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- Publications Acknowledging ARO Support in the Grant Period:
 - 1) F. M. Menger and A. L. Galloway, to be published in *J. Am. Chem. Soc.* "Contiguous vs. Segmented Hydrophobicity in Micellar Systems."
 - F. M. Menger, A. L. Galloway, M. E. Chlebowski, and R. P. Apkarian J. Am. Chem. Soc. 2004, 126, 5987. "Ultrastructure in Frozen/Etched Saline Solutions: On the Internal Cleansing of Ice."
 - F. M. Menger, A. L. Galloway, and D. G. Musaev Chem. Commun. 2003, 2370. "Relationship between Rate and Distance."
 - 4) F. M. Menger *Proc. Natl. Acad. Sci. U.S.* **2002**, *99*, 4818. "Supramolecular Chemistry and Self-assembly."
 - 5) F. M. Menger and V. A. Azov *J. Am. Chem. Soc.* **2002**, 124, 11159. "Synthesis and Properties fo Water-Soluble Asterisk Molecules."
 - 6) F. M. Menger, H. Zhang, K. L. Caran, V. A. Seredyuk, and R. P. Apkarian *J. Am. Chem. Soc.* 2002, 124, 1140.
 "Gemini-Induced Columnar Jointing in Vitreous Ice. Cryo-HRSEM as a Tool for Discovering New Colloidal Morphologies."
 - V. A. Migulin and F. M. Menger Langmuir 2001, 17, 1324.
 "Adamantane-Based Crystals with Rhythmic Morphologies."

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Report

May 31, 2004 marks the end of a 2-decade long association between my research group and the ARO. From my point of view, and I hope from the ARO's also, it has been a productive relationship. Above all, I appreciate how Drs. Ghiardelli, Seiders, and Lee have given me the freedom to let our imaginations wander and thus to produce the best chemistry we are capable of pursuing. In a spirit of nostalgia, I would like to summarize below some of the highlights of our ARO-funded research, particularly those papers directly related to decon of CWA.

1) F. M. Menger and L. G. Whitesell, *J. Am. Chem. Soc.* **1985**, *107*, 707. "A Protease Mimic with Turnover Capabilities."

This paper described the synthesis of a micellar aldehyde (dubbed "Atlanta 1") whose hydrate would attack and hydrolyze bound esters. The important point is that the ester cleavage occurred with *turnover*, i.e. with a true enzyme-like catalysis which, at that time, was a rather unique mechanism.



2) F. M. Menger, L. H. Gan, E. Johnson, and D. H. Durst J. Am. Chem. Soc. 1987, 109, 2800. "Phosphate Ester Hydrolysis Catalyzed by Metallomicelles."

A metallomicelle containing copper (dubbed "Atlanta 2") was found to hydrolyze phosphate esters with rate accelerations approaching 10⁵. The combination of micellar binding and metal catalysis led to a remarkable catalysis for nerve agent simulants.



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3) F. M. Menger and A. R. Elrington *J. Am. Chem. Soc.* **1990**, *112*, 8201. "Rapid Deactivation via Microemulsion Technology."

F. M. Menger and A. R. Elrington *L'Actualite chimique* **1991**, 177. "Formulating Microemulsions that Decontaminate a Toxic Substance."

F. M. Menger and H. Park *Recl. Trav. Chim. Pays-Bas Journal* **1994**, *113*, 176. "Microemulsions as Reaction Media. Self-organizing Assemblies in an Environmental Cleanup Problem."

F. M. Menger and M. J. Rourke *Langmuir*, **1999**, *15*, 309. "Deactivation of Mustard and Nerve Agent Models via Low-Temperature Microemulsions."

