AVIPure® dsRNA Clear OPUS® Column

User Guide

RPT-00728v1 AVIPure® dsRNA Clear OPUS Column User Guide



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Contents

2. Ak	troduction	5
	y Performance Attributes	
Proce	ss Conditions	6
	4.1 Sample Volumes and Column Loading	
	4.2 Equilibration and Binding Conditions	
	4.3 Elution Conditions	
	4.4 Sanitization and Clean-in-place (CIP) Conditions	
	4.5 Storage	
4	4.6 Neutralization	7
4	4.7 Example Chromatogram	8
5. Or	dering Information	9
6. In	dex	0
List of	f Tables	
Table	1. Explanation of User Attention Phrases	5
	2. Safety Precautions	
Table	3. Process Steps	5
Table	4. Process Development Scale	Э
	5. Large Scale	
List of	f Figures	

Abbreviations

microgram μg microliter μL micron μm CIP clean-in-place circRNA circular RNA centimeter cm Column Volume CV dsRNA double-stranded RNA dΤ deoxythymidine

FLuc firefly luciferase antibody

g gram

Gu-HCl guanadine hydrochloride

liter L molar Μ mg milligram minute min milliliter mL millimolar mM mRNA messenger RNA NaCl Sodium chloride nanometer nm

OPUS Open Platform User Specified

PPE Personal protective equipment

saRNA self-amplifying RNA

sec second

ssRNA single-stranded RNA

1. Introduction

AVIPure® dsRNA Clear OPUS® Columns remove double-stranded RNA (dsRNA) byproducts from in vitro transcribed mRNA, saRNA, and circRNA. The capacity for dsRNA is ≥0.4 mg per mL of column volume (CV). A 1 mL AVIPure dsRNA Clear OPUS Column can process >10 mg of single-stranded RNA (ssRNA) dependent on dsRNA content.

2. About This Document

This manual uses several different phrases. Each phrase should draw the following level of attention:

Table 1. Explanation of User Attention Phrases

Phrase	Description
Note:	Points out useful information.
IMPORTANT	Indicates information necessary for proper instrument operation.
PRECAUTION	Cautions users of potential physical injury or equipment damage if the information is not heeded.
WARNING!	Warns users that serious physical injury can result if warning precautions are not heeded.

3. Safety Precautions

Table 2. Safety Precautions

Symbol		Description		
WARNING	(!)	Wear standard laboratory personal protective equipment (PPE), including lab coat, protective eye wear, and gloves.		
WARNING	(1)	This product is for laboratory and manufacturing production use only. Not for administration to humans.		
IMPORTANT	(1)	This product is shipped in an $18.0 \pm 1\%$ ethanol solution, a recognized bacteriostatic agent. It is flushed from the resin during equilibration and preparation for use. Follow all local regulations for safe disposal.		
WARNING		 Flammable liquid and vapor. Keep away from heat/spark/open flame/hot surfaces. No smoking. Keep container tightly closed. Ground/bond container and receiving equipment. Store in a well-ventilated place. Keep cool. 		
IMPORTANT	(1)	Dispose of contents/container in accordance with local/regional/national/ international regulations.		
IMPORTANT	(1)	For a full list of precautionary statements, please read the <u>Safety Data Sheet</u> (SDS).		

4. Key Performance Attributes

- High specificity of the affinity ligand provides effective dsRNA binding and low non-specific binding
- Simple flow-through process with no organic solvent requirement
- High salt conditions may enable dsRNA removal prior to poly-dT chromatography
- Compatible with standard bioprocess columns and relevant process flowrates

Process Conditions

Optimal conditions for each RNA sample must be determined empirically. dsRNA removal is enhanced with high salt (>0.5 M NaCl) and pH values between pH 7.2 and 9.5. The level of dsRNA byproducts in a given RNA can vary by several orders of magnitude; therefore, the column should be loaded with approximately 10 mg RNA/mL $_{CV}$ for initial testing. Fractions should be analyzed for dsRNA breakthrough to determine the appropriate ssRNA challenge for a given feed. Depending on the feed, the column may be able to clear significantly more than 10 mg RNA/mL $_{CV}$, but loading the column to a 10% breakthrough capacity might require loading hundreds of column volumes (in the case of a low dsRNA feed) and long processing times. To avoid this, the column should be sized to allow processing in the desired time frame (e.g., 30 – 60 minutes).

The pressure limit of the resin is 3 bar. Maximum flow velocity is 1200 cm/h for PD-scale columns with 5 cm bed height, and 600 cm/h for columns with 10 cm bed height.

