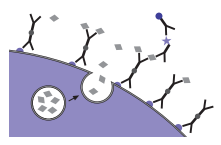


Mouse IL-10 Secretion Assay -**Cell Enrichment and Detection Kit (PE)**

For 50 tests with 10⁷ cells

Order no. 130-090-490



Unless otherwise specifically indicated, Miltenyi Biotec products and services are for research use only and not for diagnostic or therapeutic use.



Miltenyi Biotec B.V. & Co. KG Friedrich-Ebert-Straße 68, 51429 Bergisch Gladbach, Germany Phone +49 2204 8306-0, Fax +49 2204 85197 macsde@miltenyi.com, www.miltenyibiotec.com

Index	1. Descripti	on	
 Description Principle of the Mouse IL-10 Secretion Assay Background and product applications Reagent and instrument requirements Protocol overview 	Components	 ml Mouse IL-10 Catch Reagent: anti-IL-14 monoclonal antibody (rat IgG2b) conjugated to cel surface specific monoclonal antibody (rat IgG2b). ml Mouse IL-10 Detection Antibody: anti-IL-14 monoclonal antibody (rat IgG1) conjugated to PE (phycoerythrin). 	
Experimental set-up 3.1 Controls 3.2 Kinetics of restimulation and proposed time schedule 3.3 Counterstaining of cytokine secreting cells	Size	1 ml Anti-PE MicroBeads : colloidal super paramagnetic MicroBeads conjugated to monoclonal mouse anti-PE antibody (mouse IgG1). For 50 tests with 10 ⁷ cells	
3.4 Detection without prior enrichment	Product format	All components are supplied as a suspension containing 0.1% gelatine and 0.05% sodium azide.	
 Protocol for the Mouse IL-10 Secretion Assay 4.1 Cell preparation 4.2 (Antigen-specific) <i>In vitro</i> stimulation 	Storage	Store protected from light at 4°C. Do not freeze. The expiration dates are indicated on the vial labels.	
4.3 Cytokine Secretion Assay	1.1 Principle of	f the Mouse IL-10 Secretion Assay	
4.4 Magnetic labeling4.5 Magnetic separation5. Detection and analysis of IL-10 secreting cells	Secretion Assay, r cell preparations	For analysis of murine antigen-specific T cells using the Mouse IL-10 Secretion Assay, mouse spleen cells or other leukocyte containing single cell preparations are restimulated for a short period of time with specific peptide, protein or other antigen preparations.	

6. References

2

7. Appendix: Flask and dish sizes for stimulation

1. Description

Anti-PE MicroBeads IL-10 Detection Antibody (PE) IL-10 secreting cell

Subsequently, an IL-10-specific **Catch Reagent** is attached to the cell surface of all leukocytes. The cells are then incubated for a short time at 37°C to allow cytokine secretion. The secreted IL-10 binds to the IL-10 Catch Reagent on the positive cells. These cells can subsequently be labeled with a second IL-10 specific antibody, the **Mouse IL-10 Detection Antibody** conjugated to phycoerythrin (PE) for sensitive detection by flow cytometry.

The IL-10 secreting cells can now be magnetically labeled with **Anti-PE MicroBeads** and enriched over a MACS Column which is placed in the magnetic field of a MACS Separator. The magnetically labeled cells are retained in the MACS Column while the unlabeled cells run through. After the column has been removed from the magnetic field, the magnetically retained cells can be eluted as positively selected cell fraction, enriched for cytokine secreting cells. The cells can now 1. Description

5

1. Description

be used for cell culture or for analysis. Since viable cells are analyzed, non-specific background can be minimized by dead cell exclusion. This provides highest sensitivity of analysis.

1.2 Background and product applications

The Mouse IL-10 Secretion Assay is designed for the isolation, detection and analysis of viable IL-10 secreting murine leukocytes. It is specially developed for the detection and isolation of antigen-specific T cells after in vitro restimulation with specific antigen to induce secretion of IL-10. IL-10 is a cytokine predominantly secreted by CD4⁺ memory and effector T cells and antigen-presenting cells, e.g. monocytes/macrophages. It is believed to have important suppressive functions on immune responses and may also be involved in the maintenance of tolerance.

Quantitative analysis of antigen-specific T cell populations can provide important information on the natural course of immune responses. MACS enrichment of the antigen-specific T cells increases the sensitivity of analysis, allowing detection of frequencies as low as one in a million cells.

