In Vitro Screening Method for Characterization of Macrophage Activation Responses

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Abstract: Macrophage activation refers to the enhanced functionality of macrophages in response to endogenous or exogenous stimuli. Due to the existence of limitless stimuli and a multitude of receptors on macrophage surfaces, the nature of activation (or acquired functioning) can be specific to the encountering stimulus. This article describes a macrophage-activation screening platform in a 96-well format. The methodology involves the generation of bone marrow-derived macrophages, their activation into two extreme activation states, and screening of activated macrophages for expression of bonafide protein biomarkers. A high-throughput and stringent assay to determine macrophage activation markers developed in this article can be adapted for biomarker determination in pathological conditions and toxicant/drug safety screening.

Keywords: macrophage; BMDM; macrophage activation; M1; M2; alternatively-activated macrophages (AAM); classically-activated macrophages (CAM)

1. Introduction

Local alteration in homeostatic tissue environment is sensed by resident macrophages and other innate immune cells [1–5]. Based on the nature and severity of perturbations, tissue macrophages boost their functional armor via increased expression of genes involved in pathogen killing, wound healing, cytokine secretion, and phagocytic clearance [6]. Despite clear-cut categorization of activated macrophage populations into M1 (classically activated) and M2 (alternatively activated), the activation markers identified in macrophages obtained from different tissue compartments and disease models reflect a great degree of heterogeneity in macrophage activation responses [2,7].

M1 macrophage activation is induced by Interferon (IFN)-γ and lipopolysaccharide (LPS) [8]. These macrophages express pro-inflammatory cytokines including Interleukin (IL)-6, IL-1β, and TNFα, and inducible nitric oxide synthase (iNOS) [9,10]. M2 macrophages are divided into three subcategories, i.e., M2a, M2b, or M2c, based on the expression of surface markers, production of specific cytokines, and their acquired functions [11]. M2a activation is induced by IL-4 and/or IL-13 and they express markers including Arg1 and Fizz1 [8,12]. M2b or regulatory macrophages are induced by LPS and immune complexes and secrete IL-10. M2c macrophages are induced by IL-10 and secrete profibrotic, i.e., TGFβ, and anti-inflammatory, i.e., IL-10, mediators [13,14].

Identification of macrophage activation patterns can be performed in purified macrophages from healthy and diseased tissues using gene expression profiling [15,16], western blotting, immuno-cytochemical staining, and flow cytometry [17,18]. Although employed routinely, these approaches generally pose challenges, including the selection of effective reagents (antibodies), tissue/cell processing, and lack of positive controls. In our attempt to develop a robust, reliable, and consistent assay, after screening various antibody specificities using protein electrophoresis/western assays, we employed an in-cell western technology to develop a macrophage activation screening platform. In-cell western offers many advantages...
compared to traditional assays such as immuno-cytochemistry and western blotting. It provides higher sensitivity and more meaningful protein expression analyses in whole cells. This approach allows for enhanced quantitative analyses in a high throughput manner in plated whole cells. Importantly, this approach allows reduction in the number of animals used in research. Although developed using BMDM cell line, the assay can be adapted to various primary macrophage (Bronchoalveolar/peritoneal macrophages) and immortalized macrophage cell lines.

2. Materials

Table 1 summarizes the reagents and equipment utilized in this study.

