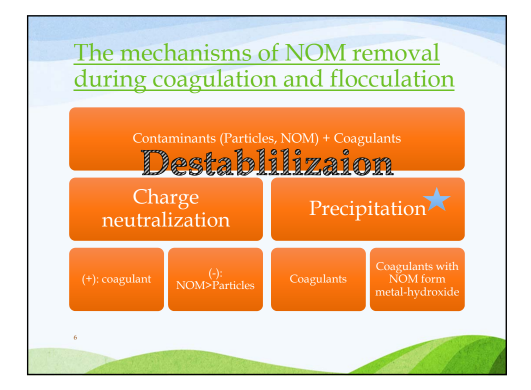


Water treatment processes

- Enhanced coagulation (low pH coagulation)
 - High SUVA water choose enhanced coagulation process
 - Removal efficiencies 25-70% (TOC)
 - The mechanisms of NOM removal during coagulation
 - Better removal of hydrophobic fraction and high molecular weight NOM
- Adsorption
 - Trace organic compounds or NOM (causes odor and tastes, synthetic organic chemicals)
 - Mechanisms: adsorption and biodegradation (depends on Size and chemical properties of NOM)
 - Lower size of NOM has better removal.
- Ion exchange
 - Electronegativity of NOM
- MF/UF
- Ozonation

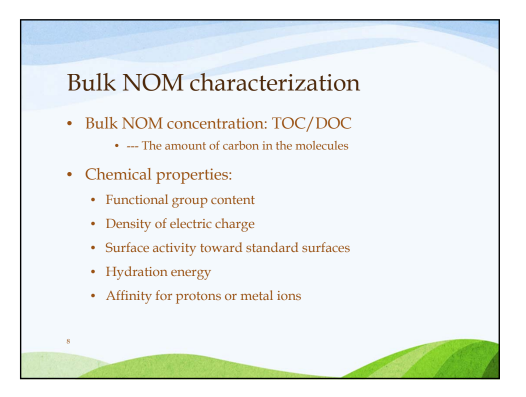


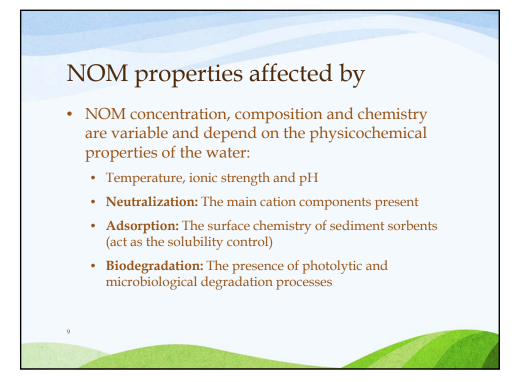
Introduction of NOM Characterization

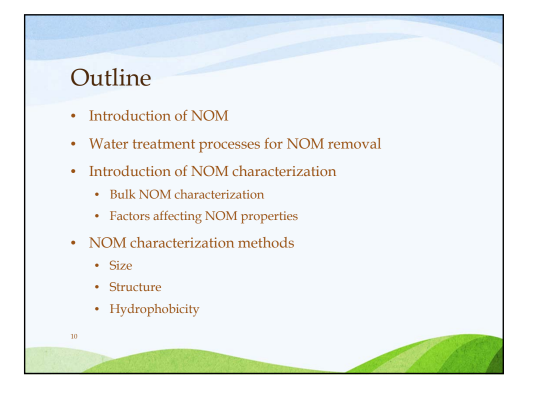
- Purpose:
 - -- predicting and perhaps controlling NOM reactivity
- Difficulty:
 - NOM includes hundreds or thousands of distinct chemical species. It is not realistic to evaluate the properties individually.
- Solution:

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- Bulk NOM properties
- Separate NOM into a limited set of categories
- Characterize the group of NOM by their similar composition and properties





