Integrated Platform for Monitoring Single-cell MAPK Kinetics in Computer-controlled Temporal Stimulations

Hyunryul Ryu¹, Minhwan Chung², Jiyoung Song², Sung Sik Lee³, Olivier Pertz⁴ & Noo Li Jeon²,⁵

Extracellular response kinase (ERK) is one of the key regulator of cell fate, such as proliferation, differentiation and cell migration. Here, we propose a novel experimental pipeline to learn ERK kinetics by temporal growth factor (GF) stimulation. High signal-to-noise ratio of genetically encoded Fluorescence resonance energy transfer (FRET) biosensor enables to get a large number of single-cell ERK activity at each time point, while computer-controlled microfluidics fine-tune the temporal stimulation. Using this platform, we observed that static Epidermal growth factor (EGF) stimulation led to transient ERK activation with a significant cell-to-cell variation, while dynamic stimulation of 3′ EGF pulse led to faster adaptation kinetics with no discrepancy. Multiple EGF pulses retriggered ERK activity with respect to frequency of stimulation. We also observed oscillation of ERK activity of each cell at basal state. Introducing of Mitogen-activated protein kinase kinase (MEK) inhibitor, U0126, was not only dropping the average of basal activity for 7.5%, but also diminishing oscillatory behavior. Activity level raised up when inhibitor was removed, followed by transient peak of ERK kinetics. We expect this platform to probe Mitogen-associated protein kinase (MAPK) signaling network for systems biology research at single cellular level.
stimulation experiments. Harvey et al. and Fritz et al. showed ERK response by EGF stimulation. Purvis et al. showed p53 dynamics at damaged cell by gamma-irradiation. Selimkhanov et al. reported complex kinetics of ERK, calcium and NF-κB by EGF, ATP and LPS. However, these methods were limited to static stimuli, which acts as a limiting factor in the analysis of the system properties of the molecular pathways. Precisely controlled temporal stimulation is to overcome these limitations, by providing a quantitative input, giving not only the mathematical characteristics about the pathway, but also enabled dynamic regulation of the gene expression.

Here, biosensor and microfluidics, were integrated to observe ERK response from the pre-defined stimulation in a real-time. EKAR2G, FRET-based ERK biosensor, was genetically encoded, providing stable and uniform expression throughout the population. Meanwhile, the medium-filled reservoirs of microfluidic device were pressurized with computer-controlled pressure pump to manipulate GF concentration in the cell chamber. By measuring the intensity of ratio-metric single cell images, we analyzed the discrepancy of individual ERK kinetics to various stimulation patterns; sustained, pulsed and multi-pulsed. Basal oscillation of ERK activity of each cell was observed. Presenting of MEK inhibitor, U0126, was able to drop the average of basal activity and also eliminate amplitude of oscillation. Taking advantage of precise control of the stimulation regimes and high-throughput imaging capability with air objects, we expect this integrated platform to be used to obtain quantitative data, establishing advanced mathematical models of MAPK dynamics.

**Experimental Section**

**Generation of sensor-expressing cell-line and cell culture.** HEK293T and HEK293 β5 cells (American Type Culture Collection) were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 1% L-glutamine, and 1% penicillin/streptomycin. We produced lentiviral vectors expressing EKAR2G1 to establish stable cell lines, as used in previous research. Briefly, HEK293T cells (Invitrogen, USA) were transfected with lentivirus and packaging constructs. Supernatant was then collected and concentrated with a Lenti-X concentrator kit (Clontech, Japan). HEK293 β5 cells were infected and selectively cultured with 1 µg/ml puromycin (Sigma, Germany). Then, cells were sorted by flow cytometry to express homogeneous and high level of biosensor. After sorting, cells were subsequently cultured in presence of 1 µg/ml puromycin. At 70% confluence, cells were passaged gently with 0.25% Trypsin-EDTA (Sigma, Germany). Note that EKAR2G measures ERK phosphorylation activity specifically in the cytosol. This readout might therefore differ slightly from pERK antibody staining that measures the absolute amount of phosphorylated ERK.

