Eastern Staining: A Simple Recombinant Protein Detection Technology Using a Small Peptide Tag and Its Counter Partner Which is a Fluorescent Compound

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Small peptide tags such as c-myc, HA, or FLAG tag have facilitated efficient Western-blotting of proteins of interest especially when specific antibodies for the proteins are not available. However, the conventional Western-blotting requires the multi-steps process taking at least several hours up to two days. With examples of various applications, here we show a convenient and time-saving method for protein detection which employs a fluorescent chemical BDED and its binding peptide RC-tag. And we propose "Eastern staining", as a standard term for protein detection method using fluorescent chemicals and their binding small peptide tags. Eastern staining may substitutes for the time-consuming "immuno-staining" in many versatile applications.

SYNOPSIS

Small peptide tags such as c-myc, HA, or FLAG tag have facilitated efficient Western-blotting of proteins of interest especially when specific antibodies for the proteins are not available. However, the conventional Western-blotting requires the multi-steps process taking at least several hours up to two days. With examples of various applications, here we show a convenient and time-saving method for protein detection which employs a fluorescent chemical BDED and its binding peptide RC-tag. And we propose "Eastern staining", as a standard term for protein detection method using fluorescent chemicals and their binding small peptide tags. Eastern staining may substitutes for the time-consuming "immuno-staining" in many versatile applications.
INTRODUCTION

Recombinant protein expression and its detection is a crucial technique for biomedical research. Conventionally, denatured proteins in cell lysate are separated according to their molecular masses by electrophoresis, blotted to a membrane and detected by antibodies which are visualized by attached fluorophores or chemical reactions. Expression of the protein of interest is judged based on the size and antigenicity of the detected protein. This Western blotting technique is, however, highly dependent on the availability, affinity and specificity of antibodies which vary significantly from one to another. Small peptide tags such as c-myc, HA, FLAG, and polyhistidine have been introduced into recombinant proteins to simplify detection and purification of proteins\(^1\). Although the antibodies for the common tags are less variable compared to specific epitope-targeting antibodies, batch to batch variations of the antibody titre and quality are still inevitable.

Bioorthogonal chemistry, which covalently labels biomolecules with exogenously delivered chemical probes in a highly specific manner in living systems, has caught attention during last decade as a strategy for simple and easy labeling of proteins\(^2\). Due to the much smaller size of chemical probes compared to that of fluororescence proteins, which often perturbs the function of conjugated proteins because of bulky size, the function and structure of proteins are normally not affected by bioorthogonal labeling. The chemical labeling also overcomes the variation problem of antibodies described above. Among the switchable fluorescent chemical probes designed to become fluorescent by binding to small peptide tags are biarsenicals FlAsH-EDT\(_2\) and dimaleimide fluorogens\(^3\) which bind to 4 and 2 cysteine residues. We recently reported a novel bodipy-diacrylate compound 4b (BDED) and its binding peptide RC-tag which has 2 cysteine residues placed next to arginines\(^4\). As the BDED compound is cell permeable and changes its emission spectrum from yellow to green by forming covalent bonds with the 2 cysteins in the RC-tag, both intracellular and extracellular proteins can be labeled with visual confirmation of proper binding to the target. We have shown that RC-tagged exogenous protein labeled by BDED is conveniently visualized in SDS-polyacrylamide gel. As summarized in Figure 1, the whole procedure of this method takes less than 2 hours while conventional Western-blotting takes at least 7 hours.

RESULTS AND DISCUSSION

Evaluation of the applicability of Eastern staining on various functional mammalian proteins

As a demonstration, we fused the RC tag to the N-terminus of several functional proteins, such as human \(\beta\)-actin (42 kDa), human \(\alpha\)-tubulin (52 kDa), and mouse fatty acid-binding protein 7 (Fapb7, 14 kDa) (Figure 2A). Recombinant proteins fused to RC tag together with c-myc tag were expressed in HEK 293A cells. After 48 hours, transfected cells were stained with BDED and the total cell extract was subjected to SDS-PAGE as depicted in Figure 1.

The size of green fluorescence bands appeared on the gel exactly corresponded to the expected molecular weight of respective proteins fused with tags (Figure 2B, left). The identities of the fluorescence bands were confirmed by the followed detection of c-myc tag by Western blotting (Figure 2B, middle). Perfect overlapping of the protein bands shown by BDED labeling and by c-myc tag Western blotting confirms the specificity and reliability of Eastern staining among the whole proteome (Figure 2B, right). Sensitivity of Eastern staining on RC-tag was comparable to that of c-myc Western blotting by an anti c-myc antibody 9E10. Small amount of proteins, which are not distinguishable by Coomassie brilliant blue staining, are efficiently visualized by Eastern staining (Figure 2B, lanes 3, 4, 5).

