

# **illustra** ProbeQuant G-50 Micro Columns

Product Booklet

cytiva.com

28903408PL AB

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# 1 Introduction

# Product code

28903408

## Important

Read these instructions carefully before using the products.

## **Intended** use

The products are intended for research use only, and shall not be used in any clinical or *in vitro* procedures for diagnostic purposes.

It is the responsibility of the user to verify the use of the illustra<sup>™</sup> ProbeQuant<sup>™</sup> G-50 Micro Columns for a specific application, as the performance characteristics of this product have not been verified for any specific organism.

## Safety

All chemicals should be considered as potentially hazardous. For use and handling of the products in a safe way, refer to the Safety Data Sheets.

## Storage

Store at ambient temperature (4°C to 30°C). Do not freeze.

# Expiry

For expiry date, refer to outer packaging label.

# 2 Components

# **Kit contents**

Identification	Pack size Product code	50 purifications 28903408
	illustra ProbeQuant G-50 micro columns	50
	Probe buffer type 1 (Blue colored cap)	10 mL
	Collection tubes	50

Refer to the Certificate of Analysis for a complete list of kit components.

Cytiva supplies a wide range of buffer types across the illustra nucleic acid purification and amplification range. The composition of each buffer has been optimized for each application and may vary between kits. Care must be taken to only use the buffers supplied in the particular kit you are using and not the buffers supplied in other illustra kits. Make sure you use the correct type of Probe buffer for your purification.

# Materials to be supplied by user

Disposables:

DNase-free 1.5 mL microcentrifuge tubes (screw caps are recommended when handling radioactivity).

Chemicals:

An excess of Probe buffer type 1 (150 mM STE buffer pH 8.0) is provided with the ProbeQuant G-50 micro columns. Should extra buffer be required, see *Buffer composition, on page 21* for buffer composition.

# **Equipment needed**

Microcentrifuge that accommodates 1.5 mL microcentrifuge tubes Vortex mixer (optional)

When measuring percent incorporation of radiolabel or determining counts per minute (cpm) per microliter (cpm/µL) of radiolabeled probe:

- Scintillation vials or microcentrifuge tube holders
- Scintillation counter
- Additional DNase-free 1.5 mL microcentrifuge tubes (screw caps are recommended when handling radioactivity).

# 3 Description

## Introduction

illustra ProbeQuant G-50 Micro Columns contain Sephadex<sup>™</sup> G-50 DNA grade F. They purify DNA by the process of gel filtration. Molecules larger than the largest pores in the Sephadex are excluded from the gel and elute first. Intermediate size molecules penetrate the matrix to varying extents, depending on their size. Penetration of the matrix retards progress through the column; very small molecules elute last. The volume required to elute these small molecules is dependent on the volume available both inside and outside the pores i.e. the bed volume.

Gel filtration resins do not exhibit a fixed exclusion limit when used in a spin-column format, as with illustra ProbeQuant G-50 micro columns. Exclusion limits of gel filtration resins are only meaningful in continuous flow processes where the molecules being purified have sufficient time to reach an equilibrium between the time spent in the gel filtration medium and the time spend in the eluent stream. In spin column chromatography, the observed exclusion properties that allow the product to pass through the gel while the smaller impurities are retained, depends on experimental factors, such as the resin used, sample volume, product size, and the g forces used during centrifugation.

The protocol provided with the illustra ProbeQuant G-50 micro columns has been optimized for the quantitative removal of unincorporated labeled nucleotides from a DNA labeling reaction only. Cytiva provides a wide range of nucleic acid purification products for other applications (see *Related products, on page 22* for more details).

illustra ProbeQuant G-50 micro columns can be used simultaneously for both preparative and analytical applications (labeled DNA must be at least 20 bases in length). The columns can be used for the following:

- Purification of a DNA labeling reaction prior to applications such as hybridization.
- Determination of the cpm/µL of labeled probe.
- Measurement of the percent incorporation of a radionucleotide precursor into a labeled probe.

illustra ProbeQuant G-50 micro columns are ready-to-use and require less than 4 minutes from Sample Application to completion of the Elution step. These columns provide a prepacked and preequilibrated alternative to Trichloroacetic Acid (TCA) precipitations. The sample elutes in 150 mM STE buffer, pH 8.0 (see *Buffer composition, on page 21* for buffer composition).

