the charge of the central cation is distributed over the surrounding O- ligands and the sum of bond valences around oxygen should be equal to the oxygen valence. Hiemstra et al. [70, 71] have used the bond valence concept in the multi-site complexation (MUSIC) model to describe the protonation of mineral surfaces. The model allows differentiating various types of surface groups in terms of formal charge and takes into account that most metal oxide surfaces have multiple types of surface oxygen groups that can react with protons. The pK$_H$ values of the groups can be predicted and the model has highlighted that the two pK$_H$ values of two successive protonation steps on one and the same O-ligand differ by about 10 pK-units ($\Delta$logK$_H \approx 10$). This implies that practically only one proton association or dissociation step per surface group can occur in the normal pH window. Other predictions are: (i) the inertness of double coordinated
surface groups on sesquioxides, which may result in non-charged perfect crystal faces for
gibbsite [72, 73] and hematite [73, 74], and (ii) the difference in proton affinity for the two
basic types of triply coordinated groups of goethite [71]. The MUSIC model may also
account for changes in temperature as shown by Machesky et al. [75]. It should further be
noted that the predicted $\Delta pK_H$ value of 10 provides evidence that the older description of the
proton charging of mineral surfaces based on two protonation steps for each surface group
($\equiv$SO$^{-}$ to $\equiv$SOH$^0$ to $\equiv$SOH$_2^+$; 2-pK model) is fundamentally wrong.

It is instructive to discuss the primitive, but relevant, situation of a homogeneous silica
and gibbsite surface; each containing singly coordinated surface hydroxyls only [76-78].
According to the bond valence concept, the silanol group has a formal charge of zero
($\equiv$SiOH$^0$), while the aluminol group has a formal charge of -1/2 ($\equiv$AlOH$^{-1/2}$). By varying
the pH of the aqueous solution the $\equiv$SiOH$^0$ can release its proton and becomes $\equiv$SiO$^{-}$, while
the $\equiv$AlOH$^{-1/2}$ group may associate with a proton to form $\equiv$AlOH$_2^{+1/2}$; therefore, the silica
surface is acidic and the gibbsite surface is amphoteric. This difference leads to a different
type of point of zero net proton charge (pznpc): the silica charge approaches the pznpc
asymptotically (all groups in the protonated form at low pH), while the charge of gibbsite
changes linearly around the pznpc (equal numbers of $\equiv$AlOH$^{-1/2}$ and $\equiv$AlOH$_2^{+1/2}$). To be
able to calculate the charge vs. pH curves at different ionic strength a further model is
required that describes the electrical double layer. For flat surfaces the diffuse part of the
electrical double layer can be described with the Debye-Hückel or the Gouy-Chapman
model [79], but for a good description of the electrical double layer of silica, gibbsite and
other mineral surfaces it is necessary to incorporate besides the diffuse layer also a Stern
layer adjacent to the surface. The Stern layer accounts for the dimensions of the counterions
close to the surface and allows the structure of water near the mineral surface to be different
from bulk water [80]. The MUSIC model in combination with the double layer model
provides the means to calculate the proton charge pH curves of silica and gibbsite. As a
consequence of protonation reactions and the different points of zero net proton charge, the shape of the proton charge curve as function of pH of silica is rather different from that of gibbsite. Also the surface potential behavior is different: the surface potential of silica is highly non-Nernstian (i.e. strongly dependent on the ionic strength and non-linear with a change in pH), while the surface potential of gibbsite is approximately Nernstian (linear with a pH change and nearly independent of the ionic strength) [76-78]. Other metal oxide surfaces have also fractional formal charges and react in a similar way as gibbsite. At constant pH also the surface potential of metal oxide surfaces is about constant and about independent of further solution conditions, therefore metal oxide surfaces are also called ‘constant (surface) potential’ surfaces. The fact that most surfaces have multiple types of surface oxygen groups that can react with protons makes the situation more complicated, but the principle difference between silica-type and gibbsite-type behavior remains. The proton charge vs. pH at different ionic strength values, as observed for gibbsite [81], silica [82] and hematite [83] is depicted in Fig. 5.

The multiple types of surface oxygen groups on silica and metal oxides and a detailed electrical double layer model are of most importance for a good understanding of ion binding other than protons, because these ions often do not react with all surface groups and/or may lose part of their hydration shell depending on the type of surface group (inner-sphere versus outer-sphere complex formation). Information on the type and structure of the adsorption complexes has to be derived from spectroscopic studies and can be used in the extended MUSIC model for ion binding, the CD-MUSIC model [84, 85], where CD stands for charge distribution of the bound ions. For goethite the available parameters allow calculations with many different ions and reliable predictions can be made based on ‘model goethite’ even in complex situations.

Concluding, silica and metal oxides have rather different surface properties: (i) silica contains both hydrophilic and hydrophobic sites (siloxanes), while the normal metal oxides...
are hydrophilic, (ii) silica has an asymptotic point of zero net proton charge and a highly
non-Nernstian surface potential, while the metal oxides charging curves are linear in the
point of zero net proton charge and their surface potentials are close to Nernstian. Metal
oxide surfaces can be called, to a good approximation, ‘constant potential’ surfaces. For ion
adsorption to goethite accurate parameters are available for many ions so that reliable
predictions can be made for soils dominated by goethite.

3.3 Silicate clays (alumino-silicates)

A very important fraction of soils are the clays or alumino-silicates. Clay particles are
crystalline and found around the world but they are more widespread in temperate areas. A
main difference with most metal oxides is that silicate clays have a layered structure [14,
86]. The basic structure is composed of a sheet of tetrahedrons of silicon atoms surrounded
by oxygen atoms and a sheet of octahedrons in which an aluminum ion is surrounded by six
hydroxyl groups or oxygen atoms. The tetrahedrons in a sheet are linked by three of the
oxygens in the tetrahedron with adjacent tetrahedra and arranged in hexagonal rings, which
allow the sheet to extend indefinitely in the plane direction; the remaining oxygen is linked
to Al in the octahedral sheet. Also apical oxygen atoms are common to adjoining tetrahedral
and octahedral sheets. Combinations of stacked tetrahedral and octahedral sheets are termed
layers. Many layers are found in each crystal and in some clays the layers are separated by
interlayers in which water and adsorbed cations are found. In nature, cations having nearly
the same radius as $\text{Si}^{4+}$ (e.g. $\text{Al}^{3+}$) can replace $\text{Si}^{4+}$ in the tetrahedral sheet by isomorphous
substitution. Isomorphous substitution can also take place in the octahedral sheet with $\text{Al}^{3+}$
being replaced by a similar-sized cation (e.g. $\text{Mg}^{2+}$). The substitution finds mostly place
with cations that have a lower valence than either $\text{Si}^{4+}$ or $\text{Al}^{3+}$ and this results in unsatisfied
negative charges within the crystal, therefore isomorphous substitution is the primary source
of the negative charge of clay surfaces. The negative charges attract cations from the soil solution.

Based on the number and arrangement of tetrahedral and octahedral sheets contained in the crystal units or layers, silicate clays are classified into two different groups: i) 1:1 type minerals in which the layers have a TO structure build up of one tetrahedral (T) sheet and one octahedral (O) sheet and ii) 2:1 type minerals with a TOT layer structure. The stacking of layers in a 1:1 type crystal leads to two different basal plates: a T-layer with surface oxygens and an O-layer with surface hydroxyls. For a 2:1 type mineral both basal plates are T-layers with surface oxygens. The surface oxygens of the tetrahedra in a T-surface layer form a siloxane surface; in the centers of the 6-membered rings of the tetrahedra a cavity is formed of about 0.23 nm that acts as a specific site for cations if their size matches the size of the cavity [87]. Although the siloxane sites make the surface hydrophobic, the charged sites that occur due to the isomorphic substitution and the presence of hydrated cations make the surface considerably less hydrophobic than a neutral siloxane surface. The basal surface of the T-layer is very common, it occurs for all clays; as different clays have different degrees of isomorphous substitution the hydrophobicity-hydrophilicity of the T-surface also varies, the higher the structural charge density, the lower the hydrophobicity.

The surface oxygens of the O-layer are similar to the double coordinated oxygens of the planar 001 face of gibbsite and form surface hydroxyls that are fully charge satisfied and unreactive with respect to protons. The structural surface charge and the presence of the hydroxyls make the O-basal plate hydrophilic.

The edges of the clay particles have unsatisfied bonds that are quite reactive with respect to protons. For the T-layer edge these are the silanol groups that can protonate: $\equiv\text{SiO}^- + \text{H}^+$ $= \equiv\text{SiOH}^0$. For the O-layers the edge face has doubly and singly coordinated oxygen groups. The doubly coordinated $\equiv\text{Al}_2\text{-OH}^0$ groups are inert in the normal pH range, the singly coordinated $\equiv\text{AlOH}^{1/2-}$ groups at the O-edges are the reactive groups; for their protonation
one can thus write $\equiv\text{AlOH}^{1/2-} + \text{H}^+ = \equiv\text{AlOH}_2^{1/2+}$. Note that the site density of $\equiv\text{SiO}^-$ at the edges for a 2:1 clay will be higher than that at the edges of a 1:1 clay. In the normal pH range the silanol sites will be partly negative, partly neutral, while the aluminol sites will be partly positive, partly negative, the edge surface is therefore amphoteric. The above is the simplest representation of the edge, in practice the situation is more complicated because of non-ideal structures and end effects that cause bond-length relaxation and charge redistribution at the edge surface [88-90].

Proton binding to clays is considerably more complex than to silica or metal oxides; protons will participate in the cation exchange at the basal planes and in the protonation or deprotonation of the edges. Because of the presence of the structural charge on the basal planes next to the variable charge of the edge, one deals with a patchwise heterogeneous surface and for such surfaces the points of zero net proton charge are dependent on the differences between the patches and the electrolyte concentration [83, 91]. One of the complications is that the proton binding to the edges can be affected by the structural charge due to ‘spillover’ of the electrostatic potential between basal and edge surface. In a relatively simple model Avena et al. [92] incorporated both the structural charges that can exhibit ion exchange and the variable edge charge that can change with pH according to a simple proton association - dissociation reaction. The spillover of the potential is taken into account by smearing-out the total particle charge over the entire surface. This model captures the basic features of the proton charging and exchange reasonably well, but the parameters have to be fitted. Fig. 6 depicts the measured proton charge curves for montmorillonite that clearly show the shifts of the points of zero net proton charge, together with the calculated curves [92].

When the spillover effect is calculated in a sophisticated way and the site structure and reactivity are modeled with the MUSIC approach with predicted pK_H values the situation is considerably more complicated. Bourg et al. [88] mention the various models that have been
used to describe the proton charging of montmorillonite and discuss the experimental results
and the most promising sophisticated models.

It is also instructive to briefly discuss how the layers are held together for the main clay
types. In the 1:1 type crystals, with kaolinite as most prominent example, the layers are held
together by hydrogen bonds; as these bonds are strong, the layers are closely held together
in stacks that are large in size, have no internal surfaces and do not swell when wetted. Also
the isomorphous substitution is low, consequently, these minerals have a relatively low
adsorption capacity for cations. The so-called cation exchange capacity or CEC is largely
due to cation adsorption on the edges of the particles and therefore relatively strongly
depending on the pH. The 2:1 clays can be subdivided in four general groups depending on
how strongly the layers are bound together: micas, chlorites, vermiculites and smectites. In
micas a relatively high fraction (~25%) of the silicon atoms in the tetrahedral sheets have
been replaced by aluminum. The resulting large negative charge is satisfied by potassium
ions held rigidly between the adjoining 2:1 layers and preventing expansion of the crystal.
As a consequence of the strong binding they have a fairly large stacking of layers and are
non-expanding. They resemble the 1:1 type clays somewhat and have a relatively low CEC
that is also fairly pH dependent. The chlorites are also non-expansive, the adjacent negative
layers are held together by a positively charged magnesium-dominated octahedral sheet in
the interlayer between the two 2:1 layers. Chlorites have particle sizes, CECs, and physical
properties similar to those of fine-grained micas. Vermiculites have a significant substitution
of aluminum for silicon in the tetrahedral sheets as well as some substitution of magnesium
for aluminum in the octahedral sheet, but lower than in micas. The individual 2:1 layers are
held together only loosely by Mg$^{2+}$ and Ca$^{2+}$ ions that act as bridges between the structural
charges in the adjoining layers. Water molecules are attracted between the layers and
vermiculites expand when wet and shrink when dry. Due to the weak attraction between the
layers the surfaces adjoining the interlayer are also available for adsorption and this leads to
a very large total interface and a large CEC. Due to the limited expansion and the high CEC, vermiculites have a high affinity for weakly hydrated cations such as $\text{K}^+$, $\text{NH}_4^+$ and $\text{Cs}^+$. The final group contains the smectites, with montmorillonite as well known member. In the smectites some magnesium has been substituted for aluminum in the octahedral sheet and some aluminum for silicon in the tetrahedral sheet, but the degree of substitution is less than in vermiculite and the interlayer is larger. Due to the relatively low structural negative charge, $\text{K}^+$ fixation is lower than in vermiculite. As in vermiculite, the cations in the interlayer are exchangeable and this gives rise to high CECs. It may be clear that also surfactant binding to non-expanding clays is far simpler than that to the expanding clays.

3.4. Concluding remarks

Comparing the ion binding models for metal oxides and silica with those of humic substances it follows that the electrostatic part of the model for metal oxides and silica is far more complicated than that of the humic substances, especially when the adsorption of ions other than protons is considered; for humic substances the heterogeneity is most important. Adequate understanding of proton adsorption to silicate clays is only possible when a distinction is made between the edges and the basal planes and considering the spillover of the electrostatic potential of the basal planes to the edges. For all systems the proton charging may change when another species is adsorbed (primary charge adjustment) due to generic electrostatic effects and/or site competition. The different particles also differ in hydrophobicity. Humic substances have many hydrophobic parts, the silica surface is partly hydrophobic and the metal oxides are hydrophilic. Furthermore, adsorption on silica should be distinguished from that on the metal oxides, because of the differences in proton charging behavior and hydrophobicity. With silicate clays the edges are hydrophilic and the basal planes of 2:1 clays are hydrophobic, but the hydrophobicity is weakened by the presence of the structural charge and the adsorbed cations. For kaolinite (1:1 clay) the basal
siloxane surface is fairly hydrophobic, but the basal aluminol surface is hydrophilic; with montmorillonite (2:1 clay) the two basal siloxane surfaces are less hydrophobic than the siloxane surface of kaolinite due to the greater degree of isomorphic substitution.

4. Surfactant binding to humic substances

4.1. Early experimental studies and isotherms measured with surfactant electrodes

The investigations of surfactant binding to humic and fulvic acids are relatively recent, still limited, and have not been reviewed. Therefore, the most important binding studies will be discussed. Early reports are of Tombacz [93, 94] who investigated alkylammonium - humate complexes with X-ray and interfacial tension measurements. Binding isotherms to dissolved humic substances were reported for the first time in 1996 by Traina et al. [95], who used an ultra-centrifugation technique and fluorescence quenching to obtain the binding characteristics of C10-, C12-, and C14-linear alkyl-benzene sulfonates (LAS; in 0.03M NaCl and 0.01M CaCl2) to dissolved humic substances. By using pulse-field gradient NMR Otto et al. [96] found that humic substances enhance the aggregation of the anionic sodium dodecylsulfate (SDS) prior to micellization and that they form ion pairs with cationic C16-trimethylammonium bromide.

The first isotherms ranging from low surfactant concentrations till the CMC were reported by Koopal et al. [97], who used a potentiometric method with a surfactant electrode to measure the equilibrium surfactant concentration. Three surfactants were used, one anionic (SDS) and two cationic (C12- and C16-pyridinium chloride or C12PC and C16PC) and the binding to purified Aldrich humic acid (PAHA) was investigated at three pH values (5, 7, 10) at 0.025 M background electrolyte concentration. No binding could be observed for SDS under the given conditions. The isotherms of the cationic surfactants were independent of the humic acid (HA) content in solution (0.2-0.5 g/L), which indicates that
the results were not affected by possible dynamic aggregation/segregation of HA. The observed isotherms for the cationic surfactants are depicted in Fig. 7; the binding is plotted vs. $c/\text{CMC}^0$, where $c$ is the surfactant concentration and $\text{CMC}^0$ is the CMC in the absence of HA. The binding of C12PC and C16PC increased with increasing pH to maxima that corresponded to the negative charge of the HA (arrows in Fig. 7). By plotting the binding vs. $\log c/\text{CMC}^0$ the isotherms of C12PC and C16PC merged largely, indicating that next to the electrostatic attraction hydrophobic bonding is important. When the charge neutralization point was reached the cationic surfactant–PAHA complexes precipitated, which occurred at approximately 10% of the CMC. For C16PC, the precipitation was complete, but in the case of C12PC, a noticeable fraction of PAHA remained in solution.

Subsequent binding studies were made by Yee et al. [98-101], Matsuada et al. [102] and Ishiguro et al. [103], who all used the potentiometric technique with a surfactant electrode to measure the equilibrium surfactant concentrations. Yee et al. [100] also investigated the binding of SDS to HA using Aso HA (AHA) derived from the Aso area of Kyushu Island, Japan. At pH 9.18 and low ionic strength (0.03M) no binding could be observed between SDS and AHA with either the potentiometric or dynamic light scattering (DLS) method, which confirmed the result of Koopal et al. [97]. However, with the DLS method some interaction could be detected at pH 3.98 and high ionic strength (0.10M). The attraction can be explained by the fact that at 0.1M the electrostatic repulsion is suppressed, but the hydrophobic attraction remains the same and the net effect is a weak attraction. Temperature studies of C12-pyridiniumbromide binding to Aso fulvic (AFA) and AHA at pH 9.18 and ionic strength of 0.03M showed that the binding of C12P$^+$ with AFA was endothermic, i.e. driven by a positive entropy [99], while the enthalpy of C12P$^+$ binding with AHA was slightly negative [101]. For AFA different binding modes were observed at two pH regions, i.e., cooperative binding at pH > 7 and non-cooperative binding at pH < 7, see Fig. 8 [99].
For the AHA-C12P+ system no cooperative binding was observed, but the C12P+ concentration at which the first binding was observed was much lower for AHA than for AFA [101]. The latter reflects that AHA is more hydrophobic than AFA because the acidic type of groups to which the surfactant head groups bind are very similar for FA and HA and the negative charge density of FA is in general larger than that of HA. The C12P+ binding to AFA did rise steeply just before the CMC, i.e. AFA-C12P+ cooperation occurred very close to the CMC. The isotherm of C12P+ binding to AHA leveled-off just before the CMC as the binding was close to its maximum value. The behavior of the AFA-C12P+ system at pH>7 could therefore also be caused by enhanced C12P+ micellization in the presence of AFA, instead of typical C12P+ binding to AFA. The stronger hydrophobicity of AHA than of AFA was confirmed by studying the effect of the aliphatic chain length (n) of the surfactant tail (n: 12, 14, 16) [98]; the results are plotted in Fig. 9. In all cases the comparable isotherms started for AHA at concentrations that were ~10x lower than for AFA. The isotherms shifted with increasing chain length roughly with the same factor as the CMC, but the shift was larger for AFA than for AHA. The results for AHA correspond with the observations of Koopal et al. [97] and Ishiguro et al. [103].

It should be realized that two different hydrophobic interactions have to be considered with surfactant-HS interaction: (i) the hydrophobic effect of transferring the hydrocarbon tail of surfactant into the hydrophobic parts of humic substances (HS) and (ii) the lateral hydrophobic attraction between bound surfactants; (i) contributes to the greater binding strength of HA than FA and (ii) may cause cooperative binding. However, when the hydrophobic effect of transfer of the tail into the hydrophobic parts of HS is large, than the extra effect of (ii) will be small. This explains why (ii) is larger for AFA than for AHA.

Yee et al. [99, 101] also investigated the role of the HS content (0.2-0.5g/L), pH and ionic strength. In accordance with Koopal et al. [97], no HS concentration dependence of binding was observed in both systems; possible self-aggregation of HS did not affect the...
The binding of C12P$^+$ increased with increasing pH, similarly as observed by Koopal et al. [97] for C12P$^+$ and C16P$^+$, and this can be explained by the increase of the negative charge of HS with increasing pH. The bound amount decreased with increasing ionic strength, due to the screening of the electrostatic attraction between surfactant ions and HS; the effect was larger for AHA than for AFA. With DLS the hydrodynamic radii of the AHA-C12PB (PB = pyridinium bromide) and AFA-C12PB complexes were measured at pH 9.2 at ionic strength values of 0.03M and 0.1M for a constant concentration of AFA or AHA of 0.05 g/L. In the absence of surfactant no reliable average size could be measured, but in the presence of surfactant good results were obtained. At low surfactant binding the aggregate diameters were about 200 nm and with increasing surfactant binding up to ~4 mmol C12PB/g the aggregate sizes increased; at 0.1M the increase in aggregate size was similar for AFA and AHA (~1400 nm), while at 0.03M the increase in size was somewhat lower for AFA (~1200 nm) but considerably less for AHA (~500 nm). The difference might be due to the fact that the particle concentration in the case of AFA is much higher; at high ionic strength rapid aggregation may occur for both HS, while at 0.03M the kinetics are important.

The binding of Cn-trimethylammonium (CnTA$^+$) ions to AFA and AHA has been studied by Yee et al. [100] and Matsuda et al. [102]. Yee et al. compared the binding of C12TA$^+$ with that of C12P$^+$ at pH 9.2 and 0.03 M at 25 °C. The binding of C12TA$^+$ to AFA or AHA was weaker than that of C12P$^+$, presumably due to the relatively large size of the headgroup of C12TA$^+$, that prevents a close approach to the negative sites of HS. The differences between the CnTA$^+$ isotherms for AFA and AHA were similar as in the case of C12P$^+$-AFA/AHA A stronger interaction with HA than with FA for C16TAB was also observed by Otto et al. [96] using NMR diffusion analysis. The surfactant head group structure also affects the aggregation behavior: the hydrodynamic diameters of AFA-C12TAB and AHA-C12TAB aggregates were smaller than those of AFA-C12PB and
AHA-C12PB aggregates. Similarly as for C12PB the aggregate size of the HS-C12TAB aggregates increased with surfactant binding and the AFA-C12TAB aggregates were larger than those of AHA-C12TAB. Thieme and Niemeyer [104] also observed an increase in size of HS-C12TAB aggregates with increasing C12TAB binding.

Matsuda et al. [102] investigated first the binding of C12TA+ and C10TA+ surfactants to PHA (derived from an HA-rich layer in Heilongjiang, China) at pH 8 and different NaCl concentrations (0-50 mM) and subsequently that in the presence of divalent metal ions (Cu, Cd, Zn, Pb, and Ca) at pH 8. With respect to C16TA+ the binding curves of C12TA+ were shifted to higher surfactant concentrations; the shift was comparable to the shift in CMC. As stated before, the stronger binding for C12TAB than C10TAB indicates the role of the hydrophobic interaction between the tail of the surfactant ions and the hydrophobic parts of PHA. The affinity of C12TA+ for PHA decreased linearly with the square root of the NaCl concentration. In the presence of 0.5 mM divalent metal salts, the PHA-CnTA+ binding decreased significantly, which can be explained by site competition and screening of the electrostatic attraction. The effect of the different metal ions on C12TAB binding to PHA was similar, but for C10TAB binding to PHA the effect decreases in the order Cd, Zn, Pb, Ca, and Cu.