Application of Eastern staining in immunoprecipitation

Eastern staining can be applied to various experiments, for example, immuno-precipitation (IP). IP is a widely used technique to isolate or enrich a specific protein from mixed proteins. The performance of IP is usually analyzed by Western-blotting using the respective antibodies against the proteins of interest. Therefore, small peptide tags such as c-myc tag, HA tag, or FLAG tag
are useful tools for IP of proteins for which the antibodies are not available. If the RC-tag is used in conjunction with those small tags, IP can be verified in a short processing time via Eastern staining. We immunoprecipitated hFabp7 fused with the c-myc tag and RC tag (RC·myc·Fabp7) using a monoclonal c-myc antibody (9E10). Transfected cells were stained with BDED and the total cell extract was subjected to IP followed by SDS-PAGE. Eastern-staining facilitated the visualization of isolated proteins without laborious procedures of Western blotting, i.e., protein transfer to membrane, incubation with antibody, and multiple times of washing.

In Western blotting for immunoprecipitated proteins, the
heavy chain and light chain of antibody are also detected as shown in Figure 3B (right), if the antibodies for Western blotting and IP are from the same species. It can make it difficult to detect the proteins when their sizes are similar with those of antibody chains. However, Eastern staining is free from such unnecessary bands resulted from the antibodies (middle). The other experiments to study protein-protein interactions, such as “Co-IP” (protein complex-immuno-precipitation) or “pull-down assay” would be benefited by the Eastern staining.

Combination of RC tag with His-tag to detect the expression of His-tagged protein in crude cell extract by Eastern staining

As an other example, we combined an RC-tag with a His-tag which is prevalently used in the affinity-purification of recombinant proteins expressed in diverse biological systems such as E. coli, yeast, insect cells, or mammalian cells. Purification is performed by a time-consuming procedure employing the costly nickel column. Therefore, the expression of the His-tagged proteins in the biological system should be confirmed before conducting the affinity purification in a large scale. When a target protein is favorably expressed in E. coli, the recombinant protein induced by the lac operon takes up a significant amount in total proteome profile. Therefore, His-tagged proteins are obviously observed by staining the total proteome resolved in a gel by Coomassie brilliant blue or silver staining in our empirical observations. More sensitive methods, however, are required when the target proteins are produced in eukaryotic cells. Besides Western blotting using antibody against His-tag, different types of reporter agents, e.g., fluorophores, horse radish peroxidase (HRP), or alkaline peroxidase (AP), conjugated to nickel-nitrilotriacetic acid (NTA) are developed for faster and easier detection. While those His-tag affinity probes are convenient to use, they are hardly applicable to the detection of small amount of His-tag fusion proteins since the transient binding between the metal ion and imidazole ring lacks in sensitivity and stability. We thus combined an RC-tag to a His-tagged protein and conducted Eastern staining for the convenient in-process detection of proteins. His-tag was additionally fused to the RC-tagged Fabp7, and the recombinant Fabp7 (RC·myc·Fabp7·His) was expressed in mammalian cells. Small amount (10 µg) of total proteins extracted from the transfected cells were incubated with BDED in vitro at room temperature. When the incubated total lysate was resolved in SDS-PAGE, His-tagged Fabp7 was

Figure 2. RC tag-mediated “Eastern staining” shows specific and reliable protein detection. (A) Chemical structure of BDED. (B) Schematic illustration for the RC-tagged recombinant proteins. (C) Protein extracts from the cells, transfected with the vectors in B and stained with BDED, were analyzed in the SDS-PAGE and α-myc Western blotting was followed to confirm the expression of tagged proteins. Coomassie brilliant blue staining data of the gel was prepared with equal amount of protein per lane for electrophoresis. M: protein size marker. N: no transfection (Gel scan Ex/Em = 488/SP 526 nm)
visualized due to the BDED bound to the RC-tag (Figure 4A). We compared the sensitivity of RC-tag mediated Easterning staining with that of a commercial dye for His-tag fusion proteins, InVision™ His-tag In-gel Stain (Invitrogen). A fluorescent dye, conjugated to Ni²⁺-NTA, was applied directly to the protein gel after SDS-PAGE. As shown in Figure 4B, the affinity dye was able to visualize microgram quantity of fusion protein purified form E. coli protein extract (Figure 4B, lanes H-Fabp7 (5) and H-Fabp7 (10)), but the His-tagged Fabp7 in mammalian cell extract was not detectable.

