Instructions for Use
CELLdisc™ - Multi Layer Device (1 - 40 Layers)

The Greiner Bio-One CELLdisc™ is a ready-to-start, multi layer device, as easy to use as a T-flask. The innovative ergonomic CELLdisc™ design provides a versatile system for the propagation of adherent mammalian cells from research scale to industrial batches. It is available either with the standard tissue culture surface (TC; red screw cap) or the Advanced TC™ surface (blue screw cap) identical to all Greiner Bio-One cell culture products to assure consistent performance from lot to lot and from format to format.

The single layer CELLdisc™ (CD1), in contrast to the larger multi layered versions, allows easy monitoring of cell growth, cell morphology and confluence. As conditions within the single layer CELLdisc™ will be identical to those within larger units, the CD1 can be used to anticipate when media changes are required and harvesting is recommended in any larger units being cultivated alongside the CD1. To ensure that the ambient conditions for the reference CD1 and the multi layer CELLdisc™ are absolutely identical, the CELLevator™ allows storage of the single layer CELLdisc™ on top of the CELLdisc™ production unit. To ensure safe usage in general, customers are advised to test Greiner Bio-One systems under the conditions defined in their own protocols.

Intended Use
The product is a cell culture disposable to be used by trained personnel in a laboratory surrounding.

General CELLdisc™ Details

- Filter port for pressure equilibration
- Integrated continuous gas support channel
- Large opening port for easy filling
- Connecting channel for liquid transfer
- Rim assures equal thermal distribution
Instructions for Use

1. **Single Layer CELLdisc™ (Order no. 678101)**

1. Unpack the single layer CELLdisc™ and place it in a laminar air flow cabinet in order to work in sterile conditions.
2. Prepare cell suspension in accordance with the concentration (cells/cm²) used with other disposables for adherent cell culture. Unscrew screw cap and transfer the cell suspension directly into the CELLdisc™ using the large opening port either by pouring or pipetting (Fig. 1). Firmly tighten the screw cap onto the CELLdisc™ to close it.
3. Tilt the single layer CELLdisc™ gently from one side to the other to assure that media and cells distribute evenly.

**Warning:** The media or cell suspension should **not touch the filter.** If the filter has absorbed any fluid, this will inhibit any gas transfer into and out of the CELLdisc. In this case the disposable has to be discarded and a new single layer CELLdisc™ has to be used.

2. **CELLdisc™ 4 - 16 Layers (Order no. 678104/-904, 674108/-908, 678116/-916)**

**Note:** To obtain equivalent cell growth in all layers, formation of air bubbles must be avoided during CELLdisc™ processing. Therefore, an exact angle of 30° and a specific position of the central filling channel (see Fig. 2 + Fig. 3) must be maintained during filling of 4-, 8- and 16-layer CELLdisc™. This handling procedure guarantees that the pressure is equalized through the central gas channel without contact with the medium (indicated in blue in Fig. 4). Thus, the air does not flow through the liquid and does not create air bubbles. In addition to the filling process, any generation of air bubbles should be avoided. Vigorous shaking of the CELLdisc™ is not recommended. To achieve this, larger volumes of liquids should be mixed outside the CELLdisc™ and then added to the disposable as described below. Small amounts can be pipetted directly into the CELLdisc™ and then distributed to all layers by repeating the equilibration process.

Filling and emptying of CELLdisc™

1. Unpack the CELLdisc™ and place it in a laminar air flow cabinet in order to work in sterile conditions.
2. Prepare cell suspension in accordance with the concentration (cells/cm²) used with other disposables for adherent cell culture.
3. Hold the CELLdisc™ at an **angle of 30°** (Fig. 2) with the screw cap at the indicated position (Fig. 3) and unscrew the screw cap. Add the cell suspension either by pouring or pipetting directly into the CELLdisc™ using the large opening port as indicated in Fig. 4 and 5. The media will fill the topmost layer first and then move slowly to each layer underneath.
Wait until the liquid is distributed into the individual layers and firmly tighten the screw cap onto the CELLdisc™ to close it (Fig. 5 + 6).

To start the liquid equilibration lay down the CELLdisc™ horizontally still keeping the screw cap in the indicated position (Fig. 7 + 8). The media will now separate equally between each layer as the media and all layers are in contact through the central filling channel.

**Note:** Small volumes of liquids (e.g. trypsin) may accumulate in the upper layers during filling and not flow through the whole filling channel. To guarantee equal distribution the liquid must be in contact with all layers and with the filling channel before the equilibration process is initiated. Therefore the CELLdisc™ must be positioned horizontally with the opening port at the lowest position.

4. **Turn the CELLdisc™ as indicated in Fig. 8 to disconnect media flow from the filling channel (Fig. 9 + 10). Do not rotate the CELLdisc™ any further as this could lead to wetting of the filter.**

**Warning:** The media or cell suspension should not touch the filter. If the filter has absorbed any fluid, this will inhibit any gas transfer into and out of the CELLdisc™. In this case the disposable has to be discarded and a new CELLdisc™ has to be used.

