

APPLICATION INFORMATION

Image Cytometry

ANALYSIS OF GPCR ACTIVITY

This application note describes the successful deployment of the NORAK Transfluor[®] assay for GPCR activity on the Cell Lab IC 100 Image Cytometer from Beckman Coulter, Inc. The Norak Transfluor assay measures GPCR activation by the formation of GFP-labeled pits, or vesicles, depending on the GPCR that is being expressed by a cell. The Cell Lab IC 100 is a novel Image Cytometry system that delivers accurate, quantitative imaging and analysis of cell populations, at high speeds, directly from microtiter plates. For accurate reading and measurement, the Cell Lab IC 100 quantifies distribution of fluorescence emanating from the GPCR-associated arrestin-GFP fusion protein and in sub-cellular compartments. Definitive measurements are quantified with a proprietary, multi-scale vesicle definition algorithm developed at Beckman Coulter, Inc.

Background and Significance

GPCRs mediate the activity of cell surface receptors and the transduction of myriad intra-cellular responses. GPCRs have proven to be a highly amenable class of targets for successful therapeutic intervention. In fact, of the approximately 500 drugs currently on the market today, more than 30% are mediated through the activation of GPCRs. Because GPCRs are membrane-bound proteins, they have been difficult to study in cell extracts, or to isolate and characterize. To directly ascertain GPCR activity, intact cell-based assays are quickly becoming a method of choice in high-throughput screening.

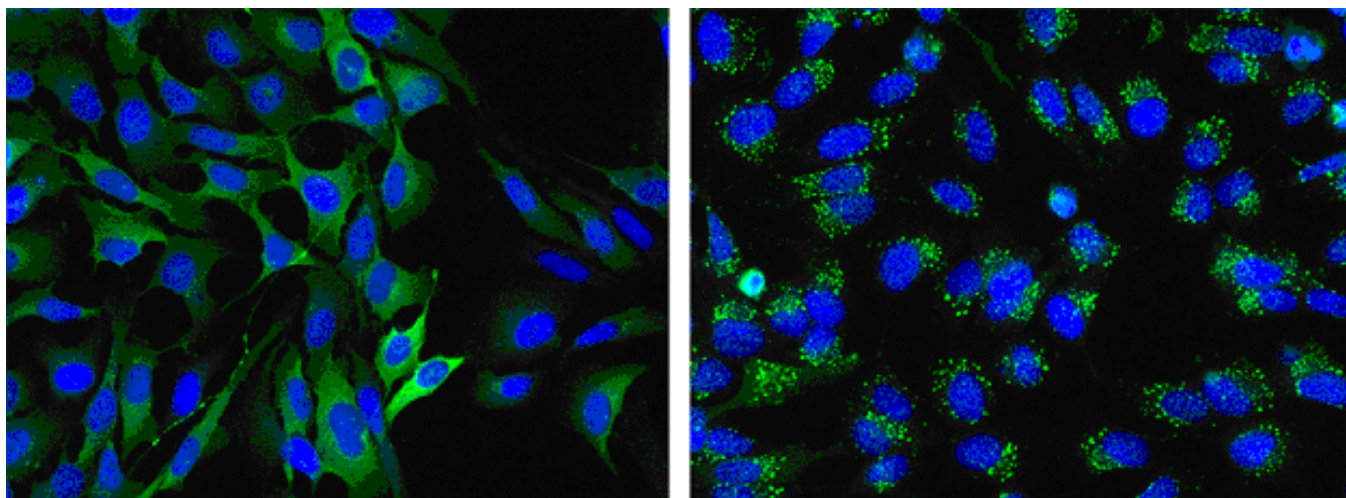


Figure 1. Vesicle response in human osteosarcoma cells (U2OS). Example 20X 0.5 N.A. fluorescent micrographs of the Norak Transfluor assay vesicle response using the Cell Lab IC 100 Image Cytometer. GFP expression (green) and Hoechst staining (blue) are visualized before (left) and after (right) vesicle formation.

GPCRs transduce extracellular signals through the formation of protein complexes that affect both activation and subsequent desensitization of a cell surface receptor. Agonist binding to a receptor at the cell surface initiates a conformational change in the intracellular domain of the receptor that results in the phosphorylation of the receptor and subsequent binding of arrestin to the receptor. The arrestin-receptor complex is then transported to clathrin-coated pits and internalized to clathrin-coated vesicles. Finally, the entire complex is delivered to the endosomes. Some GPCRs dissociate from arrestin at or near the plasma membrane, while others remain associated and traffic into endocytic vesicles.

