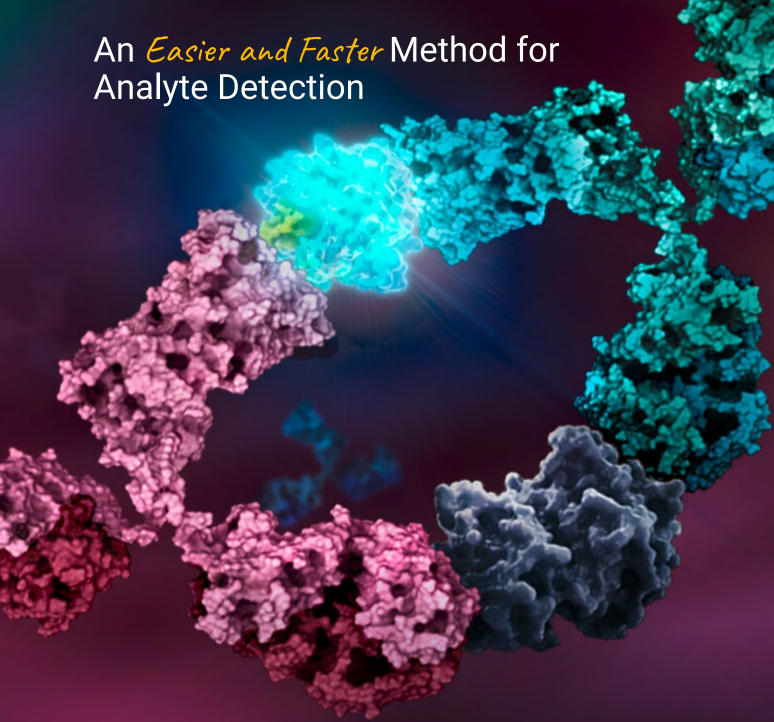
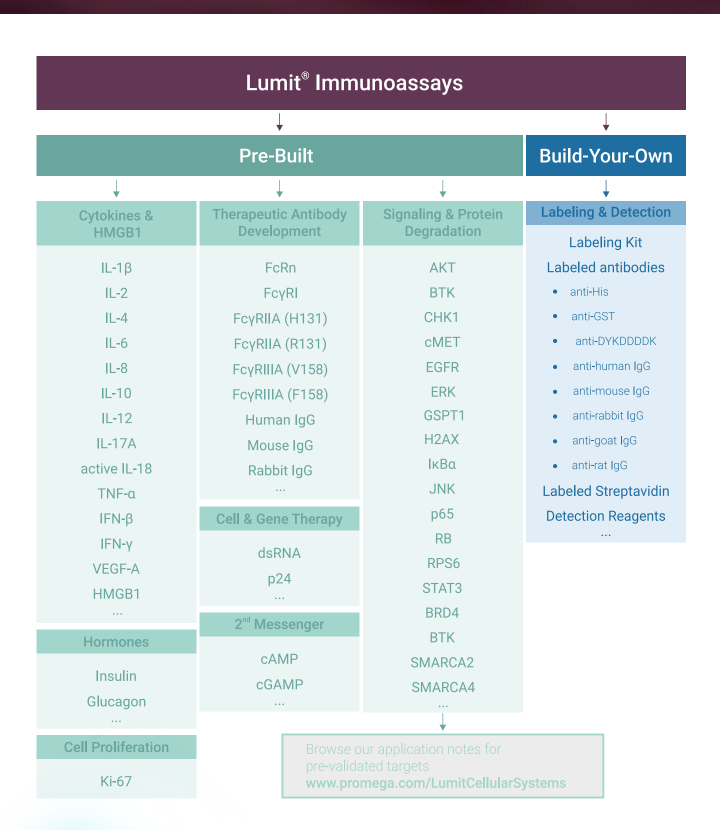


# Lumit® Immunoassays



Signaling | Metabolic Regulators | Cytokines Protein Interaction & Degradation | Cell & Gene Therapy Therapeutic Antibody Development







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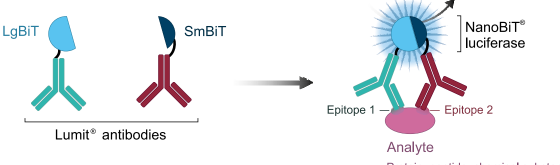
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## 1. Introduction

## 1.1 Lumit® Technology

Detection and quantification of analytes are often performed using time-consuming, multi-step methods such as Western blotting and ELISA. The Lumit® technology offers a simple and fast alternative for running homogeneous immunoassays in multi-well plate formats. Its high specificity and compatibility with high-throughput screening (HTS) make it a powerful tool for scientists working in areas ranging from basic research to drug discovery.

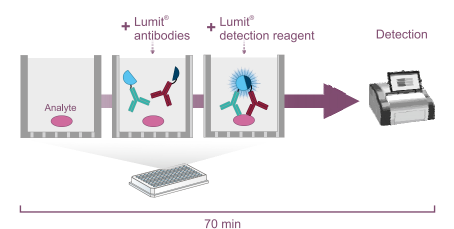
#### **Assay Principle**



Protein, peptide, chemical substance, etc.

The underlying principle of Lumit® is the NanoLuc® Binary Technology (NanoBiT®). In Lumit® Immunoassays, two antibodies are chemically labeled with the small and large engineered subunits of NanoBiT luciferase, i.e., SmBiT and LgBiT. Direct or indirect binding to the analyte brings these subunits into spatial proximity, enabling them to reconstitute a functional luciferase. In the presence of the substrate furimazine, a bright luminescence is generated that is directly proportional to the amount of analyte in the sample.

#### **Assay Workflow**



#### **Features & Benefits**

- Simple homogenous workflow
- ✓ No washing
- ✓ No blocking
- · Detection of analytes in
  - ✓ Buffer
  - ✓ Cell culture supernatants
  - ✓ Cell lysate
- Signal detection on a conventional plate-reading luminometer

#### **Advantages**

- · Simple add-and-read protocol with no washing steps
- · Fast results with reduced hands-on time
- · No immobilization to plates, beads, or other surfaces required
- · Direct analyte measurement in cell culture plates
- · Sensitive luminescence detection with a wide dynamic range
- Detection on a conventional luminometer
- · High specificity and low background signal
- Easy to automate and HTS-compatible (96-well and 384-well plates)

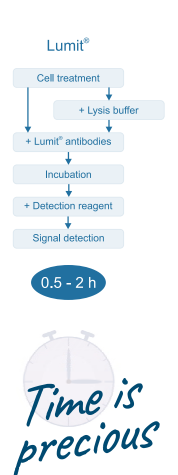
#### Lumit® vs. Conventional Immunoassays

Lumit® Immunoassays are fast, add-and-read plate-based assays. No washing steps are required making Lumit® a compelling alternative to labor-intensive methods, e. g., ELISA and Western blot.

#### Your Short-Cut to High-Quality Data







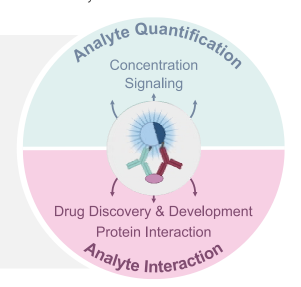
## 1. Introduction

### 1.2 Applications & Assay Formats

A multitude of applications is supported by the different Lumit® Immunoassay formats.

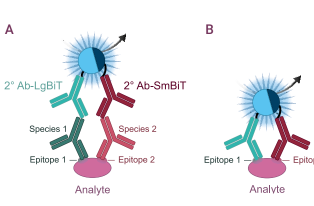
#### **Applications**

- · Quantitative analyte detection in biological samples
- Competitive binding studies for proteins and small molecules
- · Drug screening for proteins and small molecules
- · Measurement of signaling pathway activation
- · Analysis of protein interactions
- High-throughput screenings (HTS)



#### **Analyte Quantification**

### Indirect Assay Format Di



#### **Direct Assay Format**

(A) In the indirect assay format, two epitopes on the analyte are recognized by two different primary antibodies from different species. These antibodies are detected using BiT-labeled secondary antibodies.

The Lumit® technology supports various assay formats that enable the quantitative

analysis of analytes.

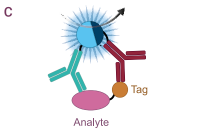
using BiT-labeled secondary antibodies. This format has been extensively validated for the analysis of post-translational modification (PTM) in cell lysates and is also referred to as Lumit® Immunoassay Cellular

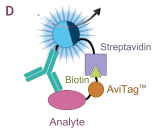
Systems.

**(B)** The direct assay format employs two BiT-labeled primary antibodies. Alternatively, antibodies can be replaced with labeled analyte-binding proteins, as demonstrated in the Lumit® Assays for IL-18 and dsRNA.

- (C) BiT-labeled anti-tag antibodies or
- **(D)** BiT-labeled streptavidin are alternative formats available for use with tagged analytes.

#### **Alternative Assay Formats**

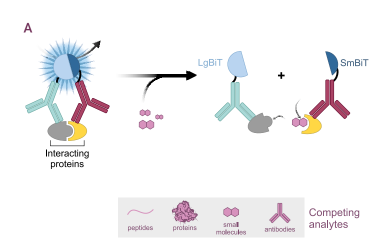




#### **Analyte Interaction**

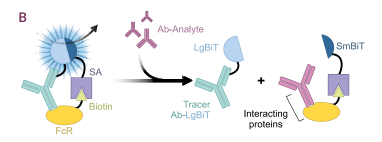
Binary interactions between analytes can be easily studied using the Lumit® technology. Different formats allow the determination and characterization of binding events, such as protein:protein and protein:ligand interactions, through competitive loss- or gain-of-signal assays.

#### Loss-of-Signal Binding Immunoassays

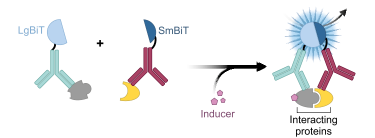


Lumit® loss-of-signal binding assays are used for determining the potency of competing analytes *in vitro*.

- (A) This biochemical assay format uses two BiT-labeled primary antibodies to target the interacting proteins of interest, enabling the determination of competing analytes' (inhibitors) potency. The addition of competing analytes reduces the luminescent signal. This assay setup is exemplified by the Lumit® SARS-CoV-2 RBD:hACE2 Assay.
- (B) The Lumit® FcR Binding Immunoassays are examples of competitive loss-of-signal assays. In this setup, a biotinylated Fc receptor is combined with SmBiT-labeled streptavidin (SA-SmBiT) and LgBiT-labeled antibody tracers (Tracer-Ab-LgBiT). This approach measures the affinity of analyte antibodies (Ab-Analyte) for the Fc receptor through the competitive displacement of Tracer-Ab-LgBiT.



#### **Gain-of-Signal Binding Immunoassays**



Lumit® gain-of-signal assays are used to analyze the potency of protein:protein interaction inducers. In this format, two BiT-labeled primary antibodies bind the protein pair of interest. The presence of a PPI inducer facilitates the interaction between the two proteins, resulting in a relative increase in luminescent signal.

### 2.1 Cytokines / HMGB1

#### **Description & Application**

Pre-built Lumit® Cytokine/HMGB1 Immunoassays are fast, plate-based, add-and-read quantification assays. Their sensitivity is emphasized by a low limit of detection (LOD) while their wide linear range mitigates the need for sample dilutions. Lumit® assays can be applied for low- or high-throughput experiments.

#### **Principle & Workflow**

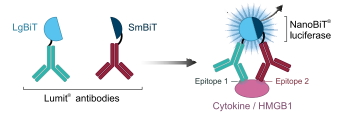
The assay kits contain analyte-specific primary antibodies, labeled with the NanoBiT® subunits LgBiT and SmBiT, an analyte standard, and detection reagent. Analyte detection within the cell culture supernatant is either performed in the presence of cells (Option #1) or upon transfer to a separate plate (Option #2). Binding of both antibodies to the analyte facilitates reconstitution of the NanoBiT® luciferase. Upon addition of detection reagent, a bright luminescent signal that is proportional to the analyte level can be recorded on a conventional plate-reading luminometer.