Ishiguro et al. [103] studied the binding of Cn-pyridinium chloride (C12PC and C16PC) to purified Aldrich humic acid (PAHA), Dando humic acid (DHA, Japan), Inogashira humic acid (IHA, Japan), Laurentian fulvic acid (LFA, Canada) and Strichen Bs fulvic acid (SFA, Netherlands) at pH 5 and 0.005M NaCl, and C16PC binding to PAHA was also studied at 0.1M NaCl. Besides surfactant isotherms also titrations of HS with surfactant were carried out using an automatic titrator combined with a Mütek particle charge detector (M-PCD) [105, 106]. This M-PCD-method is well suited to measure the charge sign of colloidal particles and in the case of titrations with a complexing agent of opposite charge the iso-electric point (IEP) can be detected. The M-PCD titrations were used to determine the
charge of the HS samples at the given conditions (pH 5 and 0.005M NaCl) with poly-DADMAC (poly-diallyldimethylammonium chloride) a strong cationic polyelectrolyte [107, 108] and with C16P⁺. For this charge determination the IEP is required. For the HA samples the charges determined with C16P⁺ were slightly higher than those observed with poly-DADMAC, this could be explained by better screening and stronger charge adaptation in the case of C16P⁺. For the two FA samples the charges obtained with C16P⁺ were significantly larger than those measured with poly-DADMAC. This difference was explained by the relatively low hydrophobicity and corresponding relatively low affinity between FA and C12P⁺. By binding of C16P⁺ a surfactant-FA complex becomes more hydrophobic, this enhances the affinity and the isotherm becomes very steep; for such conditions the rate of titration was likely too fast and leading to too high IEPs. With C12P⁺ the M-PCD method was only used to obtain at the given conditions the bound amount of surfactant at the IEP and the corresponding solution concentration of surfactant. Comparison of the C16P⁺ results with those of C12P⁺ showed that for PAHA the bound amounts (mol/g) of C16P⁺ and C12P⁺ were the same, i.e. at the IEP the charge associated with the HS was neutralized by bound surfactant ions. For the other HS samples the bound amount of C12P⁺/C12PC at the IEP was larger than that of C16P⁺. This must indicate that at the IEP some C12PC was included in the complex, as there is no reason to believe that the screening of the HS charge by C12P⁺ is better than that by C16P⁺ when the HS-C12P⁺ affinity is weaker. Chloride inclusion is most likely caused by the fact that around the IEP the C12PC concentration in solution is relatively high. The behavior of CnP⁺ surfactants often shows a dependence on the type of counterion, indicating incomplete dissociation [109]. With C16PC the affinity for HS is so high that the amount of C16PC left in solution is very low. Therefore, binding of uncharged C16PC to HS is insignificant for C16PC. The M-PCD titrations with the surfactants also reveal that the surfactant–HS complexes reached their IEP before the CMC was reached. Around the IEP some flocculation of the complexes
could be observed, and upon continuation of the titration charge reversal of the aggregated complexes occurred. This may have lead to partial re-dispersion of the aggregates. Shang and Rice [110] and Subbiah and Mishra [111] investigated HA-C16TB complexes by, respectively, small-angle X-ray scattering and synchronous and excitation emission matrix fluorescence, both found (partially) reversible flocculation around the CMC. Shang and Rice used a purified peat HA; Subbiah and Mishra, who used Aldrich HA that was purified by two filtration steps, also investigated SDS and Triton-X100 (t-Octyl-C$_6$H$_4$(OCH$_2$CH$_2$)$_x$OH, $x$=9-10), a nonionic surfactant, and found only very weak interactions for these surfactants.

Binding isotherms of C16P$^+$ to PAHA at two ionic strength values reported in [103] are reproduced in Fig. 10. The binding is plotted in panel a) vs. log $c_{eq}$(surfactant) and in panel b) the same result is depicted but now as double logarithmic plot. The horizontal dashed line indicates the bound amount of C16P$^+$ and the approximate concentration at the IEP as measured with the M-PCD. The thin vertical lines in the top of the figures indicate the CMC values. In panel a) also the proton release due to C16P$^+$ binding is included. Two striking effects can be observed: (i) the isotherms at different salt concentration intersect at the IEP of the complex and (ii) close to or at the CMC the isotherms rise steeply. The latter is an artifact; it is binding of surfactant in micelles, not to PAHA. At the common intersection point (CIP = IEP) the charge of PAHA is neutralized by C16P$^+$. The CIP occurs at a surfactant concentration smaller than the CMC and the C16P$^+$ binding continues till the CMC because of hydrophobic attraction. Binding above the CIP=IEP indicates that C16P$^+$ binds super-equivalently to PAHA. Before the CIP, where the complex is negative, the salt concentration screens the negative charge of the complex and this diminishes the head group surface attraction, at the CIP the complex is net uncharged and salt has no effect, beyond the CIP the complex is positive and the salt concentration screens the repulsion between the head groups. As a consequence the salt concentration dependence inverses at the CIP=IEP. The proton release is up to the CIP very small, C16P$^+$ binding occurs on
already dissociated groups; beyond the CIP the proton release increases and is only slightly smaller than the increase in surfactant binding, so most of the increment in surfactant charge is balanced by proton release.

For sake of comparison the binding of C12P⁺ to sodium polystyrene sulfonate (SPSS), a flexible linear anionic polyelectrolyte, is also plotted in Fig. 10, panels c) and d) [112]. The C12P⁺ isotherms of SPSS at the three different ionic strength values merge before the CMC (the measurements are not accurate enough to observe a CIP); however, for SPSS-C16P⁺ a clear CIP could be detected [112]. The double logarithmic plots (b and d) are best suited to investigate cooperativity of binding. It should be remarked that sometimes the S-shape of single logarithmic plots is used to indicate cooperativity, but this leads easily to erroneous results, because the Langmuir isotherm (no cooperativity) also has an S-shape in these plots.

Firstly, the binding of C12P⁺ to SPSS is considered as this system shows the various aspects most clearly (this is the reason that in Fig. 10 SPSS-C12P⁺ was preferred above SPSS-C16P⁺). Along the isotherm four regions can be distinguished. Region I, at the very low surfactant binding (or surfactant concentration) is the region where the slope of the isotherm is unity, this is the “Henry region” where binding and concentration are proportional; the surfactant is ideally bound on isolated sites of equal energy. For SPSS-C16P⁺, region I cannot be observed; the slope is very steep from the start due to the strong cooperativity [112]. In region II, the slope of isotherms becomes larger than unity, which clearly indicates cooperativity. The transition point of region I and II is defined as the critical association concentration (CAC); at the CAC aggregation of surfactant starts. In region II the number of aggregates along the polyelectrolyte chain increases with increasing surfactant concentration. In region III the slope of the isotherms gradually lowers and becomes smaller than unity due to three factors: 1) the attractive electrostatic interaction becomes smaller due to the decrease of the negative charge of SPSS-C12P⁺ complex, 2) the mixing entropy of “free” and C12P⁻-occupied sites decreases and 3) polymer chain
conformations become more restricted. The last factor will be of little importance for HS, which is internally structured. The CIP marks the end of region III and the start of region IV, the negative charge of the polymer is compensated by the positive charge of bound surfactant. In region IV the aggregates grow slightly further and the SPSS-C12P⁺ complex becomes increasingly positive.

Let’s now return to the PAHA-C16P⁺ system [103]; in the double logarithmic plot the ‘Henry slope’ is indicated by the dashed line. At 0.005M NaCl a CAC and cooperativity cannot be observed, but there is a slight kink in the isotherm that could be the transition point between regions I and II. Region III (up to the CIP) and region IV beyond the CIP can be clearly observed. At 0.1M the isotherm starts with a slope higher than unity, that implies some cooperativity is observed at low surfactant concentration, but the CAC cannot be established. The increase in cooperativity with increasing electrolyte concentration occurs because the electrostatic repulsion between the bound surfactants decreases.

In Fig. 11 the binding at 0.005 M NaCl of C16P⁺ is compared to that of C12P⁺. The influence of carbon chain length on binding can be clearly observed. The C16P⁺ binds at much lower surfactant concentrations than C12P⁺ and the isotherm is somewhat steeper. The larger affinity for C16P⁺ is largely due to the fact that C16P⁺ ‘hates’ the aqueous solution more than C12P⁺, because the shift of the two isotherms is similar to that of the CMC. This reveals the importance of the hydrophobic effect for the binding.

The C12P⁺ binding isotherms for the different humic and fulvic acids at 0.005 M NaCl are compared in Fig. 12. The shape of the binding isotherms to the humic acids is distinctly different from that of the fulvic acids and the C12P⁺ binding decreases in the order of PAHA ≫ IHA ≈ DHA ≫ SFA ≈ LFA. The order ranks the HS with respect to hydrophobicity, PAHA is most hydrophobic the FA’s the least. Cooperativity is not observed at pH 5 and 0.005 M NaCl because the slopes of the curves are almost unity or smaller than unity. These results correspond with the findings of Yee et al. [98, 100, 101] discussed above.
4.2. Surfactant binding measured by solid-phase micro-extraction

Recently, Chen et al. [113] have demonstrated that polyacrylate (PA) coated microfiber sorbs organic cations via ion exchange at the carboxylic groups. When the binding isotherm is known, it can be used to determine the freely dissolved concentration of the given cationic surfactants. The free surfactant concentration in a HS-surfactant system can be obtained by extracting the surfactant bound to PA and determine the extracted amount by HPLC, hereafter the calibration line gives the equilibrium surfactant concentration. The method is called solid-phase micro-extraction (SPME). The method will work well when the surfactant binding to PA is negligible compared to the binding to HS. The double logarithmic representation of the surfactant isotherm to PA fiber is used as calibration curve for the dissolved surfactant, because in this way the isotherm is linear in the low concentration range. Binding isotherms of hexadecyl-trimethylammonium chloride (C16TAC) and benzyldimethyl-dodecyl-ammonium chloride (C12BDAC) to PA are reported as examples; the isotherms of these two surfactants coincide, while their CMCs differ by more than a factor 10. Based on the difference in hydrophobicity this must imply that PA is strongly hydrophilic and that there is no hydrophobic contribution to the specific affinity for the PA-carboxylic groups, mainly the surfactant head group chemistry seems important. The linear part of the log-log isotherms of the two surfactants cover a concentration range from about $10^{-10}$ to $10^{-6.5}$ M. The upper limit of this range is $10^3$ to $10^4$ times smaller than the CMCs. This implies that accurate measurements are only possible in an equilibrium surfactant concentration range considerably below the CMC, for higher concentrations the sensitivity lowers with increasing concentration. Therefore, the method is most valuable for the linear part of the log-log surfactant HS isotherms. The preliminary conclusion is that the PA-SPME will be of substantial added value in the low surfactant concentration range, but that the surfactant electrode will be superior in the higher
concentration range (>10^3 CMC). Anyhow, this new technique is highly welcome and will promote further research of cationic surfactant binding. As example a series of binding isotherms of C12BDAC to purified Aldrich humic acid (purified AHA) at pH 6 and different salts and salt concentrations were measured. Chen et al. [114] continued the latter study in a subsequent paper in which the pH range of the measurements was extended to pH 3, some results are reproduced in Fig. 13.

4.3. Modeling surfactant - humic substance interaction

Ishiguro and Koopal [115] modeled the results of C12PC and C16PC to different humic and fulvic acids with the NICA-Donnan model [38, 43, 54-56] (see also section 7) under the assumption that the surfactant (s) was dominant: (k_s c_s)^ns ≈ \Sigma (k_i c_i)^ni with k_i the intrinsic affinity, c_i the concentration and n_i the stoichiometry of component i. This implies that only the low affinity or carboxylic groups were considered and that the surfactant binding to these groups was larger than the proton binding. For these conditions the NICA equation simplifies to a Hill type equation, but the cooperativity parameter, m, includes the cooperativity ns and the site heterogeneity parameter, p: m = ns • p. This Hill-type equation is complemented with the Donnan model that accounts in a simple way for the electrostatic interactions by assuming that for a given set of conditions there is only one electrostatic potential that governs the behavior in the Donnan phase. With the combined model the experimental results for the binding of the different surfactants to the different HAs at different salt concentration and pH could be well described. For the FAs only the linear part of the log-log isotherms could be described well.

Chen et al. [114] also modeled the surfactant isotherms with the NICA-Donnan model. As the measured isotherms were restricted to the linear range, they made the assumption that the total binding was hardly or not affected by the surfactant (s) binding: (k_s c_s)^ns \ll \Sigma (k_i c_i)^ni with k_i the intrinsic affinity, c_i the concentration and n_i the stoichiometry of...
component $i$. In this limit the NICA equation becomes a Hill-type equation ($m=ns$) for which $(k_se_s)^{ns} << 1$, which has the same form as the classical Freundlich equation. By combining this equation with the Donnan model to incorporate the electrostatics and using the generic description of HA as provided by Milne et al. [55, 56] the salt effects at pH 6 could all be explained well.

By considering the successful modeling attempts by Ishiguro and Koopal [115] and Chen et al. [114] and the fact that two different limiting forms of the NICA equation both lead to a Hill type equation also explains the success of the use of the Hill equation in the work of Yee et al. [98-101]. However, as the Donnan (electrostatic) part was omitted by Yee et al. no explanation could be provided for pH and salt effects. For more information on the NICA-Donnan equation section 7 should be consulted.

### 4.4. Sorption of organic cations and risk assessment

Droge and Goss [116] studied the sorption of organic cations to soil organic matter (SOM) using dynamic column experiments (peat mixed with silicium carbide) with different compositions of electrolytes in aqueous eluents. The sorption affinity of the organic cations to Pahokee peat soil (i) strongly decreased with increasing electrolyte concentration, (ii) was higher in NaCl solutions than in CaCl$_2$ solutions of similar ionic strength, and (iii) was more sensitive to a decrease in NaCl than to a decrease in CaCl$_2$.

Based on the results of this comprehensive study the authors made three remarks. (1) The mass action law equations for ion-exchange reactions predicted trends in a qualitative but not in a quantitative way. (2) Complex models, such as the NICA-Donnan model, are required to fully account for the contributions of ionic interactions to the sorption of organic cations to SOM. (3) The results imply that risk assessment models for organic bases should take ion-exchange processes adequately into account when estimating soil sorption coefficients and bioavailability. These findings indicate that, in general, the NICA-Donnan
5. Surfactant adsorption to metal (hydr)oxides and silicas

5.1 General aspects and reviews

In contrast to the binding of surfactants to HS, the adsorption of surfactants to mineral surfaces has received much attention, because insight in the structure that adsorbed surfactants adopt is vital for practical applications. In older studies most attention was focused on adsorption isotherms and their description. The typical isotherm of ionic surfactants on oppositely charged metal oxides, when depicted as logarithm adsorbed amount vs. logarithm concentration eq (as done in the section on surfactant binding to SPSS and HS), can be subdivided into four regions. Somasundaran and Fuerstenau [117] have introduced this “four-regions” concept and it has been followed in many studies. Typically, the four-regions isotherm extends over about three decades in concentration and adsorbed amount; two examples of the four-regions isotherm are presented in Fig. 14(a), the isotherm to the left depicts the adsorption of an anionic surfactant (sodium C9-benzene sulfonate) on rutile at low pH, the isotherm at the right shows the adsorption of a cationic surfactant (C12-pyridinium chloride) on rutile at high pH; for sake of comparison the same isotherms are presented in Fig. 14(b) as semi-logarithmic plot that provides better insight in the higher adsorbed amounts close to the CMC [118].

To interpret the four-regions isotherm, the adsorption is best described with the relation $\Gamma = Kc^n$, where $\Gamma$ is the adsorbed amount, $c$ the concentration divided by the standard state concentration (1 mol/L), $K$ the effective affinity constant and $n$ the cooperativity. In region I isolated surfactant molecules adsorb and slope-I =1 or close to unity, which implies $n_1$ =1 and ideal behavior. In region II, which starts at low amounts adsorbed, slope-II is mostly >1
and thus $n_{II} > 1$ which is due to cooperative association of adsorbed molecules caused by hydrophobic attraction. For $n_{II} > n_{I}$ the transition I/II is called the \textit{critical surface association concentration} (CSAC). At transition II/III the slope of the log-log isotherm mostly decreases, i.e. $n_{III} < n_{II}$; for $n_{III} > 1$ there is still cooperative adsorption but the cooperation is less than in region II; for $n_{III} < 1$ there is negative cooperativity likely caused by a decrease in configurational entropy due to high surfactant adsorption. Region IV begins at the CMC, beyond which the chemical potential of the monomeric surfactant remains approximately constant, therefore also the adsorption is constant. In some cases, e.g. for cationic surfactant adsorption on silica at low ionic strength, a four-regions isotherm is observed that does not confirm to the above picture: $n_{II} < n_{I}$ (=1) and $n_{II} < n_{III}$ with $n_{III} > 1$, thus region III has the steepest slope. In that case the concentration at which the transition II/III occurs is the CSAC. Although the four-region-type isotherm is well established, especially the interpretation of transition II/III is debated together with the structure of the adsorbed molecules in both adjoining regions. This discussion is enhanced by the results of modern techniques that give information on the adsorbed layer structures. An important contribution to the discussion has been the paper of Manne et al. [119] in which convincing evidence, based on atomic force microscopy (AFM), was presented that surfactants tend to self-assemble at surfaces to form analogues of the micelles in bulk solution. Since then the attention has shifted from adsorption isotherms to modern techniques that provide information on the adsorbed structures. Most reviews of ionic surfactant adsorption pay attention to the four-regions isotherms, but the debate around the four-regions isotherm is still not fully settled and in reviews different views may be expressed. We will return to the four-regions isotherm when the measured isotherms on silica and the metal oxides at constant pH and different ionic strength values will be discussed. First, the various reviews on surfactant adsorption to silica and metal oxides will be briefly introduced.
Several authors have reviewed the state of art of surfactant adsorption, mostly with emphasis to their specialism in the field. The older studies are reviewed in [120, 121] and Rosen’s book [24] provides a general review of surfactants and surfactant adsorption with emphasis on surfactants and their mixtures both in solution and at interfaces and on the modification of the surface properties by surfactants. After the pioneering work of Manne et al. [119, 122] to directly visualize the structure of aggregates formed on surfaces using high resolution scanning probe and atomic force microscopy (AFM), these techniques have been used regularly. This work has been reviewed in [123] and in [119]. Johnson and Nagarajan have presented a theory of surfactant self-assembly on isotropic hydrophobic and hydrophilic surfaces [124, 125]. The theory is extending the treatment of self-assembly in solution [28, 126] with an additional term that accounts for the replacement of the solid surface-water contact by the solid surface-aggregate contact. This contribution depends on the surfactant and surface properties and can be calculated for an assumed geometrical shape of the aggregate; for hydrophobic surfaces and nonionic surfactants it is the Gibbs energy change of replacing surface-water contact by solid surface-aggregate core contact, while for hydrophilic surfaces and ionic surfactants it is (i) the electrostatic interaction between the charged surfactant head groups and the hydrophilic surface, if the surface is charged, plus (ii) the Gibbs energy change associated with displacing water by the surfactant head groups wherever aggregate solid surface contact occurs. The CSAC and the aggregate morphology can be predicted by computing the equilibrium Gibbs energy of formation for a range of assumed geometrical structures and selecting the structure with the lowest Gibbs energy. The CSAC is always predicted to be lower than the bulk phase CMC and a variety of aggregate morphologies are predicted. Morphological transformations of surface aggregates to less energetically favorable structures will occur when the surfactant concentration increases. For hydrophobic surfaces the morphologies include hemispheres, hemicylinders, finite disks, and continuous monolayers depending upon surfactant, solid...
surface and surfactant concentration. For hydrophilic surfaces the situation is more complex
and in addition to full spheres and cylinders a series of composite structures are proposed
which are composed of monolayers oriented with head groups in contact with the surface
and covered with either hemispheres, hemicylinders, finite disks, or another monolayer
(making the full structure a bilayer). It should be noted that the authors have assumed that
monolayer structures with head groups in contact with the surface are instable and
transform in composite structures of covered monolayers. This is only correct for surfactant
concentrations close to the CMC. Stability and wetting studies have shown that beyond the
CSAC the instability and contact angle still increase up to the point where surfactant charge
and surface charge compensate each other [127], which points to largely uncovered
monolayer type structures or teepee-micelles [128]. Nevertheless, the model provides basic
insight in surface aggregation and allows a better judgment of the AMF images that are
mostly obtained around or above the CMC.

Derived from results obtained with modern measurement techniques Tiberg et al. [129]
have discussed in a comprehensive way the adsorption mechanisms and interfacial
structures of nonionic and ionic surfactants for both hydrophobic and hydrophilic surfaces.
It is an excellent review for a good orientation on surfactant adsorption.

Atkin et al. [128] have given a follow-up on this review with emphasis on the adsorption
process for ionic surfactants. Traditional equilibrium data (including the four-regions
isotherms), discrete aggregation on the substrate (mainly silica), and the morphology of the
aggregates are thoroughly discussed. As the AFM is most useful at detecting periodicity,
surfactant–substrate combinations that produce highly regular morphologies produce the
clearest AFM images. The hydrophobic cleavage plane of graphite orients the adsorbed
surfactant structures more strongly than any other substrate, and hemi-cylindrical structures
are formed for surfactant chains with more than 10 carbon atoms, smaller chains result in a
laterally featureless adsorbed layer. The hydrophilic, crystalline mica substrate also orients
the adsorbed surfactant structures, but not as strongly as graphite. The alkyl trimethyl-
ammonium surfactants form cylindrical admicelles, which were stable at relatively high
ionic strength, but less stable at low ionic strength. The morphology of adsorbed gemini
surfactants on mica depend strongly on the geometry of the surfactant monomer. On silica
surfactants are less strongly oriented, due to a lower surface charge and the amorphous state
of the substrate. The adsorbed configuration of C16TAB was concentration dependent, with
short rods at 0.9*CMC and worms at 10*CMC. Similar structures have been observed for
C14TAB and C12TAB on silica. For C16TAC, only spheroidal structures were present. In
all cases there was little or no long range ordering of the adsorbed aggregates. By
sequentially increasing the substrate hydrophobicity the structure of adsorbed nonionic
surfactants can be changed in a controlled manner from diffuse adsorbed micelles on the
least hydrophobic interface, to densely packed micelles, to a bilayer, to a monolayer on the
most hydrophobic interface. The information gained by AFM, fluorescence quenching and
neutron reflectivity is discussed and with this knowledge, kinetic data obtained from
ellipsometry and optical reflectometry, are analyzed. In the summary the likely mechanisms
of adsorption along the four-regions isotherm are proposed.

