

Application Guide for Multimode Readers.



TECHNOLOGY ORIENTATION

Cell-based assays	4
ABS - Absorbance	16
FI - Fluorescence Intensity	24
TRF - Time-Resolved Fluorescence	32
FRET - Fluorescence Resonance Energy Transfer	33
TR-FRET - Time-Resolved Fluorescence Resonance Energy Transfer	34
FP - Fluorescence Polarization	46
Alpha - Amplified Luminescent Proximity Homogeneous Assay	52
LUMI-Luminescence	56



Cell-based assays.

Imaging-based analysis of cell samples using Spark® and Spark Cyto

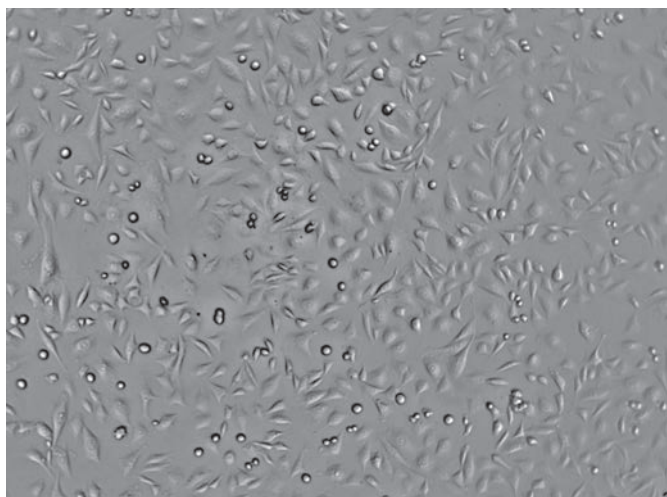


Figure 1A: Bright field cell imaging of a CHO cell with digital zoom applied, using the Spark.

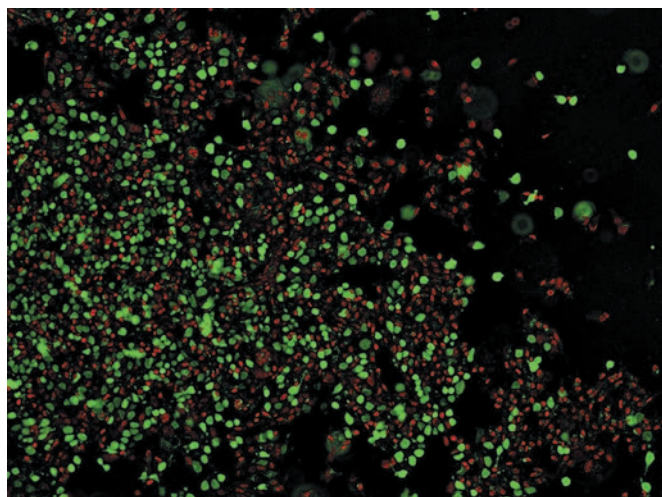


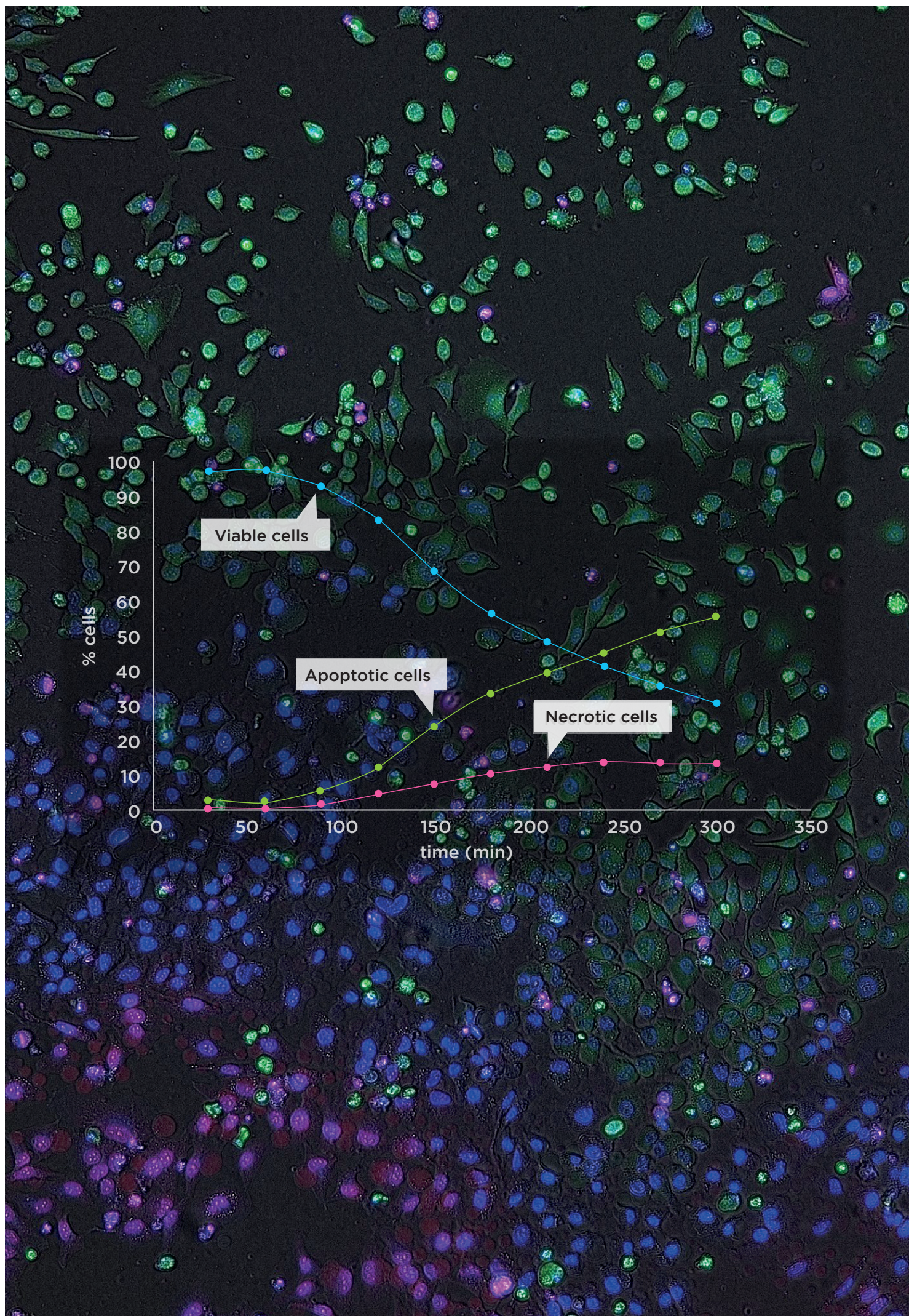
Figure 1B: Fluorescence cell imaging with real-time cytometry using the Spark Cyto; 4x image of HeLa cells stained with Calcein/PI (cell viability application).

TECHNOLOGY

Cell-based assays in microplate formats have become a vital part of life science experiments, with cell growth, proliferation, migration and cytotoxicity being central research objectives. However, confluence levels, gene expression activity and the mode of cell death are also important parameters that are routinely analyzed in cell culture laboratories. In this context, imaging-based readouts offer valuable insights and provide the basis for reproducible analysis and accurate interpretation of experimental data. Automating and standardizing these processes streamlines cell-based workflows and helps to free up time in the lab.

Tecan's Spark multimode reader has a cost-effective, built-in cell imaging module, enabling automated cell counting and viability analysis in slides (Tecan's cell chips), as well as bright field imaging and automated confluence assessment in microplates (see Figure 1A). Spark's imaging module includes LED bright field illumination, 4x objective magnification with an optical resolution of $\geq 3 \mu\text{m}$, a 1.3-megapixel CMOS camera and a robust autofocus system. It offers full environmental control, including temperature, CO_2 and O_2 , as well as evaporation protection if needed. Injectors with reagent heaters and stirrers, and integrated microplate lid-handling are also available. With these features, Spark delivers better quality and more physiologically-relevant data for your cell-based research, and opens the door to the exciting world of live cell imaging.

The Spark Cyto is the first multi-mode plate reader with live cell imaging and real-time cytometry capabilities. It contains the same detection modes, features and functions as the Spark, but is also equipped with a high performance bright field and fluorescence imaging module (see Figure 1B), unlocking new possibilities for your cell-based research. Spark Cyto uniquely brings together top-of-the-range camera components with proprietary patent-pending technology to ensure that you can truly investigate your entire cell population. It gives you the ability to record the whole well area of a 96- or 384-well microplate with just one image, without tiling or distortion. This means that you never miss a cell when investigating the total population in a microplate well. Spark Cyto includes three magnification levels, combined with four channels for fluorescence and bright field imaging, enabling high quality cell analysis for a wide variety of applications. The system is controlled by the versatile and user-friendly SparkControl™ software, which includes a number of predefined application protocols – as well as user-defined capabilities – offering both ease of use and flexibility for operators. Images acquired with the Spark Cyto can be automatically processed with Image Analyzer, Tecan's proprietary imaging software package. Image Analyzer offers you an array of customization options, making it easy to adjust and optimize your imaging parameters.



**Spark****Spark Cyto****Reader****Features**

Imaging modes	Bright field	Bright field, Digital phase Fluorescence
Sample formats (for imaging)	6 to 96-well	6 to 384-well
Camera sensor	CMOS, 1.3 megapixel	CMOS, 5.0 megapixel
Objectives	4x	2x, 4x, 10x
Max. optical resolution	3.0 μ m	1.2 μ m
Channels	Bright field	Bright field Four fluorescence channels (blue, green, red, far-red)
Autofocus	LED	Astigmatism-based LED
Field of view	Center User-defined Whole-well by tiling	Center User-defined Whole-well by tiling Whole -well, 96- and 384-well with a single image
Image Analysis	Real-time analysis; speed: <45 min for 96-well (whole-well by tiling) + confluence assessment	Real-time analysis; speed: <20 min for 96-well (whole-well by single image) + confluence assessment
Re-analysis capability	N/A.	Image Analyzer software

Application

Counting and viability in cell chips	•	
Confluence	•	•
Nuclei counting		•
Transfection efficiency		•
Cell viability		•
Cell death		•
User-defined		•
REC™	•	•

SUPPORT**Links**

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry>
- <http://lifesciences.tecan.com/live-cell-imaging>
- <http://lifesciences.tecan.com/cellbiology>

Cell counting and viability of cells in suspension

Determine the number and health of your cells using Tecan Cell Chips™

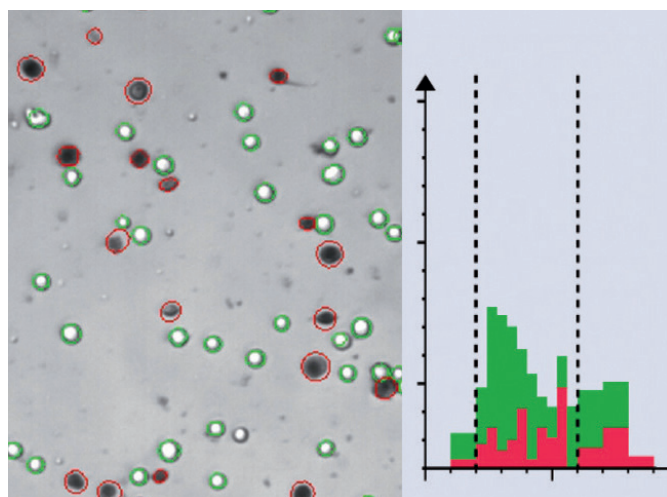


Figure 1A: Discrimination of live/dead cells in suspension using Spark.



Figure 1B: Cell Chips on adapter plate.

Technology

Cell-based assays in a microplate format have become a vital part of life science research; proliferation, cytotoxicity and gene expression studies all inevitably use cells as a working tool. Regardless of the research area, cell counting and quality checks are commonly performed before cell seeding or passaging steps. One approach is to identify if cells are alive or dead by staining with trypan blue; only dead cells or those with a damaged cell membrane will be stained blue, enabling them to be discriminated from viable, healthy cells (Figure 1A).

The Spark's cell imaging module offers automated, label-free cell counting and trypan blue-based cell viability analysis with an easy-to-use, disposable Cell Chip (Figure 1B). Each Cell Chip has two sample chambers that can accommodate independent cell suspensions or replicates of the same sample. Four Cell Chips can be loaded onto the Cell Chip adapter plate, allowing up to eight samples to be analyzed in less than two minutes.

The cell counting and cell viability applications both offer the same features, with the difference that the viability application automatically takes into account a 1:1 trypan blue dilution when calculating the results, showing the number and percentage of viable cells. Users can set the cell size to between 4 and 90 μm before starting the measurement, allowing you to tailor the assay to

your chosen cell line. For a better counting accuracy and reproducibility, the system also allows you to select one, four or eight images per chamber. This enables fast cell counting for applications with relatively high cell concentrations ($\sim 1 \times 10^6$ cells/ml) – by taking a single image at the center of each chamber. In contrast, for low cell concentrations ($\sim 1 \times 10^4$ cells/ml), eight separate images can be used per sample to dramatically improve the accuracy of cell concentration measurements. Furthermore, the two chambers of each slide can be defined as 'duplicates' for automatic evaluation. Cell counting and cell viability results are clearly presented in a table and histogram on the software dashboard, allowing straightforward size distribution analysis of the cell population. All results are then saved in a PDF report for easy and convenient data storage.

MAJOR APPLICATIONS

- Cell culture quality control
- Cell seeding
- Cell proliferation studies

SUPPORT

Link

<https://lifesciences.tecan.com/live-cell-imaging?p=tab--2>

Cell Confluence

Check the growth status of your cells

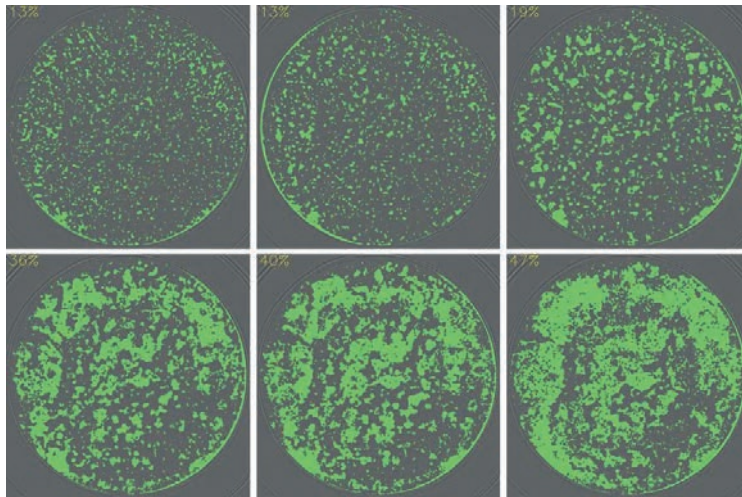


Figure 1A: Confluence assessment in the Spark: Continuous growth monitoring with the cell confluence applications (96-well images; tiled)

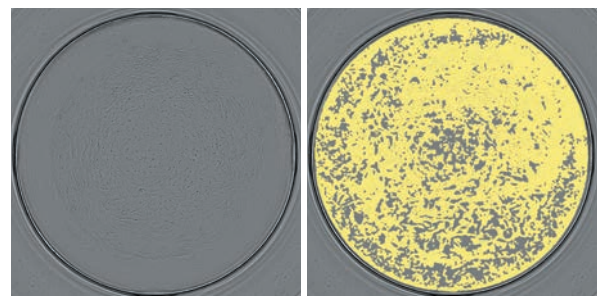


Figure 1B: Confluence in the Spark Cyto: 96-well plate imaged with unique whole-well imaging using the 2x objective and analyzed with the confluence application. Processed bright field image (left) and applied confluence mask depicted (right). No tiling or edge-to-edge optical distortion leads to superior results when analyzing cell populations in terms of confluence.

