TITLE:
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CITATION:
Yoshida, Tomoki ...(et al). BTeam, a Novel BRET-based Biosensor for the Accurate Quantification of ATP Concentration within Living Cells. Scientific Reports 2016, 6: 39618.

ISSUE DATE:
2016-12-21

URL:
http://hdl.handle.net/2433/217852

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BTeam, a Novel BRET-based Biosensor for the Accurate Quantification of ATP Concentration within Living Cells

Tomoki Yoshida, Akira Kakizuka & Hiromi Imamura

ATP levels may represent fundamental health conditions of cells. However, precise measurement of intracellular ATP levels in living cells is hindered by the lack of suitable methodologies. Here, we developed a novel ATP biosensor termed “BTeam”. BTeam comprises a yellow fluorescent protein (YFP), the ATP binding domain of the ε subunit of the bacterial ATP synthase, and an ATP-nonconsuming luciferase (NLuc). To attain emission, BTeam simply required NLuc substrate. BTeam showed elevated bioluminescence resonance energy transfer efficiency upon ATP binding, resulting in the emission spectra changes correlating with ATP concentrations. By using values of YFP/NLuc emission ratio to represent ATP levels, BTeam achieved steady signal outputs even though emission intensities were altered. With this biosensor, we succeeded in the accurate quantification of intracellular ATP concentrations of a population of living cells, as demonstrated by detecting the slight distribution in the cytosol (3.7–4.1 mM) and mitochondrial matrix (2.4–2.7 mM) within some cultured cell lines. Furthermore, BTeam allowed continuous tracing of cytosolic ATP levels of the same cells, as well as bioluminescent imaging of cytosolic ATP dynamics within individual cells. This simple and accurate technique should be an effective method for quantitative measurement of intracellular ATP concentrations.

Adenosine 5′-triphosphate (ATP) is the central energy currency of all organisms. It plays critical roles in numerous cellular processes, hence intracellular ATP levels may be closely related to functions, viability, and fate of cells. For instance, assessing cellular ATP contents would be useful for the evaluation of the cytotoxicity and proliferative effects of drugs and various biological compounds1–3. Therefore, development of a technique for precise measurement of intracellular ATP levels is imperative.

Conventionally, bioluminescence assay using firefly luciferase has been used for evaluating ATP contents in cell extracts. The luciferase emits light (λ ≈ 550 nm), following hydrolysis of ATP and oxidation of luciferin. This assay typically consists of two steps: extraction of ATP from cells, and reaction of extracted ATP with firefly luciferin-luciferase. However, there are some problems in the conventional assay. First, ATP could be hydrolyzed prior to reaction with luciferase, especially by ATPases released from the cells. Second, bioluminescence output could be affected by variations in several factors, including cell number and luciferin-luciferase concentrations. In addition, the disruptive cell lysis step of this assay makes it impossible to track ATP levels of the same cells over time. Moreover, calculation of intracellular ATP concentration is a complicated task due to the need for precise measurements of both the total cell volume and total amount of ATP.

Efforts have been made to solve the aforementioned problems. For instance, one-step cell homogenization methods could suppress the decomposition of ATP in the extraction process by using modified lysis buffer1,4. Heterologous expression or introduction of firefly luciferase inside cells enabled the detection of intracellular ATP without lysis of the cells5,6. More recently, a biosensor based on an ATP-nonconsuming luciferase succeeded in monitoring ATP dynamics inside living cells7. The improved methods offer a selection of alternative ATP assays, which provide insights into the dynamics of intracellular ATP. However, these methods still have limitations, including the inability to obtain accurate intracellular ATP levels due to the variable bioluminescence output from
luciferases. Furthermore, with the ATP-consuming luciferase-based biosensors, the ATP consumption due to the luciferase-luciferin reaction itself might carry the risk of perturbing the intracellular ATP levels.

Previously, our group developed two types of genetically encoded ratiometric fluorescence biosensors for imaging of ATP levels inside single living cells: Förster resonance energy transfer (FRET)-based ATeam8–10, and circularly permuted fluorescent protein-based QUEEN 11. Different from the luciferase-based ATP assays, the two fluorescence ATP biosensors report ATP level as a ratio value; emission ratio of two fluorescent proteins for ATeam or that of two excitation wavelength for QUEEN. By using the ratio values as output signals, intracellular ATP levels can be quantitatively visualized irrespectively of the expression levels of the biosensors8–13. However, those applications are limited due to the necessity of excitation light. For instance, autofluorescence from the background, which is inevitable when using excitation light, potentially causes lower signal-to-noise ratio than bioluminescence techniques, especially in multi-well plate-based assays. In addition, phototoxicity of excitation light to cells or organisms is unavoidable.