Zhmud and Tiberg [130] discuss applications of modern techniques (tensiometry,
ellipsometry, photon correlation spectroscopy, and neutron reflectivity) with emphasis on
the physical principles and theoretical aspects related to the adsorption and desorption
kinetics, interfacial structure development, wetting enhancement, and interfacial dynamics.
The review includes mathematical derivations that demonstrate how raw data can be
transformed into sought layer characteristics. After the discussion of the techniques the
intermolecular interactions in aggregated surfactant complexes are discussed and
experimental studies on the dynamics and structure of adsorbed surfactant layers are briefly
reviewed. The discussion of the isotherm for ionic surfactants is somewhat confusing
because two surface association concentrations are considered, the hemi-micelle
concentration and the critical surface association concentration. Although different names
have been used for the concentration at which aggregation starts, there is for a given system
only one concentration where the four-region isotherm steeply increases. In the last part of
the review wetting enhancement by surfactants and the effect of surfactants on the damping
of capillary waves are discussed.

A topical review on *in-situ* surface-enhanced infrared and non-linear vibrational
sum-frequency spectroscopy in relation to a better understanding of adsorption at
mineral–water interfaces is presented by Schrödle and Richmond [131]. Particular
consideration is given to organic adsorbates including surfactants. The authors conclude that
non-linear optical methods greatly extend the well-established linear IR techniques and
provide many opportunities to advance our understanding of the structure and dynamics of
mineral–water interfaces. This will be true, but, as the material is quite complex, help of an
expert will be required for the data analysis.

Many of the experimental results discussed in the above reviews are obtained on graphite,
mica, quartz and silica and these surfaces are different from the metal oxide surfaces.
Reviews that put more emphasis on metal oxide surfaces are those of Paria and Khilar [132]
and Zhang and Somasundaran [133]. Paria and Khilar discuss the four-regions isotherms of
ionic surfactants on oppositely charged metal oxides, and pay special attention to region IV,
that occurs for surfactant concentrations > CMC, where some anomalies have been
observed. Influences of molecular structure, temperature, salt concentration that are very
important in surfactant adsorption are reviewed for both nonionic and ionic surfactants and
the state of aggregation of the adsorbed surfactants is discussed on the basis of AFM and
fluorescence measurements. Also the adsorption behavior and mechanism of different
mixed surfactant systems (anionic–cationic, anionic–nonionic and cationic–nonionic) are
reviewed. Mixtures of surface-active materials can show synergistic interactions, which can
be manifested as enhanced surface activity, spreading, foaming and detergency.
The review of Zhang and Somasundaran [133] is concerned with adsorption of single
surfactants as well as mixtures of various types and provides lists of systems (surfactant /
surface / method) that have been studied for both ionic and nonionic surfactants. Also
zwitterionic surfactants are briefly discussed. The information derived from the isotherms,
combined with fluorescence spectroscopies, electron spin resonance (ESR), Raman
spectroscopies, small angle neutron scattering, neutron reflectivity and AFM, has been used
to discuss the structural properties of the surface aggregates along the isotherms. Compared
to the adsorption of single surfactants, adsorption of mixtures of anionic– cationic, anionic–
nonionic, cationic– nonionic, cationic– zwitterionic and nonionic– nonionic types generally
exhibit synergy at interfaces. To predict the shape of mixed aggregates of nonionic
surfactant mixtures a new model of the critical packing parameter is derived that takes into
account the mole fractions of adsorbed surfactants.

5.2 Surfactant adsorption and charge regulation

In the next sections the adsorption isotherms of ionic surfactants to rutile (TiO₂) and
silica measured at constant pH and different background 1-1 electrolyte concentrations will
be critically examined, together with the adaptation of the surface charge as a result of the
surfactant adsorption. Results for rutile are taken from [118, 134-136], those for silica from
[137, 138]. It will be shown that these surfaces behave differently with respect to surfactant
adsorption. A strictly constant pH allows the calculation of the surface charge regulation
(adaptation) and measurements at different ionic strength at constant pH can be of help with
unraveling the orientation of the adsorbed surfactant. In order to keep the pH constant, the
pH is regularly checked during the course of adsorption and restored to its initial value. The
amounts of alkali or acid to restore the pH are used to calculate the proton charge adaptation
of the surface. The surface charge adjustment is important in itself (in many studies the
effect of a change in surface charge is simply ignored), but it can also help to create a
picture of the adsorption behavior. It is also required for full understanding of specific co-ion and counterion effects on surfactant adsorption. A combination of surfactant adsorption and surface charge (adjustment) results makes it possible to differentiate between (i) adsorbed surfactant molecules with the head groups adjacent to the surface (‘head-on’ adsorption) which may occur with a counterbalancing (de)protonation of the surface, and (ii) molecules with an oppositely orientation, i.e. with their head group at the solution side (‘head-out’ adsorption) which will not affect the surface charge. To discriminate between different aggregate structures monolayer type aggregates of head-on adsorbed molecules will be called ‘teepee’-micelles [128] and local bilayer type structures (head-on + head-out adsorbed molecules) ad-micelles. Knowledge about the prevailing molecular orientation and or aggregate structure is of direct importance for wetting and stability behavior of the particles [25, 127].

5.3. Ionic surfactant adsorption on rutile.

Rutile is chosen because it allows studying anionic surfactant adsorption on positive rutile at the acidic side of the pH of zero net proton charge (pH<pH_{pzmc}) and cationic surfactant adsorption on negative rutile at the alkaline side (pH> pH_{pzmc}). Rutile is not of direct importance for soil systems but at the same ionic strength the adsorption behavior of surfactants on rutile is very similar to that on aluminum oxides and hematite when ΔpH = pH-pH_{pzmc} is considered as parameter that characterizes the charge density [134, 139-141]. At similar pH values the adsorption isotherms and admicelle structures of a given surfactant on different metal oxides surfaces will strongly depend on pH-pH_{pzmc}, i.e., on the type of metal oxide surface [142]. Nonionic adsorption on rutile will not be discussed.

In Fig. 15 adsorption isotherms of sodium nonyl-benzene-sulfonate (SNBS) on rutile at pH 4.1 and three salt concentrations are depicted as log-log (a) and lin-log (b) plots [134]. In region I the coverage is too low for the adsorbed molecules to interact with each other,
the slope is about unity. Mutual attraction starts in region II, where the coverage is still very low: roughly between 0.03 and 3\% of the maximum coverage. The concentration at the transition point between region I and II is the critical surface association concentration or CSAC. A straight line with a slope of about four fits the data points in region II, indicating a strongly positive cooperativity of the adsorption and proof for surfactant self-assembly. The slope in region III is considerably lower than that of region II and it depends notably on the salt concentration. It is less than unity for low salt concentrations and somewhat higher than unity for 0.1M salt. Salt addition promotes the adsorption in region III by screening the head group repulsion. The transition between regions II and III is more pronounced for low salt concentrations, which also indicates that at low salt concentrations the repulsion between head groups becomes important in region III. The CMC marks the transition to region IV, where the plateau is reached.

Due to the fact that isotherms at three salt concentrations are plotted another important phenomenon occurs: the three isotherms at different salt concentration intersect in a common intersection point or CIP. The presence of such a CIP was first noted by De Keizer et al. [143] who also showed that in the absence of specific adsorption of other ions and/or activity effects the CIP should correspond to the IEP of the particles covered with surfactant.

In Fig. 15(b), the lin-log plot, the strong cooperativity present at low coverage is evident at low salt concentrations as a weak shoulder in the isotherms. The lin-log isotherms show much better than the log-log isotherms that the ‘bulk’ of the adsorption occurs in region III. Also the position of the (near) CIP is clearly shown. The plateau values reached in region IV are about the same for the different salt concentrations.

The adsorption of SNBS and surface charge of rutile are shown in Fig. 16 as function of the surfactant concentration at three pH values and two salt concentrations [134]. To facilitate the comparison the surface charge is divided by F, the Faraday, so that proton excess, $\Gamma(0)$, and adsorbed amount of surfactant, $\Gamma_s$ (SNBS), are both expressed in $\mu$mol/m$^2$. 
Information about the structure of the adsorbed layer can be obtained by comparing $\Gamma(0)$ and $\Gamma_s$. For a given pH, see e.g. pH 4.1 (Fig. 16, panel a and b), head-on adsorbed surfactant ions may either have displaced adsorbed Cl$^-$ ions or caused the formation of new proton charges at the surface to counter balance the surfactant charge and the latter increases $\Gamma(0)$. Head-out adsorption will not affect the surface charge. Therefore, as long as the surface charge adjusts to the surfactant adsorption head-on adsorption occurs. Initially $\Gamma(0) > \Gamma(SNBS)$ because the surface charge is already present (best seen at 0.1 M NaCl) but at somewhat larger surfactant concentrations $\Gamma(0) < \Gamma(SNBS)$. Beyond the intersection point the difference between $\Gamma_s$ and $\Gamma(0)$ is due to head-out adsorption plus some head-on adsorption of molecules that are accompanied by a counter ion; therefore, head-out adsorption is about proportional to $\Gamma_s - \Gamma(0)$ but not necessarily equal to this difference. The proton uptake due to surfactant adsorption is the strongest at low surfactant adsorption and low salt concentrations: with almost every adsorbing surfactant molecule a proton is adsorbed, indicating that all surfactant ions will be adsorbed head-on. At higher salt concentrations the initial surface charge is higher, therefore the initial slope of the $\Gamma(0)$ curve is less, surfactant ions cause displacement of Cl$^-$ ions at the surface by NBS$^-$ ions and proton adsorption. At high amounts adsorbed the surface charge becomes almost independent of the salt concentration: the surfactant is responsible for the screening of the surface charge. Very similar results are obtained at pH 5.1 and 6.0. At pH 6 the pure rutile surface is slightly negative ($pH_{pznc}$ rutile 5.85); however, the specific affinity of the head group for the surface is sufficient to overcome the coulomb repulsion and the surfactant adsorption induces a positive charge on the surface. Therefore, only positive surface charges are observed once surfactant is present, and the two curves coincide at low surfactant concentrations.

In general, the proton adsorption tends to level off after the CIP in the isotherms. The CIP corresponds for SNBS with the IEP and with the point where the surface charge and the
surfactant charge balance, the equivalence point. The strong surface charge adjustment before the CIP in combination with the observed steep slope of region II of the isotherms indicates that teepee-micelles are formed at the surface. Teepee-micelle formation is promoted by the ‘flexibility’ of the surface charge. Changes in surface charge due to SNBS adsorption still occur beyond the CIP. This demonstrates that the head-on adsorption is not completed at the CIP. The substantial difference between $\Gamma_s$ and $\Gamma(0)$ beyond the CIP indicates that a large part of the additionally adsorbing molecules is oriented head-out: after the CIP the formation of admicelles is dominant. The rate of transition from teepee- to ad-micelles depends strongly on the salt concentration; the higher the salt concentration the sharper becomes the transition. The SNBS adsorption around the CMC depends only weakly on the salt concentration, increasing the salt concentration increases the adsorption somewhat due to the better screening of the head group charges.

Comparing the behavior at the three pH values (Fig. 16) it can be observed that the pH effect on the SNBS adsorption is most strongly at the initial part of the isotherm (region I and II): the smaller $\Delta pH = pH - pH_{pznpc}$ is, the lower is the affinity of the surfactant for the surface. However, the pH may also affect the admicelle structure because an increase in surface charge density can lead to a reduction in the effective headgroup area of the adsorbed surfactant and this may decrease the critical packing parameter [142]. Further adsorption studies on positively charged rutile with four sodium p-dodecylbenzene sulfonate isomers (3SDBS, ISDBS, 5SDBS, 6SDBS) [135] to study the effects of aliphatic chain length and architecture, revealed that the adsorption increases with chain length and decreases if the point of attachment of the aliphatic chain to the hydrophilic part of the molecule is shifted from a terminal to a medial position.

The adsorption on negative rutile was studied by using two pyridinium surfactants (C12, C14) and the results were compared with those of SNBS (see Fig. 14) [118, 136]. The four-regions isotherm and the CIP are found also for the anionic surfactant adsorption on
positive rutile, but for the weakly adsorbing CnP⁺ ions the situation is considerably more complicated than for the strongly adsorbing NBS⁻ ion. Due to the relatively weak CnP⁺ adsorption, the competition with the salt cations (K⁺) becomes noticeable and the ‘CIP’ is no longer ‘sharp’, the IEP shifts with the salt concentration and differs from the ‘CIP’. However, the ‘CIP’ remains the transition point for tepee- to ad-micelle formation. For more details and a further discussion of surfactant adsorption on rutile or the metal hydroxides in general the quoted references should be consulted.

5.4. Ionic surfactant adsorption on silica

The adsorption of cationic surfactants on silica (negative) is far greater than the adsorption of anionic surfactants. Nevskaiia et al. [144] have shown that small amounts of anionic surfactants (C₉H₁₉-phenol-(OCH₂-CH₂)ₙ-SO₄⁻ Na⁺, with n = 4 and 10) could be adsorbed on quartz but the relevance is very limited. Therefore, no further attention will be paid to anionic adsorption to silica.

The adsorption isotherms for C₁₂PC on Aerosil, measured at pH 9 and two salt concentrations are shown in Fig. 17(a) as log-log plot and in Fig. 17(b) as lin-log plot [137]. The isotherms at low and high salt concentrations have different shapes. At low salt concentration the adsorption shows a ‘hesitation’ in the both the lin-log and log-log plot. This is usually observed for adsorption of cationic surfactants on silica in the absence of salt or at low salt concentrations [145-148]. At both low and high salt concentrations four regions may be distinguished in the log-log plots and for low salt concentration the different regions can also be observed in the lin-log plot. The slope in region I of the C₁₂PC isotherms is equal to unity within experimental error. In this region specific and coloumbic attraction between surface and surfactant ions take place, but the adsorbed molecules don’t interact with each other. In region II, the slope of isotherms, measured at low salt concentration, is less than unity. This demonstrates that surfactant self-assembly still does not occur! Self-assembly is
counteracted by mutual head group repulsion and by the partial hydrophobicity of the
pyrogenic silica surface that interacts with the surfactant tails by hydrophobic attraction. This
behavior deviates strongly from that observed for adsorption of C12PC on rutile [118, 136]
and in general from surfactant adsorption on the crystalline metal oxides. On silica surfactant
molecules adsorb, at low salt concentration, without appreciable lateral interaction up to the
end of region II. In the same concentration range the surface charge adapts strongly to the
surfactant adsorption. Therefore, the mutual repulsion between the headgroups will be modest
and most likely the tail-surface interaction is the main reason for the absence of self-assembly.
Evidence for tail-surface interaction was obtained from both comparison of the effect of
C16TAB and tetramethyl-ammonium bromide on the charging behavior of Aerosil [138] and
theoretical calculations using the SCFA model [137]. The surfactant concentration has to
increase to about 0.1*CMC before region III begins and the aggregation starts. Due to lateral
hydrophobic attraction the slope of the isotherm in region III is steeper than that in II and the
CSAC is located at the transition of region II to region III.
Increasing the salt concentration increases the slope of region II due to the higher initial
surface charge and a better screening of the lateral head group repulsion. At 0.1M salt the
slope of region II is larger than that at region I; therefore, the CSAC now occurs at the
transition from region I to region II. A relatively high salt concentration leads to a relatively
high surface charge due to the presence of many counterions close to the surface, and this
reduces the surface hydrophobicity. Consequently, not only the mutual head group repulsion
is diminished, but also the tail-surface attraction, therefore surfactant self-assembly occurs at
a much earlier stage than at low salt concentration. The slope in region III depends only
weakly on the salt concentration and is now smaller than in region II. The transition to region
IV is observed at the CMC. An increase in the salt concentration leads to a rise of the plateau
value and small decrease of concentration (CMC), where plateau is reached: diminishing the
lateral repulsion allows a closer packing of the molecules and/or a higher aggregate density.
The isotherms show a CIP in region II, the concentration at which the intersection occurs, corresponds well with the point where the surfactant charge and the surface charge balance, the equivalence point. The fact that CIP and equivalence point coincide is direct evidence that the binding of C12P⁺ to Aerosil is much stronger than that of K⁺. The much stronger binding of C12PC to silica than to rutile [118, 136] is also likely due to the tail-surface affinity in the case of silica. The reasons for the CIP are the same as those mentioned for SNBS adsorption on rutile. The difference with the SNBS-rutile case is that the CIP is located in region II instead of in region III of the isotherm.

The effect of C12PC adsorption on the surface charge of Aerosil at pH 9 is shown in Fig. 18 for two salt concentrations (0.001 M and 0.1M KCl) [138, 149]. To allow easy comparison of the adsorbed amount of surfactant, Γₛ, and the surface charge the latter is divided by F, the Faraday, an indicated as, Γ(0); they are both expressed in μmol/m². The maximum change in surface charge occurs at low surfactant and low salt concentrations, where the adsorption and surface charge isotherms almost coincide. It means, that at low salt concentrations with almost every adsorbing surfactant ion a proton is desorbed and practically all adsorbed surfactant ions will adsorb head-on. After compensation of the surface charge by surfactant (the equivalence point) the surface charge continues to increase weakly whereas the surfactant adsorption increases substantially. This indicates that locally adsorption of surfactant in head-out position starts, i.e. head group directed to the solution. The weak increase of Γ(0) shows that some of the additionally adsorbing molecules are still head-on adsorbed. On top of this some other surfactant ions may adsorb head-on if they are accompanied with a counter-ion. In general, one may conclude that small admicelles are formed around the initially head-on adsorbed (isolated) molecules. This representation of the surface aggregate corresponds with that suggested by Zhu et al. [150, 151].

At high salt concentration the changes in surface charge due to the surfactant adsorption are much less, evidently C12P⁺/K⁺ exchange is important. The high salt concentration
screens the surface charge fairly well and it is difficult for the surfactant to start its
adsorption. Once the adsorption starts it increases strongly over a rather small concentration
range and a sharp intersection point with the Γ(0) isotherm is found. This equivalence point
corresponds with the CIP of the C12PC isotherms. Before the equivalence point C12P⁺
adsorption occurs by exchange of K⁺, beyond the equivalence point the adsorbing surfactant
ions bring their ‘own’ counter ions to compensate their charge. These adsorption
mechanisms have been indicated before by Bijsterbosch [148]. The super-equivalent
adsorption indicates admicelle (head-on + head-out adsorption) formation. The effect of a
high salt concentration is twofold: (1) before the CIP teepee-micelles are formed (steep
slope of region II of the log-log isotherm) and (2) the transition from head-on adsorbed
molecules to admicelles becomes very sharp. In summary, below the CIP head-on
adsorption occurs: at low salt concentration as isolated molecules, at high salt concentration
as teepee-micelles, above the CIP head-out adsorption and ad-micelle formation occurs at
all salt concentrations. Increasing the salt concentration increases the packing density of the
adsorbed surfactants, but the values of the Γ(0) at the CMC hardly depend on the salt
concentration. This indicates that the admicelles become more asymmetric and that around
the CMC the screening of the surface charge is almost entirely done by surfactant.

Goloub et al. [137] also studied the adsorption behavior of C16PC and the effect of the pH
on both C12PC and C16PC; in general the behavior for C16PC is very similar but the
isotherms are shifted to lower concentrations, which makes it difficult to study the lower
regions of the four-regions isotherms. Lowering the pH to 7 or 5 also makes the study of the
isotherm more difficult because less adsorption occurs when the negative surface charge is
relatively low. For a further discussion the original papers should be consulted.

The experimental results for both rutile and silica have also been compared with
calculations using the Self-Consistent-Field Adsorption (SCFA) model [25, 134, 135, 137,
152, 153] (see also section 7) both SCFA theory and experiments show that the course of the
isotherms is dependent on the salt concentration and the charge characteristics of the surface. Also the SCFA model isotherms obtained at different salt concentrations show a CIP, and also here the CIP marks the point where the coulombic interaction vanishes and the surfactant orientation of the newly adsorbing molecules changes form mainly head-on to predominantly head-out. Although the SCFA theory is not perfect, it definitely can help to understand surfactant adsorption and to design experiments to gain further insight in the adsorption behavior.

The surfactant orientations in the adsorbed layer as deducted from the present analysis relate very well to experimentally observed maximums in hydrophobicity [154-157] and flotation recovery [158, 159] of particles as a function of surfactant concentration: at low adsorbed amounts the particles are hydrophobized, whereas at high adsorbed amounts the presence of head groups at the solution side make the particles more hydrophilic again [1]. The results also indicate that around the CIP the colloidal stability will be at its minimum, not only because the particles plus adsorbed layer are uncharged, but also because the particles are hydrophobized [127].

5.5. Nonionic surfactant adsorption on silica

The adsorption of nonionic surfactants of the oligo(oxyethylene) n-alkyl ether \([\text{Cn(EO)m}]\) or oligo(oxyethylene) n-alkylphenyl ether \([\text{Cn\Phi(EO)m}]\) type at the solid/aqueous interface has been the subject of much experimental and theoretical research, see the review of Tiberg et al. [129] and the cited references. The vast majority of adsorption studies of nonionic surfactants have been performed on silica. A classical set of adsorption isotherms of nonionic surfactants with different polar head group is shown in Fig. 19 [160]. From Fig. 19 it follows that the affinity increases with EO chain length, but that the adsorption level around the CMC strongly decreases.

The main experimental evidence shows that the adsorbed nonionic surfactants aggregate
strongly on hydrophilic surfaces such as silica. Oxyethylene (EO) groups bind initially to the silanol groups by hydrogen bonding and this initiates strong adsorption at the CSAC through self-assembly. Contrary to ionic surfactants, nonionics do not readily make hydrophilic particles very hydrophobic because the head-on adsorption (monolayer-type) is relatively low. With limited EO chain length pronounced adsorption on silica starts at the CSAC, because the affinity of the EO segments to the surface is relatively weak. In the case of surfactants having short EO chains relative to the length of the aliphatic chain part, bilayers are formed on the surface and when the EO chain is long the adsorbed layer consists of small admicelle type aggregates.

Denoyel and Rouquerol [161] have shown that different silicas (Spherosil, Aerosil and quartz) show quite different adsorption levels (µmol/m²), but that the adsorption mechanism is the same on the silicas despite the great differences in adsorption. The initial anchoring of molecules is exothermal on the three samples (but with a different magnitude of the corresponding enthalpy) and the subsequent surfactant aggregation is endothermal and tends for all three surfaces towards a value equal to the micellization enthalpy. This two-step mechanism involves weak hydrophobization of the surface at low surfactant concentration followed by hydrophilization at concentrations still below the CMC. Such a transition has been confirmed for the quartz-nonionic surfactant system by contact angle measurements [162].

