The Influence of Sulfate Ion on the Coal-Wetting Performance of Anionic Surfactants

By Howard W. Kilau
Mission: As the Nation’s principal conservation agency, the Department of the Interior has responsibility for most of our nationally-owned public lands and natural and cultural resources. This includes fostering wise use of our land and water resources, protecting our fish and wildlife, preserving the environmental and cultural values of our national parks and historical places, and providing for the enjoyment of life through outdoor recreation. The Department assesses our energy and mineral resources and works to assure that their development is in the best interests of all our people. The Department also promotes the goals of the Take Pride in America campaign by encouraging stewardship and citizen responsibility for the public lands and promoting citizen participation in their care. The Department also has a major responsibility for American Indian reservation communities and for people who live in Island Territories under U.S. Administration.
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UNIT OF MEASURE ABBREVIATIONS USED IN THIS REPORT

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Unit Description</th>
<th>Abbreviation</th>
<th>Unit Description</th>
</tr>
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<tr>
<td>°C</td>
<td>degree Celsius</td>
<td>mmol/L</td>
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<td>dyn/cm</td>
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<td>minute</td>
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THE INFLUENCE OF SULFATE ION ON THE COAL-WETTING PERFORMANCE OF ANIONIC SURFACTANTS

By Howard W. Kilau

ABSTRACT

The U.S. Bureau of Mines is investigating surfactants added to water sprays to enhance the control of dust during coal mining operations. The objective of the present work was to establish the general applicability of adding sulfate ion to anionic surfactant solutions in order to improve wetting action. This was pursued through laboratory wettability testing of four anionic surfactants and one nonionic surfactant in combination with various concentrations of sulfate ion.

The experimental results demonstrated that the addition of sulfate ion enhanced the wetting characteristics of all the anionic surfactants when applied to hard-to-wet coals, while easy-to-wet coal showed more complex wetting behavior. Sulfate ion in the presence of nonionic surfactant enhanced wetting only slightly or not at all.

Surface tension measurements of the dilute aqueous surfactant-sulfate anion solutions indicated this property was not important in coal-wetting phenomena, as long as the value was below the critical value for coal. Sulfate ion helps surfactant reduce surface tension but primarily appears to affect wetting action by altering the adsorption characteristics of surfactant on the coal surface.

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INTRODUCTION

Coal dust arising from mining creates health and safety problems. Pneumoconiosis (black lung disease) is a serious health problem frequently afflicting coal miners exposed to respirable coal dust for long periods. Also, coal dust suspended in the underground mine atmosphere constitutes an explosion and fire hazard. In the case of methane ignition, suspended coal dust can help propagate fire rapidly throughout a mine to entrap miners before they have a chance to escape. Since 1969, the Coal Mine Health and Safety Act has limited coal mine atmospheres to a maximum of 2.0 mg/m² of respirable coal dust, or lower if the dust contains >5 µg silica. Mine operators use various methods to control dust levels, including water sprays directed at the face and bit during coal cutting. However, for many mining operations, particularly longwall sections, water spray and other conventional methods are inadequate to reduce levels below the standard threshold limit (1).²

Surfactant addition to the water used in sprays has been suggested and used in some mines for improving the effectiveness of water sprays. A surfactant added to water generally helps the wetting of coal since the coal surface is normally very hydrophobic in nature. Because of the improved wetting action with surfactant, one would predict a favorable effect on coal dust suppression. However, the mine experience using surfactants has not been consistently positive and has been frequently below expectations. Even laboratory testing of surfactant wetting agents has shown numerous inconsistencies among various surfactant products applied to different coals and a lack of correlation with field testing. Accordingly, the U.S. Bureau of Mines is investigating the fundamentals of coal wetting in the belief that a basic understanding of the wetting mechanism is needed to properly evaluate the dust suppression effectiveness of surfactants and to achieve the potential that these reagents appear to offer for reducing coal dust levels in the underground mining environment. This work is in support of the Bureau's mission to maintain a healthful and safe working environment in the mines by developing technology that enables industry to comply with current regulations in a cost-effective manner.

In earlier Bureau work (2), the coal-wetting ability of sodium di (2-ethylhexyl) sulfosuccinate anionic surfactant (SD2ES) was observed to be greatly enhanced through the addition of sodium and potassium salts of multivalent anions to the surfactant solution. These factors include the surface tension of the liquid solution to fill with coal particles when deposited on a planar surface of the particles (minus 200 mesh). There are several factors that can influence the wetting action of surfactant solutions applied to the surface of coals. These factors include the surface tension of the liquid solution, the adsorption of surfactants and ions from the solution onto the coal surface, the exchange of surfactant ions with coal surface ions, the packing of surfactant molecules on the surface, and the adsorption of surfactant on previously adsorbed surfactant to form agglomerates called hemimicelles. A theoretical description of these processes is provided in the following section.

²Italic numbers in parentheses refer to items in the list of references at the end of this report.
Figure 1.-Effect of multivalent anion addition on wetting performance of SD2ES surfactant. Drop Penetration test on hard-to-wet coal using 0.14-wt-pct SD2ES solution. Error bars shown represent 95-pct confidence level.
CRITICAL SURFACE TENSION

Pure water is normally ineffective in wetting coal because of its high surface tension of 72.8 dyn/cm. A considerable reduction in this surface tension is required for successful coal wetting. A critical surface tension was determined by Parekh and Aplan to be about 45 dyn/cm for all coals, independent of rank. Once this critical value is attained, it is believed that other factors, particularly surfactant adsorption, begin to be important and dominate the wetting action.

All the initial surfactant concentrations used in this work were adequate to provide sufficient reduced surface tension for wetting of coal. However, it is possible that for the smallest initial concentrations, adsorption of surfactant on the coal may consume the reagent in the droplet to a point where the concentration is borderline for maintenance of adequate reduced surface tension in the final stages of coal wetting in the Drop Penetration test.

SURFACTANT ADSORPTION

In addition to reducing surface tension below the critical value, the object of adding surfactants is to convert the hydrophobic areas of the coal surface to a hydrophilic state by the adsorption of the surfactant exclusively on hydrophobic sites. If this condition is accomplished, then improved wettability of the coal surface will result (if the liquid surface tension is below the critical value). The coal surface is conceived as composed of both hydrophobic and hydrophilic sites. The adsorption of surfactants on these different kinds of sites complicates the wetting behavior of coal in surfactant solutions. The situation is depicted schematically in figure 2, where the hydrophobic coal surface is shown as the shaded area and hydrophilic sites as small squares enclosing a negative charge (box A). The negatively charged hydrophilic sites stem from oxidation of organic surface groups and the presence of inorganic oxide impurities.

