

Multiplexed Analysis of Second Messenger Signaling in Live Cells Using Aequorin and GloSensor™ cAMP on the Hamamatsu μCell™

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1. Abstract

Detection of intracellular second messenger signaling is an established method for measuring G-protein-dependent GPCR activation. Although there are several technologies available for measurement of second messengers via endpoint analysis, technologies for monitoring second messengers in living cells include Promega's GloSensor™ cAMP for quantifying intracellular [cAMP] and technologies such as the photoprotein Aequorin or various fluorescence-based indicators for [Ca²⁺]. These technologies serve to quantify second messengers in live cells and in real-time following GPCR activation, providing several advantages over lytic endpoint assays. However, it may be challenging when screening for GPCR activity modulators when G-protein-dependent signaling is uncharacterized or when the desired second messenger detection format cannot be predicted (for example, in the case of orphan receptors). Furthermore, for GPCRs capable of modulating both [cAMP] and [Ca²⁺] pathways concurrently, it would be desirable to measure G protein coupling simultaneously. Few technologies exist that allow for simultaneous measurement of Ca²⁺ and cAMP in live cells, while maintaining assay robustness and high signal-to-background for use in HTS. To address this limitation, Promega has developed a live cell method for the kinetic measurement of Ca²⁺ and cAMP by multiplexing of Aequorin and GloSensor™ cAMP bioluminescent sensor technologies. Using the Hamamatsu FDSS/μCell, we report simultaneous analysis of Ca²⁺ and cAMP mobilization following agonism of Parathyroid Hormone Receptor (PTH1R) using a promiscuous compound directing both G_{α_s}+G_{α_q} signaling, as well as a biased compound specifically directing G_{α_s} coupling alone. The combination of these bioluminescence-based sensor technologies with the Hamamatsu FDSS/μCell serves as an ideal platform for the analysis of these divergent second messenger signaling events in live cells and in real time.

2. GloSensor™ cAMP Assay

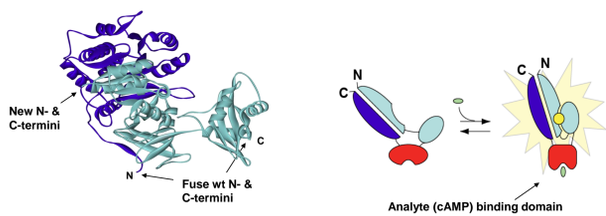
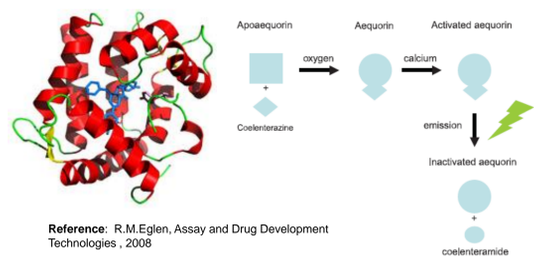


Diagram of GloSensor™ cAMP Activation. The assay is based on the GloSensor™ Technology, a genetically modified form of firefly luciferase which has been modified with a cAMP-binding protein moiety inserted into unique N and C termini. Upon binding of cAMP, conformational change is induced leading to increased luciferase activity.

- **Live Cell Assay:** Excels at kinetic and modulation studies of G_{α_s}-coupled receptors signaling through cAMP.
- **Transient or Stable Expression:** GloSensor™ cAMP Assay is utilized by transiently expressing a receptor of interest and the biosensor in the cell line of choice. Alternatively, stably transfected cell lines with both the biosensor and the receptor of interest can be made.
- **Simple Protocol:** Cells are pre-equilibrated with GloSensor™ cAMP Reagent, then cells are treated with specific agonists/antagonists or compounds, and luminescence is measured in real-time (typically 10-30 minutes).

3. Aequorin Assay for [Ca²⁺]

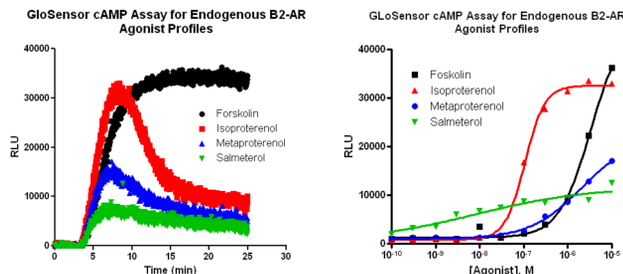


Reference: R.M.Eglen, Assay and Drug Development Technologies, 2008

Diagram of Ca²⁺-mediated Aequorin Activation. Photoproteins such as Aequorin are widely used for measuring rapid GPCR-induced, transient changes in [Ca²⁺] from G_{α_q}-coupled receptors. Aequorin is composed of two distinct units which reconstitute spontaneously, providing a method to quantify changes in [Ca²⁺] in live cells.

