Title:
Detecting Intracellular Thiol Redox State in Leukaemia and Heterogeneous Immune Cell Populations: an optimised protocol for digital flow cytometers

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Quantifying intracellular thiols by flow cytometry

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Abstract
Flow cytometric methods for detecting and quantifying reduced intracellular thiol content using fluorescein-5-maleimide in viable eukaryotic cells date back to 1983 [1]. There has been little development in these methodologies since that time, a period that has witnessed huge technological advances, particularly with the emergence of digital multi-parameter flow cytometric systems. Concurrent advancement in our understanding of redox regulation within eukaryotic cellular systems has also followed, whereby it is now accepted that cysteine thiols partake in redox reactions, which regulate protein activity and function [2,3]. Moreover, we are at the dawn of a new era in redox biology whereby the importance of ‘reductive stress’ in eukaryotic cellular systems is gathering momentum [4]. It is therefore critical that methods be continually advanced to better understand these concepts in more detail at the cellular level. Flow cytometry is a powerful technique that may be used for this purpose. Henceforth we have rejuvenated these methods to address modern scientific questions. In this paper, essential detail is provided on:

- The adaption of a protocol initially described by Durand & Olive [1] for use with modern digital flow cytometer configurations.
- Demonstration that flow cytometry can detect the gain and loss of reduced intracellular thiols in cells exposed to physiological doses of hydrogen peroxide mediated by glucose oxidase [5].
- Validation of F5M protein labelling by coupling method to confocal microscopy and downstream proteomics, thus permitting a powerful experimental platform for potential use with next generation flow cytometry e.g. CyTOF [6].

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Methods Details

Materials

1. Complete IMDM tissue culture medium: containing 10% (v/v) foetal calf serum (FCS), 2 mM L-glutamine, 100 U/ml penicillin and 100 mg/ml streptomycin (Sigma-Aldrich, Dorset, UK).
2. Hank’s balanced salt solution (Sigma-Aldrich, Dorset, UK).
3. Dulbecco’s phosphate buffered saline (D-PBS) endotoxin tested, pH 7.0 (Sigma-Aldrich, Dorset, UK).
4. Flow cytometry stain buffer (FSB): D-PBS adjusted to contain 2% (v/v) FCS, 0.1% (w/v) sodium azide and 5 mM EDTA (stored at 4 °C).
5. Ficoll-Paque PLUS (GE, Buckinghamshire, UK) or Hisopaque 1077 (Sigma-Aldrich, Dorset, UK).
6. Trypan blue solution 0.4% (Sigma-Aldrich, Dorset, UK).
7. Glucose oxidase from *Aspergillus niger* (Sigma-Aldrich, Dorset, UK): 100,000 U/g.
8. N-ethylmaleimide (NEM): 1 mM stock in D-PBS stored at 4 °C protected from light (Sigma-Aldrich, Dorset, UK).
9. Fluorescein-5 maleimide (F5M): 1 mM stock in D-PBS stored at 4 °C protected from light (Sigma-Aldrich, Dorset, UK).

   *Note*: in our laboratory we use an Accuri-C6 (BD-Accuri, Berkshire, UK); however, the method is compatible with other digital systems e.g. Guava® EasyCyte (MerkMillipore, Hertfordshire, UK)
12. NuPAGE 4-12% SDS-PAGE gel, MOPS running buffer and SDS-PAGE LDS sample loading buffer (ThermoFisher, Hampshire, UK)
Cells

1. Jurkat (ATCC®TIB-152™), human acute T-cell lymphoblastic leukaemia.

2. Donor human peripheral blood mononuclear cells (PBMCs).

Note: full ethical approval and donor consent is required for the isolation of PBMCs from human blood, in strict accordance with the declaration of Helsinki [7].
Experimental procedures

Analyzing the effect of glucose oxidase mediated H₂O₂ on intracellular thiol oxidation using flow cytometry

The protocol described is based on a previous method described in Durand & Olive [1], which incorporated various fluorescent stains for the detection of reduced intracellular thiols, including fluorescein-5 maleimide (F5M). The current method is optimised for bench-top digital flow cytometers and allows low level detection of F5M in viable cells, which proved challenging using the previous method. The optimisation of the technique described herein is of particular importance, since some reactive oxygen species (ROS) e.g. H₂O₂ have the propensity to oxidise proteins with exposed cysteine residues, in turn altering their activity [8]. In the experimental model proposed, exposure of cells to physiological doses of exogenous H₂O₂ can be modelled in vitro by adding glucose oxidase (GOX) to the culture medium [5]. The example included uses the Jurkat immortalised T-cell line to monitor cellular cysteine thiol redox state over time in response to H₂O₂. The method details are as follows:

1. Maintain Jurkat cells in 15 ml complete IMDM, at a density between 2 x 10⁵ and 10⁶ cells/ml at 37 °C and 5 % CO₂.
   Note: IMDM is particularly suited to this because of the high glucose concentration (4.5 g/l), which is a substrate for GOX.
2. Prepare a 10 mU/ml GOX solution and a 10 mg/ml catalase solution and store at 4 °C until required.
   Note: The catalase is used to confirm that H₂O₂ is mediating changes in thiol redox state.
To prepare glucose oxidase solutions for in vitro H₂O₂ generation, it is recommended to obtain high purity glucose oxidase solid (e.g. Sigma cat no. 49180 or similar quality). Cheaper preparations can contain confounding contaminants such as catalase, which
metabolises H₂O₂. In a Class 2 biological safety cabinet, dissolve solid GOX in tissue culture-grade water to a concentration of 1 U/ml (U = enzyme units). Sterilise by filtration and aliquot for storage at -20°C and avoid freeze thaw. Prepare 10 mU/ml working solutions by thawing 1 U/ml stock aliquots at 37°C for 1 min and diluting with tissue culture-grade water immediately prior to use. Discard excess. Note that H₂O₂ production generated by GOX in vitro can be measured and titrated to physiological levels using the H₂O₂-specific fluorescent probe Amplex UltraRed (Life Technologies, Hampshire, UK) [5].

3. Transfer the contents of a 24 to 48 hour Jurkat cell culture flask (60-80% confluence) to a 15 ml tube and centrifuge at 100-300 x g for 10 minutes at room temperature in a bench-top centrifuge with swing-out rotor.

4. Remove the supernatant and carefully suspend the pellet in 1 ml complete IMDM pre-warmed to 37 °C.

5. Perform a viable cell count using trypan blue and dilute the Jurkat cells to a viable density of 10⁷ cells/ml.

6. Add 100 µl of the diluted Jurkat cell suspension to five wells in a 96-well tissue culture treated plate and label ‘0h’, ‘1h’, ‘2h’, ‘3h’, ‘4h’.

7. Add 98 µl to two wells in the same plate and label; ‘0h +C’ and ‘4h +C’. Note the “+C” refers to the addition of catalase. Thus there are 7 assay wells in total.

8. Add 2 µl of the 10 mg/ml catalase stock solution (prepared in step 2) to wells labelled ‘0h +C’ and ‘4h +C’ only.

9. To start the assay add 100 µl of the GOX solution (prepared in step 2) to wells labelled ‘4h’ and ‘4h +C’ and incubate at 37 °C and 5 % CO₂.
Note: that this assay has been optimised to use IMDM because of the high glucose content (4.5g/l), rather than RPMI-1640 which has much lower glucose content (2g/l). This is important since GOX requires a glucose substrate to generate H₂O₂.

10. After 1 hour, add 100 µl of the GOX solution to the ‘3h’ well and return to the incubator.

11. Repeat after a further 1 and 2 hours for wells labelled ‘2h’ and ‘1h’ respectively.

12. Finally, 1 hour after the addition of GOX to ‘1h’, add 100 µl of the GOX solution to wells labelled ‘0h’ and ‘0h +C’ and immediately transfer the contents of all 7 assay wells to a fresh 1.5 ml microcentrifuge tube. Centrifuge all at 100-300 x g for 5 minutes at room temperature.

13. Carefully remove supernatants and suspend the pellets in 1 ml D-PBS. Centrifuge all tubes at 100-300 x g for 5 minutes at room temperature.

14. Remove supernatants and suspend pellets in 1 ml pre-chilled 0.1 µM F5M solution and incubate all tubes for 20 min, on ice and protected from light.

15. Centrifuge tubes at 100-300 x g for 5 minutes at 4 °C.

16. Remove supernatants and suspend pellets in 1 ml pre-chilled FSB and centrifuge at 100-300 x g for 5 minutes at 4 °C.

17. Remove supernatants and suspend pellets in 100 µl FSB and go to flow cytometer. Keep tubes on ice and protect from light.