We recommend to stain RC-tag with BDED in live cells for general Eastern staining of proteins in broad molecular weight range. When the total lysate of HEK293 cell was incubated with BDED in vitro, several protein bands of unknown identity were found at around 50 kDa to 90 kDa. Therefore, in vitro Eastern staining of RC-tag would be applicable to the proteins at low molecular weight range (less than 50 kDa).

CONCLUSION AND PROSPECTS

Chemical tag is an emerging technology for convenient labeling of proteins in live cell. Since the pioneering invention of FlAsH/Tc-tag system by Tsien et al., diverse chemical tags have been competitively developed with different properties and advantages. Enzyme tags such as HaloTag or SNAP tag provide high specificity with minimal background, short peptide tags such as AviTag or LAP provide specific labeling by using transacting ligases, and self-affinity peptide tags such as Tc-tag and RC-tag enable minimally invasive tagging and intracellular staining with cell-permeable dyes.
Although most tags for chemical labeling were developed to facilitate optical imaging of proteins by circumventing the drawbacks of fluorescent proteins, application of those tags can be expanded much further for various purposes. In this report, we introduced “Eastern staining using RC-tag and its binding fluorescence dye BDED” as a convenient protein detection method with examples of its application in biological experiments. This chemical tag-mediated protein detection method will significantly simplify a lot of experimental protocols which have required multi-step immuno-staining.

MATERIALS AND METHODS

Expression vectors
pc-RC·myc·Cherry was described in the previous report. pc-RC·myc·Fabp7: cDNA of mouse Fabp7 was prepared by RT-PCR using the total RNA of NS5, a mouse neural stem cell line. The open reading frame (ORF) of Fabp7 was PCR-amplified with sense and antisense primers and the resulting PCR product was cloned into the BamHI/XhoI sites of pc-RC·myc which encodes the RC tag, together with the c-myc tag. This clone was named as pc-RC·myc·Fabp7

- Fabp7
  Fabp7_S_1: CCGCGATCCATGGTAGATGCTTCTCGGC

- Fabp7 AS_1: CCCGTCGAGTGGCCTTTTCAACAGCGAACAGCA
  (Underlined are the restriction enzyme sites inserted for cloning).

- pc-RC·myc·Actin and pc-RC·myc·Tubulin: DNA fragment encoding RC·myc was prepared by digestion of the pc-RC·myc with NheI/BamH1, separated by electrophoresis and eluted from the gel. Purified DNA fragment was cloned into the NheI/BglII sites of pAcGFP1-Actin or pAcGFP1-Tubulin (Clontech) to exchange the GFP ORF with the DNA fragment encoding RC·myc.

pc-RC·myc·Fabp7·His: Fabp7 primers_2 were used to PCR-amplify the expression cassette of Fabp7 with His tag at the C-terminus. The produced PCR product was digested with BamHI/XhoI and inserted into the BamHI/XhoI site of pc-RC·myc.

- Fabp7 S_2: CCGCGATCCATGGTAGATGCTTTTCTGGGCAACCT
- Fabp7 AS_2: CCCGTCGAGTCAATGGTAGATGCTTCTCTGACAGCA

- pQE30-Fabp7: PCR-amplified ORF of Fabp7 was digested with BamH1/XhoI and inserted into the BamH1/SalI sites of

Figure 4. Eastern staining performed in in vitro provided convenient confirmation of recombinant protein expression. (A) Total protein (10 µg) from the 293A cells transfected with the pc-RC·myc·Fabp7·His was incubated with BDED (1 µM, 15 min, RT) and analyzed in SDS-PAGE. M: protein size marker, (B) 293A cells were transfected with pc-RC·myc·Fabp7·His and the total protein extracts was resolved in SDS-PAGE. The gel was visualized on a fluorescence gel scanner after staining by using the InVision™His-tag In-gel staining kit (Invitrogen) (left) and stained with Coomassie Brilliant Blue (right). Purified His-tagged Fabp7, produced in E. coli, was included as a positive control for the His-tagged protein. H-Fabp7 (5) and H-Fabp7(10) mean 5 µg and 10 µg of purified recombinant Fabp7, respectively. BM is the BenchMark™His-tagged Protein Standard (Invitrogen) as a molecular weight marker.
pQE30 (Qiagen) for the expression of N-terminal His-tagged Fabp7 in E. coli.