It is suggested that inherent moisture in the coal provides a medium for solubilization of multivalent positive ions from the coal. These cations are attracted to the negative sites on the coal surface to form a layer of positive ions on hydrophilic sites as illustrated (box B, figure 2). The moisture content of the coal, by providing a medium for diffusion through the coal, may determine the extent of the distribution of the positive ions on the sites. Also, development of additional negative sites through oxidation upon exposure to air and distribution of positive ions through an adsorbed moisture layer after exposure may occur.

ION EXCHANGE

The hydrophilic head of the surfactant molecule is also capable of being attracted to the positive layer of ions on the coal's natural hydrophilic sites. In the case of anionic surfactants, this attraction is likely to be especially strong because of the electrostatic forces between the negatively charged head of the surfactant ion and the positively charged ion layer on the coal surface. This attraction results in the displacement of monovalent negative ions previously attached to the positive ion layer in an ion-exchange type of reaction.

This ion-exchange reaction, where negative surfactant anions replace other attached anions on the positive ion layer, converts the hydrophilic sites to an undesirable nonwetting state because of the orientation of the attached surfactant with its hydrophobic tail directed toward the aqueous phase (box D, figure 2). The addition of Na₂SO₄ provides divalent sulfate anions that compete with monovalent surfactant anions for attachment to the layer of positive ions on the coal surface. Because of their higher valence charge compared with that of surfactant anions, sulfate anions will tend to be preferentially adsorbed on the positive ion layer, displacing surfactant anions following the affinity hierarchy of ion-exchange reactions. This mechanism prevents conversion of these hydrophilic sites to a hydrophobic state with the attendant decreased wettability (figure 2, box E).

\(^4\) The critical surface tension was first proposed and measured by Zisman (5) and is defined as the surface tension of a liquid required for spontaneous spreading of the liquid over a given solid surface to give a contact angle of zero.

\(^5\) The selectivity order of several common anions on anion-exchange resins has been shown (5) to be as follows:

\[\text{SO}_4^{2-} > \text{CrO}_4^{2-} > \text{citrate} > \text{tartrate} > \text{NO}_3^- > \text{AsO}_4^{3-} > \text{PO}_4^{3-} > \text{MoO}_4^{2-} > \text{acetate}, \Gamma > \text{Br} > \text{Cl} > \text{F}^-\]

Usually, higher valence species displace monovalent species.
Figure 2.—Schematic of wetting mechanism involving adsorption of anionic surfactant and sulfate ion onto hydrophobic and hydrophilic coal surface.
The affinity hierarchy may not prevail if an excess of monovalent surfactant ion species over divalent sulfate species is present in solution. In such a case, the protection of hydrophilic coal sites afforded by sulfate anion would be diminished. An analogy from water treatment technology is the process of regenerating an ion-exchange resin by using high concentrations of Na\(^+\) ions (from NaCl brine) to remove attached higher valence Ca\(^{2+}\) and Mg\(^{2+}\) ions from the resin. It is to be noted that nonionic surfactants would be unaffected by ion-exchange effects. Thus, nonionic surfactants, if adsorbed on the positive ion layer of the coal, would not necessarily be removed or affected by addition of Na\(_2\)SO\(_4\). Also, electrostatic repulsive forces would be ineffective to prevent adsorption of nonionic surfactant on top of previously attached sulfate ions.

CLOSE-PACKING EFFECTS

The addition of Na\(_2\)SO\(_4\) electrolyte provides numerous ions (Na\(^+\) and SO\(_{4}^{2-}\)), which can reduce the distance over which electrostatic forces in the double layer are effective; i.e., the Debye length is shortened. In effect, the "cloud" of ions from the electrolyte can screen or reduce the repulsive electrostatic forces between the negatively charged heads of the anionic surfactant adsorbed on the hydrophobic coal surfaces. As a result, greater density of the adsorbed surfactant ions is permitted, which tends to convert more of the coal's hydrophobic surface to a hydrophilic condition and increase wettability. This situation is depicted in an approximate manner in figure 3. Part A of the figure shows the low-density adsorption of surfactant ions on the coal surface occurring in the absence of Na\(_2\)SO\(_4\). Part B shows the increased density of surfactant adsorption occurring in the presence of Na\(_2\)SO\(_4\), caused by the reduction of repulsive forces between the negatively charged heads of the adsorbed surfactant ions.

Similar improved close packing of anionic surfactant molecules through Na\(_2\)SO\(_4\) addition can also occur at the air-liquid interface in aqueous solutions of surfactant. This phenomenon reduces the liquid surface tension of anionic surfactant solutions with increasing concentrations of Na\(_2\)SO\(_4\).

HEMIMICELLE FORMATION

An additional factor in the adsorption of surfactant molecules is the possibility of hemimicelle formation on the coal surface. Adsorption of anionic surfactant on positively charged surfaces is believed to occur in three distinct modes as surfactant concentration is increased. At low concentration, surfactant adsorbs mainly by ion exchange. At intermediate concentrations, a marked increase in adsorption occurs resulting from interaction of the hydrophobic chains of oncoming surfactant anions with those of previously adsorbed surfactant and with themselves. This aggregation of the hydrophobic groups of the surfactant has been called hemimicelle formation. An example of hemimicelle formation on a cluster of hydrophilic sites on the coal surface is depicted schematically in figure 4. A cluster of surfactant ions is shown attached to a group of adsorbed dipositive cations on the coal surface. The hydrophobic tails of these surfactant molecules extending into the solution are capable of attracting further surfactant anions to form the aggregation depicted in the figure. The effect of this kind of hemimicelle formation is to restore wettability to the coal surface lost by the first layer adsorption of surfactant in adverse orientation. The final mode of adsorption, labeled the "electrostatically hindered mode", is weak because further surfactant adsorption must overcome electrostatic repulsion between oncoming surfactant ions and similarly charged hemimicelles on the coal surface.

MATERIALS

COALS

Chemical analyses of the hard-to-wet and easy-to-wet coals tested are given in table 1. The apparent ranks of the coal samples were based on ASTM D388-84 classification system. The two samples were received in the form of 3-in lumps and were ground to minus 200 mesh for the Drop Penetration test. The grinding was done in air, but care was taken not to overgrind the samples so as to lessen oxidation of the coal surfaces. Laser light-scattering measurements of the ground particles showed 74 pet of the hard-to-wet coal and 69 pet of the easy-to-wet coal to be in the size range of 11 to 62 μm. The two coals chosen were selected on the basis of their extreme wetting properties, the one being difficult to wet and the other wetting easily at low concentrations of surfactant in the Drop Penetration test.
Figure 3.- Schematic of creation of surfactant ion close packing on coal surface.

A, Loose packing of adsorbed anionic surfactant, no $\text{Na}_2\text{SO}_4$ present.  
B, Close packing of adsorbed anionic surfactant aided by presence of $\text{Na}_2\text{SO}_4$. 