In the present study, we developed a novel genetically encoded ratiometric luminescent ATP biosensor, which is based on bioluminescence resonance energy transfer (BRET) and requires no excitation light. With this biosensor we succeeded in measurement of accurate intracellular ATP concentrations in cytosol and the mitochondrial matrix in a population of living mammalian cells. In addition, we demonstrated continuous measurement of intracellular ATP concentrations within the same cells. Of particular note, the biosensor enabled quantitative bioluminescent imaging of intracellular ATP dynamics within individual cells.

Results

Development and In Vitro Characterization of a BRET-based ATP Biosensor. We developed a BRET-based ATP biosensor termed BTeam (BRET version of ATeam). BTeam comprises mVenus14 (a variant of yellow fluorescent protein, YFP), the ATP binding domain of the ε subunit derived from Bacillus subtilis F1F-ATP synthase, and NanoLuciferase (NLuc), an ATP-nonconsuming luciferase with characteristics of high pH stability and brightness15 (Fig. 1A). BTeam emits lights in the presence of the NLuc substrate, furimazine. As shown in Fig. 1B, BTeam has emission peaks at 455 and 527 nm derived from NLuc and YFP, respectively. BTeam showed elevated BRET efficiency in the presence of ATP, as demonstrated by its emission spectra changes where ATP concentration is increased.

Figure 1. Development of the BRET-based Biosensor BTeam. (A) Schematic drawing of BTeam. The ATP-free form of the ε subunit is extended and flexible16, which separates YFP and NLuc, resulting in low BRET efficiency (left). On the other hand, binding of ATP to the ε subunit causes a conformational change16 that draws YFP and NLuc closer, increasing BRET efficiency (right). It should be noted that the ε subunit reversibly binds and dissociates ATP without its hydrolysis. (B) ATP-dependent luminescence spectral changes of purified BTeam. (C) ATP-dependent BRET ratio changes of purified BTeam. BRET ratio (YFP/NLuc luminescence ratio, mean ± SD) of purified BTeam (closed bar) and BTeamR122K/R126K (open bar) at different ATP concentrations were calculated from luminometer measurements at 25 °C (n = 3, independent repetitions). BTeamR122K/R126K is a variant of BTeam designed not to bind ATP.
increase in ATP concentration and YFP intensity is inversely proportional to increase in NLuc intensity. The response of the emission ratios of YFP/NLuc (the BRET ratio) of BTeam could represent ATP concentrations (Fig. 1C). However, the values of the BRET ratio were almost invariant for BTeamR122K/R126K, in which the ATP-binding ability was disrupted by replacing Arg-122 and Arg-126 with lysine (Fig. 1C and Supplementary Fig. S1). The results indicate that the BRET ratio is dependent on the conformational changes of the ε subunit of BTeam upon ATP binding.

Next, we examined the chemical properties of purified BTeam (Fig. 2). Among the nucleotides tested (10 mM ATP, ADP, UTP, GTP, or CTP), only ATP increased the BRET ratio of BTeam, while the other nucleotides did not (Fig. 2A). These results are in accordance with previous results regarding ATeam16. A slight decrement in the BRET ratio for the other nucleotides was observed, the cause of which remains unknown. However, considering that the decrement was minimal and that those nucleotide concentrations were 3.7–11.3 times lower than that of ATP inside cells17, the effects of other nucleotides may be negligible for the ATP measurement with BTeam. BRET ratio of BTeam was almost constant over a range of pH values from pH 7.1 to 8.3, with ATP concentrations ranging from 0–10 mM (Fig. 2B). Previous study has reported that cytosolic and mitochondrial pH is about 7.3 and 8.0, respectively18. Consistently, we determined the cytosolic pH of HeLa cell to be 7.34 ± 0.03. Thus, BTeam had appropriate pH stability for evaluating cytosolic and mitochondrial ATP levels. The affinity of BTeam for ATP was weakened with an increase in temperature, as indicated by the values of K_{0.5} which demonstrated an exponential increase from 1.7 to 4.1 mM, concurrent with a temperature increase from 25 °C to 40 °C (Fig. 2C). These results suggest that temperature control is crucial for the precise measurement of ATP with BTeam.