Table 3. Process Steps

Step	Column Volumes	Residence Time (Min)	Suggested Buffer	Notes
Sanitization (OPTIONAL)	15	1	Choose one: • 6M guanidine hydrochloride • 0.1 M phosphoric acid • 1 M acetic acid • 100 mM NaOH • 70% Ethanol	A 15 min static hold can also be used for this step
Equilibration	2 – 5	1	Match oligo-dT load buffer, OR 50 mM HEPES, 0.7 M NaCl, 2 mM EDTA, pH 7.2	pH and conductivity return to baseline
Load	>5	1	Adjust RNA to 0.5 – 1.5 M NaCl, with pH 7.2 – 9.5 buffer.	Filter RNA solution after adjusting to final pH and conductivity. Challenge with ≥10 mg/mL _{CV} . Monitor sample for time-dependent precipitation
Chase	5	1	Equilibration buffer	-
Elution/CIP	5	1	6 M guanidine hydrochloride	dsRNA elutes and column is regenerated
Re-equilibration	2-5	1	Equilibration buffer	pH and conductivity return to baseline
Long-term storage	3	1	18 – 20% ethanol	at 2 – 8° C

The quality of the RNA sample is an important factor in successful processing. Samples that display a single crisp band on a gel or CE trace give the best results (i.e., >2 log reduction of dsRNA). Samples that are smeary, degraded, or display multiple bands may have lower dsRNA clearance. For the best results, optimize the IVT (in vitro transcription) parameters until a feed has a single, sharp RNA band.

Initially, use purified RNA suspended (or diluted) in the oligo dT binding buffer used to purify the RNA. Oligo-dT binding buffers have high salt (>0.5 M), and compatibility with the specific RNA should already be established. Processing purified RNA in this manner should provide the highest dsRNA clearance (>2 log reduction of dsRNA). Some RNAs precipitate slowly (over $^{-15}$ – 60 minutes) in high-salt buffers after initially appearing clear and passing through a 0.22 μ m filter. This precipitation reduces the yield of RNA in the flow-through and increases the RNA that elutes in 6 M guanidine hydrochloride. This may be avoided by decreasing the salt, by mixing the RNA and salt immediately before the column using inline dilution techniques, or by heating the sample ($^{-30}$ – $^{-37}$ °C) to increase solubility. Because different RNA molecules have different solubility profiles, these approaches need to be tailored for individual RNAs.

To determine the feasibility of situating the AVIPure dsRNA Clear OPUS Column prior to oligo dT purification, perform a crossover study with crude IVT mixtures to determine if any components of the crude IVT reaction interfere with the AVIPure dsRNA Clear OPUS Column. Ensure the solution is filtered before loading the column, as some components of the IVT mixture may precipitate in

high salt. When using the AVIPure dsRNA Clear OPUS Column in line with an oligo dT step, bypass the AVIPure column during the oligo dT water elution step.

4.1 Sample Volumes and Column Loading

Because the column operates in flow-through mode, the sample volume should be at least several times larger than the column volume. RNA concentrations between 0.25 and 1 mg/mL are common, and higher concentrations can be tested if desired. Attempting to process very small sample volumes (<5 CV) can lead to very dilute samples due to peak broadening as there is no bind/elute step to concentrate the sample. The sample can be applied by pump or sample loop.

To test the recommended 10 mg mRNA/mL_{CV} in a 1 mL column, 10 mg of mRNA are required. With 0.5 mg/mL mRNA, the sample volume would be 20 mL, and at 1 mg/mL mRNA the sample volume would be 10 mL. Using lower quantities and volumes of RNA can lead to lower apparent recoveries due to excessive dilution from peak broadening, and non-specific binding to system tubing and fraction vessels.

To evaluate larger challenges (e.g., 30 - 100 mg mRNA/mL_{CV}), the sample volume can be increased. Alternatively, more concentrated samples can be evaluated. In general, the yield of dsRNA-depleted RNA will increase as the challenge increases. It is important to collect fractions to determine if dsRNA breakthrough occurs with increased challenge.

4.2 Equilibration and Binding Conditions

Binding of dsRNA in AVIPure dsRNA Clear OPUS Columns is optimal with 0.5 - 1.5 M NaCl and pH values between 7.2 and 9.5. The equilibration should match the composition of the RNA to be processed. Common biological buffers (HEPES, phosphate, tris, etc.) can be used interchangeably. If satisfactory results are not obtained with oligo dT binding buffer, it is recommended to increase the salt and/or pH and retest. This process may be more efficient in filter plate format with loose resin.

4.3 Elution Conditions

The bound dsRNA can be eluted with 6 M guanidine hydrochloride (GuHCl); this buffer also cleans and regenerates the column for reuse. The resin has been held for a total of 20 hours in 6 M GuHCl without loss in dsRNA capacity.

The GuHCl should be removed from the eluate before analytical testing.

4.4 Sanitization and Clean-in-place (CIP) Conditions

Testing indicates that AVIPure dsRNA affinity resin is stable for over 2.5 hours in 6 M GuHCl, 0.1 M phosphoric acid, 1 M acetic acid, 70% ethanol, and 0.1 M NaOH. Using 6 M GuHCl as an elution/CIP is recommended, adding additional CIP steps only if necessary.