Coal Solution

\[ \text{Na}^+ \text{ Na}^+ \]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]

\[ \text{Na}^+ \text{ Na}^+ \\]

\[ \text{SO}_4^{2-} \text{ Na}^+ \]
Figure 4.—Schematic of hemimicelle formation on coal surface.
Table 1.—Analyses of coal samples used in wettability experiments, weight percent (as received)

<table>
<thead>
<tr>
<th></th>
<th>Hard-to-wet,</th>
<th>Easy-to-wet,</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>moisture, volatile matter</td>
<td>moisture, volatile matter</td>
</tr>
<tr>
<td>Moisture</td>
<td>0.3</td>
<td>2.2</td>
</tr>
<tr>
<td>Volatile matter</td>
<td>21.7</td>
<td>34.3</td>
</tr>
<tr>
<td>Total C</td>
<td>78.9</td>
<td>58.5</td>
</tr>
<tr>
<td>Fixed C</td>
<td>67.3</td>
<td>44.8</td>
</tr>
<tr>
<td>Ash</td>
<td>10.7</td>
<td>18.7</td>
</tr>
<tr>
<td>S</td>
<td>.7</td>
<td>3.9</td>
</tr>
<tr>
<td>O</td>
<td>3.4</td>
<td>11.3</td>
</tr>
<tr>
<td>H</td>
<td>4.5</td>
<td>4.2</td>
</tr>
<tr>
<td>N</td>
<td>1.4</td>
<td>1.2</td>
</tr>
<tr>
<td>Al</td>
<td>1.8</td>
<td>2.0</td>
</tr>
<tr>
<td>Si</td>
<td>2.4</td>
<td>3.5</td>
</tr>
<tr>
<td>Fe</td>
<td>1.1</td>
<td>.9</td>
</tr>
<tr>
<td>Ca</td>
<td>.2</td>
<td>.4</td>
</tr>
<tr>
<td>Mg</td>
<td>.1</td>
<td>.1</td>
</tr>
<tr>
<td>Na</td>
<td>.&lt;1</td>
<td>.1</td>
</tr>
</tbody>
</table>

1Jim Walter Corp., Blue Creek #5 Mine, Alabama, Blue Creek Seam.
3By subtraction [100 - (pet moisture, ash, volatile matter)].
4By subtraction [100 - (pet C, H, S, N, ash, moisture)].

NOTE.—Coal ranks were determined by ASTM D388-84 (8).

WETTING SOLUTION REAGENTS

Four of the surfactants tested were anionic; i.e., they formed negative anions when dissolved in water. These surfactants were compared with a nonionic surfactant with and without electrolyte (Na2SO4) addition. Table 2 lists the surfactants and electrolyte reagent used in the wettability testing. The structural formulas for the surfactants are given in figure 5.

The wetting solutions were prepared by diluting the surfactant reagents to the desired concentration with pure water to obtain a 3-L quantity of solution. This solution was then divided into 500-mL aliquots in volumetric flasks. Na2SO4 was added to the aliquots to achieve the desired concentrations of added SO4^2- ion (usually 2.5, 5.0, 15.0, and 30.0 mmol/L). It is to be noted that all the surfactant concentrations given for the wetting solutions in this report are expressed in terms of active surfactant ingredient. The pure water used to prepare the surfactant wetting solutions was first distilled and then passed through a deionizer cartridge.

Table 2.—Reagents used in wettability experiments

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Abbreviation</th>
<th>Commercial name</th>
<th>Manufacturer</th>
<th>Concentration of active ingredient, wt pet (as received)</th>
<th>Comments</th>
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<tbody>
<tr>
<td>Sodium di (2-ethylhexyl) sulfosuccinate.</td>
<td>SD2ES</td>
<td>Aerosol OT-75</td>
<td>American Cyanamid.</td>
<td>75</td>
<td>Anionic, double hydrocarbon chain, in alcohol-water solution.</td>
</tr>
<tr>
<td>Sodium di hexyl sulfosuccinate.</td>
<td>SDHS</td>
<td>Aerosol MA-80</td>
<td>. . do</td>
<td>80</td>
<td>Do.</td>
</tr>
<tr>
<td>Sodium dodecyl sulfate.</td>
<td>SDS</td>
<td>Sodium dodecyl sulfate.</td>
<td>BDH Chemicals, Poole, England</td>
<td>99</td>
<td>Anionic, single hydrocarbon chain, solid powder.</td>
</tr>
<tr>
<td>Sodium dodecyl benzene sulfonate.</td>
<td>SDBS</td>
<td>Dodecyl benzene sulfonic acid sodium salt.</td>
<td>. . do</td>
<td>80</td>
<td>Anionic, single hydrocarbon chain, solid powder, 20 wt pet Na2SO4 impurity.</td>
</tr>
<tr>
<td>Ethoxylated acetylenic glycol (Ethoxylated 2,4,7,9-tetramethyl-5-decyne-4,7-diol).</td>
<td>EAG</td>
<td>Surlyn 465</td>
<td>Air Products and Chemical Corp.</td>
<td>100</td>
<td>Nonionic, single hydrocarbon chain, liquid, 65 wt pet ethylene oxide adduct.</td>
</tr>
<tr>
<td>Sodium sulfate</td>
<td>Na2SO4</td>
<td>Anhydrous sodium sulfate.</td>
<td>Baker</td>
<td>99</td>
<td>Analyzed reagent grade, solid powder.</td>
</tr>
</tbody>
</table>
Figure 5.—Structural formulas for surfactants used. A, Sodium di (2-ethylhexyl) sulfosuccinate (SD2ES). B, Sodium di hexyl sulfosuccinate (SDHS). C, Sodium dodecyl sulfate (SDS). D, Sodium dodecyl benzene sulfonate (SDBS). E, Ethoxylated acetylenic glycol (EAG) or 2,4,7,9-tetramethyl-5-decyn-4,7-diol with 10 mol ethylene oxide adduct.
EXPERIMENTAL PROCEDURE

The extent of wetting achieved with the various wetting solutions, when applied to the two wettability classes of coal, was measured with the Drop Penetration test. This test was described in earlier work (2). In this test, a 40-μL droplet of the surfactant solution to be tested is deposited (from a micropipette) onto the planar surface of a bed of minus 200-mesh coal particles. The droplet is observed through a microscope (×20), and the time for the droplet to completely fill with coal particles is recorded with a stopwatch. The coal bed was 8 mm in depth, and its top surface was smoothed and leveled with a straightedge, avoiding any packing. Determining the point of complete filling of the droplet was difficult because some wetting solutions formed thin liquid films near the endpoint, which remained long after the droplet was apparently filled with particles. It was observed that as the particles filled the droplet, a stage was reached in which only a small amount of liquid was visible, flowing in small rivulets on the surface of the filled droplet. Consistency in determining the endpoint was attained by designating complete particle filling as the point at which no coal particle movements could be observed in any of the rivulets (believed to correspond to the cessation of particle penetration into the droplet, resulting in the stopping of fluid flow in the rivulets). Generally, the Drop Penetration test was repeated 10 times for a wetting solution on a coal sample, and the average of the trials was computed. The standard deviation in the computed average time for complete particle filling was typically 10 to 15 pct. No attempt was made to test very dilute solutions of wetting agents where wetting times exceeded 12 min because of the experimental inconvenience and because evaporation of the liquid droplet might become a factor.

