

EKKO™ CD Microplate Reader

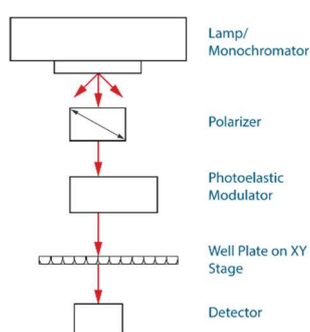
Sensitivity - Detection limits using proteins

I – INTRODUCTION

Circular dichroism (CD) refers to the differential absorption between left and right circularly polarized light is commonly used for chiral analysis. Generally, it is used for determining enantiomeric purities in asymmetric syntheses and assigning the secondary structures of proteins, both of which require the ability to do the measurements in a high-throughput fashion^{1,2}.

Recently these determinations have been expedited by the addition of robotic liquid handling systems coupled to a conventional CD spectrometer. However, transferring individual samples and cleaning the cuvette between measurements makes CD measurements with traditional CD Spectropolarimeters a time-consuming process.

The EKKO™ CD Microplate Reader uses a vertical light path allowing the CD measurements to be read from a well plate directly. As such, the time-consuming processes of 1) transferring the contents from each well of a well plate into a cuvette, and 2) cleaning the cuvette between measurements have been eliminated, significantly increasing productivity, as much as 100-fold with respect to standard CD's coupled to a robot^{1,2,3}. Even with several types



of commercially available well plates, the effective path length will be determined by the upper surface of a solution in the well. It is also expected that the presence of the meniscus would affect the CD measurement results given that regardless of volume in the well.

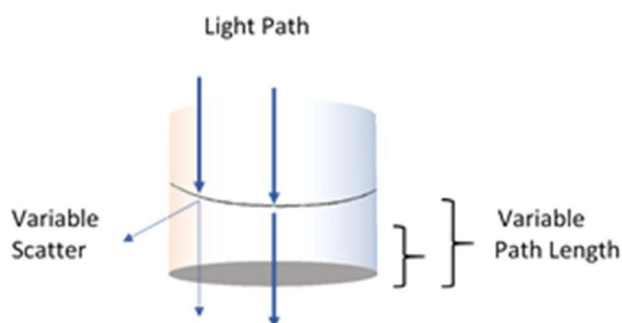


Fig. 1. Presumed effects on the light path as a function of volume and the meniscus in a well.

At a minimum, the combination of the variable path length and meniscus effects have the potential to alter CD determinations when compared to traditional technologies which have fixed path lengths and no meniscus because both events meaningfully affect the intensity of the observed CD signal.

In this note, we present CD measurements of (α)-lactoglobulin, Cytochrome C, Hemoglobin and Lysozyme with variable well volumes and concentrations to address the sensitivity of the EKKO™ CD Microplate Reader for CD measurements.

II – RESULTS & DISCUSSION

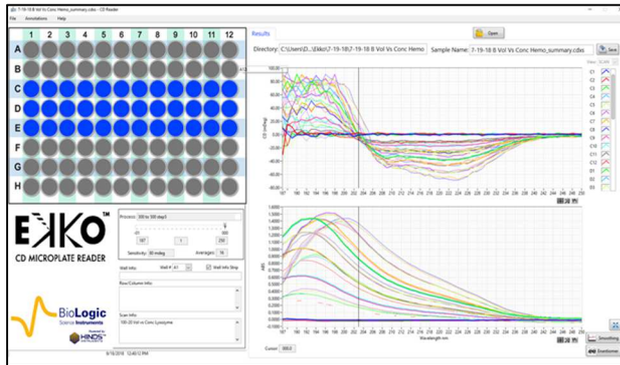


Fig. 2. CD spectra Matrix of Hemoglobin with increasing concentrations and volumes. 100, 150, or 200 μ l of Hemoglobin (Sigma) at 20 to 100 μ g/ml was in wells #C1 to #E12 of a solid fused silica 96 well plate (Hellma). No effort to reduce the noise was used to collect the raw data.