In the Norak Transfluor™ assay, a cell line is developed to monitor the interaction of a given GPCR and a GFP fused to β -arrestin. When each GPCR is activated, the β -arrestin will bind to the membrane associated GPCR. The activated β -arrestin-GPCR complex then enters clathrin-coated pits and migrates to intracellular vesicles via the endosomal pathway. Some GPCRs retain the arrestin molecule throughout this process, so that vesicles will fluoresce with GFP. Other GPCRs will dissociate from the arrestin such that the GFP remains with the pits, or is released to the intracellular space, and the receptors recycle back to the cell surface and bind to arrestin again.

For example, in cell lines expressing either the dopamine D1A receptor (D1AR) or the alpha 1b-adrenergic receptor (α 1bAR), arrestin-GFP, remains localized at the plasma membrane in pits. In contrast, arrestin-GFP is internalized into endocytic vesicles with the vasopressin V2 receptor (V2R) upon ligand binding. The Norak Transfluor assay has 2 outcomes, depending on the specific GPCR being assayed. Where gross vesicle response may be analyzed with low resolution optics, meaningful edge/pit response requires sub-micron resolution, which was achieved in this study with the Cell Lab IC 100 cytometer. This detailed image quantification provides valuable insight for use in drug discovery.

Experimental Results and Automation

The combination of automated sub-micron imaging, proprietary image processing and the Norak Transfluor assay enables accurate measurement of GPCR activity through vesicle response and pit formation. Validated results of vesicle response and pit formation using the Cell Lab IC 100 cytometer are presented with fluorescent images and dose response curves.

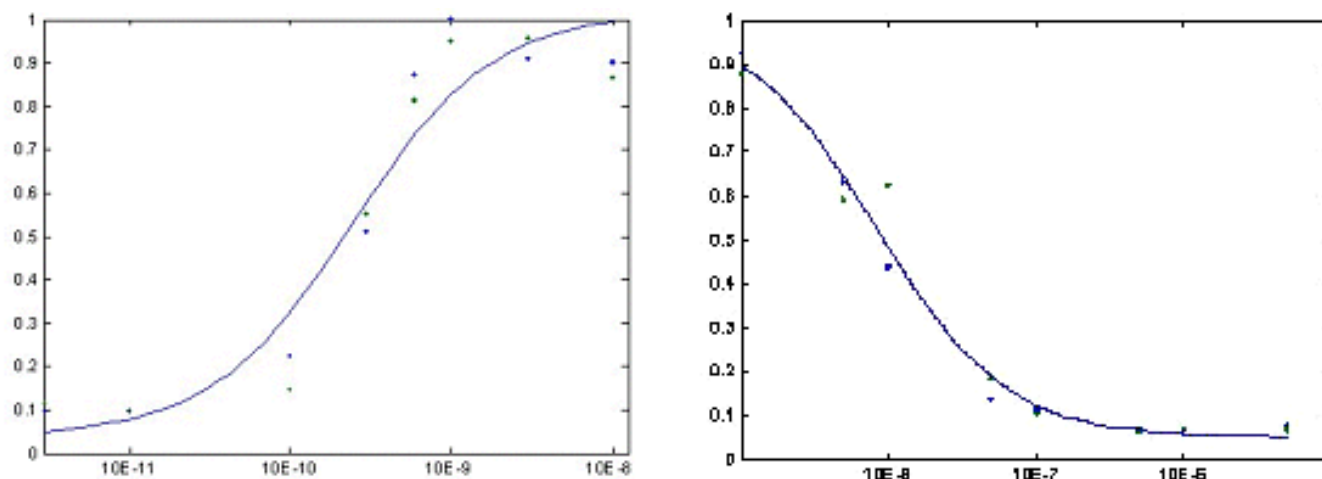


Figure 2. Dose response of GPCR activation. Agonist induction (left, $EC_{50} = 0.242$ nM) and antagonist inhibition (right, $IC_{50} = 8.386$ nM) of GPCR response with respect to dose (M). Fluorescent response is normalized and a cell based proprietary metric developed by Beckman Coulter Inc.

In the example images, GFP expression is visualized in green while the nuclear Hoechst stain is seen in blue. Response measurements were obtained from human osteosarcoma cells (U2OS) grown, dosed and imaged in 96-well plate format.