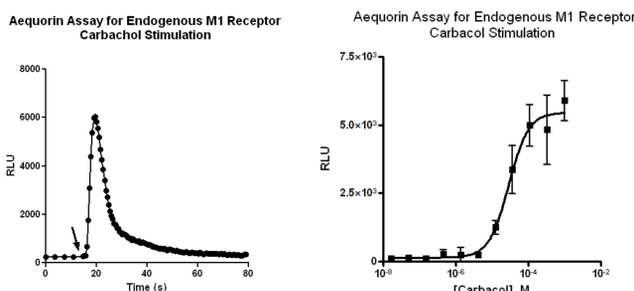
- **Live Cell Assay:** Apoaequorin binds Coelenterazine to produce Aequorin in live cells. When Ca²⁺ binds Aequorin, the protein undergoes conformational changes, resulting in oxidation of coelenterazine to the excited form coelenteramide.
- **Fast Kinetics:** As an excited coelenteramide relaxes to the ground state blue light at wavelength 470 nm is emitted. The intensity of light emission can vary but typically occurs within seconds, enabling a live cell endpoint that can be resolved over time with GloSensor™ cAMP
- **Simple Protocol:** Cells are pre-equilibrated with coelenterazine, then cells are stimulated and luminescence is measured within seconds.

4. GloSensor™ cAMP Assay: Representative Data for G_{α_s} Signaling



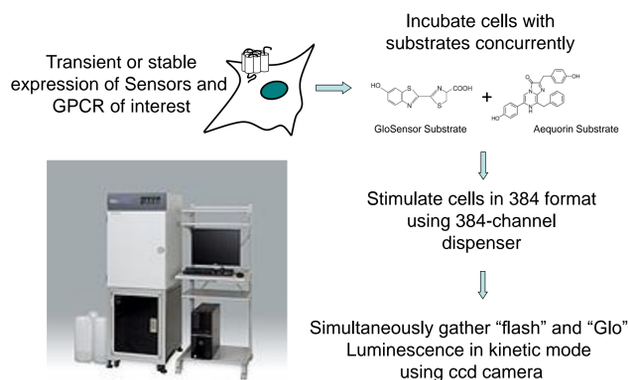
Materials and Methods. Left: Kinetic measurement of agonist-induced cAMP mobilization in HEK293 cells stably expressing GloSensor™ cAMP. (L9) In 384 format, cells were preincubated with GloSensor™ cAMP substrate for 1.5h prior to stimulation. Luminescence was measured on the Hamamatsu μCell with 3s of integration time. Right: Dose-response measurement of agonist-induced cAMP mobilization under similar conditions (7.5 minute timepoint).

5. Aequorin Assay: Representative Data for G_{α_q} Signaling

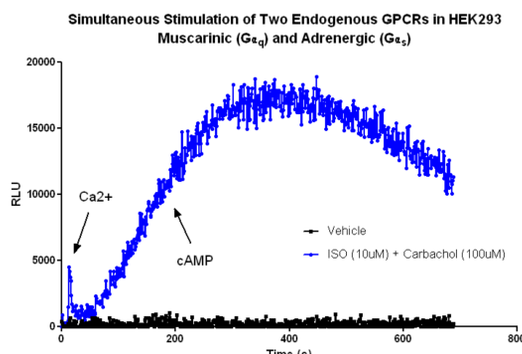


Materials and Methods. Left: Kinetic measurement of Carbachol-induced Ca²⁺ mobilization in HEK293 cells. Cells were transfected with plasmid DNA encoding Aequorin and seeded in 96-well plates. Following >24h of transfection, cells were preincubated with Aequorin substrate for 3h prior to Carbachol stimulation (0.5s of luminescence integration time). Right: Dose-response measurement of Carbachol-induced Ca²⁺ mobilization as described but at a 3s timepoint.

6. Simple Workflow for Multiplexing of Aequorin / GloSensor™ cAMP Using The Hamamatsu μCell



7. Simultaneous measurement of G_{α_q} and G_{α_s} signaling from two GPCRs



Materials and Methods: HEK293 cells were transiently-transfected with plasmid DNA encoding using Aequorin and GloSensor™ cAMP 22F and seeded in 384-well, clear-bottom plates. 24 h post-transfection, cells were preincubated with GloSensor™ cAMP substrate and coelenterazine for approximately 3 hours. Cells were then stimulated with Isoproterenol + carbachol or vehicle. Luminescence was then measured on a Hamamatsu μCell using 1s of luminescence integration.