The MACS enrichment also enables further functional characterization of the antigen-specific cells and downstream experiments, as well as the expansion of antigen-specific cells allowing research on potential future immunotherapeutical applications.

4

1. Description

Examples of applications

- Detection and enrichment of viable IL-10 secreting mouse leukocytes for phenotypic and functional characterization.
- Detection and enrichment of IL-10 secreting antigen-specific T cells for enumeration, expansion and phenotypic as well as functional characterization.
- Isolation and expansion of antigen-specific T cells for research in immunotherapy, e.g. for adoptive transfer experiments.

1.3 Reagent and instrument requirements

- Buffer (degassed): phosphate buffered saline pH 7.2, containing 0.5% bovine serum albumin (BSA) and 2 mM EDTA (e.g. 4 ml of a 0.5 M EDTA stock solution per 1 liter of buffer).
- (Optional) 0.5 M EDTA stock solution: dissolve 56 g sodium hydroxide (NaOH) in 900 ml dd H₂O. Add 146.2 g ethylenediamine-tetraacetic acid, adjust pH to 7.5, fill up to 1000 ml with dd H₂O.
- Culture medium, e.g. RPMI 1640 containing 5% murine serum (do not use BSA or FCS because of non-specific stimulation!).
- **Propidium iodide (PI)** or **7-AAD** to exclude dead cells from the analysis.
- (Optional) Staining reagents such as CD4-FITC or CD8-FITC and CD45R/B220-PerCP[™].

• MACS Columns and MACS Separators:

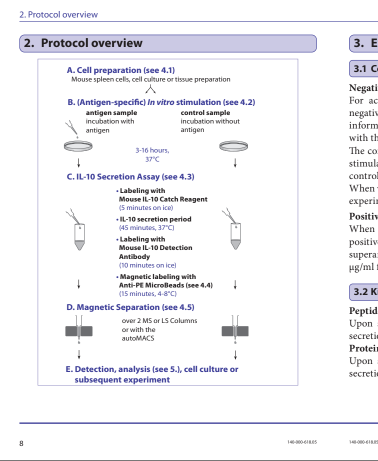
Column	max. number of labeled cells	max. number of total cells	Separator
MS	107	2 x 10 ⁸	MiniMACS, OctoMACS; with Column Adapter: VarioMACS, SuperMACS
LS	10 ⁸	2 x 10 ⁹	MidiMACS; with Column Adapter: VarioMACS, SuperMACS
autoMACS	2 x 10 ⁸	4 x 10 ⁹	autoMACS

▲ Note: Column adapters are required to insert certain columns into VarioMACS or SuperMACS. For details, see MACS Separator data sheets.

- Refrigerated centrifuge (4-8°C).
- Rotation device for tubes: MACSmix (# 130-090-753).
- (Optional) Pre-Separation Filter (# 130-041-407).

140-000-618.05

140-000-618.05



3. Experimental set-up

3. Experimental set-up

3.1 Controls

Negative control

For accurate detection of antigen-specific cells secreting IL-10, a negative control sample should always be included. This will provide information about IL-10 secretion unrelated to the in vitro stimulation with the specific antigen, e.g. due to ongoing in vivo immune response. The control sample should be treated exactly the same as the antigenstimulated sample except for the addition of antigen, or by using a control antigen.

When working with immunized mice, it could be relevant to include an experiment analyzing cells of a non-immunized mouse.

Positive control

When setting up a new experiment, it is recommended to include a positive control. As a positive control, a sample stimulated with the superantigen Staphylococcal Enterotoxin B (Sigma, St. Louis, USA) 10 μ g/ml for 3-16 hours, may be included in the experiment.

3.2 Kinetics of restimulation and proposed time schedule

Pentides

Upon stimulation with peptide, the cells can be analyzed for IL-10 secretion 3-6 hours after onset of stimulation.

Proteins

Upon stimulation with protein, the cells can be analyzed for IL-10 secretion 6-16 hours later.

3. Experimental set-up

It is possible to start the stimulation of the cells late in the afternoon, and to perform the IL-10 Secretion Assay the following morning.

Costimulation

The addition of costimulatory agents like CD28 antibody may enhance the response to the antigen. If costimulatory agents are added to the antigen sample, they also have to be included in the control sample.

3.3 Counterstaining of cytokine secreting cells

The IL-10 secreting cells are stained with PE-conjugated Mouse IL-10 Detection Antibodies. To identify cells of interest, counterstaining for T cells with e.g. CD4-FITC or CD8-FITC is important.