Technology

In cell culture applications, confluence assessment is an important quality control parameter. It is commonly used to estimate the proportion of adherent cells on a growth surface, as an indicator of the cell density inside a well or culture flask. In addition, the growth rate or gene expression of some cell lines varies depending on the degree of confluence. Consistent determination of cell confluence and estimation of cell numbers are therefore important for reproducible assays and accurate interpretation of experimental data. Image capture and confluence determination for adherent cells directly in microplates improves crucial scientific parameters, like assay variation and cell seeding uniformity.

Spark's cell imaging module allows you to precisely determine the confluence level in a well from 10 to 90 %, giving you a new level of data quality for your assays. The easy-to-use SparkControl software allows you to select different imaging areas within the well. Depending on the assay, you can either select just a single image of the well center or a full-well picture composed of

multiple images, with automatic well border detection to compensate for variations in microplate dimensions. Confluence is assessed in real time, with data displayed on the dashboard and exported directly to Excel® for further analysis. Each confluence value is linked directly to the raw and analyzed images, enabling quick and easy visual confirmation of cell confluence.

Confluence determination is completely independent of cell morphology, using an algorithm designed to offer exceptional sensitivity, even when working with low contrast cells, such as fibroblasts. This makes Spark's confluence measurement compatible with virtually any cell type, offering an easy-to-use approach to improve your assays uniformity. Label-free confluence assessment can also be used as a replacement for costly and time-consuming cell mass analyses like the quantification of protein levels or ATP content. It also enables applications like cell migration, wound healing and clonogenicity assays.

Spark Cyto allows you to image a whole-well in a 96- or 384-well plate with one single picture using a proprietary patent-pending wide field of view technology. This gives you a complete picture of your cell population in less time, allowing you to drive your research in new directions. Re-analysis and algorithm adjustment capabilities allow the user to adapt the sensitivity of the confluence assessment according to the experimental conditions. A new and extremely useful feature of the Spark Cyto is the calculation of the roughness factor within the confluence application. This is a qualitative assessment of the cells' granularity, and is expressed as the normalized mean standard deviation of pixel intensities over all separated areas. It can be used as an indicator of the relative signal variation across the cell population.

MAJOR APPLICATIONS

- Cell culture quality control
- Cell migration and wound healing studies
- Cell proliferation studies
- Cytotoxicity studies
- Single cell clone selection

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <https://lifesciences.tecan.com/live-cell-imaging?p=tab--3>

Nuclei counting

Accurately determine the number of cells in a microplate well

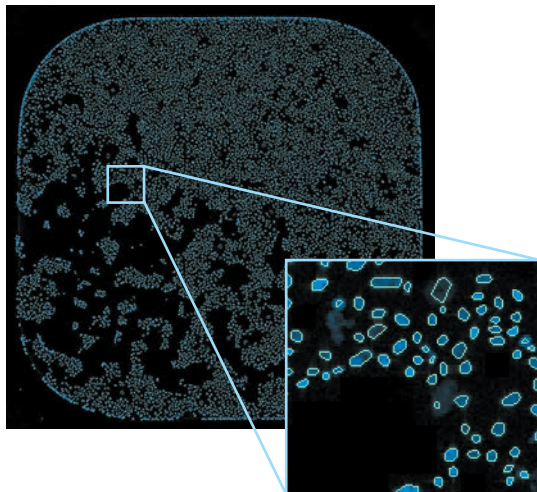


Figure 1: Whole-well image of cells stained with Hoechst 33342, using a nuclei counting mask .

Technology

Cell counting is used as a widespread measurement in tissue culture to assess density for cell-based assays, as a readout in proliferation, viability and toxicity assays, and to normalize cell-based assay results. Cell counting and viability assessment of suspension cells in slides is available with the Spark, but Spark Cyto offers a pre-defined application to image adherent cells cultured in a monolayer, using fluorometric probes that intercalate with nuclear DNA. Optimized for Hoechst 33342, this function provides an easy method for cell counting using any blue fluorescent dye with nuclear DNA binding capabilities. The ability to perform whole-well imaging of 96- or 384-well plates with one single picture ensures accurate cell counting and always reveals the full picture. Images and results can even be re-analyzed and optimized using the Image Analyzer.

MAJOR APPLICATIONS

- Cell culture quality control
- Cell proliferation studies
- Cytotoxicity studies

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <https://lifesciences.tecan.com/live-cell-imaging?p=tab--3>

Transfection efficiency

Transfection efficiency checks and quantification of protein expression in microplates

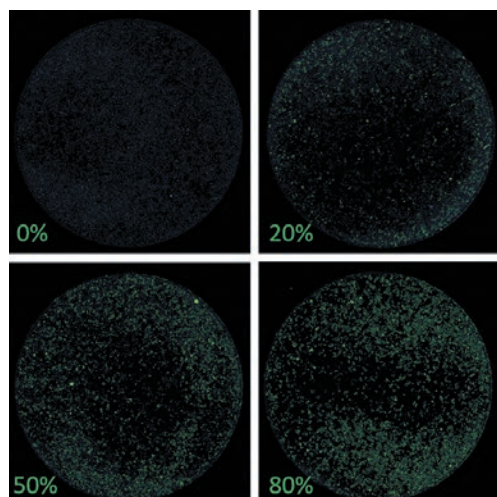


Figure 1: Representative overlays of various transfection rates imaged and calculated using the Spark Cyto.

Technology

Transfection and transduction are indispensable tools for studying biological functions and processes. Assessing transfection/transduction efficiency of cells in a qualitative and quantitative manner is therefore a frequent and important for many laboratories to ensure that cell samples are suitable for downstream applications.

Transfection efficiency is usually assessed with the aid of fluorescent labels, such as green fluorescent protein (GFP), either manually by fluorescence microscopy or using single-cell analysis techniques such as flow cytometry. However, this requires detachment of the cells from the plates. Live cell imaging permits the analysis of the cell expression *in situ*. The transfection efficiency application of the Spark Cyto is optimized for GFP-transfected cells, counter-stained with a blue nuclear stain, such as Hoechst 33342 or DAPI (4',6-diamidino-2-phenylindole).

MAJOR APPLICATIONS

- GFP transfection and transduction efficiency
- GFP reporter assays

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <http://lifesciences.tecan.com/cellbiology>

Cell viability

Quantification of live and dead cells in microplates

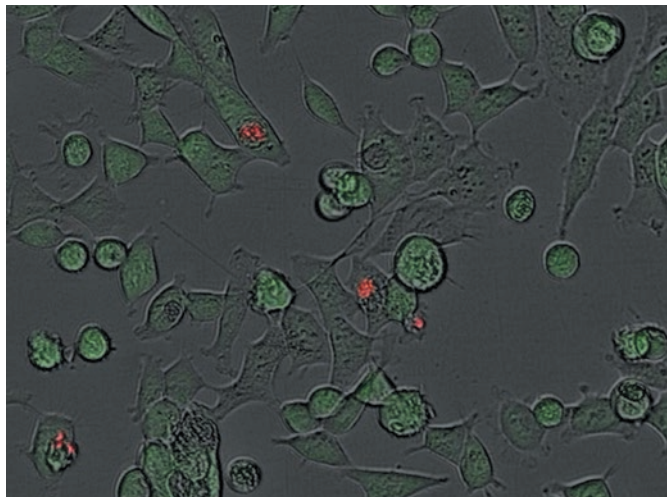


Figure 1: HeLa cells cultured in a 24-well plate, acquired with the 10x objective, showing an overlay of the bright field, green and red channels (digital zoom applied).

Technology

Viability assays are commonly used in cell biology labs as a screening tool to assess cell line response to compounds of interest or different growth conditions. There are numerous methods for determining cell viability, some providing 'well' information – eg. bioluminescent detection of ATP, or calorimetric assays using MTT, WST-1 or resazurin – while others use fluorescence-based imaging to discriminate between live and dead cells on an individual level. These fluorescent viability markers are usually combined with appropriate cell death markers, based on membrane permeability and accessibility of the nuclei.

Spark Cyto's preset cell viability application uses a common double staining approach to discriminate between live (green) and dead (red) cells in a population. Using two fluorescent dyes, such as Calcein-AM (live cells) and propidium iodide (dead cells), you can image and analyze your population in minutes. Calcein-AM is a non-fluorescent, cell-permeable derivative of calcein that becomes fluorescent upon hydrolysis within the cytosol. Propidium iodide (PI) is a fluorescent intercalating agent that can be used to stain cells that binds to DNA with little or no sequence preference. It is not membrane-permeable, making it useful to differentiate necrotic, apoptotic and healthy cells based on membrane integrity.

MAJOR APPLICATIONS

- Viability assessments
- Cytotoxicity studies

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <http://lifesciences.tecan.com/cellbiology>

Cell death

Detecting phosphatidylserine externalization by adherent cells in microplates

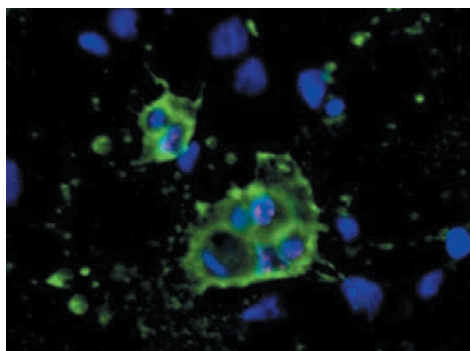


Figure 1: Apoptotic A431 cell imaged with the 10x objective and digitally zoomed-in.

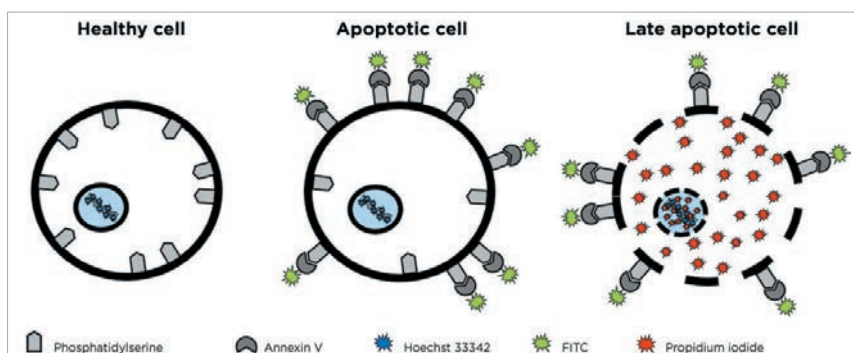


Figure 2: Schematic overview of fluorescence profiles of viable, apoptotic and necrotic cells stained with Hoechst 33342, annexin V-FITC and PI.

Technology

Apoptosis – or programmed cell death – is a central component of many biological processes, including development, tissue homeostasis, and maturity and maintenance of the immune system. The identification and discrimination of apoptotic, necrotic and viable cells is therefore important for a variety of different biological research fields.

The detection of, and discrimination between, apoptosis and necrosis can be accomplished by differential staining of markers characteristic of the relevant type of cell death, for example, with Hoechst 33342, PI and annexin V-FITC. This approach is implemented in Spark Control as the predefined cell death application.

In healthy cells, phosphatidylserine (PS) is located on the inner leaf of the plasma membrane. As one of the first steps in apoptosis, PS is externalized, which is a signal for the cells to be phagocytosed. Once present on the outer leaf, it can be detected by the calcium dependent protein annexin V conjugated to a fluorescent label, such as fluorescein isothiocyanate (FITC) or Alexa Fluor® 488. Adding a second, cell-impermeable dye, such as PI, permits the discrimination of necrotic cells, because only these cells have permeabilized outer membranes, making them accessible to the dye. Staining with PI and FITC/Alexa Fluor 488 is a standard procedure to discriminate

apoptosis and necrosis in cell cultures; early apoptotic cells are annexin V positive and PI negative, whereas late apoptotic cells are annexin V/PI double positive, as shown in Figure 2. The total number of cells can then be identified by staining with a nuclear dye, such as Hoechst 33342 or DAPI.

MAJOR APPLICATIONS

- Cell death discrimination
- Apoptosis quantification

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <http://lifesciences.tecan.com/cellbiology>

Real Time Experimental Control (REC)

Never miss a critical biological event!

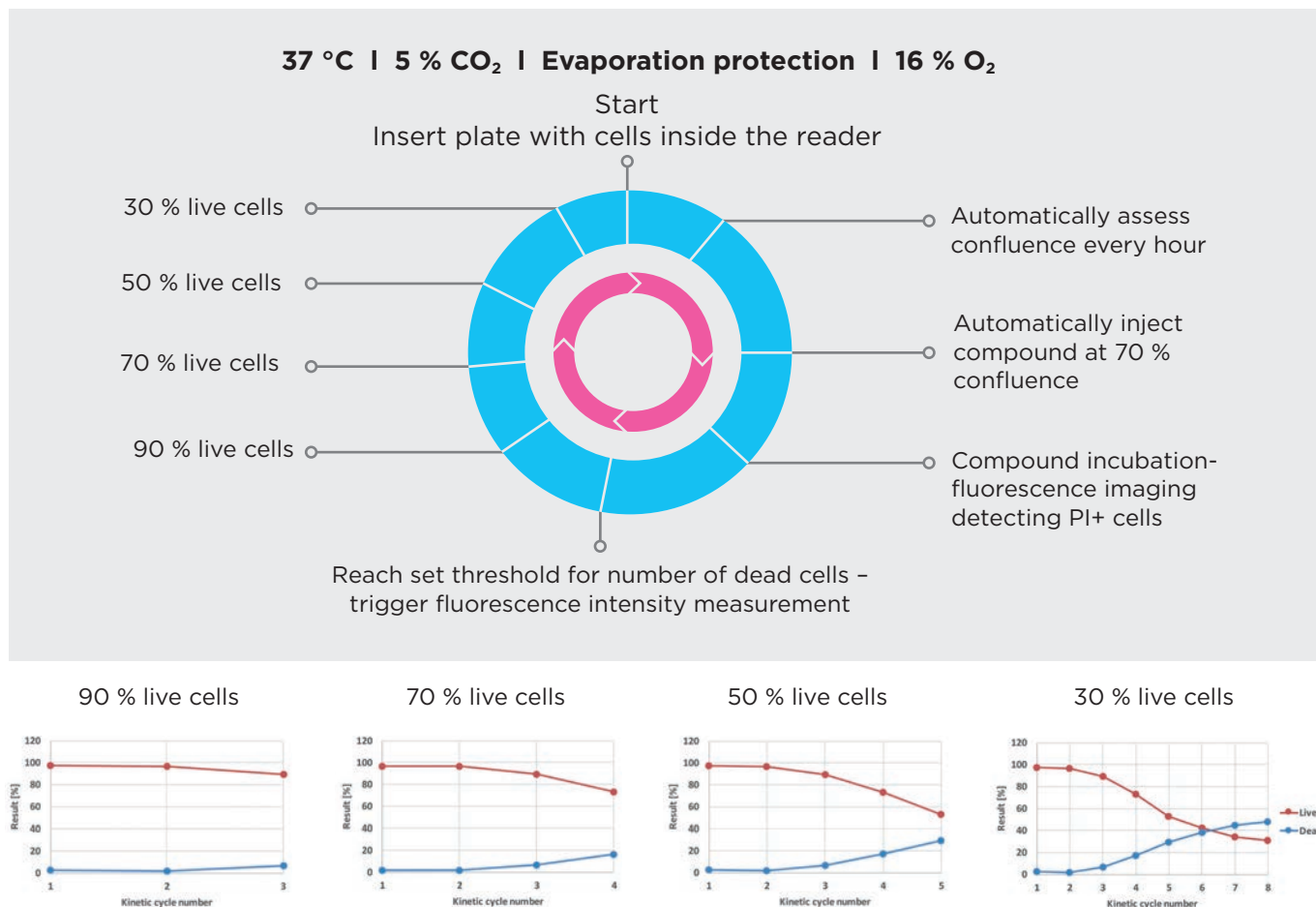


Figure 1: Schematic overview of an exemplified REC case.