#### **Assay Features**

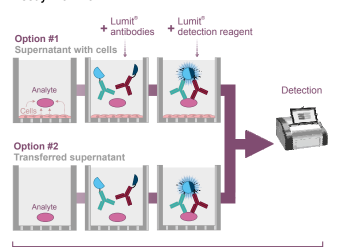
Sample material	Supernatant with cells Transferred supernatant
Sample volume	12.5 – 80 µl
Assay format	Direct assay format 96- / 384-well plates
Assay protocol	Homogeneous Add-and-read
Time required	70 minutes or less
Multiplexing option	Caspase-Glo® 1 Inflammasome Assay



#### **Assay Principle**



#### **Assay Workflow**



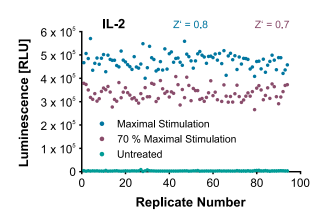
70 min

Assay	Dynamic range	LOD
Lumit® IL-2 (Human)	28.2 - 25000 pg/ml	11.2 pg/ml
Lumit® IL-4 (Human)	18.2 - 25000 pg/ml	6.7 pg/ml
Lumit® IL-6 (Human)	18.2 - 25000 pg/ml	7.5 pg/ml
Lumit® IL-10 (Human)	18.2 - 25000 pg/ml	7.4 pg/ml
Lumit® IFN-γ (Human)	7.2 - 10000 pg/ml	1.7 pg/ml
Lumit® TNF-α (Human)	18.2 - 25000 pg/ml	2.9 pg/ml
Lumit® IL-12 (Human)	18.2 - 25000 pg/ml	4.5 pg/ml
Lumit® IL-1β (Human)	22 - 40000 pg/ml	10 pg/ml
Lumit® IL-1β (Mouse)	11 - 20000 pg/ml	8 pg/ml
Lumit® HMGB1 Human/Mouse Immunoassay	4 - 1000 ng/ml (hu) 3 - 2187 ng/ml (ms)	1 ng/ml 3 ng/ml
Lumit® Active IL-18 (Human)	11 - 20000 pg/ml	≤ 10 pg/ml
Lumit® IL-8 (Human)	7.29 - 10000 pg/ml	1 pg/ml
Lumit® IL-17A (Human)	18.2 - 25000 pg/ml	3 pg/ml
Lumit® IFN-β (Human)	18.2 - 25000 pg/ml	5.3 pg/ml
Lumit® VEGF-A (Human)	18.2 - 25000 pg/ml	3 pg/ml

LOD: limit of detection (3 SD above background); hu: human; ms: mouse

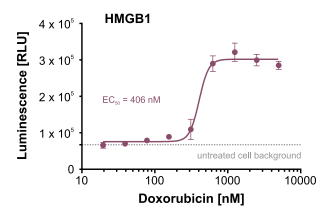
#### Validation of HTS-compatibility

Human PBMCs, plated at 10000 cells/well into a 384-well plate were treated with Cell Stimulation Cocktail for 24 hours at maximal or 70 % maximal levels of stimulation. For each condition, luminescence of 94 replicates was determined upon reagent addition. Z' factors determined for IL-2 release were substantially greater than 0.5, indicating amenability to screening applications.



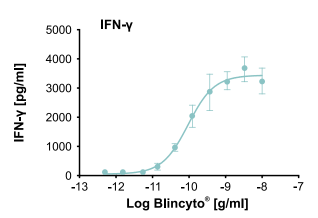
#### Drug-induced immunogenic cell death

Mouse EL4 cells were treated with doxorubicin for 24 hours. HMGB1 within the supernatant was quantified in the presence of cells (w/o transfer) by using Lumit® HMGB1 Immunoassay.



#### Detection of IFN-y as a marker of T cell activation

Purified CD8+T cells (effector cells) were combined with Raji B cells (target cells) and a serial dilution of Blincyto® (a CD3 and CD19 bispecific T cell engager). IFN-y release from effector cells into the cell culture supernatant was analyzed in the presence of cells (w/o transfer) by using Lumit® IFN-y Immunoassay.



#### **Product Box**

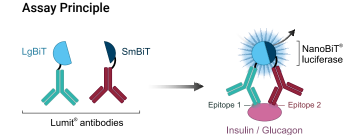
Lumit® Cytokine Immunoassays Cat.# see page 40
Lumit® HMGB1 Immunoassay Cat.# W6110, W6112



#### 2.2 Hormones

#### **Description & Application**

Pre-built Lumit® Insulin and Glucagon Immunoassays are fast, plate-based, add-and-read quantification assays that deliver results in approximately one hour. The wide linear range of these sensitive assays mitigates the need for sample dilution. Lumit® Immunoassays are amenable for low- and high-throughput applications.

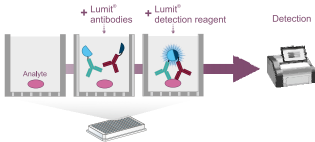


#### **Principle & Workflow**

The assay kits contain analyte-specific primary antibodies, labeled with the NanoBiT® subunits LgBiT and SmBiT, an analyte standard, and detection reagent. Analyte detection is performed in cell culture supernatant, transferred to a separate plate. Binding of both antibodies to the analyte facilitates reconstitution of the NanoBiT® luciferase. Upon addition of detection reagent, a bright luminescent signal that is proportional to analyte levels can be recorded on a conventional platereading luminometer.

**Assay Workflow** 

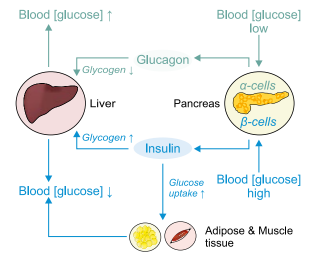
**Transferred supernatant** 



70 min

#### **Assay Features**

Sample material	rerial Transferred supernatant	
Sample volume	5 – 50 μΙ	
Species compatibility	Human, mouse, rat	
Assay format	Direct assay format 96- / 384-well plates	
Assay protocol	Homogeneous Add-and-read	
Time required	70 minutes or less	
Multiplexing option	Use both Lumit® assays and analyze insulin and glucagon side-by-side to gain more information on islet function	

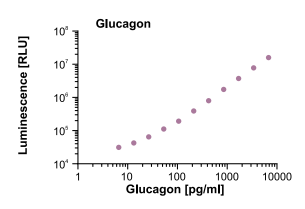


Assay	Dynamic range	LOD
Insulin	58 – 46000 pg/ml	58 pg / ml
Glucagon	3 – 7000 pg / ml	3 pg / ml

LOD: limit of detection

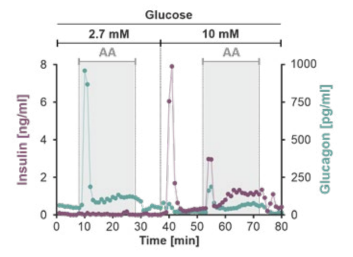
#### Broad dynamic range and picomolar sensitivity

The broad dynamic range enables sensitive detection without sample dilution. A dilution series of glucagon was detected in a 96-well plate by addition of Lumit® antibodies. After 1 hour of incubation, Lumit® detection reagent was added, and luminescence was measured. Analysis was performed in quadruplicates.



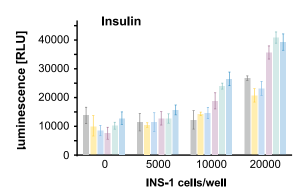
#### Measuring hormone secretion in perfusion experiments

In a perfusion chamber, 80 mouse islets were treated with glucose in combination with an amino acid (AA) mixture. Low and high blood glucose levels were simulated with 2.7 mM glucose and 10 mM glucose respectively. Perfusate aliquots were collected every minute. Insulin and glucagon levels were detected in 10  $\mu$ l perfusate in a 384-well plate. This data was kindly provided by Drs. H. Foster and M. Merrins (University of Wisconsin VA Hospital, Madison, WI).



#### Monitoring insulin secretion in response to glucose

INS-1 rat insulinoma cells plated at different cell numbers in 96-well plates were stimulated with glucose at different concentrations for 60 minutes. Insulin secretion was determined by removing 10  $\mu l$  of supernatant and assayed with the Lumit® Insulin Immunoassay Kit in 384-well plates.



#### **Product Box**

Lumit® Glucagon Immunoassay Lumit® Insulin Immunoassay Cat.# W8020, W8022

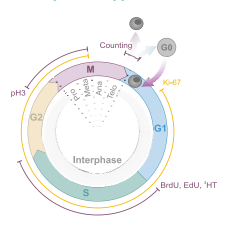
Cat.# CS3037A01, CS3037A05



### 2.3 Cell Proliferation

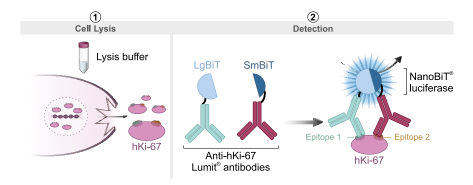
### Lumit® hKi-67 Immunoassay for Cell Proliferation

#### **Description & Application**



The Lumit® hKi-67 Immunoassay for Cell Proliferation is a no-wash, plate-based assay that detects hKi-67, a well-established nuclear marker of cell proliferation expressed in all active phases of the cell cycle and is absent in resting, non-dividing cells. This assay enables researchers to confidently track proliferation changes at early time points and can be completed in under two hours. By adopting this "add-mix-measure" bioluminescent assay, researchers can streamline their workflow, reduce prep work, and accelerate time-to-results-offering a more efficient and high-throughput-compatible alternative to traditional proliferation assays (e.g., BrdU, EdU, 3H-thymidine, metabolic activity assays, and cell counting).

#### **Assay Principle**

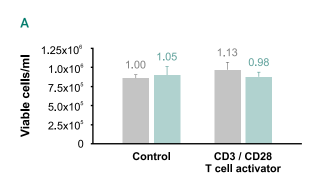


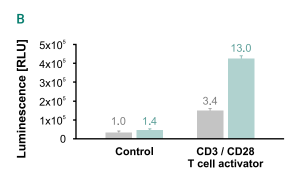
#### **Principle & Workflow**

The Lumit® hKi-67 Immunoassay utilizes primary antibodies against hKi-67, labeled with NanoBiT® luciferase subunits LgBiT and SmBiT. In the presence of hKi-67, the subunits reconstitute the functional NanoBiT® luciferase, producing a bright luminescent signal upon addition of the Lumit® Detection Substrate C. This signal is directly proportional to the hKi-67 levels, providing a sensitive and specific readout of cell proliferation. Prior to antibody addition, the nuclear hKi-67 protein is released by addition of a lysis buffer to the assay plate. Optionally. the assay can be multiplexed with the included CellTox™ Green Cytotoxicity Assay to identify antiproliferative effects without cell death.

#### **Assay Workflow**

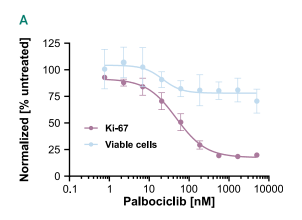


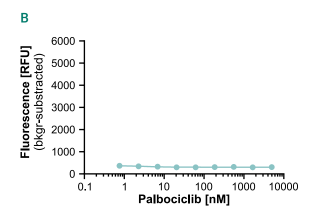




#### **Determination of mitogenic effects**

Human CD8+ T cells (STEMCELL Technologies™) were seeded at 80000 cells/well and treated with a CD3/CD28 T cell activator in the absence or presence of IL-2 (10 ng/ml) for 48 hours. The number of viable cells was determined by trypan blue staining (A) or analyzed using the Lumit® hKi-67 Immunoassay (B). hKi-67, an early marker of cell proliferation, showed a significant increase in response to T cell activation, while no changes in absolute cell number were observed.





