Jun 13, 2019

Ethanol precipitation of RNA from small or large volumes

DOI

dx.doi.org/10.17504/protocols.io.36fgrbn

Stephen Floor¹

¹University of California, San Francisco

Stephen Floor Lab



Stephen Floor

University of California, San Francisco



DOI: dx.doi.org/10.17504/protocols.io.36fgrbn

Protocol Citation: Stephen Floor 2019. Ethanol precipitation of RNA from small or large volumes. protocols.io <u>https://dx.doi.org/10.17504/protocols.io.36fgrbn</u>

License: This is an open access protocol distributed under the terms of the <u>Creative Commons Attribution License</u>, 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: June 13, 2019

Last Modified: June 13, 2019

Protocol Integer ID: 24487

Keywords: RNA, ethanol precipitation

Abstract

How to precipitate RNA with ethanol and resuspend it! Two separate protocols - one for small and one for large volumes. The procedure for large volume precipitation is similar, but begins with an initial precipitation and transfers to a small tube for the final steps to avoid loss in the large tube. Strongly recommended to move to a smaller tube.

Recommended reading: Walker & Lorsch RNA purification--precipitation methods.

Materials

🔀 70% Ethanol

X 100% Ethanol (KOPTEC) VWR International (Avantor) Catalog #89125-186

0.3 M NaOAc pH 5.2

3 M NaOAc pH 5.2

Note: the protocol requires 70% ethanol pre-chilled to -20 C and requires 100% ethanol pre-chilled to -20 C for large volumes.

Precipitating RNA (< 500 ul sample volume)

- 1 Record the sample volume. If unknown, you can estimate it using the side of the tube or meausure it by pipetting it.
- 2 Add NaOAc pH 5.2 to sample to a final concentration of 0.3M

Note

This can be accomplished by diluting the sample by at least five volumes with 0.3M NaOAc or adding 3M NaOAc to a final concentration of 0.3M.

[CRITICAL] Do not exceed 500 ul NaOAc-diluted volume per 1.5 ml tube. For large volumes, see alternate protocol below.

3 Add 1 ul glycoblue per 250 ul sample volume

Note

volume here is the volume after adding NaOAc, not the original volume.

4 Vortex briefly

5 Add 2 volumes 100% ethanol

Note

volume here is the volume after NaOAc, not the original volume. For example, add 1 ml ethanol to 500 ul NaOAc solution.

6 Vortex briefly

7 Incubate at -80 degrees for at least 1 hour or in a dry ice ethanol bath for 30 minutes.

	Note
	[PAUSE] RNA is more stable for long periods when stored in ethanol at low temperature than as a pellet. Shorter incubation times are probably ok - glycoblue protocol recommends 30 minutes at -20 C.
8	Pellet RNA at 🚯 16000 x g , in tabletop microfuge 16k g for 25 minutes at 4 °C
	Note
	Tip: orient the tubes so the hinge is pointed outwards and the pellet will be easier to identify along the hinge-side of the tube.
9	Aspirate supernatant carefully without dislodging pellet, first using a P-1000 for most volume and then a P-20 for the end. Using a vacuum aspirator is not recommended as it is easy to aspirate the pellet.
10	Gently add 500 ul of 70% ethanol at -20 °C
	Note
	the warmer the 70% ethanol is, the more RNA will re-solubilize. Take the 70% out from the -20 right before adding it and then return it to the -20.
11	Pellet RNA at 🚯 16000 x g , in tabletop microfuge 16k g for 5 minutes at 4 °C
12	Aspirate supernatant carefully without dislodging pellet, first using a P-200 for most volume and then a P-20 for the end.
13	Quick spin to bring down remaining ethanol
14	Remove as much ethanol as possible without aspirating pellet using a P-20
15	Dry for 5 minutes at RT and not longer - fully dried pellets are harder to resuspend

- 16 Add the desired volume of water or buffer
- 17 Scrape the tube along a tube rack to dislodge pellet
- 18 Quick spin
- 19 Measure concentration using nanodrop or qubit

Precipitation of RNA (large sample volume)

20 Add NaOAc to sample to a final concentration of 0.3M

Note

This can be accomplished by diluting the sample by at least five volumes with 0.3M NaOAc or adding 3M NaOAc to a final concentration of 0.3M.

- Add 1 ul glycoblue per 250 ul sample volume (after adding NaOAc)
- 22 Vortex briefly
- Add 2 volumes 100% ethanol (1 ml 100% ethanol to 500 ul NaOAc-diluted sample)
- 24 Vortex briefly
- 25 Incubate at -80 degrees 1 hour to overnight.

Note

[PAUSE] RNA is more stable for long periods when stored in ethanol at low temperature than as a pellet.

Pelleting and resuspension of RNA 26 Centrifuge in a conical tube at > ~5000g, for 25 minutes 27 Aspirate supernatant carefully with a serological pipet or vacuum aspirator Note don't worry about getting all the ethanol off - trying to get all the ethanol off will likely remove some pellet as this is a loose pellet 28 Add 750 ul of 100% ethanol at -20 °C 29 Resuspend chunks of pellet in the 100% ethanol by pipetting up and down 30 Transfer ethanol and chunks to a microfuge tube 31 Spin for 5 minutes at 16k g at 4 °C 32 **ED** go to step #12 to wash and resuspend pellet