Technology

Many live cell kinetic experiments require specific actions at different points in the experiment – for example, the addition of a compound once a certain confluence is reached. Real Time Experimental Control (REC) allows certain actions – such as injection of reagents – to be triggered automatically as part of a kinetic experiment, and offers real-time data and image analysis for complete confidence in your results. REC grants you the ability to create new experimental workflows in your lab. Combining standard detection technologies, imaging capabilities and additional unique features, such as integrated humidity and environmental controls, REC unlocks new research possibilities.

REC uses all of these features to create workflows that ensure you never miss a critical biological event – without chaining you to the bench.

MAJOR APPLICATION

Cell-based applications in general

SUPPORT

Links

- <https://lifesciences.tecan.com/plate-reader-live-cell-imaging-cytometry?p=tab--3>
- <http://lifesciences.tecan.com/cellbiology>



ABS – Absorbance.

Light is absorbed by the sample

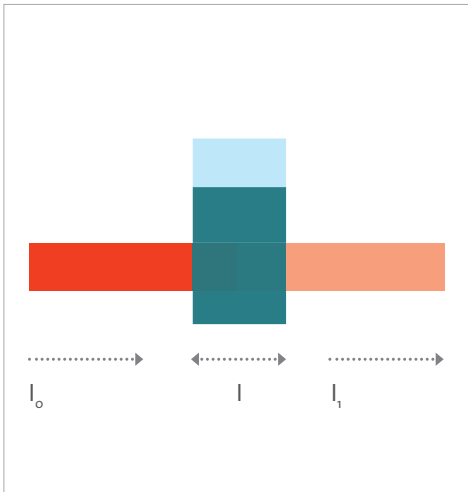


Figure 1: Schematic representation of an absorbance measurement performed in a cuvette.

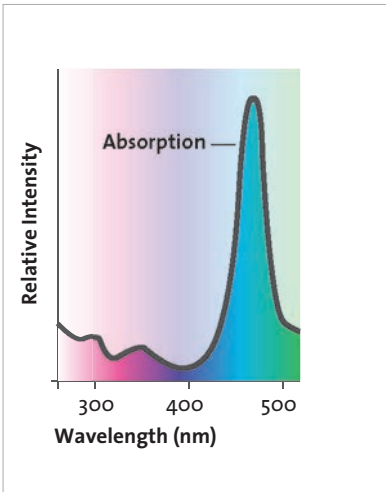


Figure 2: An absorbance spectrum shows the extent of light absorption at any specific wavelength.

Absorbance OD	Transmittance [%]
0	100
1	10
2	1
3	0.1
4	0.01

Table 1: The relationship between absorbance and transmittance values. An absorbance value of 3 means that only 0.1 % of the light is able to pass through the sample. Most multimode readers can only read samples up to an OD of 4.

TECHNOLOGY

When light shines through a turbid or colored liquid, some of its intensity is absorbed by the liquid's molecules or particles (Figure 1). The amount of light that penetrates the sample and reaches the detector is called the transmittance (T), and the light absorbed by the sample is called the absorbance (A, Abs), or optical density (OD)¹.

OD values commonly correspond to a 1 cm path length, which is the width of standard cuvettes (Figure 1). Note that absorbance is a logarithmic function (Table 1) of the transmittance, as defined by the Lambert Beer law²:

$$A = -\log_{10}(I_t/I_o)$$

The absorbance spectrum is a function of the molecule; light of different wavelengths can be absorbed differently depending on the sample, as shown in the absorbance spectrum in Figure 2.

For this reason, the absorbance is always stated together with the wavelength, for example OD₆₀₀.

Major applications

- DNA / RNA quantification (Abs)
- MTT / MTS assays
- BCA, Modified Lowry and Bradford assays - Protein quantification

¹ Bioanalytik. Von F. Lottspeich. Spektrum, Heidelberg, 1998
² 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.



DNA / RNA quantification (Abs)

DNA / RNA quantification based on absorbance



Figure 2: Tecan NanoQuant Plate™

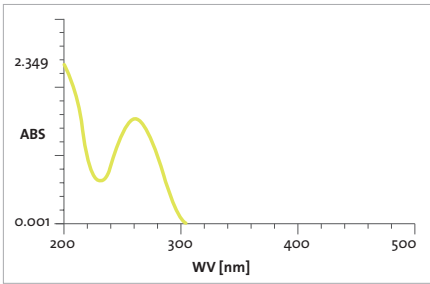


Figure 1: DNA absorbance spectrum.

1OD ₂₆₀	Concentration (µg/ml)
dsDNA	50
ssDNA	33
RNA	40
Oligonucleotides	20-30

Table 1: Nucleic acid concentrations at OD₂₆₀ = 1 and neutral pH

ASSAY OVERVIEW

Technology

Absorbance

DNA and RNA can be quantified based on absorbance at 260 nm, which is in the UV range and not visible to the human eye (Figure 1). Tecan’s multimode readers provide cuvette ports for DNA and RNA measurement (quartz or UV-transparent cuvettes must be used). Alternatively, Tecan’s patented NanoQuant Plate™ (Figure 2) is ideal for smaller volumes (2 µl), higher throughput (16 samples at once), and more economical DNA / RNA quantification. 1 OD₂₆₀ corresponds to different concentrations, depending on the type of nucleic acid being quantified (Table 1).

The ratio of absorption at 260 nm vs 280 nm is commonly used to assess DNA contamination of protein solutions, since proteins – in particular, the aromatic amino acids – absorb light at 280 nm^{3,4}. It is generally acknowledged that pure DNA has a ratio of 2, and RNA, 1.8⁵. Recently researchers found that the A₂₆₀/A₂₃₀ ratio also provides valuable information about the nucleic acid purity⁶.

Typical detection limits for absorbance based DNA quantification are within the single digit ng/µl range.

Alternative

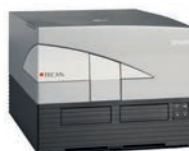
LifeTechnologies’ PicoGreen® and RiboGreen® quantification assays provide a broader dynamic range.

Sample protocol for DNA measurements

- 1. Blank the reader with the same buffer used to dilute the DNA
- 2. Take an appropriate volume of sample (NanoQuant Plate: 2µl, cuvettes: volume depends on the min. / max. filling volume)
- 3. Measure OD₂₆₀ and OD₂₈₀
- 4. If the OD₂₆₀ value is greater than two, dilute samples
- 5. To calculate the concentration, multiply the OD₂₆₀ by the concentration factor shown in Table 2 and your dilution factor, if applicable
- 6. Determine the purity by dividing the value for OD₂₆₀ by the value for OD₂₈₀

³ a b c d e Sambrook and Russell (2001). Molecular Cloning: A Laboratory Manual (3rd ed.). Cold Spring Harbor Laboratory Press, ISBN 978-0-87969-577-4.
⁴ (Sambrook and Russell cites the original paper: Warburg, O. and Christian W. (1942). "Isolierung und Kristallisation des Gärungsferments Enolase". Biochem. Z. 310: 384-421.)
⁵ Glasel, J.A. (1995) Validity of Nucleic Acid Purities Monitored by A260/A280 Absorbance Ratios, Biotechniques 18:62-63
⁶ <http://www.qiagen.com/literature/benchguide/default.aspx>

Instrument parameters



Reader	Infinite® 200 PRO	Spark	Spark Cyto
Assay	DNA concentration	DNA concentration	DNA concentration
Measurement Mode	Absorbance	Absorbance	Absorbance
Wavelength	260 nm	260 nm	260 nm
Bandwidth	5 nm	default	default
Flashes	25	25	25
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

MTT⁷/MTS⁸ assay

Cell viability / cytotoxicity assays with absorbance readout

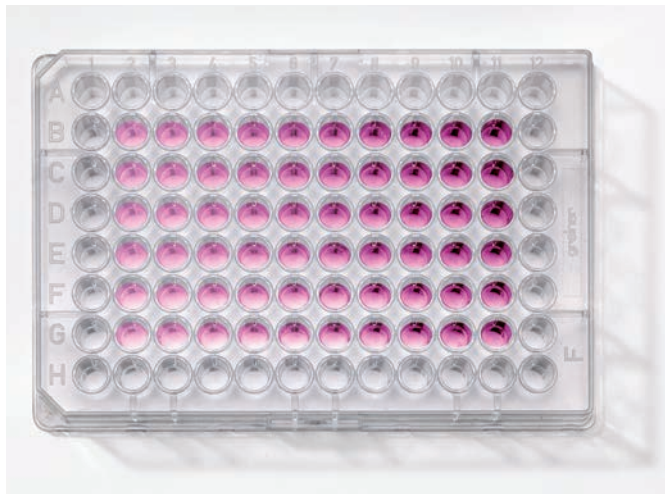


Figure 1: MTT assay in a 96-well format

ASSAY OVERVIEW

Technology

Absorbance

Principle, Major application

The MTT and MTS assays are colorimetric (Figure 1) – and hence absorbance – assays that measure cell viability. MTT / MTS is taken up by the cells and processed to varying extents depending on the cell viability. Viable cells reduce more MTS / MTT to formazan, yielding a more intense purple color.

Mechanism

While MTT assays need to use a reagent to make the formazan generated soluble, MTS assays yield water-soluble products, potentially making them homogeneous assays.

MTT assays use a solubilization reagent, such as dimethyl sulfoxide (DMSO) or isopropanol, to dissolve the non-water-soluble formazan product, yielding a colored solution that can be quantified by absorbance measurement at approximately 565 nm, dependent on the solvent employed⁹.

MTS assays are an improved version of the MTT assay. Its reagents are reduced more efficiently within the cell than MTT, and the resulting product is water-soluble and less cytotoxic than the insoluble formazan used in the MTT assay. This makes it a one-step (homogeneous) assay, with the convenience of adding the reagent directly to the cell culture without the intermittent steps required in the MTT assay.

However, when MTS is used in a homogeneous way the assay becomes susceptible to colorimetric interference, as traces of colored compounds may remain in the microplate¹⁰.

Provider

MTT, MTS and similar reagents are available from various chemistry distributors, or as the CellTiter Glo® 2.0 Assay (Promega)¹¹.

Alternatives

An alternative to absorbance-based cell viability assays, are fluorescence-based systems using the redox indicator Resazurin to detect the cell's metabolic activity. These assays are characterized by a more convenient assay handling and higher sensitivity levels.

⁷ MTT (3-(4,5-Dimethylthiazol-2-yl)-2,5-phenylditetrazolium bromide, a yellow tetrazole)

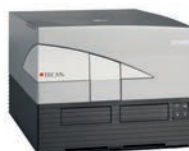
⁸ MTS (3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium)

⁹ Mosmann T (1983). Rapid colorimetric assay for cellular growth and survival: application to proliferation and cytotoxicity assays. *Journal of immunological methods* 65 (1-2): 55-63.

¹⁰ Cory AH, Owen TC, Barltrop JA, Cory JG (1991). Use of an aqueous soluble tetrazolium/formazan assay for cell growth assays in culture. *Cancer communications* 3 (7): 207-12.

¹¹ https://www.promega.com/products/cell-health-assays/cell-viability-and-cytotoxicity-assays/celltiter_glo-2_0-assay/?catNum=G9241

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	MTT assay/CellTiter Assay	MTT assay/CellTiter Assay	MTT assay/CellTiter Assay
Measurement Mode	Absorbance	Absorbance	Absorbance
Wavelength	565 nm	565 nm	565 nm
Bandwidth	9 nm	3.5 nm	3.5 nm
Flashes	25	10	10
Settle time	0 ms	50 ms	50 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

BCA, Modified Lowry and Bradford assays – Protein quantification

Protein quantification assays with absorbance readout



Figure 1: Bradford Protein Assay measured in cuvettes, showing increasing protein concentrations.

ASSAY OVERVIEW

Technology

Absorbance

Principle, Major application

All three assays are designed to determine the protein concentration of a sample. For detection, a liquid reagent needs to be added to the samples. This reagent interacts with the proteins, leading to a visible color change (Figure 1) that is directly proportional to the concentration. Absolute concentrations are calculated using a standard curve.

Provider

Various companies have established their own assays for this purpose. The main differences between the various assays are the dynamic range and the measurement wavelength.

Mechanism

The **BCA™** Protein Assay (Thermo Scientific Pierce) uses bicinchoninic acid (BCA) for colorimetric quantification of total protein in a sample¹³. The method is based on the reduction of Cu^{2+} to Cu^+ by protein in an alkaline

medium to form a colored watersoluble chelate that can be measured at its absorption maximum of 562 nm. The linear working range for BSA is 20 to 2000 $\mu\text{g/ml}$ ¹⁴.

The **Bradford** Protein Assay (BioRad) is based on the Coomassie® Brilliant Blue G-250 dye which binds to basic and aromatic amino acid residues, particularly arginine. This induces a shift of the dye's absorbance maximum from 465 nm to 595 nm. The Bradford assay can be performed as a microassay procedure, with a linearity range of 125 to 1,000 $\mu\text{g/ml}$ BSA¹⁵.

In the **Modified Lowry** Protein Assay (Thermo Scientific Pierce), the protein reacts with cupric sulfate and tartrate in an alkaline solution, which results in formation of a tetradentate copper-protein complex, reducing the Folin-Ciocalteu Reagent. The absorbance of the blue, water-soluble product can be measured at 750 nm. The assay – tested with BSA protein¹⁶ – exhibits good linearity in the range of 1 to 1500 $\mu\text{g/ml}$.

Alternatives

Potential alternatives for protein quantification reach from absorbance-based methods using the protein extinction coefficient¹⁷ to fluorescence based assays like NanoOrange® to even dedicated¹⁸ devices.

¹³ Smith, P.K., et al.: Measurement of protein using bicinchoninic acid. Anal Biochem., 150, 76-85, 1985

¹⁴ <https://www.thermofisher.com/us/en/home/life-science/protein-biology/protein-assays-analysis/protein-assays.html>

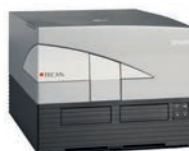
¹⁵ <http://www.bio-rad.com/webroot/web/pdf/lsr/literature/4110065A.pdf>

¹⁶ <https://www.thermofisher.com/us/en/home/life-science/protein-biology/protein-assays-analysis/protein-assays.html>

¹⁷ <http://web.expasy.org/protparam/protparam-doc.html>

¹⁸ <http://www.millipore.com/techpublications/tech1/an2222en>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	BCA Assay	BCA Assay	BCA Assay
Measurement Mode	Absorbance	Absorbance	Absorbance
Wavelength	565 nm	562 nm	562 nm
Bandwidth	9 nm	default	default
Flashes	25	25	25
Settle time	0 ms	0 ms	0 ms

Assay	Modified Lowry assay	Modified Lowry assay	Modified Lowry assay
Measurement Mode	Absorbance	Absorbance	Absorbance
Wavelength	750 nm	750 nm	750 nm
Bandwidth	9 nm	default	default
Flashes	25	25	25
Settle time	0 ms	0 ms	0 ms

Assay	Bradford assay	Bradford assay	Bradford assay
Measurement Mode	Absorbance	Absorbance	Absorbance
Wavelength	595 nm	595 nm	595 nm
Bandwidth	9 nm	default	default
Flashes	25	25	25
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

FI – Fluorescence Intensity.