Conclusion

Beckman Coulter's proprietary metric FLIV (fractional localized intensity in vesicles) provides a sensitive and accurate measurement of GPCR activation, and enables the rapid and precise quantification of fluorescent signals imaged by the Cell Lab IC 100 Image cytometer. Use of the Cell Lab IC 100 in combination with the Norak Transfluo assay presents a powerful quantification tool for GPCR activity.

The Cell Lab IC 100 Image Cytometer from Beckman Coulter, Inc. delivers an unmatched combination of speed, accuracy and detail to quantitative imaging and analysis of cell populations. The Cell Lab IC 100 is used to accelerate drug discovery, clinical diagnostic research and basic research.

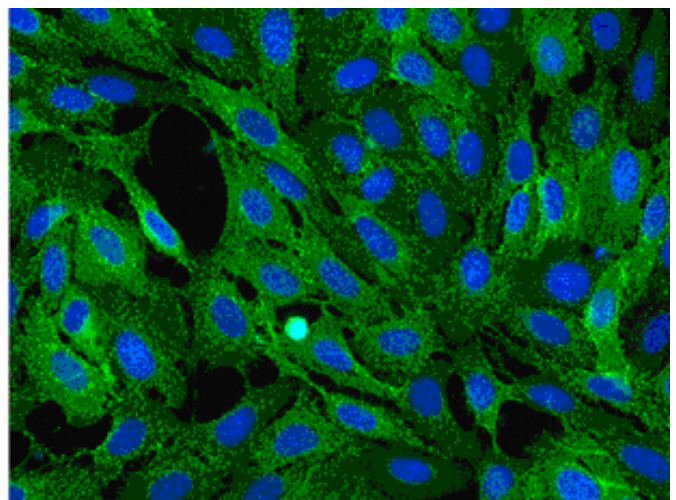
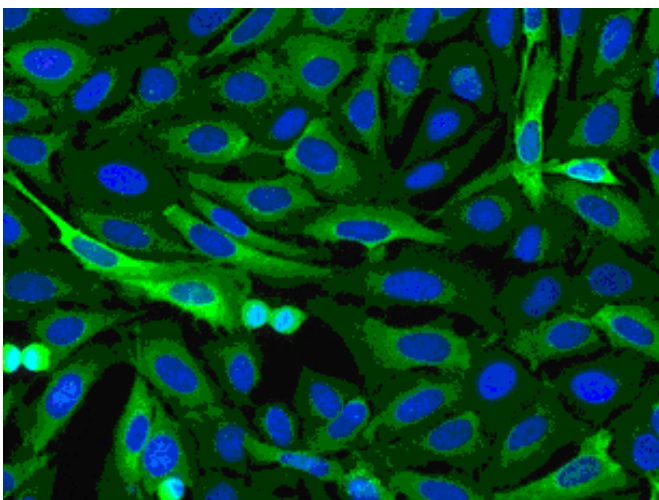
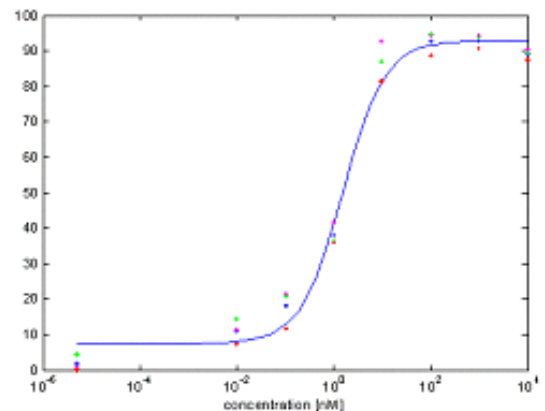


Figure 3. Pit formation in human osteosarcoma cells (U2OS). Example 20X 0.5 N.A. fluorescent micrographs of the Norak Transfluo assay pit formation using the Cell Lab IC 100 Image Cytometer. GFP expression (green) and Hoechst staining (blue) are visualized before (left) and after (lower right) pit formation. Dose response curve (upper right) shows fluorescence response with respect to isoproterenol concentration (M) as derived from 20X 0.5 N.A. images. $EC_{50} = 1.49 \pm 1.01 nM$.

* Access to the Norak Transfluo assay requires a separate patent license, which can be secured from Norak Biosciences, Inc. (Tel. 919-248-8000) for an additional fee. Beckman Coulter, Inc. does not have the right or authority to grant any license under patents covering Transfluo. No license to make, use, sell, offer for sale, or import Transfluo is granted by the purchase or license of Beckman Coulter. All trademarks are the property of their respective owners



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B2004-6612

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