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Version 1

Mammalian Cell Culture: Subculturing V.1



Forked from Mammalian Cell Culture: Subculturing

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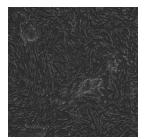
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Protocol status: Working

We use this protocol and it's working

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Abstract

This protocol details how to subculture nearly confluent mammalian cells grown in a T-75 flask.

Materials

- Gloves
- 0.05% or 0.25% warmed Trypsin-EDTA
- Warmed cell culture Media (e.g. DMEM:F12, EMEM)
- DPBS
- 15 mL centrifuge tube
- Serological pipette and tips
- 1000 μL pipette and tips

Troubleshooting

Safety warnings



Gloves must be worn at all times. Perform all work within biosafety cabinet.

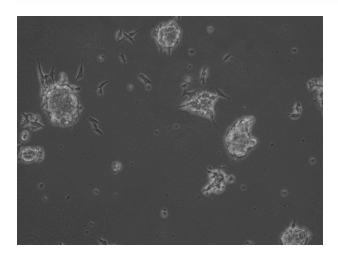


Assess Cell Confluency

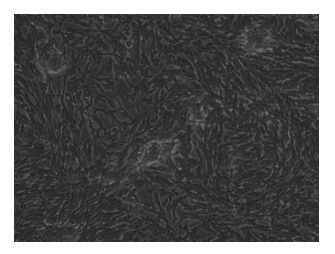
Under light microscope, look at the cells and assess level of confluency. This is how you will determine the need to subculture.

Note

Confluency can be estimated by evaluating the percentage of surface covered by cells.



Low confluency SH-SY5Y



High confluency SH-SY5Y

Wash Cells



- 2 Remove media from flask.
- 3 Using serological pipette, add <u>A 1 mL</u> DPBS to T-25 flask.
- 4 Using serological pipette, remove DPBS and dispose into waste beaker.
- 5 Repeat the above 2 steps, so that you will wash the cells twice.

Note

Always use a fresh pipette tip when drawing liquid from a stock solution.

Trypsinize

- 6 Add 4 mL warmed trypsin-EDTA to T-25 flask.
- 7 Wait 00:05:00 for trypsin-EDTA to detach the cells.

Note

This time will vary in practice, and depends on cell type and trypsin concentration (i.e. 0.05% vs 0.25%). Some cell types may take up to 15 minutes. In those cases, assess detachment progress using a light microscope.

8 Add 4 mL cell culture media.

Note

Trypsin-EDTA is neutralized by adding a volume of cell culture media equal to that of trypsin-EDTA.



Spin Down

- 9 Using a serological pipette, transfer the cell suspension (cells, trypsin-EDTA, and cell culture media) into a 15 mL centrifuge tube.
- Add 4 9.5 mL fresh cell culture media to T-25 flask, this will preserve any remaining cells.
- 11 Centrifuge the cell suspension on 1.5 kRPM for 00:03:00.

Resuspend and Reseed

12 Remove supernatant, dispose into waste beaker.

Note

You can leave a small amount with the serological pipette, the rest will be taken off in the next step.

13 Using a 1000 μL pipette, carefully remove the remaining supernatant, being cautious not to disturb the cell pellet.

Safety information

Always dispose of pipette tips in sharps container. Do not use the same tip twice.

- Add \perp 1000 μ L cell culture media to the cell pellet, and allow to sit for \bigcirc 00:01:00 .
- 15 Gently pipette mix the cell pellet until the pellet is resuspended.

Note

Pipette mixing is done by slowly drawing in solution and pushing it out several times, all without removing the pipette tip from the solution. Ask T.A. for help on this if you need some pointers.



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Note

You may seed more than 2 flasks, just use smaller volumes in each.

17 Label flask with updated passage number along with the date.

Incubate

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