Light is absorbed and released (emitted)

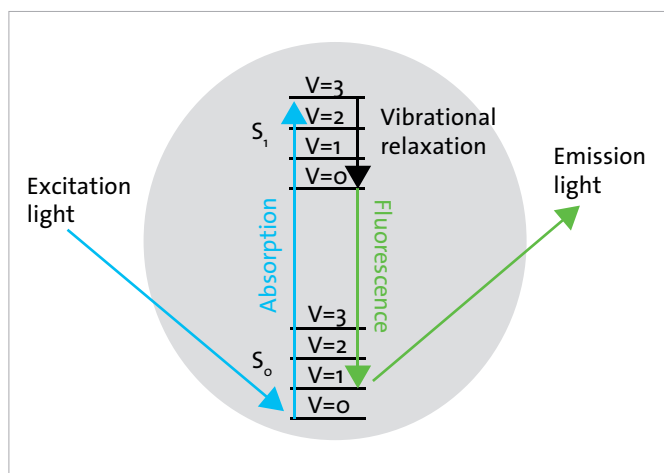


Figure 1: Jablonski diagram. S = electronic state, V = vibrational level. After photon absorption (= excitation), the molecule adopts a state of higher energy S_1 (= excited state) including several vibrationally excited substates. By vibrational relaxation, the molecule relaxes to the lowest excited S_1 state (black arrow). From this state the molecule relaxes into the vibrational states of S_0 by emitting light.

TECHNOLOGY

Fluorescence describes a molecule's ability to emit (release) previously absorbed light (Figure 1). The emission occurs almost instantly (within 1 ns = nano second) and, according to the laws of physics, the emitted light will always have a higher wavelength and hence a lower energy. A fluorescence spectrum consists out of an absorption (excitation) and emission spectrum (Figure 2).

Fluorescence labels (fluorophores) can be attached to any available biomolecule and used to answer quantitative, as well as qualitative, questions. For example, 'does the sample contain the fluorophore?' (qualitative), and 'how much of the fluorophore is in the sample?' (quantitative). Signals are quantified as Relative Fluorescence Units [RFU].

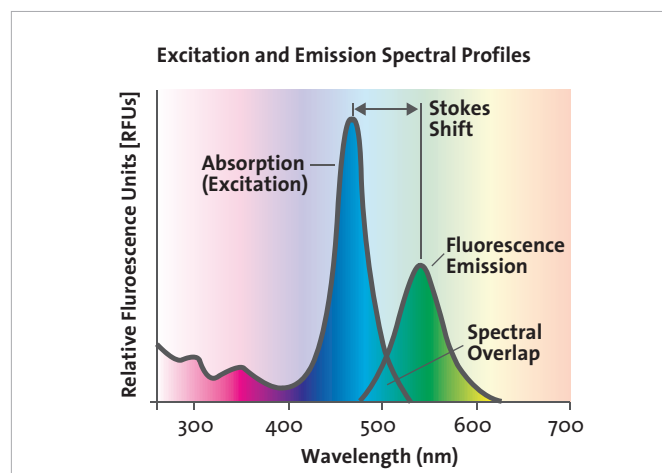


Figure 2: While the excitation spectrum describes how efficient it is to excite the fluorophore at a specific wavelength, the emission spectrum describes how efficient it is to detect the emitted light at any given wavelength. The Stokes shift describes the distance between the excitation and emission maximum, and is given in nanometers (nm).

Major applications

- PicoGreen® and RiboGreen® DNA / RNA quantification
- Resazurin-based cell viability assays
- Fluorescent proteins (GFP, RFP, YFP, etc.)
- ORAC (Oxygen Radical Absorbance Capacity) assay



PicoGreen® and RiboGreen® DNA / RNA quantification



High sensitivity, fluorescence-based DNA / RNA quantification



Figure 1: Green DNA

ASSAY OVERVIEW

Technology

Fluorescence Intensity

Assay design and provider

LifeTechnologies' PicoGreen¹⁹ and RiboGreen²⁰ (Figure 1) quantification assays use a fluorescence approach to determine DNA and RNA concentrations. Using the Quant-iT PicoGreen dsDNA Assay Kit, you can selectively detect as little as 25 pg/ml of dsDNA in the presence of ssDNA, RNA, and free nucleotides.

The assay is linear over three orders of magnitude, and has little sequence dependence, allowing you to accurately measure DNA from many sources.

RiboGreen RNA reagent is one of the most sensitive detection dyes for the quantification of RNA in solution, offering linear fluorescence detection in the range of 1 to 200 ng/ml of RNA.

Mechanism

Both assays are easy to use; simply add the dye to the sample, wait five minutes, and detect the fluorescent signal.

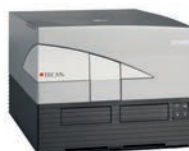
Alternatives

If sensitivity is not a major issue, it may be possible to perform DNA quantification using absorbance at 260 nm.

¹⁹ <http://products.invitrogen.com/ivgn/product/P7589>

²⁰ <http://products.invitrogen.com/ivgn/product/R11490>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	PicoGreen/RiboGreen	PicoGreen/RiboGreen	PicoGreen/RiboGreen
Measurement Mode	FI Top	FI Top	FI Top
Excitation wavelength Donor	485 (9) nm	485 (20) nm	485 (20) nm
Emission wavelength Donor	535 (20) nm	535 (25) nm	535 (25) nm
Lag time	0	0	0
Integration time	20 μ s	default	default
Flashes	25	10	10
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

Resazurin assay

A Fluorescence Intensity-based cell proliferation assay

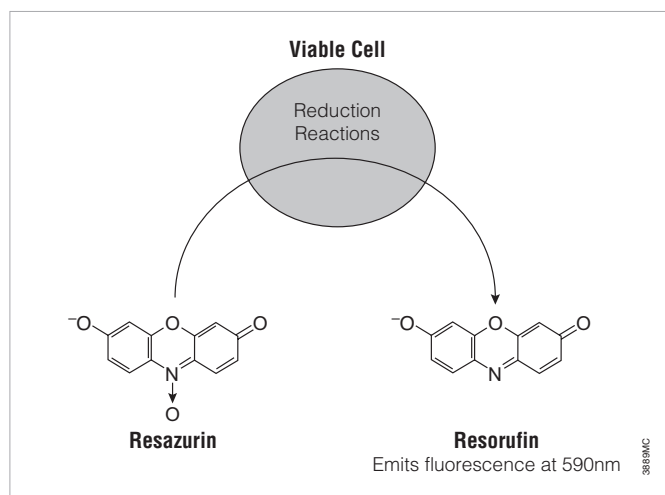


Figure 1: Viability dependent conversion of Resazurin to Resorufin (Promega)

ASSAY OVERVIEW

Technology

Fluorescence Intensity

Major application

Resazurin is a chemical compound that is frequently used for cell viability assays.

Provider

Resazurin was initially used for bacterial studies, but is now also available for eukaryotic cell-based applications under brand names such as the alamarBlue® assay²¹ (Life Technologies) and CellTiter-Blue® Cell Viability Assay²² (Promega).

Mechanism

Resazurin is a redox indicator that can be added directly to cells. Viable cells convert the dark blue, oxidized form of the dye (resazurin) into a red, fluorescent reduced form called resorufin (Ex: 570 nm; Em: 590 nm). The amount of fluorescence or absorbance is proportional to the number of living cells, and corresponds to the cell's metabolic activity. Damaged and non-viable cells have lower innate metabolic activity, and therefore generate a proportionally lower signal than healthy cells. The system is specific for cell viability as non-viable cells rapidly lose metabolic capacity and do not reduce resazurin. Consequently, a fluorescent signal²³ is not generated.

Alternatives

PrestoBlue Cell Viability Reagent²⁴, a new development from Invitrogen that offers much shorter incubation times.

The absorbance based MTT/MTS assay.

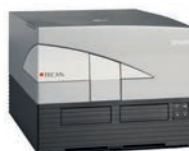
²¹ <http://www.invitrogen.com/site/us/en/home/brands/Molecular-Probes/Key-Molecular-Probes-Products/alamarBlue-Rapid-and-Accurate-Cell-Health-Indicator.html>

²² https://www.promega.com/products/cell-health-assays/cell-viability-and-cytotoxicity-assays/celltiter_blue-cell-viability-assay/?catNum=G8080

²³ O'Brien, J.; Wilson, I.; Orton, T.; Pognan, F. Investigation of the Alamar Blue (resazurin) fluorescent dye for the assessment of mammalian cell cytotoxicity. Eur. J. Biochem. 2000, 267, 5421-5426.

²⁴ <http://www.invitrogen.com/site/us/en/home/brands/Molecular-Probes/Key-Molecular-Probes-Products/PrestoBlue-Cell-Viability-Reagent.html>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	alamarBlue, CellTiter-Blue	alamarBlue, CellTiter-Blue	alamarBlue, CellTiter-Blue
Measurement Mode	FI Bottom	FI Bottom	FI Bottom
Excitation wavelength Donor	560 (9) nm	560 (20) nm	560 (20) nm
Emission wavelength Donor	600 (20) nm	600 (25) nm	600 (25) nm
Lag time	0	0	0
Integration time	20 μ s	default	default
Flashes	25	10	10
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

GFP (Green Fluorescent Protein)

Fluorescent protein frequently used as an expression / activation reporter

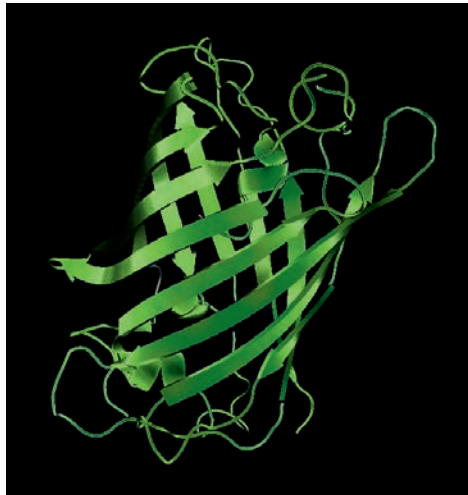


Figure 1: Protein structure of GFP

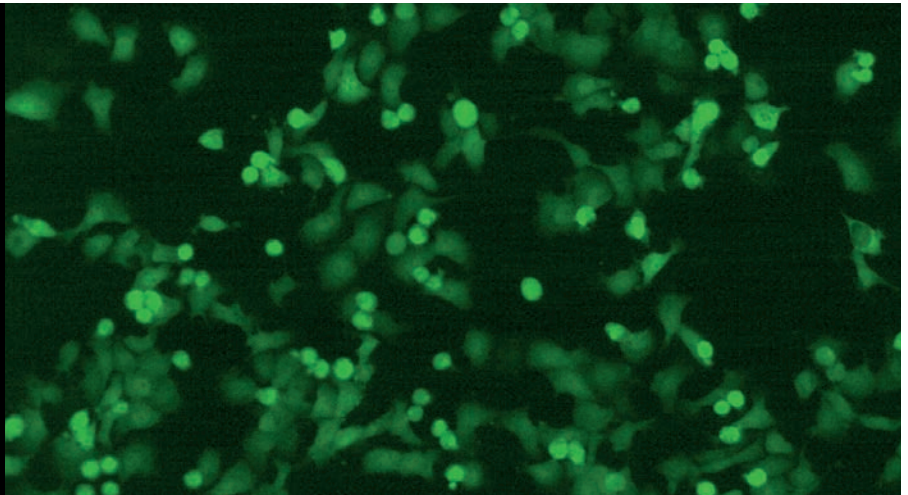


Figure 2: GFP-transfected eukaryotic cells

ASSAY OVERVIEW

Technology

Fluorescence Intensity

Principle

GFP (Figure 1) is a protein derived from a jellyfish which has the ability to fluoresce in the green wavelength range and can be detected using standard FI measurements.

Major applications

GFP can be used in an almost unlimited number of ways, for example as a BRET / FRET partner in binding studies, or for gene activation, where it is often fused / cloned to a gene of interest and co-expressed once the gene is activated (Figure 2). Commonly, it is used to differentiate between constitutive (permanent) and temporary expression. Constitutive expression is mostly used to monitor growth or proliferation of cells or bacteria, while temporary expression is used for gene activation studies.

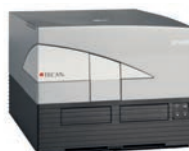
Format, provider

Due to multiple engineering efforts, an almost unlimited number of mutants exist, resulting in a large bandwidth of excitation and emission values. Some of these variants are commercially available, while others are published and therefore not protected. Consequently, only a selection of measurement parameters can be given, since the wavelength depends on the mutant type of the protein.

Alternatives

Technology-wise, there are a lot of alternative fluorescent proteins available, such as CFP (cyan), YFP (yellow) and RFP (red). From an assay perspective, the alternative selected depends on the application. For gene expression studies, DLR® (Dual luciferase reporter assay) or GeneBLAzer® assays may be suitable. For FRET / BRET studies, fluorescent labels might be an alternative.

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	GFP	GFP	GFP
Measurement Mode	FI Bottom	FI Bottom	FI Bottom
Excitation wavelength Donor	485 (9) nm	485(20) nm	485(20) nm
Emission wavelength Donor	535 (20) nm	535(25)	535(25)
Lag time	0 μ s	0 μ s	0 μ s
Integration time	20 μ s	default	default
Flashes	25	15 (5x3, optimal read)	15 (5x3, optimal read)
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

TRF – Time-Resolved Fluorescence.

Light is absorbed and emitted for a relatively long period of time

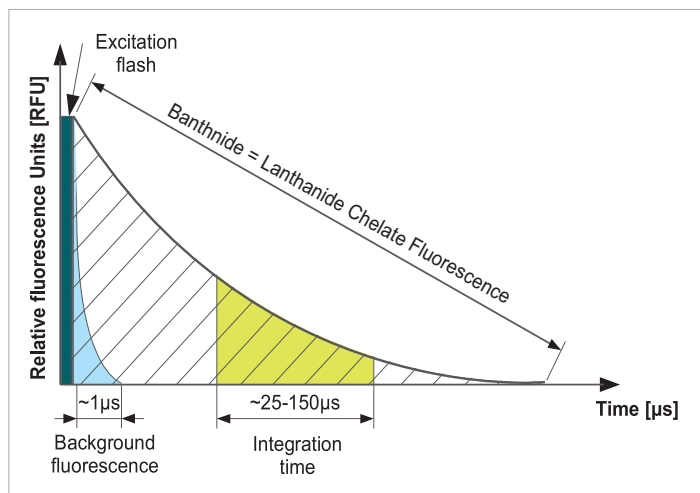


Figure 1: Schematic drawing of a time resolved emission spectrum

TECHNOLOGY

TRF is similar to standard fluorescence, except that the light is emitted for a much longer period of time (Figure 1). The advantage of this is that the signal can be measured once all the background fluorescence (noise) has subsided, increasing the signal to noise ratio, and hence the sensitivity. Only lanthanides – also called rare earth metals – are capable of this kind of fluorescence²⁵.

In most cases, it is possible to substitute fluorescence applications with TRF to achieve higher sensitivity and / or lower background noise.

Major applications

- DELFIA® – Dissociation-Enhanced Lanthanide Fluorescent Immunoassay

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

²⁵ Lakowicz, J. R. (1999). Principles of Fluorescence Spectroscopy. Kluwer Academic / Plenum Publishers

FRET – Fluorescence Resonance Energy Transfer.

