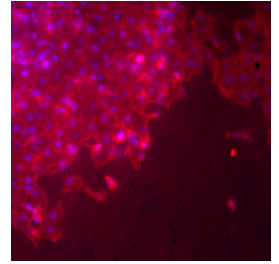


Feb 18, 2019 Version 3

Mammalian Cell Staining V.3

DOI

dx.doi.org/10.17504/protocols.io.x95fr86



Kenneth Schackart¹, Kattika Kaarj¹

¹University of Arizona

481b Laboratory



Kenneth Schackart

University of Arizona

OPEN  ACCESS



DOI: dx.doi.org/10.17504/protocols.io.x95fr86

Protocol Citation: Kenneth Schackart, Kattika Kaarj 2019. Mammalian Cell Staining. **protocols.io**

<https://dx.doi.org/10.17504/protocols.io.x95fr86>

License: This is an open access protocol distributed under the terms of the **Creative Commons Attribution License**, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Protocol status: Working

We use this protocol and it's working

Created: February 18, 2019

Last Modified: February 18, 2019

Protocol Integer ID: 20509

Abstract

This protocol details how to stain mammalian cells cultured on a 96-well plate. Actin filaments, focal adhesion sites (as indicated by the presence of vinculin), and nuclei will be stained.






Materials


- 4% Paraformaldehyde solution
- 0.1% Triton X-100 solution in PBS
- Blocking buffer (PBS + 1% bovine serum albumin)
- Washing buffer (PBS + 0.05% Tween-20)
- Anti-vinculin solution (1:500 in blocking buffer)
- TRITC-conjugated phalloidin and FITC-conjugated antivinculin secondary antibody solution (1:1:248, TRITC:FITC:blocking buffer) referred to as FITC:TRITC
- DAPI solution (1:249 DAPI:blocking buffer)
- Phosphate buffered saline (PBS)






Fix the cells

- 1 Remove cell culture media.
- 2 Add  100 μL of  4 % volume paraformaldehyde solution.
- 3 Incubate for  00:05:00 .


Perforate cell membrane

- 4 Remove paraformaldehyde solution.
- 5 Wash twice with  100 μL washing buffer.


Note

Washing buffer is PBS with the detergent Tween-20.
- 6 Add  100 μL of  0.1 % volume Triton X-100.
- 7 Incubate for  00:05:00 .

Block unspecific binding


- 8 Remove Triton X-100.
- 9 Wash twice with  100 μL washing buffer.



10 Add  100 μL blocking buffer.

Note

Blocking buffer is PBS with BSA (bovine serum albumin) and is used to prevent unspecific binding.

11 Incubate for  00:10:00 .

Bind anti-vinculin to vinculin

12 Remove blocking buffer.


Bind anti-vinculin to vinculin

13 Wash twice with washing buffer.

Bind anti-vinculin to vinculin

14 Add  250 μL of Anti-Vinculin and blocking buffer mixture.

Bind anti-vinculin to vinculin



15 Incubate for  00:20:00 .

Stain actin filaments and focal adhesion sites




16 Remove anti-vinculin blocking buffer mixture.

Stain actin filaments and focal adhesion sites



- 17 Wash twice.
- 18 Add  100 μ L FITC:TRITC solution, cover in foil.
- 19 Incubate for  00:30:00 .

Stain nuclei

- 20 Remove stains.
- 21 Add  100 μ L of DAPI solution, cover in foil.
- 22 Incubate for  00:05:00
- 23 Remove DAPI solution.
- 24 Add  100 μ L PBS.

Image

- 25 Image your cells using UV, Blue, and Green excitation.