Review

- 1. Surfaces of materials have unique descriptive properties:
 - Excess surface free energy
 - Atomic / Molecular composition (vs. bulk)
 - Chemical composition (reactivity vs. bulk)
 - Topography (vs. shape)
- 2. There are numerous surface specific characterization techniques the most prominent of these for evaluating biomaterial surfaces are:
 - 2. Contact Angles
 - 3. ESCA, SIMS
 - 4. SPM (AFM, etc)

These techniques provide information about surface energetics, atomic and molecular composition, surface chemistry, and topography.

VdW interactions between objects

Found by integrating the VdW atom-atom interactions between two bodies and lumping the strength of interaction into geometry-independent constant: HAMAKER constant, *A*.







Protein structure energetics

A balance of free energy contributions determine the stability of protein structure.

Type of interaction	$\Delta_{\text{compact-unfolded}}G$	Remarks
Coulomb	≥ 0	Depending on the pH relative to the isoelectric point of the protein/sorbent complex.
Hydrogen bond	≈ 0	Formation of protein-protein and water-water
Dipole	≈ 0	bonds compensated by loss of protein-water bonds.
Dispersion	≲ 0	Atomic packing densities in compact protein molecules higher than in water.
Hydrophobic dehydration	<< 0	Entropy increase in water released from con- tact with hydrophobic components.
Distortion of bond lengths and angles	> 0	Some bonds are under stress in the folded structure.
Rotational freedom along the poly- peptide chain	>> 0	Folding reduces the conformational entropy of the polypeptide chain and, possibly, the side groups.







Scheme 1 Interdependency of the major subprocesses that are involved in the overall protein adsorption process. Adsorption-promotion is denoted by + and adsorption-opposition by -.

Favorabl	e an	d irr	eversi	ble
Good, e	ol' ∆G =	Δ <i>Η -Τ</i> Δ	.S	
	Lysozyme at pH 10 $(Z_{H} = +5)$			ol)
Brad million and	ΔG (kJ/i	ΔH mol)	ΔS (kJ/kmol)	
Overall protein adsorption process	<< 0	-90	> 0	
Why pH 10? It's close to What kind of surface? Po	pH(iep) lystyren	of prote e with s	ein. What is some negat	pH(ie ive cl

-	$(Z_{\rm H} = +5)$				
	ΔG (kJ/:	∆H mol)	ΔS (kJ/kmol)		
Overall protein adsorption process	<< 0 -90	>0			
Dissociation of H+	-20	0	0.07		
Overlap of electric fields	-10	-20	-0.03		
Change in the chemical medium of the incorporated ions	30	-80	-0.37		
Dehydration of the sorbent surface	-220	-40	0.60		
Rearrangements in the protein structure	< 0	50	> 0		













Incremental, Dynamic Process

Protein adsorption to surfaces is followed by higher order interactions.

- Table 1 Important Processes in the Formation of an Adsorbed Layer of Cells or Proteins
- Approach—The transport of cells, proteins, and other biomolecules to the surface. The surface is only briefly clean and is quickly "conditioned" beginning with a layer of small, abundant proteins or nonprotein macromolecules.
- Initial attachment—Biofluid components adsorb/adhere to a clean surface by a reversibly bound contact point.
- Arrangement—Bound components increase their strength and/or number of surface bonds, while decreasing the reversibility of their attachment. Conformational, positional, and orientational changes occur. Denaturation allows normally hidden hydrophobic groups to seek out nonpolar regions on the surface.
- Interactions—Competition, cooperation, displacement, and exchange lead to a steadystate surface composition markedly different from that in the biological fluid source.

A Short History

It has long been noted that blood coagulated more rapidly on negatively charged glass than on hydrophobically modified glass, or on polymers.

This affect was first attributed to a simple relationship of charge up until ~1960. The idea was that negatively charged surfaces decreased coagulation times in a way that is analogous to the proposed action of negatively charged heparin, an anticoagulant.

Proteins largely have an overall negative charge and were thought to avoid negatively charged surfaces.

The discovery of the surface coagulation activation properties of the negativelycharged protein "Hageman Factor" left some doubt about this theory. It turns out that Hageman Factor was activated on negatively charged surfaces, leading to coagulation.

(Hence begins the study in earnest of proteins on biomaterial surfaces...)

The Search for Heuristics

Using the method of "critical surface energy" developed by Zisman, researchers were able to measure a specific surface property and correlate it to biologic activity.



THE ROLE OF SURFACE ENERGY IN THROMBOGENESIS*

ROBERT E. BAIER, Ph.D. Principal Physicise Cornell Aeronautical Laboratory of Cornell University Bartine, N.Y.

Thus present situation with respect to the evaluation of biomedical basily exploring mechanisms for exis from their own independent circular maxes. After final herakthrough of the walks of any independent circular maxe we discover only that we have all solup been within a much larger, more intricate maze—the whole complex maze of biomedical problems—and that there are many other investigators still within isolated circular loops like the one from which we have recently contenged.

Figure 1 illustrates an overview of three such confining rings, one labeled surface texture; one, surface charge; and the other, surface chemistry. These rings represent three of the primary surface characteristics that are now being carefully examined in the hope of disovering 'magic numbers' which night papily in determining the ultimate thrombogenicity or thromboresistance of a candidate biomaterial. The figure many of the investigators contributing to this program is to find that specific area where their own special interest, their own "magic number" if you will, overlaps any or all of the others, the general goal ommany twint a specific set of parameters that are nguide them in the formulation of new and better blood-constating materials. Still, it is recognizable that the surface qualities of the materials are not the only factors to be considered. They can be contrasted with all of those

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Low Critical Surface Energy Hypothesis



Surface ? Free Energy ? Interfacial ?

D. Lyman (MSE Utah) argued that the surface free energies (vs. critical surface energy) drives protein adsorption and therefore biological activation (as in the case of Hageman Factor). Thus highly charged surfaces are less biocompatible. Examples are glass and blood activation. (New Method: Fowkes)

Andrade (BIOEN, Utah) argued that the free energy of a polymer-water interface is what governs protein adsorption – so as the solid looks more and more like water there is an increase in biocompatibility. Examples are hydrogels and PEO-modified surfaces have reduced coagulation effects. (New Materials: Hydrogels)

Vogler (U Penn) proposed an extension to the free energy theories – that protein adsorption is mediated by water structure at the interface. Thus, Baier's "zone of biocompatibility" exists at the limit between hydrophobic (i.e. Teflon) and hydrophilic (i.e. PEO) materials. (Not sure on the result of this one...)

(New Method: SFG)

What we want to know...

What properties of a biomaterial surface mediate biological response?

To what extent?



Example: Bacterial Adhesion

Bacteria take advantage of surface effects to gain a foothold – then they rework the surface!







Protein adsorption to surfaces

Does it even matter? Yes and no, in a great deal of cases!

Nonetheless, it plays a significant role in:

- Complement activation (IgG, IgM)
- · Coagulation activation (Hageman Factor)
- · Fouling of contact lenses (albumin, lysozyme)
- Interesting scientific pursuits
- Initial response to implants
- Where transport is important (drug delivery)
- etc.

The goal has shifted from understanding the adsorption properties of unmodified materials to intelligent design of materials to mediate the adsorption process (or highjack it entirely.)

