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Adsorption and Wetting Mechanisms at the Surface of Aqueous Hydrocarbon Solutions as a Possible Source of Atmospheric Pollution

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Abstract — Adsorption and Wetting Mechanisms at the Surface of Aqueous Hydrocarbon Solutions as a Possible Source of Atmospheric Pollution — Hydrocarbons in solutions have been the subject of very few investigations despite their relevance, in particular, in situations where environmental mechanisms are involved.

We present, here, a synthesis of several studies conducted within our laboratory about the adsorption, at the water surface, of benzene and cyclohexane from under-saturated solutions. The coadsorption of lead nitrate is also evidenced, though it does not adsorb in absence of organic molecules in the surface. Most of the data reported here were collected from series of measurements made with a very uncommon method: the bubble column; this technique, though being very time-demanding and lacking of user-friendliness, proved its usefulness and relevance in the study of such weakly adsorbable, surface-tension inactive compounds. The study of mixtures is simple and requires no model, which is precious in environmental research. The gathered data demonstrate that, through mechanical mechanisms such as bubbling, coadsorption can lead to the passing from water to the atmosphere of harmful non-soluble, poorly surface-active, components.

Résumé — Mécanismes d’adsorption et de mouillage à la surface de solutions aqueuses d’hydrocarbures comme source possible de pollution atmosphérique — Les solutions d’hydrocarbures ne sont pas très souvent étudiées. Cependant, leur comportement est impliqué dans de nombreux mécanismes, en particulier les mécanismes environnementaux. Dans le présent article, nous présentons la synthèse de plusieurs travaux ayant trait au comportement superficiel, en particulier d’adsorption, de solutions non saturées de benzène et de cyclohexane. Il y est mis en évidence la coadsorption de nitrate de plomb, bien que celui-ci, en l’absence de molécules organiques dans la surface, ne s’adsorbe pas du tout. Ces données ont été obtenues essentiellement au moyen d’une technique rarement utilisée, la colonne à bulles, brièvement décrite ci-dessous. Celle-ci, en dépit des contraintes de sa mise en œuvre, s’est révélée très utile dans l’étude de ces composés faiblement adsorbables, et peu actifs sur la tension de surface. L’étude des mélanges se fait directement, sans avoir à utiliser de modèle, ce qui est précieux pour des études environnementales. Les résultats obtenus établissent de façon convaincante que des mécanismes, tels que le pétilllement, joints à la coadsorption, peuvent entraîner le passage de composés nocifs pour l’environnement, peu solubles, peu actifs sur la surface quand ils sont isolés, depuis l’eau jusque dans l’atmosphère.
LIST OF MAIN SYMBOLS

\( \alpha \)  vapour phase in equilibrium with the liquid phase(s).
\( \beta \)  water-rich liquid phase.
\( \gamma \)  hydrocarbon-rich liquid phase.
\( \gamma^\circ \)  contact angle in a drop of \( \gamma \) lying on \( \beta \) phase in contact with \( \alpha \) phase.
\( J \)  solute concentration profile in the bubble column.
\( J_i \)  solute concentration profile of \( i \) component, in the bubble column.
\( K \)  proportionality constant in an adsorption isotherm of Freundlich type.
\( K_i \)  proportionality constant in adsorption isotherm of \( i \) component (Freundlich type).

INTRODUCTION

It is commonly admitted that the light hydrocarbons present in the atmosphere [1] come from industrial activities, road traffic, or evaporation from accidentally-spread layers. But, the contribution of the atmospheric dissolution of such species through vapour pressure equilibrium or surface adsorption and bubbling has been rarely considered in atmospheric pollution mechanisms [2-4]. However, in the vicinity of wrecked tankers with huge amounts of oil progressively leaking from their tanks and dissolving in the seawater, the passing of hydrocarbons from water to the atmosphere through surface mechanisms is worth being taken into account. This is also true near industrial wastelands where interstitial waters charged with dissolved hydrocarbons may feed with them ground waters, springs and surface waters. Moreover, adsorption is of a much greater concern whenever the presence of hydrocarbons induces the coadsorption of other harmful species unable to adsorb and be transferred otherwise.

In previous studies [5-6], we showed that, once adsorbed at the air/water interface, benzene and cyclohexane can induce the coadsorption of a heavy metal salt, namely lead nitrate. We selected this salt because of its solubility properties; in addition, it constitutes an excellent model for ions of concern in environmental problems [7]. The present report focuses on the specific role played by hydrocarbons in the experimental data collected over our experiments; it also gives more insight into the specific method imposed by the nature of our investigations.

Indeed, the study of hydrocarbon adsorption is uneasy because the method classically used to get air/water adsorption relies on the treatment of surface tension data by applying the Gibbs isotherm formula. Unfortunately, despite the well-established adsorption of hydrocarbons in noticeable amounts [9], the very low activity demonstrated by most of them on water surface tension imposes the use of different methods for studying surface properties. For instance, the

wetting of hydrocarbons has been investigated by ellipsometry [10-12].

However, in the study of benzene in solutions reported here the first parameter to be measured was surface tension because it can, conversely to other hydrocarbons, be determined close to the solubility limits. On the other hand, for very dilute solutions and hexane we applied the bubble column method [13], which is in fact a variant of the depletion method used to study adsorption on solid suspensions; in the case of a three-component mixture, this method allows one to determine the adsorption of each solute. But, since its implementation is quite difficult, it is worth being employed only in very specific cases where no other method is relevant. Whenever surface tensions are employed for three-component mixtures, one must be aware that their use requires that Gibb’s formula be written in a convenient form. The study of coadsorption is then a rather difficult task, however simplified in dilute solutions, which is the case here. As the mutual wetting properties of water-rich and hydrocarbon-rich phases in equilibrium at the solubility limit are now known to significantly affect adsorption process [14-15], the first part of this study will introduce such data.

1 THEORETICAL ASPECTS

1.1 Adsorption and Wetting

Cahn’s theory [14-16] shows that adsorption layers and complete wetting can be described in terms of phase nucleation. This is particularly valuable near a critical point where one of the coexisting phases completely wets the other one at the contact with vapour.

