

BioTek Synergy H1 Microplate Reader for Absorbance Measurements

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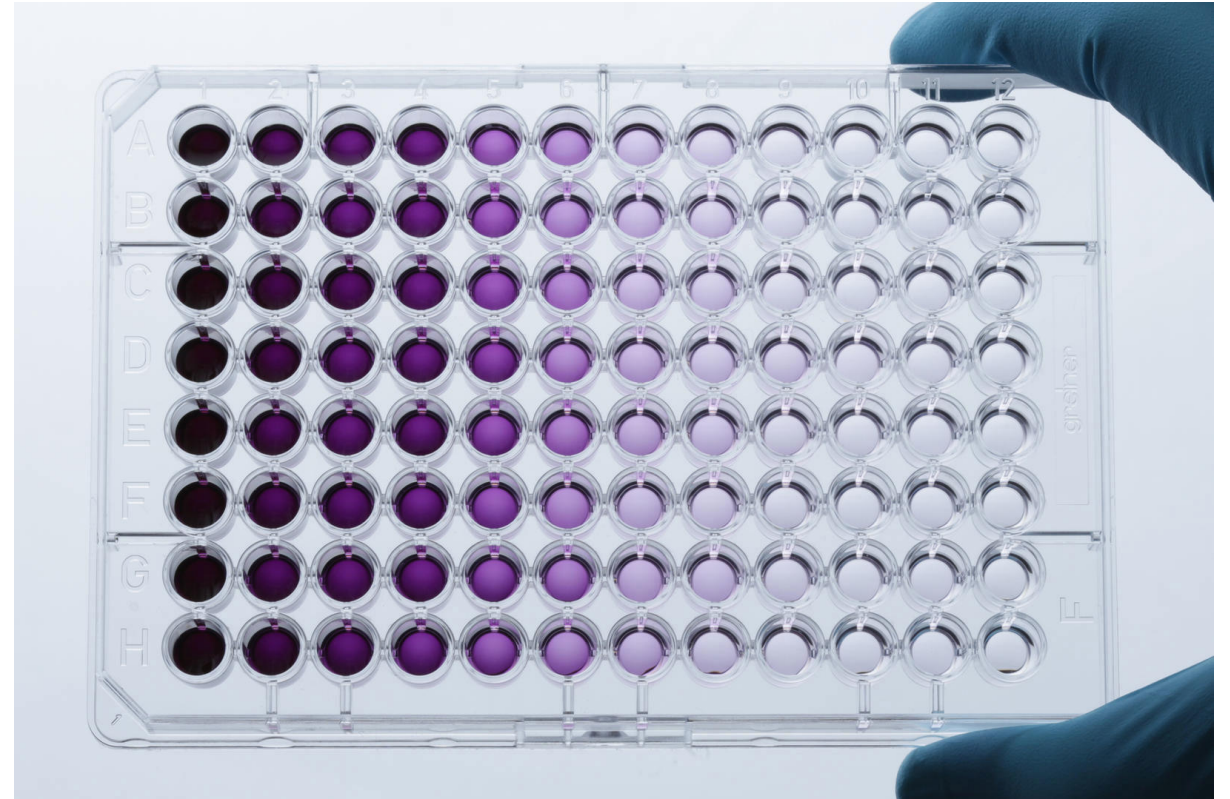
Surface and Colloid Science

Beer-Lambert law relates absorbance and solution concentration linearly

- **Intensity** I - measure of how much light
- **Transmittance** T - ratio of transmitted light vs. incident light at a wavelength λ
 - $T = \frac{I}{I_0}$
- **Absorbance** A - capacity of a substance to absorb light at a wavelength λ
 - $A = -\log_{10}(T) = -\log_{10}\left(\frac{I}{I_0}\right)$
- **Beer-Lambert law** - absorbance varies linearly with solution concentration and path length
 - $A = \epsilon b C$
 - ϵ - extinction coefficient
 - b - path length
 - C - solution concentration