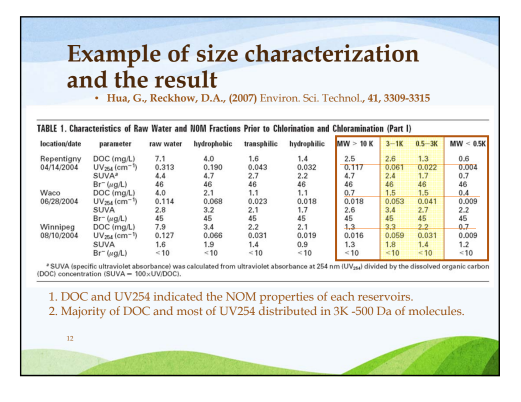


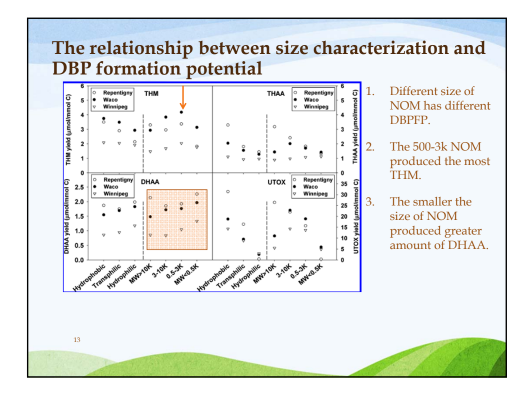
Size characterization of NOM – introduction

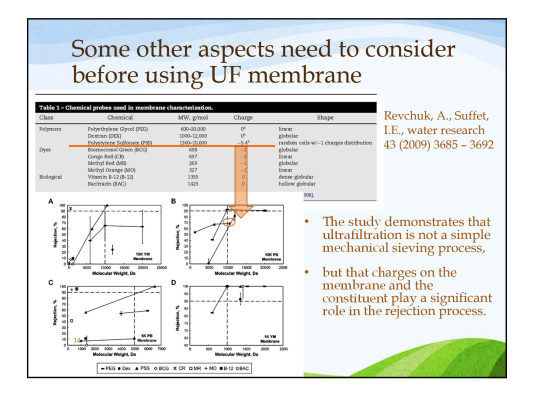
- Most dissolved humic substances have a molecular weight of a few hundred to a few thousand atomic mass units.
- Low-resolution separations:
 - Ultrafiltration using membranes have a specific nominal molecular weight cutoff
 - MW cut-offs of 10KDa, 3kDa and 0.5kDa
- High-resolution separations:

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• size exclusion chromatography (SEC)





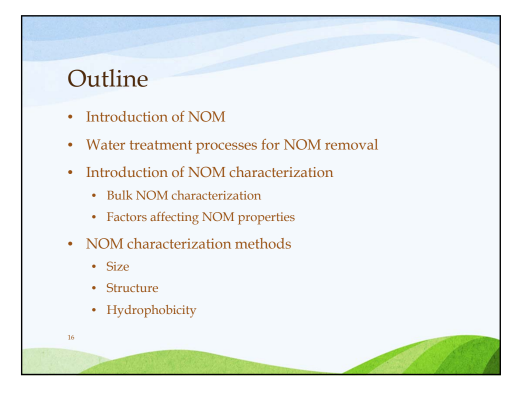


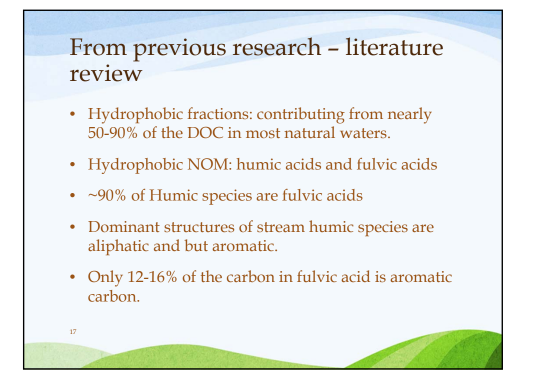
NOM characterization by Structure - introduction

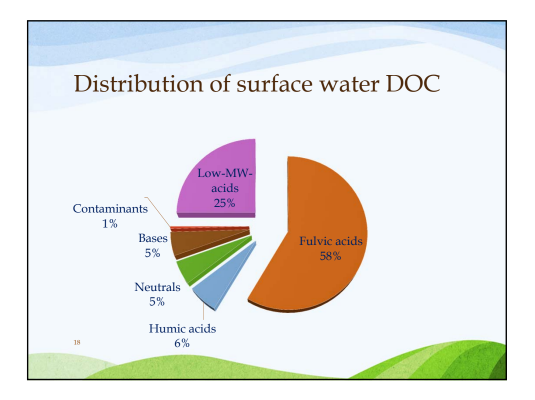
- In general, this is a kind of NOM characterization approach without fractionation.
 - ¹³C-NMR,
 - Fourier Transform Infrared (FTIR) spectroscopy and
 - pyrolysis-gas chromatography-mass spectrometry (Pyr-GC-MS).
 - UV absorbance •
 - Fluorescence spectroscopy
- We should consider: •