7. Biased Agonism at PTH1R

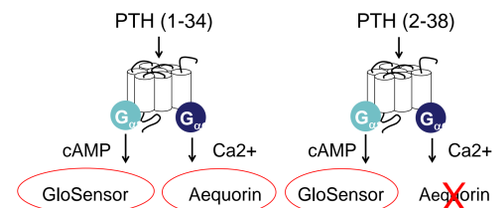
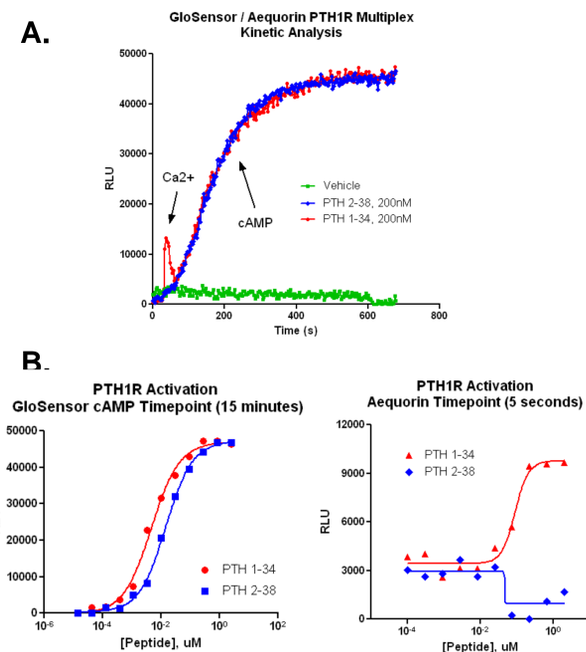


Diagram of biased agonism at Parathyroid Hormone Receptor (PTH1R) using a non-selective and G_{α_s}-selective peptide agonist. Peptide agonist PTH(1-34) is expected to activate both G_{α_s} and G_{α_q} pathways, leading to concurrent activation of both cAMP and Ca²⁺. Amino-terminal truncation of PTH peptide (PTH-2-38) is expected to induce cAMP pathway selectively. GloSensor™ cAMP and Aequorin can therefore be used to query these distinct signaling mechanisms.

Reference: Takasu H et al. Biochemistry. 1999 Oct 12;38(41):13453-60. Amino-terminal modifications of human parathyroid hormone (PTH) selectively alter phospholipase C signaling via the type 1 PTH receptor: implications for design of signal-specific PTH ligands.

8. Multiplexing of [Ca²⁺] and [cAMP] Signaling from PTH1R



Simultaneous multiplexing of Ca²⁺ and cAMP mobilization using GloSensor™ cAMP and Aequorin. Materials and Methods: HEK293 cells were triple-transfected with plasmid DNAs encoding PTH1R, Aequorin, and GloSensor™ cAMP 22F using Fugene HD, and seeded in clear-bottom 384-well plates. 24h posttransfection, cells were preincubated with GloSensor™ substrate and coelenterazine for 3h. A. Kinetic analysis of PTH1R activation using a peptide agonist activating G_{α_s}+G_{α_q} or an agonist selectively activating G_{α_s} only. Cells were then stimulated with 200 nM PTH(1-34) peptide, PTH(2-38) peptide, or vehicle. Kinetic analysis of luminescence was performed on the Hamamatsu μCell using 3s of luminescence integration time. B. Simultaneous dose-response profiles for PTH1R activation of cAMP after 15 min (left) or Ca²⁺ after 5s (right).

9. Conclusions

GloSensor™ cAMP and Aequorin are complementary bioluminescent technologies for multiplexing of second messenger signaling.

- Kinetic measurements of Ca²⁺ and cAMP can easily be multiplexed on the Hamamatsu FDSS/μCell in an HTS-compatible format
- The method described is compatible via transient transfection of DNA encoding the biosensors and the GPCR of interest (no stable cell lines required)
- Two endogenous GPCRs can be measured in a single well, offering a unique solution for analysis of GPCR signaling when G-protein couplings are not fully characterized
- Multiplexing of Aequorin/GloSensor™ cAMP enables a novel approach to GPCR functional selectivity studies

Questions or comments?
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