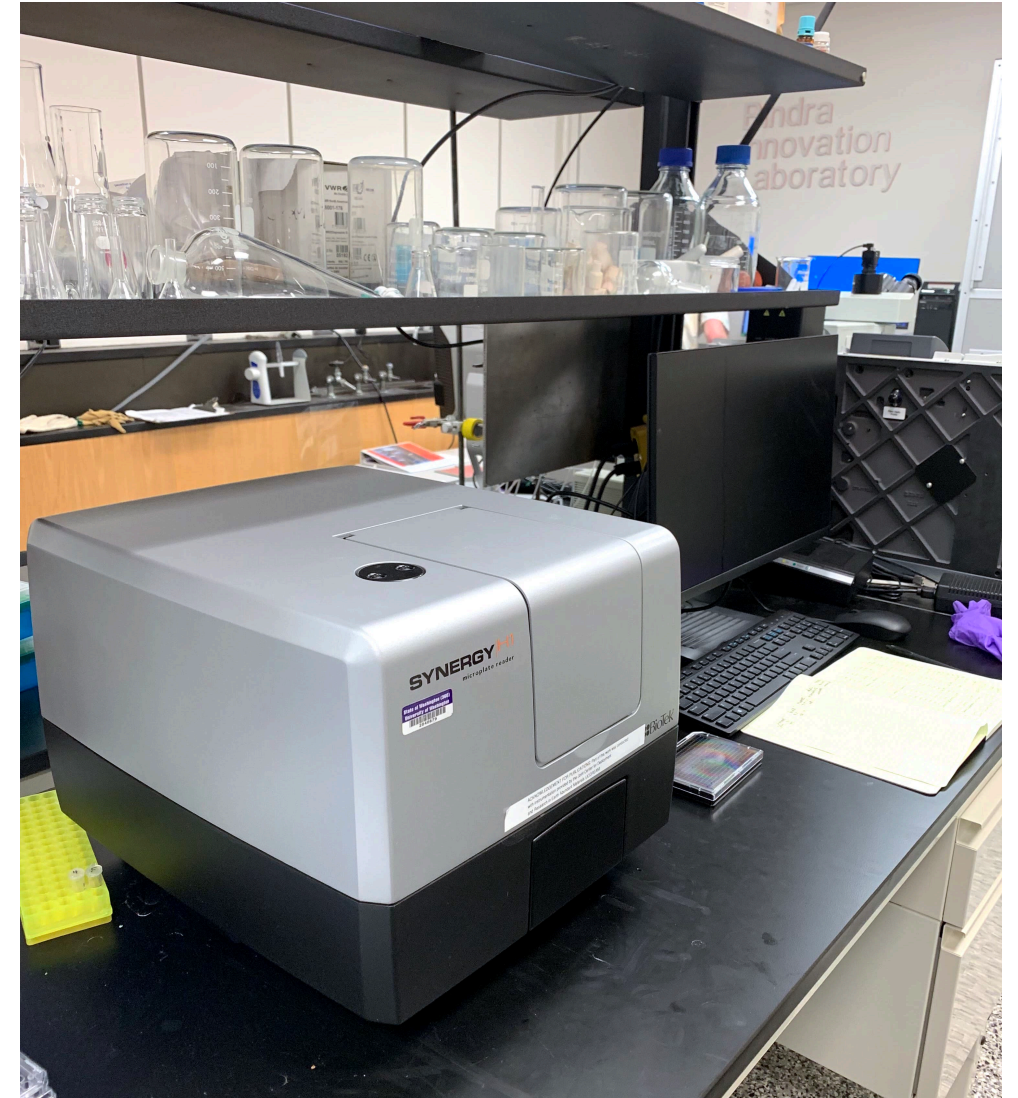
96-well plates contains 200 μL sample of interest in each well

- 200 μL sample in each well
- At least one blank control
- Avoid air bubble (reverse pipetting)
- Label the wells



Microplate reader location, access, and training

- The microplate reader (BioTek Synergy H1) is located in the Bindra Innovation Laboratory (Benson Hall 121).
- Book usage time on shared Google Calendar
- Log usage time on logbook



Microplate reader startup

- Turn on the microplate reader
 - Wait for self-diagnosis
 - Create empty Google Sheet
- Open "Gen5 3.09" software