| Reagent                        | Company/Institute          | Cat. Number | Comments                              |
|--------------------------------|-----------------------------|-------------|---------------------------------------|
| C57BL/6j                        | Jackson Laboratory         | 000664      | Mouse Strain                          |
| L-929 Cell line                 | ATCC                        | CCL-1       | Mouse connective tissue fibroblast cell line |
| Dulbecco’s Modified Eagle Medium (DMEM) | Gibco (Life Technologies) | 11995-065   | With Glucose, L-Glutamine and Sodium Pyruvate |
| RPMI 1640                       | Gibco (Life Technologies)  | 22400-089   | With L-Glutamine and HEPES            |
| Fetal Bovine Serum              | Atlanta Biologicals        | S11550H     | Heat Inactivated                      |
| Penicillin/Streptomycin (100X)  | Sigma-Aldrich              | P4333       | Working concentration (1X)            |
| Phosphate Buffered Saline (PBS) | Sigma-Aldrich              | P5368       | 0.01 M PBS; pH 7.5                    |
| Interleukin 4                   | EMD Millipore              | IL016       | Recombinant (Murine)                  |
| Interferon Gamma (IFN-γ)        | EMD Millipore              | IF005       | Recombinant (Murine)                  |
| Lipopolysaccharide (LPS)        | EMD Millipore              | LPS25       | LPS, *E. Coli* O111:B4                |
| 96-well plates                  | Thermo Scientific          | 165305      | Black with polymer base               |
| Filtration Unit                 | Genesee                    | 25-227      | 0.22 µm                               |
| Betadine                        | Purdue Products            | 67618-151-17| Povidone-iodine, 7.5%                 |
| Cell strainer                   | Corning Life Sciences      | 352350      | 70 µm                                 |
| T25 culture flask               | Genesee                    | 25-207      |                                       |
| T75 culture flasks              | Genesee                    | 25-209      |                                       |
| 100 mm Petri-dishes             | Thermo Scientific          | 130182      |                                       |
| 10% Buffered Formalin           | Fisher Scientific          | SF100-20    | pH 7.0                                |
| Triton x-100                    | Fisher Bioreagents         | BP151-100   | Electrophoresis Grade                 |
| Tween-20                        | Fisher Bioreagents         | BP337-100   | Electrophoresis Grade                 |
| FIZZ1A                          | ABCAM                      | AB39626     | Rabbit Polyclonal                     |
| YM1/2                           | National Institute of Health | Rabbit Polyclonal | A kind gift from Dr. Shioko Kimura |
| INOS                           | ABCAM                      | AB15326     | Rabbit Polyclonal                     |
| HIF2a                           | ABCAM                      | AB199       | Rabbit Polyclonal                     |
| COX 1                           | Cell Signaling             | 4841        | Rabbit Polyclonal                     |
3. Procedure

3.1. Generation of Macrophage-Colony Stimulating Factor (M-CSF) Containing L929 Medium

An overview of experimental procedure describing generation of BMDMs is shown in Figure 1.

1. Plate 10^6 L929 cells (M-CSF producing cell line, CCL-1, was purchased from ATCC) in 20 mL of DMEM (Gibco, #11995-065) supplemented with 10% heat inactivated fetal bovine serum (FBS), and 1% penicillin/streptomycin (Lonza, #17-602E) in a T75 cell culture flask.

2. Grow cells for one week (or until the flask is ~90% confluent) at 37°C, 5% CO_2. It is important to keep the seeding density and harvesting schedule consistent to avoid variation in L929 concentration is the L929 medium.

3. Collect the medium and spin at 400×g for 5 min to remove any floating cells and cellular debris.

4. Filter the supernatant through a 0.22 µm filter. Store the filtered L-929 medium in 150 mL aliquots at -20 °C.

5. L929-conditioned macrophage media is then prepared by mixing L-929 medium (30%) and RPMI 1640 (Caisson Labs, #RPL09-500 mL) complete medium (70%).

![Figure 1. BMDM Generation.](image)

3.2. Collection of Bone Marrow Cells

1. Euthanize six-week-old male mice on C57BL/6J background by CO_2 inhalation and cervical dislocation. Disinfect euthanized mice with 70% ethanol and betadine (this step can be performed on bench-top).

2. Using sterile surgical instruments, isolate femur and tibial bones, and transfer to 100 mm cell culture dish containing 5 mL of RPMI 1640 complete media (10% FBS, and 1% pen/strep).

3. Transfer cell culture dish containing bones to cell culture hood for further processing. Using new set of surgical instruments, cut bones from both ends to expose the marrow cavity. Flush the bone marrow into new cell culture dish with 10 mL syringe (containing RPMI 1640 complete media) fitted with 22G needle.

4. Dissociate bone marrow with repeated passing through 25G needle. Strain the cell suspension through 70 µm cell strainer in to 50 mL sterilized conical tube.
(5) Spin the cell suspension at 400 \( \times \) g for 5 min. Resuspend the cell pellet in 20 mL of L929-conditioned macrophage medium and plate in T25 cell culture flask for overnight incubation. Next morning (~12 h wait), replate non-adherent cells from T25 flask into 100 mm petri dishes (20 dishes per mouse) for differentiation of bone marrow cells to macrophages.

(6) Add 3 mL fresh L929-conditioned macrophage medium every day. Switch L929-conditioned macrophage medium at 3-day intervals (7 mL per dish). Collect adherent cells (mature macrophages) on 10th day of culture and proceed to in vitro macrophage activation step.