Quantitative Measurement of ATP Levels in Living Mammalian Cells. Further characterization of BTeam was carried out at 37 °C using live HeLa cells stably expressing BTeam. When BTeam without any targeting signal sequence (cyt-BTeam) was expressed in the cells, it was localized primarily in the cytosol with a slight fraction in the nucleus, as examined by YFP fluorescence (Fig. 3A). After addition of the NLuc substrate reagent to the medium, luminescence was emitted from the cells. The luminescence intensity decreased as time elapsed. In contrast, the BRET ratio from the cells remained almost constant for 90 min (Fig. 3B). Furthermore, neither the concentration of the substrate reagent in the medium (Fig. 3C) nor the density of the cells (Fig. 3D) affected the values of the BRET ratio from the cells. These characteristics mark a clear improvement over other luciferase-based assays, whose output is sensitive to cell number and luciferin concentrations. These distinct advantages of BTeam as a ratiometric biosensor should translate to more accurate quantification of intracellular ATP levels within living cells. Additionally, the absence of ATP hydrolysis for emission processes of BTeam should allow for more precise evaluation of ATP levels compared to other methods using ATP-consuming luciferases19.

Next, we demonstrated the measurement of basal intracellular ATP levels in several cell lines with BTeam (Fig. 4). Cells transiently expressing BTeam were incubated in a 96-well plate, and the basal intracellular ATP levels at 60–70% confluence were measured. A regression model using Hill’s equations was obtained, based on the BRET ratios of purified BTeam versus ATP concentrations at 37 °C as a standard curve for the calculation of ATP concentrations (Fig. 4A). The BRET ratio values were in the range from 0.385 to 0.409 (Fig. 4B). The calculated intracellular ATP concentrations were 3.8 ± 0.1 mM in HeLa, 3.7 ± 0.1 mM in Cos7, 4.1 ± 0.2 mM in HepG2, 3.7 ± 0.04 mM in HEK293A, 3.9 ± 0.3 mM in PC12, 3.7 ± 0.2 mM in B16F10 cell lines (Fig. 4C). ATP concentrations calculated from BTeam measurements were nearly equal to those estimated by a conventional firefly luciferase assay (see Materials and Methods for detail). The calculated ATP concentrations are also consistent with the previously reported values (2.5 ± 1.2 and 3.1 ± 2.1 mM for normal and tumor cells, respectively)17. Considering these results, BTeam exhibits promising potential as an accurate and effective tool for quantitative measurement of intracellular ATP concentrations. ATP levels in the mitochondrial matrix were also measured with mit-BTeam,
a variant BTeam targeted to the mitochondrial matrix (Fig. 3A and Supplementary Fig. S1). The values of the BRET ratio from four transiently transfected cell lines were in a narrow range from 0.297 to 0.318 (Fig. 4D). The calculated mitochondrial ATP concentrations were 2.5 ± 0.1 mM in HeLa, 2.4 ± 0.03 mM in HepG2, 2.5 ± 0.1 mM in PC12, and 2.7 ± 0.1 mM in B16F10 cells (Fig. 4E). Mitochondrial ATP levels were approximately 1.5 mM lower than cytosolic levels, a consistent trend reported in previous studies. A potential explanation for the disparity between cytosolic and mitochondrial ATP levels is a rapid exchange of ATP and ADP between the cellular compartments via adenine nucleotide translocator. Additionally, ADP is a primary building block for ATP synthesis, and evidence of higher mitochondrial ADP levels has been previously reported. Our results further corroborate the difference between cytosolic and mitochondrial ATP levels.

We used BTeam to trace changes in cytosolic ATP levels of the same cells. It was clearly evident that the BRET ratio of HeLa cells stably expressing BTeam gradually decreased in the starved condition, and rapidly recovered after replenishment of nutrients (Fig. 5E). The result indicates that BTeam is adequate for continuous measurements of accurate intracellular ATP of the same cells.