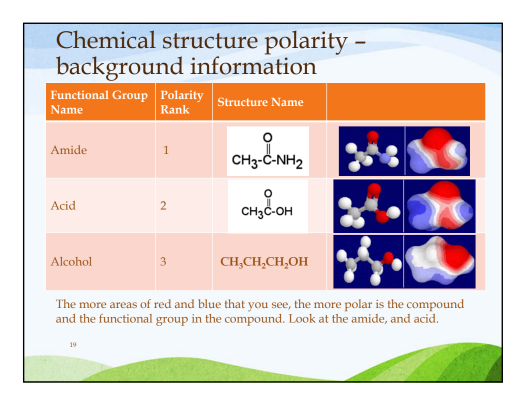
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- Concentrated NOM
- Change of NOM structure during these processes

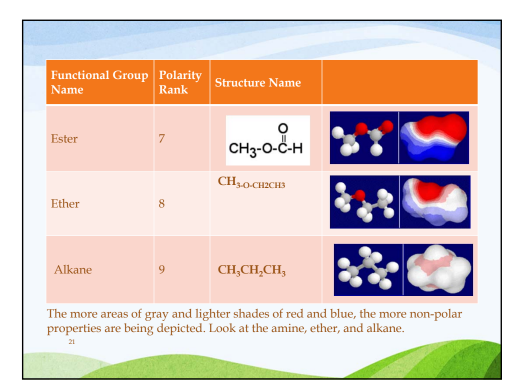


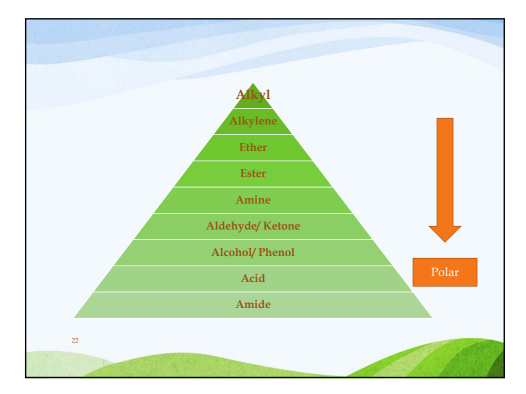


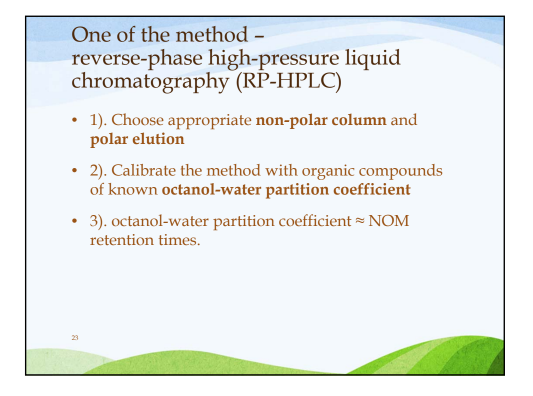


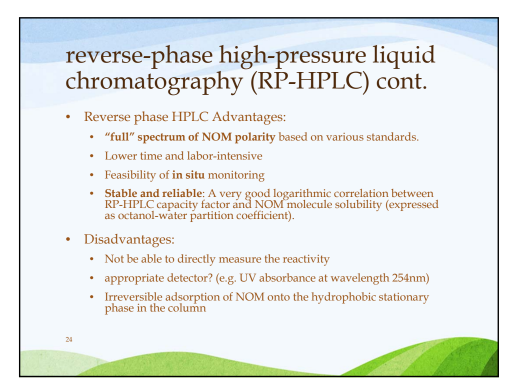


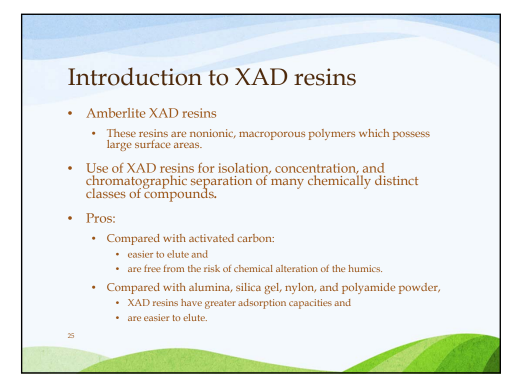
Functional Group Name	Polarity Rank	Structure Name	
Ketone	4, 5	о сн ₃ ссн ₃	*** 🚔
Aldehyde	4, 5	о ен ₃ сн ₂ с-н	*** **
Amine	6	CH ₃ CH ₂ CH ₂ NH ₂	