# The basic principle

Use of illustra ProbeQuant G-50 Micro Columns involves the following steps:



Step	Comments	Component
1. Sample Preparation	Radiolabeled sample is	Probe buffer type 1
	prepared and sample volume is made up to 50 µL with Probe buffer type 1.	A range of labeling productsare available from Cytiva (see <i>Related</i> <i>products, on page 22</i> ).
2. Column Preparation	Sephadex G-50 resin is re-suspended and excess storage buffer is removed by centrifugation.	illustra ProbeQuant G-50 micro column
3. Sample Application	Labeling reaction is applied to the center of the resin bed.	
4. Elution	Purified radiolabeled probe is eluted from the column, ready for downstream applications.	

# **Product specifications**

Table 1. illustra ProbeQuant G-50 Micro Column specifications

Sample Type	DNA radiolabeling reaction
Principle	Gelfiltration
Column matrix	Sephadex G-50 DNA grade F
Column storage buffer	150 mM STE containing 0.15% Kathon™ CG/ICP Biocide as preservative.
Input sample volume	25-50 µL
Percent sample recovery	>80%
Maximum column loading capacity	10 µg
Length of labeled DNA recovered	> 20 bases (N.B. there is no maximum length of probe that can be purified).

Sample Type	DNA radiolabeling reaction
Nuclease Testing	Column components are tested in nickase, single and double-stranded exonuclease and RNase assays.
Major subsequent applications	Determination of the cpm/µL of labeled probe.
	Measurement of the percent incorporation of a radionucleotide precursor into a labeled probe.
	Hybridizations, such as Southern blots.

Although tests have shown low levels of RNase activity for these columns, they are not specifically treated to be RNasefree. Customers regularly use our ProbeQuant G-50 Micro Columns for purification of RNA and are very satisfied. However, as the columns have not been treated for absence of RNase we cannot guarantee that RNA will not be degradedbut in principle it works well.

Biotinylated probes can be purified using the illustra ProbeQuant G-50 Micro Columns, as it is the size of the probe and not the modification that is important for purification purposes.

For purification of labeled DNA less than 20 bases in length, we recommend use of illustra MicroSpin<sup>™</sup> G-25 Columns. These columns are suitable for removal of unincorporated nucleotides from end-labeled oligonucleotides and small DNA fragments at least 10 bases long. Note that illustra MicroSpin G-25 Columns are supplied in double distilled water containing 0.05% Kathon and for optimal results may need to be equilibrated in 150 mM STE buffer pH 8.0 before use. If the use of a microcentrifuge with radiolabeled probes is an issue, consider use of the gravity flow system provided by illustra NICK Columns. Note that with illustra NICK Columns the final elution volume will be 400  $\mu$ L.

# 4 Protocol

**Note:** Columns are NOT transferable between Cytiva kits, e.g., the composition of the ProbeQuant G-50 micro columns is not the same as the composition of the MicroSpin G-50 columns.

See Materials to be supplied by user, on page 5 and Equipment needed, on page 5 for Materials & Equipment to be supplied by user.

# Definitions

This user documentation contains safety notices concerning the safe use of the product. See definitions below.



## NOTICE

**NOTICE** indicates instructions that must be followed to avoid damage to the product or other equipment.

- **Note:** A note is used to indicate information that is important for trouble-free and optimal use of the product.
- **Tip:** A tip contains useful information that can improve or optimize your procedures.

# Protocol for purification of radiolabeled probe

## **Column Preparation**

#### Step Action

1 Perform the radiolabeling reaction according to instructions (A range of labeling products are available from Cytiva).

#### Note:

Determination of  $cpm/\mu L$  of labeled probe and percent incorporation of radiolabel:

Reserve a 2  $\mu$ L aliquot of the radiolabeling reaction for analysis at this stage. This is the TOTAL sample. See Determination of cpm/ $\mu$ L of labeled probe, on page 16 & Determination of percent incorporation of radiolabel, on page 17 for processing and analysis of this sample.

- 2 Adjust the volume of the sample to 50 µL with Probe buffer type 1.
- **Tip:** If the sample volume is greater than 50 μL, use multiple columns, and apply a 50 μL aliquot to each column. Do not re-use columns. If more than 50 μL is loaded onto a column, unincorporated labeled nucleotides may elute off.