For a good understanding of the behavior accurate experiments come in the first place, but modeling experiments can help to explain the behavior and reveal trends that govern the adsorption. SCFA model calculations [152] have been used to analyze the results of Levitz et al. [160, 163, 164] on the adsorption of a series of alkyl-phenol polyoxyethylene glycols on silica. This work was followed by a detailed experimental study of the adsorption of C12(EO)6, C12(EO)25 and the oligomer (EO)122 on different silicas [153]. To be able to apply the various techniques, different silica surfaces had to be used. To some extent this
was a disadvantage, but by comparing the results, trends emerged, and these could be studied with the 1D and 2D versions of the SCFA theory and in this way a better insight was gained. Recently the SCFA theory has been applied to investigate the experimentally observed pH and salt concentration dependence of Cn(EO)m adsorption to silica [165]. The parameter choices and some aspects of the model were refined as compared to the older SCFA modeling; the calculations contributed to detailed insight in the mechanisms of the pH and salt effects. At high ionic strength, the solvent quality for the surfactant head groups is affected, which changes both bulk solution and adsorption behavior of the surfactant. Nonionic surfactants adsorb above the CSAC, which is a function of surfactant and surface properties. Therefore, the CSAC varies with both the ionic strength and the pH. The model predicts that with increasing ionic strength, the CSAC will first slightly increase but then drop substantially. The charge on the surface is pH dependent, and as the head groups bind through H-bonding to the silanol groups, the CSAC increases with increasing pH. The predictions follow experimental findings, which shows that the molecularly realistic SCFA model can reveal a rich interfacial behavior.

6. Surfactant adsorption to clays

6.1 General aspects and outline

An analysis of surfactant adsorption on clays is faced with several difficulties: (1) clay surfaces have basal plates and edges that make the surfaces patchwise heterogeneous, (2) for swelling clays surfactants may also intercalate between the layers and (3) the anionic surfactants tend to precipitate in the presence of multivalent cations. The patchwise nature of the clay surface is a fact that has to be considered in all cases of surfactant adsorption to clays. For expanding clays the adsorption is far more complicated than for non-expanding clays. For instance, the adsorption of anionic (SDS), nonionic (Triton X100) and cationic
(C18TAB) surfactants did not modify the X-ray diffraction diagrams of kaolinite and sepiolite (non-expanding clay types), but the interlayer space of montmorillonite (expanding clay type) increased through nonionic and cationic surfactant adsorption [166]. When the interlayer spacing expands upon surfactant adsorption the adsorption capacity becomes a function of the amount adsorbed and this leads to complex shaped adsorption isotherms that do not follow classical behavior. A further complication with anionic surfactant adsorption is that cations bound to the clay surface may exchange with the counterions of the anionic surfactant and this can lead to precipitation of the anionic surfactant, especially when the clay charge is (partially) compensated by multivalent cations [167]. Furthermore, at relatively low pH leaching of Al$^{3+}$ ions from the clay may occur and can be promoted by the presence of anionic surfactants through Al$^{3+}$ complexation with the anionic surfactant followed by surfactant precipitation [168-170]. Any precipitation of the surfactant obscures the pure adsorption results.

The most studied non-expanding clay with respect to surfactant adsorption is kaolinite and cat-, an- and nonionic surfactant adsorption to kaolinite will be discussed in the next sections, with emphasis to cationic surfactants. Subsequently, surfactant adsorption to montmorillonite will be discussed at surfactant concentrations up to the CMC. The situation of surfactant binding to expanding clays to produce organo-clays for various specific applications is a field in itself. For reviews of this field we refer to [171-174] and to some original papers [175-178].

### 6.2 Cationic Surfactant adsorption on kaolinite

De Keizer et al. [143, 179, 180] have made a very thorough study of the adsorption of cationic surfactant adsorption (C16TAB and C12PC) on kaolinite (Sigma Chemicals) by investigating the adsorption behavior at different pH values and salt concentrations as a function of temperature. Adsorption isotherms (lin-log) of C16TAB and C12PC at different
electrolyte concentrations show a well-defined CIP where the adsorption is independent of
the electrolyte concentration [143, 179], see Fig. 20. The CIP coincided with the IEP of the
particles with adsorbed surfactant, indicating that at this point the surface charge is
compensated by the surfactant charge, including any counter-charge that may be present
within the slip layer. The presence of a CIP and the correspondence with the IEP seems to
be a general property for adsorption of organic ions on a surface with an opposite charge.
Very similar behavior occurred for the surfactant adsorption on silica and rutile (metal
oxides). Also for kaolinite head-out (second layer) adsorption driven by hydrophobic
attraction starts after the CIP. This view is supported by the effect of the temperature on the
adsorption. Below the CIP, adsorption is independent of temperature, whereas after the CIP
it has a maximum as a function of temperature. With increasing temperature the adsorption
changes from endothermic to exothermic, as for micellization. The CIP occurred for both
surfactants at about 32 μmol/g, roughly 40% of the maximally attained value. In the
CIP=IEP the charged surfactant groups just compensate the surface charge. Therefore, the
above value may also be compared with the cation-exchange capacity (CEC). The CEC
according to the silver-thiourea method was 57 μmol/g (exchange on plates plus edges),
while the CEC with the ammonium acetate method at pH 7 was 30 μmol/g (cation exchange
on the plates). This provided evidence that C16TAB and C12PC adsorption took place
essentially on the plate surfaces.

In part 3 of the study [180] the investigations were complemented with
micro-calorimetric measurements of the adsorption enthalpies of C12PC on Na-kaolinite at
two electrolyte concentrations (5 and 100 mM NaCl) and three temperatures (6, 20 and
60 °C). Curves of cumulative adsorption enthalpies against amount adsorbed show a break
around the CIP adsorption. The curves are about linear both before and after the break, and
result in partial molar adsorption enthalpy values that are independent of surface coverage,
confirming that the surface is homogeneous with respect to the C12PC adsorption. This is
additional evidence that the surfactant is hardly adsorbed on the edges. The isotherms are
independent of temperature up to CIP coverage, implying a zero isosteric adsorption
enthalpy, but upon formation of the second layer (head-out mode) the adsorption has a
maximum at about 23 °C, implying a transition from an endothermic to an exothermic
process. The calorimetric adsorption enthalpies change sign at T ≈ 24-30 °C. In this respect,
the adsorption process is very similar to micellization [181]. The electrolyte concentration
has a minor effect on the adsorption enthalpies of the first (head-on adsorption) and second
(head-out adsorption) layers. The directly measured heats agree qualitatively with the
isosteric heats.

In the last paper [180] the isotherms were also modeled with a simple two-layer model
based on the Frumkin-Fowler-Guggenheim model that takes into account both the
interaction with the surface and the lateral interaction, therefore the standard Gibbs energies
have two contributions: $\Delta G^0_{\text{surface}}$ and $\Delta G^0_{\text{lat}}$. For the first layer it is assumed that the
molecules adsorb head-on to the mineral surface with their tails at the solution side, for
adsorption in the second layer the ‘surface’ is made-up of the tails of the first layer and the
molecules adsorb head-out with their tails contacting the new ‘surface’. Since the
adsorption on kaolinite particles involves chemical and electrical contributions, the changes
in the standard Gibbs energies can be written as $\Delta G^0 = \Delta G^0_{\text{chem}} + \Delta G^0_{\text{el}}$. $\Delta G^0_{\text{el}}$ accounts for
the generic interactions due to the ‘smeared-out’ electrostatic potentials, $\Delta G^0_{\text{chem}}$ contains
chemical (specific), hydrophobic and ‘local’ electrostatic interactions (local self-atmosphere
potentials). Because of their special nature and important role the hydrophobic attractions
are separately indicated. The standard Gibbs energy of adsorption in contact with the
surface (layer: 1, surface: s), $\Delta G^0_{1,s}$ can then be written as $\Delta G^0_{1,s} = \Delta G^0_{1,s,\text{spec}} +$
$\Delta G^0_{1,s,\text{hydrophobic}} + \Delta G^0_{1,s,\text{el}}$; $\Delta G^0_{1,s}$ becomes less negative with increasing electrolyte
concentration, because salt addition decreases the electrostatic attraction part $\Delta G^0_{1,s,\text{el}}$. The
effect of salt addition on $\Delta G^0_{1,s}$ is opposite to that on $\Delta G^0_{1,\text{lat}}$, because for $\Delta G^0_{1,\text{lat}}$
(=ΔG_{1,lat-hydrophobic}^0 + ΔG_{1,lat-el}^0) the salt effect is favorable (repulsion between surfactant head
groups decreases). The two salt effects compensate each other at the CIP. The ΔG_{2,s}^0
(second layer) values are hardly dependent on the salt level, because the second layer starts
when the particle potential is close to zero (CIP ≈ IEP); therefore, to a good approximation,
ΔG_{2,s}^0 ≈ ΔG_{2,s-chem}^0 ≈ ΔG_{2,s-hydrophobic}^0, the standard hydrophobic attraction between layer 2
and layer 1 (which is the new s). However, the lateral attraction in layer 2, ΔG_{2,lat}^0,
(=ΔG_{2,lat-hydrophobic}^0 + ΔG_{2,lat-el}^0) also contributes and increasing the salt concentration from
20 to 100 mM causes a decrease of ΔG_{2,lat-el}^0 as a result of a decrease in the electrostatic
repulsion between the head groups and, in turn, this increases net attraction and hence the
adsorption.

The sum (ΔG_{1,s}^0 + ΔG_{1,lat}^0) turns out to be hardly temperature dependent. This shows that
the independence of the temperature of the adsorption in the first layer is caused by internal
compensation. As a result, according to the Gibbs-Helmholtz equation, ΔH_{1,s}^0 ≈ 0. However, for the second layer the sum (ΔG_{2,s}^0 + ΔG_{2,lat}^0) is smaller at 20 °C than at 60 °C, in line with
the exothermic nature of the process over this temperature range. At 20 °C the adsorption
enthalpies, ΔH_{1,s}, are much smaller than ΔG_{1,s}^0 indicating that entropic contributions dominate
at both electrolyte concentrations. Apparently, for both layers the enthalpies of hydrophobic
bonding of the tails and those due to interactions of the head group and/or change of hydration
at the surface compensate. At 60 °C the entropy has decreased, but the (exothermic) enthalpy
renders the standard Gibbs energies virtually temperature independent. This is another
example of enthalpy entropy compensation which is characteristic for hydrophobic bonding
and which has also been found for the solubility of alkanes in aqueous solution [182, 183].

Xu and Boyd [184] have obtained very similar C16TAC isotherms (lin-log) on kaolinite
(Source Clay Minerals Repository, University of Missouri, Columbia) at pH 6.5 and two
salt concentrations (2 and 42 mM), see Fig. 21. The (C)IP of the isotherms is found at about
20 μmol/g (0.5CEC). The reported CEC value at pH 6.5 was 40 μmol/g; this would imply
that about 50 % of the CEC is due to the plate surfaces. Although the authors measured
electrophoretic mobilities and dispersion stability of montmorillonite, it is a pity that they
did not state such results for kaolinite, because this might have shown the correspondence
of their (C)IP with the IEP or the minimum in dispersion stability.

Wang et al. [185] studied the adsorption of a series of alkyl-trimethylammonium
bromides (RR\_N(CH\_3)Br) with different chain lengths on sodium kaolinite (Source Clay
Minerals Repository, University of Missouri, Columbia) by micro-electrophoresis and
adsorption isotherms. The isotherms (lin-log) were measured over a relatively large
concentration range and have a similar shape than those measured by de Keizer et al. [143],
Mehrian et al. [179] and Xu and Boyd [184]. At very low concentrations the adsorbed
amount increased only slowly with increasing surfactant concentration; the second stage of
adsorption occurred at higher surfactant concentrations below the CMC and a sharp increase
of adsorption was observed. Adsorption in the second stage is accompanied by a sharp
change in the $\zeta$-potential, charge reversal and second layer formation. The third region is
located around the CMC with a constant adsorption above the CMC, where the monomer
concentration and the adsorption remain constant. For all surfactants the amount adsorbed at
the IEP is about 10-12 $\mu$mol/g. Though kaolinite came from the same source as that of Xu
and Boyd [184], this value is about half the value reported by Xu and Boyd [184]. The CEC
of the kaolinite is not reported in the paper so no further comparison can be made between
the adsorption at the IEP and the CEC. An increase in the number of CH\_2 groups in the
shorter alkyl chain of the double chain surfactant, or the introduction of another kind of
alkyl group (phenyl group), was found to have much less effect on the adsorption behavior
than changing the length of the alkyl chain in the single-chain surfactants. This corresponds
with the trend in CMCs and with the findings of Böhmer et al. [135] for surfactant
adsorption on rutile. Intra-molecular association reduces the effective hydrophobicity of the
shorter chains in the double-chain surfactants.
A further study on chain architecture and adsorption on kaolinite was made by Qi et al. [186] who compared the adsorption of C16TAB with a C16-2-C16 gemini surfactant of similar structure (2 CH₂ segments as spacer) at pH 6 and a ionic strength of about 0.02 M salt. Compared to C16TAB the affinity of the gemini surfactant was higher and the isotherm was steeper, but the maximum adsorptions in µmol/g at the CMC were rather similar. The maximum adsorption was about 1.4CEC, so well above the charge compensation point, which provides evidence for the presence of the second layer.

An alternative approach to detect the onset of the head-out adsorption and the ratio head-out / head-on adsorption (maximum adsorbed amount in the second ‘layer’ relative to that in the first ‘layer’) has been introduced by Li et al. [187-189]. The somewhat simplified notion of the adsorption behavior is that first complete cation exchange occurs due to head-on surfactant adsorption (maximum first layer surfactant adsorption = CEC) and that subsequently further surfactant is adsorbed in the head-out mode by hydrophobic attraction (second layer adsorption). In the absence of added background electrolyte, surfactant bound head-on by ion exchange will loose its counterion and the cations that were adsorbed to the negative sites on basal planes will be released. For the second layer, formed by hydrophobic attraction, the mechanism is different: it leads to head-out surfactant adsorption and these surfactant ions will still require counter ions in the diffuse layer to compensate their charge and as the surfactant counterion is the only negative ion present except for some OH⁻, the surfactant ion adsorption will be accompanied by adsorption of its counterion. Thus, by comparing the surfactant adsorption with the counterion release from the clay (or with the CEC) and with the surfactant counterion co-adsorption a distinction can be made between first layer and second layer surfactant adsorption. This type of investigation goes back to the classical study of Bijsterbosch [148] of the counterion binding to C16TAB and C12TAB on silica.
Li and Bowman [187] studied kaolinite (KGa-1; Source Clay Minerals Repository, University of Missouri, Columbia, MO) and a zeolite, clinoptilolite, and observed (1) that the maximum adsorption of C16TAX (X= Br⁻, Cl⁻ or HSO₄⁻) was in all cases larger than the CECs of the minerals, which implied that the adsorption was in excess of ion exchange and (2) that the maximum adsorption increased in the order HSO₄⁻<Cl⁻<Br⁻ which showed the surfactant counterion effect for the adsorption in the second layer. At the adsorption maximum the ratio surfactant counterion adsorption / surfactant adsorption ranged from 0.33 to 0.55 in going from HSO₄⁻ to Br⁻, which is a clear indication of the formation of a second layer and that the counterion affects the amount of surfactant that is adsorbed head-out.

In a subsequent study Li and Gallus [188, 189] investigated C16TAB adsorption on two kaolinites, KGa-1b and KGa-2 (Clay Mineral Repository; Purdue University, West Lafayette, IN, USA); the CECs determined by an ammonia method were 30 and 37 mmol/g, respectively. Relatively crude surfactant and counterion isotherms were measured, together with the accumulative cation desorption. The discussion of the results is not always convincing, but the cation release as function of the surfactant adsorption provides clear information about the situation up to the CMC. Around the CMC the maximum amounts of C16TA⁺ adsorbed of about 58 and 85 µmol/g were reached and the corresponding amounts of cation released were about 18±2 µmol/g and 11±2 µmol/g for KGa-1b and KGa-2, respectively. However, at about 10 µmol/g surfactant adsorption the total cation release was already about 90% of the maximum release, and at 30-40 µmol/g nearly all the cations were released. This indicates that the adsorption in the second layer started at adsorption values of about 10-15 µmol/g; upon further surfactant adsorption a small fraction of the ions will still be head-on adsorbed, but the majority of surfactant ions was adsorbed head-out (second layer). Therefore, the head-out adsorption started at surfactant concentrations clearly below the CMC and the contribution of the head-out adsorption (by hydrophobic attraction) was
far greater than that of the head-on adsorption (by cation exchange) and this means that asymmetric admicelles were formed. In the kinetic experiments the adsorption of bromide ions differed by about 15 µmol/g from the surfactant adsorption, which is close to the above estimate of the head-on adsorption. FTIR measurements showed a consistent shift of C–C symmetric and asymmetric vibration from high to low frequencies, which supported the view of a transition from head-on adsorbed monomers to admicelles as the C16TAB adsorption increased.

Li and Gallus [189] also investigated the effect of surfactant chain length (C12TAB and C16TAB) and surfactant mixtures on kaolinite. As expected the surfactant chain length had no effect on the head-on adsorption (= cation exchange or first layer adsorption), but it did affect the adsorbed amount in the head-out mode (second layer). For C16TAB the head-out adsorption was considerably larger than the head-on adsorption, but for C12TAB the situation was reversed. For the 50/50 mixture the adsorbed amounts in head-on and head-out mode were comparable. Regardless of how much alkyl-ammonium adsorbed at the adsorption plateau, the amounts of cations desorbed were 15-18 µmol/g for KGa-1b and 9-11 µmol/g for KGa-2, respectively. Therefore, the CEC of KGa-1b and KGa-2 as determined by alkyl-ammonium adsorption/cation desorption will be 15-18 µmol/g, and 9-11 µmol/g, respectively. These values are much less than the values determined by an ammonia method [190] or the reference values listed on Clay Mineralogical Society website (http://cms.lanl.gov/chem.htm). However, the value of cations desorbed from KGa-1b through surfactant adsorption was similar to the CEC values of KGa-1 determined by a complexation method using copper bisethylenediamine and copper triethylenetetramine in the presence and absence of the buffer tris(hydroxymethyl)- aminomethane [191]. Comparing this with the results of Mehrian et al. [179] would imply that the difference should be related to the CEC of the plates as compared to the total CEC (plates + edges).
Recently, Malek and Ramli. [192] investigated C16PB on kaolinite with respect to antibacterial activity. Based on X-ray diffraction they first concluded that the structure of kaolinite was not affected by the surfactant adsorption. Although the interpretation of the results by the authors contains several errors, it is interesting that the antibacterial activity started after the IEP, i.e., when the surfactant started to adsorb in the head-out mode that made the overall particle charge positive, which must have caused the action of C16P⁺ against the bacteria cells.

6.3 Anionic surfactant adsorption on kaolinite

Adsorption of anionic surfactants to kaolinite also occurs and for this adsorption the amphoteric edge sites are crucial. The point of zero net proton charge of the edge of kaolinite depends on the ionic strength and kaolinite composition, and is found to be in the pH range 4 to 7 [193]. Therefore, the net proton charge of the edge is positive at relatively low pH and negative at relatively high pH. The investigations of anion adsorption have mainly been carried out at relatively low pH, which can lead to some dissolution of kaolinite and surfactant precipitation so that next to surfactant adsorption also surfactant (surface) precipitation occurs. The total amount of surfactant removed from the solution is therefore called the surfactant abstraction. Somasundaran et al. [168-170, 194, 195] have investigated these aspects in detail in their studies of sodium dodecyl benzenesulfonate and sodium dodecyl sulfonate abstraction by kaolinite. Results obtained under various ionic strengths and pH values indicated a complex abstraction mechanism involving surfactant adsorption by ion exchange and hydrophobic attraction and metal activated adsorption and surface precipitation of surfactant, all depending on the solution pH. The precipitation of sulfonate with dissolved mineral species results in an adsorption maximum in micellar solutions. The dissolution of kaolinite and the resulting solution speciation can be found in [169]. Results for the abstraction of sodium dodecyl benzenesulfonate on Na-kaolinite
(Georgia kaolinite from the clay repository at University of Missouri subjected to an ion-exchange treatment) as a function of pH at 0.1 M NaCl and 25 °C as obtained by Siracusa and Somasundaran [195] are summarized in Fig. 22. The authors made the following remarks concerning the abstraction (adsorption) mechanisms. (1) At acidic pH levels below pH 3.7 significant precipitation of the dodecylbenzene sulfonate with cationic dissolved aluminum species occurs and subsequent re-dissolution of the aluminum-sulfonate precipitates above the CMC produces an abstraction maximum. (2) At or near pH 4.4 (the pznpc of the sample) a reduction in the concentration of dissolved cationic aluminum species results in decreased precipitation such that the adsorption maximum is no longer observed. However, some contribution of aluminum-sulfonate precipitation is still present. (3) At pH levels above pH 4.4, both in the neutral region (pH 7.9) and the alkaline region (pH 10.8), incongruent dissolution of kaolinite can result in phase transformations to gibbsite; therefore, adsorption occurs at levels governed not only by the surface of the kaolinite, but also by that of gibbsite.

Very similar pH dependent results have been obtained by Torn et al. [196] for the dodecylbenzenesulfonate / Na-Kaolinite (Sigma Company) system at 0.01 M NaCl and 25 °C, but in this case the abstraction maximum occurred at pH 4.8-4.9. Proton titrations of the kaolinite sample suggested a pznpc of about 7 [197], which is higher than the pznpc of the kaolinite used by Siracusa and this might explain the shift of the maximum. Torn et al. [196] concluded that both electrostatic and hydrophobic interactions play a role in the adsorption of the sulfonate surfactant to kaolinite.

Poirer and Cases [167] have studied the effect of chain length (C14 and C12) and the presence of a benzene ring in the apolar chain (C10Φ) of the sulfonate surfactants on their adsorption to kaolinite. The adsorption isotherms at 28 °C showed a sharp increase in abstraction close to the saturation concentration of the surfactants, which was due to the precipitation of the sulfonates. The isotherms were shifted toward lower equilibrium
surfactant concentrations as the hydrophobic moiety of the surfactant increased. The shifts were due to the increase in hydrophobic attraction and corresponded with the differences in CMC with surfactant structure. The benzene ring was about equivalent to three methylene groups. Del Hoyo et al. [166] investigated the adsorption of SDS on kaolinite by FTIR measurements and observed modification of the CH stretching wavenumbers of SDS indicating a re-arrangement of the adsorbed surfactant molecules and hydrophobic attraction between the hydrocarbon chains of the surfactant.

How the structure of the adsorbed surfactant layer evolved as a function of the surfactant concentration has not been discussed in the above mentioned studies, but most likely also in this case head-on adsorption occurs at low adsorption values and head-out adsorption at surfactant concentrations closer to the CMC.