18. At the flow cytometer, acquire the ‘0h’ tube first, followed by the others in chronological order. Draw a region around the Jurkat cell population as seen in Fig. 1A and create an FL1 (λ488nm; λ530/30nm) histogram gated on the region as seen in Fig. 1B.

19. Normalise the FL1 mean fluorescent intensity (MFI) values to ‘0h’ in order interpret the flow cytometry data as shown in Fig. 1C and in Table 1.

20. Viable cell counts should be performed on GOX treated cells before or after flow cytometry acquisition as shown in Fig. 2A.
To demonstrate that the GOX generated H$_2$O$_2$ is mediating the loss in reduced thiols, the effect of catalase in ‘4h’ vs ‘4h +C’ treated cells can be observed on normalised data as shown in Fig. 2B. The data suggest that H$_2$O$_2$ mediates the loss in reduced intracellular thiols as measured by a loss in F5M (FL1) fluorescent signal.

Confocal microscopy confirmation that the F5M signal is intracellular using primary T cells.

It is unclear from the flow cytometry data whether the increase in MFI following F5M treated Jurkat immortalised T-cells is ‘intracellular’ or due to F5M binding to the plasma membrane. To verify this, T-cells from healthy donor peripheral blood mononuclear cells (PBMC) were analysed by flow cytometry and then confocal microscopy. Primary T-cells were surface stained for CD3 at this stage to demonstrate the versatility of the method within heterogeneous cell populations. Venous blood from healthy donors following informed consent was collected by a trained phlebotomist into a 6 ml lavender stopper K2EDTA Vacationer® tube (BD). All procedures from this point should be performed inside a Class 2 biological safety cabinet:

1. Transfer 6 ml venous blood to a fresh sterile universal tube and add an equal volume of Hank’s balanced salt solution (HBS); 12 ml total volume.
2. Add 3 ml Ficoll-Paque PLUS to three sterile 15 ml centrifuge tubes and carefully layer 4 ml of the diluted venous blood on top of the Ficoll-Paque PLUS.
3. Transfer the tubes to a bench-top centrifuge with swing-out rotor and centrifuge at 400 x g for 40 minutes at room temperature (18-20 °C) with the brake off.
4. Carefully remove the tubes from the centrifuge and return to the biological safety cabinet. The PBMC layer (white and delicate) will be visible at the Ficoll-Paque PLUS interface. Carefully remove the upper layer (plasma) to waste with a fresh sterile Pasteur pipette without disturbing the PBMC layer. Then using a fresh sterile Pasteur pipette, transfer the
PBMC layer to a fresh 15 ml centrifuge tube. For each tube, remove the entire PBMC interface with as little of the Ficoll-Paque PLUS layer as possible.

5. Add 8 ml HBS to each tube and centrifuge at 100-300 x g for 10 minutes at room temperature.

6. Using a fresh sterile pipette, remove the supernatant from each tube taking great care not to disturb the PBMC pellet and suspend in 1 ml HBS. The PBMC’s can be pooled at this stage. Add sufficient HBS to the pooled cells to make 10 ml.

7. Centrifuge pooled PBMC’s at 100-300 x g for 10 minutes at room temperature.

8. Remove the supernatant taking great care not to disturb the PBMC pellet and suspend the pellet in 2 ml pre-warmed (37 °C) complete IMDM. Transfer the cells to an incubator set at 37 °C and 5% CO2 for subsequent analysis.

9. Prepare the F5M working solution for flow cytometric use in advance. Protect from light and store at 4 °C (up to 1 month) or at -20 °C for up to 1 year.

10. Remove the PBMC’s from the CO2 incubator and centrifuge at 100-300 x g in a benchtop refrigerated centrifuge with swing-out rotor for 5 minutes at room temperature.

11. Remove supernatant and suspend the pellet in 10 ml D-PBS (equilibrated to room temperature) and centrifuge at 100-300 x g for 10 minutes.

12. Remove supernatant and suspend the pellet in 1 ml FSB. Prepare a 1 in 20 dilution of the PBMC suspension for cell viability counting (5 μl PBMC in 95 μl FSB is sufficient) and transfer the remaining suspension to ice.

13. Mix 10 μl of the 1 in 20 PBMC dilution with 10 μl trypan blue solution and count viable cells using a haemocytometer.

14. Adjust the PBMC suspension to a density of $10^7$ viable cells/ml with pre-chilled (4 °C) FSB and add 100 μl (10^6 cells) to three fresh 1.5 ml microcentrifuge tubes and centrifuge for 5 minutes at 100-300 x g and 4 °C.
Note: At this point it is preferable to pellet cells using a microcentrifuge.