Surface tensions of all wetting solutions at 28 °C were also measured with a du Nouy ring-type instrument. The particle size distributions of the ground coal samples were measured with a Microtrac Particle-Size Monitor, model 7981, Leeds and Northrup Co.6

EXPERIMENTAL RESULTS

PURE WATER

No wetting action was observed in the Drop Penetration test on either coal sample by pure water. This expected result is a consequence of the high surface tension of water, which must be lowered below the critical value before coal wetting can proceed.

SD2ES SURFACTANT ON HARD-TO-WET COAL

For SD2ES surfactant solutions without sulfate additive, applied to the hard-to-wet coal sample, the wettability increased as the surfactant concentration was increased. A wetting time limit was reached at about 10 s for 0.61 wt pct SD2ES concentration (fig. 6), at zero concentration of added Na2SO4. Increasing the SD2ES concentration to 0.82 wt pct did not significantly lower the wetting time at zero concentration of added Na2SO4. In general, it was found in this and other work (2) that 7 to 10 s wetting time is a minimum value in the Drop Penetration test (when employed as described) regardless of the wetting agent or coal employed.

Certain characteristics visible in figure 6 reveal the dominant influence of adsorption phenomena on the wetting action at the coal-liquid interface compared with the influence of the liquid's bulk properties. Figure 6 also shows remarkable improvement in wetting time with only a slight increase in SD2ES concentration from 0.14 to 0.20 wt pct at zero concentration of added Na2SO4. This wetting response suggests that adsorption of surfactant on hydrophilic coal sites is a major factor contributing to the wetting behavior. This is inferred from the almost negligible difference in liquid surface tensions of these two wetting solutions. The small difference in the two measured values (28.4 dyn/cm for 0.14 wt pct SD2ES and 28.0 dyn/cm for 0.20 wt pct SD2ES) indicates a lack of surfactant molecule close-packing change at the air-liquid interface, which probably reflects a similar situation at the coal-liquid interface. Further improvements in wettability occurred with further increased SD2ES concentration but with increasingly less effect, up to the wettability limit at 0.61 wt pct SD2ES (curves C and D).

6Reference to specific products does not imply endorsement by the U.S. Bureau of Mines.
Figure 6.—Effect of Na$_2$SO$_4$ on wetting performance of SD2ES surfactant applied to hard-to-wet coal. Error bars shown represent 95-pct confidence level.
Hard-to-wet coal is conceived as having a preponderance of hydrophobic surfaces and a smaller fraction of hydrophilic sites, compared with easy-to-wet coal. It is suggested that adsorption of surfactant (in the absence of Na₂SO₄) occurs simultaneously on hydrophobic and hydrophilic surfaces at low surfactant concentrations. For low concentrations, adsorption of surfactant on hydrophilic coal sites would have a significant negative effect on wettability since not only are hydrophilic coal sites converted to a hydrophobic state, but scarce surfactant anions are consumed in the process, leaving less reagent available for desirable adsorption on hydrophilic sites to improve the total coal wettability (figure 2, box D). Although simultaneous adsorption occurs, the attraction of surfactant to hydrophilic sites is believed somewhat greater than to hydrophilic sites. This results in a net wetting improvement as surfactant concentration is increased, as seen in figure 6. For hard-to-wet coal, a point is reached at which adsorption on all available hydrophilic sites has reached saturation, so that any further adsorption occurs solely on the remaining hydrophobic sites, which constitute the majority of available adsorption sites for hard-to-wet coal.

It is suggested this saturation point in the adsorption of surfactant onto hydrophilic coal sites is reached between 0.14 and 0.20 wt pet SD2ES. The great change in wetting time between these concentrations is believed due to the transition from competing, simultaneous adsorption on the two kinds of sites (with its canceling effect on wettability) to a stage of exclusive, noncompetitive adsorption on hydrophobic coal sites. When the wetting solution is at or below 0.14 wt pet SD2ES, wetting improvement by van der Waals' adsorption on hydrophobic surface is offset partially by concurrent adsorption of surfactant ions on hydrophilic sites (by electrostatic forces) in opposite orientation to produce hydrophobicity as shown in figure 2 (box D).

At concentrations of 0.20 to 0.61 wt pet SD2ES, adsorption can occur exclusively on hydrophilic sites in the later stages of wetting, resulting in greatly reduced total wetting time. At higher concentrations of surfactant, the wetting improvement change as a function of surfactant concentration is ultimately limited. This may be because a monolayer of adsorbed surfactant has been achieved or because kinetics factors (such as diffusion of particles across the coal-liquid interface) control the adsorption at faster wetting times.

There is the possibility of hemimicelle formation on the coal surface in the later stages of surfactant adsorption, which could also contribute to the sharp drop in wetting time between curves A and B (fig. 6). In this case, after saturation of hydrophilic sites with surfactant, hemimicelle formation occurs atop the surfactant layer to restore hydrophilicity, as in figure 4. This occurrence, together with the attendant conversion of hydrophobic coal surface to a wettable state from van der Waals' adsorption, could further contribute to the enhanced wetting improvement observed between 0.14 and 0.20 wt pet SD2ES surfactant solutions in figure 6.

As shown in figure 2 (box E), addition of Na₂SO₄ protects hydrophilic sites from conversion to hydrophobicity by preventing surfactant adsorption in adverse orientation. Figure 6 shows that the greatest improvement from addition of Na₂SO₄ occurred for a low concentration of surfactant (curve A). At this SD2ES concentration (which is just below the transition concentration where hydrophilic coal sites are believed to become nearly or completely covered with adsorbed surfactant), the Na₂SO₄ addition acts to preserve hydrophilic sites and reduces the number of sites available for surfactant adsorption in adverse orientation (2). In effect then, Na₂SO₄ lowers the transition concentration for SD2ES, so that the wetting improvement through Na₂SO₄ additions is nearly as beneficial as that when SD2ES concentration alone was increased (curve A). This occurs because scarce surfactant anions are not only prevented from adsorbing improperly (eliminating hydrophilic surface), but surfactant also then becomes available for adsorption on hydrophobic sites and possibly for hemimicelle formation on any remaining unprotected hydrophilic sites. Fast wetting times can be achieved with high concentrations of surfactant alone as well as with lower concentrations of surfactant in combination with Na₂SO₄ reagent. Practically speaking, however, it is preferable from an economic standpoint to add inexpensive Na₂SO₄ reagent to low concentrations of more costly surfactant to improve wetting rather than to increase surfactant concentration.