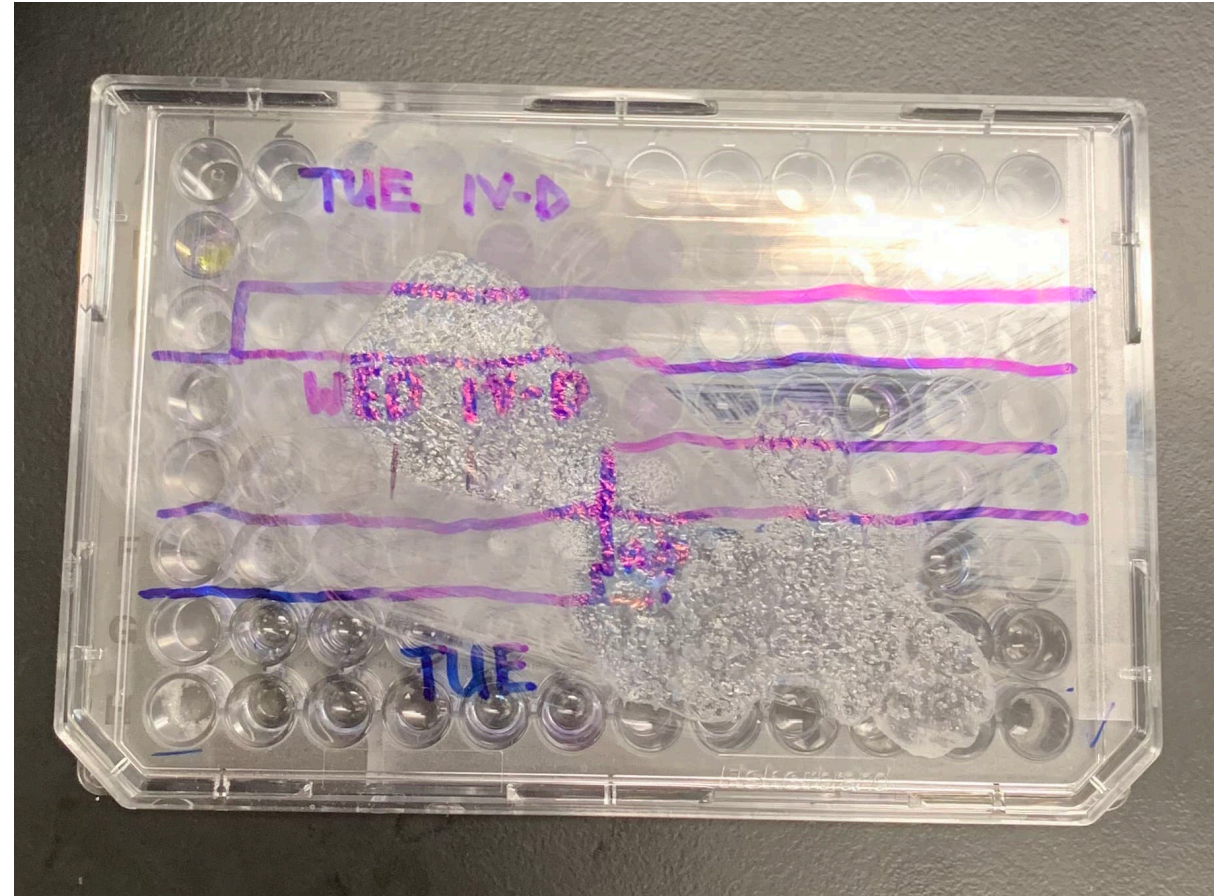


Plate reading settings - absorbance wavelength

- Measurement mode (default)
 - Read method: Absorbance
 - Read type: Endpoint/Kinetic
 - Optics type: Monochromators
- Absorbance wavelength
 - Lab 3-1: CMC by Dye Titration
 - $\lambda = 615$ nm for pinacyanol chloride
 - Lab 3-3: Aggregation of Clay
 - $\lambda = 860$ nm for turbidity
 - Lab 4-4: Bubble Fractionation
 - $\lambda = 590$ nm for crystal violet

Plate reading settings - plate layout

- Plate Layout
 - Select “Blanks” and “Samples”



Absorbance measurement

- Place the plate into the plate holder
 - A1 well is on the top right.
- Read the plate using computer software
- Remove the plate from the plate holder.



Data recording

- Export the data in both matrix and stats form
 - Use blank-subtracted absorbance

Microplate reader shutdown

- Close all programs on the computer.
- Push the IN/OUT button for the plate holder so the plate holder retracts.
- Push the ON/OFF button for the plate reader so the instrument is turned off.

Absorbance measurements for Surface and Colloid Science Laboratory

- Lab 3-1: CMC by Dye Titration
 - $\lambda = 615$ nm for pinacyanol chloride
 - Look for changes in linear trend of absorbance
- Lab 3-3: Aggregation of Clay
 - $\lambda = 860$ nm for turbidity
 - Need calibration curve
- Lab 4-4: Bubble Fractionation
 - $\lambda = 590$ nm for crystal violet
 - Need calibration curve