**Ratiometric Luminescent Imaging of Intracellular ATP Levels of Single Living Cells.** Finally, ATP levels in individual living cells were imaged with BTeam using a microscope. NLuc and YFP emissions from a BTeam-expressing cell were separated with a beam splitter and were simultaneously captured with an EMCCD camera. Even at single-cell level, the BRET ratio of the cytosol was significantly higher than that of mitochondria (Fig. SCD), consistent with the data obtained from a population of cells using a luminometer (Fig. 4). It should be noted that, due to the filter settings, the different basal values in the BRET ratio of the cells were observed between imaging assays and luminometer assays (see Material and Methods for details). We then imaged cytosolic ATP levels of single living cells over time. When 2-deoxyglucose and oligomycin A, inhibitors of glycolysis and oxidative phosphorylation, respectively, were added to the medium, a rapid decrease in the BRET ratios of the cells was observed (Fig. 6A,B and Supplementary video 1). The result indicates that BTeam is not only adequate...
for quantitative measurement of intracellular ATP of a population of cells, but also is capable of exploring ATP dynamics at the single-cell level.

**Discussion**

In the present study, we developed a novel BRET-based ATP biosensor, termed BTeam. BTeam presents ATP concentrations as a value of the BRET ratio (emission ratio of YFP/Nluc). Based on this property, BTeam can steadily measure ATP even though emission intensities are altered, unlike other bioluminescent ATP assays. With this biosensor, we demonstrated accurate measurements of intracellular ATP concentrations of a population of living cells, as well as longitudinal monitoring of cytosolic ATP levels of the same cells. The intracellular ATP levels measured with BTeam were nearly equal to those estimated from cell extracts using firefly luciferase assay, which was consistent with previous report17. Our results further corroborate the importance of intracellular ATP levels in the health conditions of cells, since assessing ATP contents in cell extracts has been used for potential explanations of the cytotoxicity and proliferative effects of drugs1–4. Due to its simple and accurate properties for measurements of cytosolic and/or mitochondrial ATP of multiple cell populations, BTeam would be useful for high-throughput drug screenings that result in changes in ATP levels.

Fluorescence biosensors have recently been applied to imaging of ATP levels, or the ATP/ADP ratio, in living cells8–11,23. In general, temporal and spatial resolutions are better in imaging with fluorescence biosensors than with luminescent biosensors because fluorescent biosensors can emit stronger signals. However, BTeam may also have some advantages in live cell ATP imaging over fluorescence-based biosensors. For instance, BTeam may be potentially useful for investigating ATP dynamics in plants, because unlike fluorescence biosensors it...
does not require excitation light, which would perturb photosynthetic activity and induce strong autofluorescence from chlorophyll. In addition, BTeam may be used in conjunction with optogenetics techniques that utilize light-sensitive actuators such as channelrhodopsin \(^{24}\), which could be skewed by excitation light for traditional fluorescent biosensors.

In conclusion, BTeam is as an effective tool for accurate measurement of intracellular ATP concentrations, and it should be useful in a wide variety of applications.

### Methods

#### Chemicals and Cells.
Cells were routinely cultured at 37 °C in 5% CO\(_2\) in the following growth medium: Dulbecco's modified Eagle's medium (DMEM) with 10% fetal bovine serum (FBS) for HeLa, Cos7, HEK293A, and HepG2; DMEM with 10% FBS and 5% horse serum for PC12; and Minimum Essential Media (MEM) with 10% FBS for B16F10 cells. Cells expressing BTeam were generated by transfection with plasmids carrying BTeam cDNA, using Lipofectamin® 2000 reagent (Invitrogen). The transfected cells were assayed between 1 and 3 days after transfection. To establish stable cell lines expressing BTeam, cells were cultured in growth medium containing 0.75 mg/mL G418 for over 2 weeks after transfection. Subsequently, cells stably expressing BTeam were isolated. All chemical reagents and other solvents used were analytical grade. Nucleotides, including ATP, were pre-mixed with equimolar magnesium chloride before use; the term ATP in the present experimental results represents Mg-ATP.

#### Gene Construction.
A modified cDNAs for mVenus, with an in-frame deletion eliminating 11 amino acids from the C-termius, and NLuc (Promega) with a deletion of 3 amino acids from the N-terminus (with permission from Promega), were amplified by PCR, respectively. The PCR products were ligated into the XhoI-ClaI and EcoRI-HindIII sites in pRSET-AT1.038, respectively, to generate pPRSET-BTeam for expression of BTeam in \(E. coli\). The cDNA of BTeam was excised from pRSET-BTeam with XhoI and HindIII (Thermo Scientific). The restriction fragment was then ligated into the XhoI-HindIII sites of pcDNA3.1 (−) (Invitrogen) for mammalian expression. The constructs used in the present study are illustrated in Supplementary Fig. S1.