Light is absorbed, transferred to another fluorophore and then emitted

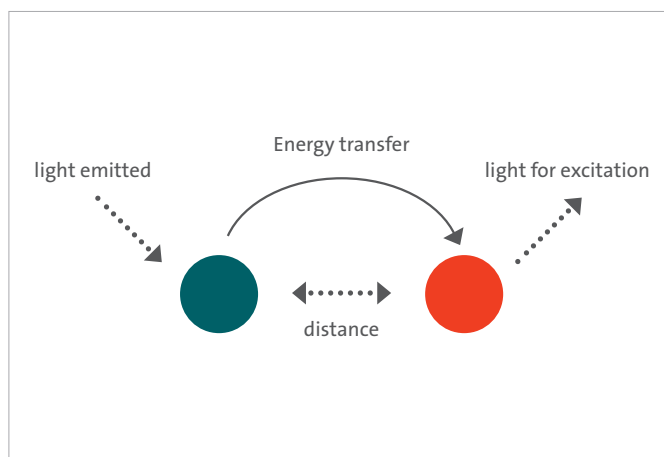


Figure 1: Schematic principle of FRET. Light emitted from the green molecule is used to excite the red molecule.

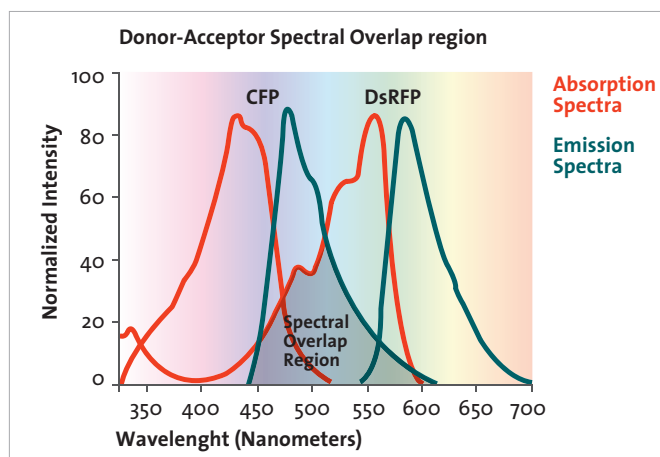


Figure 2: FRET is possible because the emission spectrum of CFP and the excitation spectrum of DsRFP overlap between 450 and 600 nm.

TECHNOLOGY

As the name implies, FRET involves energy transfer between two fluorescent molecules (Figure 1). However, there are some specific requirements for this transfer to take place. Firstly, the emission spectrum of the donor fluorophore and the excitation spectrum of the acceptor fluorophore need to overlap (Figure 2), as the emission light of the donor fluorophore is used to excite the acceptor fluorophore. Secondly, the distance between the two fluorophores – the Förster radius – should be less than 10 nm (Figure 3)²⁸.

One way in which FRET is used is to determine if two biomolecules are in close proximity. In this case, both biomolecules must be labeled with fluorophores and then combined. After an incubation period, the assay is performed. Samples are excited at the donor excitation wavelength, and measured at the donor and acceptor emission wavelength. To compensate for well-to-well variation, for example from pipetting errors, the ratio of both values is calculated (ratiometric assay). If donor and acceptor are in close proximity, FRET will take place, otherwise only the emission signal of the donor is measurable.

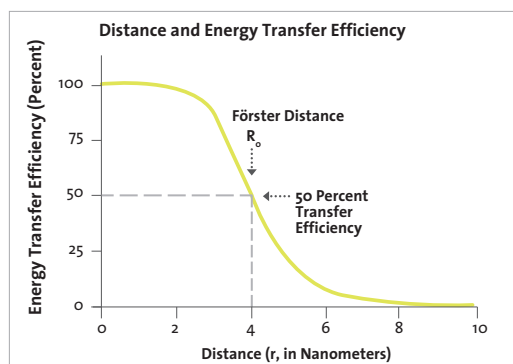


Figure 3: Förster radius – the distance where the FRET signal intensity is reduced to 50 %.

Major applications

- GeneBLAzer and Tango™ GPCR Assay System

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

²⁸ Lakowicz, J. R. (1999). Principles of Fluorescence Spectroscopy. Kluwer Academic / Plenum Publishers

TR-FRET – Time-Resolved Fluorescence Resonance Energy Transfer.

FRET with a longer lifetime and hence a lower background

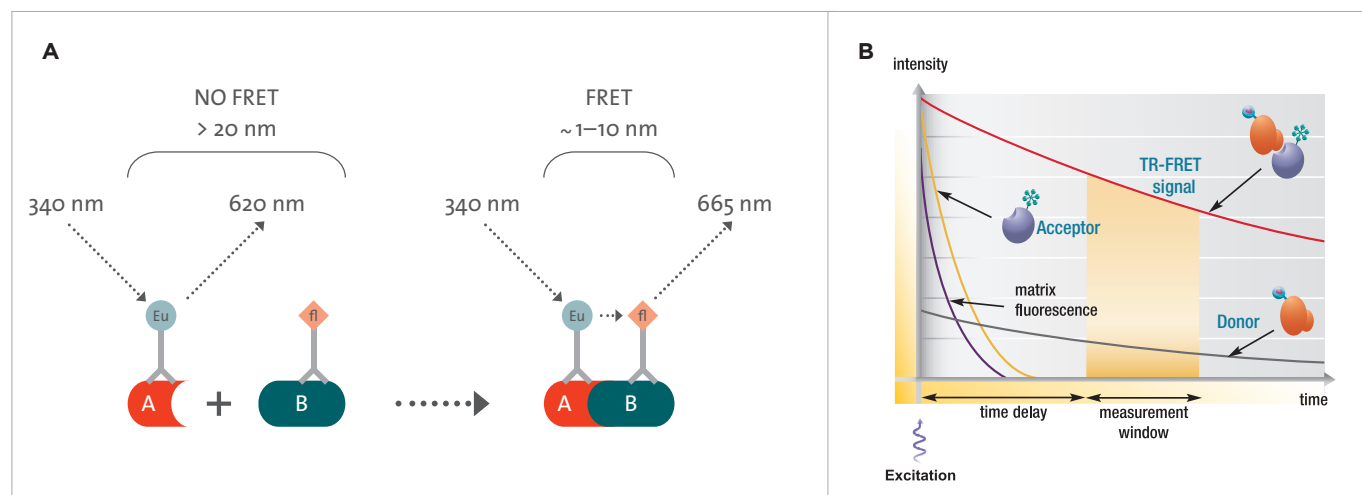


Figure 1: From a technology perspective TR-FRET is equivalent to FRET. In order to reduce the background of the assay, a lanthanide label (e.g. Eu) and a standard fluorescence molecule (fi) as acceptor molecule.

Figure 2: Signal vs. time plot showing the advantages of using lanthanide labels in a FRET process molecule.

TECHNOLOGY

TR-FRET combines the advantages of Time-Resolved Fluorescence (TRF) with the functionality of Fluorescence Resonance Energy Transfer (FRET) (Figure 1). While most standard FRET assays have difficulties with high background resulting from the excitation of matrix molecules. This issue can be resolved by using time-resolved donor molecules. As shown in Figure 2, by the time donor and acceptor emission signals are measured all the unspecific background noise has vanished, giving a high signal to background noise ratio and therefore greater sensitivity.

Assay design

TR-FRET assays are commonly designed to detect whether molecules are in close proximity. This can be exploited to determine if, for example, a protein, peptide substrate, small molecule, phosphorylation, or acetylation is present, or if binding has occurred, for example a receptor-ligand interaction. A major limitation of this technology is that the maximum distance between the donor and the acceptor molecule cannot exceed 10 nm³².

Major applications

- HTRF® – Homogeneous Time-Resolved Fluorescence
- Adapta® Universal Kinase Assay and Substrates
- LanthaScreen™ Kinase Activity Assays
- Transcreener® TR-FRET Assays

³² Lakowicz, J. R. (1999). Principles of Fluorescence Spectroscopy. Kluwer Academic / Plenum Publishers



DELFI[®] - Dissociation-Enhanced Lanthanide Fluorescent Immunoassay²⁶



TRF-based alternative to absorbance-based ELISA

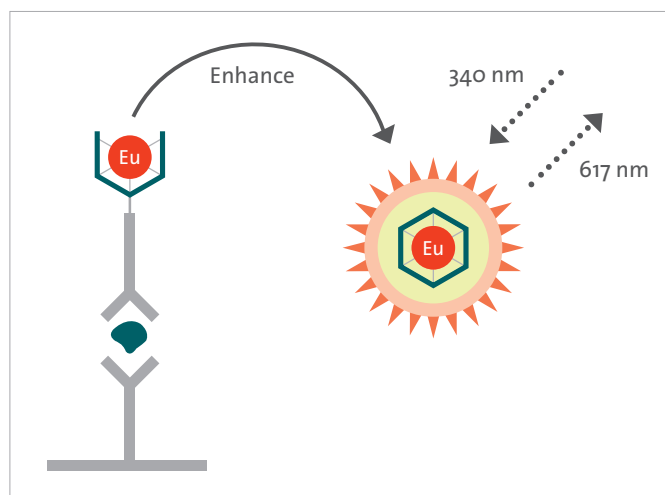


Figure 1: Schematic principle of DELFIA

ASSAY OVERVIEW

Technology

TRF - Time-Resolved Fluorescence

Principle, Provider

PerkinElmer offers the most common lanthanide chelates, including Europium (Eu), Samarium (Sm), Terbium (Tb) and Dysprosium (Dy), under the brand name DELFIA, making DELFIA a technology rather than a single assay.

Major applications

In addition to the self-labeling kits, which allow users to label almost any biomolecule with the lanthanide chelates, PerkinElmer offers pre-coupled antibodies and DNA probes. DELFIA is also available as a ready-to-go assay for cytotoxicity or cell proliferation studies. Other major applications include: receptor-ligand binding, enzyme assays, protein-protein and protein-DNA interaction studies.

Alternatives

As a common application, it is used as an alternative approach to the well-established, absorbance-based ELISA²⁷.

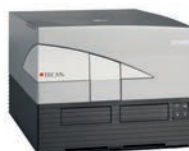
Mechanism

The biomolecule (antibody, DNA probe, etc.) used for detection is labeled with one of the lanthanide chelates. Assays are performed in an endpoint manner and only need to be read once, when all pipetting steps are complete. All steps are performed according to a standard ELISA protocol. Instead of a substrate an enhancement solution is added, that disconnects the chelate lanthanide-chelate complex from the antibody to increase the signal intensity.

²⁶ <http://www.perkinelmer.com/Catalog/Category/ID/delfia%20trf%20assays%20and%20reagents>

²⁷ Enzyme linked Immunosorbent assay

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	Delfia	Delfia	Delfia
Measurement Mode	FI Top (2 labels)	FI Top (2 labels)	FI Top (2 labels)
Excitation wavelength	340(35)	340(35)	340(35)
Emission wavelength	612(10)	612(10)	612(10)
Lag time	200 μ s	100 μ s	100 μ s
Integration time	400 μ s	400 μ s	400 μ s
Flashes	25	150	150
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

GeneBLAzer and Tango™ GPCR²⁹ Assay System³⁰

Gene activator assay with FRET readout

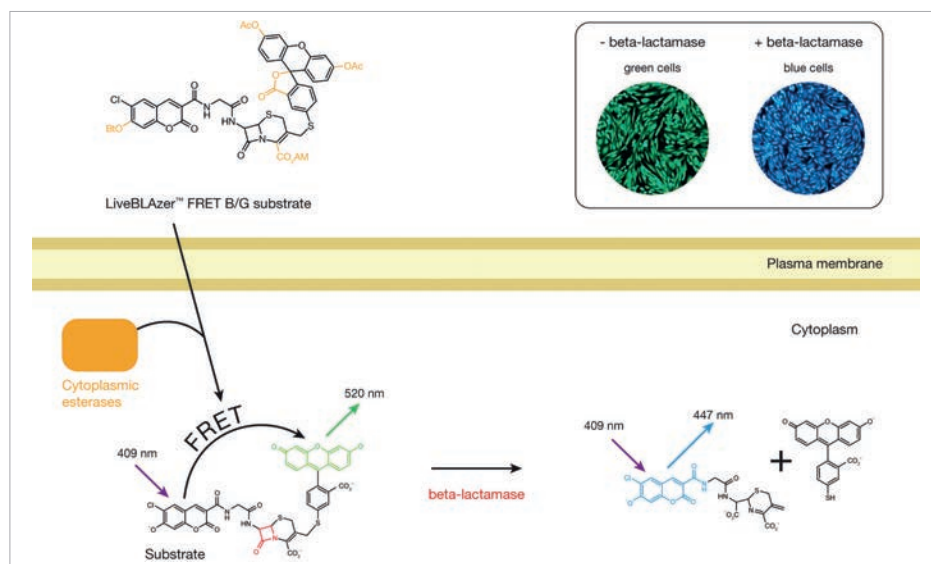


Figure 1: Principle of the GeneBLAzer assay. If the substrate is cleaved, the FRET signal is disrupted.

ASSAY OVERVIEW

Technology

Fluorescence Resonance Energy Transfer (FRET),
ratiometric

Major application, Principle

Life Technologies' GeneBLAzer assays are designed to monitor the activation of genes, including surface and intracellular reporters, a wide range of signal transduction pathways, ion channels and other transporters. The basis for the GeneBLAzer assay are cell lines possessing a β -lactamase³¹ (BLA) gene under the control of a promotor which is downstream of the monitored target protein.

Provider

The β -lactamase-transfected cell lines can either be purchased from Life Technologies or self-transfected. Tango cell lines are also based on the GeneBLAzer technology, but are designed exclusively for GPCR activation assays.

Mechanism

As shown in Figure 1, after the transfection, cells are loaded with an engineered fluorescent substrate which is an assembly of two fluorophores: coumarin and fluorescein. If the target protein is inactive, BLA is not expressed and the substrate molecule remains intact. In this state, excitation of the coumarin results in FRET to the fluorescein moiety and emission of green light.

However, in the presence of BLA expression, the substrate is cleaved, causing the separation of the fluorophores, and FRET cannot occur. This results in the emission of a blue fluorescence signal from coumarin.

Reporter assays are often measured over several hours, or even days. During this time period the plate can either be shuttled between the incubator and the reader, or a temperature and gas controlled multimode reader such as the Infinite 200 PRO or Spark may be used.

Alternatives

LiveBLAzer™, which is a combination of GeneBLAzer and resazurin.

²⁹ G-Protein Coupled Receptors: important group of cell surface receptors for cell signalling

³⁰ http://www.invitrogen.com/site/us/en/home/Products-and-Services/Applications/Drug-Discovery/Target-and-Lead-Identification-and-Validation/g-protein_coupled.html/GPCR-Cell-Based-Assays/GeneBLAzer-Theory.html

³¹ β -lactamase is an enzyme that can cleave specific substrates

Instrument parameters



Reader	Infinite 200 PRO
Assay	GeneBLAzer
Measurement Mode	FI Bottom (2 labels)
Excitation wavelength Donor	415(20)
Emission wavelength Donor	460(20)
Lag time	0 μ s
Integration time	40 μ s
Flashes	25
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms
Excitation wavelength Acceptor	415(20)
Emission wavelength Acceptor	535(25)
Lag time	0 μ s
Integration time	40 μ s
Flashes	25
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

HTRF® – Homogeneous Time-Resolved Fluorescence³³



TR-FRET-based assay platform

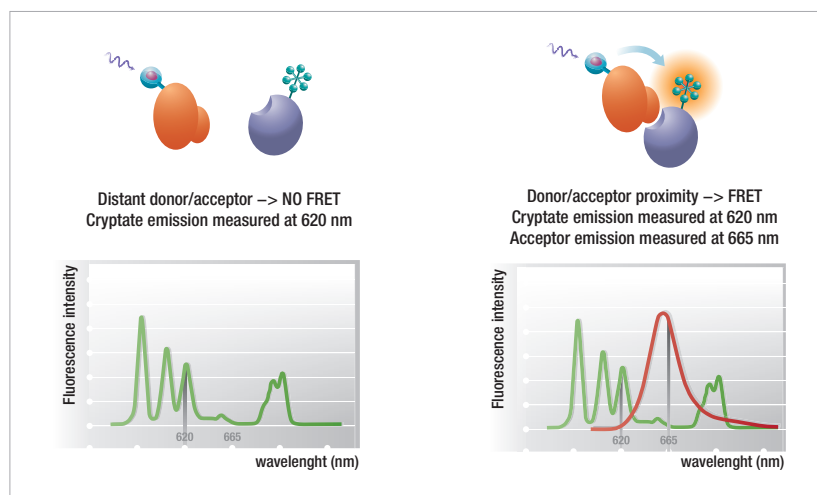


Figure 1: Mechanistic principle of the HTRF technology and fluorescence spectrum

ASSAY OVERVIEW

Technology

TR-FRET – Time-Resolved Fluorescence Resonance Energy Transfer

Provider

HTRF is Cisbio's TR-FRET-based assay platform, which provides a broad range of solutions.