It is not certain what caused the lack of improved wetting response when Na₂SO₄ was added to surfactant solutions of higher concentration (figure 6, curves B, C, and D). It may be that the advantageous valence-controlled affinity hierarchy for ion-exchange adsorption of sulfate ion (on the hydrophilic positive layer of the coal) becomes inoperative at these higher concentrations of surfactant. When valence no longer controls adsorption, surfactant ions, because of their high concentration in solution, remain firmly attached to hydrophilic sites on the coal despite the presence of sulfate anion. Alternatively, if hemimicelle formation has taken place on hydrophilic coal surface at higher surfactant concentrations to restore hydrophilicity, the addition of Na₂SO₄ is seen to be superfluous and consequently ineffective.
SD2ES SURFACTANT ON EASY-TO-WET COAL

The wettability behavior of SD2ES surfactant and Na$_2$SO$_4$ additive on easy-to-wet coal was different in several respects from their behavior on hard-to-wet coal. Figure 7 displays the wetting times measured on easy-to-wet coal as a function of SD2ES and added Na$_2$SO$_4$ concentrations. As with hard-to-wet coal, increasing the surfactant concentration improved the wetting until a limit was reached (curve D). However, unlike with hard-to-wet coal, further increase of SD2ES concentration did not maintain wetting at the limiting value but instead decreased the wetting, the excess SD2ES concentration apparently harming the wetting action above the limiting concentration (curves E and F).

To understand this wetting behavior for easy-to-wet coal, it is suggested that the coal surface again be envisioned as possessing both hydrophobic and hydrophilic surface sites, but in this case the hydrophilic sites outnumber the hydrophobic sites. For surfactant in the absence of Na$_2$SO$_4$ addition, it is likely that the smaller number of hydrophobic coal sites are almost covered by adsorbed surfactant long before the hydrophilic sites, as surfactant concentration is increased. If sufficient surfactant is available, adsorption proceeds exclusively on hydrophilic sites after hydrophobic surface has reached adsorption saturation. However, this final stage of adsorption begins to create increased hydrophobic character on the coal surface because of adsorption of surfactant on the hydrophilic sites in adverse orientation (figure 2, box D). The point at which the net adsorption on the easy-to-wet coal sample (hydrophobic minus hydrophilic adsorption) begins to favor attachment to hydrophilic surfaces to create hydrophobicity is seen to occur in figure 7 at a concentration of about 0.41 wt pct SD2ES. This transition point is marked by a substantial decrease in wettability (longer wetting time, curve E at zero Na$_2$SO$_4$ concentration). Further increase in SD2ES concentration only results in further deterioration of wettability because of further adverse adsorption on hydrophilic sites to create hydrophobicity (curve F).

The possibility of hemimicelle formation was previously mentioned for hard-to-wet coal, but there appears to be no evidence of such phenomena for easy-to-wet coal in figure 7. The absence of restoration of wettability at high concentrations of surfactant (curve F) leads to the conclusion that hemimicelles are not formed on easy-to-wet coal. Furthermore, one may argue that since hemimicelle formation is a property related to surfactant characteristics, the wetting phenomena observed for hard-to-wet coal that was previously attributed to possible hemimicelle formation is due to other causes. However, considerably more surfactant molecules are required to form hemimicelles on easy-to-wet coal because of the greater hydrophilic surface area and, therefore, greater numbers of adsorbed surfactant molecules in adverse orientation. Hence, greater

Figure 7.—Effect of Na$_2$SO$_4$ on wetting performance of SD2ES surfactant applied to easy-to-wet coal.
surfcatant concentration is required to form hemimicelles on easy-to-wet coal compared with hard-to-wet coal, to offset the hydroporphic effect of adversely adsorbed surfac-
tant. The hemimicelle effect may not be noticeable until concentrations of >0.82 wt pet SD2ES are attained. How-
ever, concentrations of >0.82 wt pet SD2ES are not prac-
tical because of solubility limitations.

In figure 7 (curves A, B, C, and D), the addition of 
Na2SO4 generally improved the wettability, but not remark-
ably except for SD2ES concentrations above the transition 
concentration for maximum wettability (curves E and F). 
It is suggested the subdued wetting response at lower 
surfactant concentration results because much of the sul-
fate anion adsorbs onto free hydrophilic sites without 
displacing adsorbed surfactant anions, since surfactant 
coverage of the numerous hydrophilic sites on easy-to-wet 
coal is not likely to be extensive at lower concentrations of 
surfactant. At higher surfactant concentration, above the 
transition concentration, the addition of Na2SO4 is seen to 
be much more effective in improving wetting, presumably 
because the sulfate anion has a greater probability of dis-
placing adsorbed surfactant ions, since the population den-
sity of adsorbed surfactant on the hydrophilic surface has 
probably become substantial at that point. Low concen-
trations of Na2SO4 (<5.0 mmol/L) appeared ineffective 
above the surfactant transition concentration (figure 7, 
curves E and F). This region may indicate there is insuf-
ficient sulfate concentration to adequately displace ad-
sorbed surfactant from the hydrophilic surface to give 
significantly visible wetting effects.

SURFACE TENSION OF SD2ES SOLUTIONS

Comparison of the surface tension of the surfactant 
solutions (fig. 8) with their wetting abilities on hard-to-wet 
coal (fig. 6) reveals few similarities other than as surfactant 
and Na2SO4 concentrations are increased, wettability tends 
to increase and surface tension to decrease. However, the 
large difference in wetting performance between 0.14 and 
0.20 wt pet SD2ES is not reflected by a similar large differ-
ence in surface tension between these two concentrations 
(figure 8, curves A and B). Also, figure 8 shows a con-
sistent and similar decrease in surface tension as Na2SO4 
is added to 0.14, 0.20, and 0.41 wt pet SD2ES surfactant 
solutions, while the wettability improvement from Na2SO4 
addition was less effective from dilute to higher concen-
trations of surfactant (figure 6, curves A, B, and C). For 
easy-to-wet coal, the impairment of wettability at higher 
concentrations of surfactant (fig. 7) is not predicted by 
figure 8, where surface tension declines gradually or 
remains constant as surfactant concentration is increased.

OTHER ANIONIC SURFAC TANT S

Figure 9 demonstrates that anionic SDS surfactant was 
effective in wetting hard-to-wet coal, the wetting response 
increasing as the concentration of the surfactant was in-
creased (curves A and B at zero concentration of added 
Na2SO4). As with SD2ES, a small increase in surfactant 
concentration (0.74 to 0.99 wt pet) resulted in a remark-
able decrease in wetting time. Again, the addition of 
Na2SO4 improved the wettability most effectively at the 
lower concentration of surfactant for hard-to-wet coal 
(figure 9, A compared with curve B).