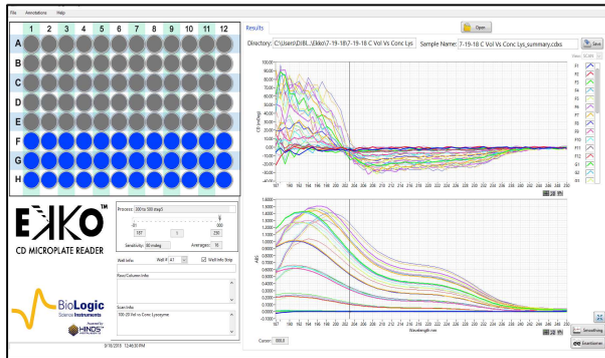


Fig. 3. CD spectra Matrix of Lysozyme with increasing concentrations and volumes. 100, 150, or 200 μ l of Lysozyme (Sigma) at 20 to 100 μ g/ml was in wells #F1 to #H12 of a solid fused silica 96 well plate (Hellma). No effort to reduce the noise was used to collect the raw data.

Figures 2 and 3 are representative matrices of the CD and Absorbance data obtained for increasing concentrations of protein at loads of 100, 150 & 200 μ l's per well. As expected, increasing concentrations of protein distorted the CD spectra at the higher ends of the matrices regardless of volume added within the well when compared to known spectral characteristics of the proteins involved. Similar results were obtained with α -Lactoglobulin and Cytochrome C.

To determine where the deviation from linearity in both the CD and Absorbance in the measurements occurred, Absorbance and CD were plotted as a function of increasing concentration at multiple wavelengths for each of the matrices.

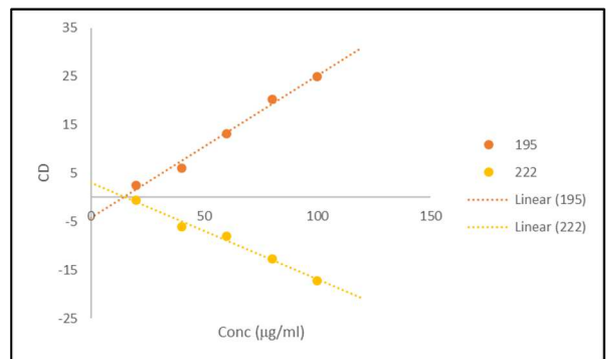
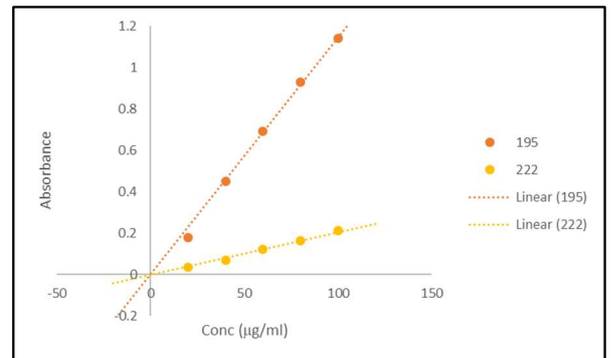
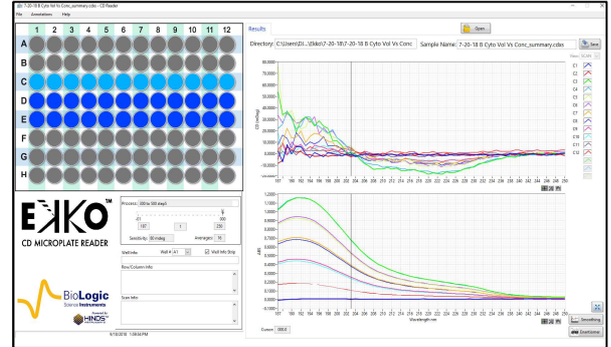


Fig. 4. Signal versus Concentration for Cytochrome C Matrix. 100 μ l of Cytochrome C (Sigma) at 20 to 100 μ g/ml was in wells #C1 to #C12 of a solid fused silica 96 well plate (Hellma). No effort to reduce the noise was used to collect the raw data (top). Replots of the Absorbance 100 μ l of Cytochrome C as a function of concentration for 20 to 100 μ g/ml at 195 and 222 nm (middle). Replots of the CD 100 μ l of Cytochrome C as a function of concentration for 20 to 100 μ g/ml at 195 and 222 nm (bottom).