6.4 Nonionic surfactant adsorption on kaolinite

The adsorption of nonionic surfactants on kaolinite resembles that on silica. Also on kaolinite the surfactant monomers adsorb by hydrogen bonding between the polyoxyethylene segments and the hydroxyl groups on the surface. In addition interaction may occur with the hydrated cations on the siloxane basal plate as is observed for montmorillonites [198]. The energy involved in the initial adsorption process is somewhat stronger than with silica. Similar as for silica, the energetic balance at the beginning of the adsorption is exothermic, because contacts between the surface and the ethoxy segments of the polar chains are enthalpically favorable. When the coverage of the surface increases, the enthalpy decreases down to the appearance of an endothermic regime. In the latter regime the molar enthalpy of adsorption is comparable to that of micellization in aqueous solution and this is characteristic for the presence of surfactant admicelles (or two adsorption layers) on the surface [161]. The temperature dependence of the adsorption [161, 199] is in agreement with this adsorption mechanism. Two steps can be observed along the adsorption
isotherms [144, 161, 199, 200], some typical isotherms are depicted in Fig. 23. The first
pseudo-step occurs at low coverage and is followed by a rapid but gradual rise of the
adsorption up to the CMC, somewhat above the CMC the adsorption plateau is reached.
Due to the polydispersity of most nonionic surfactants the plateau starts somewhat beyond
the CMC. The second step corresponds with a cooperative adsorption mechanism and the
formation of admicelles. The relative importance of the first step increases with the EO
chain length (stronger adsorption), but the adsorption level at the CMC strongly decreases
with EO chain length because the large head group makes it more difficult for the apolar
chains to associate by hydrophobic attraction. The plateau adsorption is reached at higher
surfactant concentrations for longer EO chains. The reason for this is that when oxyethylene
length rises, the monomer solubility in water is higher and, thus, the CMC is also higher.

For different kaolinite samples (Supreme, Charentes, Aldrich, Sigma) the isotherms
have a similar shape, but the adsorption levels of the first step and the final plateau around
the CMC differ [200], which implies that the structural organization of the surface
aggregates is dependent on the surface heterogeneity of the kaolin crystals. Based on the
characteristics of the kaolinite samples and the adsorption results no final conclusion could
be reached regarding the role of the edge surfaces and the basal surfaces.

6.5 Anionic surfactant adsorption on montmorillonite

The adsorption of surfactants to montmorillonite differs from that of kaolinite for two
reasons. (1) Kaolinite has two different basal plate surfaces and a relatively large edge
surface, while the basal plate surfaces for montmorillonite are the same and the edge area is
relatively small. (2) Montmorillonite is an expanding clay-type, which implies that the
adsorption capacity may increase with increasing surfactant adsorption, depending on the
conditions. Anionic surfactant adsorption will mainly occur on the edges and the anions will
not easily intercalate between the clay platelets, this and the relatively small edge area of
montmorillonite make anionic surfactant adsorption on pure montmorillonite relatively unimportant. Similarly as for kaolinite, adsorption of anionic surfactants to montmorillonite is complicated by surfactant precipitation when multivalent ions are present as counterions of the negative clay charge and by partial clay dissolution and precipitation phenomena as described for kaolinite. For instance, Yang et al. [201] have shown that the abstraction (adsorption + precipitation) of sodium dodecylbenzene sulfonate (SDBS) by montmorillonite was affected by the presence of Ca$^{2+}$ ions. They observed that: (i) SDBS was abstracted significantly by montmorillonite saturated with Ca$^{2+}$, but little by Na-saturated montmorillonite; (ii) the amount of SDBS abstracted by Ca-montmorillonite was enhanced by NaCl; and (iii) no significant intercalation of SDBS into Ca-montmorillonite was observed by X-ray diffraction analysis. Therefore the abstraction for the Ca-montmorillonite was largely due to surfactant precipitation with Ca released by ion exchange.

6.6 Cationic surfactant adsorption on montmorillonite

The binding of cationic surfactants on montmorillonite is dominated by adsorption at the plate surfaces, therefore, for these surfactants adsorption in the interlayers plays an important role. The adsorption behavior at the external surface of montmorillonite will be similar to that on silica and/or kaolinite, but the adsorbed amount and the conformation of adsorbed molecules in the interlayer space are affected by the two neighboring surfaces and the distance between these surfaces (the interlayer distance). In fact one deals with adsorption in a narrow slit pore with the complication that the slit width increases step wise with increasing surfactant adsorption. The interaction between the platelet surfaces and, therefore, the interlayer distance, are affected by the surfactant adsorption and vice versa. In other words, the interlayer distance, the adsorbed amount of surfactant and the conformation
of the adsorbed surfactant molecules are mutually related and this makes the adsorption behavior complicated.

To unravel cationic surfactant adsorption in practice, next to information on the adsorbed amount, at least, also information is required on the interlayer distance and/or the conformation of the adsorbed surfactant molecules in the interlayer. Somewhat similar to Mehrian et al. [180], Pan et al. [202] measured C12TA+/C12TAB isotherms and adsorption enthalpies for Na- and Al-montmorillonites in a wide pH range; the results support the two-stage adsorption above the IEP. Very illustrative experimental results on the adsorption of hexadecyl-trimethylammonium (C16TA+ = HDTMA) on montmorillonite (Wyoming montmorillonite, SWy-1; CEC = 900 µmol/g) and kaolinite (Na-kaolinite, CEC = 40 µmol/g at pH 6.5) are provided by Xu and Boyd [184, 203]. They studied the adsorption by combining adsorption isotherms, electrophoretic mobility, colloidal stability and X-ray diffraction to determine the effects of clay type and solution composition on the adsorption of cationic surfactants and to unravel the orientation and/or conformation of the adsorbed surfactants on the external and internal (interlayer) surfaces and the dispersion stability of surfactant-clay complexes. Some of their results are depicted in Figs. 24 and 25. Fig. 24 shows the isotherms of C16TAC on three different clays; isotherms provide information on the total surfactant adsorption. With a single isotherm further information is required to make a distinction between the adsorptions on external and internal surfaces. The C16TAC adsorption isotherm on Ca-montmorillonite (Ca-SWy-1) follows the ‘normal’ isotherm pattern and is similar to that on Na-kaolinite (non-swelling), but the adsorption isotherm on Na-montmorillonite (Na-SWy-1) differs from the other isotherms with its peculiar S-shape at relatively low C16TAC adsorption (< 1CEC). The fact that the shape of the C16TAC isotherms on Na-kaolinite and Ca-SWy-1 are rather similar indicates that the swelling of C16TA/Ca-SWy-1 due to surfactant adsorption is negligible small. The peculiar S-shape of the of C16TA/Na-SWy-1 system with different adsorption values for one equilibrium
concentration must be due to interlayer expansion due to surfactant adsorption. The behavior of the C16TA+/Na+-montmorillonite clay is further illustrated in Fig. 25 to provide proof for this behavior.

The C16TA+/C16TAC isotherm on Na-SWy-1 (Fig. 25a) can be divided into four distinct regions. Region 1 (0-0.75 CEC) is non-monotonic and characterized by an equivalence of Na release and C16TA⁺ adsorption, resulting in superimposable C16TA⁺ adsorption and cation release curves. In regions 2 and 3, the adsorption isotherm is monotonic and the adsorbed amount of C16TA⁺/C16TAC is greater than the corresponding Na⁺ release; this implies that the head-on adsorption (ion exchange) slows down and the head-out adsorption increases (difference between the C16TA⁺/C16TAC and Na⁺ isotherms). The head-out adsorption is due to surfactant adsorption to already adsorbed surfactants by hydrophobic attraction between the surfactant tails. The slope of the C16TA⁺/C16TAC adsorption isotherm in region 3 is much larger than in region 2, which indicates cooperative adsorption in region 3. Moreover, the difference between the C16TA⁺/C16TAC and the Na isotherms increases strongly, indicating that head-out adsorption to already adsorbed surfactants increases much stronger than the ion exchange. At the CMC the C16TA⁺/C16TAC and Na isotherms reach a plateau (region 4).

Fig. 25b shows the behavior of the electrophoretic mobility with increasing surfactant adsorption; it reflects the situation at the external surface of the clay. In region 1 Na⁺ exchanges for C16TA⁺, therefore it is to be expected that the electrophoretic mobility of the C16TA/Na-SWy-1 particles remains the same as for homoionic Na-SWy-1; this is indeed the case. In region 2 (0.75 CEC to 1.0 CEC) the net particle charge is relatively low; more C16TA⁺ is adsorbed than Na⁺ is released and the electrophoretic mobility increases rapidly, it passes the IEP (particle charge changes sign) and reaches a maximum value at 1CEC, the end of region 2. The fact that the IEP is reached before C16TA⁺ (ads) = CEC implies that the Na⁺ ions also contribute to the charge neutralization at the IEP. The fact that the
mobility changes sign confirms that the surfactant adsorption is super-equivalent and this implies that beyond the IEP head-out adsorption by hydrophobic attraction starts. The mobility is very sensitive to changes in charge around the IEP where the particle charge is low, therefore the information in region 2 is directly related to the charge situation of the clay complex and provides valuable information. In region 3 the mobility decreases slightly and stays constant in region 4. Somewhat outside the IEP (zeta potentials > 50 mV) retardation and relaxation effects can easily lead to only small changes in mobility with particle charge [204]. Moreover, the changing particle (aggregate) sizes (see the stability) further complicate the situation. Therefore, further interpretation of the mobilities in regions 1 and 3 or 4 is not feasible.

The adsorption behavior at the external surface was further investigated by dispersion stability measurements (optical density of the clay suspension), see Fig. 25b. The degree of clay dispersion decreases with increasing surfactant adsorption till the IEP is reached. This is due to (i) the decrease of the net negative charge of the C16TA/Na-SWy-1 particles, so that the electrostatic repulsion between the particles decreases and (ii) at the same time the hydrophobicity of the particles increases because the surfactant tails point to the solution (ion exchange and head-on surfactant adsorption) and this increases the inter-particle (hydrophobic) attraction. The stability has a minimum around the IEP because the net particle charge is zero and the amount of unscreened surfactant tails pointing towards the solution is at its maximum, which makes the particles maximally hydrophobic. Beyond the IEP further C16TA+ adsorption in the head-out mode makes the C16TA+-SWy-1 particles positive and increases the particle hydrophilicity, together this results in a strong increase in the dispersion stability.

In Fig. 25c d(001)-spacings obtained by X-ray diffraction are shown for water saturated, partially wet (dried at 95% relative humidity, RH), and dry (dried at 5% RH) C16TA+/Na+-SWy-1 as function of the C16TA+ adsorption. These results reflect the
situation at the interlayer surfaces. The clays are nearly identical at high C16TA$^+$ adsorption, but completely different at low C16TA$^+$ loadings (< 1CEC). As the C16TA$^+$ loading increased from 0.1 CEC to 0.75 CEC (region 1), the d-spacings of wet (100% RH) clay samples (C16TA$^+$/Na$^+$-SWy-1) increased gradually from 1.76 to 2.21 nm. No further change in the d-spacing was observed at higher C16TA$^+$ adsorption. Partial air-drying (95% RH) or drying (5% RH) of the clays decreases the d-spacings and results in stepwise d-spacing vs. C16TA$^+$ loading curves. At loading levels up to 0.3 CEC, the d-spacing was around 1.4 nm. The first step occurs between 0.3 and 0.5 CEC and raises the spacing to about 1.8 nm, the second step occurs at C16TA$^+$ loadings of 0.5 to 0.8 CEC leads to a final d-spacing of about 2.2 nm. The steps are sharp for the dry clay. The step-wise behavior of the d-spacing indicates that the conformation of the adsorbed surfactant ions changes stepwise. Confining the interlayer to lower values of the d-spacing by drying allows only specific surfactant conformations and sudden conformational changes result. However, at 100% RH the d-spacing increases gradually, therefore the capacity for surfactant adsorption increases gradually and the actual conformation of the adsorbed surfactant is more random than for the (partially) dried clays.

With the C16TA/Ca-SWy-1 system and adsorption values of about 0.25 CEC to 0.8 CEC the d-spacing stays around 1.8 nm and then increases gradually with higher loadings to 2.16 nm (not shown). The higher charge of the Ca$^{2+}$ ions keeps the two interlayer surfaces better together and only close to C16TA$^+$ adsorption values = 1CEC some swelling is observed and the isotherm has a similar shape as the C16TA$^+$ isotherm to kaolinite. The peculiar shape of the C16TA$^+$ isotherm for Na-SWy-1 and the ‘normal’ shape of the isotherm for Ca-SWy-1 is thus due to the fact that in the adsorption range 0-0.8 CEC swelling occurs for Na-SWy-1 and no swelling for Ca-SWy-1. Due to swelling and a concomitant change of the adsorption capacity in the case of the C16TA/Na-SWy-1 system different adsorption values can occur at one equilibrium surfactant concentration.
When for the C16TA/Na-SWy-1 system the ionic strength is increased by increasing the NaCl concentration, the isotherms gradually change to the ‘normal’ shape at about 42 mmol/L NaCl. At this ionic strength the d-spacing stays about constant up to 0.8 CEC. Around loadings close to the CEC the isotherms at different NaCl concentration intersect and after the intersection point (loadings larger than the CEC) the adsorption increases with increasing NaCl concentration. The residual water in the interlayer allows salt ions to enter the interlayer, and similarly as at the external surface, this leads to an increased screening of the lateral repulsion between the charged head-groups of super-equivalently bound surfactant ions and an increase in adsorption. For the same reason also surfactant counterions that partially associate with the surfactant ions increase the surfactant adsorption for loadings beyond the CEC. The NaCl concentration has little effect on the C16TA/Ca-SWy-1 system at C16TA⁺ loadings up to the CEC and after the CEC the behavior is similar to than of the C16TA/Na-SWy-1 system.

Concluding, based on the results of Xu and Boyd [184] it follows that the swelling of expanding clays due to cationic surfactant adsorption depends on the strength at which the clay layers are held together. It is well known that the attraction between the clay layers depends on the degree of isomorphic substitution, the valence and size of the counterions of the clay charge and the ionic strength [205]. Therefore, also the swelling of clays due to cationic surfactant adsorption is depending on these variables. Swelling occurs when the surfactant isotherm shows a peculiar S-shape for surfactant loadings up to 0.8 CEC; the adsorption capacity is a function of the adsorbed amount and this makes it possible that different adsorption values may occur at one equilibrium surfactant concentration. The surfactant adsorption at the external surface of expanding clays shows a very similar behavior to that at kaolinite. Up to loadings close to the IEP of the external particle surface (roughly corresponding with a loading approaching 1 CEC) the surfactant adsorption occurs by ion exchange and head-on adsorption. Beyond the IEP of the external surface (or
loadings $\geq 1$ CEC), adsorption of the surfactant by hydrophobic attraction starts and this adsorption increases strongly with further surfactant loading. The adsorption beyond the 1 CEC increases with increasing ionic strength and is sensitive to the kind of surfactant counterion. The clay stability is strongly affected by this adsorption behavior. The minimum dispersion (colloidal) stability occurs at the IEP of the external surface where the net particle charge is zero and the hydrophobicity of the particles is at its maximum due to the unscreened surfactant tails protruding into the solution. The results of Xu and Boyd are largely supported by Tahani et al. [206] who investigated the adsorption of the benzyldimethyl dodecyl ammonium chloride on a Na-montmorillonite by various techniques.

An interesting complementary study, specifically oriented towards the structural characteristics of the surfactant adsorbed in clay interlayers, has been presented by Zhu et al. [207] who studied the interlayer aggregates of C16TA$^+$/C16TAB in a bentonite (> 95% pure montmorillonite, < 5% quartz; CEC 1084 $\mu$mol/g) for different surfactant loading levels (0.20 CEC to 2.56 CEC), using in situ X-ray diffraction (XRD) and Fourier Transform Infra Red (FTIR) spectroscopy on samples in the hydrated state. For the dried and hydrated organo-bentonites, the measured basal spacings depended on surfactant loadings and indicated that the water molecules present play an important role in regulating the microenvironment, even at high surfactant loads. In the presence of water the surfactant tails formed aggregate structures in the interlayer space due to hydrophobic attraction. As a result, the values of the d001 spacing increased gradually with increase in surfactant loading. In situ FTIR spectra indicated that the surfactant intercalated at high loading had a more ordered and compact structure than the associates at low surfactant sorption density: the surfactant state in wet organic bentonites changed from a liquid-like (disordered) state to a solid-like (ordered) state.
Based on the FTIR results, the XRD data, and the dimensions of the C16TAB surfactant in moist organo-bentonites with different surfactant loading, Zhu et al. proposed that C16TA+/C16TAB arrangements in wet organo-bentonites were different from that in dried organo-bentonites. For the wet organo-bentonites the interlayer spacing around 0.2 CEC is 1.02 nm and the C16TA+ ions lie about parallel to the silicate interlayer surfaces. Then a smooth step in the interlayer spacing occurs and at 0.4 to 1.2 CEC the spacings of 2.0 - 2.5 nm are roughly the length of C16TA+. Therefore, a possible arrangement could involve the C16TA+ head groups being adsorbed on the two interlayer surfaces and the ‘interfingered’ long alkyl chains roughly perpendicular to the interlayer surfaces. When the loading level further increases (1.2 to 2.6 CEC), a next smooth step in interlayer spacing occurs to about 3.3 nm and the C16TA+ ions (ion exchanged) and C16TAB molecules (hydrophobically attracted) form a partly ‘interfingered’ admicellar type layer. With these surfactant conformations at loadings > 0.4 CEC the surfactant phase in the interlayer is divided into three regions: region 1 and 3 are adjacent to the two interlayer surfaces, each extending to about 0.5 nm from the surface, containing H2O, surfactant head groups and, if loaded in excess of the CEC, also the surfactant counterions (Br-), and region 2 with a thickness of about 2 nm in the center of the interlayer space, containing the aggregated tails of the alkyl chains and residual water. The total structure is bilayer-like and/or resembles flattened admicelles. The core region with the tails becomes more aliphatic and less aqueous as the surfactant loading increases from 0.40 to 2.6 CEC. With increasing surfactant loading the core region becomes more hydrophobic, but induced steric hindrances increase and the space left for ad-solubilization decreases; therefore, the partitioning of organic contaminants shows a maximum around loading levels of 1 CEC. This maximum corresponds roughly with the maximum hydrophobicity of the surfactant layer at the external surfaces of the clay around loadings of 1 CEC, but the structure of the external surfactant layer is rather
different (head-on monolayer type and/or teepee-like micelles) from that in the interlayer (flattened admicelle-like).

The specific behavior of surfactant adsorption in confined space has also been investigated theoretically by Molecular Dynamics studies. Initially these studies modeled dry organo-clays [177, 208-210] but in later studies water was included in the simulations [211, 212]. The latter are most relevant with respect to understanding the adsorbed surfactant conformations in the interlayer space. The introduction of the paper by Liu et al. [178] provides a brief overview of the history of MD calculations on organo-clays and the force fields used; in that of Zhou et al. [213] the most relevant experimental results and the recent MD simulations are mentioned. Liu et al. [178] discuss the structure and dynamics of alkylammonium-intercalated wet SWy-type montmorillonite with emphasis to the alkyl chain length; the paper is complementary to the recent paper of Zhou et al. [213], which will be discussed here. The study of Zhou et al. [213] on C16TA+/C16TAB intercalated Ca-montmorillonites (Mt) in the water-saturated condition largely mimics the experimental system of Zhu et al. [207] discussed above, and provides a good illustration of the information that can be gained by MD simulations.

The simulation results of Zhou et al. [213] show that, as the surfactant adsorption on montmorillonite (CEC 1060 µmol/g) increases, the arrangement of C16TA+/C16TAB transforms from flat bilayer-type to inclined paraffin-type with a large amount of water in the interlayer space. This configuration is different from that in dry systems; for dry samples the surfactant conformation changes from monolayer to bilayer and to pseudo-trilayer as surfactant loading increases [171, 175]. Compared to the dry models, the percentage of gauche conformations of C16TA+ decreases in the water saturated condition. The head groups of C16TA+ are located close to the centers of the six-member rings of the silicate surfaces and coordinated with 4–6 water molecules. Their mobility is low due to the electrostatic interactions, while the alkyl chains show a higher mobility. Montmorillonite
with a high CEC (1310 μmol/g) has a stronger confining effect on both C16TA+/C16TAB and water, which reduces the mobility of alkyl chains and water molecules within the interlayer space.

A general impression of the surfactant conformations in the interlayer space as function of surfactant loading is provided by the density distributions for ammonium N, alkyl C, Ca²⁺, Br⁻ and water O [213]. The calculated density distributions for montmorillonite with a CEC of 1060 μmol/g are depicted in Fig. 26 for different loading levels relative to the CEC as indicated in the figure. Corresponding snapshots of C16TA⁺ intercalated in the interlayer that further visualize the distribution are presented in Fig. 27. For Mt-0.25CEC-1nm (= montmorillonite with a C16TA⁺/C16TAB loading equal to 0.25 CEC and an interlayer spacing of about 1 nm) both ammonium N and alkyl C are arranged in two layers according to the two clear peaks of the density distributions, indicating a bilayer structure, see Figs. 26a and 27a. As the surfactant loading level increases (Mt-0.5CEC-2nm; Figs. 26b and 27b), the density distribution of ammonium N remains two peaks close to the silicate surface due to the strong electrostatic attraction between the negative charge site and the cationic ammonium N. The distribution of alkyl C is rather different from that at 0.25 CEC, it transforms to one broad peak in the center and the arrangement of alkyl chains is inclined to the Mt surface. In these two incomplete cation exchange cases Ca²⁺ displays a symmetrical distribution with ammonium N to balance the residual negative charge. Ca²⁺ plays no role for surfactant loadings > 1.0 CEC, but in this case the negative surfactant counterion (Br⁻) comes into play. For Mt-1.0CEC-2.2nm (Figs. 26c and 27c), the situation is similar as for Mt-0.5CEC-2nm, but more pronounced. The broad central peak of alkyl is larger, indicating strong association of the alkyl chains in the interlayer space; the arrangement of alkyl chains is still inclined to the Mt surface. In the excessive surfactant loading case (Mt-2.0CEC-3nm; Figs. 26d and 27d), most N is still close to the surfaces, but part of N is distributed over the core of the interlayer. The N distribution is roughly paralleled by the Br distribution, but
close to the surface N and Br occupy separate layers. The magnitudes of the N and Br peaks
close to the surface indicate that a large part of head groups that did not participate in the
ion exchange is also located close to the surface. The small peaks of ammonium N in the
interlayer space are balanced with Br in the core of the interlayer space. The alkyl chains are
connected closely, reflecting the association by hydrophobic attraction. In all cases, the
density of water near the Mt surfaces is somewhat higher than in the center, which is
probably related to the hydration of the accumulated ions and ammonium head groups close
to the surface. The hydrophilicity of the interlayer surfaces is somewhat questionable,
because a siloxane surface without charge is fairly hydrophobic. The density distributions of
interlayer species for Mt with a high CEC (1310 µmol/g) are in most cases similar as those
shown here. However, for the Mt-0.25CEC-1nm system, the alkyl C forms several peaks
near the center of the interlayer space instead of two peaks. The ammonium head groups
close to the surface cause most likely obstruction of the orientation of the alkyl chains
parallel to the surface. To sum up, under the water saturated condition and low surfactant
loading the surfactant alkyl chains are oriented roughly parallel to the surface; at loadings ≥
0.5 CEC alkyl chains are concentrated in the center of the Mt interlayer space and most
head groups are close to the surface. The calculated results are consistent with the
experimental results of Zhu et al. [207] and the density profiles are subtler versions of the
schematic models suggested by Zhu et al.