15. Remove supernatants and suspend each pellet in FSB to a final volume of 90 μl.

16. Add 10 μl of mouse anti-human CD3 (clone SK7) APC conjugate to two tubes and 10 μl of mouse IgG1 APC conjugate isotype control to the other. This step will surface stain the T-cells allowing intracellular vs extracellular differentiation during confocal microscopy.

17. Vortex all tubes briefly (1 to 2 seconds) and incubate on ice for 30 minutes protected from the light.

18. Add 1 ml D-PBS (chilled to 4 °C) to each tube and centrifuge for 5 minutes at 100-300 x g and 4 °C. Remove supernatants and then repeat this step.

19. Remove supernatants and add 1 ml D-PBS to the isotype control tube and 1 ml D-PBS to one of the anti-human CD3 tubes and label it ‘F5M’.

20. Add 1 ml of 1 μM NEM solution to the remaining anti-human CD3 tube. These cells will be used to set the background for the F5M T-cell stain. Label this tube ‘NEM’.

   Note: This concentration of NEM is based on 99+% purity. A lower grade can be used, which may require titrating to achieve the optimal concentration.

21. Incubate all tubes for 20 minutes on ice, protected from the light.

22. Centrifuge for 5 minutes at 100-300 x g and 4 °C.

23. Remove supernatants and suspend each pellet in 1 ml D-PBS and centrifuge for 5 minutes at 100-300 x g and 4 °C.

24. Remove supernatants and add 1 ml chilled D-PBS to the isotype control tube. These cells will be used to set the negative T-cell gate. Add 1 ml 0.1 μM F5M to both ‘NEM’ and ‘F5M’ tubes.

25. Incubate all three tubes for 20 minutes on ice, protected from the light.

26. Centrifuge for 5 minutes at 100-300 x g and 4 °C.
27. Remove supernatants and suspend each pellet in 1 ml pre-chilled FSB and centrifuge for 5 minutes at 100-300 x g and 4 °C. Repeat this step.

28. Finally remove supernatants and suspend each pellet in 100 μl pre-chilled FSB, keep on ice, protect from light and go to flow cytometer.

29. At the flow cytometer, acquire the ‘Ig-control’ cells first. These cells can be used to set the negative T-cell gate, which will appear in the FL4 parameter (laser: λ640 nm, filter: λ675/25 nm). Fig. 3A and 3B illustrates the gating strategy.

30. Next acquire the ‘NEM’ cells and create an FL1 (laser: λ488 nm, filter: λ530/30 nm) histogram plot gated on the T-cell population as seen in Fig. 3B. The peak shown on the FL1 histogram is representative of the F5M background staining and can be seen in Fig. 3C.

31. Finally acquire the ‘F5M’ cells. As shown in Fig. 3C, these cells fluoresce at an order of magnitude greater than ‘NEM’ cells.

32. To confirm viability, perform cell counts either before or after flow cytometry acquisition as described in steps 3 and 4. Fig. 3D shows little loss in cell viability post F5M treatment.

33. As shown in Fig. 3E and 3F, confocal microscopy is used to confirm that the F5M is entering the T-cells and therefore that the flow cytometric FL1 signal generated is representative of intracellular thiol redox status. To visualise F5M labelled CD3+ T-cells by confocal microscopy, carefully affix a glass microscope slide to the microscope and pipette 50-100 μl of the final stained cell suspension as a droplet onto the upper surface. To limit evaporation gently place a thin coverslip (19 mm) on top of the droplet. Focus cells using a 63x oil immersion objective. Use both the argon (F5M) and helium-neon (CD3-APC) lasers at a previously determined appropriate power setting, and adjust the offset, gain, zoom and Z-position as necessary for optimal image quality. If available, opt
for sequential scanning modes to allow simultaneous visualisation of both signals with minimal crosstalk between channels. DNA stains such as DAPI can be added to the staining protocol to identify nuclear regions where the appropriate excitation lines and channels are available.