The \( B \) and \( C \) phase diagrams presented on Figure 1 are characteristic of binary systems each of them composed of two partly miscible liquids (e.g. water and hydrocarbon); these systems exhibit a wetting transition at a temperature \( T_W \), and \( x \) is the molar fraction of the hydrocarbon. Water-rich and hydrocarbon-rich solutions are respectively on the left and right sides of the diagram; solutions of both types coexist within the miscibility gap. Conversely to \( A \) diagram (which could be, for instance, nicotine/water or dimethylpyridine/water diagram) lower critical points are obviously missing on \( B \) and \( C \) diagrams; however, the shapes of both diagrams suggest that their occurrence has been prevented by crystallisation.

On this figure, it is worth focusing on the physical processes occurring along a horizontal path (at \( T < T_W \)) starting from the water-rich phase (\( \beta \)) below the wetting transition temperature: indeed, while moving closer to the coexistence line, the surface layer is progressively enriched in hydrocarbon until its composition and that of the \( \gamma \) phase become alike. The \( \beta \) phase is then totally wetted by the forming hydrocarbon-rich phase (\( \gamma \)). Because of similarities
between the nearly alike compositions of \( \beta \) and \( \gamma \) phases, most of the time this complete wetting is observed near the critical point (whenever it exists); let us denote \( \alpha \) the vapour, then the relation between the different surface tensions can be expressed as follows [16]:

\[
\sigma_{\alpha \beta} = \sigma_{\beta \gamma} + \sigma_{\gamma \alpha}.
\]

(1)

This situation corresponds to a null spreading coefficient, \( S = \sigma_{\alpha \beta} - \sigma_{\beta \gamma} - \sigma_{\gamma \alpha} = 0 \).

Far from the critical point, relation (1) becomes an inequality:

\[
\sigma_{\alpha \beta} > \sigma_{\beta \gamma} + \sigma_{\gamma \alpha}.
\]

(2)

This inequality always corresponds to the complete wetting of \( \beta \) by \( \gamma \) but, here, it corresponds to a positive spreading coefficient, \( S = \sigma_{\alpha \beta} - \sigma_{\beta \gamma} - \sigma_{\gamma \alpha} > 0 \). It is worth recalling that spreading coefficients are positive, negative or null depending on their definitions and on the wetting properties of the system under study [17-19].

But, one should note that the above reasoning about adsorption (see Fig. 1) is always valid under complete wetting condition and whatever the relation (1) or (2) between the surface tensions.

The reverse inequality:

\[
\sigma_{\alpha \beta} < \sigma_{\beta \gamma} + \sigma_{\gamma \alpha}.
\]

(3)

corresponds to a partial wetting of \( \beta \) by \( \gamma \). Let us consider a drop from \( \gamma \)-phase and denote the angle between the tangents to the three interfaces at the triple contact line; this angle is related to interfacial tensions by [16]:

\[
M = \frac{\left[ (\sigma_{\alpha \beta})^2 - (\sigma_{\beta \gamma})^2 - (\sigma_{\gamma \alpha})^2 \right]}{2\sigma_{\beta \gamma} \sigma_{\gamma \alpha}} (= \cos \hat{\gamma})
\]

(4)

where \( M \) is the wetting coefficient of \( \beta \) phase by \( \gamma \) phase in contact with \( \alpha \). \( M \) equals \( \cos \hat{\gamma} \) only when Equation (3) is proved. Relation 4 is the equivalent of the Young law for the wetting of solids. Equation (2) corresponds to a wetting coefficient \( M > 1 \). Passing from complete to partial wetting constitutes a wetting transition, which affects adsorption behaviour.

### 1.2 Bubble Column Theory

In the bubble column filled with the solution under study, numerous rising bubbles of small size carry upwards the species adsorbed at their surface, and then burst at the top (Fig. 2). The resulting concentration gradient is analysed at the time when the steady state is established.

According to the theory developed by Lemlich [20] and applied by Wan and Tokunaga [13], the fundamental hypothesis is Equation (5) where \( K \) is the proportionality constant; this relation assumes a linear variation of adsorption, \( \Gamma \), versus the bulk concentration, \( C \). In environmental studies [21], this isotherm equation is often termed the Freundlich isotherm (with exponent parameter equal to 1); but, in fact it is rather the limit form of the Langmuir isotherm at very low concentrations [8]. This approximate form can be used for almost all of the systems with very small adsorption values:

\[
\Gamma = K C.
\]

(5)

In this report, for better readability, the factor \( K \) will be named later “adsorption coefficient”.

At steady state and with respect to the species concentration at the column bottom, \( C_b \), the concentration profile \( C(z) \) along the column is:

\[
\frac{C(z)}{C_b} = \exp(Jz)
\]

(6)

with

\[
J = \frac{a/K}{AD}
\]

(7)
where \( A \) is the column cross-section, \( a \) is the surface area per bubble, \( f \) is the bubble generation rate (calculated from the measured gas flow and the bubble volume mean value), and \( D \) is the eddy dispersion coefficient; moreover, the gas within each bubble is supposed to be at atmospheric pressure. The parameter \( J \) characterises the separation of the solute from the solvent induced by the adsorption upon the ascending bubbles.

Moreover, integrating Equation (6) along the column height and expressing the mass conservation during the building of concentration gradient lead finally to:

\[
\frac{C(z)}{C_0} = \left[ \frac{HJ}{\exp(HJ) - 1} \right] \exp(Jz)
\]  

where \( C_0 \) is the species homogeneous initial concentration, and \( H \) is the height of the liquid column [13]. For each species present in the bubble column, the measurement of \( C(z) \) at steady state along the column versus the altitude \( z \) gives \( J \) from Equation (8). With this value of \( J \), additional measurements of \( A, a, f \) and \( D \) lead to \( K \) by solving Equation (7), and finally to \( \Gamma \) by using Equation (5). Usually, \( D \) can be determined from the expression of the time-dependent concentration profile and measurement of such profiles by different methods [13, 22]. Actually, we followed Wan and Tokunaga and took always the same value of \( D \) \((6.1 \times 10^{-4} \text{ m s}^{-1})\) and the same relative uncertainty, 0.1, because the characteristics of our column and system were alike theirs.

### 1.3 Adsorption from Surface Tension in the Case of Two Solutes

#### 1.3.1 Lead Nitrate Surface Content in Presence of Hydrocarbon (Benzene)

Let us consider a system of surface tension \( \sigma \) containing a surface layer; it consists of a salt (denoted \( s \)), composed of cations (+) and anions (-), benzene (B) and water (W).