## **Column Preparation**

#### Step Action

1 Re-suspend the resin in the column by vortexing.

Step	Action
2	Loosen the cap one-quarter turn and twist off the bottom closure.
3	Place the column in the supplied Collection tube for support.
4	Spin for 1 minute at 735 × g.
5	Proceed immediately to <i>Sample Application, on page 14.</i>

*Tip:* See RPM calculation from RCF, on page 15 for RPM calculation from RCF.



# NOTICE

Use columns immediately after preparation to avoid drying out of the resin. If the column resin appears dry, displaced or cracked after the first spin, this is usually indicative of overcentrifugation (too fast or too long). Re-hydrate the column with  $250 \,\mu$ L of Probe buffer type 1, vortex and recentrifuge, checking the settings. Spin speed can be reduced by 20% if necessary. Do not use the pulse button on the microcentrifuge as this may override the speed setting.

# **Sample Application**

Step	Action
1	Place the column into a fresh DNase-free 1.5 mL microcentrifuge tube (user supplied).
2	Slowly apply 50 $\mu L$ of sample to the top-center of the resin, being careful not to disturb the resin bed.



# NOTICE

The resin will have come away from the column slightly to form a pillar. It is essential that the sample being purified is applied slowly and is not allowed to run down the sides of the resin bed. Avoid touching the resin bed with the pipet tip.

# Elution

#### Step Action

1 Spin for 2 minutes at 735 × g. The purified sample is collected in the bottom of the 1.5 mL microcentrifuge tube.

#### Step Action

2 Cap the microcentrifuge tube.

#### Note:

Determination of cpm/ $\mu$ L of labeled probe and percent incorporation of radiolabel:

Reserve a 2 µL aliquot of the purified sample for analysis at this stage. This is the COLUMN sample. See Determination of cpm/µL of labeled probe, on page 16 & Determination of percent incorporation of radiolabel, on page 17 for processing and analysis of this sample.

- 3 Proceed to Chapter 6 Quick Reference Protocol Card, on page 24 to determine the cpm/µL of labeled probe or Determination of percent incorporation of radiolabel, on page 17 to determine the percent incorporation of the radiolabel.
- 4 Store the purified probe at -20°C.

# 5 Appendices

# **RPM calculation from RCF**

The appropriate centrifugation speed for a specific rotor can be calculated from the following formula:

RPM = 1 000 × √(RCF/1.12r)

Where RCF = relative centrifugal force, r = radius in mm measured from the center of the spindle to the bottom of the rotor bucket, and RPM = revolutions per minute. For example, if an RCF of 735 × g is required using a rotor with a radius of 73 mm, the corresponding RPM would be 3 000.

The table below shows appropriate RPM for various microcentrifuges.

Microcentrifuge	Appropriate RPM for an RCF of 735 × g
Heraeus Biofuge 15	2 800
Beckman GS15R	2 100
Hettich Mikro 24-48	2 630
Hettich Mikro EBA12	2 700
Eppendorf Centrifuge 5415C	3 000
Eppendorf Centrifuge 5417C	2 700

Table 2. Appropriate RPM for an RCF of 735 × g

# Determination of cpm/µL of labeled probe

Step	Action
1	Mix 2 $\mu L$ of the "COLUMN" sample with 98 $\mu L$ of Probe buffer type 1 in a screw cap microcentrifuge tube.
2	Cap the tube and vortex gently to mix. Pulse spin in a microcentrifuge to collect the sample in the bottom of the microcentrifuge tube.
3	Dispense $2 \times 50 \mu\text{L}$ aliquots of the diluted sample into duplicate screw capped microcentrifuge tubes.
4	Transfer each tube to a scintillation vial or suitable tube holder for placement into the scintillation counter. Alternatively, the samples may be added directly to scintillation vials containing scintillant; cap and invert to mix.

#### Step Action

5 Count the samples using an appropriate program.

#### Tip:

<sup>32</sup>P-labeled samples may be counted using a 1 minute Cerenkov counting program if no scintillant is used, or a standard counting program if scintillant is used.

6 Add the counts per minute obtained for the duplicate samples and divide by 2. This value is the average cpm or cpm/μL of the labeled probe.

#### Tip:

To determine the total cpm incorporated into the purified sample, multiply cpm/µL by the volume of the sample recovered from the column.