Format, major applications

Biomolecules for detection can either be self-labeled or purchased pre-labeled. Additionally, ready-to-go assays and pre-coupled antibodies are available for major targets, including GPCRs, with second messengers and binding assays, kinases, epigenetic enzymes, protein-protein interactions and biomarkers.

Mechanism

HTRF is based on Eu^{3+} / Tb^{2+} cryptate donors and XL665 or d2 acceptors, which can be coupled to almost any biomolecule desired, including proteins, peptides, DNA and small molecules. The technology is based on no wash assay procedure detecting proximity events between donor and acceptor dyes.

The assay detection is obtained upon dispensing acceptor and donor conjugates to the sample to be assessed (e.g. enzymatic reaction mixture, cell lysate, or supernatant). No washing steps are required (homogeneous assay), and detection is performed after the completion of incubation, by measuring both specific donor and acceptor fluorescence (Figure 1).

To compensate for well-to-well variation, the ratio of both values is calculated (hence ratiometric assay). Donor fluorescence will always be detected and used as an internal control, while an emission signal from the acceptor is only detected if both biomolecules are in close proximity and FRET occurs³³.

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

³³ <http://www.cisbio.com/drug-discovery/htrf-technology>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	HTRF (Europium)	HTRF (Europium)	HTRF (Europium)
Measurement Mode	FI Top (2 labels)	FI Top (2 labels)	FI Top (2 labels)
Excitation wavelength Donor	320(25)	320(25)	320(25)
Emission wavelength Donor	620(10)	620(10)	620(10)
Lag time	150 µs	100 µs	100 µs
Integration time	500 µs	400 µs	400 µs
Flashes	50	50	50
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms
Excitation wavelength Acceptor	320(25)	320(25)	320(25)
Emission wavelength Acceptor	665 (8)	665(8)	665(8)
Lag time	150 µs	100 µs	100 µs
Integration time	500 µs	400 µs	400 µs
Flashes	50	50	50
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	same as Label 1	same as Label 1	same as Label 1
Settle time	0 ms	0 ms	0 ms



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	HTRF (Terbium)	HTRF (Terbium)	HTRF (Terbium)
Measurement Mode	FI Top (2 labels)	FI Top (2 labels)	FI Top (2 labels)
Excitation wavelength Donor	340(35)	340(35)	340(35)
Emission wavelength Donor	620(10)	620(10)	620(10)
Lag time	150 µs	100 µs	100 µs
Integration time	500 µs	300 µs	300 µs
Flashes	50	50	50
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms
Excitation wavelength Acceptor	340(35)	340(35)	340(35)
Emission wavelength Acceptor	665 (8)	665 (8)	665 (8)
Lag time	150 µs	100 µs	100 µs
Integration time	500 µs	400 µs	400 µs
Flashes	50	50	50
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	same as Label 1	same as Label 1	same as Label 1
Settle time	0 ms	0 ms	0 ms

Adapta® Universal Kinase Assay and Substrates

Life Technologies' version of the ADP detection assay

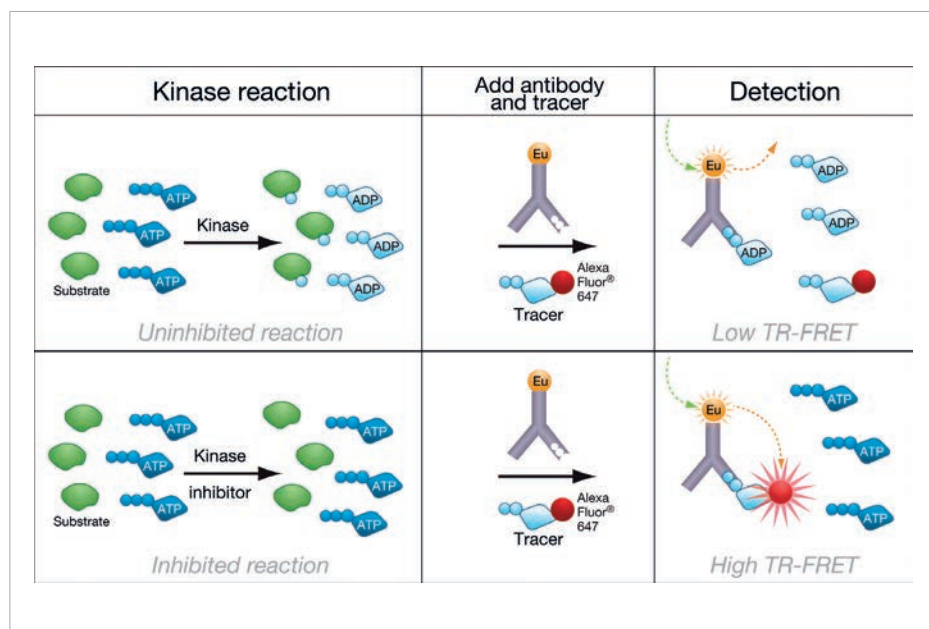


Figure 1: Schematic principle of the Adapta assay

ASSAY OVERVIEW

Technology

Time-Resolved Fluorescence Resonance Energy Transfer (TR-FRET), ratiometric

Principle, provider, major application

Life Technologies' Adapta Universal Kinase Assay Kit is a homogeneous, fluorescence-based immunoassay for measuring the activity of ADP-producing enzymes, mainly kinases. Additionally, the Adapta assay is available for a selection of lipid- and peptide-based substrates. Life Technologies supplies europium-coupled antibody specific for ADP. In contrast to the Transcreener® platform, Adapta is only available for ADP.

Mechanism

The ADP-specific antibody and the tracer are added to the sample. In an inhibited reaction (Figure 1), the monitored kinase produces no ADP and only the added, tracer-bound ADP molecule binds to the antibody, causing a high FRET signal.

Active kinases convert ATP to ADP. The free ADP competes with the tracer-bound ADP to bind to the antibody, resulting in a low FRET signal. Hence, the signal intensity is indirectly proportional to the activity of the kinase.

Instrument parameters



Reader	Infinite 200 PRO
Assay	Adapta Assay
Measurement Mode	FI Top (2 labels)
Excitation wavelength Donor	340(35)
Emission wavelength Donor	620(10)
Lag time	100 μ s
Integration time	200 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms
Excitation wavelength Acceptor	340(35)
Emission wavelength Acceptor	665(8)
Lag time	100 μ s
Integration time	200 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

LanthaScreen™ Kinase Activity Assays³⁴



Kinase activity assay with TR-FRET readout

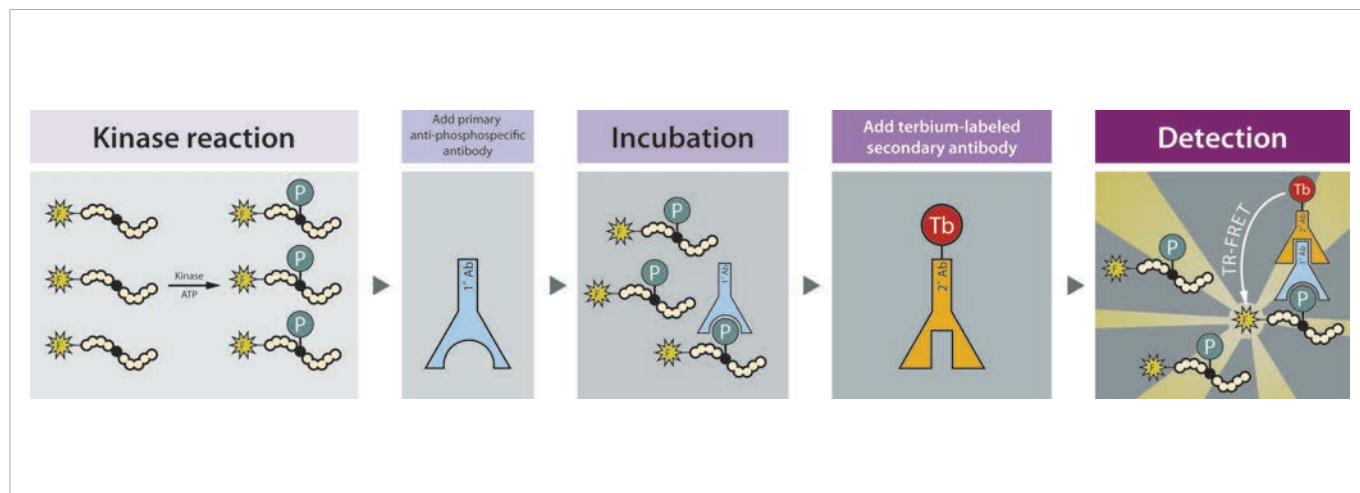


Figure 1: Schematic principle of the LanthaScreen assay

ASSAY OVERVIEW

Technology

Time-Resolved Fluorescence Resonance Energy Transfer (TR-FRET), ratiometric

Principle, provider, major applications

LanthaScreen is a kinase activity assay sold by Life Technologies. Kinases are important cellular enzymes, and their major function is to add phosphate groups to peptide substrates. For researchers, it is important to know how active kinases are in the presence of certain inhibitors. LanthaScreen quantifies kinase activity by measuring the amount of phosphorylated substrate.

Format

Life Technologies supplies a broad panel of fluorescein-labeled substrates and the corresponding lanthanide-labeled antibody specifically for the detection of phosphorylated substrates.

Mechanism

Kinase and fluorescein-labeled substrates are incubated to enable phosphorylation. After incubation, a terbium-labeled antibody is added to the reaction (Figure 1).

Scenario 1 – kinase is active

The substrate was phosphorylated, allowing the phospho-specific antibody to bind. The fluorescein and terbium labels are now in close proximity, resulting in a high FRET signal.

Scenario 2 – kinase is inactive

No phosphorylation occurred, and therefore the antibody could not bind to the substrate. FRET cannot occur. The final result is a dimensionless number that is calculated as the ratio of the acceptor (fluorescein) signal to the donor (terbium) signal.

³⁴ www.invitrogen.com/lanthascreen

Instrument parameters



Reader	Infinite 200 PRO
Assay	LanthaScreen
Measurement Mode	FI Top (2 labels)
Excitation wavelength Donor	340(35)
Emission wavelength Donor	495(10)
Lag time	100 μ s
Integration time	200 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms
Excitation wavelength Acceptor	340(35)
Emission wavelength Acceptor	520(10)
Lag time	100 μ s
Integration time	200 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

SUPPORT

Links

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FP – Fluorescence Polarization.

Binding assay for biomolecules

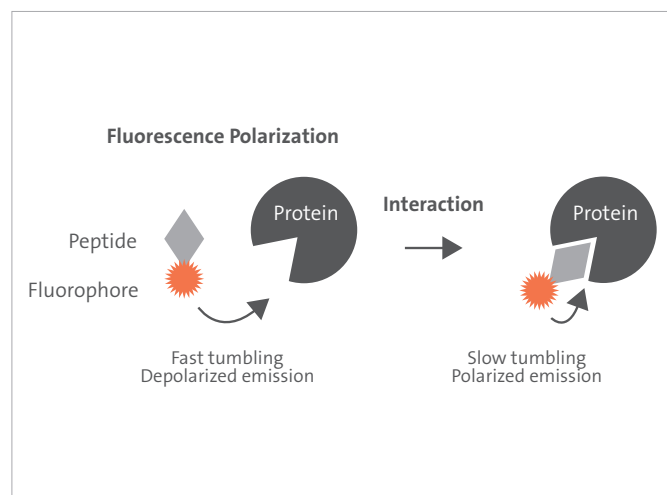


Figure 1: Schematic representation of Fluorescence Polarization

$$mP = 1000 \times \frac{I_{\parallel} - I_{\perp}}{I_{\parallel} + I_{\perp}}$$

Equation 1: Calculation of the polarization value. $I_{\parallel}$ = light parallel to the polarization plane. $I_{\perp}$ = light perpendicular to the polarization plane

TECHNOLOGY

Fluorescence anisotropy is colloquially referred to as Fluorescence Polarization. On excitation with polarized light, the emission from many samples is also polarized. Rotational movements of the excited molecule destroy this correlation. The extent of polarization remaining depends on the size of the molecules measured: the bigger the molecules, the slower they rotate and the higher the conservation of the original polarization. Other influences include solvent viscosity, temperature and the lifetime of the excited state.³⁵

The following metaphor is an easy way to explain FP. Imagine a little child playing in a field. While it is free, it can twist and turn as much as it wants in any direction. Once it is 'attached' to its mother's hand, the movements will slow down and get direction. This comparison can be used to show how FP detects molecular interactions (Figure 1). The little child represents the smaller of the interaction partners, and the mother the larger one. A fluorescent probe is attached to the small molecule to observe the turning and twisting movements. As long as there is no interaction between the small and the large molecule, the rotation of the fluorophore is fast and the emitted light depolarized. Once it binds to a larger

interaction partner, its movements will slow down and the emitted light will preserve more and more of the original polarization.

Assay design

A major application of FP is the detection of molecular interactions. FP assays require interaction partners to be different sizes, and the smaller molecule to be labeled with a fluorophore. Commercial assays commonly provide these labeled partners or substrates. The final result is a ratio of the polarization values, measured before and after addition of the suspected interaction partner. The polarization is calculated using the equation given below, measuring the intensity of emitted light in perpendicular and parallel planes.

Changes in polarization give information about the creation of interactions and their strength. A higher mP (milli-polarization) value represents a stronger interaction between the two molecules.