**Design and fabrication of microfluidic device.** For mammalian cell experiment, microfluidic device was modified from the previous research. Previously, device was designed to stimulate Saccharomyces cerevisiae for systems biology research. The height of cell culture chamber was 40 µm. However, for mammalian cell with 20 µm diameter, this was not appropriate to provide healthy micro-environment. On the other hand, the higher height of the device the more medium between control part and cell chamber, which cause time delay on temporal stimulation. We separate the microfluidic device into two part with two-layered system; 40 µm for micro-channels of control part and 100 µm for cell chamber region. Cells were stabled within cell culture region, while cellular environment switched between 30 seconds between on-and-off states.

Microfluidic device was replicated from a Silicon wafer with SU-8 micro-structures. Silicon master mold was composed of 40 µm and 100 µm thickness layers of photoresist (PR). First, the plasma treated Silicon wafer was spin-coated with SU-8 100 (Microchem, USA) negative PR for 40 µm thick. After baking at 65 °C for 5 minutes and 95 °C for 20 minutes, wafer was masked by the negative film mask (Han & All Tech, Korea), and exposed to 250 mJ of 405 nm ultraviolet light. (Shinu MST, Korea) Wafer was, then, baked again at 65 °C for a minute and 95°C for 10 minutes. SU-8 developer (Microchem, USA) was used to remove unexposed part. The second layer of PR was spin-coated for 100 µm thick, and baked at 65°C for 10 minutes and 95°C for 30 minutes. Film mask for the second layer was aligned using alignment pattern on the first developed layer. Wafer was exposed to 500 mJ of UV light. After the baking step at 65°C for a minute and 95°C for 10 minutes, wafer was dipped into the developer, and baked to evaporate the residual solvents on the top.

Poly-dimethylsiloxane (PDMS) was used to replicate the master. Elastomer base and curing agent (Sylgard 184, Dow Corning) was mixed at a 10:1 ratio and degassed in a vacuum chamber for 5 minutes. Precursor was poured on the top of Silicon mold for 7 grams, and solidified at 80°C for 30 minutes. Plastic reservoirs from 8-well strip (Evergreen sci, USA) were glued with precursor. Additional 30 g of precursor was poured to seal reservoirs. The replica was cut and punched as shown in Fig. S1A. PDMS replica and coverslip (Tasumi, Japan) were plasma treated and bonded irreversibly. To enhance the bonding strength, device was heated for 5 minutes at 80°C hot plate. Microfluidic device was immediately filled with PBS to avoid bubble trapping.

**Preparation of microfluidic device and cell seeding.** Prior to cell culture, microfluidic device was coated with poly-D-lysine (PDL, Sigma, Germany). Reservoirs on control part was filled with 2 µg/ml of PDL solution and kept in room temperature for at least 6 hours. PDL solution was washed out before cell culture. HEK293 β5/EKAR2G1 cell-line were prepared with concentration of 2×10^6 cells/ml. Outlet reservoir connected to cell chamber was filled with 50 µl of suspension. Cells were flowing toward the cell chamber by hydrostatic pressure. After 30 minutes incubation, residual cells in the outlet were removed and replenished with fresh medium.

**Live cell imaging and monitoring of ERK biosensor.** All experiments were performed on an Eclipse Ti inverted fluorescence microscope (Nikon, Japan) with Plan Apo oil 20x (NA: 0.75) objectives controlled by Metamorph. (Molecular Devices, USA) Hamamatsu Orca R2 camera was used to acquire images at a 16-bit depth. Donor and FRET images were acquired sequentially using motorized filter wheels with the following excitation, dichroic mirrors, and emission filters (Chroma, USA): donor channel: 430/24 ×, Q465LP, 480/40 m; FRET channel: 430/24 ×, Q465LP, 535/30 m; mCherry channel: ET572/35, 89000bs, 632/60 m for dextran imaging.
exposition settings were used throughout the experiments. 440 nm, donor and FRET channel excitation, and 565 nm, red channel, LED lamps were used as light sources (Lumencor, USA), with 1.1% for 440 nm and 1.5% for 565 nm of LED power. To minimize photo-damage, exposure time was 300 ms for donor channel and 300 ms for FRET at binning $2 \times 2$.