Figure 10 demonstrates the effect of Na2SO4 addition on 
the Drop Penetration wettability of SDBS surfactant. This 
surfactant showed wetting characteristics similar to those 
of SD2ES and SDS on hard-to-wet coals. Greatly im-
proved wetting resulted from small increases in surfactant 
concentration, similar to the results for SD2ES and SDS 
(curves A and B). Wetting was improved with Na2SO4 
addition at low concentrations of surfactant (curve A), but 
when surfactant concentration was increased, the effect of 
Na2SO4 addition was small (curve B).

Figure 11 presents the experimental wetting results for 
anionic surfactant SDHS, curves B and B1 (the figure also 
includes the nonionic surfactant EAG, curves A and A1, 
which will be discussed in the next section). The anionic 
surfactant SDHS demonstrated improved ability to wet 
hard-to-wet coal as Na2SO4 was added (curve B), a result 
similar to that for the other anionic surfactants tested in 
the slow-wetting range (i.e., for surfactant concentrations 
yielding 300- to 600-s wetting times before Na2SO4 is 
added).

Thus, two main characteristics of wetting behavior are 
visible for hard-to-wet coal treated with anionic surfactants 
and Na2SO4:

1. Greatly improved wetting as Na2SO4 is added to 
surfactant solutions in the slow-wetting range (curve A of 
figures 6, 9-10, and curve B of figure 11); and
2. Diminished effect on wetting as Na2SO4 is added to 
surfactant solutions in the fast-wetting range, i.e., those 
solutions that yield, in the absence of Na2SO4, wetting 
times in the range 10 to 120 s (curves B, C, D, and E, of 
figure 6, and curve B of figures 9-10).

A plot of the surface tension of SDS and SDBS at 
various concentrations of surfactant and Na2SO4 additive 
(fig. 12) shows little difference in surface tension between 
surfactant concentrations, while wetting times can vary 
widely with concentration (figures 9-10). For example, 
there is little surface tension difference between curve A 
and curve A1 of figure 12, while there is a great disparity in 
wetting times on hard-to-wet coal between these same 
two concentrations of SDS (figure 9, curves A and B). 
Similarly, although there is little difference between the 
surface tension curves for various concentrations of SDBS 
in figure 12 (curves B, B1, and B2), the wetting times for 
hard-to-wet coal were greatly different for 0.44 and 0.60 wt 
pct SDBS (fig. 10).

The effect of added Na2SO4 on SDS, SDBS, and SDHS 
for easy-to-wet coals (figure 9, curve C; figure 10, curve 
C; figure 11, curve B) resembled to some degree the small 
wetting response displayed by SD2ES on easy-to-wet coal 
when SD2ES concentration was below the transition con-
centration (figure 7, curves A, B, C, and D). These anionic 
surfactants displayed only slight wetting improvement with
Figure 8. Surface tension at 28°C for wetting solutions containing SD2ES plus Na₂SO₄.
Figure 9.—Effect of Na₂SO₄ on wetting performance of SDS surfactant.
Figure 10—Effect of Na$_2$SO$_4$ on wetting performance of SDBS surfactant.
Figure 11.—Effect of $\text{Na}_2\text{SO}_4$ on wetting performance of SDHS and EAG surfactants.
Figure 12.-Surface tension at 28° C for wetting solutions containing SDS and SDBS plus Na₂SO₄.
increased Na$_2$SO$_4$ concentration, a result in harmony with their evidently weaker adsorbing properties on coal, compared with SD$_2$ES. Thus, it is suggested that there is a smaller number of these weaker absorbing surfactant anions to be displaced from hydrophilic sites by sulfate anion, and once these anions are displaced, adsorption onto hydrophobic sites to produce more wettability proceeds less effectively.

**NONIONIC SURFACTANT**

For EAG nonionic surfactant applied to hard-to-wet coal (figure 11, curve $A$), the wetting characteristics are quite different from those of the anionic surfactant. The effect of adding Na$_2$SO$_4$ to this surfactant was minor, with only a small improvement in wetting recorded up to 15 mmol/L Na$_2$SO$_4$ added, and no further improvement at greater concentrations of Na$_2$SO$_4$. For easy-to-wet coal, improved wetting response was completely absent in the presence of Na$_2$SO$_4$ (figure 11, curve $A_1$). This result is expected in light of the proposed wetting mechanisms, since these uncharged surfactant species should not be particularly affected by the presence of ionic species in solution.

Comparisons of the wetting response for SDHS and EAG surfactant in figure 11 with the surface tension results presented in figure 13 reveal poor correlation between surface tension and coal wettability by these surfactants. It is evident that the class of surfactant (anionic or nonionic) employed in combination with Na$_2$SO$_4$ is much more important than the surface tension reduction afforded by the wetting agent. For example, in figure 13, the surface tensions are almost identical for 0.42-wt pct EAG and 0.77-wt pct SDHS surfactant solutions containing 22 mmol/L of added Na$_2$SO$_4$, while wetting times of the same two solutions differed immensely (by 7 min, curves $A$ and $B$ in fig. 11).

![Figure 13](image_url)
DISCUSSION

COMPARISON OF SURFACTANT WETTING PERFORMANCE

Figure 14 compares wetting curves obtained for the five surfactants on hard-to-wet coal in the slow-wetting range. It is evident that SD2ES is the most effective wetting agent (per unit weight of active ingredient) of all the surfactants tested. SD2ES was observed to have shorter wetting time at 0.14 wt-pct concentration than the other surfactants at much higher concentrations. Apparently SD2ES has a greater affinity for hydrophobic coal sites and possibly greater ability to close-pack molecules on the coal surface than the other reagents. This wetting superiority extended also to the observed wettabilities obtained after Na₂SO₄ addition. The response of SD2ES to the electrolyte addition was sufficient to maintain faster wetting times over the other wetting agents. However, the response of SDBS and SDHS surfactants to Na₂SO₄ addition was particularly strong, enabling these reagents to perform nearly as well as SD2ES in this wetting range, although much higher concentrations of surfactant were required. The response of SDBS to Na₂SO₄ addition was weaker than that of the other anionic surfactants, which may be related to excess Na₂SO₄ concentration in SDBS solution resulting from the additional Na₂SO₄ supplied by the surfactant's impurity content. The poorest wetting response resulting from Na₂SO₄ addition was observed for EAG surfactant, a consequence of its nonionic properties.

The coal wetting superiority of SD2ES on hard-to-wet coal was also evident for surfactant solution concentrations in the fast-wetting range, but to a lesser degree (fig. 15). For easy-to-wet coal, the wetting responses were rather similar among the anionic surfactants, although SD2ES was still slightly more effective (fig. 16). In the wetting range shown, Na₂SO₄ addition was not particularly effective for any of the surfactants.