▲ Do **not use** tandem conjugates of phycoerythrin, like Cy-Chrome^{*} (PharMingen), PE-Cv5 (Serotec), ECD, PC5 (Coulter-Immunotech) etc., they may also be recognized by the Anti-PE MicroBeads.

▲ Upon activation of T cells, TCR and some associated molecules, like CD3, might be down-regulated.

▲ The samples should be stained with propidium iodide (PI) or 7-AAD prior to acquisition, to exclude dead cells from analysis. This will reduce non-specific background staining and increase sensitivity.

▲ For optimal sensitivity, we recommend labeling of undesired non-T cells such as B cells with antibodies conjugated to PerCPTM, e.g. CD45R/ B220-PerCPTM. These cells can then be excluded together with PI stained dead cells by gating.

4. Protocol for the Mouse IL-10 Secretion Assay

3.4 Detection without prior enrichment

(Optional) If the sample contains more than 0.01-0.1% of IL-10 secreting cells, the analysis can also be performed without prior magnetic enrichment (see also: Mouse IL-10 Secretion Assay - Detection Kit (PE) # 130-090-489).

4. Protocol for the Mouse IL-10 Secretion Assay

4.1 Cell preparation

Mouse spleen preparation

Prepare fresh mouse spleen cells or other leukocyte containing single cell preparations under sterile conditions according to standard protocols. Avoid excess of dead cells.

4.2 In vitro stimulation

Always include a negative control in the experiment. A positive control may also be included (see 3.1).

▲ Do **not use** media containing any **non-murine** proteins, like BSA or FCS, because of non-specific stimulation.

4. Protocol for the Mouse IL-10 Secretion Assay

4. Protocol for the Mouse IL-10 Secretion Assay



Protocol for in vitro stimulation

- 1. Wash cells by adding medium, centrifuge at 200xg for 10 minutes. Pipette off supernatant.
- Resuspend cells in culture medium at 10⁷ cells/ml and 5x10⁶ cells/ cm² (see 7. Appendix: Flask and dish sizes for stimulation).
- 3. Add antigen or control reagent:

peptide:	3-6 hours at 37°C, 7% CO_2 , e.g.	1-10 μg/ml
protein:	6-16 hours at 37°C, 7% CO ₂ , e.g.	10 µg/ml
SEB:	3-16 hours at 37°C, 5-7% CO ₂ , e.g.	10 µg/ml

For comparison of different experiments, the stimulation time should be kept constant (see 3.2).

4. Collect cells carefully by using a cell scraper, or by pipetting up and down when working with smaller volumes. Rinse the dish with cold buffer. Check microscopically for any remaining cells, if necessary, rinse the dish again.

4.3 Cytokine Secretion Assay

General considerations

▲ The assay is optimized for cell samples containing < 5% of total IL-10 secreting cells. If ≥ 5% of IL-10 secreting cells are expected, it is necessary to dilute the cells further during the cytokine secretion period, and therefore a larger test tube will be needed (see table below). The dilution avoids non-specific staining of cells not secreting IL-10 during this period.

 For each test with 10⁷ total cells, prepare: 100 ml of cold buffer (4-8°C)
 100 μl of cold medium (4-8°C)
 10 ml (or 100 ml; see table below) of warm medium (37°C).

▲ Work fast, keep the cells cold, use pre-cooled solutions which will prevent capping of antibodies on the cell surface and a non-specific cell labeling (exception: warm medium during secretion period).

▲ Volumes shown below are for 10^7 total cells. When working with less than 10^7 cells, use the same volumes as indicated. When working with higher cell numbers, scale up all reagent volumes and total volumes, accordingly (e.g. for $2x10^7$ total cells, use twice the volume of all indicated reagent volumes and total volumes).

▲ Do not remove supernatant by decanting. This will lead to cell loss and incorrect incubation volumes. Pipette off or aspirate supernatant.

▲ Dead cells may bind non-specifically to MACS MicroBeads or antibodies. Therefore, when working with cell preparations containing large amounts of dead cells, they should be removed before starting the IL-10 Secretion Assay, e.g. by density gradient centrifugation or by using the Dead Cell Removal Kit (# 130-090-101).