#### **Determination of antiproliferative effects**

Jurkat cells (20000/well) were treated with a dilution series of the CDK4/6 inhibitor palbociclib for 24 hours. The hKi-67 level was assessed using the Lumit® hKi-67 Immunoassay (A, purple), while changes in the viable cell number were determined with the CyQUANT™ Direct Assay (Thermo Fisher Scientific) (A, blue) in a parallel plate. Cell death was monitored by multiplexing the Lumit® assay with the CellTox™ Green Cytotoxicity Assay (B). Without inducing cell death, palbociclib elicited an antiproliferative response, evident from a dose-dependent decrease in both the hKi-67 level and viable cell number. Compared to the DNA-stain-based measurement of viable cells, the hKi-67 assay provided a much more pronounced assay window, enabling an earlier readout with reduced incubation times.

**Product Box** 

Lumit® hKi-67 Immunoassay for Cell Proliferation

Cat.# CS3076A01



## 2.4 Second Messengers Lumit® cAMP Immunoassay

#### **Description & Application**

The Lumit® cAMP Immunoassay is a homogeneous, no-wash competition assay for measuring 3',5'-cyclic adenosine monophosphate (cAMP). This key second messenger regulates metabolism, gene expression, and signal transduction. cAMP is synthesized by adenylyl cyclase upon GPCR activation and hydrolyzed by phosphodiesterases (PDEs). The assay uses competitive binding to detect cAMP level changes with high specificity and sensitivity. It is ideal for studying GPCR signaling, PDE activity, and cAMP modulators in biochemical and cellular formats.

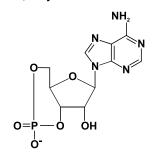
#### **Principle & Workflow**

The assay includes cAMP linked to SmBiT (Tracer cAMP SmBiT), a cAMP-specific primary antibody, and an anti-mouse antibody conjugated to LgBiT. These components form a luminescent complex via NanoBiT® luciferase reconstitution. cAMP binding displaces the tracer in a dose-dependent manner, reducing luminescence. Enzymatic reactions are stopped (and cells lysed) by adding 2 % TCA. After a 30-minute incubation with the antibody-tracer mixture, the Lumit® Detection Reagent is added, and luminescence is measured on a plate reader.

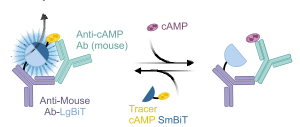
#### **Assay Features**

Sample material	Purifed enzyme, cells
Sample volume	12 – 120 µl
Assay format	Competitive assay format 96- / 384-well plates
Assay protocol	Homogeneous Add-and-read
Time required	< 1 hour
Dynamic range	1.37 – 132 nM
LOD	0.325 nM

#### 3',5'-cyclic adenosine monophosphate (cAMP)



#### **Principle**



#### **Assay Workflow**

# Treated sample + 2% TCA + Antibody Tracer Mixture @RT 30 min + Lumit® **Detection Reagen** @RT 3 min Detection

#### **Assay Specificity**

Serial dilutions of cAMP and other nucleotides (cGMP, 3'3'cGAMP, and 2'3'cGAMP) were analyzed for their ability to compete with the cAMP tracer of the Lumit® cAMP Immunoassay. Except for cAMP, none of the tested nucleotides yielded a signification decrease in luminescence emphasizing assay specificity.

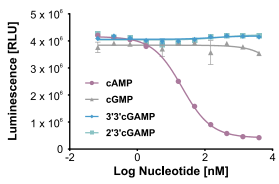
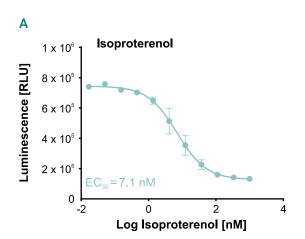
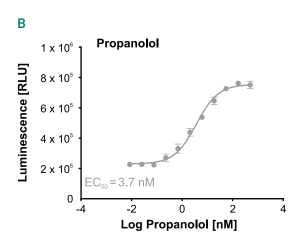


Figure adapted from Mikheil et al. Scientific Reports. 2024;14(1):4440





#### **Antagonist Competition of Agonist-Induced cAMP Response**

A375 cells, expressing endogenous  $\beta$ 2-adrenergic receptors, were seeded at 1000 cells/well in 384-well plates. The following day, cells were treated with a concentration series of (A) the  $\beta$ -adrenergic receptor agonist isoproterenol alone, or (B) the antagonist propranolol in combination with 50 nM isoproterenol. After a 5-minute incubation at room temperature, the reactions were stopped with 2% trichloroacetic acid (TCA), and cAMP levels were measured using the Lumit® cAMP Immunoassay.



# 2.4 Second Messengers Lumit® cGAMP Immunoassay

#### **Description & Application**

The Lumit® cGAMP Immunoassay is a homogeneous, no-wash competition assay for measuring 2',3'-cGAMP. This cyclic dinucleotide, composed of guanine and adenine monophosphate, activates the STING pathway, driving pro-inflammatory cytokine production in innate immunity. cGAMP is synthesized by cGAS upon recognition of cytosolic double-stranded DNA (dsDNA). The assay enables the identification and characterization of cGAMP modulators in biochemical and cellular formats.

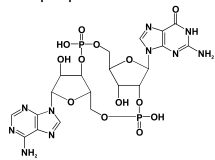
#### **Principle & Workflow**

The assay includes 2',3'-cGAMP linked to SmBiT (Tracer cGAMP SmBiT), a mouse-derived anti-cGAMP antibody, and an anti-mouse antibody conjugated to LgBiT. These components form a bioluminescent complex via NanoBiT® luciferase reconstitution. cGAMP binding displaces the tracer in a dose-dependent manner, reducing luminescence. Enzymatic reactions are stopped (and cells lysed) by adding 0.5 N HCl. After a 90-minute incubation with the antibody tracer mixture, the Lumit® Detection Reagent is added, and luminescence is measured on a plate reader.

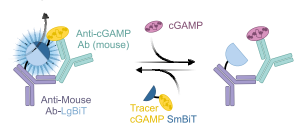
#### **Assav Features**

Adday i catalco	
Sample material	Purified enzyme, cells,
	human serum
Sample volume	4 – 40 µl
Assay format	Competitive assay format
	96- / 384-well plates
Assay protocol	Homogeneous
	Add-and-read
Time required	2 hours
Dynamic range	< 1 nM - 10000 nM
LOD	0.6 nM

# 2',3'- cyclic guanosine monophosphate-adenosine monophosphate

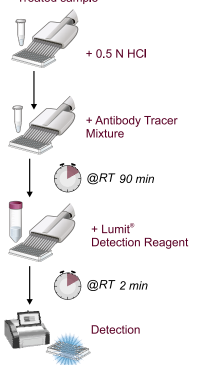


#### **Principle**



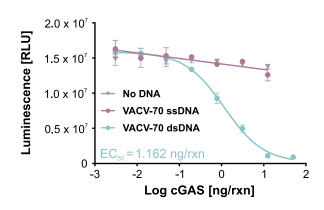
#### **Assay Workflow**

#### Treated sample



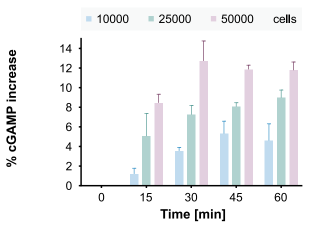
#### Biochemical assay of cGAS activity

cGAS reactions (40  $\mu$ I) were carried out in presence of ATP (100  $\mu$ M), rGTP (100  $\mu$ M), and VACV-70 DNA (1 ng). After incubation for 30 minutes at 37°C the reaction was terminated with 0.5 N HCl. The cGAMP level was determined by using the Lumit® cGAMP Immunoassay. Activation of of cGAS requires dsDNA while as no activation was detected with ssDNA and the no DNA control.



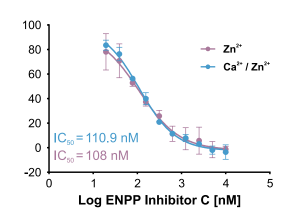
#### Monitoring cGAMP levels in RAW 264.7 cells

Different numbers of RAW 264.7 cells were treated with ISD dsDNA (4  $\mu g/ml$ ) to stimulate endogenous cGAS activity in the presence of ENPP1 Inhibitor C (60  $\mu M$ ). Reactions were terminated at indicated time points using 0.5 N HCl and cGAMP levels analyzed with the Lumit  $^{\circ}$  cGAMP Immunoassay



#### Biochemical assay of ENPP1 activity/inhibition

Enzymatic ENPP1 (10 ng) reactions (40  $\mu$ l) were carried out in presence of Zn²+ (5  $\mu$ M) and cGAMP (10 nM) with and without Ca²+ (0.5 mM) at 37 °C. After 30 minutes, 0.5 N HCl was added to stop the reaction and cGAMP levels were quantified using the Lumit® cGAMP Immunoassay.



Figures adapted from Hsiao et al. Scientific Reports. 2024;14:31165

#### **Product Box**

Lumit® cGAMP Immunoassay

Cat.# CS3592A05

% Activity



# 2.5 Cell & Gene Therapy Lumit® dsRNA Detection Assay

#### **Description & Application**

The Lumit® dsRNA Detection Assay detects and quantifies double-stranded RNA (dsRNA) contamination in *in vitro* transcription (IVT) mRNA samples, which can impact RNA therapeutic development. This assay is specific, showing no cross-reactivity with single-stranded RNA (ssRNA) or DNA, and works with dsRNA as short as 30 base pairs. The assay generates a luminescent signal proportional to the dsRNA concentration. It offers a simple, washfree workflow, suitable for high-throughput applications. Compared to traditional ELISA kits, the Lumit® dsRNA Detection Assay provides better sensitivity, accurately quantifying dsRNA in a range of 0.04 – 2.5 ng/ml.

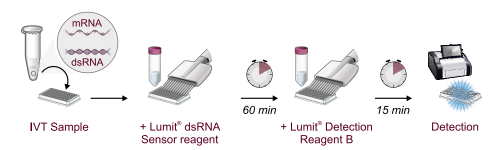
#### **Assay Principle**



#### **Principle & Workflow**

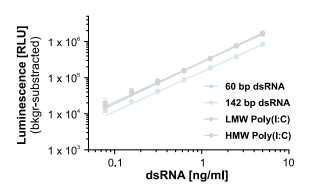
The Lumit® dsRNA Detection Assay utilizes recombinant dsRNA-binding proteins fused to the SmBiT and LgBiT subunits of NanoBiT® luciferase, respectively. In the presence of dsRNA, the two subunits reconstitute an active luciferase whose luminescent signal correlates with the dsRNA concentration. To perform the assay, dsR-NA-containing samples are mixed with the Lumit® reagents in a homogeneous, solution-based format. No wash steps or immobilization are required. The resulting luminescent signal is proportional to the dsRNA content and can be easily measured using a conventional plate luminometer, making the workflow simple and high-throughput compatible.

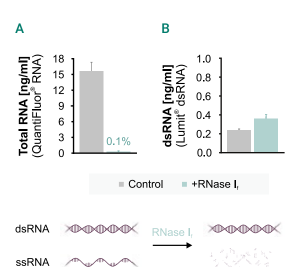
#### **Assay Workflow**

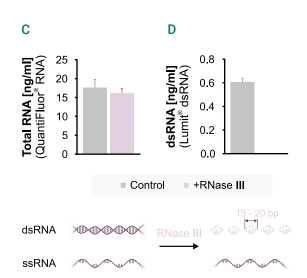


#### Independence of sequence and size

Titrations of different dsRNAs that vary in size and sequence were analyzed with the Lumit® dsRNA Detection Assay. The data shows that the assay can sensitively quantify dsRNA independent of sequence and size. The following dsRNAs were used: (1) 60 bp dsRNA (synthesized dsRNA comprised of the first 60 bp from the SARS-CoV-2 spike ORF), (2)142 bp dsRNA (commercially available "standard"), (3) LMW Poly(I:C) (synthetic dsRNA analog; 0.2 – 1 kb), (4) HMW Poly(I:C) (synthetic dsRNA analog; 1.5 – 8 kb).