4.5 Storage

Store columns in 18 - 20% ethanol at $2 - 8^\circ$ C. Benzyl alcohol is not recommended because it absorbs at 260 nm and can interfere with RNA quantitation.

4.6 Neutralization

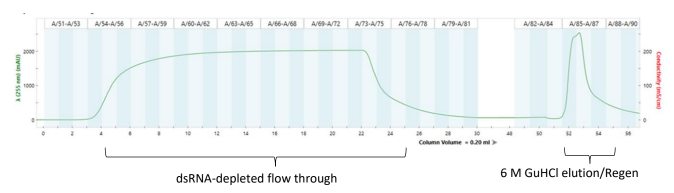
With higher pH values, neutralization buffer should be added to the post-column fractions. This should be tested to ensure compatibility with the specific RNA. A 10% volume of 1 M sodium acetate, pH 4 can be used to neutralize samples run with 50 mM of pH 9.5 carbonate running buffer.

4.7 Example Chromatogram

The absorbance trace on a chromatogram will show a broad peak during loading as the mRNA is flowing through (Figure 1).

- Column: 0.2 mL AVIPure dsRNA Clear OPUS Column
- Buffer: 10 mM Tris pH 8, 0.5 M NaCl, 0.5 mM EDTA
- Elution/CIP: 6 M guanidine HCl
- Flow rate: 0.4 mL/min (30 sec RT)
- Feed: 0.51 mg/mL FLuc with 1.4% dsRNA (by ELISA)
- Challenge: 10.2 mg FLuc/mL_{CV}
- Sample application method: Dynamic loop
- Sample volume: 20 CV

Figure 1. Example Chromatogram



5. Ordering Information

Contact your account manager to place an order, or contact your regional customer service using the email addresses below:

North America: $\underline{customerservice US@repligen.com}$

Europe: customerserviceEU@repligen.com
China: customerserviceEU@repligen.com

Table 4. AVIPure dsRNA Clear Affinity Resin - Bulk

Volume	Part #	Format	dsRNA capacity	Approx. mRNA to load	Notes
5 mL	100RNA-5	5 mL loose resin in 18% EtOH	0.4 mg dsRNA per mL of resin	10 mg per mL of resin	Ideal for high throughput screening in 96-well plate format or small drip columns

Table 5. AVIPure dsRNA Clear OPUS Columns

Column Volume (mL)	Part #	Format	dsRNA capacity	Approx. mRNA to load	Notes	
8 x 200 μL	23052208R	Strip of 8 Robocolumns®	80 μg dsRNA per column	2 mg per column	For use with Tecan fluid	
8 x 600 μL	23052208R-30	Strip of 8 Robocolumns	240 μg dsRNA per column	6 mg per column	management system for rapid process development	
0.5 mL	23052205	0.5 x 2.5 cm Minichrom® column	0.2 mg	5 mg		
1 mL	23052206	0.5 x 5 cm MiniChrom column	0.4 mg	10 mg	For small scale process	
5 mL	23052204-100	0.8 x 10 cm MiniChrom column	2 mg	50 mg	development	
10 mL	23052207-100	1.13 x 10 cm MiniChrom column	4 mg	100 mg		
50 mL	23052210V-100	2.5 x 10 cm Valichrom® column	25 mg	500 mg	Excellent tool for scale up and process validation	
50 mL	BC-025-DRNA-100-G	2.5 cm ID x 10 cm bed height	20 mg	500 mg	Process 3 L per hour	
200 mL	BC-050-DRNA-100-G	5 cm ID x 10 cm bed height	80 mg	2 g	Process 12 L per hour	
500 mL	BC-081-DRNA-100-G	8.1 cm ID x 10 cm bed height	200 mg	5 g	Process 31 L per hour	
800 mL	BC-100-DRNA-100-G	10 cm ID x 10 cm bed height	320 mg	8 g	Process 47 L per hour	
1200 mL (1.2L)	BC-126-DRNA-100-G	12.6 cm ID x 10 cm bed height	480 mg	12 g	Process 75 L per hour	
1500 mL (1.5 L)	BC-140-DRNA-100-G	14 cm ID x 10 cm bed height	600 mg	15 g	Process 92 L per hour	
3000 mL (3.0 L)	BC-200-DRNA-100-G	20 cm ID x 10 cm bed height	1200 mg (1.2 g)	30 g	Process 188 L per hour	
5000 mL (5.0 L)	BC-250-DRNA-100-G	25 cm ID x 10 cm bed height	2000 mg (2.0 g)	50 g	Process 295 L per hour	

6. Index

CIP	3, 6, 7, 8
Elution	6, 7, 8
Equilibration	6, 7
Flow Velocity	6
Load	6
Note	5
Ordering	9

Precautions	5
Pressure Limit	ε
Process	6, 9
Safety	
Storage	6, 7
Volumes	6, 7
Marning	-

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