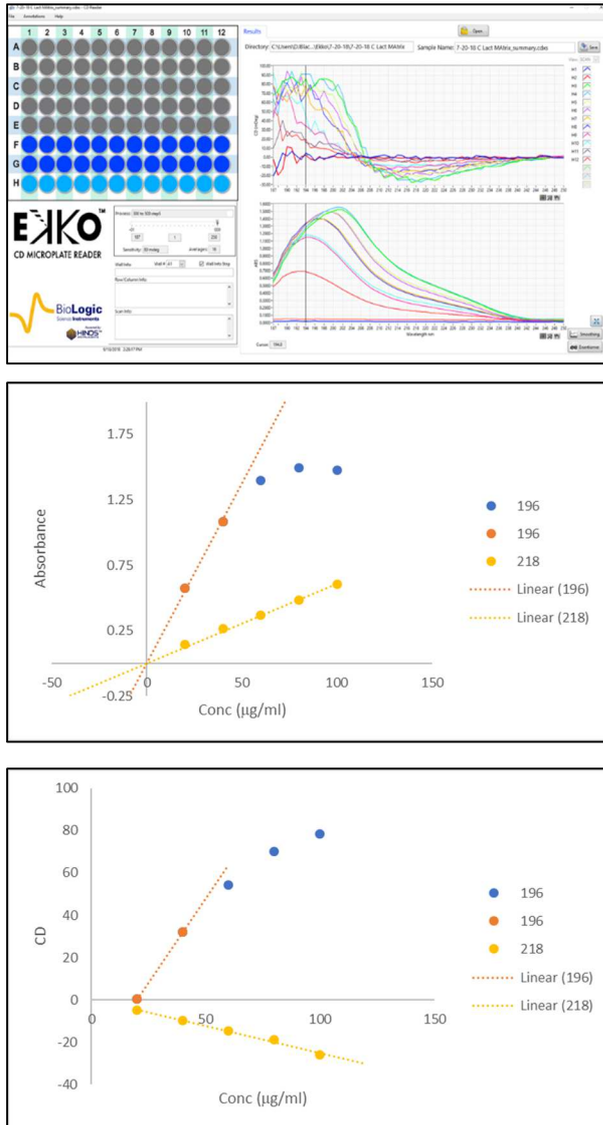


Fig. 5. Signal versus Concentration for α -lactoglobulin Matrix. 200 μ l of α -lactoglobulin (Sigma) at 20 to 100 μ g/ml was in wells #H1 to #H12 of a solid fused silica 96 well plate (Hellma). No effort to reduce the noise was used to collect the raw data (top). Replots of the Absorbance 100 μ l of α -lactoglobulin as a function of concentration for 20 to 100 μ g/ml at 196 and 218 nm (middle). Replots of the CD 100 μ l of α -lactoglobulin as a function of concentration for 20 to 100 μ g/ml at 196 and 218 nm (bottom).

Figures 4 and 5 are representative replots of the signal's verses concentration for the various volumes included in the protein matrices. They illustrate that the signal intensities for the differing secondary structure composition of the protein will have

a profound effect on the usable concentration for any CD measurement as expected. Furthermore, they demonstrate that signals are usable up to an absorbance value of approximately 1.2. The mean value for the last usable absorbance point before deviation from linearity in both absorbance and CD is 1.2 ± 0.1 Abs units. This equates to an approximate concentration of 50 to 100 μ g/ml per mm path length (approximated as the volume in a column for any given addition) for a fully helical protein.

As with conventional technologies for the measurement of circular dichroism, any sample which has an absorbance value below ~ 1.2 will result in a reliable measurement and be within the linear range of the EKKO™ CD Microplate Reader.

III – SUMMARY & RECOMENDATIONS

1. The EKKO™ CD Microplate Reader is linear for the determinations of absorbance and CD up to a value of ~ 1.2 equating to a protein concentration of 50 to 100 μ g/ml for a helical protein.
2. The EKKO™ CD Microplate Reader has similar detection limits as standard CD Technologies.
3. It is recommended that the precise concentration and volume needed be empirically determined under normal operations given the potential meniscus effects resulting from the physical characteristics of the buffer components.

REFERENCES

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- 2) Jo, H.H., Cao, X. You, L., Anslyn, E.V., and Krische, M.J., Application of high-throughput enantiomeric excess optical assay involving a dynamic covalent assembly: parallel asymmetric allylation and ee sensing of homoallylic alcohols. *Chem. Science*. 6, 6747-6753 (2015).
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- 4) Hussain, R., Javorfi, T., Rudd, T. R., and Siligardi, G., High-throughput SRCD using multi-well plates and its applications. *Nature, Sci Reports* 6, 38028 (2016).