Application of Gibbs Duheim’s relation gives:

\[
-d\sigma = \Gamma_s d\mu_s + \Gamma_\text{B} d\mu_\text{B} + \Gamma_W d\mu_W
\]

where \( \Gamma_i \) is the adsorption of \( i \) species.

Gibbs Duheim’s equation for the bulk writes:

\[
x_i d\mu_i + x_\text{B} d\mu_\text{B} + x_W d\mu_W = 0
\]

where \( x_i \) is i mole fraction.

Eliminating \( d\mu_W \) between (9) and (10) leads to Gibbs’ equation:

\[
-x_i d\mu_s + x_\text{B} d\mu_\text{B} + x_\text{W} d\mu_\text{W} = 0
\]

where \( x_i \) is the relative adsorption of \( i \) component with respect to water.

\[
\Gamma_{i,w} = \Gamma - \left( \frac{x_i}{x_\text{W}} \right) \Gamma_W
\]

Let us write \( \Gamma_{s,w} = \nu_+ \Gamma_{s,W} \) and \( \Gamma_{\text{B},w} = \nu_- \Gamma_{\text{B},W} \) where \( \nu_+ \) and \( \nu_- \) are the number of cations and anions in one molecule of salt, and then:

\[
d\mu_s = \nu_+ d\mu_+ + \nu_- d\mu_-
\]

Gibbs’ equation becomes:

\[
-x_s d\mu_s + x_B d\mu_B + x_W d\mu_W = 0
\]

With lead nitrate, \( \nu_+ = 1, \nu_- = 2 \), and the corresponding term \( \nu = \nu_+ + \nu_- = 3 \).

The next step is the evaluation of \( d\mu_B \) as it may manifest in an experiment where the molarity \( c_B \) has been kept constant contrarily to the formality \( c_S \). All chemical potentials in the solution are related through the bulk Gibbs Duheim equation (10), which writes:

\[
x_s d\mu_s + x_B d\mu_B + x_W d\mu_W = 0
\]
where:
\[
\mu_s = g_s^0 + RT \ln \left[ \gamma_y^{v_y} \gamma_y^{v_y} \nu \nu \cdot c_y^{(v_y)} \right]
\]
or:
\[
\mu_s = g_s^0 + RT \ln \left[ \gamma_y^{v_y} \nu \nu \cdot c_y \right]
\]
and:
\[
d\xi = RT d \ln \left[ \gamma_y^{v_y} c_y \right] = \left( \nu \nu + \nu \nu \right) RT d \ln \left( \gamma_y c_s \right) \tag{14}
\]
where:
\[
\gamma_y = \gamma_y^{v_y} \gamma_y^{v_y} \nu \nu
\]

One should keep in mind that the solvent is water; by using an unsymmetrical reference system for the activities, (13) becomes:
\[
x_v R T d \ln \left( \gamma_y c_s \right) + x_B R T d \ln \left( \gamma_y c_B \right) + x_W R T d \ln \left( f_w-x_w \right) = 0 \tag{15}
\]

Equation (15) is general for aqueous solutions containing a neutral molecule, B, together with an \( (\nu \nu - \nu \nu) \) electrolyte. By taking into account that \( c_s \) and \( c_B \) are around \( 10^{-2} \) mol L\(^{-1}\), one litre contains \( c_s \) moles of salt, \( c_B \) moles of B and about 55.5 moles of water. So, \( x_s = \alpha = c_s/55.5 \) and \( x_B = \beta = c_B/55.5 \) and \( x_W = 1 \) whereas \( f_w = 1 \) because, in very dilute solutions, the solvent obeys Raoult’s law. Finally, as experiments were made at constant \( c_B \), equation (15) becomes:
\[
\alpha v R T d \ln \left( \gamma_y c_s \right) + \beta R T d \ln \gamma_B + RT d \ln x_W = 0 \tag{16}
\]

Further approximations are still possible. First, according to the Debye Hückel law, \( \gamma_n \) can be written as:
\[
\log \gamma_n = -B \left[ c_s z_+ \left( \frac{1}{2} \sum c_s z_+^2 \right) \right]^{1/2}
\]
\[
= -B \left[ c_s z_+ \left( \frac{1}{2} \left( \nu \nu + \nu \nu \right) c_s \right) \right]^{1/2}
\]
where \( B \) is a solvent- and temperature-dependent constant.

Introducing for further use the term allows us to write:
\[
d \ln \gamma_n = -\frac{2.303}{2} B \left[ c_s z_+ \left( \frac{1}{2} \left( \nu \nu + \nu \nu \right) c_s \right) \right]^{1/2} \left( \frac{dc_s}{c_s} \right) \tag{17}
\]

In addition, \( d \ln (x_W) = d \ln (1-\alpha-\beta) = d (-\alpha-\beta) = -d \alpha \) as \( \beta = x_B = c_B/55.5 \) is constant in our experiments.

Consequently:
\[
\alpha v R T d \ln \gamma_y + \beta R T d \ln \gamma_B + RT d \ln x_W = 0 \tag{18}
\]

and finally, for a 1-2 electrolyte at 25°C, in an aqueous solution where \( B = 0.5 \), we get:
\[
\beta d \ln \gamma_B = -2 + 6 \left( 3 c_s^2 \right) \frac{d \alpha}{\alpha} \tag{19}
\]

Equation (19) is valid for dilute solutions where \( c_B \) is constant. It expresses the “salt effect” exerted upon the neutral molecule B in solution by the salt at the concentration \( \alpha \) (molar fraction). The induced variation of \( \ln \gamma_y \) directly displaces the solubility limit of B; this constitutes the best evidence of the “salt effect”.