12

140-000-618.05 140-000-618.05

13

4. Protocol for the Mouse IL-10 Secretion Assay

Labeling cells with Mouse IL-10 Catch Reagent

- Use 10⁷ total cells in a 15 ml closable tube per sample.
 ▲ Note: For larger cell numbers, scale up all volumes accordingly. For less than 10⁷ cells, use same volumes.
- Wash cells by adding 10 ml of cold buffer, centrifuge at 300xg for 10 minutes at 4-8°C, pipette off supernatant completely.
 Note: Do not remove supernatant by decanting. This will lead to cell loss and

incorrect incubation volumes.

- 3. Repeat wash step, pipette off supernatant completely.
- 4. Resuspend cell pellet in 80 μl of $cold\ medium\ per\ 10^7$ total cells.
- 5. Add 20 μl of Mouse IL-10 Catch Reagent per 10^7 total cells and incubate for 5 minutes on ice.

EL-10 sec

IL-10 secretion period

 Add warm (37°C) medium to dilute the cells according to the following table:

Expected number of IL-10 secreting cells		
< 5 %	10 ⁶ cells/ml	10 ml
≥ 5 %	≤ 10 ⁵ cells/ml	100 ml

▲ Note: For frequencies of cytokine secreting cells >> 20% the cells need to be further diluted, e.g. by a factor of 5.

4. Protocol for the Mouse IL-10 Secretion Assay

 Incubate cells in closed tube for 45 minutes at 37°C under slow continuous rotation by using the MACSmix (# 130-090-753), or turn tube every 5 minutes to resuspend settled cells.

 \blacktriangle Note: During this step it is crucial to prevent contact of cells to avoid cross contamination.



Labeling cells with Mouse IL-10 Detection Antibody

- 1. Put the tube on ice.
- Wash the cell by filling up the tube with cold buffer, centrifuge at 300xg for 10 minutes at 4-8°C. Pipette off supernatant completely.
- 3. Repeat wash step, pipette off supernatant completely.
- 4. Resuspend cell pellet in 80 µl of **cold buffer** per 10⁷ total cells.
- 5. Add 20 μl of Mouse IL-10 Detection Antibody (PE) per 10^7 total cells.
- (Optional) Add additional staining antibodies, e.g. CD4-FITC or CD8-FITC and CD45R/B220-PerCPTM.
- 7. Mix well and incubate for 10 minutes on ice.
- Wash cells by adding 10 ml of cold buffer, centrifuge at 300xg for 10 minutes at 4-8°C. Pipette of supernatant.

4. Protocol for the Mouse IL-10 Secretion Assay

4. Protocol for the Mouse IL-10 Secretion Assay

1. Prepare two columns per sample by rinsing with cold buffer:

 Place the first column into the magnetic field of a MACS Separator (use column adapter with VarioMACS or SuperMACS).

(Optional) Pass the cells through Pre-Separation Filter

Collect unlabeled cells which pass through and wash with

appropriate amount of cold buffer. Perform washing steps by adding buffer successively once the column reservoir is empty.

Remove the first column from separator, place the second column

into the separator, and put the first column on top of the second

Pipette appropriate amount of cold buffer onto the first column.

Immediately and firmly flush out fraction with the magnetically labeled cells using the plunger supplied with the column, directly

LS: 5 ml

8. Collect unlabeled cells that pass through and wash with appropriate

LS: 3x3 ml

successively once the column reservoir is empty.

amount of cold buffer. Perform washing steps by adding buffer

LS: 3x3 ml

Collect total effluent. This is the unlabeled cell fraction.

LS Column: 3 ml

MS: 500 µl

5.

7.

one.

and discard effluent.

MS:3x500 ul

onto the second column. MS: 1 ml

MS: 3x500 µl

(# 130-041-407) to remove clumps.

4. Apply cell suspension onto the column.

4.4 Magnetic labeling

Magnetic labeling with Anti-PE MicroBeads

- 1. Resuspend cell pellet in 80 μ l of **cold buffer** per 10⁷ total cells.
- Add 20 μl of Anti-PE MicroBeads per 10⁷ total cells, mix well and incubate for 15 minutes at 4-8°C.

▲ Note: Incubate in refrigerator at 4-8°C, do not work on ice during this step.

- 3. Wash cells by adding 10 ml of **cold buffer**, centrifuge at 300xg for 10 minutes at **4-8°C**, pipette off supernatant.
- 4. Resuspend cell pellet in 500 μ l of **cold buffer** per 10⁷ cells, for higher cell numbers use a dilution of 10⁸ cells/ml.
- 5. (Optional) Take an aliquot for flow cytometric analysis and cell count of the fraction before enrichment.
- 6. Proceed to magnetic separation (see 4.5).