3.3. BMDM Macrophage Activation

A step-by-step protocol for generation of activated macrophages is shown in Figure 2.

(1) Seed BMDMs in designated wells (seeding density: 40,000 per well in 200 \( \mu \)L volume) of 96-well cell culture plate. Incubate the plate at 37 °C for 1 h.

(2) To prevent the likely influence of serum constituents on the macrophage responsiveness at baseline, replace the media with serum-free RPMI 1640 complete medium, and incubate the plate for overnight serum starvation. Of note, in case of possible loss of viability of cells due to serum-free environment, RPMI 1640 complete medium with 1–2% serum could be used.

(3) Next day, label wells for M1, M2, and M0 (non-activated). Replace media (24 wells/treatment) with fresh serum-free RPMI 1640 complete medium containing IFN-\( \gamma \) (215 U/mL) + LPS (10 ng/mL; added 8 h after the start of IFN-\( \gamma \) treatment) for M1 activation or IL-4 (20 U/mL) for M2 activation. Wells assigned to M0 group will receive fresh serum-free RPMI1640 complete medium with no additives [8].

3.4. Screening for Macrophage Activation Markers

A detailed overview of macrophage activation screening is shown in Figure 3.

(1) Wash once with 1 \( \times \) PBS buffer and fix with 100 \( \mu \)L of 10% neutral buffered formalin (Fisher Scientific) for 20 min, at room temperature. Use of multichannel pipette is highly recommended. To avoid dislodgment of cells, it is important to add buffers/solutions alongside the walls.

(2) Centrifuge the 96 well plate at 500 \( \times \) g for 5 min. Aspirate the formalin fixative and wash cells once with 1 \( \times \) PBS.

(3) Wash cells three times (5 min for each wash) with 0.1% Triton X-100 PBS (1 \( \times \) PBStr) to permeabilize cells.

(4) Wash cells once with 1 \( \times \) PBS. Add 200 \( \mu \)L of LI-COR Odyssey blocking buffer per well to block cells. Incubate on a rocker for 1 h at room temperature.
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(5) Add 50 µL of primary antibody solution (prepared in Odyssey Blocking Buffer) to each well and incubate overnight on rocker in a cold room.

(6) Wash the plate three times with 0.1% Tween-20 PBS (1 × PBSTw) for 5 min at room temperature.

(7) Incubate with 50 µL of goat anti rabbit IRDYE 800CW secondary antibody solution (1:1000) for 1 h at room temperature. At this step, include DNA labeling dye (1:10,000 DRAQ5 stain) for normalizing cell numbers in each well.

(8) Wash with 1 × PBSTw for 5 min, three times.

(9) Wash with 1 × PBS for 5 min. Discard wash solution and dry the plate on paper towels.

(10) Scan the plate in both the 700 nm and 800 nm detection channels using Odyssey Imager (Odyssey CLx Infrared Imaging System, Lincoln, NE). Normalize the primary antibody signal by dividing value obtained on 800 nm channel with values obtained on 700 nm channel for each well.

**Figure 3.** Macrophage Activation Marker Analyses.

**4. Results**

First, we tested the specificity of antibodies for bonafide M1 and M2 markers by performing western blots (Figure 4). The protein expression of INOS, a bonafide marker to test M1 activation, was enhanced in classically activated macrophages, whereas the protein expression of FIZZ1, a bonafide marker to test M2 activation, was enhanced in alternatively activated macrophages.

**Figure 4.** Western blot analyses on Naïve (M0), M1-activated, and M2-activated BMDMs.

INOS (M1 marker) and FIZZ1 (M2 marker) expression status was interrogated using in cell western approach (Figure 5). Similar to the results obtained with western blots (Figure 4), INOS was enhanced in classically activated macrophages while FIZZ1 was enhanced in alternatively activated macrophages in in cell western assays.
Antibody specificity for other activation markers (ARG1, COX1, MMP12, ALOX12/15, and YM1/2) was confirmed using western blot (data not shown). Finally, activated macrophages were interrogated for the expression levels of selective activation markers (Figure 6).