By taking into account relations (13), (17) and (20), we can rewrite Equation (12) as:
\[
-d \alpha = RT \times \left[ \frac{\ln \alpha - \alpha v B}{c_s} \left[ \frac{1}{2} \left( \nu \nu + \nu \nu \right) c_s \right] \right] \tag{21}
\]

or:
\[
\Gamma_{B(NO_3)_2, w} = \left( 1 - 2 c_{B(NO_3)_2} \right)^{1/2} \times \left[ -\left( \frac{1}{3RT} \right) \left( \frac{d \alpha}{d \ln c_{B(NO_3)_2}} \right) + \left( \frac{2}{3} - 2 c_{B(NO_3)_2} \right) \frac{1}{2} \frac{c_{B(NO_3)_2}}{c_B} \Gamma_{B(w)} \right] \tag{22}
\]

### 1.3.2 Determination of Benzene Surface Content in Presence of Lead Salt

At constant \( c_s \), and with the same approximations about water as above, (16) leads to:
\[
\alpha v R T d \ln \gamma_y + \beta R T d \ln \gamma_B c_B + RT d \ln x_W = 0 \tag{23}
\]

So, (12) can be expressed as:
\[
-d \alpha = \Gamma_{S,W} \times R T d \ln \gamma_y + \Gamma_{B, W} R T d \ln \gamma_B c_B \tag{24}
\]

and with \( \log \gamma_y = -2 \left( 3 c_s^2 \right)^{1/2} = \text{const} \)
relation (19) leads to:

\[ c_B d \ln \gamma_B = \left[ -2 + 6 \left( \frac{c_S}{\gamma} \right)^2 \right] d c_S \]  \hspace{1cm} (26)

If the approximation contained in Equations (25) and (26), namely the Debye-Hückel constant, \( B \), is independent of benzene concentration, \( \gamma_s \) and \( \gamma_B \) are both constant, and Equation (24) very simply writes:

\[ -d\sigma = \Gamma_{B,W} RT d c_B \]  or \[ \Gamma_{B,W} = - \left( \frac{1}{RT} \right) \left( \frac{\partial \sigma}{\delta \ln c_B} \right)_{c_s,T} \] \hspace{1cm} (27)

2 MATERIALS AND METHODS

2.1 Bubble Column

Most of the device characteristics depicted on Figure 2 were taken from the description by Wan and Tokunaga [13]. The column consists of a 2-m-high tube of Pyrex; its inner diameter is 19 mm. Eight septum-capped sampling ports are distributed along the length of the tube with shorter intervals toward the top end. Its bottom end is a 5-mm high gas chamber, whose ceiling is made of fifth grade fritted glass plate; to maximise bubble concentrations, its diameter is greater than the column cross-section. When the column is filled with a solution, injection of Argon gas into the gas chamber below the fritted glass plate generates bubbles, which rise up across the column of liquid. The gas flow is controlled and measured with a smart DMFC metal-sealed ultra high purity mass flow-meter and -controller, model 6256S by Brooks Instruments. As benzene and cyclohexane are rather volatile solutes, to avoid any loss of vapour at the top of the column during experiments, we added there a 15-cm-high double walled glass cooled at 6°C to be just below the freezing point of benzene. Finally, to get a fairly constant inner temperature in the column we coiled up a plastic and thermoregulated water-fed tube around the tube of Pyrex to set the temperature within the column at 25.0 ± 0.1°C.

Under the above conditions for the column, a gas flow rate of about \( 1.66 \times 10^{-7} \text{ m}^3 \text{ s}^{-1} \) (measured under standard conditions) produced relatively high populations and fast streams of bubbles of small sizes. The diameters of bubbles were measured photographically with a DXC-107A/107AP Color Video Camera (Sony) equipped with a manual macro zoom with a 10 \( \times \frac{1}{2} \) C frame and 18/180 mm focal length. The camera was connected to a computer for image acquisition. For scaling purpose, a 15-mm-diameter flat glass showing a micrometer was installed at the mid column section away from the wall for photographing. Photographs were taken at a fast shutter speed (1/2000 s) under the standard gas rate and registered by the Capturix VideoSpy software. The sizes of bubbles were determined, bubble after bubble, with the Jasc Paint Shop Pro7 software. For a given solution, the total surface area of 100 photographed bubbles was calculated on assuming spherical symmetry; the average bubble diameter was found to be \( 260 \pm 11 \mu \text{m} \) and led to \( \sigma = 0.22 \pm 0.02 \text{ mm}^2 \) as average area.

The time required to get a stable profile was tested by repeating the experiments with different solutions always at the same concentrations and varying time. We observed a concentration gradient only with very diluted solutions, in the range \( 10^{-5}-10^{-4} \text{ mol L}^{-1} \). Correctness for time and concentration was simply tested by verifying that a constant concentration was obtained at the upper septum after a certain period of time. Then, the concentration profile was determined. At steady state, 1.5 ml samples were withdrawn with syringes through the sampling ports in descending order. Samples of benzene solution were poured into closed 20 cm\(^3\) flasks containing each 18 cm\(^3\) of a 50-50 (v/v) water-methanol mixture prior to their measurement by UV spectroscopy. Samples of cyclohexane were introduced in vapour-tight, septum-closed, 10 cm\(^3\) flasks. According to the nature of samples, the flasks were either used directly (head space method for cyclohexane) or to transfer the samples to the electrochemical device (lead ion determination).

2.2 Analytical Methods

2.2.1 Hydrocarbons

We checked the benzene content of solutions by UV spectroscopy according to the method described in [23]. But, as this method was inappropriate for cyclohexane, for this hydrocarbon we successfully used head-space gas chromatography [24].

A 5-cm\(^3\) sample of aqueous hydrocarbon solution was placed in a previously washed and dried 10-cm\(^3\) vapour-tight flask closed with a septum. For every sample, the flask was systematically stored for 20 min in a thermostatically regulated oven heated at 60°C to reach equilibrium within the flask headspace between dissolved hydrocarbon and vapour. Then, we withdrew a 1-cm\(^3\) aliquot of vapour and injected it immediately in a DANI Educational type gas chromatograph equipped with a splitless injector at 250°C and a 300°C FID type detector; the vector gas was nitrogen. The oven was maintained at 100°C. The column used for separation was a 30-m long, 0.32-mm-diameter SPB1 apolar one with a 0.25-μm-thick film of poly-dimethyl-siloxane as stationary phase.

After standardisation and adjustment of the method, a stock solution was made by shaking at 25°C for 3 weeks a known weight of cyclohexane in water; it is worth noting that dissolved quantity was ten-fold smaller than the solubility limit. Then, this solution was diluted to prepare a series of standard solutions; aliquots were treated as described above,
and vapour was analysed by chromatography to make a
calibration curve by plotting a straight line between the chro-
matographic peak area and the cyclohexane concentration.