Prior to the 1990's, the emphasis had been on micellar catalysis, but it became evident that micelles have a major disadvantage: their solublization capacity is low. It was for this reason that we switched to oil-in-water microemulsion systems that had a much greater ability to solubilize water-insoluble organics such as mustard. Probably our most important achievement in this area was a complex system that contained 11 components: propylene glycol, oil, stabilizer, two nerve agent simulants, and a mustard simulant. Reactions at 25° took place in seconds or over many minutes. Only by virtue of self-assembly does this extraordinarly complex mixture adopt a useful molecular organization. Since the microemulsion has the important property of resisting freezing, it does not have to be kept warm in a cold climate. Note that our microemulsion has similarities to DeconGreen, developed recently at Aberdeen. It too uses propylene glycol as a major component of the solvent system. In contrast to our household bleach being the reactant, DeconGreen uses fairly concentrated hydrogen peroxide (35%). Hydrogen peroxide is less reactive than hypochlorite, thereby requiring the presence of a metal catalyst. Both our low-temperature microemulsion and the DeconGreen must be prepared prior to use (in our case at least a week), thus skirting the desirability (and in my view impossibility) of having a non-caustic single-solution with a shelf-life of 5 years.



Figure 1. The 8-component oil-in-water microemulsion used in this study. NPDPP, TDP, and R_2S are three contaminants in the oil droplet (see text) which are destroyed by hypochlorite in the aqueous phase. Surfactant (heavy dots with tails) and cosurfactant (alcohol) stabilize the oil droplet.

3.

4) The preceding lists papers directly relate to the decon problem. But there were many other ARO-supported publications that delved into related subjects such as the mechanistic source of fast catalytic rates; the construction of new catalytic polymers (i.e. high-porosity, copper-containing polymers); packing behavior in monomolecular films; ion conductance in lipid monolayers; giant vesicles; transport through liquid membranes; synthesis of organic self-assembled fibers; synthesis of new polyhydroxylated surfactants; enzyme-vesicle interactions for detection and release; remote steric effects; and synthesis of asterisk molecules.

ARO-supported papers in these areas have been listed in previous reports and will not be repeated here. The point here is that ARO funding over the years has been critical to the diversity and richness of our research program area, and both my students and I greatly appreciate this assistance.

Report Summary of Current Grant

1) Ester-Modified Surfactants

On the next page are listed six new ester-containing cationic surfactants and a few of their colloidal properties. The project had two purposes: (a) We wanted to determine if endowing surfactant chains with ester groups could improve the solubilizing capacity of their micelles (much as ethyl acetate is a better solvent than hexane). (b) We wanted to compare the hydrophobicity of a contiguous hydrocarbon chain (e.g. a dodecyl chain) with that of an equivalent number of carbons but segmented (e.g. two ester-separated six-carbon chains).

We found that in three of the surfactants (A, B, and E) segmentation was deleterious to colloidal properties such as aggregation number and solubilization power. On the other hand, the three other surfactants (C, D, and F) were much more conventional in their colloidal properties. In fact, D solubilized mesitylene more effectively than did DTAB and TTAB (see Table). A long discussion of the surfactants will appear in a *J. Am. Chem. Soc.* paper that is now in press. Its Abstract summarizes are main findings:

This paper addressed a question not yet posed systematically in surfactant chemistry: How do the colloidal properties of surfactants respond to insertion of nonhydrocarbon functionalities (i.e. ester groups) within chains that are normally entirely hydrocarbon? In answering this question, two classes of such chain-modified surfactants were discovered. One class forms only small aggregates with noncooperative self-assembly, low foaming, high areas of occupancy at the air/water interfaces, and weak solid adsorption and solubilization properties. The other class is much more normal with regard to their properties and, in fact, can even exceed conventional surfactants in mesitylene solubilization. Differences between the two

Ester-Modified Surfactants

Label	$#CH_2$'s	Structure
Α	12	O O O N ⁺ Me ₃ Br
B	14	O O O N ⁺ Me ₃ Br
С	16	0 0 0 N ⁺ Me ₃ Br [−]
D	18	0 0 0 N ⁺ Me ₃ Br ⁻
E	15	O O N ⁺ Me ₃ Br [−]
F	13	O O N ⁺ Me ₃ Br [−]
DTAB	11	∩ N ⁺ Me ₃ Br ⁻
TTAB	13	N ⁺ Me₃Br ⁻