4.5 Magnetic separation

Magnetic separation using MS or LS Columns

▲ Choose an appropriate MACS Column and MACS Separator according to the number of total cells (see table in 1.3).

▲ When enriching antigen-specific T cells, **always perform two consecutive column runs** to achieve best results.

16

Ó

140-000-618.05 140-000-618.05

4. Protocol for the Mouse IL-10 Secretion Assay

- 9. Remove the second column from separator, place the column on a suitable collection tube.
- $10. Pipette appropriate amount of cold buffer onto the column. Immediately flush out the fraction with the magnetically labeled cells by firmly applying the plunger supplied with the column. MS: 500 \mu l LS: 5 ml$

▲ Note: For subsequent cell culture, the cells can also be eluted with medium. If part of the cells are analyzed by flow cytometry, the medium should **not contain** phenol red.

11. Proceed to analysis (see section 5.), cell culture or other subsequent experiment.



Magnetic separation using the autoMACS

▲ Refer to the autoMACS User Manual for instructions on how to use the autoMACS Separator.

- 1. Prepare and prime autoMACS.
- 2. (Optional) Pass cells through Pre-Separation Filter (# 130-041-407) to remove clumps.
- Place tube containing magnetically labeled cells in autoMACS. Choose separation program "Posseld". Collect the separated fractions from outlet port "pos2".
- 4. Proceed to analysis (see section 5.), cell culture or other subsequent experiment.

5. Detection and analysis of IL-10 secreting T cells

5. Detection and analysis of IL-10 secreting T cells

Add propidium iodide (PI) or 7-AAD to a final concentration of 0.5 μ g/ml **just prior** to acquisition for exclusion of dead cells from flow cytometric analysis. Incubating with PI for longer periods will affect the viability of the cells.

Do not fix the cells when using PI or 7-AAD.

▲ For optimized sensitivity, an appropriate number of viable cells has to be acquired from the antigen stimulated sample as well as from the control sample.

- Acquire 2x10⁵ events from the fraction before enrichment (see 4.4 step 5.).
- For **enumeration** of low frequent IL-10 secreting cells, acquire all of the positive fraction. For **preparative purposes**, acquire an aliquot of the positive fraction to determine the performance of the cell enrichment.

To illustrate the analysis, we describe the detection of IL-10 secreting T cells by using the Mouse IL-10 Secretion Assay. The detailed description, including how to set gates, may serve as a model for the analysis of your own sample.

5. Detection and analysis of IL-10 secreting T cells

5. Detection and analysis of IL-10 secreting T cells

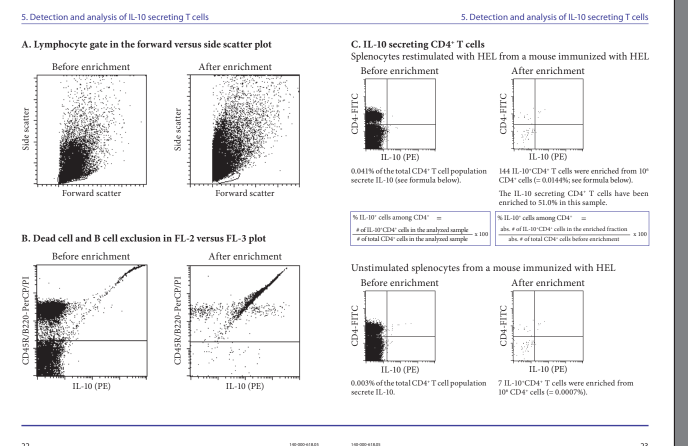
- 1. BALB/c mice were intraperitoneally (i.p.) immunized with 100 µg Henn eggwhite lysozyme (HEL) in incomplete Freund's adjuvant with 200 ng Pertussis Toxin. 200 ng Pertussis Toxin in PBS was i.p. injected again 24 hours later.
- After 3 weeks 107 mouse spleen cells of the immunized mouse were 2. incubated in vitro for 16 hours with or without 100 µg/ml HEL.
- The Mouse IL-10 Secretion Assay was performed on the stimulated 3 and the unstimulated sample.
- Counterstaining of T cells was performed by using CD4-FITC. 4.
- B lymphocytes were stained with CD45R/B220-PerCPTM. 5.
- Dead cells were stained with propidium iodide (PI), which was 6. added just prior to flow cytometric analysis to a final concentration of 0.5 µg/ml.
- $200{,}000$ viable cells of the original fractions and the complete 7. enriched fractions were acquired by flow cytometry, from the stimulated and the unstimulated samples.
- A lymphocyte gate based on forward and side scatter (FSC/SSC) 8. properties was activated prior to further gating to exclude B cells and debris (see A.).