For cyclohexane and benzene, repeatability was assessed
and found to be about 1% by successively withdrawing
6 samples from the bubble column through the same
mid-column septum.

2.2.2 Lead Ions (Pb++)

Solutions containing either pure lead nitrate salt or lead
nitrate and one among the hydrocarbons under study were
taken from the bubble column through septa as previously
described. Our experimental conditions, i.e. hydrocarbons far
from their solubility limit, made us assume that their adsorp-
tion was unaffected by the presence of salt; we, therefore,
only analysed the salt content. One should, however, note
that cyclohexane and salt were simultaneously determined
only in one sample. This will be detailed in the Results
section to demonstrate that our hypothesis stands.

Pb++ concentrations were determined by Differential Pulse
Anodic Stripping Voltammetry, DPASV [25]. The electro-
chemical system consisted of a Hanging Mercury Drop
Electrode (663 VA stand, Metrohm) connected to a voltam-
metric analyser (microAutolab, Eco Chemie). Current and
potential were sampled with a personal computer. All poten-
tials were referred to the Ag/AgCl, 3 mol L-1 KCl electrode.
A platinum wire was used as a counter electrode. The quartz
voltammetric cell was thermostatted at 25.0 ± 0.1°C. The
concentration in salt being about 10-5 mol L-1, the samples
were systematically ten-fold diluted with a 10-2 mol L-1
nitric acid solution to slightly acidify the medium and pre-
vent metal from adsorbing on cell walls. The samples were
deoxygenated for 10 min with water-saturated nitrogen. The
duration for accumulation was 120 s under a −0.6 V potential,
and the equilibration time was 10 s. The voltammetric modu-
lation time was 50 ms, with 1 s as interval time, and the mod-
ulation amplitude was 50 mV with a step potential of
20 mV.

2.3 Surface Tensions and Wetting

Surface tensions were measured with the Wilhelmy method
while taking great care about cleanliness and control of tem-
perature. The plate was always burnt prior to measurement to
be sure of the quality of platinum surface. During measure-
ments, the solutions were kept within a double-walled glass
cell thermostatted at ± 0.1°C. The cell was placed in an
even thermostatted at the same temperature at more or less
0.5°C. Tensions were measured with a LAUDA TE1C ten-
siometer by taking the maximum force value, which means
making the measurement at the exact time the liquid surface
touched the plate while the cell was moving up. Reproducibility of measurements was carefully checked on
taking account of all among the other possible factors, i.e.
benzene volatility, adsorption kinetics; etc. Consequently, the
error on surface tension was estimated to be 0.001 σ.

To study the reciprocal wetting of mutually saturated
water-hydrocarbon mixtures, in agreement with equations
(1 to 4) we measured, at first, the surface tension of both
phases against air, then their interfacial tension. After equili-
bration of both phases (see below) and resting, we pipetted
30 cm³ of the β, water-rich, denser phase, poured them in
the cell and measured the surface tension, σbc. Then, the
Wilhelmy plate was totally sunk in the aqueous phase; we
gently poured 30 cm³ of the γ, hydrocarbon-rich, lighter phase
onto the aqueous phase, and measured the interfacial tension
σγb by taking out the plate from β to γ. Finally, after careful
cleaning of the plate, we determined the surface tension of the
γ upper phase, σγc.

2.4 Chemicals

To get true molecular solutions of hydrocarbons, we first pre-
pared saturated solutions by gently stirring with water just a
little more than the necessary quantity of benzene [5] or
cyclohexane. For benzene, the solubility determined at 25°C
was 2.37 × 10-2 mol L-1 [5]; this value agrees with literature
data [26-27]. According to [28], the one for cyclohexane is
6.55 ± 0.27 × 10-4 mol L-1; this value is very close to our
own estimation.

Water was purified on a milliQ device from Millipore and
its surface tension was measured every day. Benzene, Normapur minimum 99.7 wt%, was supplied by Prolabo.
Cyclohexane Chromanorm 99.7 wt% for HPLC and lead
nitrate, pro analysis, were supplied by Merck. These com-
pounds were used with no further purification. The surfactant
used to standardise the column was dodecylbenzenesulfonic
acid, sodium salt, technical 88% from ACROS Organics;
considering its strong adsorption ability made us use it with
no additional purification.

Under our experimental conditions, all the verifications
we made demonstrated the lack of complexation for Pb++
ions [5-6].

3 RESULTS AND DISCUSSION

3.1 Wetting

Table 1 lists the interfacial tensions measured by tensiometry
at 25 and 30°C in saturated biphasic mixtures of water—
benzene as well as in water—cyclohexane; α means vapour,
β denotes aqueous phase, γ refers to oily phase and ɣ is the
angle defined above.

One should underline that all measurements were made on
the same day and on same biphasic samples. Moreover, the
errors made about the benzene system (Table 1) were
estimated from a *repeatability* assessment conducted only on the aqueous phase. We should have done the same for the other interfaces, but it was too difficult; so, the error values about these data are likely slightly biased. But the conclusion about wetting is obviously valid.

The errors about the cyclohexane system were calculated from *reproducibility* estimation since the measurements had been made on the same day for each set of $\sigma$ ($\sigma_{\text{ftr}}$, $\sigma_{\text{fbr}}$, $\sigma_{\text{rt}}$) and repeated several times on different samples and days. However, one should note that the lack of systematic check of their concentrations contrarily to what was done for aqueous solutions of benzene may explain the larger dispersion of data that prevents us from concluding whether wetting is just complete, or slightly above, or slightly below; we can only say that wetting is very close to complete. In both cases, the oil-rich phase spreads totally, or nearly totally, onto the water-rich phase; it occurs more easily with the benzene system because of the greater similarity between the concentrations of both phases. It is worth noting that the difference in wetting between systems results from the outstanding difference between the values of interfacial tension. Indeed, $\sigma_{\text{fbr}}$ was always found to be lower with benzene, likely because it vanishes in the close vicinity of a critical point where both concentrations are nearly alike [29-34]. In this study, since we were very far from a critical point, the system with closer concentrations are nearly alike [29-34]. In this study, since we were very far from a critical point, the system with closer concentrations are nearly alike [29-34].