An alternative way of modeling surfactant binding from aqueous solutions in confined
space is by Self-Consistent Field calculations with the SCFA model [214-217]. With the
SCFA model surfactant isotherms can be calculated for two surfaces at a given distance and
the interlayer spacing can only be changed by discrete step of about the diameter of a water
molecule; simultaneous with the adsorption isotherm the interaction force between the two
surfaces can be calculated. Calculations made [214-217] have not addressed the specific
clay-surfactant systems, but some knowledge of the obtained results is still useful for a
better understanding of the complications that occur with surfactant adsorption in confined space. For charged hydrophilic surfaces the SCFA calculations have provided insight in the adsorption behavior below and above the charge compensation point [214, 215]. The confined space drives the adsorbed surfactant layer toward a structure similar to that obtained at the charge compensation point. At solid-solid separations where the surfactant layers on each surface begin to merge with each other (proximal adsorption), there is a confinement-induced phase transition leading to complex adsorption and interaction behavior at very small separations. In addition, calculations show that the interaction between surfactant tails is a key contributor to both the magnitude of proximal adsorption and the exchange between surfactant and co-ions at the surface. For (charged) hydrophobic surfaces [216, 217] a first-order phase transition takes place when the slit width approaches the thickness of the two surfactant layers; the transition is driven by the unfavorable hydrophobic-water contacts. At the transition, the average orientation of the surfactants switches from a high concentration of tails at the surface and head groups in the center to a bilayer configuration where the tail profiles from both sides merge in the center and the head groups are in the vicinity but slightly away from the surface. For more information the original papers should be consulted.

6.7 Nonionic surfactant adsorption on montmorillonite

Similarly as with cationic surfactants the binding of nonionic surfactants on montmorillonite is dominated by adsorption at the plate surfaces, therefore, also for these surfactants adsorption in the interlayers plays an important role. Detailed results on the adsorption of nonionic surfactants on montmorillonite at low surfactant concentrations are scarce; recent results mainly concentrate on concentrations around and above the CMC because the interest is on preparation of organo-clays made with nonionic surfactants. Rheinländer et al. [218] have investigated C12(EO)8 on four different clays
(Na-montmorillonite = Na-Mt, Ca-bentonite (95% montmorillonite) = Ca-Mt, Na-kaolinite and Na-illite) based on adsorption isotherms and for the swelling clay minerals (Na-Mt and Ca-Mt) microcalorimetry and X-ray diffraction. The measured isotherms were presented as double logarithmic plots that are reproduced in Fig. 28. The isotherm on Na-kaolinite runs below that on Na-illite, and both isotherms (mmol/g) are clearly below those on the swelling minerals. However, when the adsorbed amount is expressed per m² the maximum adsorption values on the Na-clays are similar and about twice as high as that on Ca-Mt. The isotherms for Na-Mt and Ca-Mt are similar; however, the adsorption for Ca-Mt is stronger at low concentrations than that of Na-Mt, but at high concentrations the situation is reversed (maximum adsorptions Na-Mt and Ca-Mt, respectively, 0.94 and 0.35 mmol/g). A greater (exothermic) initial differential molar enthalpy (heat) of adsorption confirms the higher affinity of C12(EO)8 for Ca-Mt than for Na-Mt. The heat of adsorption of the two clays decreases about linearly with surfactant loading and becomes zero above 0.26 mmol/g, which is below the adsorption maximum of the two clays. The behavior of the heat of adsorption indicates two steps in the adsorption process: (i) adsorption due to interaction with the surface and (ii) adsorption by mainly hydrophobic attraction with already adsorbed surfactant. The observations that zero heat of adsorption to Ca-Mt and Na-Mt is reached at the same loading (mmol/g), but that the maximum amounts adsorbed are rather different indicates that the second adsorption step is larger for Na-Mt than for Ca-Mt. Although the adsorption process contains two steps, the shape of the log-log isotherms indicates that there is no critical surface association concentration (CSAC), all isotherms show a gradual increase in adsorption and the slope is of the isotherms is always ≤ 1. Therefore, the transition of the first to the second adsorption process is gradual. This is the case for all four clay samples studied.

The X-ray diffraction results showed that the basal (d001) spacing of pure air-dried Na-Mt of about 1.25 nm differed significantly from that of Ca-Mt of 1.5 nm due to stronger hydration of the Ca²⁺ than of Na⁺. The basal spacing of Na-Mt with surfactant goes from
1.25 nm to 1.4 nm for 0.1-0.35 mmol/g and to about 1.8 nm for higher adsorption values.

For Ca-Mt the spacing changes from 1.5 to 1.8 nm between 0.2 and 0.3 mmol/g. These values are for both swelling clay minerals indicative of the formation of flat-lying C12(EO)8 molecules in the first adsorption step and of the formation of a ‘flat’ bilayer structure in the second adsorption step. As the spacings are derived from air-dried samples, the distances will be likely different in the water saturated situation (and therefore the surfactant conformation), but the two steps in the adsorption process are reflected in the measured values. The stronger hydration of Ca\textsuperscript{2+} than of Na\textsuperscript{+} has, most likely, also caused the higher affinity of C12(EO)8 for Ca-Mt than for Na-Mt.

Sonon and Thompson [219] investigated the adsorption of C12(EO)<23> (=Brij 35; the notation <n> indicates that the EO chain length is polydisperse) on two smectites (Wyoming montmorillonite and Panther Creek smectite) that were brought in the K and Ca form. For both smectites the adsorption affinity and adsorption plateau were somewhat larger for the Ca-smectite than the K-smectites. The d001 values of both air-dried K-saturated smectites increased in two steps from 1.11 nm to about 1.7 nm as more surfactant was sorbed, while the d001 spacing of the air-dried Ca-smectites increased gradually from 1.47 nm to about 1.7 nm. These results confirm those of Rheinländer et al. [218].

Backhaus et al. [220] continued the work of Rheinländer and investigated the adsorption of the octylphenol poly(ethylene oxide)s [(C8Φ(EO)<n>], TX100 (<n> = 9.5), TX-165 (<n> = 16) and TX-305 (<n> = 30), on Ca-Mt and silica. Note that the hydrophobic character of a C8Φ group is very similar to that of a C12 group. The adsorption isotherms of C8Φ(EO)<9.5> (= TX100) on Ca-Mt at pH 4 and pH 9 in 10\textsuperscript{-2} mol/L CaCl\textsubscript{2} showed only small differences, which indicated that the share of the edge surfaces to the surfactant adsorption on Ca-Mt is small. Furthermore, the affinity (measured by the initial slope of the isotherm) of C8Φ(EO)<9.5> at pH 4 to Ca-Mt was higher than that on silica, which point to a different binding mechanism. With silica the EO segments mainly bind to the silanol groups
by H-bonds, but with expanding layered silicates these groups are hardly present and the
binding mechanism is most likely H-bonds with the hydration water of the bound counterions
on the clay surfaces. The interlayer spacings of Ca-Mt with adsorbed C8Φ(EO)<9.5> in the
dried and moist state show a similar behavior as in the case of C12(EO)8 [218], but the
spacings for the moist sample are even at the largest adsorption values considerably (2nm)
larger than the spacings of the air-dried sample. This indicates again that in aqueous solution
the interlayer spacings will be different from that of the air-dried samples.

The isotherms (non-logarithmic) of the three surfactants (TX100, TX-165, TX-305) [220]
were all L2-type [221], i.e. a steep rise towards the maximum adsorption. For TX-100
adsorption on montmorillonite or bentonite the L2 type isotherm has also been observed by
[218, 222, 223]. Somewhat above the CMC the isotherms show a plateau that decreases with
increasing EO head group length. The fact that the plateau adsorption is reached beyond the
CMC is due to the fact that head group polydispersity leads to a relatively low CMC and a
maximum adsorption at relatively high concentration [224]. The lack of the S-shaped course
and thus of cooperative effects suggests that, unlike in the case of silica gel, no aggregate
formation took place on the surface at low surfactant concentrations. This behavior is to be
expected when the affinity of the surfactant molecules to the surface (hydrated Ca ions) is
greater than the hydrophobic attraction between the surfactant molecules. With the relatively
small interlayer spacing, the second step caused by hydrophobic attraction will be largely
lateral (tail-tail) attraction. For steric reasons the lateral hydrophobic attraction decreases by
the head group size and this explains the decreasing plateau adsorption with increasing head
group size.

A decrease in the adsorption with increasing head group size for a given alkyl chain length
has also been observed by Shen [225] who measured the adsorption of C9(EO)<10>,
C9(EO)<20> and C18(EO)<20> (= Brij 78) on bentonite at pH 6-7 and 0.01 M NaCl.
Doubling the alkyl chain length at a given degree of ethoxylation increased the adsorption
maximum also but the effect was much smaller than that of doubling the head group size, especially when the adsorption was expressed as mmol/g. Deng et al. [226] who studied the adsorption of C16(EO)<10> (= Brij-56) and C9Φ(EO)<12> (= Igepal-CO720) on Ca-bentonite found somewhat larger adsorption for the surfactant with the smaller head group in line with the results of Shen [225]. However, the proposed surfactant conformation for the freeze-dried composites with EO segments in the interlayer and the alkyl chains sticking-out in air, is highly unlikely for water saturated samples and for the situation of the intercalated smectite in contact with an aqueous surfactant solution, because in this case hydrophobic attraction comes into play and the proposed conformation would lead to many unfavorable CH2-water contacts which can be avoided when the alkyl chains associate in the interlayer space. Therefore, it is to be expected that with adsorption from solution the conformation will be somewhat similar to that of cationic surfactants in the interlayer as described in the previous section.

To unravel the EO binding mechanism and the EO chain conformation of intercalated polymeric nonionic surfactants Deng et al. [198] also investigated the intercalation of bentonites (with different cations) by: C16(EO)<10> (= Brij 56), C18(EO)<100> (= Brij 700), C2(CH2-CH2)<15>(EO)<40> (= polyethylene-polyethylene oxide or PE-PEO) and (EO)<76> (= PEO). The surfactants and the organo-bentonites were characterized with X-ray diffraction (XRD) and Fourier transform infrared (FT-IR) spectroscopy. The surfactants intercalated the bentonite and expanded the d001 spacing of the freeze-dried samples to nearly 1.8 nm. The shapes and positions of the IR bands of interlayer surfactants resembled amorphous PEO. The EO segments of the surfactants were arranged in a distorted and extended form in the interlayer, instead of in the favored helical conformation of the crystalline state. The shifts of the C–O–C stretching bands of the intercalated surfactants pointed to H-bonding of the EO oxygen atom with water in the hydration shell of the exchangeable cations, or to direct coordination or ion–dipole interaction between the
exchangeable cations and the oxygen atoms. The type of exchangeable cations (Na\(^{+}\), K\(^{+}\), Ca\(^{2+}\), Mg\(^{2+}\), Cu\(^{2+}\), Ni\(^{2+}\), H\(^{+}\)) in the interlayer of smectite did not affect the conformation of the EO chain of the intercalated surfactants. These results give valuable information on the binding mechanisms and indicate clearly that the (EO)<n> conformations in the interlayer of the freeze-dried samples are more disordered than in the crystalline state. However, as indicated above, in the case of water-saturated samples the conformation of the EO chains is likely more random and affected by the presence of the apolar segments in the interlayer, because the apolar segments will locally associate due to hydrophobic attraction and not protrude into the aqueous solution.

The studies by Guégan [227, 228] on the intercalation of C10(EO)3 in the interlayers of Ca- and Na-montmorillonite (SWy-2) also deserve some attention because this surfactant easily forms in solution somewhat above the CMC a lamellar phase that consists of the stacking of molecules by hydrophobic attraction and it is well known that relatively small Cn(EO)m surfactants with m<n, such as C12(EO)6, adsorb on silica with a condensation step just before the CMC in a bilayer arrangement [153, 164]. With Ca-Mt the adsorption isotherm of C10(EO)8 was high affinity with a pseudo plateau reached at about the CMC. X-ray diffraction and FT-IR measurements on samples dried at 70 °C indicated an expansion of the basal spacing to 1.7 nm and adsorption in two adsorbed monolayers parallel to the clay surface [227]. In a subsequent study [228] intercalation of Na-Mt by C10(EO)3 was investigated, the Na-Mt suspension was well swollen and showed a stable hydration state in which silicate layers were widely expanded and already opened when the surfactant solution was added, thus allowing an easier adsorption. Moreover initial solution concentrations of C10(EO)3 were selected that where several times the CMC which already displayed the Lα-lamellar phase. The adsorption values that where reached were similar and somewhat higher (40%) than those in the Ca-Mt system. The results obtained with small angle X-ray scattering and with FT-IR (samples dried at 70 °C) however revealed d001
spacings from 1.1 nm for Na-Mt and 3.8 nm for the loaded samples and in the interlayer space a well-ordered surfactant bilayer, with a structure similar to that of the lamellar phase. The X-ray diffraction patterns further indicated that the C10E3 molecules were not aggregated on the external clay surfaces. Although the translation of these results to the surfactant conformations that are adopted with adsorption from solution is not straightforward, it seems plausible that a kind of bilayer with a hydrophobic core in the interlayer space might also be possible with simple C10(EO)3 adsorption. Calculations with the SCFA model might provide further insight; calculations for nonionic surfaces have been made before [152, 153, 165] and can be extended to adsorption in a confined space as has been done for ionic surfactants [214-217].

7. Surfactant adsorption modeling

7.1 General remarks

Surfactant adsorption isotherms have been qualitatively and (semi-) quantitatively interpreted using several adsorption models, ranging from the Langmuir equation to very sophisticated models. For a review of the models used to describe surfactant adsorption, including those specifically designed for surfactant adsorption to solid-liquid interfaces, the reader is referred to [25], where also references can be found. Here some classical models are briefly discussed in relation to surfactant adsorption. In general two types of modeling can be distinguished, (i) mean-field models in which the adsorbed layer is assumed to be smeared-out homogeneously over the surface and (ii) aggregation models in which the surfactants are adsorbed as aggregated, micellar type structures. Simple examples of the first category are the Langmuir-, the Frumkin-Fowler-Guggenheim- and the Freundlich -equation; examples of the aggregate models are the Hill- and the Gu-equation.

7.2 Mean-field models of monocomponent solute adsorption
The famous Langmuir model applies to sorption from an ideally dilute solution of non-interacting molecules on homogeneous binding sites. The binding sites should be attached to particles, but the geometry of the particles and precise location of the sites is not relevant, only the homogeneity of the sites is important. The Langmuir (L) Eq. can be written as:

$$\frac{\theta}{1-\theta} = Kc$$

(1)

where $\theta$ is the fraction of the sites covered with the adsorbate, $c$ the equilibrium concentration and $K$ the equilibrium affinity constant accounting for all types of interactions with the sites. Preferably, the concentration, $c$, is expressed as normalized concentration, i.e., with respect to a chosen standard state: $c = c' / c^*$, therefore, as both $c'$ and $c^*$ should have the same units, $c$ is dimensionless. When the concentration $c'$ is expressed in mol/L a convenient standard state is $c^* = 1$ mol/L. For a good understanding of the obtained parameter values the standard state should be indicated. In the case of a normalized concentration also the affinity constant $K$ is dimensionless and can be expressed as a Boltzmann factor containing the standard molar Gibbs energy of adsorption: $K = \exp(-\Delta G^0/RT)$, the surface covered with water (considered as a continuum) is taken as the standard state for the energy difference, $R$ is the universal gas constant and $T$ the absolute temperature. The site coverage can be expressed as the covered amount of sites divided by the total amount of sites, or, as is done most often in practice, the adsorption, $\Gamma$, divided by the maximum adsorption, $\Gamma_m$, and in that case the L-equation becomes

$$\Gamma = \frac{\Gamma_m Kc}{1 + Kc}$$

(2)

As the L-eq. applies to ideal sorption it will, in general, not be able to describe surfactant adsorption, but when it does, the obtained parameters $K$ and $\Gamma_m$ are conditional, that is to say, depending on the conditions of the experiment (surfactant concentration range and environmental conditions). Literature examples of reasonable fits of the L-eq. for surfactant
adsorption are mostly based a limited surfactant concentration range around the CMC where
the plateau adsorption is reached.

The Frumkin-Fowler-Guggenheim (FFG) equation is an extension of the L-eq. in which
lateral interaction between the adsorbed molecules is taken into account; it can be written as
\[
\frac{\theta}{1-\theta} = K_c \exp\left( -b \theta \right) \quad \text{with} \quad \theta = \frac{\Gamma}{\Gamma_m} \quad \tag{3}
\]
where \( b \) is a dimensionless lateral interaction parameter: \( b = \frac{\Delta G_{\text{lat}}^0}{RT} \) with \( \Delta G_{\text{lat}}^0 \) the
standard Gibbs energy of lateral interaction. This equation is more realistic for surfactant
adsorption than the L-eq. because next to the interaction with the surface (\( K \)), the lateral
hydrophobic attraction is included explicitly; the parameter \( b \) can be expressed as linearly
dependent on the alkyl chain length. In many studies of surfactant adsorption FFG-type
models have been used to qualitatively understand the role of surface-surfactant attraction
as compared to the lateral hydrophobic attraction.

When the binding sites are heterogeneous and randomly or regularly mixed the
adsorption isotherm can be obtained by solving the integral adsorption equation that
contains both the local isotherm for binding on equal energy sites and the heterogeneity
distribution. A Freundlich-type isotherm equation is obtained when the local isotherm is
L-type and the heterogeneity distribution is given by a quasi-Gaussian distribution function
[57]:
\[
\frac{\theta}{1-\theta} = \left( K_c \right)^p \quad \text{with} \quad \theta = \frac{\Gamma}{\Gamma_m} \quad \tag{4}
\]
where \( K_c \) is the median value of the affinity and \( p (0 < p \leq 1) \) is a measure of the width of
the distribution, for \( p =1 \) the surface is homogeneous. The quasi-Gaussian distribution
function is called ‘Sips-distribution’ after Sips [229, 230] who derived Eq. 4. Eq. 4 is called
the Langmuir-Freundlich equation, for \( p = 1 \) it reduces to the L-eq.. Because of the presence
of an exponent in the equation the use of a dimensionless solute concentration is a must, it
makes the interpretation of \( K \) possible. For sorbed amounts much lower than \( \Gamma_m \) Eq. 4
reduces to the common Freundlich equation, \( \Gamma = \Gamma_m \left( \frac{K c}{c^*} \right)^{1/\beta} = K^* c^{\beta'/\beta} \), where \( K^* \) is a composite parameter that can be indicated as the effective affinity constant; it has the same dimension as \( \Gamma \) (provided \( c \) is normalized) but information on \( \Gamma_m \) cannot be obtained. Freundlich type equations are more ‘flexible’ than the L-eq.; therefore, they have been mainly applied when the L-eq. did not fit the results well. The interpretation of the obtained parameters is however not so easy. Below we return to this aspect.

So far electrostatic interactions are not considered explicitly; in mean-field models these interactions can be formally introduced in the mentioned models by replacing the (normalized) equilibrium concentration of solute by the equilibrium solute concentration adjacent to the adsorption site:

\[
C_s = C_b \exp \left( \frac{-zF\psi_s}{RT} \right) \tag{5}
\]

where \( c_s \) is the solute-ion concentration at the location of the binding site, \( c_b \) the normalized equilibrium solute concentration in the dilute bulk solution, the Boltzmann factor accounts for the smeared-out electrostatic interaction at the location of the binding sites (mostly the surface) that the solute ion experiences, \( z \) is the valence of the solute ion and \( \psi_s \) is the electrostatic potential at the location of the site. The value of \( c_b \) can be multiplied by the activity coefficient calculated with, for instance the Debye-Hückel or Davis model [231], when the ionic strength is such that activity corrections are necessary.

A problem with Eq. 5 is that mostly no adequate information is present on \( \psi_s \). For low and moderate ionic strength values the zeta-potential, obtained with one of the electrokinetic methods [204], can be used as approximation for \( \psi_s \). Alternatively, a model can be used that relates the particle charge to \( \psi_s \). For flat surfaces and not too high surface potentials the Debye-Hückel model can be used [79], it relates \( \psi_s \) to the total (net) surface charge density, \( \sigma_t (\text{C/m}^2) \):
where $\kappa$ is the inverse Debye length, $\varepsilon_0$ the permittivity in vacuum and $\varepsilon_r$ the relative permittivity of water. The value of $\kappa$ is determined by the ionic strength of the solution, at room temperature and a $z$-$z$ electrolyte of concentration $c_z$,

$$\kappa = \left(10c_z z^2 \lambda^2\right)^{0.5} \text{ with } \kappa \text{ in nm}^{-1} \text{ and } c_z \text{ in mol/L.} \quad (7)$$

Substitution of Eqs. 5-7 in Eq. 3 provides an equation that explicitly takes into account the specific interaction with the surface, the electrostatic interactions both with the surface and in the adsorbed layer and the lateral interactions:

$$\frac{\theta}{1-\theta} = Kc \exp \left( -b\theta \frac{zF\sigma_p + z^2F^2I_m\theta}{RT\kappa} \right) \quad (8)$$

where use has been made of the fact that the total (net) particle charge density $\sigma_t = \sigma_p + zF^2I_m\theta$ with $\sigma_p$ the primary surface charge density and $zF^2I_m\theta$ the charge density contribution due to the ionic solute adsorption. This type of equation can be used to explain the presence of a common intersection point (CIP) of ionic surfactant isotherms measured at different ionic strength values. At the CIP the electrostatic interaction vanishes, as there is no effect of the salt concentration on the adsorption. This must mean that $zF\sigma_p + z^2F^2I_m\theta = 0$, i.e., the adsorbed surfactant charge just compensates the primary surface charge. For concentrations below the CIP the primary particle charge dominates and an increase in ionic strength (increase in $\kappa$) decreases the electrostatic surface-surfactant attraction, therefore the adsorption decreases with increasing ionic strength. For concentrations beyond the CIP the surfactant adsorption dominates the electrostatic interaction and with an increase in ionic strength the repulsion between the charged surfactant head groups decreases, therefore the adsorption increases with increasing ionic strength. From the above equations Eq. 8 is best equipped to understand qualitative or
(semi)-quantitative ionic surfactant adsorption behavior. Eq. 8 is a FFG-type equation this can be shown by regrouping the various terms:

\[
\frac{\theta}{1-\theta} = K_{e1} c \exp\left(-b\theta - b_d \theta\right)
\]

where \( K_{e1} = \exp\left(-zF\sigma_p/RT\epsilon\kappa\right) \) and \( b_d = \exp\left(-z^2F^2I_m/RT\epsilon\kappa\right) \). Eq. 9 shows that when the electrostatic interactions are not explicitly considered with ionic surfactant adsorption and the simple FFG- Eq. (3) is applied, that both the affinity and the lateral interaction parameter contain an electrostatic contribution. Further extensions of Eqs. 3, 6, 8 and 9 are discussed in [232, 233].