9 Dead cells and B cells were excluded according to PI- and CD45R/ B220-PerCPTM-staining in a fluorescence 2 versus fluorescence 3 plot (see B.).

The dead cell exclusion is crucial for the analysis of rare antigenspecific T cells, as immunoglobulins or MicroBeads may bind nonspecifically to dead cells. This could lead to false positive events. The sensitivity of the detection will further be enhanced by exclusion of undesired non-T cells which may cause non-specific background staining.

10. For analysis IL-10 (PE) versus CD4-FITC staining of viable lymphocytes is displayed (see C.).

20



140-000-618.05

140-000-618.05

6. References

7. Appendix: Flask and dish sizes for stimulation

6. References

- Manz, R; Assenmacher, M; Pflüger, E; Miltenyi, S; Radbruch, A (1995) Analysis and Sorting of Live cells According to Secreted Molecules Relocated to a Cell-Surface Affinity Matrix. Proc.Natl.Acad.Sci. USA 92: 1921-1925. [139]
- Assenmacher, M; Löhning, M; Scheffold, A; Manz, RA; Schmitz, J; Radbruch, A (1998) Sequential production of IL-2, IFN-γ and IL-10 by individual staphylococcal enterotoxin B-activated T helper lymphocytes. Eur. J. Immunol. 28: 1534-1543. [483]
- Brosterhus, H; Brings, S; Leyendeckers, H; Manz, RA; Miltenyi, S; Radbruch, A; Assenmacher, M; Schmitz, J (1999) Enrichment and detection of live antigen-specific CD4⁺ and CD8⁺ T cells based on cytokine secretion. Eur. J. Immunol. 29: 4053-4059. [573]
- Ouyang, W; Löhning, M; Gao, Z; Assenmacher, M; Ranganath, S; Radbruch, A; Murphy, KM (2000) Stat6-Independent GATA-3 Autoactivation Directs IL-4-Independent Th2 Development and Commitment. Immunity 12: 27-37. [597]
- Hu-Li, J.; Pannetier, C; Guo, L; Löhning, M; Gu, H; Watson, C; Assenmacher, M; Radbruch, A; Paul, W (2001) Regulation of Expression of IL-4 Alleles: Analysis Using a Chimeric GFP/ Gene. Immunity 14: 1-11. [971]
- Hayakawa, Y; Takeda, K; Yagita, H; Kakuta, S; Iwakura, Y; Van Kaer, L; Saiki, I; Okumura, K (2001) Critical Contribution of IFN-γ and NK cells, but not perforinmediated cytotoxicity, to anti-metastatic effect of a-galactosylceramide. Eur. J. Immunol. 31: 1720-1727. [1073]
- Becker, C; Pohla, H; Frankenberger, F; Schüler, T; Assenmacher, M; Schendel, DJ; Blankenstein,T (2001) Adoptive tumor therapy with T lymphocytes enriched through an IFN-γ capture assay. Nature Medicine 7: 10. [1207]

For further references visit our website www.miltenyibiotec.com.

7. Appendix: Flask and dish sizes for stimulation

For (antigen-specific) stimulation (see 4.2 step 2.) the cells should be resuspended in culture medium at 10^7 cells/ml and $5x10^6$ cells/cm². Both the dilution and the cell density are important to assure optimum stimulation.

The following table lists culture plate, dish and flask sizes suitable for different cell numbers. It also indicates the appropriate amount of medium to add.

total cell number	medium volume to add	culture plate	well diameter
0.15 x 10 ⁷	0.15 ml	96 well	0.64 cm
0.5 x 10 ⁷	0.5 ml	48 well	1.13 cm
1 x 10 ⁷	1 ml	24 well	1.6 cn
2 x 10 ⁷	2 ml	12 well	2.26 cn
5 x 10 ⁷	5 ml	6 well	3.5 cn
total cell number	medium volume to add	culture dish	disl diamete
4.5 x 10 ⁷	4.5 ml	small	3.5 cn
10 x 10 ⁷	10 ml	medium	6 cn
25 x 10 ⁷	25 ml	large	10 cm
50 x 10 ⁷	50 ml	extra large	15 cr
total cell number	medium volume to add	culture flask	growt are
12 x 10 ⁷	12 ml	50 ml	25 cm
40 x 10 ⁷	40 ml	250 ml	75 cm
80 x 10 ⁷	80 ml	720 ml	162 cm
120 x 10 ⁷	120 ml	900 ml	225 cm