### 3.2 Adsorptions

To test the bubble column, we used SDBS and compared the results with those issued from surface tension measurements. SDBS is very surface-active, even in very dilute solutions where the use of the bubble column is relevant. At 25°C and concentrations for solutions in the range $10^{-5}$-2 $\times 10^{-4}$ mol L$^{-1}$, the surface tension varies linearly with $C$, the solution concentration, between 70 and 50 mN m$^{-1}$. Then, the adsorption provided by the Langmuir isotherm formula can be simply expressed as:

$$ \Gamma = K_C C \quad (28) $$

where $K_C = 4.703 \times 10^{-5}$ m.

The experiment made at $C_0 = 1.99 \times 10^{-5}$ mol L$^{-1}$ *with the bubble column* under conditions already described (namely, $f = 1.666 \times 10^{-7}$ m$^3$ s$^{-1}$, $D = 6.10 \times 10^{-2}$ m s$^{-1}$, $A = 3.14 \times 10^{-4}$ m$^2$) led to $J = 0.850$ m$^{-1}$, and from Equation (5) to $K = 4.780 \times 10^{-5}$ m, which is very close to the value found for $K_C$.

The excellent agreement between these data mainly results from the good adsorbability and separability of the surfactant that led to a high $J$ and permitted its very precise determination (0.35%). In the other experiments, the values found for $J$ were far much lower.

The adsorption constants of hydrocarbons in water, $K_C$ and $K_B$ for cyclohexane and benzene, respectively, were determined at 25°C under various experimental conditions and for different concentrations of hydrocarbon. On the other hand, for the determination, at 25°C, of $K_{Pb,C}$ and $K_{Pb,B}$ in the case of lead nitrate in aqueous solutions of cyclohexane and benzene, respectively, the only factor to be varied was the lead salt concentration. Even though the concentration in hydrocarbon was kept unchanged, it needed to meet two unavoidable constraints:

- produce a surface content high enough to induce metallic salt adsorption;
be sufficiently low to permit the observation of a concentration gradient in the bubble column.

As previously underlined, such an observation is impossible when the mean concentrations in the column are too high. The narrowness of the operating window made us carry out our investigations in presence of salt at a single hydrocarbon concentration.

Table 2 introduces our results, and the experimental data at the origin of these values are reported in Figures 3 to 6 as bubble column-normalised concentration profiles ($C(z)/C_0$), with $i$ indexing the species under study. The profiles displayed on Figures 3 and 4 are those of cyclohexane and benzene in solutions, respectively; Figures 5 and 6 show those of lead salt in presence of cyclohexane and benzene, respectively. At similar $a$ and $f$ values, such a plot allows one to check the consistency of data with the hypothesis of Equation (5), namely the adsorption proportionality to bulk concentration (constant $K$). Equation (8) then leads to the $J$ parameter, which was exploited through Equation (7) to get $K_i$ where $i$ was $C$ or $B$, with or without a co-adsorbate. In Equation (7), $A$ and $f$ were given the following values $A = 3.14 \times 10^{-4} \text{m}^2$, $f = 1.666 \times 10^{7} \text{m}^3 \text{s}^{-1}$. After determination, the $a$ parameter was considered to be $a \approx 0.22 \text{mm}^2$ with a dispersion of 12% for every experimental run. Figures 2 to 5 also display the lines representing the data issued from Equation (7) and highlight a quite good agreement between lines and experimental points. Reproducibility on $J$ was assessed to be 10% through six complete runs of experiments made at the same concentration of benzene, but on different days. A numerical analysis of the confidence degree on $J$ according to the numerical procedure at the origin of its calculation gave also 10%.

As $J$ was assumed to be constant, it was calculated numerically from Equation (8) under the linearised form where the slope is exactly $J$:

$$\ln \frac{C(z)}{C_0} = Jz + \ln \left[ \frac{HJ}{\exp(HJ) - 1} \right]$$  \hspace{1cm} (29)

All the series of data corresponding to different runs on the same system can be drawn on the same straight line; So, $J$ was determined with a regression method to minimise the sum of square differences between the experimental values of:

$$\ln \frac{C(z)}{C_0}$$

To estimate the confidence degree on $J$ from the numerical determination, we used a Gauss-Newton method [35]. For SDBS and benzene, $\Delta J/J = 0.35\%$ and $\Delta J/J = 3.5\%$, respectively; for cyclohexane and lead nitrate in any hydrocarbon mixture $\Delta J/J = 4.5\%$.

| Species $i$ symbols | $K_i/m$ | $J_i/m^3$ | $n$ | $\Gamma_i$ or $\Delta \Gamma_i$/mol $m^2$ | $\Gamma_H$/mol $L^1$ | $\Delta \Gamma_{Pb,H}$/mol $L^1$ |
|---------------------|---------|-----------|-----|----------------------------------------|---------------------|-------------------------|
| SDBS               | $(4.78 \pm 0.55) \times 10^{-5}$ | $0.850 \pm 0.003$ | (1) | $9.35 \times 10^{-7}$ | $1.99 \times 10^{-5}$ | |
| C                  | $(3.73 \pm 0.97) \times 10^{-6}$ | $0.09 \pm 0.004$ | (3+1) | $(0.09-1.49) \times 10^{-7}$ | $(2.47-40.0) \times 10^{-6}$ | |
| B                  | $(1.05 \pm 0.26) \times 10^{-6}$ | $0.023 \pm 0.0008$ | (6) | $(3.15-6.30) \times 10^{-7}$ | $(3.0-6.0) \times 10^{-4}$ | |
| Pb,C               | $(3.27 \pm 0.85) \times 10^{-7}$ | $0.0079 \pm 0.0004$ | (2) | $(+1$ at $C_{Pb,C}=2.0 \times 10^{-5})$ | $1.49 \times 10^{-7}$ | $(3.13-4.41) \times 10^{-9}$ |
| Pb,B               | $(0.29 \pm 2.16) \times 10^{-7}$ | $0.02 \pm 0.0007$ | (2) | $4.0 \times 10^{-5}$ | $(0.96-1.35) \times 10^{-5}$ | $5.0 \times 10^{-4}$ | $(1.5-2.35) \times 10^{-5}$ |
However, the difference between $K_c$ limits, cyclohexane adsorbs more than benzene. Of water [36] responsible for an enhancement of benzene electron-rich population of the cyclohexane results from possible interactions between the (slightly) greater affinity of water for benzene compared to the same origin as the difference in solubility limits: the marked effects upon adsorption are worth being discussed. Explained, because of the uncertainties on ligible and the other ones being estimated as previously estimated uncertainties.

