



Oct 09, 2022

SDS-PAGE

Ana Belem García González¹, Georgina Diego¹, Irán Alessandra Chaparro Rodríguez¹, Jair Alexis Gardea Sáenz¹

¹Tecnologico de Monterrey Campus Chihuahua

1 Works for me Share

This protocol is published without a DOI.



ABSTRACT

This protocol shows the steps carried out by team Tec-Chihuahua to perform SDS-PAGE

PROTOCOL CITATION

Ana Belem García González, Georgina Diego, Irán Alessandra Chaparro Rodríguez, Jair Alexis Gardea Sáenz 2022. SDS-PAGE. **protocols.io** https://protocols.io/view/sds-page-cezqtf5w

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

CREATED

Aug 08, 2022

LAST MODIFIED

Oct 09, 2022

PROTOCOL INTEGER ID

68368

MATERIALS TEXT

SOLUTIONS:

- Acrylamide/Bis 30%
- 29.2 g of Acrylamide.
- 0.8 g of Bis Acrylamide
- Dissolve in 50 mL of distilled water in constant agitation, gauge to a volume of 100 mL.
- Store at 4 °C degrees while protected from light.
- Tris-HCl/ SDS 4X pH 8.8 (1.5 M Tris-HCl, 0.4% SDS)



- 18.17 g of TRIZMA® base
- 0.4 g of SDS
- Dissolve in 80 mL of distilled water
- Adjust to a pH de 8.8 with HCl and gauge to a 100 mL
- Filtrate the solution
- Store at 4 °C degrees

Tris-HCl/ SDS 4X pH 6.8 (0.5 M Tris-HCl, 0.4% SDS)

- 6.06 g de TRIZMA® base
- 0.4 q de SDS
- Dissolve en 80 mL of distilled water.
- Adjust to a pH de 6.8 with HCl and gauge to a 100 mL
- Filtrate the solution
- Store at 4 °C degrees

■ 10% Ammonium Persulfate

- 500 mg of APS (Ammonium Persulfate)
- Dissolve en 5 mL of distilled ultra pure water
- Store at -20 °C for up to two weeks
- Store at 4 °C degrees for one use only

2X Loading Buffer

- 2 mL of glycerol
- 400 uL of Mercapto
- 0.02 g of bromophenol blue
- 0.4 g of SDS
- 2.5 mL of 8.6 pH buffer

1X Running Buffer

- 14 g of glycine.
- 3 g of Trizma base.
- 1 g of SDS.
- Gauge to a 1 L.

Staining Solution

- 1.25 g of brilliant blue R in 250 mL of methanol
- 200 mL of distilled water.
- 50 mL of Glacial Acetic Acid
- Store in room temperature while protected from light

Destaining Solution

- 250 mL of methanol
- 62.5 mL of Glacial Acetic Acid
- 312.5 mL of distilled water.
- Store in room temperature

Usage:Pour solution into a container and allow to stir until the SDS gel is clear or electrophoresis bands are visible.

Protocol 3h 45m

- 1 Clean the components of the electrophoresis camera with 70% ethanol and gauze 's.
- 2 Assemble the chamber and check that there are no leaks by pouring distilled water between the glasses.
- 3 Prepare polyacrylamide gels:

Α	В
Reagents	2 minigels
Distilled water	3.4 mL
Acrylamide/Bis 30%	4 mL
Tris-HCI/SDS	2.5 mL
4X pH 8.8 (1.5	
M Tris-HCl,	
0.4% SDS)	
ASP 10%	100 μL
TEMED	4 μL

Separation gel (12%)

4 Pour the solution between the glasses with a 1 mL micropipette, leaving a space of 1.5 cm for the concentrating gel. To level, distilled water is added and allowed to settle for © 00:30:00 or until a line is seen between the gel and the water.

Α	В
Reagents	2 minigels
Distilled water	2.7 mL
Acrylamide/ Bis 30%	1 mL
Tris-HCI/ SDS 4X pH 8.8 (1.5 M Tris-HCI, 0.4% SDS)	1.3 mL
ASP 10%	50 μL
TEMED	4 μL

Concentrating gel (6%)

5	Pour solution onto separating gel using a 1 mL micropipette.
6	1h 30m © 00:30:00 © 00:30:00 Insert the comb (carefully avoiding the formation of bubbles) and leave to solidify for © 00:30:00
7	When the gels are polymerized, prepare an electrophoresis chamber with 1X running buffer until it covers the gels and it reaches the line of two gels.
8	Sample PTake the pellets contained in Eppendorf tubes ■ Take the pellets contained in Eppendorf tubes ■ Add □300 µL of 1X loading buffer and resuspend the pellet.reparation:
9	To denature proteins, heat samples in boiling water for 5 min.
10	Load gels with the hot sample.
11	In the first well add \$\boxed{\subseteq 7} \mu L\$ of the molecular weight marker.
12	Once the samples are loaded, run the gel at 80 V for © 00:20:00 and then at 180 V for © 00:45:00
13	Turn off the camera and disarm it.
14	Remove the gels from the glasses and place in a container with staining solution. Leave stirring for one hour.

Remove staining solution after one hour, add destaining solution and leave stirring for 40m

© 00:20:00 Change the destaining solution and leave again for © 00:20:00

16 Leave stirring until the gel is transparent.

17 Analyze the gels.

15