EFFECT OF SURFACTANT CONCENTRATION ON PROMOTION OF WETTING BY SULFATE

Hard-to-Wet Coal

It was observed earlier that the wetting response of anionic surfactant solutions to Na₂SO₄ addition is greatly dependent on surfactant concentration. Figure 17 compares this concentration dependency for three anionic surfactants (SD2ES, SDBS, and SDS) and the nonionic EAG surfactant when applied to hard-to-wet coal in the presence of 15 mmol/L added Na₂SO₄. The ordinate in the figure is given in percentage units of wetting improvement (as defined earlier) to compress the scale for better ease of comparison. Generally, it is seen from the figure that the promotion of wetting from Na₂SO₄ addition (on hard-to-wet coal) was decreased for anionic surfactants as the surfactant concentration was increased. On the other hand, changing the concentration of EAG surfactant had only a small effect on promotion of wetting by Na₂SO₄.

SDBS surfactant demonstrated less response to concentration changes than SD2ES and SDS when compared in the concentration ranges of about 0.4 to 0.6 and 0.7 to 0.8 wt-pct of SD2ES and SDS surfactant, respectively. However, for SDBS surfactant, the concentrations of sulfate used to compute percent wetting improvements are not accurate since this surfactant contains 20 wt-pct Na₂SO₄ impurity. In reality, the SDBS contribution of Na₂SO₄ impurity provides considerable Na₂SO₄ in solution (depending on surfactant concentration) before any Na₂SO₄ is added separately. For example, there are 5.6 and 11.3 mmol/L of Na₂SO₄ in 0.4- and 0.8-wt-pct SDBS surfactant solutions, respectively, before any addition of electrolyte. Thus, the calculation of the percent wetting improvement after adding 15 mmol/L of Na₂SO₄ to an 0.8-wt-pct SDBS surfactant solution is really the difference between solutions containing 11.3 and 26.3 mmol/L of Na₂SO₄. This is likely an important wetting response factor, since the greatest response to Na₂SO₄ addition by anionic surfactant tends to occur at the lowest range of Na₂SO₄ concentrations. For example, in figure 6, for 0.14 wt-pct SD2ES, a much greater wetting improvement occurs between zero concentration of Na₂SO₄ and 15.0 mmol/L Na₂SO₄ than between 11.0 and 26.0 mmol/L Na₂SO₄. This factor also explains the feeble wetting response of high concentrations of SDBS in figure 10 (curve A). High SDBS concentration results in excessive Na₂SO₄ concentration and, therefore, reduced wetting response.

Similar circumstances are also visible in figure 9 for SDS (curve A) and in figure 11 for SDHS (curve B). Therefore, it is safe to say that the slope of curve C in figure 17 would likely have been greater and aligned better with the SDS and SD2ES curves if a purer sample of SDBS had been available for testing.

Easy-to-Wet Coal

Figure 18 displays the wetting promotion of Na₂SO₄ as a function of the concentration of SD2ES, SDS, and SDBS anionic surfactants on easy-to-wet coal. As in figure 17, the wetting dependencies are expressed in percent wetting improvement of surfactant solution with added Na₂SO₄ (15.0 mmol/L) compared with the same surfactant solution without added Na₂SO₄. Figure 18 reveals that the promotion of wetting by 15 mmol/L of Na₂SO₄ was greatly different depending on the anionic surfactant used and its concentration. This result was dissimilar to results when anionic surfactant was applied to hard-to-wet coal, which showed a wetting promotion dependent chiefly on surfactant concentration and not on the anionic surfactant used (figure 17, curves B, C, and D).
Figure 14.—Comparison of surfactant performance on hard-to-wet coal in slow-wetting range.
Figure 15.—Comparison of surfactant performance on hard-to-wet coal in fast-wetting range.

Figure 16.—Comparison of surfactant performance on easy-to-wet coal in fast-wetting range.
Figure 17.--Influence of surfactant concentration on wetting promotion by sulfate for hard-to-wet coal. Percentage wetting improvement for surfactant solutions containing 15 mmol/L added Na₂SO₄ compared with the same surfactant solutions without added Na₂SO₄.
SD2ES surfactant showed a maximum percent wetting improvement at about 0.61 wt pct surfactant concentration, while wetting improvement dropped off abruptly at lower and higher concentrations (fig. 18). On the other hand, SDS surfactant showed a gradual wetting improvement as its concentration was increased (fig. 18). For SDBS, the percent wetting improvement was greatest at low concentrations but dropped off rapidly until, at about 0.44 wt pct surfactant, wetting improvement leveled off, and higher concentrations had little effect on it.

In the case of SD2ES, higher surfactant concentration had little effect on wetting beyond the maximum. The wetting maximum for SD2ES suggests that 15 mmol/L of Na₂SO₄ may not be sufficient to compete with a high concentration of SD2ES surfactant ions for hydrophilic coal sites despite the valence advantage of sulfate ion in the ion-exchange reaction.

It is evident, then, that application of Na₂SO₄ to aid the performance of anionic surfactants applied to easy-to-wet coal is a complicated matter. Wettability testing is
required to determine the optimum concentrations of surfactant and additive for a given easy-to-wet coal. In contrast, for hard-to-wet coal, application of Na$_2$SO$_4$ is always effective at lower concentrations of anionic surfactant.

**RELATION BETWEEN SURFACTANT COAL-WETTING PERFORMANCE AND SURFACE TENSION**

Comparison of the ranking of wetting ability for the various surfactants with their ranked ability to reduce the surface tension of water indicates poor correlation except for SD2ES surfactant, which performed better than the other reagents in both reducing surface tension and wetting time (compare figure 19 with figure 14).

Hence, coal-wetting improvement through surfactant addition is not merely a function of a surfactant's ability to reduce the surface tension of water. The surface tension does need to be reduced below 45 dyn/cm, but most surfactants will reduce the surface tension below this critical value easily at low concentrations. For example, less than 0.05 wt pct was required for SD2ES. If spontaneous wetting is achieved at 45 dyn/cm, why should further reduction in surface tension aid wetting beyond this point? Figures 6-7 and 9 affirm that higher concentrations of surfactant and Na$_2$SO$_4$ tend to improve coal wetting up to a concentration limit, but the limit occurs far beyond the concentrations giving critical surface tension.

Surface tension measurements of the wetting solutions do suggest the possible occurrence of close-packing phenomena at the coal-liquid interface. The reduced liquid surface tension from adding Na$_2$SO$_4$ to anionic surfactant solutions indicates that analogous close packing of surfactant molecules is occurring at the liquid-air interface. Table 3 shows the decrease in surface tension of anionic surfactant solutions as Na$_2$SO$_4$ is added, compared with the constancy of surface tension for EAG nonionic surfactant solution regardless of the presence of Na$_2$SO$_4$ at a given surfactant concentration. The surface tension reduction achieved for anionic surfactant through adding Na$_2$SO$_4$ is derived from improved close packing of surfactant at the air-liquid interface, caused by reduced electrostatic repulsion forces. On the other hand, the surface tension of nonionic surfactant solution and its wetting performance are unaffected by Na$_2$SO$_4$ since electrostatic forces are not operating in this case, and electrolyte addition will not affect close packing of molecules at either the air-liquid or coal-liquid interfaces. If anything, excessive Na$_2$SO$_4$ addition would tend to increase surface tension in this case, since concentrated electrolyte solutions (in the absence of surfactant) are known to increase surface tension above that for pure water (9).