According to Equation (30), the errors on $A$ and $f$ being negligible and the other ones being estimated as previously explained, because of the uncertainties on $K$ values, only the marked effects upon adsorption are worth being discussed. The discussion about $K_c$ and $K_b$ values in the case of only two components is much easier than under other conditions. Indeed, their determination on concentration ranges centred at $10^{-5}$ and $5 \times 10^{-4}$ mol L$^{-1}$ for cyclohexane and benzene, respectively, ensures the validity of the values found for $K$ because it corresponds to a true linear behaviour at the start of adsorption isotherms as assumed in Equation (5). Though the bulk concentrations were closer to the solubility limit of cyclohexane ($((6.55 \pm 0.27) \times 10^{-4}$ mol L$^{-1}$) than to that of benzene ($((2.37 \pm 0.02) \times 10^{-2}$ mol L$^{-1}$), they were very far from them. Finally, as $K_c$ is almost four times $K_b$, it means that, in very dilute solutions and far from the influence of solubility limits, cyclohexane adsorbs more than benzene.

However, the difference between $K$ values has likely the same origin as the difference in solubility limits: the (slightly) greater affinity of water for benzene compared to cyclohexane results from possible interactions between the electron-rich population of the π system and hydrogen atoms of water [36] responsible for an enhancement of benzene solubility. In dilute solutions, solute-solvent interactions are known to play a pre-eminent role.

Our measurements of lead nitrate adsorption were made at a single bulk initial concentration for each hydrocarbon: namely $4 \times 10^{-5}$ and $5 \times 10^{-4}$ mol L$^{-1}$ for cyclohexane and benzene, respectively. These bulk concentrations correspond to adsorptions of $1.57 \times 10^{-7}$ and $5.31 \times 10^{-7}$ mol m$^{-2}$, respectively, imposed by experimental constraints and calculated by using the $K_c$ and $K_b$ values determined in hydrocarbon solutions. However, a determination of $K_c$ made in presence of lead nitrate (Table 1) gave the same value and evidenced the correctness of our estimated values. Thus:

- very far from hydrocarbon solubility limits, the salt effect on adsorption is quite negligible,
- in these composite systems, the establishment of parallel concentration profiles, both governed by the same rules (Equations 5 to 9) seems unaffected by the bubble-induced perturbations.

The more general conclusion drawn from Table 2 data about lead salt adsorption is that this process occurs even with water-dissolved traces of hydrocarbon. Lead adsorption constants are about ten-fold less than that of cyclohexane and alike that of benzene:

$$\frac{K_{pb,c}}{K_c} = 0.084 \quad \text{and} \quad \frac{K_{pb,b}}{K_b} = 0.86$$

(31)

The adsorption ratios between a hydrocarbon and a metallic salt for these dilute solutions can be calculated from the ratios of adsorption constants as defined by Equation (5).
Assuming that $K_{p_b,H}$ is independent of the $H$ concentrations, which has not yet been experimentally proven, leads to:

$$\frac{\Gamma_{p_b,C}}{\Gamma_C} = 0.084 \frac{C_{p_b,C}}{C_C} \tag{32}$$

and:

$$\frac{\Gamma_{p_b,B}}{\Gamma_B} = 0.86 \frac{C_{p_b,B}}{C_B} \tag{33}$$

Equations (31)-(33) can then be read as follows: for the same bulk concentration ratio ($C_{p_b,H}/C_{p_b,H}$), the adsorption ratio ($\Gamma_{p_b,H}/\Gamma_H$) for benzene is ten-fold that of cyclohexane; thus, one mole of benzene attracts ten times more moles of lead salt than cyclohexane does.

Figure 7 summarises Table 2 under the form of adsorption isotherms for pure benzene, cyclohexane and lead salt in presence of one of these hydrocarbons. The resulting graph highlights the concentration ranges where measurements were possible and facilitates a visual comparison of the different values of $K$ corresponding to the slopes of straight lines constituting the isotherms. Figures 8 and 9 gather both types of data presented uniquely versus hydrocarbon concentrations, and allow one to visually compare both systems. Figure 8 compares not only the adsorption of the pure hydrocarbons, but also the salt one at constant salt concentration versus the corresponding concentration of hydrocarbon. These isotherms are simply the linear interpolation between zero and a point calculated from Table 2 for a salt content equal to the highest concentration of hydrocarbon used in our experiments. Under these conditions and as expected from equations (32) and (33), salt adsorption is stronger in the presence of benzene than in the presence of cyclohexane. The linear interpolation is rather hard, but data recently obtained within our laboratory on another system showed the relevance of this hypothesis [37].

Figure 9 was obtained by drawing salt adsorption isotherms for the same concentrations of salt, but different ones for hydrocarbons (see the experimental constraints already mentioned, which restricted the ranges of values given in Table 2). At low hydrocarbon content, which corresponds to the case where the comparison is realistic because of the solubility limit of cyclohexane, salt adsorption for identical values of $C_H$ (along a vertical line) is reduced when cyclohexane is replaced by benzene; this observation is puzzling because it disagrees with the above analysis. This discrepancy comes from either a wrong building of the salt isotherms, or the behaviour observed here means that that adsorption is more affected by the $C_{p_b,H}/C_{salt}$ ratio than by the nature of the hydrocarbon under study. One should note that Figure 9 allows such a comparison only under certain conditions; in the case of cyclohexane, because of its solubility limit, the ratio is far much higher.