Fabp7 S_1: CCCGATCATGGTAGCTTTCTGC
Fabp7 AS_1: CCCCTCGAGTGCCTTTTCAACAGCGA-
CAGCA

Expression vectors used are pc-RC·myc·Cherry, pc-RC·myc·Fabp7, pc-RC·myc·Actin, pc-RC·myc·Tubulin, and pQE30-Fabp7. The generated clones were confirmed by DNA sequencing and purified by Qiagen Midiprep column for transfection.

Cell culture and transfection
HEK293A cells, an immortalized line of primary human embryonic kidney cells, were purchased from Invitrogen and maintained in DMEM containing 10% fetal bovine serum, 1% antibiotics-antimycotics reagent. Cell culture materials were purchased from Invitrogen. For transient transfection, cells were plated at a density of 2 × 10^5 cells/well in 12 well plates and 500 ng of plasmid DNA were transfected with Lipofectamine 2,000 (Invitrogen). After incubation for two days, the transfected cells were harvested for experiments.

Eastern staining
BDED was reconstituted in DMSO as 1 mM stock solution, and stored at -20°C. Immediately before staining, medium in the wells were drained and the BDED diluted in the pre-warmed serum-free medium was added directly onto the cells. After incubation (20 min, 37°C), cells were washed with PBS and subjected to SDS-PAGE, IP, or Western-blotting.

SDS-PAGE, gel-scanning, Western blotting, and Coomassie Brilliant Blue staining
Total protein was extracted by using CellLyticM™ cell lysis solution (Sigma). Generally 10 μg of the protein/well was loaded in SDS-PAGE gel for gel-scanning. NuPAGE Novex Bis-Tris Gels (Invitrogen) were used for PAGE and the gel was scanned in Typhoon 9410 Gel Scanner (GE Healthcare) using a 488 nm laser and 526SP emission filter for the RC-tag: BDED observation. After gel scanning, proteins were transferred onto PVDF membrane for Western blotting. The mouse monoclonal anti-c-myc (clone 9E10., Santa Cruz. Sc-40) antibody was visualized by secondary goat anti-mouse IgG conjugated with Cy5 (Invitrogen, A10524) which was detected by a 633 nm laser and 670BP emission filter on the Typhoon 9410 Gel Scanner. For Coomassie bright blue staining, the gel was stained in a staining solution (50% MeOH, 40% water, 10% glacial acetic acid, 0.25% Coomassie Brilliant Blue R-250) for 2-4 hours. Then, the background staining was removed by washing in destaining solution (50% MeOH, 40% water, 10% glacial acetic acid) by changing the destaining solution several times.

Recombinant His-Fabp7 preparation
pQE30-Fabp7 was transformed into E. coli strain SG13009 (Qiagen), which provides very stringent control of protein induction. Single colony of SG13009 (pQE30-Fabp7) was inoculated in 5 mL of LB medium containing ampicillin (100 μg/mL) and cultured for overnight. On the following day, 1 mL of the seed culture was put in 250 mL of LB medium and incubated until the optical density reaches 0.6 at 600 nm (OD600). Isopropyl β-D-1 thiogalactopyranoside (IPTG) was added to the medium at the working concentration of 0.5 mg/mL and recombinant protein was induced for 5 hours at 30°C. Cells were harvested by centrifugation and the His-tagged Fabp7 was purified using the QIAexpress kit (Qiagen) according to the user manual. Purify of the recombinant protein was checked by SDS-PAGE and Coomassie brilliant blue staining.

Immuno-precipitation
Mammalian cell expression vector pc-RC·myc·Fabp7 was transfected into 293A cells. After two days incubation, cells were stained with BDED and 100 μg total lysate, prepared by using CellLyticM™ (Sigma), was incubated with 200 ng of monoclonal c-myc antibody (9E10) for overnight at 4°C. On the following day, 20 μL of Protein G-conjugated Sepharose beads (Pierce) were added to each of the lysate-antibody mixtures and incubated for 4 hours at 4°C with agitation. After three times repeated centrifugation and washing, 100 μL of 1 × LDS gel loading buffer (Invitrogen) was added to the beads and heated at 70°C for protein denaturation. IP samples were resolved in SDS-PAGE and the gel was observed on the gel scanner. After scanning, proteins in gel were transferred to PVDF membrane and Western blotting was performed against myc tag using the 9E10 antibody. The protein samples before and after immuno-precipitation were compared by SDS-PAGE and Coomassie brilliant blue staining (Figure 3).