5. Discussion

In this article, we report a well-standardized and highly adaptive in vitro method to assess macrophage activation patterns based on the expression of activation-specific protein markers. Although the current study is focused on BMDM activation responses to M1 and M2 treatments, the method can be easily modified for any other macrophages, including other primary macrophages and macrophage cell-lines. In addition, the method offers multiple advantages in research. First, the 96-well format allows simultaneous screening of large number of test samples. Additional ly, few wells of 96-well plates can be seeded with known activated and naïve macrophages that can serve as positive and negative controls during analyses of unknown samples. Second, the method can be adapted for high throughput screening of drugs/toxicants for safety assessment. Third, the method can be used for other types of macrophages, including human monocytes derived from peripheral blood mononuclear cells and macrophages derived from other species. However, differentiation of mononuclear cells into macrophages would require species-specific protocols.

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of large number of test samples. Additionally, few wells of 96-well plates can be seeded with known activated and naïve macrophages that can serve as positive and negative controls during analyses of unknown samples. Second, the method can be adapted for high throughput screening of drugs/toxicants for safety assessment. Third, the method can be adapted for phenotypic investigation of macrophage populations from various murine models of diseases.

One caveat of this method is the inability to simultaneous label cells for more than one marker. This limitation is commonly encountered in most of the current techniques except flow cytometry. However, the in vitro method described in this study presents a robust tool as a first line of investigation on macrophage responses in toxicant/drug screening as well as disease investigation. While this protocol was established using murine BMDMs, it can be used for other types of macrophages, including human monocytes derived from peripheral blood mononuclear cells and macrophages derived from other species. However, differentiation of mononuclear cells into macrophages would require species-specific protocols.

Inducible nitric oxide synthase (INOS) is the signature M1-activation marker that is characteristic of classical activation response [19]. As expected, INOS expression was only observed in classically activated macrophages. For M2 activation, we specifically focused on M2a subset that is induced by IL-4. Our method can be adapted to macrophages stimulated with TFGβ and IL-10 (M2c). For M2-specific signatures, we used found in inflammatory zone 1 (FIZZ1) and chitinase 3 like 4 (CHI3L/4) markers [12,20]. FIZZ1, a well-standardized M2 marker, was only seen in M2 macrophages. Although baseline expression of CHI3L3/4 (also known as YM1/2) was observed in M0 as well as M1 macrophages, only M2 macrophages showed significantly upregulated expression. The assessment of the expression levels of these three markers would identify M1 or M2-activated macrophages.

In summary, the assay developed in this article is highly adaptable, high-throughput, and informative and can be utilized in toxicological screening as well as disease phenotyping studies.

**Author Contributions:** S.P. and Y.S. conceived and designed the study; Y.S., S.P. and B.W.L. performed the experiments. Y.S., S.P. and B.W.L. wrote and reviewed the manuscript for intellectual contents. All authors have read and agreed to the published version of the manuscript.

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**Institutional Review Board Statement:** The animal study protocol was approved by the Louisiana State University-Institutional Animal Care and Use Committee (IACUC).

**Informed Consent Statement:** Not applicable.

**Data Availability Statement:** The data used and analyzed during the current study are included in this article and are available from the corresponding author on reasonable request.

**Conflicts of Interest:** The authors declare no conflict of interest.

**Abbreviations**

AAM: Alternatively-activated Macrophages, CAM: Classically-Activated Macrophages, BMDM: Bone Marrow-Derived Macrophages, 1X PBSTw: 0.1% Tween-20 PBS, IL-4: Interleukin-4, IFN-γ, Interferon-γ, 1X PBSTr: 0.1% Triton X-100 PBS.