#### Quantification of dsRNA in IVT samples with mixed RNA species

In vitro transcription (IVT) samples were treated with either RNase If (ssRNA-specific) or RNase III (dsRNA-specific). Total RNA and dsRNA were quantified using the QuantiFluor® RNA System and the Lumit® dsRNA Detection Assay. Digestion of ssRNA with RNase If significantly reduced the total RNA but did not affect the amount of dsRNA detected (A and B), while digestion of dsRNA with RNase III into fragments ≤20 bp resulted in a complete loss of signal in the Lumit® dsRNA Detection Assay without significantly affecting total RNA levels (C and D).

#### **Product Box**

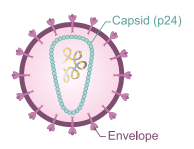
Lumit® dsRNA Detection AssayCat.# W2041, W2042Lumit® dsRNA Standard, 5-methylcytidineCat.# CS355716Lumit® dsRNA Standard, N1-methylpseudouridineCat.# CS355712Lumit® dsRNA Standard, PseudouridineCat.# CS355710

Lumit® dsRNA Standard, 5-methoxyuridine Cat.# CS355714



# 2.5 Cell & Gene Therapy Lumit® p24 Immunoassay

#### **Description & Application**

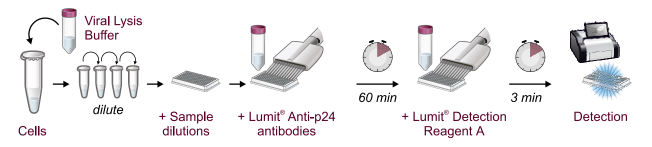


Lentiviruses are a widely used research tool because of their ability to deliver genetic material into both dividing and non-dividing cells. Accurate titer determination is essential to ensure consistent and reliable experimental outcomes, as viral dose influences transduction efficiency and biological effects. The lentiviral capsid protein p24 is a commonly measured marker for viral titer determination and its amount directly correlates with viral particle number.

#### **Principle & Workflow**

The Lumit® p24 Immunoassay quantifies p24 in a plate-based, washing-free, add-and-read format. The assay uses two anti-p24 antibodies with different epitope specificities that are labeled with either LgBiT or SmBiT. Binding of both antibodies to soluble p24 reconstitutes the functional NanoBiT® luciferase. Lentiviral samples are lysed with Viral Lysis Buffer and serially diluted. Upon addition of the anti-p24 Lumit® antibody mixture the samples are incubated for 60 minutes at room temperature to enable antibody binding. Following addition of the Lumit® Detection Reagent A, the luminescent signal is recorded on a conventional plate-reading luminometer.

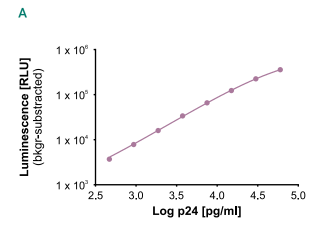
#### **Assay Workflow**

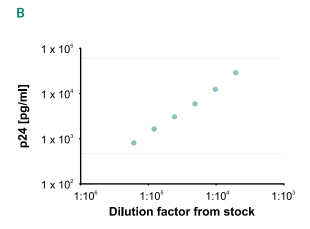


#### **Assay Features**

Sample material	Cell culture supernatant and purified p24 protein	
Sample volume	10 – 20 µl	
Assay format	Gain-of-signal assay 96- / 384-well plates	
Assay protocol	Homogeneous Add-and-read	

Time required	90 minutes
Dynamic range	2 – 60000 pg/ml
LOD	3 pg/ml





#### **Lentiviral Titer Assessment**

A two-fold dilution series of the p24 standard (A) and a lentivirus sample (B) was prepared in Viral Lysis Buffer and analyzed using the Lumit® p24 Immunoassay. The standard curve (n = 2) was fitted to a 4-parameter logistic regression model and used to calculate the concentrations of the lentivirus samples. Average concentrations and standard deviations of the sample dilutions within the standard curve range (n = 4) are shown. Dashed lines indicate the upper and lower boundaries of the standard curve.

#### **Product Box**

Lumit® p24 Immunoassay

Cat.# CS2039B25







### One Platform. Endless Possiblities.

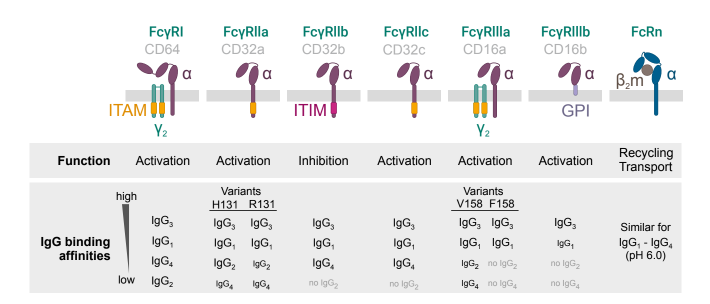
- · Luminescence, fluorescence, absorbance
- High sensitivity and broad dynamic range
- · Kinetic measurements and FRET/BRET capabilities
- · Intuitive interface with onboard data analysis

For more information visit: www.promega.com/glomax-comparison



### 2.6 Therapeutic Antibody Development

The efficacy of therapeutic antibodies depends not only on the Fab fragment's binding to the target antigen but also on interactions between the Fc fragment and Fc receptors. For example, the Fc fragment's affinity for the neonatal Fc receptor (FcRn) influences antibody half-life, while its interaction with Fc gamma receptors (FcγR) determines its ability to trigger effector functions such as antibody-dependent cellular cytotoxicity (ADCC) and antibody-dependent cellular phagocytosis (ADCP). To ensure optimal function, candidate therapeutic antibodies should be evaluated for their Fc receptor interactions during drug development.



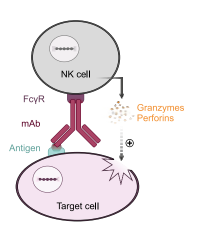
Representation of IgG-binding Fc receptors (Fc $\gamma$ R, FcRn). Binding affinity varies among IgG subclasses. ITAM = immunoreceptor tyrosine-based activation motif; ITIM = immunoreceptor tyrosine-based inhibition motif;  $\gamma$ 2 = FcR gamma subunit dimer;  $\beta_2$ m = beta-2 macroglobulin. Adapted from Bruhns, P. (2012) Blood 14;119(24):5640-9; Smith, KGC. (2010) Nat Rev Immunol May; 10(5):328-43; Hogarth, PM. (2012) Nat Rev Drug Discov Mar 30;11(4):311-31.

The neonatal Fc receptor (FcRn) is an atypical Fc receptor expressed in diverse tissues, including epithelial, endothelial, and hematopoietic cells. It binds the Fc region of IgG at acidic pH within endosomes, facilitating IgG transport across polarized barriers (e.g., epithelia) and protecting IgG from degradation, thereby extending its serum half-life. Optimizing FcRn interaction with therapeutic IgG is a critical factor in drug development, as extended half-life enhances efficacy and allows for longer dosing intervals.

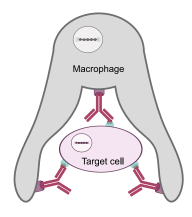
Fc gamma receptors (FcγR) play a key role in immune responses by binding to the Fc region of IgG. These interactions mediate processes such as ADCC, endocytosis, phagocytosis, inflammatory mediator release, and enhanced antigen presentation. In humans, three classes of FcγRs—FcγRI (CD64), FcγRIIa/b/c (CD32a/b/c), and FcγRIIIa/b (CD16a/b)—have been identified on immune cells. FcγRI is a high-affinity receptor (nM range), while FcγRII and FcγRIII exhibit low to intermediate affinity (μM range). Genetic variations in Fc receptors significantly influence Fc-mediated immune responses. For instance, the FcγRIIIa polymorphism V158 exhibits higher IgG binding affinity compared to the F158 variant, impacting ADCC activity in NK cells.

FcyR- / FcRn-mediated Cellular Functions

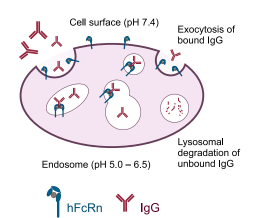
#### FcyR-mediated ADCC



#### FcyR-mediated ADCP



#### FcRn-mediated IgG recycling



FcyR-mediated ADCC: NK cells trigger antibody-dependent cellular cytotoxicity (ADCC) upon binding the Fc region of IgG via FcyR. This activation leads to the release of cytotoxic molecules, resulting in target cell death.

FcyR-mediated ADCP: Macrophages initiate antibody-dependent cellular phagocytosis (ADCP) upon engaging the Fc region of IgG, enhancing their ability to engulf and degrade target cells.

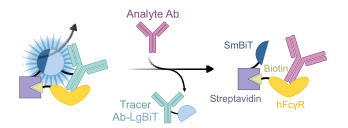
FcRn-mediated IgG recycling: FcRn binding occurs in acidic endosomal conditions, promoting IgG uptake via pinocytosis. Unbound IgG is degraded in Iysosomes, whereas FcRn-bound IgG is recycled and released through exocytosis.

# 2.6 Therapeutic Antibody Development Lumit® FcyR Binding Immunoassays

#### **Description & Application**

The Lumit® FcγR Binding Immunoassays are novel homogeneous, no-wash competition assays to measure the interaction between human Fc receptors and antibodies or Fc fusion proteins. Importantly, the in-solution format circumvents experimental artifacts caused by immobilization. These assays are used in therapeutic antibody development for antibody optimization and for testing antibody potency.

#### **Assay Principle**



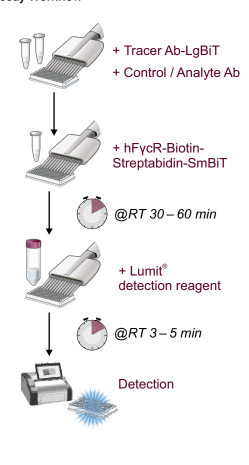
#### **Principle & Workflow**

FcγR assays consist of a LgBiT-labeled human IgG1 (Tracer Ab-LgBiT) and a biotinylated human FcγR (extracellular domain) bound to SmBiT-labeled streptavidin (hFcγR-Biotin-Streptavidin-SmBiT). In the absence of an analyte antibody (Analyte Ab), tracer binding to labeled hFcγR results in maximum luminescent signal. Binding of analyte antibodies is evident from a concentration-dependent decrease in luminescence due to competition with the tracer. These easy-to-use biochemical assays can complement and provide orthogonal data to support results from cell-based functional bioassays.

#### **Assay Features**

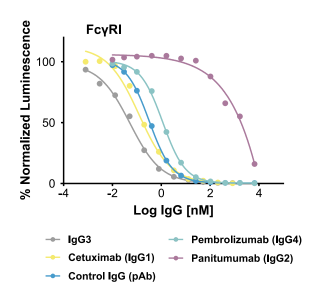
Sample material	Antibodies Fc proteins	
Sample volume	25 μl antibody	
Concentration range	4 – 4000 ng/ml	
Assay format	Loss-of-signal assay 96- / 384-well plates	
Assay protocol	Homogeneous Add-and-read	
Time required	70 minutes or less	

#### **Assay Workflow**



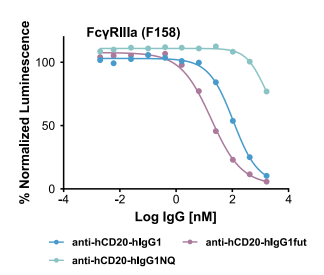
#### Lumit® FcγRI Binding Immunoassay

Lumit® Fc $\gamma$ RI Binding Immunoassays are used as potency assays to complement cell-based functional Fc effector activity assays. Fc $\gamma$ RI binds to IgG in a subclass-specific manner with IC $_{50}$  values reflective of relative IgG affinity (IgG $_3$  > IgG $_1$  > IgG $_4$  >>> IgG $_2$ ).



#### Lumit® FcγRIIIa (F158) Binding Immunoassay

Lumit® Fc $\gamma$ R Binding Immunoassays are used to assess antibody glycan status. IC $_{50}$  shifts in non-fucosylated (anti-hCD20-hIgG1fut) or non-glycosylated (anti-hCD20-hIgG1NQ) antibodies were detected.