#### Purification of BTeam.
The BTeam protein was purified with a MagneHis™ Protein Purification kit (Promega) with minor modifications. \(E. coli\) strain Rosetta (DE3) carrying the BTeam plasmid was cultured in 2 × YT medium at 25 °C for 40–60 h. Cells were collected by centrifugation at 3,200 \(\times g\) for 5 min and re-suspended in 10 mM PBS (pH 7.4) containing 0.5 mM MgCl\(_2\) and 0.05% Triton X-100 (Buffer A). The suspension was disrupted by sonication, and centrifuged at 20,000 \(\times g\) at 4 °C for 10 min. The supernatant was applied to MagneHis™ Ni-Particles pre-equilibrated with PBS. After two washes with the manufacturer's wash buffer, the purified BTeam protein was eluted with 200 mM imidazole. The purified proteins were stored at 4 °C, and used within 2 days after purification.

#### Characterization of Purified BTeam.
Luminescence spectra of the purified BTeam were investigated at 25 °C in 50 mM Mops–KOH buffer (pH 7.3), containing 50 mM KCl, 0.5 mM MgCl\(_2\), and 0.05% Triton X-100 (Buffer A), using a FP-6500 spectrofluorometer (Jasco). Luminescence spectra from 370 to 720 nm at 200 nm/ min of scanning speed were immediately scanned after addition of 2 μL Nano-Glo® Luciferase Assay Substrate (Promega) to a 400 μL reaction solution of BTeam and 0–8 mM ATP.

Emissions of YFP and NLuc from purified BTeam were measured for 0.5 sec using an ARVO-X3 (PerkinElmer) equipped with the following emission filters (Semrock): FF01–515/LP for YFP emission; and FF01–450/70 for NLuc emission. After addition of the substrate reagent to a reaction mixture composed of purified BTeam and 0–12 mM ATP in Buffer A, the BTeam YFP/NLuc emission ratio was measured at 25–40 °C. For measurements...
To avoid disturbance from the BTeam released from dead cells, after addition of NLuc substrate reagent to the Mops-KOH (pH 6.3–7.5) or Hepes-KOH (pH 7.7–8.3), 50 mM KCl, 0.5 mM MgCl₂, and 0.05% Triton X-100.

Cell numbers and diameters were measured using a CDA-500 particle analyzer (Sysmex). and cell numbers in the aliquot, respectively, and CV represents cell volume estimated from the diameters of the following equation: Cellular ATP level

The temperature dependence of the ATP affinity ($K_0.5$) of BTeam was investigated at 25–40 °C with a range of 0–12 mM ATP in Buffer A. The values of $K_0.5$ were calculated by using Hill equations; $R = (R_{max} - R_{min}) \times [S]^n/([S]^n + K_{0.5}^n) + R_{min}$, where $n$ is the Hill coefficient, $R_{max}$ and $R_{min}$ are the maximum and minimum BRET ratios, respectively.

**Luminometer assays.** Cells expressing BTeam were seeded into a 96-well plate, and incubated for 24–36 h at 37 °C in 5% CO₂ in growth medium. After washing with DMEM containing 5% FBS, cells were incubated for 30 min in phenol red-free DMEM containing 5% FBS, 20 μM HEPES (pH 7.5), and 30 μM NLuc inhibitor to avoid disturbance from the BTeam released from dead cells. After addition of NLuc substrate reagent to the medium, microplates were incubated at 37 °C for 20 min to stabilize the temperature in a CO₂ incubator. Then, luminescence emissions from the cells were measured at 37 °C for 0.5 sec using an ARVO-X3 equipped with the same emission filter set for the assay of the purified BTeam as described above. Figure 4 (B,D) illustrates the basal BRET ratio of cells transiently expressing cyt-BTeam or mit-BTeam measured 30–36 h after passage around 70% confluency. To calculate ATP concentrations from the BRET ratios of the cells, a regression model was obtained by using Hill equations for standard curve. In this study, the following coefficient values were used for the standard curve; 0.521 for $R_{max}$, 0.178 for $R_{min}$, 3.11 for $K_0.5$, and 2.58 for $n$, respectively, which were calculated from the plots of the BRET ratio values of purified BTeam against ATP concentrations at 37 °C.

