

SmartPath[®] and High Absorbance Technical Note 120

Introduction

The DeNovix[®] DS-11 spectrophotometer utilizes proprietary SmartPath[®] Technology in conjunction with an innovative microvolume design (patent pending) to enable measurements of samples with absorbance values as high as 750 AU (at a 1cm equivalent pathlength). This means that bovine serum albumin (BSA) samples up to 1125 mg/ml and dsDNA samples up to 37,500 ng/µl can be accurately and reproducibly quantified using the DS-11.

This technical note will provide some background on the relationship between the terms transmittance and absorbance as well as describe the Beer-Lambert law. This information will then be used to explain how the DS-11 enables measurement of samples with ultra high concentrations.

Transmittance

Transmittance (T) is defined as the fraction of incident light (radiant power) at a specified wavelength that passes through a sample as represented by equation 1.

eq 1.
$$T = (I/I_0)$$

I is the light intensity after it passes through the sample and I_0 is the initial light intensity.

For example, completely transparent samples will have $I = I_0$, and therefore the percent transmittance will be 100%. Samples which permit no light at a specified wavelength to pass through will have a percent transmittance of 0%.

Absorbance

Absorbance is defined as the the capacity of a sample to absorb light (radiation) and is expressed as the negative log ratio of transmittance (T) as shown in equation 2.

eq 2.
$$A = -\log T$$
 or $A = -\log (I/I_0)$

Note: The I/I_0 in equation 1 refers to light passed though a single sample. The I/I_0 in equation 2 refers to light transmitted though a sample as a ratio of the the light transmitted through the blank solution.

The table below correlates some absorbance and transmittance values as a reference.

Absorbance (A)	Transmittance (% T)	
0	100	
0.1	79	
0.25	56	
0.5	32	
0.75	18	
1	10	
1.5	3	

In equation 2, if A = 0, then no photons are absorbed and T=100%. If A = 1.00, then 90% of the photons are absorbed and 10% reaching the detector (T=10%).

A practical upper absorbance limit for most spectrometers is 1.5A or 3% T. An absorbance value of 1.5 using the SmartPath 0.02 mm pathlength is equivalent to a 1 cm pathlength absorbance value of 750.



Beer-Lambert Law

The Beer–Lambert law (or Beer's Law) combines separate concepts first described by August Beer⁽¹⁾ and JoHann Lambert^{(2).} Beer's law stated that absorbance is proportional to the concentration of the sample. Lambert's law stated that absorbance is directly proportional to the thickness (or pathlength) of the sample. The combined law correlates the absorbance to both the concentration as well as the thickness (pathlength) of the sample ^[3] and is generally written as:

A=ɛbc

- A: absorbance
- ε: absorptivity coefficient with units of L /mol*cm
- b: pathlength of the sample expressed in terms of cm.c: concentration of the sample in solution, expressed in mol/L.

Note: 1 cm equals 10 mm. All references to pathlengths used on the DeNovix DS-11 and DS-11+ are expressed in terms of mm.

Absorbance Vs Pathlength

As mentioned above, there is a linear relationship between the the absorbance of a sample and the distance (pathlength) the light travels though the sample. Using shorter pathlengths enables samples with higher absorbances (when expressed as 10 mm equivalent values) to be accurately measured.

The DS-11 uses real-time absorbance data to determine the optimal pathlength for each sample. If a sample absorbance detected at the 0.5 mm microvolume pathlength is too high, the DS-11 software will automatically move the arm down as needed to ensure that the measurement stays within the optimal detection range of the instrument.

The DS-11 microvolume mode uses pathlengths ranging from 0.5 mm down to 0.02 mm. Even when the remarkably short pathlength is used, the typical CV associated with the very high concentrated samples is within 3%.

Short Pathlengths, High Concentrations

The DS-11 software uses the Beer-Lambert equation to calculate concentrations based upon absorbance values at specific analysis wavelengths. Keeping in mind the relationships between absorbance, pathlength and concentration described by Beer's law, it is easy to understand that a decrease in pathlength (b) permits a corresponding increase in the measurable concentration (c).

The table below highlights the maximum concentration for two commonly measured biomolecules using the DS-11 microvolume mode as compared to a traditional 10 mm cuvette based system.

Table 2: Maximum Concentration as a Function of Pathlength Abs = 1.5 AU			
		5	
	10 mm	0.5 mm	0.02 mm
dsDNA BSA	75 ng/mL 2.25 mg/mL	0.1	37,500 ng/mL 1,125 mg/mL

The SmartPath® Technology enables the DS-11 to measure samples more concentrated than any other spectrophotometer on the market today making it the ideal choice life science research and manufacturing laboratories.

References

1. Beer (1852) "Bestimmung der Absorption des rothen Lichts in farbigen Flüssigkeiten" (Determination of the absorption of red light in colored liquids), Annalen der Physik und Chemie, vol. 86, pp. 78–88.

2. J.H. Lambert, Photometria sive de mensura et gradibus luminis, colorum et umbrae [Photometry, or, On the measure and gradations of light, colors, and shade] (Augsburg ("Augusta Vindelicorum"), Germany: Eberhardt Klett, 1760). See especially p. 391.

3. Tissue, B. 2013. Basics of Analytical Chemistry and Chemical Equilibria John Wiley & Sons Retrieved from http://books.google.com.

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