5. **From this position raise the CELLdisc™ upright (Fig. 11) and place the disposable on a horizontal surface inside an incubator (Fig. 12). Proceed with the cultivation based on the appropriate protocol.**

To stack individual CELLdisc™ or a single layer CELLdisc™ on top of a production CELLdisc™ use the CELLelevator™ (Fig. 13). More details on CELLelevator™ can be found on the reverse side.
**3. CELLdisc™ 40 Layers (Order no. 678140/-940)**

**Filling and emptying of CELLdisc™**

1. As with the CELldisc™ 4-16 layers, air bubble formation must be avoided during processing to obtain even cell growth in all layers. While the position of the screw cap is identical (Fig. 3) a smaller angle of 20° must be maintained during filling of a 40-layer CELLdisc™ (Fig. 16). This handling procedure guarantees that the pressure is equalized through the central gas channel without contact with the medium (see Fig. 17, indicated in blue). Thus, the air does not flow through the liquid and does not create air bubbles.

2. To start the liquid equilibration lay down the CELLdisc™ (Fig. 19) still keeping the screw cap in the indicated position and turn CELLdisc™ as indicated in Fig. 20. From this position raise CELLdisc™ upright and place the disposable on a horizontal surface inside an incubator (Fig. 21).

3. The medium is removed identically to CELLdisc™ 4-16 layers (Fig. 14 + 15).
Harvesting of cells

The protocol for harvesting cells from a multilayer device such as CELLdisc™ which does not allow for direct pipette access varies slightly from your standard protocol. Especially as the detachment of cells can be visualized microscopically only for the bottom layer of a CELLdisc™ with one, four or maximum eight layers. This cell harvest protocol refers to standard techniques and specific suggestions to gain maximum cell yields.

In general, we recommend using the same dissociating solution and concentrations for enzymatic detachment that is used to harvest these cells from a standard Greiner Bio-One cell culture disposable. As the surface treatments and basic materials are identical for these vessels, standard protocols can be converted. The only adaption required is based on growth area and cell numbers per layer. The use of chelating agents such as EDTA in addition to the enzyme (trypsin, papain etc.) may improve cellular detachment.

Remove cultivation media either by pouring or aspiration as described above. Wash cells once with PBS or an equivalent buffer using approx. 20 ml per layer. Follow the same protocol as for CELLdisc™ filling to distribute the buffer through all layers. Then tilt the CELLdisc™ slowly back and forth to gently rinse each cell layer and remove all traces of media.

Remove PBS either by pouring or aspiration. Thereafter add 7-10 ml of your enzymatic dissociation agent per layer. For a CD4 this results in 28-40 ml which has to be equilibrated as described above. Incubate CELLdisc™ at 37 °C and 5 % CO₂ for 3-5 minutes. Tapping CELLdisc™ from the side can accelerate cellular detachment. Strong adherent cells might need longer incubation or stronger tapping.

Stop the enzymatic reaction by adding the same amount of serum-containing media or an appropriate inhibitor to the CELLdisc™, equilibrate the liquids and mix the solutions by gently tilting the CELLdisc™ back and forth. Thereafter cell suspension can be harvested for further processing by pouring or aspiration.

If removal of the enzymatic dissociation agent is desired, then spin cell suspension at 100xg for 5 minutes. Remove the dissociation agent containing media and resuspend cell pellet in fresh media.

### CELLdisc™ Product Overview

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<th>Order no.</th>
<th>Layers</th>
<th>Surface treatment</th>
<th>Surface [cm²]</th>
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<th>Max working volume [ml]</th>
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CELLdisc™ is also available with external filter, triple packed:

4 Layers: 678104-EXF  
8 Layers: 678108-EXF  
16 Layers: 678116-EXF
**CELLdisc™ Accessories**

**CELLlevator™**
Stacking device for CELLdisc™

Order no. 878071

Easy and secured CELLdisc™ stacking
CELLlevator™ is positioned within the circumferential rim of
the top plate of the CELLdisc™. Another CELLdisc™ can be
positioned on top of the CELLlevator™.

**CELLring™**
Levelling ring

Order no. 878075

CELLring™ assures an exact horizontal position of CELL-
disc™ compensating surface irregularities, e.g. of a working
bench or an incubator.

**CELLhandle™**
Gripping device

Order no. 878074

CELLhandle™ facilitates secure and easy transportation of
particularly large-sized CELLdisc™ formats as well as conve-
nient emptying of these.

A video animation showing the handling of CELLdisc™ is available on our website:
www.gbo.com/celldisc

For further information please visit our website www.gbo.com or contact us.

Greiner Bio-One:

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