Total cellular ATP determination with firefly luciferase was carried out using a firefly luciferase kit (Toyo B-net) with slight modifications. Cells were trypsinized, and suspended in growth medium. Aliquots of the suspension were collected by centrifugation at 6,000 × g for 3 min at 4 °C. Pellets were disrupted with Glo Lysis Buffer (Promega) for 5 min followed by centrifugation at 20,000 × g for 5 min at 4 °C. ATP concentration in the supernatant was quantified with the reagents provided in the kit. The cellular ATP levels were calculated using the following equation: Cellular ATP level = [ATP]/(CN × CV), where [ATP] and CN represent the ATP amount and cell numbers in the aliquot, respectively, and CV represents cell volume estimated from the diameters of the trypsinized cells. Cell numbers and diameters were measured using a CDA-500 particle analyzer (Sysmex).

**Microscopy assay.** The transfected HeLa cells were plated on a collagen coated glass bottom dish, and subjected to imaging. Prior to imaging, the medium was changed to phenol red-free DMEM containing 5% FBS and 30 μM NLuc inhibitor (Promega), and the cells were incubated for 20 min at 37 °C in 5% CO₂. BRET imaging was immediately carried out after addition of the substrate reagent to the medium. Imaging was performed on a Nikon Eclipse Ti-E inverted microscope with a perfect focus system (Nikon Instruments) using a CFI Plan Apo VC 60x oil-immersion objective lens, 1.40 numerical aperture (Nikon Instruments). Dual-luminescence emission ratio imaging was achieved by using a W-View Gemini optical splitting device (Hamamatsu Photonics) equipped with FF01–450/70 (Semrock) for the NLuc emission filter and FF01–542/27 for the YFP emission filter. Luminescence emission from BTeam was obtained with a 5 sec exposure time using an EMCCD camera (Andor Technology). During imaging, the cells were maintained at 37 °C in 5% CO₂ on a microscope by using a top-stage incubator (Tokai Hit). Analysis of images was performed using MetaMorph (Molecular Devices). The YFP/NLuc emission ratio was calculated by using the integrated intensity of YFP emission and NLuc emission within the area of the whole cell (cytosol) or the area surrounding the mitochondria inside individual cells, respectively. The
ATP levels were illustrated as pseudo color images generated by the emission ratio of YFP/NLuc calculated from YFP and NLuc luminescence images.

Cytosolic pH measurement. The cytosolic pH of HeLa cells was measured with BCECF (2′,7′-bis(carboxyethyl)-5(6)-carboxyfluorescein) as the following protocol. The cells were treated with 10 μM BCECF acetoxymethyl ester (Molecular probes) in Hank’s balanced salt solution (HBSS) for 15 min at 37°C, followed by washing with HBSS twice. Fluorescence intensity of BCECF of the cells (n = 6) was measured at 37°C in DEMEM using a Nikon TE2000 (Nikon Instruments) equipped a CFI Plan Apo VC 60x oil-immersion objective lens, 1.40 numerical aperture (Nikon Instruments). The following filters (Semrock) were used for the measurement; excitation filters (FF01-445/20 and FF01-482/18), an emission filter (FF01-520/35), and a dichroic mirror (FF495-Di03). Images were captured with an ORCA-AG cooled CCD camera (Hamamatsu photonics). Calibration curve was obtained by imaging BCECF-loaded cells in pH-controlled buffers containing 150 mM KCl, 20 mM NaCl, 0.5 mM MgCl2, 0.5 mM CaCl2, 10 mM MES, 10 mM HEPES, 10μM monensin and 10μM nigericin.

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Acknowledgements
We thank Brock Binkowski for technical advice and for NLuc inhibitor, James Alan Hejna, Promise Ogor and Soaad Alfaqaan for critical assessment of this manuscript. The work was supported by JSPS KAKENHI Grant Number 26291027, 16K14709 (to H.I.), and by a grant from Nagase Science Technology Foundation (to H.I.), and by the Platform Project for Supporting Drug Discovery and Life Science Research (Platform for Dynamic Approaches to Living Systems) from the Ministry of Education, Culture, Science and Technology (MEXT) and the Japan Agency for Medical Research and Development (AMED).

Author Contributions
T.Y. and H.I. conceived the experiments; T.Y. conducted the experiments; T.Y., H.I. and A.K. analyzed the results and wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep
Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Yoshida, T. et al. BTeam, a Novel BRET-based Biosensor for the Accurate Quantification of ATP Concentration within Living Cells. Sci. Rep. 6, 39618; doi: 10.1038/srep39618 (2016).

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