Ionic surfactant adsorption is often at least partly due to ion exchange and/or accompanied by adaptation of the primary surface charge. None of the above equations considers the multicomponent nature or the competition. This implies that when the equations are applied in practice they are conditional, or stated differently, the fitted parameters are no longer adequate when the concentration of the competing ion (e.g., H\(^+\) or OH\(^-\)) is changed. This can be illustrated by using the multicomponent Langmuir equation:

\[
\theta_i = \frac{K_i c_i}{1 + K_i c_i + K_2 c_2} = \frac{K_i^c c_i}{1 + K_i^c c_i} \quad \text{with} \quad K_i^c = \frac{K_i}{1 + K_2 c_2}
\]

where \( K_i^c \) is the conditional affinity, which is only constant at constant \( c_2 \). Eq. 10 is also Langmuir-type but it provides information on how \( K_i^c \) changes with changing conditions. A condition for the application of Eq. 10 is that \( I_m \) should be the same for both components; otherwise also the stoichiometry of the exchange has to be taken into account.

With ionic surfactants and charged surfaces adsorption is often a two-step process. The first step is binding to the charged surface, where some kind ion exchange and lateral hydrophobic attraction play a role, and the second step is governed by hydrophobic attraction to already adsorbed molecules and lateral hydrophobic and electrostatic
interaction in the second layer. With the above single layer models this distinction is not
made, which implies that the second layer adsorption can only be due to lateral interaction.
As a result the $b$ parameter has a composite character, it accounts for both the layer-layer
and the lateral hydrophobic attraction. A simple model that tries to avoid this problem is the
FFG two-layer model of Mehrian et al. [180] in which not only the affinities for the first and
second layer are different but also the lateral interactions, however the electrostatic
interactions are not made explicit. The results of this model are discussed in the section on
cationic surfactant adsorption to kaolinite. Scamehorn et al. [234] also used a two-layer
FFG equation without treating the electrostatic interactions explicitly. With their parameter
choice the affinity for the first and second layer differed, but the lateral interactions were
assumed to be the same. With the chosen parameter values the model predicted a strong 2D
phase transition at low surfactant concentrations and heterogeneity was invoked to explain
the experimental isotherms. As explained above for the FFG-eq., also a two-layer FFG
model, that does not treat electrostatics explicitly, is only suited for the description of
experiments done at constant pH and constant salt concentration.

7.3 Aggregation models for monocomponent solute adsorption

The second group of adsorption models starts with the Hill equation that describes the
formation of homogeneous aggregates containing $n$ solute molecules on $N_s$ homogeneous
binding sites:

$$\theta_n = \left( Kc \right)^n \quad \text{with} \quad \theta_n = \frac{\Gamma_n}{\Gamma_t} = \frac{\Gamma_n}{n N_s} \quad (11)$$

where $\theta_n$ is the fraction of sites covered with $n$ molecules, $K$ is the affinity of the binding of
1 molecule, $K^n$ that of the $n$ molecules, $c$ the normalized solute concentration, the actual
adsorption equals $\Gamma_n$, and the total (maximum) adsorption $\Gamma_t$ is equal to $n N_s$. The process of
adsorption is *cooperative* because one adsorption step brings $n$ molecules to a site, $n$ is
therefore also called the cooperativity parameter. The Hill-eq. reduces to the L-eq. for $n = 1$;
isotherms for $n > 1$ are somewhat S-shaped and the isotherm is steeper than the L-isotherm
(*positive cooperativity*); for high $n$ the steepness of the adsorption isotherm is large and
close to a two-dimensional (2D) phase transition. It is also possible to derive the Hill-eq. for
the situation that one molecule occupies $n^*$ sites ($n^*$-dentate adsorption), so that on one site
only $1/n^*$ molecules are adsorbed. In this case $n = 1/n^*$ or $n < 1$, i.e., for $n = 1/2$ the
adsorption is bidentate. For $n < 1$ there is negative cooperativity and the isotherm is less
steep than the Langmuir isotherm because it more difficult to find $n^*$ sites than 1 site.
Instead of calling $n$ the cooperativity parameter, $n$ can thus also be called the *stoichiometry
parameter*: $n > 1$ more than 1 molecule per site, $n < 1$ more than 1 site required to adsorb 1
molecule.

It should be noted that the mathematical forms of the Hill-eq. and the
Langmuir-Freundlich-eq. are identical; in terms of the Hill-eq. heterogeneity is a form of
negative cooperativity, where adsorption becomes more difficult when the coverage
increases. When the Hill-eq. or the LF-equation is applied in a practical situation the
cooperativity parameter might therefore be affected by site heterogeneity, and the
heterogeneity parameter by the cooperativity.

Zhu and Gu [150, 151] have proposed an extension of the Hill-eq., they make a
distinction between a first and second step in the adsorption process. In the first step solute
molecules are bound to the sites and in the second steps these bound molecules act as nuclei
for an aggregation step or positive cooperative adsorption. The adsorbed amount in the first
state equals

$$I_1 = N_s \theta_1 = N_s \frac{\kappa K_c}{1 + \kappa K_c + \kappa (Kc)^n}$$

that in the second state
and the total adsorption is obtained by summation of the two contributions

\[ I'_n = N_s n_\theta_n = N_s \frac{nK(Kc)^n}{1 + Kc + \kappa(Kc)^n} \]

In these equations \( k \) is the ratio between the affinities of the first and second state, \( K \) the affinity of the second state, therefore, \( kk \kappa \) is the binding affinity for the first state. For \( k = 1 \) the affinities for the first and the second state are equal, but the nuclei formation is initially stronger than the aggregation because there are more free sites than nuclei. For \( k \gg 1 \) the isotherm is typical stepwise, the first step is pronounced at low concentration and the second step occurs at somewhat higher concentrations. The parameter \( n \) is in this case equal to the ratio of the first adsorption plateau over the second. For \( k << 1 \) the equation becomes similar to the Hill equation and the isotherm depicts one step. Eq. 12 is often called the Gu-equation, it is a very ‘flexible’ equation and regularly used to describe quantitatively the adsorption of both nonionic and ionic surfactants.

In the case of ionic surfactants the description is conditional, because for each change in solution conditions (pH and ionic strength) new parameters are required. Moreover, the affinity for the first step is for ionic surfactants determined by both the specific and electrostatic interaction and lateral interactions, the parameter \( k \) accounts for the first two interactions, \( K \) for the lateral attraction. Ion competition for the surface sites is not considered. The affinity in the second step comprises the hydrophobic affinity for the nuclei and the lateral interactions; the latter contain for ionic surfactants not only the hydrophobic attraction but also the electrostatic repulsion. Part of the lateral interaction for the second step is however also incorporated in the value of \( n \). It is therefore difficult to give a more detailed interpretation of the parameter values. The main advantage of the Eq. 12 is that a distinction can be made between the overall affinities for the first and the second step.
When it is assumed that the adsorbed states are governed by the same electrostatic adsorption potential it is possible to incorporate the electrostatic interactions in the Gu-eq. in the same way as illustrated above for the FFG-eq., but this has not been done.

Two sophisticated models for surfactant adsorption that take aggregation into account are that of Rudzinski et al. [235, 236] and Li and Ruckenstein [237]. The model by Li and Ruckenstein [237] is most advanced, it has much in common with the description of micellization of ionic surfactants but it also reflects the scaled particle theory used by Rudzinski et al. [235, 236]. Solvent molecules, surfactant monomers and two types of surfactant aggregates of various sizes cover the surface. The competition between the enthalpic and entropic contributions to the Gibbs energy in the adsorbed phase is responsible for the composition of the adsorbed phase; the standard Gibbs energy change in going from the solution to the surface aggregates is calculated by considering five contributions: hydrophobic, conformational, electrostatic, steric and interfacial. The electrostatic contributions are treated within the framework of the Poisson-Boltzmann equation for flat plates. The model can well predict the four-region isotherms, but no predictions have been made of ionic strength effects.

7.4 SCFA model for surfactant adsorption

The SCFA or Self-Consistent-Field theory for Adsorption model is a sophisticated model in which the structure and properties of the components present in the system are explicitly taken into account by considering them as composed of different segments that are linked together. The type and number of segments of a molecule, their charge and the way they are linked determine the structure and the properties of a molecule. Most molecules in the system are composed of just one or a few segments, but the molecular structure of the surfactant molecules is mimicked as closely as possible. The flat surface is also composed of segments that characterize the surface, for charged surfaces a part of the
segments can associate or dissociate a proton. The model is an extension of the theory of polymer solutions by Flory [107] to systems that have inhomogeneities in one direction (1D SCFA), i.e. perpendicular to the surface or the center of the lattice, or in two directions (2D SCFA), i.e. perpendicular to the surface and in concentric rings parallel to the surface. In the 1D SCFA option a mean field approximation is used in every lattice layer around to the center of the lattice or parallel to the surface. In the 2D case the mean field approximation is used in every ring. The interactions between the segments are calculated using Flory-Huggins interaction parameters, similarly as in the Flory theory. Starting from system characteristics such as the structure of the surface, the number of different types of molecules, the amount of each of them, their structure and properties, and the interactions between the various segments, the equilibrium distribution of molecules in the adsorbed layer is calculated. The conformational statistics of the chains in the layer are evaluated using Boltzmann statistics, i.e., each step of the walk is weighted with a Boltzmann factor that accounts for the interactions that the segment experiences in the adsorbed layer. The electrostatic interactions in the adsorbed layer are calculated using a multi-plate condenser model. In the absence of surfactants the electrostatic potential profiles correspond with the potential calculated with the Stern-Gouy-Chapman model. The detailed equilibrium distribution of surfactant segments within the adsorbed layer, i.e. within a potential field exerted by the presence of the surfactant molecules themselves and the surface, is calculated by an iterative minimization of the Gibbs energy of the system. At equilibrium the segment density distribution in the adsorbed layer and the potential field exerted by the segments and the surface are fully in accordance with each other (field is self-consistent). Once this is achieved the structure of the adsorbed layer and the thermodynamic properties of the system can be obtained. Similarly as in the model of Li and Ruckenstein [237], the SCFA theory has a closely related variant for self-assembly in solution [238, 239] so that the surfactant micellization can also be predicted. When micellization of a given surfactant is predicted
with a certain set of parameters, the same set of parameters should be used to predict the adsorption. The only new parameters for the adsorption calculation are in that case those that describe the interactions of the segments with the solid surface. As mentioned in the text on several places, the SCFA model has been very helpful for a better understanding of the surfactant adsorption behavior. Reference [25] is a fairly recent review in which experimental surfactant adsorption results obtained for metal oxides and silicas are compared with SCFA calculations. More recent results for surfactant adsorption in confined space can be found in the section on cationic surfactant adsorption on montmorillonite. For the SCF calculations a computer code, the SF-box, is available; for further information and/or application of the SF-Box see [240].

7.5 **NICA-Donnan model applied to surfactant sorption on humic and fulvic acids**

Surfactant sorption to humic and fulvic acids can be described with the NICA-Donnan model as an ion exchange process. In this model the humic substance is considered as a collection of particles that contain the binding sites and that are permeable for solvent and solute. The particles form the Donnan phase in which the electrostatic interactions are governed by a smeared-out electrostatic potential that is the result of the fixed charges and the mobile charges in the Donnan phase. The binding to the heterogeneous sites is described by the NICA model, which is based on the multicomponent Hill equation as local isotherm in combination with the Sips distribution and can be written as [57, 58]

\[
Q_i = Q_{H \text{max}}^{i} \left( \frac{\tilde{h}_i}{\tilde{h}_i^{\text{H}}} \right) \left( \frac{\tilde{K}_i^{C_D_j} \tilde{h}_i^{C_D_j}}{\sum_j (\tilde{K}_i^{C_D_j} \tilde{h}_i^{C_D_j})^{y_j}} \right)^p \left( 1 + \left( \sum_j (\tilde{K}_i^{C_D_j} \tilde{h}_i^{C_D_j})^{y_j} \right)^p \right)^{-1}
\]

(13)

where \(Q_{H \text{max}}^{i}\) is the maximum sorption of protons, \(\tilde{h}_i\) the Hill-parameter that accounts for the average stoichiometry of component \(i\) with the proton sites, \(\tilde{K}_i\) is the median affinity of
component $i$ for the sites and $p$ the heterogeneity parameter. It is assumed that the first
characterization of the humic substance is done through proton sorption, therefore this ion is
used as reference for the total number of sites ($Q_H^{\text{max}}$) and the factor $\tilde{n}_i / \tilde{n}_H$ accounts for
the fact that the maximum sorption of $i$ can be different from that of the proton. The
concentration $c_{D,i}$ is the concentration of $i$ in the Donnan phase that can be obtained from
the bulk solution concentration using Eq. 5 with $\psi_s = \psi_D$. In the Donnan model the charge
of the humic particles is neutralized by counter- and co-ions within the Donnan volume, the
latter is calculated with a simple empirical relation. Based on this electroneutrality the
Donnan potential can be calculated and $c_{D,i}$ can be obtained for all ions present [38]. The
NICA equation is a master equation that reduces to the multicomponent Langmuir equation
for all $\tilde{n}_i = 1$, and $p = 1$, to the multicomponent Langmuir - Freundlich eq. for all $\tilde{n}_i = 1$ and
to the multicomponent Hill eq. for $p = 1$.

The last quotient in Eq. 13 is the total coverage of the sites with all types of ions present
in the system, the second quotient is the fraction of the total coverage that is occupied by
component $i$. Therefore, in the case of sorption of a trace component, $j$,
\[
(\tilde{K}_j c_{D,j})^j < < \sum_i (\tilde{K}_i c_{D,i})^\tilde{n}_i
\]
the total coverage is hardly affected by the binding of
species $j$ and the NICA eq. reduces to
\[
Q_j = \Omega(\tilde{K}_j c_{D,j})^j \quad \text{with} \quad \Omega = \frac{Q_H^{\text{max}}}{\tilde{n}_H} \left( \frac{1}{\sum_i (\tilde{K}_i c_{D,i})^\tilde{n}_i} + \left( \sum_i (\tilde{K}_i c_{D,i})^\tilde{n}_i \right)^p \right)
\]
where $\Omega$ is considered to be constant. When the total adsorption is constant, then also the
Donnan potential will be constant and the electrostatic interactions do not significantly
change with sorption of $j$, thus also $c_{D,j}$ can be replaced by $c_j$ (Boltzmann factor becomes
included in $\Omega$). Therefore, at low solute concentrations in a multicomponent system the
sorption equation of a trace ionic component to the active sites of the humic substance, at
otherwise constant solution conditions (pH, ionic composition and ionic strength) is ruled by a simple equation that resembles the monocomponent Hill-eq. for low loadings. Because the sorption of $j$ is low, the site heterogeneity is not detected but the cooperativity / stoichiometry ($\tilde{\nu}_j$) is relevant. When the humic substance is well studied, $\Omega$ is known and the changes of $\Omega$ with changing solution conditions (e.g., change of pH) can be calculated, therefore the behavior of the trace component with changing solution conditions can be predicted, once $\tilde{K}_j$ and $\tilde{\nu}_j$ are known. When no information is present on the humic acid, the generic humic acid proposed by Milne et al. [55, 56] can be used to represent the unknown humic acid and by using the parameters collected by Milne the total sorption can be calculated under the given conditions. Thus also in this case Eq. 14 can be used in combination with Eq. 5 to investigate trace component sorption and how it is affected by, e.g., pH.

In the case of metal ion binding a double NICA-Donnan equation has been used, where the first equation accounts for the low proton affinity sites (‘carboxylic’) and the second for the high affinity proton sites (‘phenolic’). The above way of analyzing trace component binding is still relevant as long as the total sorption on both the phenolic and carboxylic type of groups is constant. In most cases the trace sorption will occur with either the carboxylic or the phenolic groups and in this case the type of groups that do not participate can be neglected.

For the surfactant binding only a limiting cases of the single NICA-Donnan equation have been used. As discussed in the section ‘Modeling surfactant - humic substance interaction’ Chen et al. [114] used Eq. 14 to unravel the surfactant binding and the Hill equations used by Yee et al. [98-101] are identical to Eq. 14. Ishiguro and Koopal [115] made a different simplification for their analysis of surfactant adsorption to various humic
Eq. 15 is mathematically equivalent to the Langmuir-Freundlich equation, but here the exponent is a composite parameter that reflects both the cooperativity and the heterogeneity. Because the surfactant sorption covered the entire sorption range up to $Q_{j}^{\text{max}}$ the exponent in Eq. 15 is also affected by the heterogeneity. For the calculation of the Donnan potential the total (net) charge of the humic acids was used and the proton charge adjustment due to the surfactant adsorption was accounted for. When the charge adjustments are not known and a computer code for NICA-Donnan calculations is available, a good method is to use the full (single) NICA-Donnan equation together with humic substance specific parameters or generic humic substance parameters [55, 56], because then the charge adjustments are automatically taken into account. Computer codes that include the NICA-Donnan model are ECOSAT [241] and Visual MINTEQ ver. 3.0 / 3.1 [242]. Furthermore, the computer code 'Fit' [243] is designed to fit experimental results to a range of adsorption isotherm equations.

8. Surfactant adsorption on soils

8.1 General considerations and trends

Soils are composed of organic and inorganic components each with different adsorption characteristics as discussed in the previous sections. Therefore, the adsorption of a surfactant on a soil sample will depend on both the types of soil constituents and their relative proportions. This has also been stated in literature [244]. The overall surfactant
adsorption to a given soil can be thus be approximated by a weighted summation of the
adsorptions to the individual constituents, or oppositely, the soil composition has to be
known to be able to understand the surfactant adsorption behavior. Yet, the overall
adsorption is not necessarily equal to the sum of the adsorptions to the different components
because the soil components may mutually interact and this interaction will affect the
adsorption. The mutual interaction is especially relevant and strong in the case of net
positively charged metal oxides or layered silicates and the negatively charged humic and
fulvic acids [245-249]. A similar situation is encountered with trace element binding to soils
[50].

Some general trends have also been observed for surfactant adsorption to soils.
Hydrophobic interaction is important with surfactant adsorption. Due to increasing
hydrophobic attraction surfactant adsorption increases, in general, with increasing organic
matter content of the soils [250-256]. Treatment of a soil with H₂O₂ to remove the organic
matter will therefore, in general, decrease the surfactant adsorption [228, 229], but the
contrary has also been observed [231]. Nonionic surfactants bind to soil surfaces with
hydrogen bonding on the –OH sites of soils besides hydrophobic interaction [257], but
hydrogen bonding is also possible to the hydrated cations on siloxane basal planes [198].
Longer alkyl chains lead to stronger lateral hydrophobic attraction, this explains that there is
also a positive correlation between aliphatic chain length and surfactant adsorption on soils.
An example of this behavior has been presented by Westall et al. [255] who showed that
linear alkylbenzene sulfonates with longer carbon chains adsorbed stronger on a sediment
(Fig. 29). Adsorption studies comparing linear and branched aliphatic chain structures have
also been performed for soils and alumina [258-261]. For surfactants with the same number
of CH₂ segments, branching leads to lower adsorbed amounts at the same surfactant
concentration. This is largely due to a decrease of the hydrophobic attraction; the decrease
in adsorption corresponds with the increase in CMC with branching [261].
Soils possess both permanent and pH-dependent charges [262, 263]. The contents of those charges differ among soils and affect the adsorption of charged surfactants strongly. Soils and sediments generally have negative charges, therefore, the adsorption of anionic surfactants on soils and sediments is much lower than that of nonionic and cationic surfactants [251, 255, 256, 264, 265]. Bera et al. [266] showed that surfactant adsorption on a sand was in the order of cationic > nonionic > anionic (Fig. 30). For surfactants with a similar alkyl chain the adsorption of the cationic surfactant is largest because of the electrostatic attraction between surface and surfactant and that of the anionic surfactant is smallest because of the electrostatic repulsion. An example where the relatively low anionic surfactant adsorption to sand is applied is surfactant-enhanced soil remediation. With surfactant-enhanced soil remediation, surfactant adsorption to the soil particles increases the surfactant use and this decreases the efficiency. Therefore, anionic surfactants have usually been chosen in these technologies [267-269]. However, nonionic surfactants are also often used for the remediation because of their lower CMC compared to ionic surfactants, higher degree of surface tension reduction, and relatively constant properties in the presence of salt [268, 269].

A positive correlation is observed between cationic surfactant adsorption and the soil CEC because the CEC characterizes the soil negative charge density and the latter generates the electrostatic attraction with cationic surfactants [270]. For the soils which possess variable charges the pH also affects the adsorption as shown in Fig. 31 [266]. Anionic surfactant adsorption decreases as pH increases [254, 255, 258-260, 266, 271] because the soil negative charge increases and/or the positive charge decreases. On the other hand, the opposite occurs for cationic surfactant adsorption as shown in Fig. 31 [266]. When after sodium citrate - bicarbonate - dithionite treatment the fraction amorphous oxides in soils is decreased, the adsorption of anionic surfactant also decreases because the density of positively charged sites of the soils decreases [260]. Nonionic surfactant adsorption
increases as pH decreases at low pH and remains almost constant at high pH (Fig. 31) [266]. The increase at low pH resulted from the increase of the surface hydroxyl sites (due to protonation of the surface oxygens) to which the EO segments of the nonionics can bind by hydrogen bonding.

The electrolyte concentration also affects ionic surfactant adsorption. It is observed that anionic surfactant adsorption on negatively charged soils increases, in general, with the increase of electrolyte concentration [255, 258, 266]. This must be due to a larger screening of both the electrostatic repulsion between surface and surfactant and the mutual electrostatic repulsion between the adsorbed surfactant ions. Increase of cationic surfactant adsorption on negatively charged soils with the increase of electrolyte concentration is also observed [272, 273]. This is caused by the screening of the electrostatic repulsion between the head-out adsorbed head groups of the cationic surfactants. This situation occurs for bilayer and/or admicelle adsorption where part of the head groups are directed to the soil solution; this part of the surfactants is adsorbed by hydrophobic attraction to the carbon chains of the surfactants adsorbed in the first layer. In this case, the cationic surfactant adsorption is also affected by the type of anion of the electrolyte. The surfactant adsorption increases in the order of $SO_4^{2-} > Br^- > Cl^-$ because the screening effect increases in this order [272]. The influence of the electrolyte concentration becomes reverse for ionic surfactants adsorbed in direct contact with the oppositely charged soil surface (head-on adsorption). The adsorption decreases with the increase of electrolyte concentration due to the screening of the electrostatic surfactant-surface attraction.