Major applications

- PolarScreen™ Assays
- Transcreeper® Assays
- Predictor™ hERG assay

³⁵ Lakowicz, J. R. (1999). Principles of Fluorescence Spectroscopy. Kluwer Academic / Plenum Publishers



PolarScreen™

FP-based kinase activity assay; FP-equivalent to LanthaScreen

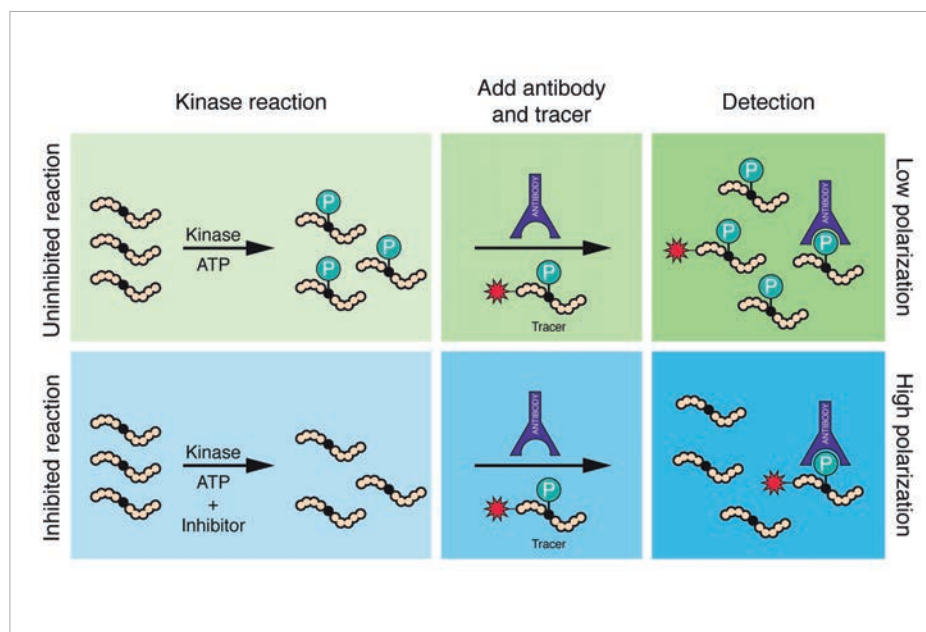


Figure 1: Schematic principle of the PolarScreen assay.

ASSAY OVERVIEW

Technology

Fluorescence Polarization (FP)

Provider, major application

PolarScreen³⁶ is Life Technologies' version of a FP-based kinase activity assay. Life Technologies offers a panel of phospho-specific antibodies which detect peptide substrates when phosphorylated by protein kinases.

Mechanism

The target kinase is incubated with a dedicated, unlabeled substrate (Figure 1). Antibody – specific for the phosphorylated- phosphorylation site of the substrate – and additional, tracer-bound substrate are added to the sample. If the unlabeled substrate's phosphorylation site remains unphosphorylated, for example due to an inactive enzyme, the antibody will only bind to the added, tracer-bound substrate, causing a low FP signal.

Active kinases will phosphorylate the non-tracer bound substrate, which then competes with the tracer-bound substrate for binding to the antibody, resulting in a high FP signal. Hence, the FP signal is directly proportional to the amount of phosphorylated substrate.

Format

Fluorescence labels are available for green, red or far-red detection. Red fluorescence readouts help to reduce autofluorescence.

³⁶ <https://www.thermofisher.com/us/en/home/industrial/pharma-biopharma/drug-discovery-development/target-and-lead-identification-and-validation/nuclear-receptor-biology/nuclear-receptor-biochemical-assays/polarscreen-competition-assays.html>

Instrument parameters



PolarScreen Green

Reader	Infinite 200 PRO
Assay	PolarScreen
Measurement Mode	FP
Excitation wavelength Donor	485(20)
Emission wavelength Donor	535(25)
Lag time	0
Integration time	20 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

PolarScreen Red

Assay	PolarScreen
Measurement Mode	FP
Excitation wavelength Donor	535(25)
Emission wavelength Donor	590(20)
Lag time	0
Integration time	20 μ s
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

PolarScreen Far Red

Assay	PolarScreen
Measurement Mode	FP
Excitation wavelength Donor	610(20)
Emission wavelength Donor	670(40)
Lag time	0
Integration time	default
Flashes	10
Mirror	automatic
Gain	optimal
Z-position	calculated from well
Settle time	0 ms

SUPPORT

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Transcreener®

Nucleotide (ADP, GDP, etc.) detection assay

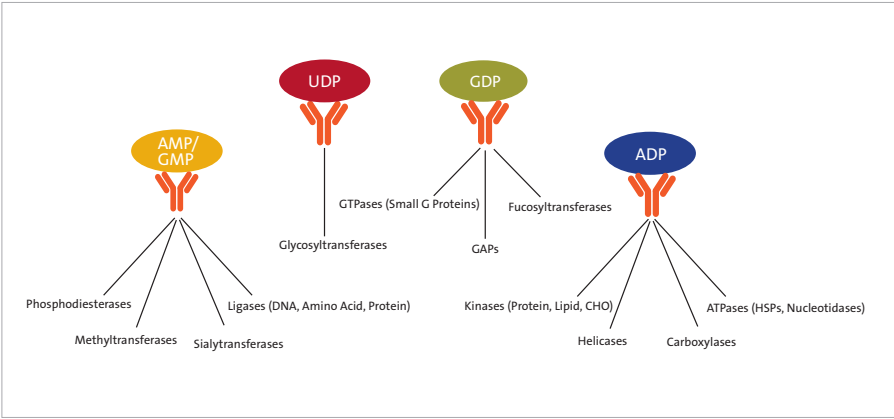


Figure 1: Transcreener targets

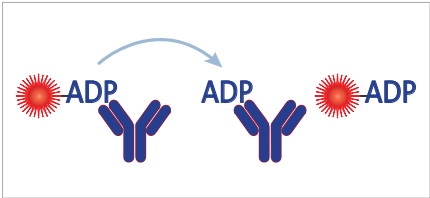


Figure 2: Transcreener principle

Assay	Readout
Transcreener ADP ² Assays	FP, FI, TR-FRET
Transcreener AMP/GMP Assay	FP, FI
Transcreener GDP Assays	FP
Transcreener UDP Assays	FP

Table 1: Alternative readouts

ASSAY OVERVIEW

Technology

- Fluorescence polarization (FP)
- Fluorescence intensity (FI)
- Time-resolved fluorescence resonance energy transfer (TR-FRET)

Principle, provider

Bellbrook’s Transcreener assays³⁷ are designed to detect various mono- and dinucleotides using FP, TR-FRET or FI detection mode. Four assays (Table 1) cover thousands of target enzymes, including any kinase, ATPase or GTPase. Transcreener is a universal assay method that can be used across entire families of nucleotide-dependent enzymes. All assays are based on different antibodies that show a high affinity for one specific nucleotide (Figure 1).

Mechanism

The mechanism is the same for all Transcreener assays. The antibody is preloaded with the corresponding nucleotide, which is conjugated to a tracer molecule. All assays use a far red tracer that minimizes compound interference. For example, in the ADP² FP assay the detection mixture comprises of Alexa 633 ADP and a highly selective ADP monoclonal antibody. The Transcreener ADP² FP assay measures the progress of any enzyme that produces ADP by displacing the tracer by ADP thereby causing a decrease in fluorescence polarization. (Figure 2)

³⁷ <https://www.bellbrooklabs.com/technical-resources/transcreener-faq/>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	Transcreeper FP	Transcreeper FP	Transcreeper FP
Measurement Mode	FP	FP	FP
Excitation wavelength	610(20)	620(10)	620(10)
Emission wavelength	670(25)	670(25)	670(25)
Integration time	default	default	default
Flashes	10	30	30
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	50 ms	100 ms	100 ms

Transcreeper FI			
Assay	Transcreeper FI	Transcreeper FI	Transcreeper FI
Measurement Mode	FI Top	FI Top	FI Top
Excitation wavelength	580(20)	580(20)	580(20)
Emission wavelength	620(20)	620(20)	620(20)
Lag time	n.a.	n.a.	n.a.
Integration time	20 μ s	default	default
Flashes	25-100	30-100	30-100
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	50 ms	100 ms	100 ms

Transcreeper TR-FRET			
Assay	Transcreeper TR-FRET	Transcreeper TR-FRET	Transcreeper TR-FRET
Measurement Mode	FI Top (2 labels)	FI Top (2 labels)	FI Top (2 labels)
Excitation wavelength Donor	320(25)	320(25)	320(25)
Emission wavelength Donor	620(10)	620(10)	620(10)
Lag time	150 μ s	150 μ s	150 μ s
Integration time	500 μ s	500 μ s	500 μ s
Flashes	10	30	30
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	0 ms	0 ms
Excitation wavelength Acceptor	320(25)	320(25)	320(25)
Emission wavelength Acceptor	665(8)	665(8)	665(8)
Lag time	150 μ s	150 μ s	150 μ s
Integration time	500 μ s	500 μ s	500 μ s
Flashes	10	30	30
Mirror	automatic	automatic	automatic
Gain	optimal	optimal	optimal
Z-position	calculated from well	calculated from well	calculated from well
Settle time	0 ms	n.a.	n.a.

SUPPORT

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- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
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Learn more at www.tecan.com

Alpha – Amplified Luminescent Proximity Homogeneous Assay.

FRET equivalent with reduced distance limitations and amplification

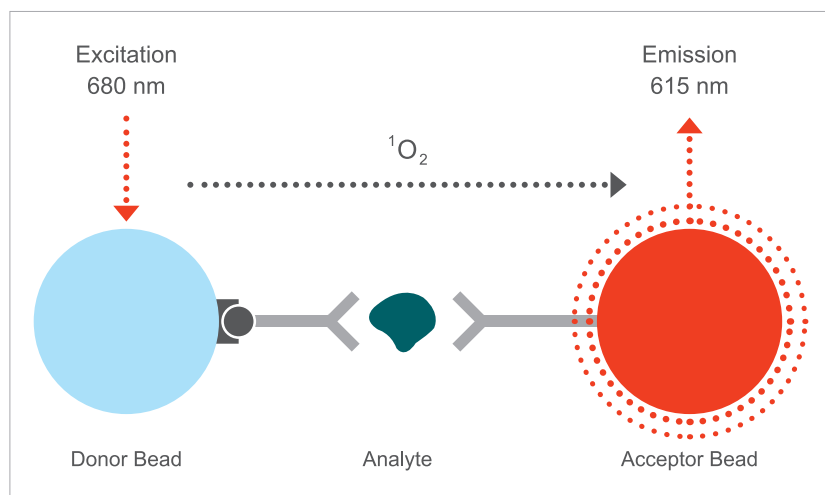


Figure 1: Schematic function of the Alpha technology.

TECHNOLOGY

Mechanism

Alpha³⁸ is a technology invented by PerkinElmer. The principle is similar to (TR-)FRET, since the Alpha Technology also relies on the interaction of an acceptor and a donor to yield a signal (Figure 1). However, instead of using simple fluorophores, chemically reactive beads are used, and the chemistry that produces the signal is also different. The main advantage of AlphaScreen® and AlphaLISA® is that the distance between the interaction partners can be up to 200 nm, compared to 10 nm for (TR-)FRET. Furthermore, there is an amplification effect which increases the sensitivity of the assay. Background is reduced because the emission wavelength is lower than the excitation wavelength.

Assay design

AlphaScreen (Amplified Luminescent Proximity Homogeneous Assay) is a bead-based screening technology developed by PerkinElmer for fast and reliable detection of biological interactions. The AlphaScreen chemistry employs donor and acceptor beads that can be linked to various types of biologically

relevant molecules. The phthalocyanine photosensitizer molecules contained in the AlphaScreen donor beads convert ambient oxygen into large quantities of singlet oxygen when excited by a high energy light source at a wavelength of 680 nm. The singlet oxygen molecules are able to cover a distance of up to 200 nm during their half-life of approximately 4 μs. If AlphaScreen acceptor beads are in close proximity to the donor beads, due to a biological binding event between their coupling partners, the singlet oxygen molecules are able to initiate a cascade of energy transfer steps in the acceptor beads, ultimately resulting in the generation of a strong light emission in the range of 520-620 nm. Due to the amplified signal generation, even small amounts of biological analytes can be detected.

AlphaLISA is a homogeneous, no-wash alternative to conventional ELISA assays based on PerkinElmer's bead-based Alpha (Amplified Luminescent Proximity Homogeneous Assay) technology. AlphaLISAs can be set up as sandwich or competitive immunoassays to detect and quantify molecules of interest in biological samples. High energy excitation of photosensitizer molecules within the AlphaLISA donor beads at 680 nm converts

³⁸ <http://www.perkinelmer.com/catalog/category/id/alphatech>



ambient oxygen to singlet oxygen, which is able to react with the chemistry in the acceptor beads if these are in close proximity. A cascade of energy transfer steps ultimately results in the generation of a strong output signal at 615 nm, indicating specific binding between the molecules attached to the two bead types.

The fluorophores embedded in the AlphaLISA acceptor beads produce a narrower bandwidth signal than the acceptor beads used for classical AlphaScreen assays. This makes AlphaLISAs less prone to signal interference at wavelengths of <600 nm, increasing the sensitivity and robustness of the assay. The no-wash nature of this assay makes it easy to use, and the use of dedicated AlphaLISA

optics permits the analysis of target molecules in blood and serum by drastically reducing the effect of hemoglobin within a sample.

The Alpha Technology's versatility offers the possibility to assay many biological targets, including enzymes, receptor-ligand interactions, low affinity interactions, second messenger levels, DNA, RNA, proteins, protein-protein interactions, peptides, sugars and small molecules³⁸.

Major applications

- AlphaScreen / AlphaLISA

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

AlphaScreen / AlphaLISA

Alpha Technology assay platforms

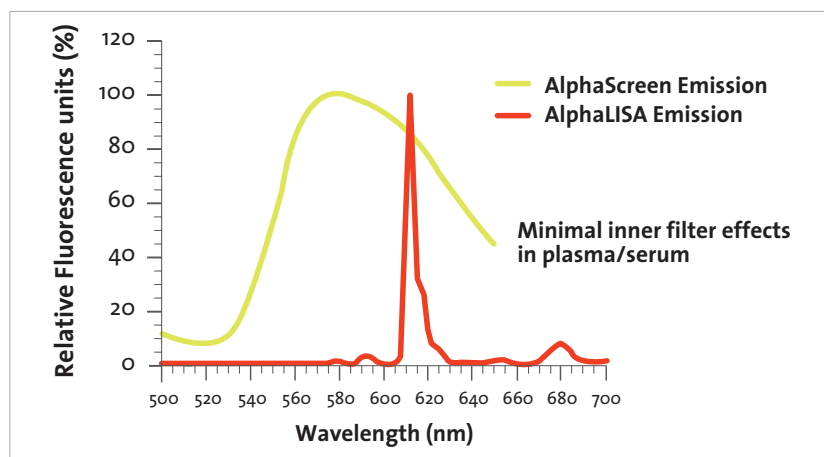


Figure 1: Emission spectra of TAR (AlphaScreen) and europium (AlphaLISA) acceptor beads.

ASSAY OVERVIEW

Technology

Alpha Technology

Provider, Format

AlphaScreen and AlphaLISA are PerkinElmer's Alpha Technology³⁹ assay / reagent platforms. PerkinElmer supplies donor beads coupled to streptavidin, and blank acceptor beads for self-labeling. Additionally, acceptor beads can be purchased precoupled to antibodies specific for a broad range of targets.

Principle

AlphaLISA is a development targeting laboratories working with crude blood samples, as the autofluorescence of hemoglobin overlaps with the emission peak (Figure 1) of the AlphaScreen acceptor. The main difference between AlphaLISA and AlphaScreen is that the AlphaLISA emission peak (europium emission) of the acceptor bead is smaller than that of the AlphaScreen assay (rubrene emission).

Mechanism

For a closer description of the assay mechanism, please refer to the Alpha Technology section.