#### **Product Box**

Lumit® FcyRI Binding Immunoassay

W7080, W7081

Lumit® FcγRIIa (H131) Binding Immunoassay

W7070, W7071

Lumit® FcγRIIa (R131) Binding Immunoassay

W7060, W7061

Lumit® FcyRIIb Binding Immunoassay

W7030, W7031

**Lumit® FcγRIIIa (F158) Binding Immunoassay** W7040, W7041

umit® FcvRIIIa (V158) Binding Immun

**Lumit® FcγRIIIa (V158) Binding Immunoassay** W7050, W7051

**Lumit® FcγRIIIb Binding Immunoassay** 

W7020, W7021



# 2.6 Therapeutic Antibody Development Lumit® FcRn Binding Immunoassay

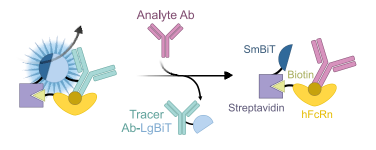
#### **Description & Application**

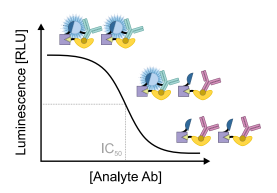
The Lumit® FcRn Binding Immunoassay is a homogeneous, no-wash competition assay to measure the interaction between human neonatal FcRn and Fc proteins, including antibodies. Importantly, the in-solution format circumvents experimental artifacts caused by immobilization. This assay is used in therapeutic antibody development to assess and tune the half-life of antibodies due to optimized binding to FcRn. In addition, it is used for determining antibody oxidation status and for detecting anti-FcRn blocking antibodies.

#### **Principle & Workflow**

The assay consists of a LgBiT-labeled human IgG1 (Tracer Ab-LgBiT) and a biotinylated human FcRn (extracellular domain) bound to SmBiT-labeled streptavidin (hFcRn-Biotin-Streptavidin-SmBiT). In the absence of an analyte antibody (Analyte Ab), tracer binding to labeled hFcRn results in maximum luminescent signal. Binding of analyte antibodies is evident from a concentration-dependent decrease in luminescence due to competition with the tracer.

#### **Assay Principle**





#### **Assay Features**

•		
Sample material	Antibodies Fc proteins	
Sample volume	25 μl antibody	
Concentration range	4 – 4000 ng/ml	
Assay format	Loss-of-signal assay 96- / 384-well plates	
Assay protocol	Homogeneous Add-and-read	
Time required	70 minutes or less	

#### References

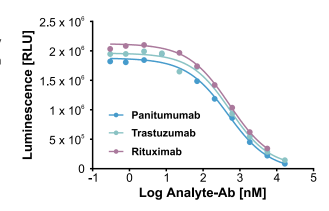
Nath, N. et al. (2021) Deciphering the interaction between neonatal Fc receptor and antibodies using a homogeneous bioluminescent immunoassay.

#### J Immunol. 207(4), 1211-1221

Tian, Z. et al. (2021) Harnessing the power of antibodies to fight bone metastasis. Sci Adv. 7(26), eabf2051

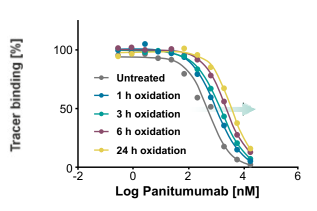
#### FcRn binding to a panel of therapeutic antibodies

A panel of therapeutic antibodies were tested for their affinity to FcRn by using the Lumit® FcRn Binding Immunoassay. An excellent assay window is observed for IgG/FcRn binding.



#### Oxidation-based loss in antibody-FcRn affinity

A therapeutic antibody was incubated with  $0.3\,\%\ H_2O_2$  for 1–24 hours inducing methionine oxidation. Dose-dependent, oxidation-based loss in antibody-FcRn affinity was readily detected by using the Lumit® FcRn Binding Immunoassay.



#### **Product Box**

Lumit® FcRn Binding Immunoassay

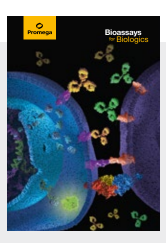
Lumit® Canine FcRn Binding Immunoassay

Lumit® Feline FcRn Binding Immunoassay

Cat.# W1151, W1152 Cat.# CS3019A02

Cat.# CS3631B01, CS3631B05





# Interested in Cell-Based Reporter Bioassays for Biologics Characterization?

Find more information about mechanism of action (MOA) assays such as ADCC and ADCP in Promega's Bioassay Guide for biologics characterization and lot-release.

www.promega.com/BioassaysForBiologics



# 2.6 Therapeutic Antibody Development Lumit® IgG Titer Immunoassay

#### **Description & Application**

The Lumit® IgG Titer Immunoassay is a homogeneous, no-wash assay designed for rapid and accurate quantification of monoclonal antibodies (mAbs). Traditional methods like ELISA and HPLC require multiple washing and incubation steps, limiting throughput and efficiency.

By utilizing a simple add-mix-measure workflow, the Lumit® IgG Titer Assay provides high sensitivity, broad dynamic range, and excellent reproducibility for IgG quantification across all IgG subclasses and other immunoglobulin types (e.g., IgA, IgM, and IgE). This makes it ideal for bioprocess development, therapeutic antibody research, and large-scale screening applications.

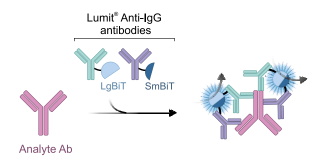
#### **Principle & Workflow**

The Lumit® IgG Titer Immunoassay contains two anti-IgG antibodies, pre-labelled with the NanoBiT® sub-units LgBiT and SmBiT that recognize immunoglobulins from different species (human, mouse, rabbit). Binding of both antibodies to the analyte IgG antibody facilitates reconstitution of the NanoBiT® luciferase. Upon addition of the detection reagent, a bright luminescent signal that is proportional to the analyte level can be recorded on a conventional plate-reading luminometer.

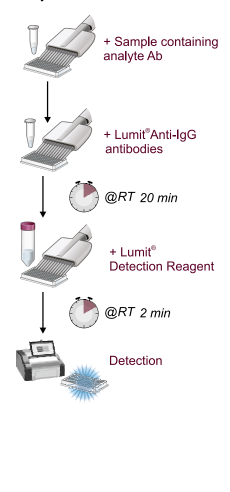
#### **Assay Features**

Sample material	Cell culture supernatant, purified IgG
Sample volume	5 μl (96-well) 1 – 2.5 μl (384-well) antibody
Assay format	Gain-of-signal assay 96- / 384-well plates
Assay protocol	Homogeneous Add-and-read
Time required	20 minutes
Dynamic range	1.6 (hu) / 3.4 (mu) / 4 (rb) – 1000 ng/ml
LOD	0.4 (hu) / 0.9 (mu) / 0.9 (rb) ng/ml

#### **Assay Principle**

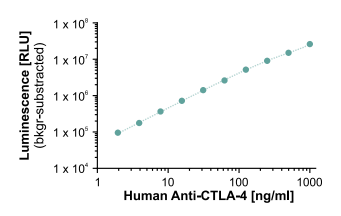


#### **Assay Workflow**



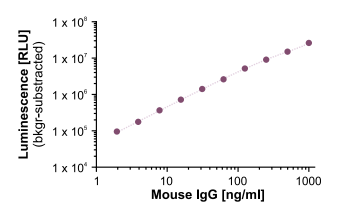
#### Human IgG standard curve

A two-fold dilution series of a human anti-CTLA-4 antibody was performed (1000 – 1.95 ng/ml) and analyzed using the Lumit® Human IgG Titer Immunoassay. The standard curve was generated using GraphPad® Prism by non-linear regression analysis with a fourth polynomial curve fitting model.



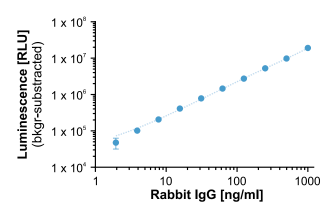
#### Mouse IgG standard curve

A two-fold dilution series of a mouse IgG antibody was performed (1000 – 1.95 ng/ml) and analyzed using the Lumit Mouse IgG Titer Immunoassay. The standard curve was generated using GraphPad Prism by non-linear regression analysis with a fourth polynomial curve fitting model.



#### Rabbit IgG standard curve

A two-fold dilution series of a mouse IgG antibody was performed ( $1000-1.95\,\text{ng/ml}$ ) and analyzed using the Lumit® Rabbit IgG Titer Immunoassay. The standard curve was generated using GraphPad® Prism by non-linear regression analysis with a fourth polynomial curve fitting model.



#### **Product Box**

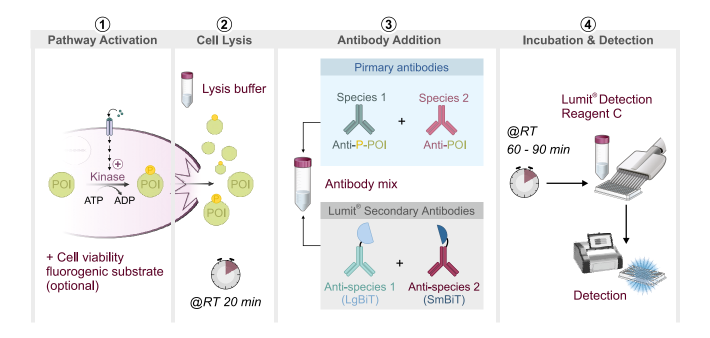
Lumit® Human IgG Titer Immunoassay Lumit® Mouse IgG Titer Immunoassay Lumit® Rabbit IgG Titer Immunoassay Cat.# CS3716A07, CS3716A13 Cat.# CS3716A03, CS3716A10 Cat.# CS3716A09, CS3716A14



# 2.7 Signaling & Protein Degradation Lumit® Immunoassay Cellular Systems

#### **Description & Application**

The Lumit® Immunoassay Cellular Systems have been extensively validated for analyzing kinase signaling pathways and targeted protein degradation in cell lysates. The complete kits include two different lysis buffers, analyte-specific primary antibodies, matched pre-labeled Lumit® Secondary Antibodies, and detection reagent. Kits containing only secondary antibodies and detection reagents are also available, allowing detection of analytes with your chosen primary antibodies. A growing list of pre-evaluated target proteins is available (see page 33), along with downloadable application notes detailing the primary antibodies used.



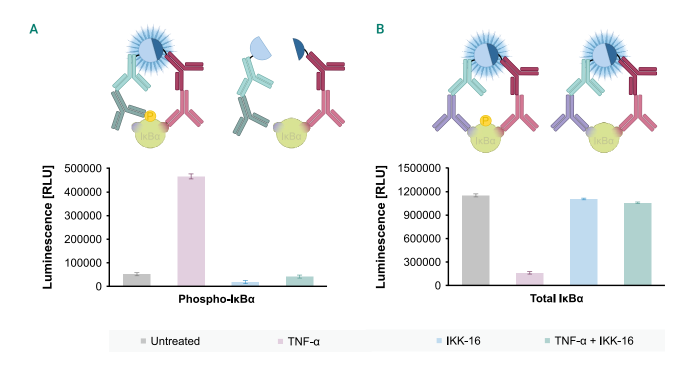
#### **Principle & Workflow**

The Lumit® Immunoassay Cellular Systems are based on the indirect Lumit® detection format in which two primary antibodies are mixed with two matching secondary antibodies that were pre-labeled with either the LgBiT or SmBiT subunit. To date, this format has been extensively validated for cell signaling analysis in cell lysates with a focus on phosphorylation as well as for measuring targeted protein degradation. The experimental workflow starts with (1) the treatment of cells to activate the signaling pathway of interest. At this point, the included cell viability fluorogenic substrate can be applied if normalization to viable cell number is desired. (2) Cells are lysed in-well by addition of a NanoBiT® compatible lysis buffer (Digitonin or Lumit® Lysis Buffer II). (3) Following addition of the antibody mix (primary and Lumit® Secondary Antibodies) and (4) incubation at room temperature for 60 to 90 minutes, the luminescent signal of the assay is determined by addition of the Lumit® detection reagent.