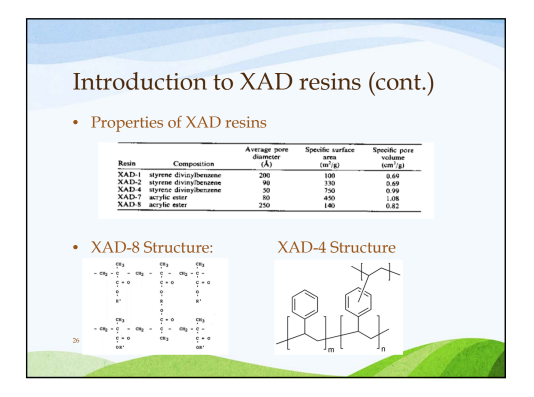


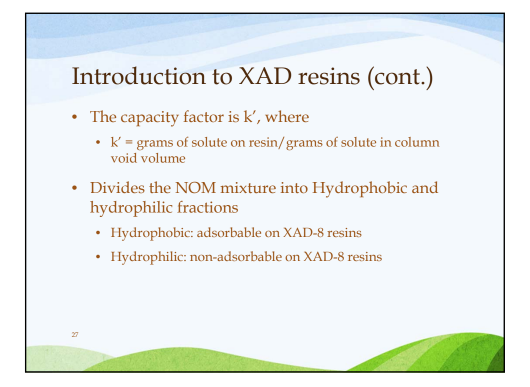


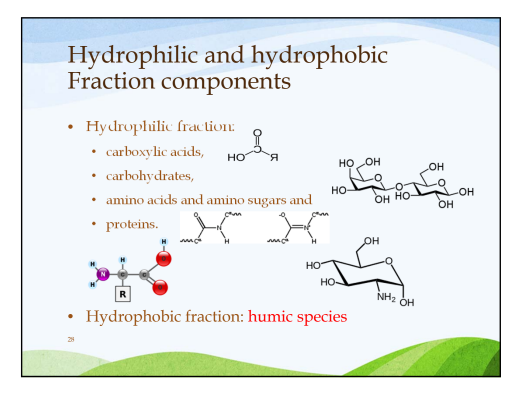


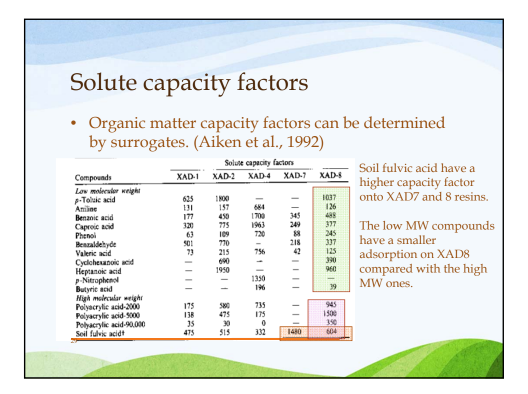


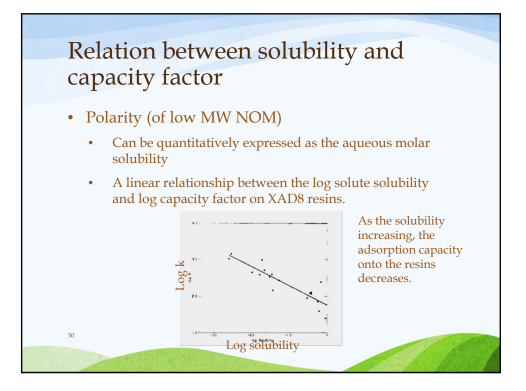


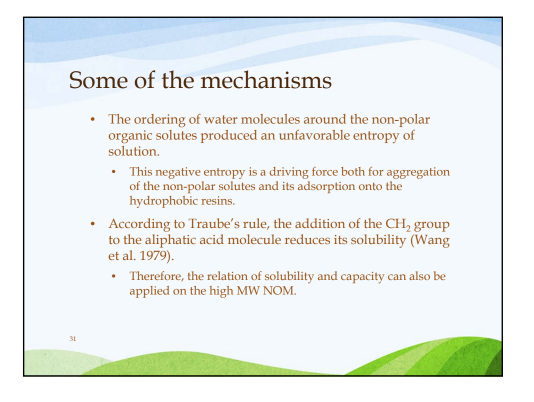






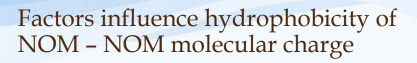






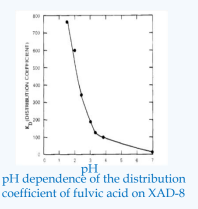
Factors influence hydrophobicity of NOM – MW Size

Resin Composi		Average pore diameter (Å)	Spec	tific surface area (m²/g)	vol	ic pore ume ³ /g)
XAD-1 styrene divinylbe XAD-2 styrene divinylbe XAD-4 styrene divinylbe	nzene	200		100 330	0.	69 69
XAD-7 acrylic ester XAD-8 acrylic ester		80 250		450 140		08 82
		Solute	capacity fi			VAD4 maxima have the smallest man
Compounds	XAD-1	XAD-2	XAD4	XAD-7	XAD-8	XAD4 resins have the smallest pore size, but highest specific surface area
Low moleculor weight p-Toluic acid Aniline Benzoic acid Caproic acid Phenol Benzaldehyde Valeric acid Cyclohexanoic acid Heptanoic acid p-Nitrophenol Butyric acid	625 131 177 320 63 501 73 — —	1800 157 450 775 109 770 215 690 1950 —	 684 1700 1963 720 756 1350 196		1037 126 488 377 245 337 125 390 960 39	low molecular weight solutes have the greatest capacity factors in XAD 4. For high molecular weight organic solutes, size exclusion occurs on XAD resins
High molecular weight Polyacrylic acid-2000 Polyacrylic acid-5000 Polyacrylic acid-90,000 Soil fulvic acid†	175 138 35 475	580 475 30 515	735 175 0 332	 1480	945 1500 350 604	XAD-8 resins were suggested to be used before the smaller pore size resins (XAD-4) to prevent the organ matters clogging.