## Determination of percent incorporation of radiolabel

Step	Action
1	Mix 2 $\mu L$ of the "COLUMN" sample with 98 $\mu L$ of Probe buffer type 1.
2	Mix 2 $\mu L$ of the "TOTAL" sample with 98 $\mu L$ of Probe buffer type 1.
3	Cap the tubes and vortex gently to mix. Collect sample in bottom of microcentrifuge tube by pulse centrifugation.
4	Dispense 2 × 50 $\mu$ L aliquots of diluted COLUMN and 2 × 50 $\mu$ L aliquots of diluted TOTAL sample into 4 microcentrifuge tubes.

#### Step Action

- 5 Transfer each tube to a scintillation vial or suitable tube holder for placement into the scintillation counter. Alternatively, the samples may be added directly to scintillation vials containing scintillant; cap and invert to mix.
- 6 Count the samples using an appropriate program.

#### Tip:

<sup>32</sup>P-labeled samples may be counted using a 1 minute Cerenkov counting program if no scintillant is used, or a standard counting program if scintillant is used.

- 7 Add the cpm obtained for diluted "COLUMN" samples and divide by 2 to obtain the average. Add the cpm obtained for diluted TOTAL samples and divide by 2 to obtain the average.
- 8 Percent label incorporation may be calculated using the formula:

% label incorporartion =  $\frac{\text{average cpm for "COLUMN" sample}}{\text{average cpm for "TOTAL" sample}} \times 100$ 

#### Tip:

Percent incorporation values obtained using this product may vary from values previously obtained by standard TCA analysis or ethanol precipitation. This product is not designed to replace these methods, but is meant to offer a fast alternative to standard techniques.

# Troubleshooting guide

This guide may be helpful in the first instance, however if problems persist or for further information, contact Cytiva technical services. Telephone numbers are on the back page. Alternatively log onto cytiva.com/illustra.

# Problem: Resin appears dry and cracked after Column Preparation step.

Possible causes	Suggestions
Centrifugation steps too long or too fast.	• Make sure that columns are spun at the correct speed and for the appropriate time. For conversion of RCF to RPM see <i>RPM calculation from RCF, on page 15.</i>
	<ul> <li>If problems persist, reduce spin speed by 20%.</li> </ul>
	<ul> <li>If column is considered too dry, or centrifugation is known to have been too fast or too long, add 250 µL Probe buffer type 1 to the column and recentrifuge using the correct settings.</li> </ul>
Use of the pulse button on the microcentrifuge.	• Do not use the pulse button on the microcentrifuge as this could over-ride the speed setting.
There is nothing wrong; this is normal.	<ul> <li>After centrifugation to remove the storage buffer, the resin will appear dry and will pull away from the sides of the column. However, excessive cracking must always be investigated.</li> </ul>

# Problem: Final sample volume is greater than 50 µL

Possible causes	Suggestions
Spin for Column Preparation step too short or too slow.	• Make sure that columns are spun at the correct speed and for the appropriate time. For conversion of RCF to RPM see <i>RPM calculation from RCF, on page 15.</i>
Spin for Elution step too long or too fast.	• Make sure that columns are spun at the correct speed and for the appropriate time. For conversion of RCF to RPM see <i>RPM calculation from RCF, on page 15.</i>
	• If problems persist, reduce centrifugation step by 20%.
Use of the pulse button on the microcentrifuge.	• Do not use the pulse button on the microcentrifuge as this could over-ride the speed setting.

# Problem: Final sample volume is much less than 50 µl

Possible causes	Suggestions
Spin for Column Preparation step too fast or too long and/or Elution step too slow or too short.	• Make sure that columns are spun at the correct speed and for the appropriate time. For conversion of RCF to RPM see <i>RPM calculation from RCF, on page 15.</i>
	• If problems persist, reduce centrifugation speed by 20%.
	<ul> <li>If column is considered too dry, or centrifugation is known to have been too fast or too long, add 250 µL Probe buffer type 1 to the column and re- centrifuge using the correct settings.</li> </ul>
Use of the pulse button on the microcentrifuge when spinning ProbeQuant G-50 micro columns.	Do not use the pulse button on the microcentrifuge as this could over-ride the speed setting.