**References**

1. Hussell, T.; Bell, T.J. Alveolar macrophages: Plasticity in a tissue-specific context. *Nat. Rev. Immunol.* 2014, 14, 81–93. [CrossRef] [PubMed]
2. Mosser, D.M.; Edwards, J.P. Exploring the full spectrum of macrophage activation. *Nat. Rev. Immunol.* 2008, 8, 958–969. [CrossRef] [PubMed]
3. Glass, C.K.; Natoli, G. Molecular control of activation and priming in macrophages. *Nat. Immunol.* 2016, 17, 26–33. [CrossRef] [PubMed]
4. Bain, C.C.; MacDonald, A.S. The impact of the lung environment on macrophage development, activation and function: Diversity in the face of adversity. *Mucosal. Immunol.* 2022, 15, 223–234. [CrossRef] [PubMed]
5. Blieriot, C.; Chakarov, S.; Ginhoux, F. Determinants of Resident Tissue Macrophage Identity and Function. *Immunity* 2020, 52, 957–970. [CrossRef] [PubMed]
6. Murray, P.J.; Wynn, T.A. Protective and pathogenic functions of macrophage subsets. *Nat. Rev. Immunol.* 2011, 11, 723–737. [CrossRef] [PubMed]
7. Gordon, S.; Taylor, P.R. Monocyte and macrophage heterogeneity. *Nat. Rev. Immunol.* 2005, 5, 953–964. [CrossRef] [PubMed]
8. Mosser, D.M.; Zhang, X. Activation of murine macrophages. *Curr. Protoc. Immunol.* 2008, 83, 14.2.1–14.2.8. [CrossRef] [PubMed]
9. Edin, S.; Wikberg, M.L.; Dahlén, A.M.; Rutegard, J.; Öberg, A.; Oldenborg, P.A.; Palmqvist, R. The distribution of macrophages with a M1 or M2 phenotype in relation to prognosis and the molecular characteristics of colorectal cancer. *PLoS ONE* 2012, 7, e47045. [CrossRef]
10. Abdelaziz, M.H.; Abdelwahab, S.F.; Wan, J.; Cai, W.; Huixuan, W.; Jianjun, C.; Kumar, K.D.; Vasudevan, A.; Sadek, A.; Su, Z.; et al. Alternatively activated macrophages; a double-edged sword in allergic asthma. *J. Transl. Med.* 2020, 18, 58. [CrossRef] [PubMed]
11. Yao, Y.; Xu, X.H.; Jin, L. Macrophage Polarization in Physiological and Pathological Pregnancy. *Front. Immunol.* 2019, 10, 792. [CrossRef] [PubMed]
12. Raes, G.; Noel, W.; Beschin, A.; Brys, L.; de Baetselier, P.; Hassanzadeh, G.H. FIZZ1 and Ym as tools to discriminate between differentially activated macrophages. *Dev. Immunol.* 2002, 9, 151–159. [CrossRef] [PubMed]
13. Wang, L.X.; Zhang, S.X.; Wu, H.J.; Rong, X.L.; Guo, J. M2b macrophage polarization and its roles in diseases. *J. Leukoc. Biol.* 2019, 106, 345–358. [CrossRef] [PubMed]
14. Arora, S.; Dev, K.; Agarwal, B.; Das, P.; Syed, M.A. Macrophages: Their role, activation and polarization in pulmonary diseases. *Immunobiology* 2018, 223, 383–396. [CrossRef] [PubMed]
15. Saini, Y.; Dang, H.; Livraghi-Butrico, A.; Kelly, E.J.; Jones, L.C.; O’Neal, W.K.; Boucher, R.C. Gene expression in whole lung and pulmonary macrophages reflects the dynamic pathology associated with airway surface dehydration. *BMC Genom.* 2014, 15, 726. [CrossRef] [PubMed]
16. Choudhary, I.; Vo, T.; Paudel, K.; Patial, S.; Saini, Y. Compartment-specific transcriptomics of ozone-exposed murine lungs reveals sex- and cell type-associated perturbations relevant to mucoinflammatory lung diseases. *Am. J. Physiol. Lung Cell. Mol. Physiol.* 2021, 320, L99–L125. [CrossRef] [PubMed]
17. Kradin, R.L.; McCarthy, K.M.; Preffer, F.I.; Schneeberger, E.E. Flow-cytometric and ultrastructural analysis of alveolar macrophage maturation. *J. Leukoc. Biol.* 1986, 40, 407–417. [CrossRef] [PubMed]
18. Choudhary, I.; Vo, T.; Paudel, K.; Wen, X.; Gupta, R.; Kesimer, M.; Patial, S.; Saini, Y. Vesicular and extravesicular protein analyses from the airspaces of ozone-exposed mice revealed signatures associated with mucoinflammatory lung disease. *Sci. Rep.* 2021, 11, 23203. [CrossRef] [PubMed]
19. MacMicking, J.; Xie, Q.W.; Nathan, C. Nitric oxide and macrophage function. *Annu. Rev. Immunol.* 1997, 15, 323–350. [CrossRef] [PubMed]
20. Raes, G.; De Baetselier, P.; Noel, W.; Beschin, A.; Brombacher, F.; Hassanzadeh Gh, G. Differential expression of FIZZ1 and Ym1 in alternatively versus classically activated macrophages. *J. Leukoc. Biol.* 2002, 71, 597–602. [PubMed]