An example of the influence of the electrolyte concentration on the sorption of dodecyl benzene sulfonate (DBS) on a humic soil is depicted in Fig. 32 [258]. The adsorption decreases with decreasing electrolyte concentration because the electrostatic repulsion increases. The influence of the (smeared-out) electrostatic potential at the location of the adsorption sites can be evaluated by using the Langmuir-Donnan model. In this model it is
assumed that the soil particles are covered with humic substances and that the surfactant binds to the sites in the humic layer according to a Langmuir equation that is extended by including the electrostatic attraction by a Boltzmann factor, $\exp(-zF\psi_D/RT)$, where $\psi_D$ is the Donnan potential in the humic layer (see section 7):

$$Q_j = Q_j^{\text{max}} \frac{K'c_{D,j}}{1 + K'c_{D,j}}$$  \hspace{1cm} (16)

$$c_{D,j} = c_{b,j} \exp\left(-\frac{z_jF\psi_D}{RT}\right)$$  \hspace{1cm} (17)

where, $Q_j$ is the surfactant adsorption, $Q_j^{\text{max}}$ the maximum surfactant adsorption, $K'$ the specific or chemical affinity constant (surface-surfactant), $z_j$ the valence of the surfactant ion, $c_{D,j}$ the surfactant concentration in the Donnan phase and $c_{b,j}$ the surfactant concentration in the bulk solution. When for all salt concentrations the value of $Q_j^{\text{max}}$ is put equal to the surfactant sorption at the CMC (estimated to be 20 $\mu$mol/g), the affinity constant, $K'$, and the Donnan potentials, $\psi_D$, can be obtained by fitting Eq. 16+17 to the experimental data in the region where the slope of the double logarithmic plots is about equal to unity (region where Eq. 16 applies). The fitted adsorption isotherms and obtained Donnan potentials are depicted in Fig. 32. The negative value of Donnan potential increases at constant salt concentration due to the fact that DBS sorption increases the negative charge of the particles. Increasing the salt concentration at a given DBS sorption has the opposite effect: the salt ions screen the particle charge; therefore, the repulsive Donnan potential decreases with increasing the salt concentration. At the lowest surfactant concentrations the isotherms rise very steeply; this likely indicates strong sorption to some highly active sites of the soil sample.

8.2 Fate of surfactants in soils; degradation, hysteresis and precipitation
When the fate of surfactants in soil and water environments is considered, surfactant degradation is important, as well as adsorption. However, different classes of surfactants have different degradation behavior in the environment [264]. Sodium dodecylsulfate (SDS) easily degrades, especially in humic soil. Therefore, the influence of degradation on adsorption experiments is significant. Results of some batch experiments of SDS in the presence of a highly humic non-allophanic andosol are depicted in Fig. 33 [274]; the sum of adsorption and degradation corresponds to the SDS decay in the soil solution. The “adsorption + degradation” amount at intermediate SDS concentrations increases with time due to increasing of degradation. The initial steep increase is somewhat puzzling; the irregular behavior suggests that either or both degradation and cooperative adsorption are important in this region [274].

The environment also affects the degradation; sodium dodecylbenzenesulfonate with a linear carbon chain (L-DBS) degrades in river water with a half-life of less than 3 days, but its degradation in aerobic soil with a half-life of 7–33 days is much longer [264]. In general, adsorption restricts L-DBS degradation, especially in soils with allophane and amorphous metal oxides [260], which adsorb the anionic surfactant by electrostatic attraction.

Hysteresis of adsorption, which implies different adsorbed amounts for the adsorption and the desorption isotherm [168, 169, 275], must be also considered as a distinctive phenomenon. In general, the adsorbed amount with desorption is larger than that of adsorption [169, 275]. The hysteresis can, in general, be due to kinetic and/or multicomponent adsorption effects. An important kinetic reason for the difference between the adsorption and desorption isotherm is that the kinetics of desorption become very slow when the adsorption isotherm is rather steep [276]. In this case the adsorption isotherm is the true isotherm, the desorption isotherm is an artifact. In multicomponent systems the adsorption conditions are often not the same as the desorption conditions. This is especially the case in the presence of impurities adsorbed to the solid and when the solid area to
solution volume ratio is different in the ad- and desorption experiment. In the case of adsorption by ion exchange (‘first layer’ adsorption) and hydrophobic attraction (admicelle formation or ‘second layer’ adsorption), desorption of the ‘second layer’ shows no hysteresis. This has, for instance, been observed for cationic surfactant adsorption on vermiculitic soil with a negative charge [272].

Ionic surfactant insolubility must be considered when the Krafft temperature is larger than the temperature under the conditions of the adsorption [201, 277, 278]. The Krafft temperature or *Krafft point* is the minimum temperature at which surfactants can form micelles, below the Krafft temperature surfactant added to a solution with a monomer concentration close to the CMC remains in the crystalline form. Therefore, micelles are not formed when the temperature is lower than the Krafft point. The Krafft point becomes higher when the coion of the surfactant is multivalent.

### 8.3 Influence of surfactant adsorption on soil structure and permeability

Because surfactants modify the surface characteristics of solids, they affect the structure of soils and this affects the water permeability and gas diffusion in soils [279-283]. Proper permeability and gas diffusion are required for crop growth; therefore, a good soil structure must be maintained in crop fields. When a soil disperses and swells during the irrigation or rainfall, dispersed soil particles clog the macro-pores (water transmission pores) in the soil and permeability is decreased. A change in soil structure that leads to a decrease in permeability is a negative factor in relation to agriculture and soil-water environment. Moreover, sometimes flooded water conveys soils and causes erosion; important soil is lost and the nutrients or chemicals in the eroded soil cause water contamination. The soil dispersion is generally caused by clays and the reduction of hydraulic conductivity is larger when the clay content in the soil is larger [280]. When the soil organic matter content is
reduced, the reduction of hydraulic conductivity becomes larger because clays in soils with
a high organic content are not easily dispersed [279].

A review on soil structure can be found in [284]. Soil and clay structural changes with
changing inter-particle interaction have been described in [263, 285-288]. In general,
repulsive inter-particle interaction allows the particles to slip along each other, which leads
to relatively dense structures; with attractive inter-particle interactions the particles stick
together which retards further individual movement and the clustering leads to a relatively
open structure. It is general knowledge that a settling stable dispersion forms a compact
structure and a settling unstable dispersion forms an open structure. The use of surfactants
for soil washing to remove low-solubility organic contaminants, such as oils, also changes
the inter-particle interactions and can result in a significant change in the hydraulic
conductivity of porous media [279, 283]. Electrolyte concentration, pH and specific ions
that all likely affect the particle-particle interaction may also change the soil permeability
[263, 288-297]. An example of the effect of surfactant adsorption on soil stability is
depicted in Fig. 34 [272], which shows the adsorption isotherm of
hexadecyltrimethylammonium (HDTMA=C16TA+), on two vermiculitic soils (Na and Ca),
and the corresponding effects on the electrophoretic mobility of the soil particles and the
optical density of the soil suspension. When the soil is better dispersed, the optical density
becomes larger. For the Na-soil, the shape of the isotherm in region I indicates that the
vermiculite is expanding upon adsorption. Up to about adsorbed amounts equal to 1.0 CEC,
the soil gradually flocculates with increasing surfactant adsorption and the electrophoretic
mobility changes from negative to zero at the IEP, where the surfactant adsorption is about
equal to the CEC. The negative charge of the soil is gradually compensated with the
adsorbed surfactant charge; therefore, the electrostatic repulsion between the soil particles
gradually vanishes and the dispersion is completely flocculated. With a vermiculitic soil and
a cationic surfactant the adsorption occurs initially by ion exchange, this implies surfactant
head groups in contact with the surface and tails contacting the solution; therefore, also the
hydrophobicity of the soil particles increases and this is likely a second contribution to the
decrease in dispersion stability. When the adsorption becomes larger than the soil CEC
further adsorption will occur by hydrophobic attraction and the formation of an admicellar
layer (second layer) on the surface. With further adsorption the soil particles become
positive and less hydrophobic and the soil disperses again. The positive charge is indicated
by the positive electrophoretic mobilities, For the Ca-vermiculite-soil, the behavior is
different. The shape of the isotherm indicates that there is negligible swelling of the
vermiculite. Moreover, the electrophoretic mobilities are rather low and this explains the
poor initial degree of dispersion. Surfactant adsorption by ion exchange hardly changes the
external particle charge and hence the mobility stays low as does the degree of dispersion.
The Ca-soil begins to disperse after passing the IEP where the particles become positive and
the second adsorption layer starts, but the stability of dispersion stays somewhat lower than
for the Na-soil. The latter may be due to the effect of the released Ca ions on the ionic
strength.

Allred and Brown [279] showed that the soil hydraulic conductivity significantly
decreased with the infiltration of ionic and nonionic surfactant solutions at high
concentration (1 mol/kg); the behavior is depicted in Fig. 35. The curves of N1, N2 and N3
in Fig. 35A represent the nonionic surfactants, the curves of A1 and A2 in Fig. 35B
represent the anionic surfactants, the curves of C1 and C2 in Fig. 35B represent the cationic
surfactants, and the curve of AM1 in Fig. 35B represents the amphoteric surfactant. The
hydraulic conductivity reduction was larger for ionic surfactants than nonionic surfactants.

When an anionic surfactant is adsorbed on a negatively charged soil, the negative charge of
the soil surface becomes larger and the soil particles disperse easier, and the particles slip
easier along each other; therefore, the hydraulic conductivity decreases. When a cationic
surfactant is adsorbed on the negatively charged soil, the situation is somewhat similar to
that described in the previous section for the vermiculite soils. The stability of dispersion
goes through a minimum, but when the positive charge due to surfactant adsorption is
sufficient the soil particles become well dispersive and the hydraulic conductivity tends to
decrease. When the electrolyte concentration is large, the influence of ionic surfactant
adsorption on the decrease of hydraulic conductivity is restricted [281], because the soil
dispersion ability is inhibited by the strong screening of the particle-particle repulsion.

As a nonionic surfactant does not change the particle charge its effect on the dispersion
stability can only result from the way the surfactant is adsorbed. On most natural surfaces
the hydrophilic head group will be in contact with the surface and similarly as for cationic
surfactants a second layer may become adsorbed at surfactant concentrations around the
CMC. For short EO chains the adsorption in the ‘first layer’ is moderate and the transition
to the ‘second layer’ occurs in a narrow concentration range close to the CMC. For long EO
chains the adsorption in the ‘first layer’ is larger, but due to the large head group it is
difficult for the alkyl chains to associate and only when the alkyl chains are also fairly long
a sufficient ‘second layer’ will form. Also with kaolinite the edge surface probably
contributes, but on montmorillonite this contribution is relatively small. Therefore, the
behavior with respect to hydraulic conductivity will strongly depend on the type of nonionic
surfactant and the type of soil surface. Mustafa and Letey [298] indicated for a hydrophobic
sandy loam, a hydrophilic sandy loam and a hydrophilic clay loam that nonionic surfactant
adsorption induced soil dispersion by decreasing the aggregate size.

Water infiltration into dry soils with a hydrophobic surface is also affected by surfactant
adsorption. Fig. 36 shows the upward water infiltration into a peat moss column [299]. The
bottom of the air-dried peat moss column was set 20 cm below the water table. A 70-mM
SDS solution infiltrated faster than pure water because the surface became wettable after the
adsorption of SDS. On the other hand, infiltration of the surfactant solution into dry sand
became slower compared to pure water infiltration, because the sand was already wettable
without surfactant. In this case, they explained that the surface tension of surfactant solution was smaller than that of pure water and the viscosity of surfactant solution was larger than pure water and both effects decreased the rate of capillary infiltration.

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Figure captions

Fig. 1. Some examples of synthetic and natural surfactants.

Fig. 2. Schematic illustration of the effect of surfactants on the surface tension of dilute aqueous solutions (a, c, d) and the adsorption isotherm (b) that can be obtained from figure (a) through application of the Gibbs equation. The kink in the surface tension plots (a, c, d) is the CMC.

Fig. 3. Set of surface tension vs. $\ln c$ curves for aqueous solutions of sodium dodecyl pyridinium chloride (DPC) in the presence of different concentrations of NaCl ($\bullet$ 0 mmol/L; □ 5 mmol/L; ○ 20 mmol/L; △ 100 mmol/L).

Fig. 4. Example of proton binding to a humic acid at four KCl concentrations. Data from ref. [45].

Fig. 5. Comparison of the charge density curves vs. pH and ionic strength of gibbsite, silica and hematite. The ionic strength values of the curves of hematite are (△) 0.002 M, (X) 0.01 M, and (O) 0.1 M. Data from ref. [81-83].

Fig. 6. Proton charge density of the montmorillonite for three values of the 1-1 electrolyte concentration (symbols) and model predictions (curves). Data from ref. [92].

Fig. 7. Binding of C12PC and C16PC to PAHA (mmol surfactant/g PAHA) at I = 0.025 M and (a) pH 5, (b) pH 7, and (c) pH 10. The reduced monomer concentration, $c/CMC^*$, normalizes the concentration differences between C12PC and C16PC. The arrows indicate the amount of negative charge (mmol/g) of PAHA in 0.02 M KNO₃ at the given pH values. Data from ref. [97].

Fig. 8. Binding isotherms for C12P⁺–AFA system as a function of pH at: (a) I = 0.03 mol dm⁻³ and (b) I = 0.10 mol dm⁻³, (●) pH 3.97, (○) pH 7.41, (□) pH 9.18, (△) pH 10.01. Data from ref. [99].

Fig. 9. Binding isotherms for (a) AHA system and (b) AFA system. Data from ref. [98].

Fig. 10. Binding isotherms of C16P⁺ to humic acid (PAHA) at two ionic strengths, (a)
log-normal plot with $H^+$ release, (b) double logarithmic plot, and binding isotherms of C12P$^+$ to anionic polyelectrolyte (SPSS) at three ionic strengths, (c) log-normal plot, (d) double logarithmic plot. Data from ref.[103, 112].

Fig. 11. Influence of cationic surfactant carbon-chain length on binding to purified Aldrich humic acid (PAHA). C16P$^+$ = cetylpyridinium; C12P$^+$ = dodecylpyridinium; 0.005 M NaCl. (a) log-normal plot, (b) double logarithmic plot. Data from ref. [103].

Fig. 12. Binding of cationic surfactant (dodecylpyridinium chloride, C12PC) to various humic and fulvic acids at 0.005 M NaCl and pH 5. PAHA, Aldrich humic acid; IHA, Inogashira humic acid; DHA, Dando humic acid; SFA, Strichen fulvic acid; LFA, Laurentian fulvic acid. (a) log-normal plot, (b) double logarithmic plot. Data from ref.[103].

Fig. 13. Benzyltrimethyldecyl-ammonium chloride (C12-BAC) sorption to purified Aldrich humic acid (AHA) determined at different background electrolytes at pH 6 (A) and at different pH in the presence of 5 mM NaCl or CaCl$_2$ (B). The isotherms were fitted with Freundlich equation with Freundlich exponent ($n_F$) of 0.8. Data from ref. [114].

Fig. 14. Adsorption isotherms of SNBS (sodium C9-benzene sulfonate; pH 4.1) and DPC (C12-pyridinium chloride; pH 8.0) on rutile at 0.01 M NaCl. Panel (a) is a double logarithmic representation of the isotherms and depicts the typical four-regions isotherms; panel (b) depicts the same results in a semi-logarithmic plot. Data from ref. [118].

Fig. 15. Adsorption isotherms of sodium nonyl-benzene-sulfonate (SNBS) on rutile at pH 4.1 and three salt concentrations are depicted as log-log (a) and lin-log (b) plots. Data from ref.[134].

Fig. 16. Adsorption of SNBS and the net proton adsorption, $\Gamma(0)$, on rutile as a function of the equilibrium SNBS concentration at three pH values and two NaCl concentrations; the pH$_{pzc}$ of rutile is 5.85. Data from ref. [134].

Fig. 17. Adsorption isotherms for C12PC on Aerosil, measured at pH 9 and two salt concentrations (a) as log-log plot and (b) as lin-log plot. Data from ref. [137].

Fig. 18. Effect of DPC adsorption on the surface charge of Aerosil, $\Gamma(0)$, at pH 9. Data from ref. [138].
**Fig. 19.** Adsorption of tertiary-octylbenzene polyoxyethylene glycol surfactants on Spherosil silica at 25 °C and pH 6.5. The TBE surfactants are monodisperse \([\text{TBE} = \text{C}_8\Phi(\text{EO})_{\text{NP}}]\), the TX surfactants are polydisperse \([\text{TX} = \text{C}_8\Phi(\text{EO})_{<\text{NP}}]\). NP is indicated in the figure. Data from ref. [160].

**Fig. 20.** Adsorption of C12PC on sodium kaolinite as a function of the NaCl concentration at pH 5 and 20 °C; the horizontal line indicates the isoelectric situation and the axis on the right side indicates the sign of the electrophoretic mobility. Data from ref. [179].

**Fig. 21.** Adsorption isotherms of C16TAC (=HDTMA) on kaolinite at pH 6.5 and two NaCl concentrations. The adsorption is expressed relative to the CEC at pH 6.5, which is 40 µmol/g. Data from ref. [184].

**Fig. 22.** Abstraction (adsorption + precipitation) isotherms of sodium dodecylbenzene sulfonate (SDBS) - kaolinite system at different pH (indicated), 0.1M NaCl and 25 °C. Data from ref. [195].

**Fig. 23.** Adsorption isotherms of three nonionic surfactants on kaolinite at pH 3.8 and T=25°C. The three surfactants are polydisperse alkylbenzene polyoxyethelenes form the Triton series: TX100 = C$_8$H$_{17}$ΦEO$_{9.5}$, TX165 = C$_8$H$_{17}$ΦEO$_{16}$ and TX305 = C$_8$H$_{17}$ΦEO$_{30}$. Data from ref. [161].

**Fig. 24.** Characteristic C16TAC (=HDTMA) adsorption isotherms on Na- and Ca-saturated montmorillonite (Wyoming montmorillonite, SWy-1; CEC = 900 µmol/g), compared with that on Na-kaolinite (Source Clays Repository, Dept. of Geology, University of Missouri, Columbia; CEC = 40 µmol/g at pH 6.5). The adsorbed amount is expressed relative to the CEC, the concentration of C16TAC (=HDTMA) is in mol/L. The clays were free of organic matter and homo-ionic (washing first with a NaCl solution and for Ca-clays subsequently with a CaCl$_2$ solution). Data from ref. [203].

**Fig. 25.** (a) Adsorption of C16TAC (=HDTMA) on Na-SWy-1 in 5 mmol/L NaCl solution and the concomitant Na release. (b) Electrophoretic mobility (solid symbols) and degree of dispersion stability (open symbols) of C16TAC-Na-SWy-1 as a function of the C16TAC adsorption; (c) d(001)-spacings of C16TAC-Na-SWy-1 obtained by X-ray diffraction as a function of C16TAC adsorption at three different water contents (dried at different relative humidity, the RH values are indicated). The C16TAC adsorption is expressed relative to the CEC (900 µmol/g). Data from ref. [184].
Fig. 26. Calculated density distributions for ammonium N, alkyl C, Ca$^{2+}$, Br$^{-}$ and water O for montmorillonite (Mt) with a CEC of 1060 $\mu$mol/g and different loading levels of C16TA$^+$/C16TAB relative to the CEC; the loadings are indicated in the figure. Data from ref. [213].

Fig. 27. Snapshots of the surfactant configurations in the Mt-C16TA+/C16TAB system. The panels a, b, c and d correspond with, respectively, Figs. 25a, 25b, 25c, and 25d. The stick layers represents the Mt-layers, the atoms in the interlayer are colored: Ca = green, N = blue, C = grey, Br = brown, H = white, O water = red. Data from ref. [213].

Fig. 28. Adsorption isotherms of C12(EO)8 on different layer silicates: (▼) Na-kaolinite, (■)Na-illite, (◆) Na-montmorillonite (Na-Mt) and (▲) Ca-bentonite (Ca-Mt). Data adapted from ref. [218].

Fig. 29. Effect of carbon chain length on linear alkylbenzenesulfonate (LAS) adsorption in a sediment. Data from ref. [255].

Fig. 30. Adsorption isotherms of cationic (C16TAB), nonionic (Tergitol 15-S-7 = C11-15(EO)7; secondary alcohol ethoxylate) and anionic (SDS) surfactants for sand. Data from ref. [266]

Fig. 31. Influence of pH on cationic (C16TAB), nonionic (Tergitol 15-S-7= C11-15(EO)7) and anionic (SDS) surfactant adsorption on sand. Data from ref. [266]

Fig. 32. (a) Influence of electrolyte concentration on the adsorption of the anionic surfactant, sodium dodecylbenzenesulfonate on highly humic non-allophanic andosol. The NaCl concentrations are indicated in the figure. Symbols are measured values. Solid lines are calculated with the Langmuir-Donnan model. (b) Donnan potentials obtained with the Langmuir-Donnan model. Data from ref. [258].

Fig. 33. Sodium dodecylsulfate (SDS) adsorption and degradation in a soil suspension (highly humic nonallophanic andosol) in relation to mixing time. Data from ref. [274].

Fig. 34. The adsorption of hexadecyltrimethylammonium (HDTMA=C16TA$^+$) on two vermiculitic soils (Na and Ca version) and the corresponding electrophoretic mobilities and optical densities of the soil suspensions. (a) Adsorption isotherms,
(b) electrophoretic mobilities of the soils and (c) optical densities of the soils. Data from [272].

**Fig. 35.** Soil hydraulic conductivity during percolation with a 1 mol/kg surfactant solution (A) nonionic surfactants (N1, N2, N3), (B) anionic surfactants (A1, A2), cationic surfactants (C1, C2) and amphoteric surfactant (Am). Data from ref. [279].

**Fig. 36.** Comparison of upward infiltration of sodium dodecylsulfate (SDS) solution at 70 mmol/L with that of pure water in a peat moss column. The water table connected with the column was set at height 0 cm. Data from ref. [299]
Fig. 1.
Fig. 2.
Fig. 3.
Fig. 4.
Fig. 5.
Fig. 7.
Fig. 9.
Fig. 10.

(a) C16P+–PAHA Binding or H+ Release (mol/kg) vs. C16PC Concentration (M)

(b) C16P+–PAHA Binding (mol/kg) vs. C16PC Concentration (M)

(c) C12P–SPSS Binding (mol/kg) vs. C12PC Concentration (M)

(d) C12P–SPSS Binding (mol/kg) vs. C12PC Concentration (M) with Critical Association Concentration (CAC) indicated.
Fig. 11.
Fig. 12.
Fig. 13.
Fig. 14.
Fig. 15.
Fig. 17.
Fig. 18.
Fig. 19.
Fig. 20.

Surfactant Adsorption ($10^{-6}$ mol/g) vs. log $C_{12}$

- NaCl 100 mM
- NaCl 20 mM
- NaCl 5 mM

Mobility

log $C_{12}$
Fig. 23.
Fig. 24.
Fig. 25.
Fig. 26.
Fig. 29
Fig. 32

(a) Adsorption (mmol/kg) vs. Surfactant Concentration (mM)

(b) Donnan Potential, \( \phi \) (mV) vs. Surfactant Concentration (mM)

- 500mM
- 100mM
- 10mM
- 1mM
- \( \phi = 0 \)
- NaCl
Fig. 33

Graph showing adsorption and degradation (mmol/kg) against surfactant concentration (mM) for different time periods: 26 hours (diamonds), 18 hours (circles), 8 hours (triangles). The x-axis represents surfactant concentration (mM) ranging from 0.001 to 100, and the y-axis represents adsorption/degredation (mmol/kg) ranging from 0 to 160.
Fig. 35