24

140-000-618.05 140-000-618.05

25

Refer to **www.miltenyibiotec.com** for all data sheets and protocols. Miltenyi Biotec provides technical support worldwide. Visit www.miltenyibiotec.com for local Miltenyi Biotec Technical Support contact information.

Warnings

Reagents contain sodium azide. Under acidic conditions sodium azide yields hydrazoic acid, which is extremely toxic. Azide compounds should be diluted with running water before discarding. These precautions are recommended to avoid deposits in plumbing where explosive conditions may develop.

Legal notices

Limited product warranty

Miltenyi Biotec B.V. & Co. KG and/or its affiliate(s) warrant this product to be free from material defects in workmanship and materials and to conform substantially with Miltenyi Biotec's published specifications for the product at the time of order, under normal use and conditions in accordance with its applicable documentation, for a period beginning on the date of delivery of the product by Miltenyi Biotec or its authorized distributor and ending on the expiration date of the product's applicable shelf life stated on the product label, packaging or documentation (as applicable) or, in the absence thereof, ONE (1) YEAR from date of delivery ("Product Warranty"). Miltenyi Biotec's General Terms and Conditions for the Sale of Products and Services available on Miltenyi Biotec's website at www.miltenyibiotec.com, as in effect at the time of order ("Product Warranty"). Additional terms may apply. BY USE OF THIS PRODUCT, THE CUSTOMER AGREES TO BE BOUND BY THESE TERMS.

THE CUSTOMER IS SOLELY RESPONSIBLE FOR DETERMINING IF A PRODUCT IS SUITABLE FOR CUSTOMER'S PARTICULAR PURPOSE AND APPLICATION METHODS.

Technical information

The technical information, data, protocols, and other statements provided by Miltenyi Biotec in this document are based on information, tests, or experience which Miltenyi Biotec believes to be reliable, but the accuracy or completeness of such information is not guaranteed. Such technical information and data are intended for persons with knowledge and technical skills sufficient to assess and apply their own informed judgment to the information. Miltenyi Biotec shall not be liable for any technical or editorial errors or omissions contained herein.

All information and specifications are subject to change without prior notice. Please contact Miltenyi Biotec Technical Support or visit www.miltenyibiotec.com for the most up-to-date information on Miltenyi Biotec products.

Licenses

This product and/or its use may be covered by one or more pending or issued patents and/or may have certain limitations. Certain uses may be excluded by separate terms and conditions. Please contact your local Miltenyi Biotec representative or visit Miltenyi Biotec's website at www.miltenyibiotec.com for more information.

The purchase of this product conveys to the customer the non-transferable right to use the purchased amount of the product in research conducted by the customer (whether the customer is an academic or for-profit entity). This product may not be further sold. Additional terms and conditions (including the terms of a Limited Use Label License) may apply.

CUSTOMER'S USE OF THIS PRODUCT MAY REQUIRE ADDITIONAL LICENSES DEPENDING ON THE SPECIFIC APPLICATION. THE CUSTOMER IS SOLELY RESPONSIBLE FOR DETERMINING FOR ITSELF WHETHER IT HAS ALL APPROPRIATE LICENSES IN PLACE. Miltenyi Biotec provides no warranty that customer's use of this product does not and will not infringe intellectual property rights owned by a third party. BY USE OF THIS PRODUCT, THE CUSTOMER AGREES TO BE BOUND BY THESE TERMS.

Trademarks

autoMACS, MACS, MACSmix, MidiMACS, the Miltenyi Biotec logo, MiniMACS, OctoMACS, SuperMACS, and VarioMACS are registered trademarks or trademarks of Miltenyi Biotec and/or its affi liates in various countries worldwide. All other trademarks mentioned in this publication are the property of their respective owners and are used for identification purposes only.

Cy is a registered trademark of GE Healthcare UK Limited.

Copyright © 2021 Miltenyi Biotec and/or its affiliates. All rights reserved.

28

140-000-618.05

140-000-618.05