**Analysing F5M labelled proteins extracted from viable cells**

The data in the previous section demonstrated that treating viable cells with F5M results in the labelling of solvent accessible intracellular *reduced* cysteine thiols [9]. This will include low molecular weight thiols such as glutathione [10], and solvent accessible cysteine thiols present in the native conformation of redox active proteins [11]. It would be expected that if a *reduced* cysteine thiol is modified through oxidation, F5M labelling will not occur, since F5M preferentially reacts with *reduced* cysteine thiols via the ‘Michael addition’ [12]. Thus identifying individual proteins using proteomic platforms e.g. mass spectrometry will add greater understanding of how individual cysteine thiols can modulate biological processes. To demonstrate this principle, proteins from Jurkat cells exposed to GOX-mediated H₂O₂ were separated by gel electrophoresis. Untreated vs. F5M SDS-PAGE separated proteins were then dissected for analysis by mass spectrometry. This protocol has potential to be adapted for use with next generation flow cytometers e.g. CyTOF [6, 13]. The details of the experiment used are below:

1. Set up two treatments in 15 ml centrifuge tubes; (i) 2 x 10⁶ Jurkat cells with 1 μM NEM for 20 minutes followed by 20 μM F5M for 20 minutes, (ii) 2 x 10⁶ Jurkat cells with F5M only for 20 minutes. Carry out all F5M incubations as described in the above sections. **Fig. 4A** illustrates the level of F5M fluorescence for these treatments compared to untreated controls as analysed by flow cytometry.
2. Centrifuge the NEM and F5M treated Jurkat cells at 100-300 x g for 5 minutes at 4 °C.

3. Remove supernatants and add 10 ml pre-chilled FSB and centrifuge at 100-300 x g for 10 minutes at 4 °C.

4. Remove supernatants and suspend both pellets in 400 μl pre-chilled (4 °C) PBS and transfer to fresh microcentrifuge tubes for cell lysis by sonication.
   Note: Keep cell lysates on ice and protected from light at all times. This will minimise protein degradation and protect F5M from photo-beaching.

5. Centrifuge tubes at 12 000 x g for 15 minutes at 4 °C and harvest supernatants into fresh microcentrifuge tubes on ice and protect from light. Perform protein quantitation on both cell extracts (Bradford assay [14] or equivalent is appropriate).

6. Add 20 μg of each protein extract to non-reducing NuPAGE® LDS SDS-PAGE sample loading buffer made up to a final volume of 20 μl and heat at 70 °C for 10 minutes, protected from light. Do not add reducing agent e.g. 2-mercaptoethanol. Allow samples to cool before loading the Novex® 4-20% gradient pre-cast tris-glycine gel.

7. Assemble the SDS-PAGE gel tank with the gel and add 1 x NuPAGE® MOPS SDS-PAGE running buffer to upper and lower buffer chambers.

8. Load samples along with molecular weight marker and electrophorese at 180 V for 60-80 minutes, or until the dye front has run off. It is important to protect the gel and SDS-PAGE gel tank from light during this procedure.

9. Remove gel to a small tray containing dH2O and scan immediately using a Typhoon-9400 variable mode imager (or equivalent) set at λ490-495 nm for excitation and λ515-520 nm for emission.

10. Remove gel from Typhoon directly into SDS-PAGE gel fix solution and incubate for 15 minutes at room temperature with gentle agitation.
11. Rinse gel in dH$_2$O and add G-250 colloidal coomassie blue reagent. Incubate overnight (10-20 hours) at room temperature with gentle agitation.

12. Remove G-250 colloidal coomassie blue reagent and de-stain with dH$_2$O for 2 hours at room temperature with gentle agitation. Change the dH$_2$O every 20 minutes.

13. Scan gel using a standard flatbed SDS-PAGE gel scanner and analyse the data along with the respective Typhoon fluorescence image data using ImageQuant (v8.1) or equivalent high resolution software. **Fig. 4B** shows Typhoon and respective coomassie gel images of proteins extracted from the NEM + F5M treated (i) and F5M alone treated (ii) Jurkat cells. Fig 3C illustrates pixel peak profiles for NEM + F5M treated (i) and F5M alone treated (ii) extracted proteins. **Fig 4C** illustrates pixel density analysis of NEM + F5M and F5M alone treated Jurkat cells.

14. Since F5M irreversible modifies cysteine thiols, identification of redox active proteins is permitted via mass spectrometry based on incremental mass shifts relating to F5M binding. **Table 2** shows various intracellular redox active proteins identified from F5M treated Jurkat cells using LC/MS analysis. For the F5M mass shift, the formula C$_{24}$H$_{13}$NO$_7$ (+427.069 Da) was used, since this relates to the mass difference expected for a Cys-thiol or Cys-thiolate modification by F5M.