Finally, for solutions of pure benzene, Figure 10 highlights a marked decrease of surface tension near the solubility limit responsible for a strong enhancement of adsorption as expected from wetting data. Figure 11 illustrates the changes in surface tension induced by the presence of lead nitrate in the solutions of benzene, and Figure 12 gives the corresponding salt adsorption values calculated as reported in Section 1.3. As already observed in dilute solutions, the adsorption of salt is dependent on benzene and salt concentrations.
The influence of organic molecules on ionic adsorption can be crudely explained from the premises of Samaras and Onsager’s theoretical treatment. As already described in the literature (see for example, reference [38]) or observed here on lead nitrate solutions, electrolyte solutions in water show the depletion of electrolyte from the surface. To explain this phenomenon Onsager and Samaras [39], themselves inspired by Wagner [40] (see also [41]) begun their model by using the classical method of electrostatic images. This method states that each charge, $q$, in a medium, $A$, of dielectric constant, $\varepsilon$, situated at the distance, $z$, from the plane boundary between $A$ and a second medium, $B$, of dielectric constant, $\varepsilon'$, (Fig. 13) can be considered as being under the influence of an image charge situated in $B$ at the position $-z$ and whose value $q'$ is:

$$q' = [(\varepsilon - \varepsilon')/(\varepsilon + \varepsilon')] q$$ \hspace{1cm} (34)

Let water and air be respectively $A$ and $B$, it leads to $\varepsilon = 78$ and $\varepsilon' = 1$, and thus:

$$q' = q$$ \hspace{1cm} (35)
As \( q' \) and \( q \) signs are alike, the interaction force is repulsive and leads to ion depletion near the surface.

Let us, now, suppose that \( q \) enters a surface layer where \( \varepsilon = 2 \), i.e. about the dielectric constant of organic solvents like cyclohexane or benzene.

Then:

\[
q' = \frac{q}{3}
\]

(36)

The coulombic force between \( q \) and \( q' \) is still repulsive (both signs are still alike), but strongly reduced. Consequently, a state of the system where \( q \) is embedded in the organic layer is less energetic and must be observed.

Then, as charge \( q \) and the charges of similar sign will be roughly situated closer to the surface, the number measuring their adsorption will become larger. It is clear that other effects such as the screening effect on \( q \) of the other charges present in the solution or the dispersion forces due to species polarisability as underlined in reference [42] have to be taken into account.

Our experiments obviously showed a reduction of ion depletion in presence of benzene and cyclohexane; but, it is worth noting that, despite their almost identical dielectric constants, salt depletion is different in both cases. Either the simple argument of image forces proposed above needs to be used in a more elaborated way, or it is insufficient to explain the experimental observations as suggested by references [41] and [42].

However, it is worth keeping in mind that a thick and dense organic molecular layer at the surface will favour ionic adsorption; it will be all the more true as its properties will be closer to those of an organic liquid. This finding explains why, on Figure 8, ion adsorption and that of the organic solvent are concomitantly increasing.

### 3.3 Environmental Impact

To discuss the impact of this physical behaviour on the environment, one should, first, take into account the order of magnitude of concentrations. In this study, the solute contents were rather close to those usually found in the environment: for example, lead was at \( 10^{-5} \) mol L\(^{-1} \), i.e. 2.07 mg L\(^{-1} \), i.e. just four times the upper limit found in natural fresh waters (0.5 mg L\(^{-1} \) in quite fresh water, 0.03 mg L\(^{-1} \) in hard water and 0.0039 mg L\(^{-1} \) as the mean value for natural water in continental North America according to [43]).
Adsorption isotherm of lead nitrate at 25°C from solutions using three different “solvents”, i.e. water with three different contents of benzene:

\[ c_B^1 = 0.915 c_0B \] (•);
\[ c_B^2 = 0.83 c_0B \] (+) and
\[ c_B^3 = 0.75 c_0B \] (▲).

The adsorption values were obtained from Equation (22) by using the surface tensions displayed on Figure 11 and the \( \Gamma_B \) values on Figure 10. The dashed line shows the salt concentration that sets the solubility limit of benzene at concentration \( c_B^1 \).

Let us now consider the role of coadsorption in bubbling and via-aerosol pollutant transfer. Bubbling leads to the formation of aerosols through the four-step process described in Figure 14:

- inclusion of air bubbles close to the water surface under the action of wind;
- bursting of bubbles giving birth to very small drops blown away by wind;
- streaming of the surface layer along the surface of the hole remaining after the bubble bursting;
- formation of a jet drop while this hole is closing.

Step 1 leads to the creation of a surface that holds an adsorption layer containing the adsorbable pollutants present in solution. The drops created in step 2 are formed from the bubble wall mainly constituted of the adsorbed layer: these droplets enriched by adsorption will later form polluted aerosols. Step 3 explains how the jet drop formed in step 4 is in turn enriched in adsorption layer components. “Tearing”, which leads to the formation of “spray drops” at the wave edge or at its breaking, also contributes to aerosol formation and implies adsorption layers.

So, in ordinary environmental conditions it is paramount to consider the process of coadsorption and its associates, bubbling and via-aerosol transfer because of the significance of adsorption values shown in the present study.

Very few investigations have been focused on aqueous solutions of hydrocarbons though in industrial wastelands, water sheets polluted with hydrocarbons at trace levels and variable quantities of heavy metal salts frequently exist. Their contribution to, at least, local atmospheric pollution should be investigated. It is likely that the case of salted water sheets is worth being considered. Even though NaCl was missing in this study, the results reported here give some precious clues about the general behaviour of pollutants, all the more as salt is known to affect the solubility limit of hydrocarbons and, consequently, their coadsorption.
CONCLUSION

Despite its unavoidable experimental limits, this study clearly shows how the adsorption of two cyclic, aliphatic or aromatic hydrocarbons with the same number of carbons is affected by:

– the molecular structure;
– the solubility limit;
– the reciprocal wetting between hydrocarbon-rich phase and solvent-rich phase at the solubility limit. It gives a new insight into adsorption process by evidencing the induction of heavy metal salt coadsorption by hydrocarbon adsorption. It showed that the extent of coadsorption depends on:
– the nature and solubility limit of the hydrocarbons under study;
– hydrocarbon adsorption;
– the ratio between hydrocarbon and salt concentrations.

However, the main conclusion is that hydrocarbon adsorption is responsible for coadsorption, since no absorption of salt took place when hydrocarbon was missing. Metal could be also adsorbed as a complexed species, but we showed that it formed no complex in solution under our experimental conditions.

A part of these data was produced with the bubble column, a method scarcely used, which proved its relevance and usefulness in the study of adsorption process with surface tension-inactive systems.

Finally, the coadsorption mechanism described here may explain the passing of scarcely soluble, non adsorbable compounds from fresh water to the atmosphere. It is likely that such investigations should be relevant in the case of salted waters.

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