"Unmatched simplicity and reliability, with unbelievably fast results."

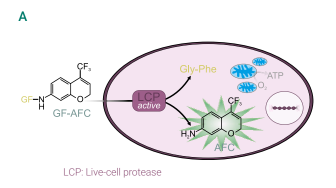
Scientist at Arctoris

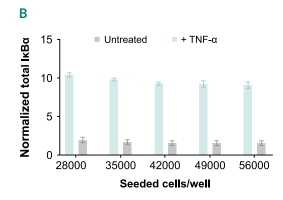
#### **Representative Data Signaling**



#### Detection of phosphorylated and total proteins within the NFkB signaling pathway

MCF-7 cells were seeded at 50000 cells/well and treated with TNF- $\alpha$  (20 ng/ml) for 30 minutes to induce the NF $\kappa$ B pathway. Cells in specificity control wells were pretreated with an IKK complex specific inhibitor, IKK16 (10  $\mu$ M, 1 hour). Subsequently, (A) phosphorylated I $\kappa$ B $\alpha$  (S32) or (B) total I $\kappa$ B $\alpha$  levels were determined in separate wells using two sets of primary antibodies.





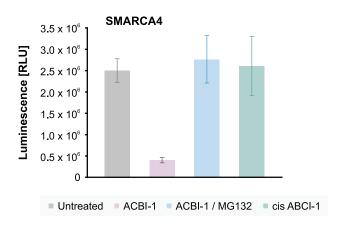
#### Normalization of luminescence data to viable cell number per well

Different cell numbers were treated with 50 ng/ml TNF- $\alpha$  for 30 minutes. (A) Cell viability fluorogenic substrate GF-AFC was added to all cells 30 minutes before lysis. This assay is based on the proteolytic processing of GF-AFC by Live-cell protease within viable cells to form a fluorescent product. Following cell lysis, an antibody mixture to detect total IkB $\alpha$  was added. (B) Luminescence and fluorescence were read at the end of the experiment. Normalization of Lumit® Immunoassay data to cell viability allows to easily account for well-to-well differences in cell number.

#### Representative Data Targeted Protein Degradation

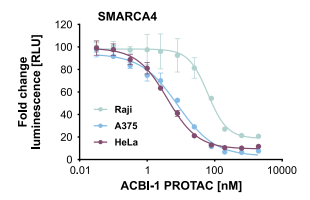
#### Degradation of SMARCA4 upon PROTAC® treatment

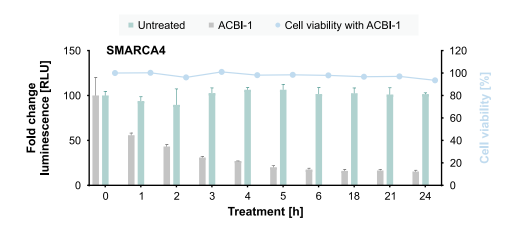
A375 cells (50000 cells/well) were treated with a single dose of ACBI-1 (250 nM) for 5 hours and targeted degradation of native SMARCA4 was assessed using the Lumit® Total SMARCA4 Immunoassay Cellular System. No degradation was observed when the stereoisomer cis ACBI-1 (250 nM) was used or when cells were co-treated with the proteasomal inhibitor MG132 (20  $\mu$ M).



#### Cell line-dependent SMARCA4 degradation profiles

Degradation potency of the PROTAC ACBI-1 was determined in three different cell lines, i.e., Raji, A375, and HeLa. Cells were seeded (50000 cells/well) and treated with a dilution series of the PROTAC ACBI-1 for 24 hours. Different  $DC_{50}$  values were observed for Raji (62.9 nM), A375 (8.0 nM), and HeLa (3.9 nM) cells.





#### Time course analysis of PROTAC-induced protein degradation of SMARCA4

Targeted degradation of native SMARCA4 in A375 cells (50000 cells/well) in the presence of ACBI-1 (100 nM) was monitored every 60 minutes over a period of 24 hours. Cell viability was determined by multiplexing with the fluorogenic substrate GF-AFC.

#### **Pre-validated Targets**

A list of application notes for pre-validated targets is available to save you time and labor. These notes contain target-specific protocols, representative data, and information on the commercially available antibodies used. The following table lists pre-validated targets for which no Complete Assays are available.

Cellular Target	Total protein	Phospho-Protein	Set ID	Lysis Buffer
Phospho-4E-BP1		Yes (Ser 65)	Set 2	Digitonin lysis solution
AKT	Yes		Set 1	Lumit® Lysis Buffer II
B-cell lymphoma 6 protein (BCL-6)	Yes		Set 1	Digitonin lysis solution
β-catenin (human)	Yes	Yes (Thr 41/Ser 45)	Set 2	Digitonin lysis solution
c-Jun		Yes (Ser 63)	Set 1	Digitonin lysis solution
c-MET		Yes (Tyr 1349)	Set 1	Lumit® Lysis Buffer II
CREB	Yes	Yes (Ser 133)	Set 1	Digitonin lysis solution
EGFR		Yes (Tyr 1173)	Set 1	Lumit® Lysis Buffer II
ER (Estrogen Receptor)	Yes		Set 1	Digitonin lysis solution
GSK1-3B		Yes (Ser 9)	Set 1	Lumit® Lysis Buffer II
HER2		Yes (Tyr 1196) and (Tyr 1221/1222)	Set 1	Lumit® Lysis Buffer II
p65	Yes		Set 2	Lumit® Lysis Buffer II
Rb (Retinoblastoma Tumor Suppressor)	Yes	Yes (Ser 780)	Set 2	Digitonin lysis solution
Ribosomal Protein S6	Yes		Set 1	Lumit® Lysis Buffer II
Smad1	Yes	Yes (Ser 463/465)	Set 1	Digitonin lysis solution
Smad2	Yes	Yes (Ser 465/467)	Set 2	Digitonin lysis solution
STAT1	Yes	Yes (Ser 727) and (Tyr 701)	Set 1	Digitonin lysis solution
Phospho-STAT2		Yes (Tyr 690)	Set 1	Digitonin lysis solution

Visit our website to browse the most recent application notes!

#### www.promega.com/LumitCellularSystems

#### **Product Box** Lumit® Secondary Antibodies of Set 1 and Set 2 Lumit® Immunoassay Cellular System - Starter Kit Cat.# W1220 Lumit® Anti-Mouse Ab-LgBiT Lumit® Immunoassay Cellular System - Set 1 Lumit® Anti-Rabbit Ab-SmBiT Cat.# W1201, W1202, W1203 Lumit® Anti-Mouse Ab-SmBiT Lumit® Immunoassay Cellular System - Set 2 Lumit® Anti-Rabbit Ab-LgBiT Cat.# W1331, W1332, W1333 Primary antibodies with matching Lumit® Immunoassay Cellular System Complete Assays Lumit® Secondary Antibodies Cat.# see page 44

# 3. Build-Your-Own Lumit® Immunoassays

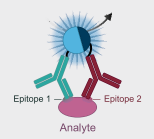
## 3.1 Lumit® Immunoassay Toolbox

Explore in chapter 3 how the Lumit® toolbox supports you in building an assay tailored to your needs!

#### **Antibody / Protein Labeling Kit**

Set up your own Lumit® assay by rapidly and efficiently labeling antibodies/proteins using the Lumit® Immuno-assay Labeling Kit.

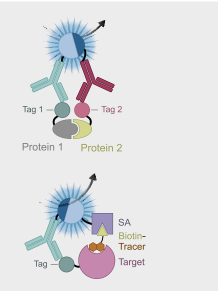
see page 35



#### Pre-Labeled Anti-Tag Antibodies / Streptavidin

Build your own protein:protein or protein:small molecule interaction assay using Lumit® anti-tag antibodies (for His-, GST-, FLAG®-, and human Fc-tagged proteins) and Lumit® Streptavidin (for AviTag®-tagged and biotinylated proteins).

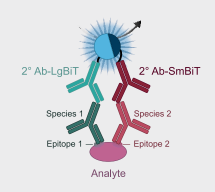
see page 36



#### **Pre-Labeled Secondary Antibodies**

Create an indirect Lumit® Immunoassay by using prelabeled secondary antibodies against IgGs from mouse, rabbit, and goat.

see page 38



#### **Detection Reagents**

Choose between different detection reagents optimized for distinct applications, e. g., biochemical or cell-based assays.

see page 39



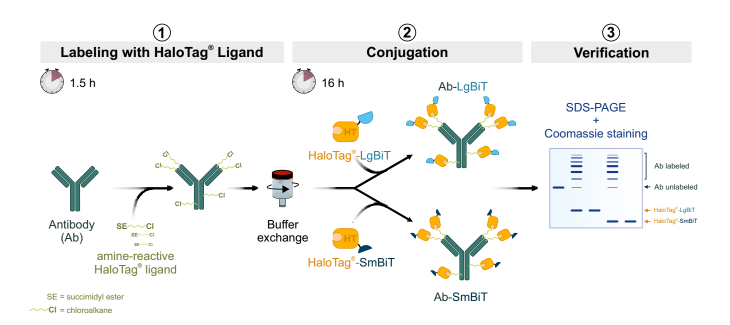
## 3.2 Lumit® Immunoassay Labeling Kit

#### **Description & Application**

The labeling kit is designed to conjugate antibodies / proteins to SmBiT and LgBiT supporting the development of your own Lumit® Immunoassay.

#### **Principle & Workflow**

The chemical labeling reaction is based on the HaloTag® technology. HaloTag® is a protein that covalently binds chloroalkane ligands (HaloTag® ligand) under physiological conditions and is used in a variety of applications, including antibody labeling. Labeling is a two-step process in which amine-reactive HaloTag® Succinimidyl Ester (O4) ligand reacts with primary amines of lysine amino acids on the antibodies/proteins (1). For this reaction, antibodies should be in an amine-free buffer without any protein preservative. Antibodies labeled with the HaloTag® ligand are then incubated with HaloTag®-LgBiT or HaloTag®-SmBiT fusion protein to make a covalent conjugate of antibody-HaloTag®-LgBiT or antibody-HaloTag®-SmBiT (2). The success of the labeling reaction is confirmed by SDS-PAGE and Coomassie staining (3). Guidance for the labeling procedure is provided in the technical manual #TM602.





#### References

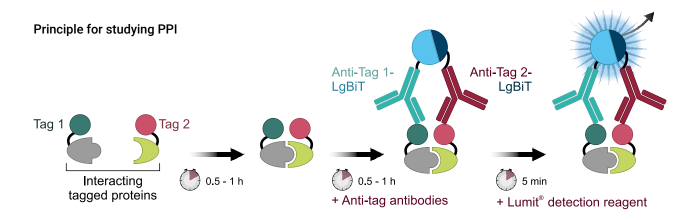
Alsulami, T. et al. (2021) Development of a novel homogeneous immunoassay using the engineered luminescent enzyme NanoLuc for the quantification of the mycotoxin fumonisin B1. Biosens Bioelectron. 177, 112939

## 3. Build-Your-Own Lumit® Immunoassays

## 3.3 Lumit® Anti-Tag Protein Interaction Reagents

#### **Description & Application**

Lumit® Anti-Tag Protein Interaction Reagents include a selection of BiT-labeled antibodies targeting common protein tags (e.g., His-, Flag®-, GST-tag, and human Fc) as well as BiT-labeled streptavidin. These reagents simplify the setup of biochemical assays to study protein:protein interactions (PPI) and screen for modulators of these interactions. Additionally, they enable the investigation of protein:small molecule interactions in a straightforward, competition-based, and HTS-compatible format.