Note: LC/MS analysis was outsourced to the Bristol Proteomics Facility (University of Bristol, UK).

**Acknowledgements**

AJW and SJC designed and performed all experiments and analysed all data. RGM assisted with the confocal microscopy. KJH assisted with mass spectrometry. PSH assisted with the GOX assays and H$_2$O$_2$ quantitation. All authors contributed to the drafting of this manuscript.
References


Figure Legends

Figure 1: The effect of exogenous GOX-generated \( \text{H}_2\text{O}_2 \) on the loss of intracellular reduced thiol using an \textit{in vitro} immortalised Jurkat T-cell model. (A) Representative FSC vs SSC flow cytometric profile of GOX treated Jurkat cells (P2). (B) Histogram (gated on P2) illustrates loss of F5M signal in Jurkat cells following incubation with 5 mU/ml GOX for 0-4 hours (h). Samples were taken for flow cytometric analysis after the following times; 0h, 1h, 2h, 3h and 4h. Data were compared with 0h to normalise the loss of F5M signal (see Table 1). (C) Chart illustrates normalised FL1 (laser: \( \lambda 488 \) nm, filter: \( \lambda 530/30 \) nm) MFI. An increase in F5M signal is noted at 1h (suggesting a reductive spike), followed by progressive decrease in signal at 2h to 4h (suggesting a loss of reduced thiols). Data shown are mean +/- SD of the mean (n=3).

Figure 2: The evaluation of exogenous GOX-derived \( \text{H}_2\text{O}_2 \) on Jurkat T-cell viability using trypan blue staining. (A) Plot illustrates the mean Jurkat viability (%) following incubation with 5 mU/ml GOX at 0h to 4h. To confirm that GOX-derived \( \text{H}_2\text{O}_2 \) was mediating the loss in reduced intracellular thiol, similar experiments were performed in the presence of 100 \( \mu \text{g/ml} \) catalase. (B) Plot illustrates the normalised mean loss of reduced intracellular thiol at 0h compared with 4h (GOX treated) and 4h +C (GOX with catalase treated). Data illustrate that in the presence of catalase, intracellular thiol remains reduced. Data shown are mean +/- SD of the mean (n=3).

Figure 3: Flow cytometric analysis of intracellular thiol redox status in primary human T-cells. PBMC’s were prepared from venous blood using Ficoll-Paque density centrifugation and washed in D-PBS. Washed PBMC’s (10^6 cells in 100 \( \mu \text{l} \)) were labelled with anti-human
CD3-APC. PBMC’s were then treated with either 1 μM NEM followed by 0.1 μM F5M, or 0.1 μM F5M alone and analysed by flow cytometry. (A) Representative forward light scatter (FSC) vs side light scatter (SSC) flow cytometric profile with the treated PBMC’s identified in P2. (B) Gated on P2, CD3+ T-cells are shown in the FL4 parameter (laser: λ640 nm, filter: λ675/25 nm) in R1 (C). Panel shows representative fluorescence histograms depicting F5M fluorescence in control and treated T-cells (gated on R1). (D) T-cell viability was monitored before and after NEM and F5M treatment using trypan blue staining. Confocal microscopy was used to confirm (E) F5M cellular uptake in PBMCs and (F) surface staining for T-cells with anti-human CD3-APC.

Figure 4: Analysis of thiol containing redox active intracellular protein targets following NEM and F5M treatment of viable Jurkat T-cells. (A) Panel shows a representative overlaid FL1 (laser: λ488 nm, filter: λ530/30 nm) histogram of Jurkat cells treated with 1 μM NEM + 0.1 μM F5M (i) and 0.1 μM F5M alone (ii). Respective protein extracts were prepared and separated by SDS-PAGE (protected from the light). Fluorescent protein bands were imaged using a Typhoon-9400 variable mode imager set at λ490-495 nm for excitation and λ515-520 nm for emission. The SDS-PAGE gel was counterstained with G-250 colloidal coomassie blue to confirm equal gel well loading. (B) Representative gel images for proteins extracted from flow cytometry FL1 peaks (i) and (ii). Typhoon images are presented along with respective Coomassie counterstained images. (C) Representative pixel peak profiles comparing fluorescent protein band intensity between (i) and (ii), lower and upper profiles respectively. Five corresponding gel bands (pixel peaks) are highlighted that differ in fluorescence intensity (no difference was observed for the respective coomassie counterstain).