

Using ULS24 CMOS Bio-imager as a Readout Sensor for Chemiluminescence Immunoassay and DNA Hybridization Assay

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Introduction

Immunoassay is a widely used method for detecting the presence and quantifying a macromolecule in a solution. This is accomplished through the use of an antibody molecule. The macromolecule, or often referred as "analyte", that is detected by the immunoassay is most often a protein molecule. Analytes in biological liquids such as serum or urine are frequently measured using immunoassays for medical and research purposes. One of the most popular and effective immunoassay methods is the enzymelinked immunosorbent assays, or ELISA

Chemiluminescent immunoassays are variations of the standard ELISA. An enzyme converts a substrate to a reaction product that emits photons of light instead of developing a visible color.

Chemiluminescence is light produced by a chemical reaction. The chemiluminescent substance is excited by the oxidation and catalysis forming intermediates. When the excited intermediates return back to their stable ground state, a photon is released, which is detected by the luminescent signal instrument.

Chemiluminescent assays, in particular certain enhanced chemiluminescent assays, are very sensitive and have a wide dynamic range. It is believed that chemiluminescence is the most sensitive detection method currently in use due to the ability of signal multiplication and amplification. Chemiluminescent reactions are measured in relative light units (RLU) that are typically proportionate to the amount of analyte present in a sample.

In order to measure light emission from chemiluminescence, photon multiplier tube (PMT) devices are typically employed in the read-out instrument. PMTs are very sensitive light detection devices.

Anitoa Systems, LLC., 1717 Embarcadero Rd., Palo Alto, CA 94303, USA (contact: info@anitoa.com)



However, PMT, along with the necessary electronic circuits to make PMT work, is very bulky and expensive. As such, chemiluminescence read-out instruments are also bulky and expensive.

Here we introduce the use of highly integrated and ultra-low-light sensitive CMOS bio-imager sensor chip to detect chemiluminescence emission from chemiluminescence immunoassay. CMOS bio-imager device described here is a true image sensor. It can potentially be used to detect the emission of chemiluminescence emission from multiple reaction sites at the same time.



Materials and method

Chemiluminescence immunoassay material: Akaline Phosphadase (AP) (part number: ALPI12G), from BBI Solutions (info@bbisolutions.com).

Luminol Substrate (part number: SR2001), Jiangsu ZECEN Biotechnology Co., Ltd, PR. China.

Plate: White opaque 96 well microtiter plate, from Thermo Scientific.

Sensor: Anitoa ULS24 Ultra-low-light CMOS bio-imager. ULS24 Solution Kit 1-Channel System.

Lens: a F1.0, 16mm CS mount lens (part number: GMF1610CIR). The aperture of the lens is also 16mm. Supplier: Yian Optics Co. Ltd, PR. China (http://www.yagx.net).

Software and settings: Anitoa ULVision software. The integration time is set to 5s to 60s. Resolution is set to 12x12. Binning mode pattern is 0xF (all 4 sensors within a big pixel are turned on).

Ambient temperature: 15-25 degree C.

Below picture shows the setup of the experiment. The CMOS sensor is mounted above the plate and it images the reaction tube from top. The enzyme mix, when activated, emits blue light centered at ~460nm wavelength. We do not use any filters for this setup.

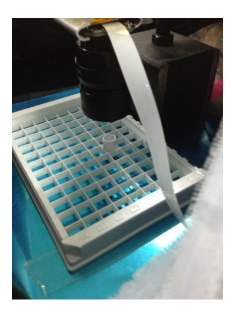


Figure 1, Chemiluminescence measurement setup



Procedure

First of all, the stock AP enzyme master mix is diluted 2000 times. This is done by taking 0.5ul of 100ug/mL master mix; add it to 1000ul Tris Buffer (ph8.0) in a reaction tube. The tube is vortexes well to ensure a good mix. Then the tube is allowed to set for 15min. We then take 200ul diluted mix and put into 800ul Tris Buffer (pH8.0), shake to mix, to further achieve another 5x dilution.

Use same method above to create 5 different concentration of reaction mixture: 0.05, 0.01, 0.002, 0.0004, 0.00008ug/mL).

After all set up is complete, fetch 1ul of enzyme mix, add to tubes that contain 50ul of luminal substrate. Start with lower concentration samples first and finish with the highest concentration sample. This is because the light attenuation is faster with higher concentration samples.