#### **Principle & Workflow**

To study protein:protein interactions, two differentially tagged proteins are incubated, with the option to include a compound that modulates the PPI of interest. After incubation, BiT-labeled Lumit® anti-tag antibodies are added. Following the addition of the Lumit® detection reagent, the luminescent signal is recorded using a plate-reading luminometer. For AviTag™-tagged or biotinylated proteins, one of the BiT-labeled Lumit® anti-tag antibodies can be replaced with Lumit® Streptavidin-LgBiT or -SmBiT.For protein:small molecule interaction studies, a biotinylated small molecule (tracer) and a tagged protein are required. The interaction between the tracer and target protein is detected using a BiT-labeled anti-tag antibody and Lumit® Streptavidin-LgBiT or -SmBiT. Target engagement by unlabeled test compounds is reflected by a decrease in the luminescent signal, caused by competitive displacement of the tracer.



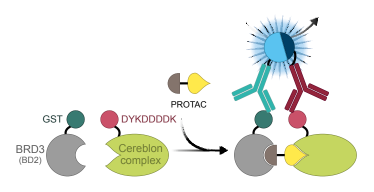
### Looking for a Source of Tagged Kinases to Quantitatively Measure Test Compound Affinity In Vitro?

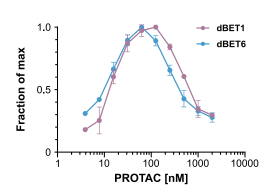
Promega offers more than 370 different tagged and purified kinases as part of the Kinase Enzyme Systems, ready-to-use assays to determine kinase activity *in vitro*.





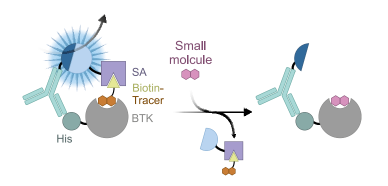
#### Representative Data

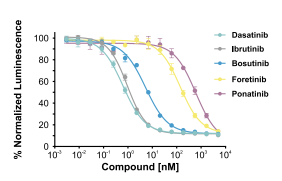




#### Monitor PROTAC®-induced protein:protein interactions

The ability of the PROTACs dBET1 and dBET6 to induce ternary complex formation between the Cereblon E3 ligase and BRD3 (BD2) was evaluated in a Lumit® Immunoassay. Recombinant GST-tagged BRD3 (BD2) (6.25 nM) and FLAG®-tagged Cereblon (6.25 nM) were incubated with different concentrations of PROTACs for 60 minutes. Detection was accomplished using Lumit® Anti-GST-LgBiT, Lumit® Anti-DYKDDDDK-SmBiT and Lumit® Immunoassay Detection Reagent A.





#### Detection and characterization of protein:small molecule interactions

The relative binding affinities of different kinase inhibitors for Bruton's tyrosine kinase (BTK) were determined in a competitive Lumit® Immunoassay. Recombinant His-tagged BTK (5 nM) was incubated with a biotinylated Ibrutinib (tracer; 37.5 nM) and kinase inhibitors at different concentrations ( $0.003 - 5000 \,\mu\text{M}$ ) for 60 minutes with mild agitation. Samples were incubated for 30 minutes with a mix of Streptavidin-LgBiT and Lumit® Anti-6His-SmBiT. The fraction of tracer-bound BTK at equilibrium was detected upon addition of Lumit® Immunoassay Detection Reagent A.

#### **Product Box**

Lumit® Anti-6His-LgBiT and -SmBiT

Cat.# W1600

Lumit® Anti-GST-LgBiT and -SmBiT

Cat.# W1620

Lumit® Anti-DYKDDDDK-LgBiT and SmBiT

Cat.# W1640

Lumit® Anti-Human IgG-LgBiT and -SmBiT

Cat.# CS332214

Lumit® Streptavidin-LgBiT and -SmBiT

Cat.# W1660

Combine with Lumit® Detection Reagent A



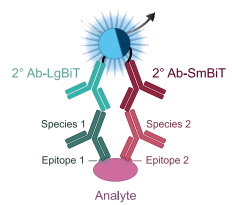
# 3. Build-Your-Own Lumit® Immunoassays

# 3.4 Lumit® Secondary Antibodies

#### **Description & Application**

Lumit® Secondary Antibodies are polyclonal BiT-labeled antibodies targeting IgG, enabling the development of Lumit® assays using the indirect immunoassay format. These antibodies are raised in donkey, immunoaffinity-purified using immobilized antigens, and conjugated to either SmBiT or LgBiT. Specificity against heavy and light chains of IgG (mouse, rabbit, and goat) has been confirmed.

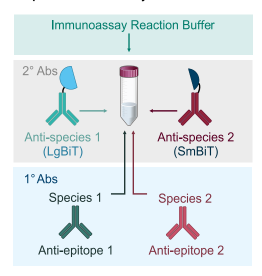
#### **Indirect Assay Format**



#### **Principle & Workflow**

To establish an indirect Lumit® Immunoassay for a new target, refer to the technical manual #TM613. In brief, users need to provide two analyte-specific primary antibodies raised in different species (e.g., mouse anti-analyte and rabbit anti-analyte). These are combined with matching Lumit® Secondary Antibodies (e.g., Lumit® Anti-Mouse Ab-LgBiT and Lumit® Anti-Rabbit Ab-SmBiT) to yield an antibody mix. Initially, it is recommended to test two or three primary antibody pairs. The best combination and concentration to use in an assay are determined through checkerboard experiments.

#### Preparation of Antibody Mix



#### **Product Box**

Lumit® Anti-Mouse Ab-LgBiT Cat.# W1021, W1022

Lumit® Anti-Rabbit Ab-LgBiT Cat.# W1041, W1042

Lumit® Anti-Rat Ab-LgBiT Cat.# CS366187, CS366188 Lumit® Anti-Mouse Ab-SmBiT Cat.# W1051, W1052

Lumit® Anti-Rabbit Ab-SmBiT

Cat.# W1031, W1032

Lumit® Anti-Rat Ab-SmBiT Cat.# CS366185, CS366186

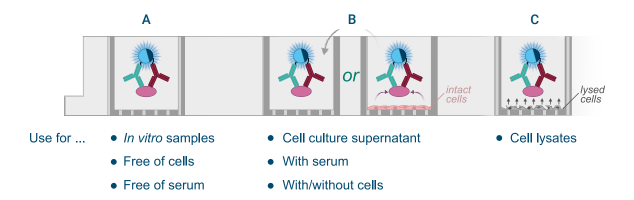


## 3.5 Lumit® Immunoassay Detection Reagents

#### **Description & Application**

Lumit® Immunoassay Detection Reagents support the development of customized Lumit® Immunoassays. Three reagent types (A, B, C) are available, enabling analyte detection under varying conditions. Detection Reagent A is intended for use in the absence of cells and serum, while Detection Reagent B is optimized for cell culture supernatant containing up to 10 % FBS and/or in the presence of intact cells. Detection Reagent C is included in the Lumit® Immunoassay Lysis and Detection Kit and is recommended for cell lysates.

#### Lumit® Immunoassay Detection Reagents



#### **Principle & Workflow**

Detailed guidelines for setting up Lumit® Immunoassays are available in technical manuals #TM613, #TM602, and #TM614. These manuals provide descriptions on how to optimize Lumit® Immunoassays, including tips for overcoming matrix effects and performing checkerboard experiments.

# Product Box Lumit® Immunoassay Detection Reagent A Cat.# VB2010, VB2020, VB2030 Lumit® Immunoassay Detection Reagent B Cat.# VB4050, VB4060 Lumit® Immunoassay Lysis and Detection Kit Cat.# W1231, W1232, W1233

# Pre-Built Lumit® Immunoassays

## Cytokines / HMGB1

Product	Cat. #	Size
Lumit® IL-1β (Human) Immunoassay	W6010 W6012 W6011	100 assays 500 assays 1000 assays
Lumit® IL-1β (Human) Standard	W116A-C	25 μl (10 μg/ml)
Lumit® IL-1β (Mouse) Immunoassay	W7010 W7012 W7011	100 assays 500 assays 1000 assays
Lumit® IL-1β (Mouse) Standard	W119A-C	25 μl (10 μg/ml)
Lumit® IL-2 (Human) Immunoassay	W6020 W6022 W6021	100 assays 500 assays 1000 assays
Lumit® IL-2 (Human) Standard	W122A-C	25 μl (10 μg/ml)
Lumit® IL-4 (Human) Immunoassay	W6060 W6062 W6061	100 assays 500 assays 1000 assays
Lumit® IL-4 (Human) Standard	W125A-C	25 μl (10 μg/ml)
Lumit® IL-6 (Human) Immunoassay	W6030 W6032 W6031	100 assays 500 assays 1000 assays
Lumit® IL-6 (Human) Standard	W128A-C	25 μl (10 μg/ml)
Lumit® IL-8 (Human) Immunoassay	CS2032C02*	100 assays
Lumit® IL-10 (Human) Immunoassay	W6070 W6072 W6071	100 assays 500 assays 1000 assays
Lumit® IL-10 (Human) Standard	W131A-C	25 μl (10 μg/ml)
Lumit® IL-12 (Human) Immunoassay	CS2032C04*	100 assays
Lumit® Active IL-18 (Human) Immunoassay	CS3291A01*	100 assays
Lumit® IL-17A (Human) Immunoassay	W1430 W1432 W1431	100 assays 500 assays 1000 assays
Lumit® IL-17A (Human) Standard	W143A-C	25 μl (10 μg/ml)

<sup>\*</sup> This is an Early Access Material. Please inquire for more information.

## Cytokines / HMGB1

Product	Cat. #	Size
Lumit® TNF-α (Human) Immunoassay	W6050 W6052 W6051	100 assays 500 assays 1000 assays
Lumit® TNF-α (Human) Standard	W137A-C	25 μl (10 μg/ml)
Lumit® IFN-β (Human) Immunoassay	W1810 W1812 W1811	100 assays 500 assays 1000 assays
Lumit® IFN-β (Human) Standard	W149A-C	25 μl (10 μg/ml)
Lumit® IFN-γ (Human) Immunoassay	W6040 W6042 W6041	100 assays 500 assays 1000 assays
Lumit® IFN-γ (Human) Standard	W134A-C	25 μl (10 μg/ml)
Lumit® VEGF-A (Human) Immunoassay	CS2032C07*	100 assays
Lumit® HMGB1 (Human/Mouse) Immunoassay	W6110 W6112	100 assays 500 assays
Lumit® HMGB1 (Human) Positive Control	W140A-C	20 μl (500 μg/ml)

#### Hormones

Product	Cat. #	Size
Lumit® Insulin Immunoassay Kit	CS3037A05* CS3037A07*	$100^{1)} - 400^{2)}$ assays $500^{1)} - 2000^{2)}$ assays
Lumit® Glucagon Immunoassay Kit	W8020 W8022	100 <sup>1)</sup> – 400 <sup>2)</sup> assays 500 <sup>1)</sup> – 2000 <sup>2)</sup> assays

<sup>1)</sup> 96-well; <sup>2)</sup> 384-well

#### **Cell Proliferation**

Product	Cat. #	Size
Lumit® hKi-67 Immunoassay for Cell Proliferation	CS3076A01*	100 assays

<sup>\*</sup> This is an Early Access Material. Please inquire for more information.

# Pre-Built Lumit® Immunoassays

#### **Second Messenger**

Product	Cat. #	Size
Lumit® cAMP Immunoassay	CS3592A08*	100 assays
Lumit® cGAMP Immunoassay	CS3592A05*	100 assays

## Cell & Gene Therapy

Product	Cat. #	Size
Lumit® dsRNA Detection Assay	W2041 W2042	100 assays 500 assays
Lumit® dsRNA Assay Buffer (5x)	W2010	8 ml
Lumit® dsRNA Standard	W2040	15 $\mu$ l (100 $\mu$ g/ml)
Lumit® dsRNA Standard, 5-methylcytidine	CS355716*	15 μl (100 μg/ml)
Lumit® dsRNA Standard, N1-methylpseudouridine	CS355712*	15 μl (100 μg/ml)
Lumit® dsRNA Standard, Pseudouridine	CS355710*	15 μl (100 μg/ml)
Lumit® dsRNA Standard, 5-methoxyuridine	CS355714*	15 μl (100 μg/ml)
Lumit® p24 Immunoassay	CS2039B25*	100 assays

#### **Therapeutic Antibody Development**

Product	Cat. #	Size
Lumit® FcγRI Binding Immunoassay	W7080 W7081	100 assays 1000 assays
Lumit® FcγRIIa (H131) Binding Immunoassay	W7070 W7071	100 assays 1000 assays
Lumit® FcγRIIa (R131) Binding Immunoassay	W7060 W7061	100 assays 1000 assays
Lumit® FcyRIIb Binding Immunoassay	W7030 W7031	100 assays 1000 assays
Lumit® FcγRIIIa (F158) Binding Immunoassay	W7040 W7041	100 assays 1000 assays
Lumit® FcγRIIIa (V158) Binding Immunoassay	W7050 W7051	100 assays 1000 assays
Lumit® FcγRIIIb Binding Immunoassay	W7020 W7021	100 assays 1000 assays

<sup>\*</sup> This is an Early Access Material. Please inquire for more information.