Surfactant	CMC, mM	Area, Å ²	Foam Vol., mL	TMS, mM	Mesitylene, mM
Α	2.4	81	0.01	0	1.7
В	2.5	123	0.48	0.026	5.3
С	2.2	75 [·]	2.5	1.2	38
D	1.4	75	3.2	5.2	93
E	1.0	334	0.04	0	0
F	4.6	54	3.5	5.4	27
DTAB	13.3	64	1.9	0.51	13
TTAB	3.3	52	3.3	7.9	36

categories of chain-modified surfactants originate from the degree of segmentation of the hydrocarbon and, in particular, upon the location of the longest segment. Segmented hydrophobicity, having in priniciple a similar "hydrophobic potential" as a contiguous hydrophobicity of equal length, can induce aggregation but, concurrently, alter the modes of assembly into films and micelles.

2) The Ultrastructure of Frozen Saline Solutions

The instant the temperature drops below 32° F the Armed Forces must confront a serious problem: ice. In the decon area, for example, an aqueous system must be kept above its freezing point if it is to be ready for use at a moment's notice. It was to address this problem that we developed a decon containing an anti-freeze, propylene glycol, and apparently DeconGreen utilizes the same strategy. This led us, quite naturally, to wonder about the fate of water containing simple additives, such as salt and surfactants, when it is frozen. What happens to the salt when saline freezes rapidly? Thus, we used a method, cryo-etch-high resolution scanning electron microscopy or cryo-etch-HRSEM (which Emory is fortunate to possess in our chemistry department), to investigate the freezing of solutions and the fate of the solutes within them.

In a typical experiment, 10 uL of a 2-5 wt % aqueous NaCl solution was sandwiched between two gold planchettes. The sample was then loaded into a Balzers HPM 010 high-pressure freezer where liquid nitrogen was sprayed from two sides of a pressurized chamber (2.1 kbar). Freezing the sample from 20 to -105° C took place in 5-6 msec (22000-25000° C/s). After freezing, the sample was immediately immersed in liquid nitrogen to maintain a temperature below -170° C. Next, the sample was placed into a Gatan 3500 CT cryostage precooled to -170° C. A cold blade was then used to fracture the specimen and expose a fresh surface. Etching and coating of this surface were carried out after transferring the cryostage to a Denton DV-602 chromium coater where a vacuum of 2 x 10^{-7} Torr was maintained. Thus, the temperature was raised from -170° to -125° or -105° C for time periods ranging from 5 to 120 min to allow the exposed ice to vaporized by sublimation ("etching"). Once the etching was completed, the sample was cooled back to -170° C and sputter-coated with chromium at a rate of 0.3 A/s using a current and voltage of 50 mA and 300 V, respectively, under a 5 x 10⁻³ Torr atmosphere of argon. The resulting 2-nm Cr film has the advantage of possessing smaller grain sizes that do the precious metals. Finally, the sample was removed from the coater and transferred to a DS-130F field emission scanning electron microscope (beam diameter = 1 nm) fitted with dual coldfinger anti-contamination traps and a Varian 860 cold-cathode gauge for monitoring the vacuum at the specimen. Images (4.8 MB) recorded in 16 s at -120° C with a Pentium-based GW video capture board, were processed using Adobe Photoshop 6.0 as TIF files.

The upshot of this rather technical method was that granular icy "fences" were seen surrounding empty areas where amorphous ice had originally resided (see picture below). The presence of such fences suggests that, during freezing, saline can purge itself of salt with remarkable speed (5-6 msec). Alternatively, channels (perhaps routed around submicroscopic crystallites of cubic ice (I_c) embedded in the amorphous ice at -105° C) can guide the migration of salt to the periphery of ice patches. Aside from the fundamental interest of these results, salt morphologies are important to biologists who may encounter fences when examining buffered samples by cryo-etch methods. The results also pertain to the rapid freezing of sea water on cold metal surfaces such as presented by ships and aircraft. In general, the study of frozen solutions merits considerable more attention.



NaCl, 2 wt %, subjected to HPF and etched for 10 min at -105°C

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