³⁹ <http://www.perkinelmer.com/catalog/category/id/alphatech>

Instrument parameters



Reader	Spark	Spark Cyto
Assay	AlphaScreen	AlphaScreen
Measurement Mode	Alpha Technology	Alpha Technology
Excitation wavelength	n.a.	n.a.
Emission wavelength	n.a.	n.a.
Excitation time	100 ms	100 ms
Integration time	300 ms	300 ms
Wavelength	520-620 nm	520-620 nm
Gain	n.a.	n.a.
Settle time	0 ms	0 ms
Temperature correction	activated	activated
Assay	AlphaLISA	AlphaLISA
Measurement Mode	Alpha Technology	Alpha Technology
Excitation wavelength	n.a.	n.a.
Emission wavelength	n.a.	n.a.
Excitation time	100 ms	100 ms
Integration time	300 ms	300 ms
Wavelength	610-635 nm	610-635 nm
Gain	n.a.	n.a.
Settle time	0 ms	0 ms
Temperature correction	activated	activated
Assay	AlphaPlex	AlphaPlex
Measurement Mode	Alpha Technology	Alpha Technology
Excitation wavelength	n.a.	n.a.
Emission wavelength	n.a.	n.a.
Excitation time	100 ms	100 ms
Integration time 1	300 ms	300 ms
Integration time 2	300 ms	300 ms
Wavelength 1	610-635 nm	610-635 nm
Wavelength 2	535-560 nm	535-560 nm
Gain	n.a.	n.a.
Settle time	0 ms	0 ms
Temperature correction	activated	activated

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

LUMI-Luminescence.

Light is emitted from the sample



Figure 1: Firefly

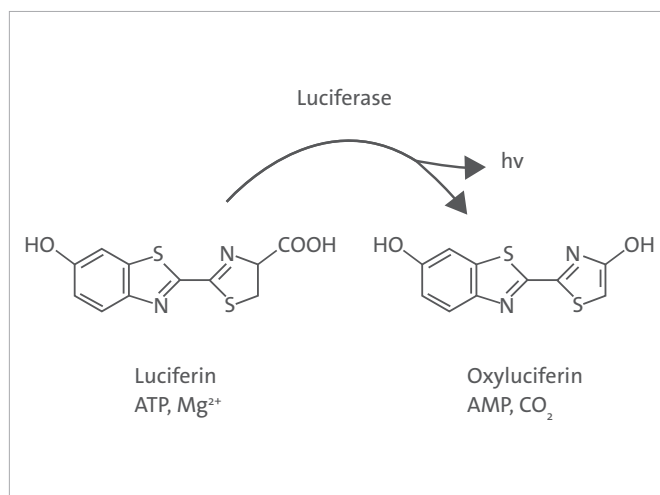


Figure 2: Substrate cleavage by luciferase

TECHNOLOGY

Luminescence is widely known as a reaction that causes the release of light. It can be caused by chemical reactions, electrical energy, subatomic motion, or stress on a crystal. For molecular biology, bioluminescence is the most important of the various luminescence reactions. At the core of this technology is the luciferase enzyme. Luciferases (Firefly, Renilla) (Figure 1) convert a substrate into an excited state (Figure 2). When returning to the ground state, a photon (light) is released (emitted).

One striking difference between fluorescence and luminescence is that luminescence requires no excitation light. This reduces the background to almost zero, resulting in better sensitivity.

Various forms of luminescence can be differentiated.

1. Glow luminescence, which generates stable and measurable light up to several hours, for example the BioThema ATP assay
2. Flash luminescence, which is characterized by rapid, but short-lived, light generation, for example DLR and Aequorin (injectors are required for flash luminescence)
3. Multicolor luminescence, such as BRET^{1/2/3/e} and Chroma-Glow™

Major applications

- Dual-Luciferase® Reporter Assay (DLR™)
- Bio Thema ATP detection kit
- BRET (Bioluminescence Resonance Energy Transfer)
- NanoBRET™



Bio Thema ATP detection kit⁴²



ATP detection kit based on luminescence

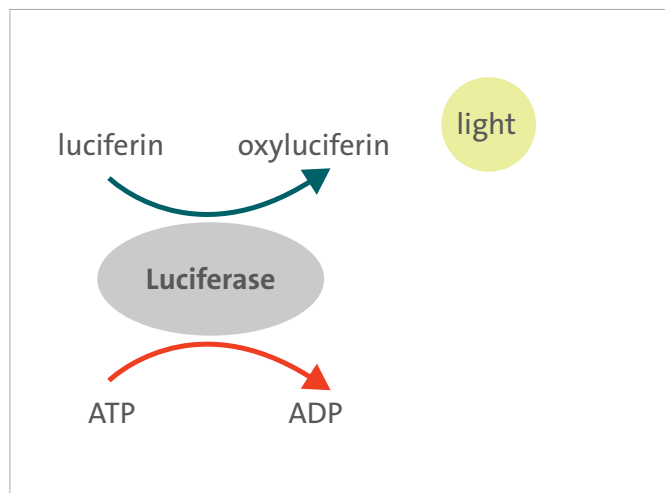


Figure 1: Schematic principle of luminescence caused by a luciferase based on the conversion of ATP.

ASSAY OVERVIEW

Technology

Luminescence (Glow)

Principle, provider

ATP (Adenosine-Tri-Phosphate) is a molecule for short time energy storage and required for almost any reaction in living organisms. Biothema's ATP detection kit measure and quantifies the level of ATP and correlates it to the activity of an enzyme or cells.

Format

The kit consists of a luciferase, the substrate and an ATP standard to quantify the signal of the sample.

Mechanism

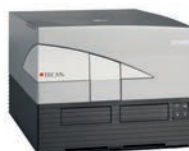
The luciferase and substrate are added to the sample. If ATP is present, the luciferase converts the substrate and light is released (Figure 1). If no ATP is present, no light is released. The more ATP is present in the sample the stronger is the signal. In a last step the signal is quantified by comparing it to a standard ATP curve.

Major applications

Major applications include ATP detection, cell proliferation, cytotoxicity, enzymatic monitoring.

⁴² <http://biothema.se/products/kits/>

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	Biothema ATP	Biothema ATP	Biothema ATP
Measurement Mode	Luminescence	Luminescence	Luminescence
Integration time	1000 ms	1000 ms	1000 ms
Attenuation	automatic	automatic	automatic
Settle time	0 ms	0 ms	0 ms

SUPPORT

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- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

Dual-Luciferase® Reporter Assay (DLR™)⁴⁰



Luminescence-based normalizable gene activator assay

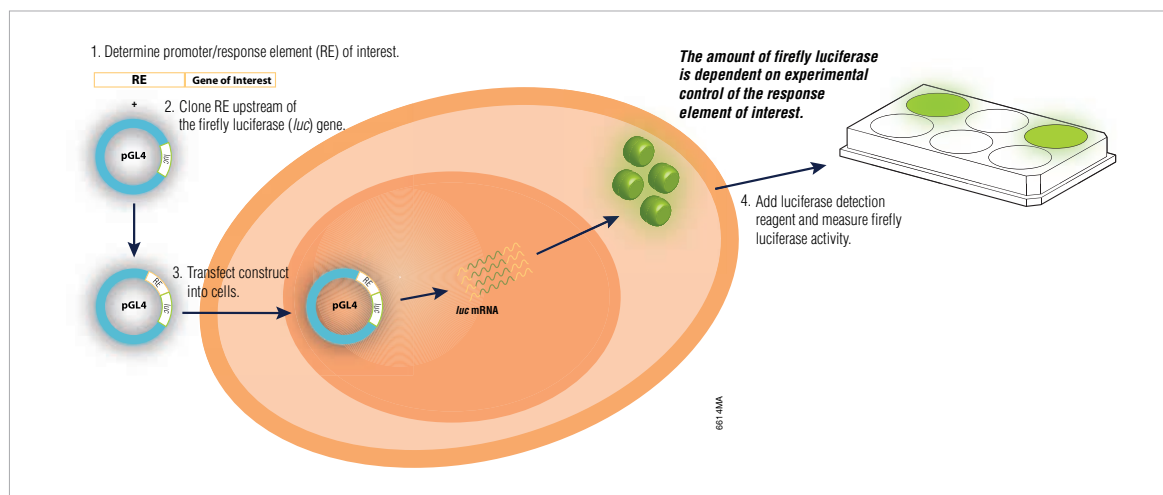


Figure 1: DLR assay mechanism

ASSAY OVERVIEW

Technology

Luminescence (flash)

Purpose, provider, major application

Promega's DLR assay measures gene activation / expression using a luminescence-based readout. One particular challenge of gene activator assays is quantification of the result. The DLR assay resolves this issue by normalizing the output using two luciferases. While one luciferase measures the expression of the gene of interest, the second luciferase measures the expression of a housekeeping gene⁴¹, which is used to normalize the signal. Promega has two series of firefly and Renilla luciferase vectors – pGL4 and pRL – designed for use with the DLR assay systems.

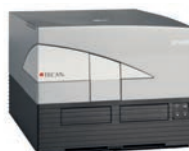
Mechanism

Cells need to be transfected with both luciferase reporter genes (Figure 1). The firefly luciferase reporter is measured first by adding Luciferase Assay Reagent II (LAR II) to generate a 'glow-type' luminescent signal. After quantifying the firefly luminescence, the reaction is quenched, and the Renilla luciferase reaction simultaneously initiated, by adding Stop & Glo® Reagent to the tube. The Stop & Glo Reagent produces a 'glow-type' signal from the Renilla luciferase, which decays slowly over the course of the measurement.

⁴⁰ http://www.promega.com/products/reporter-assays-and-transfection/reporterassays/dual_luciferase-reporter-assay-system/

⁴¹ Housekeeping genes are proteins or enzymes that are constitutively expressed in most cells as for instance the DNA polymerase

Instrument parameters



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	DLR	DLR	DLR
Measurement Mode	Luminescence	Luminescence	Luminescence
	well-wise	well-wise	well-wise
Integration time	10,000 ms	10,000 ms	10,000 ms
Output	n.a.	counts/sec	counts/sec
Attenuation	automatic	automatic	automatic
Settle time	0 ms	0 ms	0 ms
Injections	Injector A: 100 µl (refill for every injection), wait 3 s	Injector A: 100 µl (refill for every injection), wait 3 s	Injector A: 100 µl (refill for every injection), wait 3 s
	Injector B: 100 µl (refill for every injection), wait 3 s	Injector B: 100 µl (refill for every injection), wait 3 s	Injector B: 100 µl (refill for every injection), wait 3 s

SUPPORT

Links

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BRET (Bioluminescence Resonance Energy Transfer).

A FRET modification using the donor fluorophore as the light source

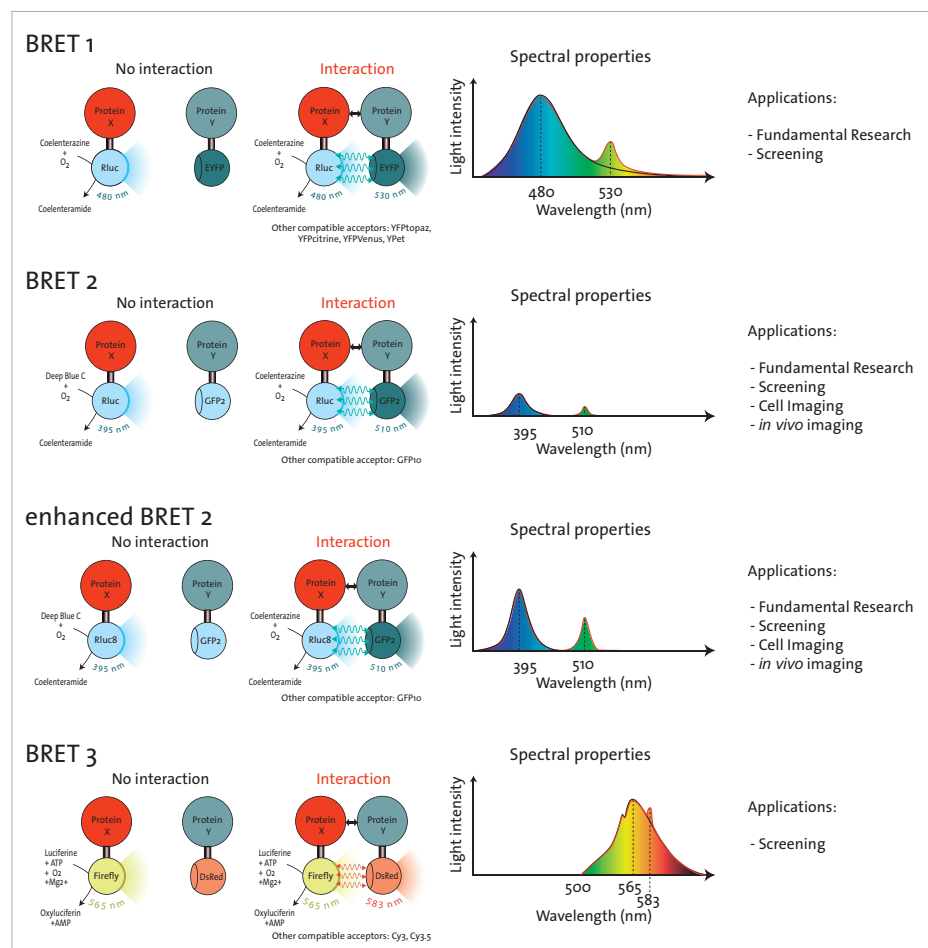


Figure 1: Summary of the most relevant BRET technologies.

TECHNOLOGY

BRET is a modification of FRET, the main difference being that the donor fluorophore is substituted by a luciferase. A major application for BRET is interaction studies. The advantage is that no excitation light is needed, and hence the background is much lower. BRET itself is a technology, and therefore can't be commercialized. However, various companies offer proprietary assay components for this technology, for example BRET^{1/3/e} are an unlicensed technology, while BRET² is a licensed product of PerkinElmer.

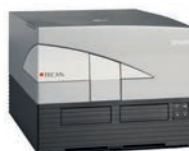
The difference between BRET¹ and BRET² lies in the selection of the donor and acceptor proteins / fluorophores (Figure 1). This results in a shift of the excitation and emission wavelength.⁴²

A new improved chemistry version of BRET, NanoBRET[™], combines an extremely bright NanoLuc[®] luciferase with a means for tagging intracellular proteins with a long-wavelength fluorophore (HaloTag), providing a better dynamic range and sensitivity.⁴³

⁴² Bacart J, Corbel C, Jockers R, Bach S, Couturier C: The BRET technology and its application to screening assays. *Biotechnol J* 2008, 3:311-324.

⁴³ Machleit et al. NanoBRET-A novel BRET platform for the analysis of protein-protein interactions. *ACS Chem. Biol.*, 2015, 10 (8), pp 1797-1804.

Compatible readers



Reader	Infinite 200 PRO	Spark	Spark Cyto
Assay	NanoBRET-based assays	NanoBRET-based assays	NanoBRET-based assays
Measurement Mode	Dual Color Luminescence	Luminescence Multi Color	Luminescence Multi Color
Filter 1 / Wavelength Range 1	Blue1_NB	445-470 nm	445-470 nm
Integration time	300 ms	300 ms	300 ms
Filter 2 / Wavelength Range 2	Red_NB	610-635 nm	610-635 nm
Integration time	300 ms	300 ms	300 ms
Settle time	0 ms	0 ms	0 ms

SUPPORT

Links

- <http://lifesciences.tecan.com/multimode-plate-reader?p=Technology>
- http://lifesciences.tecan.com/infinite200pro_readmodes

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Founded in Switzerland in 1980, Tecan has manufacturing, research and development sites in both North America and Europe, and maintains a sales and service network in 52 countries. To date, Tecan has distributed approximately 20,000 microplate readers worldwide, and is committed to continuous technological improvements and compliance with the highest global quality standards.

IMPORTANT INFORMATION

Tecan has not independently validated the methods described in the document with all possible sample types or analytical uses and is providing this example method as a convenience to users. Consistent with USP recommendations and good laboratory practice, the user must independently evaluate and validate: (a) the suitability of the method for their use, (b) their ability to process samples of their choosing following the method; and (c) their ability to proficiently perform the method in their facility with their personnel.

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