1. Description

This product is for research use only.

Components
- 2× 45 mL Debris Removal Solution

Capacity
- For 50 applications with 1 g of tissue per application.

Storage
- Store protected from light at 2–8 °C. Do not freeze. The expiration date is indicated on the vial label.

1.1 Background information

Cell debris often occurs after tissue dissociation and impairs downstream applications. The Debris Removal Solution has been developed for efficient removal of cell debris from all kind of tissue after dissociation. It has been particularly tested for debris removal from dissociated adult rat and mouse heart tissue and mouse tumor tissue. In case of adult mouse or rat heart tissue, the Neonatal Heart Dissociation Kit, mouse and rat has been used for preparation of single-cell suspensions. For dissociation of mouse tumor tissue, the Tumor Dissociation Kit, mouse has been used. The debris-free single-cell suspensions can be analyzed in vitro for phenotype distributions, and other functional, genetic, or proteomic studies. Furthermore, cells can be subsequently cultured or used for cell separation with MACS Technology.

1.2 Applications

- Debris removal from dissociated tissue to improve downstream applications, such as cell separation, cell culture, cellular or molecular analysis.

1.3 Reagent and instrument requirements

- Centrifuge with a swinging bucket rotor
- Phosphate-buffered saline buffers, e.g., PBS, D-PBS
- 15 mL reagent tubes
- (Optional) gentleMACS™ Dissociator (# 130-093-235), gentleMACS Octo Dissociator (# 130-095-937), or gentleMACS Octo Dissociator with Heaters (# 130-096-427)

- (Optional) gentleMACS C Tubs (# 130-093-237, # 130-096-334)
- (Optional) Multi Tissue Dissociation Kit 2 (# 130-110-203)
- (Optional) Red Blood Cell Lysis Solution (10×) (# 130-094-183)

Figure 1: Debris removal protocol overview, steps 6–13
2. Protocol

▲ Volumes given below are for debris removal from a cell suspension resulting from up to 1 g of starting material. When working with more than 1 g starting material, split the single-cell suspension in several reagent tubes. When working with less than 1 g, use the same volumes as indicated. When working with less than 0.5 g of tissue scale down all volumes according to the table below.

▲ For subsequent cell separation and cultivation it is recommended to dissociate at least 0.5 g of tissue.

▲ Always use pre-cooled buffers and solutions (4 °C). Both buffers, PBS as well as D-PBS, can be used.

▲ The use of a centrifuge with a swinging bucket rotor is highly recommended for an optimal debris removal.

<table>
<thead>
<tr>
<th>Resuspension (PBS or D-PBS)</th>
<th>Debris Removal Solution</th>
<th>Overlay (PBS or D-PBS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5–1 g tissue</td>
<td>6200 μL</td>
<td>1800 μL</td>
</tr>
<tr>
<td>&lt;0.5 g tissue</td>
<td>3100 μL</td>
<td>900 μL</td>
</tr>
</tbody>
</table>

▲ Note: In case of very small amount of tissue (<100 mg) cell debris removal can be performed in a 2 mL tube using 300 μL of the Debris Removal Solution, 1000 μL of PBS/D-PBS for resuspension of the cell suspension, and ~1000 μL PBS/D-PBS for overlay.

1. Centrifuge the cell suspension at 300×g for 10 minutes at 4 °C.
2. Aspirate supernatant completely.
3. Resuspend cell suspension carefully with the appropriate volume of cold buffer according to the table above and transfer cell suspension to a 15 mL tube. Do not vortex.
4. Add appropriate volume of cold Debris Removal Solution.
5. Mix well by pipetting 10 times slowly up and down using a 5 mL pipette.
6. Overlay very gently with 4 mL of cold buffer.
   ▲ Note: Tilt tube and pipette very slowly to ensure that the PBS/D-PBS phase overlays the cell suspension and phases are not mixed.
7. Centrifuge at 4 °C and 3000×g for 10 minutes with full acceleration and full brake. Three phases are formed.
   ▲ Note: If centrifuges give suboptimal centrifugation, the acceleration and brake can be reduced.
8. Aspirate the two top phases completely and discard them.
9. Fill up with cold buffer to a final volume of 15 mL.
10. Gently invert the tube three times. Do not vortex!
11. Centrifuge at 4 °C and 1000×g for 10 minutes with full acceleration and full brake.
12. Aspirate supernatant completely.
13. Resuspend cells carefully in the appropriate buffer or medium by pipetting slowly up and down. Do not vortex!
14. (Optional) If the cell pellet contains a high amount of erythrocytes it is recommended to remove them by using the Red Blood Cell Lysis Solution.
15. Cells should be processed immediately for further applications.

3. Example of debris removal after adult rat heart dissociation using the Debris Removal Solution

One total adult rat heart was dissociated using the Multi Tissue Dissociation Kit 2 in combination with the gentleMACS™ Octo Dissociator with Heaters. Subsequently, debris was depleted using the Debris Removal Solution and erythrocytes were removed using the Red Blood Cell Lysis Solution (10×). Both debris removal and erythrocytes lysis steps are described in the protocol for the dissociation of adult mouse and rat heart using the Multi Tissue Dissociation Kit 2. Cells were analyzed by flow cytometry using the MACSQuant™ Analyzer 10 based on scatter signals to demonstrate absence of debris after debris removal.

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