- K_D decreases as pH raising from 1.5 to 3.
- K_D = mg material adsorbed by resin per gram of resin/mg material in solution per mL of solution
- pH of solution can affect the charge of the organic acids, and consequentially has an effect on the partitioning
- ³³ coefficient.

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Factors influence hydrophobicity of NOM – NOM charge (cont.)

- Adsorption of fulvic or humic acids on XAD-8 resins is favored when they are in the **undissociated form** under low pH condition.
 - XAD-8 resins, composed by acrylic ester, have a measurable cation exchange capacity (Aiken et al., 1992).
- The adsorption result, in part, from intermolecular forces between the undissociated acid molecules and the acrylic ester resins.
- There is also an excellent elution efficiency (approximate 90%) of the XAD-8 resins for humic substances using 0.1N sodium hydroxide as the elution solvent (Aiken et al., 1979).
 - This is attributed to **the charge repulsion** when both the resin and the fulvic acid are anionic at pH 13 (Aiken et al., 1992).
- The similar carbon arrangement could be another reason for humic acids to be favoringly absorbed on XAD-8 resins based on "like dissolves like" principle.
 - Malcolm (1985) indicated that the dominant structures of stream humic species are aliphatic and not aromatic. ¹⁵C NMR analysis in Aiken (1992)'s research proved this, and found the carboxyl groups were the third largest components after aliphatic and aromatic carbons in fulvic acids.