#### **Therapeutic Antibody Development**

Product	Cat. #	Size
Lumit® FcRn Binding Immunoassay	W1151 W1152	100 assays 1000 assays
Lumit® Canine FcRn Binding Immunoassay	CS3019A02*	100 assays
Lumit® Feline FcRn Binding Immunoassay	CS3631B01* CS3631B05*	100 assays 1000 assays
Lumit® Human IgG Titer Immunoassay	CS3716A07* CS3716A13*	100 assays 1000 assays
Lumit® Mouse IgG Titer Immunoassay	CS3716A03* CS3716A10*	100 assays 1000 assays
Lumit® Rabbit IgG Titer Immunoassay	CS3716A09* CS3716A14*	100 assays 1000 assays

## Signaling & Protein Degradation

Product	Cat. #	Size
Lumit® Immunoassay Cellular System – Starter Kit	W1220	200 assays
Lumit® Immunoassay Cellular System – Set 1	W1201 W1202 W1203	100 assays 1000 assays 10000 assays
Lumit® Immunoassay Cellular System – Set 2	W1331 W1332 W1333	100 assays 1000 assays 10000 assays

<sup>\*</sup> This is an Early Access Material. Please inquire for more information.



## **Need Technical Assistance?**

## Please contact our technical service:



www.promega.com/support/tech-support



# Pre-Built Lumit® Immunoassays

## Signaling & Protein Degradation -

Lumit® Immunoassay Cellular Systems (ICS) Complete Assays

Product	Cat. #	Size
Lumit® Phospho-AKT (Ser 473) ICS	CS3397A07* CS3397A08*	100 assays 1000 assays
Lumit® Total BTK ICS	CS366152* CS366156*	100 assays 1000 assays
Lumit® Phospho-BTK (Tyr 223) ICS	CS3397A19* CS3397A20*	100 assays 1000 assays
Lumit® Total BRD4 ICS	CS366168* CS366172*	100 assays 1000 assays
Lumit® Phospho-CHK1 (Ser 317) ICS	CS366144* CS366148*	100 assays 1000 assays
Lumit® Total CK1α ICS	please enquire	100 assays 1000 assays
Lumit® p-cMET (Tyr 1234/1235) ICS	CS366136* CS366140*	100 assays 1000 assays
Lumit® Total EGFR ICS	CS366120* CS366124*	100 assays 1000 assays
Lumit® Phospho-EGFR (Tyr 1068) ICS	CS366104* CS366108*	100 assays 1000 assays
Lumit® Phospho-ERK (Thr 202) ICS	CS3397A03* CS3397A04*	100 assays 1000 assays
Lumit® Total GSPT1 ICS	CS366904* CS366908*	100 assays 1000 assays
Lumit® Phospho-H2AX (Ser 139) ICS	CS366112* CS366116*	100 assays 1000 assays
Lumit® Total IκBα ICS	CS3397A58* CS3397A62*	100 assays 1000 assays
Lumit® Phospho-IκBα (Ser 32) ICS	CS3397A15* CS3397A16*	100 assays 1000 assays
Lumit® Total IKFZ1 ICS	please enquire	100 assays 1000 assays
Lumit® Total IKFZ2 ICS	please enquire	100 assays 1000 assays

<sup>\*</sup> This is an Early Access Material. Please inquire for more information.

## Signaling & Protein Degradation -

Lumit® Immunoassay Cellular Systems (ICS) Complete Assays

Product	Cat. #	Size
Lumit® Total IKFZ3 ICS	please enquire please enquire	100 assays 1000 assays
Lumit® Phospho-JNK (Thr 183/Tyr 185) ICS	CS3397A34* CS3397A38*	100 assays 1000 assays
Lumit® Phospho-p65 (Ser 536) ICS	CS3397A42* CS3397A46*	100 assays 1000 assays
Lumit® Phospho-RB (Ser 807/811)	CS3397A26* CS3397A30*	100 assays 1000 assays
Lumit® Phospho-RPS6 (Ser 240/244) ICS	CS366176* CS366180*	100 assays 1000 assays
Lumit® Total SALL4 ICS	please enquire please enquire	100 assays 1000 assays
Lumit® Total SMARCA2 ICS	CS366128* CS366132*	100 assays 1000 assays
Lumit® Total SMARCA4 ICS	CS366160* CS366164*	100 assays
Lumit® Total STAT3 ICS	CS3397A50* CS3397A54*	100 assays
Lumit® Phospho-STAT3 (Tyr 705) ICS	CS3397A11* CS3397A12*	100 assays

<sup>\*</sup>This is an Early Access Material. Please inquire for more information.

# **Build-Your-Own Lumit® Immunoassays**

#### Lumit® Immunoassay Labeling Kit

Product	Cat. #	Size
Lumit® Immunoassay Labeling Kit	VB2500	1 kit

#### **Lumit® Anti-Tag Protein Interaction Reagents**

Product	Cat. #	Size
Lumit® Anti-6His-LgBiT and -SmBiT  • Lumit® Anti-6His-LgBiT  • Lumit® Anti-6His-SmBiT	W1600	20 μl each
Lumit® Anti-6His-LgBiT	W1601	200 μΙ
Lumit® Anti-6His-SmBiT	W1611	200 μΙ
Lumit® Anti-GST-LgBiT and -SmBiT  • Lumit® Anti-GST-LgBiT  • Lumit® Anti-GST-SmBiT	W1620	20 μl each
Lumit® Anti-GST-LgBiT	W1621	200 μΙ
Lumit® Anti-GST-SmBiT	W1631	200 μΙ
Lumit® Anti-DYKDDDDK-LgBiT and -SmBiT  • Lumit® Anti-DYKDDDDK-LgBiT  • Lumit® Anti-DYKDDDDK-SmBiT	W1640	20 μl each
Lumit® Anti-DYKDDDDK-LgBiT	W1641	200 μΙ
Lumit® Anti-DYKDDDDK-SmBiT	W1651	200 μΙ
Lumit® Anti-Human IgG-LgBiT and -SmBiT  • Lumit® Anti-Human IgG-LgBiT  • Lumit® Anti-Human IgG-SmBiT	CS332214*	20 μl each
Lumit® Anti-Human IgG-LgBiT	CS332222*	200 μΙ
Lumit® Anti-Human IgG-SmBiT	CS332223*	200 μΙ
Lumit® Streptavidin-LgBiT and -SmBiT 1)  • Lumit® Streptavidin-LgBiT  • Lumit® Streptavidin-SmBiT	W1660	20 μl each
Lumit® Streptavidin-LgBiT	W1661	200 μΙ
Lumit® Streptavidin-SmBiT	W1671	200 μΙ
Lumit® Anti-Mouse Ab-LgBiT and -SmBiT	refer to page 47	
Lumit® Anti-Rabbit Ab-LgBiT and -SmBiT	refer to page 47	

<sup>&</sup>lt;sup>1)</sup> Can be used with AviTag<sup>™</sup>-tagged proteins or biotinylated proteins/small molecules.

If not indicated otherwise, the number of assays is given for 96-well plates.

 $<sup>\</sup>mbox{\ensuremath{\star}}$  This is an Early Access Material. Please inquire for more information.

## **Lumit® Secondary Antibodies**

Product	Cat. #	Size
Lumit® Immunoassay Cellular System – Set 1	W1201 W1202 W1203	100 assays 1000 assays 10000 assays
Lumit® Immunoassay Cellular System – Set 2	W1331 W1332 W1333	100 assays 1000 assays 10000 assays
Lumit® Anti-Mouse Ab-LgBiT	W1021 W1022	30 µl 300 µl
Lumit® Anti-Mouse Ab-SmBiT	W1051 W1052	30 µl 300 µl
Lumit® Anti-Rabbit Ab-LgBiT	W1041 W1042	30 µl 300 µl
Lumit® Anti-Rabbit Ab-SmBiT	W1031 W1032	30 µl 300 µl
Lumit® Anti-Rat Ab-LgBiT	CS366187* CS366188*	30 µl 300 µl
Lumit® Anti-Rat Ab-SmBiT	CS366185* CS366186*	30 µl 300 µl

## Lumit® Immunoassay Detection Reagents

Product	Cat. #	Size
Lumit® Immunoassay Detection Reagent A	VB2010 VB2020 VB2030	500 assays 5000 assays 50000 assays
Lumit® Immunoassay Detection Reagent B	VB4050 VB4060	100 Assays 1000 Assays
Lumit® Immunoassay Lysis and Detection Kit	W1231 W1232 W1233	100 assays 1000 assays 10000 assays

 $<sup>\</sup>mbox{\ensuremath{\star}}$  This is an Early Access Material. Please inquire for more information.

For Research Use Only. Not for Use in Diagnostic Procedures.

## 5. References

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#### Lumit® cAMP Immunoassay

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# Are You Looking for More Information on Lumit® Immunoassays?

Download the whitepaper here: www.promega.com/lumitwp



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# Contact

- Headquarters
- R&D and Production
- Branches
- Distributors

#### **North America**

#### **USA/Canada**

Tel +1 608 274 4330 Fax +1 608 277 2516 custserv@promega.com

#### **South America**

#### Brazil

Tel +55 11 5090 3780 Fax +55 11 5096 3780 promega.brasil@promega.com

#### **Europe**

#### Germany/Austria

Tel +49 6227 6906 0 Fax +49 6227 6906 222 de\_custserv@promega.com

#### Switzerland

Tel +41 44 878 90 00 ch\_custserv@promega.com

#### France

Tel +33 4 37 22 50 00 Fax +33 4 37 22 50 00 fr.contact@promega.com

#### Italy

Tel +39 2 54 05 01 94 Fax +39 2 56 56 16 45 customerservice.italia@promega.com

#### Poland

Tel +48 22 531 0667 Fax +48 22 531 0669 pl\_custserv@promega.com

#### Central/Eastern Europe

Tel +49 6227 6906 0 Fax +49 6227 6906 222 de\_ceesales@promega.com

#### Belgium/Luxembourg/Netherlands

Tel +31 71 532 42 44 Fax +31 71 532 49 07 benelux@promega.com

### Spain/Portugal

Tel +34 916 62 11 26 esp\_custserv@promega.com

# Denmark, Estonia, Finland, Iceland, Norway, Sweden

Tel +46 8 452 2450 sweorder@promega.com

#### **United Kingdom**

Tel +44 23 8076 0225 Fax +44 23 8076 7014 ukcustserve@promega.com





#### Asia

#### China

Tel +86 10 5825 6268 Fax +86 10 5825 6160 info@promega.com.cn

#### Japan

Tel +81 3 3669 7981 Fax +81 3 3669 7982 jpmktg@jp.promega.com

#### Korea

Tel +82 2 1588 3718 Fax +82 2 2638 5418 CustServiceKR@promega.com

#### Singapore

Tel +65 6513 3450 Fax +65 6773 5210 sg\_custserv@promega.com

#### India

Tel +91 11 43005814/15/16/17 Fax +91 11 41035028 ind\_custserv@promega.com

#### **Australia and Oceania**

#### Australia

Tel +61 2 8338 3800 Fax +61 2 8338 3855 auscustserv@promega.com

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