Before the experiment, medium in each device was changed into the starvation medium, DMEM with 0.2% FBS. Each reservoir was filled with GF-containing or starvation medium for up to 200 $\mu$L. Microfluidic device was placed on the stage of microscope which is covered and controlled with heating source to stabilize the temperature at 37 °C. To prevent evaporation during the experiment, cell inlet port was sealed with transparent adhesive tape. Custom-made syringe connector was connected to the device (Fig. S1A). Built-in software of ONIX pressure pump (Millipore, USA) was used to control the valve sequence and pressure. Valve pressure was consistently set to 1.5 psi. All the protocols for stimulation experiments included 80 minutes of flow adaptation time in starvation medium to stabilize the baseline of ERK activity (Fig. S2A). Changing the opening sequence of the pressure valve, stimulation profile was temporally controlled (Fig. S1B,C).

Ratio-metric analysis of each single cell was calculated with Metamorph and ImageJ. Donor and FRET images were background-subtracted image by image. Image from FRET channel divided by the one from donor channel, and multiplied by 1000 to produce a 16-bit ratio-metric image. Projected ratio-image of time series was used to segment each coordinate of single cell. Cell clumps were discarded. Emission ratio of each cell through time was measured. The average emission ratio of 5 time points around 80-minute time point was set as the basal ERK activity level. Temporal GF stimulation can be identified using Rhodamin-dextran obtained from the time series of mCherry channel.

Results and Discussion

Experimental setup. FRET based ratio-metric biosensor have been a powerful tool for research on cell signaling dynamics. We used genetically encoded EKAR2G1 biosensor, which was verified previously (Fig. 1A)\(^3\). Biosensor was stably and homogeneously expressed in HEK293 cell-line. Meanwhile, microfluidic device was connected to computer-controlled pressure pump, and arranged to live-cell imaging system as shown in Fig. 1B. It could be stimulated precisely in temporal manner, by toggling the pressure valve on-and-off. As shown in Fig. S1A, control part was connected to each pressure valve (V1 and V2). Each reservoir of control part contains
medium with or without GF. On-and-off state of cell chamber is controlled by switching V1 and V2 (Fig. S1B). Rhodamin-dextran have been introduced in GF medium to verify temporal stimulation. By toggling two state through time, we could dynamically control microenvironment of cell (Fig. S1C). To set the baseline of our platform, ERK activity was measured in microfluidic device without any GF medium (Fig. S2A). Immediate ERK excitation was detected when the connected reservoir was pressurized, and stabilized after 80 minutes. It was reported previously that shear stress was capable to activate MAPK signaling pathway24. Stimulation was taking place after at least 100 minutes to stabilize (Fig. S2B). ERK activity of the previous 80 minutes was considered irrelevant to GF stimulation. We used the average of emission ratio intensity for 5 time points around 80-minute to normalize each single cell kinetics. Due to high signal-to-noise ratio, a large number of single cell could be captured in each microscopic view (Fig. 2C). Transient ERK excitation was evoked immediately when EGF was introduced in the cell chamber (Fig. 1D). Ultra-sensitivity, which is well-known characteristics of ERK signaling pathway25. EGF stimulation of 1 ng/ml already gave 97.7% of saturated maximum peak intensity, while 0.2 ng/ml barely excite the pathway. Figure 1E shows three different single cell kinetics from the same field of view. ERK activity of Cell 1 sustained longer than 30 minute after stimulation, while Cell 2 and 3 experienced adaptive ERK kinetics with different decaying time. This result suggest that average response analysis might miss these minor, but significant number of cell which give considerably varied behavior.