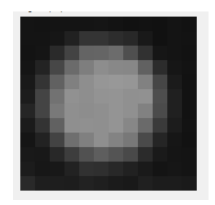
Figure 2 is a sample image taken by the sensor from the top of the test tube. As can be seen, the whole tube is imaged in the 12x12 pixel array area. To calculate the read out, we just simply add the output from all 144 pixels.

We also image the tube with luminol substrate, but with no enzyme added. This serves as negative control. When we calculate the light output from the sample images, we always substract negative control image from the image taken from the sample.



Results

Below is the read out light level from samples of different enzyme concentration:



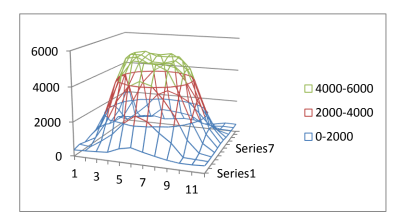


Figure 2, Sample image and its 3D histogram, from the image sensor that is performing the read out

Enzyme concentration µg/mL	Light signal output
0.05	1611321.5
0.01	306904.0
0.002	66950.4
0.0004	27796.9
0.00008	8958.3



Table 1, Chemiluminescence light readout vs. enzyme concentration

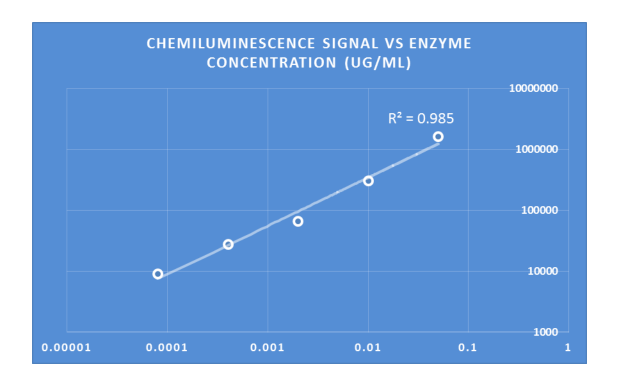


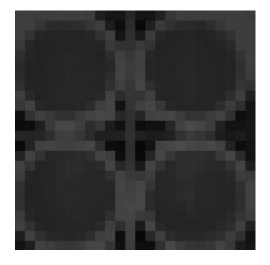
Chart 1, Chemiluminescence light readout vs. enzyme concentration in logarithm scale, fitted with a linear interpolation curve.



Test with multiple samples

With some small adjustment of the optics setting, namely object distance and lens focus, we were able to test multiple samples with one single image shot. Below are the procedures and results of this test.

First we imaged 4 empty test tube in ambient light with 10-20ms integration time. We did both 24x24 and 12x12 shots, while adjust the lens focus.



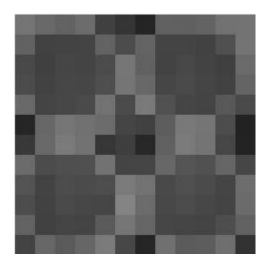


Figure 3a Image of 4 empty test tube in ambient light, 24x24 resolution mode

Figure 3b Image of 4 empty test tube in ambient light, 24x24 resolution mode

As can be seen in the screenshot, even in 12x12 mode, we can reliably distinguish the 4 test tubes.

We then placed the system in dark enclosure and imaged the tubes with longer integration time. We added ALP enzyme with luminol substrate in the up-left corner. The other 3 tunes contain only luminol substrate, but no enzyme. We performs a series of test and the result is shown below.



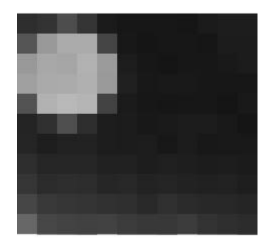


Figure 4. Screen shot of the image with upper left corner test tube filled with ALP enzyme and luminol substrate, while the other 3 tubes contain only luminol substrate, with no enzyme. The enzyme concentration from this screen shot is 0.002ug/ml.