**Table 3.—Liquid surface tensions, (air-liquid interface) of the wetting solutions at 28° C, dyn/cm**

<table>
<thead>
<tr>
<th>Surfactant</th>
<th>Added Na$_2$SO$_4$ conc</th>
<th>Reagent conc, wt pct</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 mmol/L</td>
<td>1.0 mmol/L</td>
</tr>
<tr>
<td>SD2ES</td>
<td>.004</td>
<td>47.3</td>
</tr>
<tr>
<td></td>
<td>.008</td>
<td>46.4</td>
</tr>
<tr>
<td></td>
<td>.041</td>
<td>36.4</td>
</tr>
<tr>
<td></td>
<td>.14</td>
<td>28.4</td>
</tr>
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<td></td>
<td>.20</td>
<td>28.0</td>
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<tr>
<td></td>
<td>.41</td>
<td>27.5</td>
</tr>
<tr>
<td></td>
<td>.61</td>
<td>26.5</td>
</tr>
<tr>
<td></td>
<td>.82</td>
<td>25.9</td>
</tr>
<tr>
<td>SDHS</td>
<td>.11</td>
<td>47.2</td>
</tr>
<tr>
<td></td>
<td>.23</td>
<td>40.5</td>
</tr>
<tr>
<td></td>
<td>.77</td>
<td>34.3</td>
</tr>
<tr>
<td>SDS</td>
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<tr>
<td></td>
<td>.74</td>
<td>38.4</td>
</tr>
<tr>
<td></td>
<td>.99</td>
<td>37.5</td>
</tr>
<tr>
<td>SDBS</td>
<td>.24</td>
<td>33.5</td>
</tr>
<tr>
<td></td>
<td>.44</td>
<td>32.5</td>
</tr>
<tr>
<td></td>
<td>.60</td>
<td>32.0</td>
</tr>
<tr>
<td>EAG</td>
<td>.16</td>
<td>35.4</td>
</tr>
<tr>
<td></td>
<td>.42</td>
<td>30.8</td>
</tr>
</tbody>
</table>

ND Not determined.
Figure 19.—Surface tension ranking at 28° C for various solutions containing surfactants plus Na₂SO₄ (at concentrations corresponding to wetting measurements made in the slow-wetting range on hard-to-wet coal).
It might be argued that close packing of surfactant molecules at the coal-liquid interface is wholly responsible for the improved wettability observed with Na$_2$SO$_4$ addition. However, close-packing phenomena on the coal surface would be predicted to affect wettability most noticeably at high anionic surfactant concentrations. But such was not the case in figure 6 (curves B, C, D, and E), where wetting improvements for high surfactant concentrations were modest or nonexistent upon increasing the Na$_2$SO$_4$ concentration.

A minimum surface tension of about 24.8 dyn/cm was achieved at about 0.20 wt pet SD2ES plus 30 mmol/L Na$_2$SO$_4$ (table 3). This value likely represents the point of ultimate close packing of surfactant at the air-liquid interface. Significantly, further SD2ES addition affected the wetting of the coals (positively or negatively) despite the likelihood of maximum close packing of adsorbed surfactant molecules at the coal-liquid interface. Therefore, these wetting changes must originate from the adsorption phenomena described earlier rather than from close-packing phenomena.

It is interesting to note that the minimum surface tension obtainable with SD2ES and Na$_2$SO$_4$ is approaching that of pure hydrocarbons of similar chain length, such as n-decane. One might surmise that the packing of surfactant molecules at the liquid-air interface is approximating molecular spacing. Thus, n-octane has a surface tension of 21.8 dyn/cm (10), while n-decane and n-dodecane are reported as 23.9 and 25.4 dyn/cm, respectively, at 20° C (11).

CONCLUSIONS

It was shown in this work that the favorable coal-wetting response previously achieved with SD2ES anionic surfactant, when adding a multivalent anion such as sulfate, was not an isolated case depending on a singularity of the SD2ES surfactant. Experiments with other anionic and nonionic surfactants indicated that improved wetting response for anionic surfactants in the presence of Na$_2$SO$_4$ was a consequence of their negatively charged character and, therefore, generically applicable to all anionic surfactants as a class. The limited response to Na$_2$SO$_4$ addition by nonionic surfactant further supported this conclusion.

Several factors were identified as having the potential to affect the wettability of coal by surfactants and by combinations of surfactant with electrolyte additive containing a multivalent anion. The factors are

1. Reduction of the surface tension of water by surfactant or surfactant-additive combination;
2. Adsorption of surfactant on hydrophobic sites of the coal surface to create hydrophilicity;
3. Adsorption of surfactant on hydrophilic sites of the coal surface in reversed orientation to create undesirable hydrophilicity;
4. Adsorption of multivalent anions, such as sulfate, on hydrophilic sites of the coal surface to preserve hydrophilicity;
5. Close packing of surfactant molecules on hydrophobic sites of the coal surface to improve hydrophilicity; and
6. Adsorption of surfactant to form hemimicelles on top of previously adsorbed surfactant to restore hydrophilicity.

The first factor was determined to be important only in reducing surface tension below a critical value for coal (45 dyn/cm). Below this value, liquid surface tension ceases to be important except that it may reflect the situation at the coal-liquid interface with respect to close packing of adsorbed surfactant molecules. However, in extensive testing with SD2ES surfactant, a maximum close packing appeared to be achieved on the coal surface (indicated by a relatively constant liquid surface tension), and yet wettability changed with further surfactant and Na$_2$SO$_4$ addition. Therefore, the close-packing factor 5 is considered of less importance than the adsorption factors 2, 3, and 4, which appear to dominate the wetting action of anionic surfactants on coal.

The hemimicelle formation factor 6 may also have some validity, but the applicability is tenuous since it is difficult to attribute certain experimental observations on hard-to-wet coal to this phenomenon while being unable to detect any presence of the phenomenon on easy-to-wet coal.

The anionic class of surfactants was more effective than nonionic reagents in wetting hard-to-wet coal. However, the performance of anionic surfactants can be inconsistent because of the electrostatic interactions between the charged coal surface and surfactant ions. Nevertheless, the employment of nonionic surfactants to avoid electrostatic interactions is not recommended. If nonionic surfactant adsorbs in adverse orientation or with poor close packing on the coal surface, nothing can be done to ameliorate the situation except to increase concentration (economically disadvantageous) or try a new surfactant. However, for anionic reagents, both of these conditions can be corrected by addition of an appropriate, inexpensive electrolyte such as Na$_2$SO$_4$. Nonionic reagents are regarded as ultimately more limited than anions, since the electrostatic properties of the anionics can be manipulated advantageously with additives.
REFERENCES