**ERK dynamics by temporally defined stimulus.** From 60 curves of sustained 25 ng/ml EGF stimulation experiment, we randomly choose 8 kinetics (Fig. 2A). In average, which might be resulted by usual biochemical method, only transient excitation kinetics could be observed. However, as shown in Fig. 2A, the single cell behavior clearly varied from cell to cell. Figure 2B showed ERK activity distribution for each time point by sustained EGF stimulation. Maximum emission ratio was shown insignificant difference amongst cells. However, cells were gradually wider the range of ERK activity distribution.

Taking advantage of computer controlled microfluidic-pump system, we looked at how temporally-varied stimulation patterns alter the ERK dynamics. EGF stimulation was applied in pulsatile regime to the cell chamber. In contrast to sustained EGF stimulation, we observed the immediate de-activation of ERK. As shown in Fig. 2C, randomly chosen curves were shown no significant difference in 3’ pulsed stimulation, compare to sustained stimulation experiment. Distribution of ERK kinetics was also remarkably coordinated (Fig. 2D). Pulsatile stimulation gave synchronized ERK kinetics throughout the population.

ERK response to the frequent pulsatile stimulation was observed. High frequency stimulation was filtered out, giving transient excitation curves similar to sustained experiments (Fig. 3A). In contrast, with low frequency stimulus, ERK activity was triggered according to input timing (Fig. 3B and C). This result suggest that ERK signaling pathway show as low-pass filter to an external stimulus.
ERK dynamics by temporally controlled MEK inhibitor. Not only GF, pharmaceutical inhibitor could be introduced to cell chamber in temporal manner (Fig. 4A). In our previous study, it was limited to observe the detailed oscillatory behavior of ERK in PC-12 cell due to the relatively larger interval time, e.g. 2 minutes. However, in this research, using HEK 293β cell line, which is more flat and relatively bigger cell, we can obtain a higher sensitivity of the biosensor signal to lower the time between frames to 30 seconds. Figure 4B showed the basal level activity of the ERK captured in every 30 second. Basal ERK oscillation of ERK was observed as reported previously. Since these oscillations cancel each other between cells, average plot could not capture this phenomena. At 0', medium containing 10 µM of MEK inhibitor, U0126, was introduced into cell chamber. We treated the cells with MEK inhibitor, U0126, to validate that the fluctuation was coming from the cascade, itself, not the artifacts of the assay. Presenting of inhibitor was not only dropping the average of basal activity for ~7.5%, but also diminishing the oscillation itself (Fig. 4B). Additionally, we could monitor the ERK activity while microfluidic device washed out the inhibitor from cell chamber. Cells were transiently excited above the original basal level, and gradually recovered to the initial state. As shown in Fig. 4C, ERK activity distribution dropped in presence of U0126, and raised up after washing out.
Conclusion

In this study, we integrated two well-characterized techniques; FRET biosensor and microfluidics. Highly sensitive and stable biosensor and computer-controlled microfluidics enable to monitor a individual ERK kinetics by various GF stimulation pattern in a high throughput manner. It was observed that sustained EGF stimulation induced a wide range of variation on ERK activities of each cell, while pulsatile stimulus synchronized kinetics over population. Low frequency of EGF pulse could retrigger the kinetics, however, high frequency stimulation was filtered out. Basal ERK oscillation was observed, and could be diminished by introducing MEK inhibitor. Removal of MEK inhibitor provoke immediate transient ERK excitation above the baseline, and gradually recover to original state. By observing a variety of ERK dynamics by temporal stimulation, we expect to gather quantitative data for single cell kinetics from different cell types, enhancing the knowledge about the pathway coordination. Exploring cellular responses by defined stimulation, this will produce 'a common language' that could lead a better interaction between experimentalists and theoreticians.

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**Author Contributions**

H.R. and S.L. designed devices; H.R., M.C., O.P. and N.J. designed the experiments; H.R., M.C. and J.S performed experiments; H.R., M.C. and S.L. analyzed experimental data; H.R., M.C. and O.P. wrote the main manuscript text. All authors reviewed the manuscript.

**Additional Information**

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