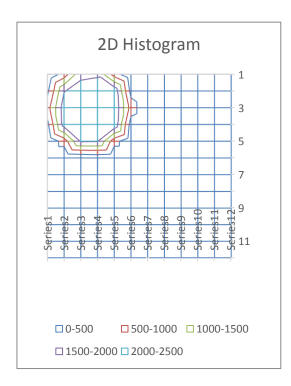


Figure 5a 2D histogram of the test described above

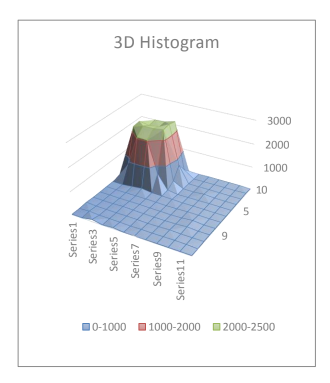


Figure 5b 3D histogram of the test described above



Note that when the histogram is produced, we performed dark subtraction to reduce background noise.

We performed the 3rd, 4th and 5th latter concentrations (0.00008ug/ml, 0.0004ug/ml, and 0.002ug/ml)

Enzyme concentration (µg/mL)	Chemiluminescence Readout
0.002	54244.93
0.0004	19897.19
0.00008	4181.16
0	81.23

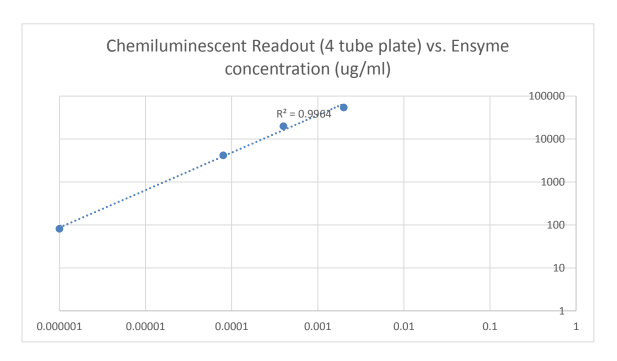


Chart 2, Chemiluminescence readout vs. enzyme concentration in logarithm scale, in a 4 tube plate format

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Directly couple Anitoa CMOS Bio-optical sensor to the assay

It is also possible to directly couple Anitoa CMOS Bio-optical sensor to the chmeluminescence assay. This method has the highest light collection efficacy and the advantage of simplicity. Below is an example of ULS24 interfacing with a microfluidic chip.

With this method, it is possible to detect ng/ML level of chemiluminescence analyte with shorter integration time, for example 1-5 second of integration time.



Figure 7 ULS24 sensor directly coupled to a microfluidic chip running beads based chemiluminescence immunoassay (Image provided by Micronit BV).

In a separate experiment (performed also in collaboration with Micronit BV), the ULS24 CMOS biosensor is used in conjunction with a microfluidic chip in a lens-less configuration (Figure 8a,b) for beads-based DNA hybridization. The capture DNA probes are attached to magnetic beads, and the detection probes were coupled with HRP enzyme. As the target DNA in a sample (e.g. PCR product) hybridize with the probes, they are trapped at the detection point in the microfluidic chip due to a small magnet.



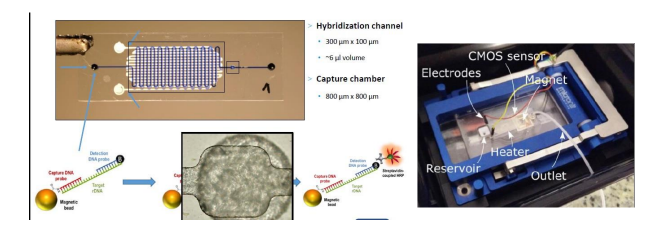


Figure 8a, b Microfluidic and beads based DNA hybridization set up, where the ULS24 CMOS biosensor is tightly coupled to the microfluidic chip in a lens-less configuration to detect chemiluminescence

To detect the hybridized target DNA, a luminol substrate is passed through the channel. As the HRP-conjugated probes react with the substrate, chemiluminescence emission is generated and detected (Figure 8c).

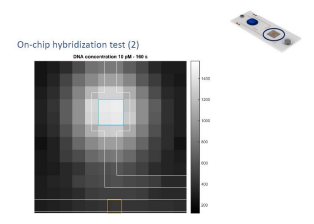


Figure 8c Chemiluminescence image captured by ULS24 CMOS Biosensor