# Problem: Unincorporated nucleotides remain in purified sample

Possible causes	Suggestions
Column not used immediately after preparation, allowing the resin to dry and crack. Sample will run through the cracks.	Use column immediately after preparation.
Sample not pipetted onto top of resin bed, but allowed to run down the sides.	CAREFULLY pipet the sample onto the top center of the resin bed.
Sample volume added greater than 50 µL.	• If sample to be purified is greater than 50 µL, use multiple columns. purified.
Spin for Elution step too long or too fast.	• Make sure that columns are spun at the correct speed and for the appropriate time. For conversion of RCF to RPM see <i>RPM calculation from RCF, on page 15</i>
	• If problems persist, reduce centrifugation speed by 20%.

## **Buffer composition**

Use nuclease free water and sterile plasticware for buffer preparation. Filter sterilize before use.

To prepare 50 mL of additional Probe buffer type 1, dissolve 0.438 g of NaCl in 40 mL of TE buffer (10 mM Tris-HCl, pH 8.0, and 1 mM EDTA) with stirring. Titrate the solution to pH 8.0 using dilute HCl or NaOH, as required. Transfer the solution to a 50 mL graduated cylinder and adjust the volume to 50 mL with TE buffer, pH 8.0.

# **Related products**

A full range of Molecular Biology reagents can be found in the Cytiva catalog and on the web site cytiva.com/illustra

A full range of Detection Products and available pack sizes can be found in the Cytiva catalog and on the web site cytiva.com

Application	Product	Product code	Packsize
Blotting	Hybond™-N+ (82 mm)	RPN82B	50 discs
	Hybond-N+ (15 × 20 cm)	RPN1520B	10 sheets
	Hybond-NX (82 mm)	RPN82T	50 discs
	Hybond-NX (15 × 20 cm)	RPN1520T	10 sheets
	Hybond-N (82 mm)	RPN82N	50 discs
	Hybond-N (15 × 20 cm)	RPN1520N	10 sheets
	Hybond-XL (82 mm)	RPN82S	50 discs
	Hybond-XL (15 × 20 cm)	RPN1520S	10 sheets
Hybond blotting RPN610 paper (20 × 20 cm)		RPN6101M	100 sheets
Radioactive labeling	Rediprime II DNA Labeling System	RPN1633	30 reactions
	Ready-To-Go™ DNA Labeling Beads (-dCTP)	27924001	1 kit
	Megaprime DNA Labeling System, dNTP	RPN1604	30 reactions

Application	Product	Product code	Pack size
Radioactive labeling	Megaprime DNA Labeling System, dCTP	RPN1606	30 reactions
	Nick Translation Kit, dNTP	N5500	20 reactions
	Nick Translation Kit, dCTP	N5000	20 reactions
	5'-End Labeling Kit	RPN1509	20 reactions
	Rapid-Hyb Buffer	RPN1635	125 mL
Detection	Hyperfilm™ MP (18 × 24 cm)	28906843	50 sheets
	Hyperfilm MP Enveloped (18 × 24 cm)	28906850	50 sheets
	Hypercassette	RPN11642	1
Purification of DNA probes and	illustra MicroSpin G-50 Columns	27533001	50 purifications
oligonucleotides	illustra MicroSpin G-25 Columns	27532501	50 purifications
	illustra ProbeQuant G-50 Micro Columns	28903408	50 purifications
	illustra NICK Columns	17085502	50 purifications
	illustra NAP™-5 Columns	17085302	50 purifications

# 6 Quick Reference Protocol Card

# A. Protocol for purification of radiolabeled probe



<ul> <li>Mix 2 µl COLUMNsample with 98 µl Probebuffer type 1</li> <li>Vortex. Pulse centrifuge</li> <li>Vortex. Pulse contrifuge</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>Dispense 2 × 50 µl aliquots into suitable tubes or vials</li> <li>This is the cpm/µl of radiolabeled probe</li> </ul>	<ul> <li>Mix 2 μl of the TOTALsample with 98 μl Probebuffer type 1</li> <li>Vortex. Pulse centrifuge</li> <li>Vortex. Pulse centrifuge</li> <li>Dispense 2 × 50 μl aliquots into scintillation vials</li> <li>Count as appropriate</li> <li>Determine mean cpm of the TOTAL sample.</li> <li>Use formula below:</li> <li>Mabel incorporation = <u>average cpm for "COLUMN''sample</u> × 100</li> </ul>
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# B. Determination of cpm/ $\mu$ L of labeled probe and percent incorporation of radiolabel



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28903408PL AB V:7 02/2021