In vitro Eastern staining of His-tagged Fabp7 and comparison with the InVision™ His-tag In-gel staining kit
His-tagged Fabp7 was expressed in 293A cells by transfecting the pc-RC·myc·Fabp7·His.Total cell lysates (10 μg) from transfected and non-transfected cells were incubated with 10 μM of BDED in tubes for 10 minutes at room temperature. After incubation, equal volume of 2 × LDS gel loading buffer was added and the mixture was boiled. Denatured protein samples were resolved in SDS-PAGE and the gel was scanned (Figure 4A).

In the meantime, plasmids pc-RC·myc·Fabp7 and pc-
RC·myc·Fabp7·His were transfected into 293 A cells, and 10 μL of total cell lysates from each sample were resolved in SDS-
PAGE. Equal amount of lysate from non-transfected cells and purified His-Fabp7 prepared by bacterial expression was loaded together for negative and positive control, respectively. After SDS-PAGE, the gel was stained with the InVision™ His-tag In-gel staining kit (Invitrogen) according to the manufacturer’s protocol and scanned by a 532 nm laser and 560 nm long pass filter on the Typhoon 9410 Gel Scanner. Immediately after scanning, the gel was stained with Coomassie brilliant blue (Figure 4B).

**Protocol for Eastern staining**

**A. Transfection**

i) HEK293A cells were maintained in DMEM (Invitrogen, cat. no. 11960) with 10% fetal bovine serum (Invitrogen, cat. no. 10099-141), 1% penicillin/streptomycin (Invitrogen, cat. no. 15140-122).

ii) Cells in 10 cm plates were washed with PBS three times after aspirating the culture medium.

iii) Cells were incubating with 1 mL of Trypsin/1 mM EDTA solution (Invitrogen, cat. no. 25200-056) for 3 minutes and adding 2 mL of culture medium for neutralizing trypsin.

iv) Medium with cells were transferred to 15 mL conical tube. Supernatant was removing after centrifugation at 1,500 rpm during 3 minutes and 10 mL of fresh culture medium were re-suspending with pellets.

v) Cells were plated at a density of 2 × 10^5 cells/well in 12 well plates.

vi) Next day, 500 ng of DNA was mixing with Lipofectamine 2000 (Invitrogen, cat. no. 11668-019) for 30 minutes in 1 mL of Opti-MEM I reduced-serum medium (Invitrogen, cat. no. 31985-062).

vii) HEK293A were incubating with Opti-MEM I reduced-serum medium containing DNA and lipofectamine 2000 for 2 days.

**B. Staining**

i) BDED was dissolved in DMSO (Sigma, cat. no. D2650) and kept at -20°C as 1 mM stock. Culture medium in the wells was exchanged to fresh medium including 500 nM of BDED.

ii) After incubation with 500 nM BDED at 37°C for 20 minutes, cells were washing with PBS three times.

**C. Protein extraction**

i) Cells were incubating with CelLyticMTM cell lysis solution (Sigma, cat. no. C29780) including protease inhibitor cocktail (Roche, cat. no. 11 836 170 001) at room temperature for 2 minutes.

ii) Cell lysates were collecting into 1.5 mL tube and supernatants were transferred to new 1.5 mL tube after centrifugation at 14,000 rpm for 10 minutes.

iii) Concentration of total proteins was decided by Brad ford assay method. 10 µg of the total proteins were mixed with Laemmli Sample Buffer (Bio-rad, cat. no. 161-0737) and boiled at 100°C for 5 minute.

iv) Boiled samples were loaded in NuPAGE Novex Bis-Tris Gels (Invitrogen) and run under 200 V for 30 minutes for fractionating total proteins.

**D. Visualization**

i) Gel was scanned in Typhoon 9410 Gel Scanner (GE Healthcare) using a 488 nm laser and 526SP emission filter for visualizing of the RC-tag: BDE signals.

**E. Confirmation of protein expression**

After scanning, proteins were transferred onto PVDF membrane by wet transfer device (Bio-rad, cat. no. 165-8029) for checking expressed proteins. PVDF membrane was incubating with 1X TTBS solution including 5% skim milk. Finally, protein was visualized after incubating with mouse monoclonal anti c-myc (clone 9E10., Santa Cruz. Sc-40) antibody and secondary goat anti-mouse IgG conjugated with Cy5 (Invitrogen, A10524) which was detected by 633 nm laser and 670BP

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**REFERENCES**