

# User Guide

## RNA (R1) Cartridge Kit (C105110/C105210/C105810)

### A. Specifications

Specification	Description
L.O.D	5 ng/μl
Sample Number	100 runs
Shelf Life	4 months

### B. Kit Components and Storage Conditions

Item	Storage Condition
RNA Cartridge (C105110/C105210/C105810)	15-27°C (Do Not Freeze)
5X Lower Marker (C109120-100A, 100μL)	Short-Term (≤ 3 months): 4-27°C Long-Term (> 3 months): -20°C
10X Separation Buffer (C104409-10X, 15mL/C104412-10X, 50mL)	4-27°C
10X Dilution Buffer (C104408-10X, 8mL/C104411-10X, 25mL)	4-27°C
Mineral Oil (C104404, 8mL)	4-27°C

### C. Cartridge Unpacking Preparation

New cartridge must pass HV check and calibration before use. Please follow unpacking guide to unpack and use 5-times diluted 5X Lower Marker to do calibration.

### D. Buffer and Marker preparation

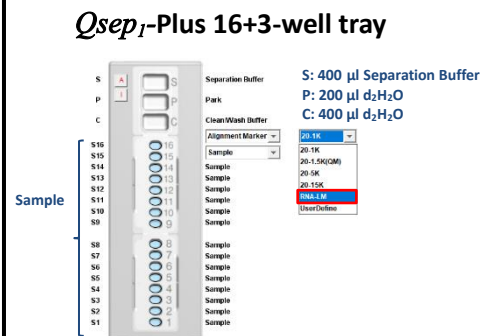
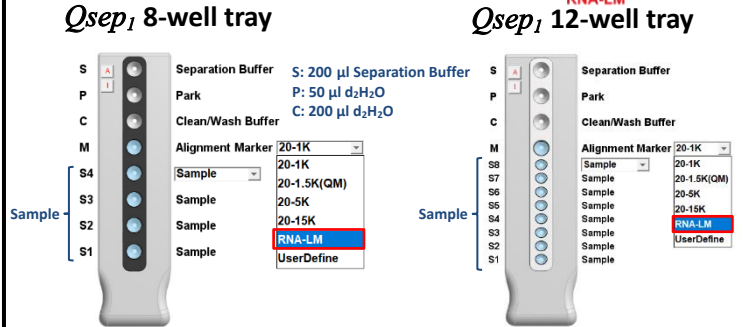
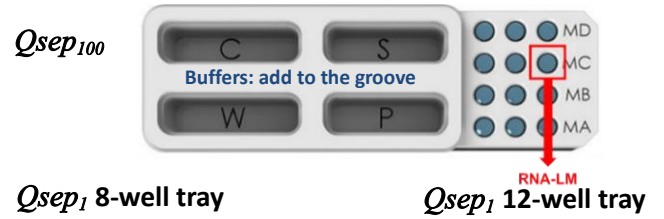
#### Buffer Preparation

- Separation Buffer (1X): 10X dilution from the stock with DEPC-treated water.
- Dilution Buffer (1X): 10X dilution from the stock with DEPC-treated water.

#### Marker Preparation

- Lower Marker (1X): 5X dilution with 1X dilution buffer.

Material	Volume (μl)
5X Lower Marker (C109120-500A)	5
1X Dilution Buffer	20
<b>Total Volume</b>	<b>25</b>



### E. Sample Preparation

#### Sample Pre-treatment

Heat-denature RNA samples at **70°C for 2 minutes** and put on ice for at least 5 minutes.

#### Sample Volume Requirements

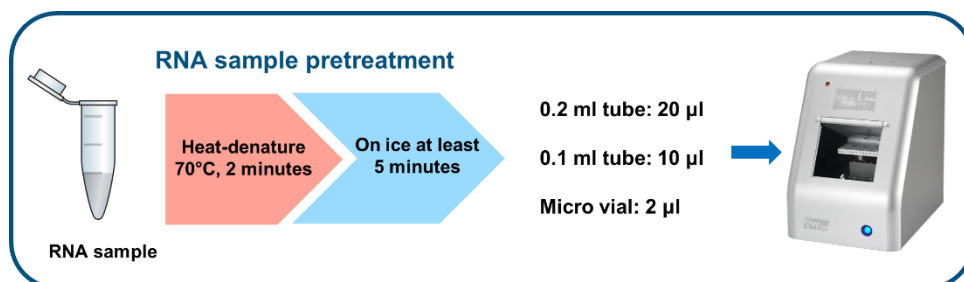
0.2 ml tube: 20 μL      Micro Vial (C104250): 2 μL  
0.1 ml tube: 10 μL      16+3-Well Tube (C104254): 10 μL

#### Recommended Sample Concentration

RNA sample: 5-50 ng/μL

**\*NOTE: If sample concentration is over 50 ng/μL, dilute sample with 0.1-1X dilution buffer based on buffer condition.**

**\*NOTE: If sample is eluted in RNase free water, add dilution buffer to make the sample into ≥ 0.1X dilution buffer condition.**



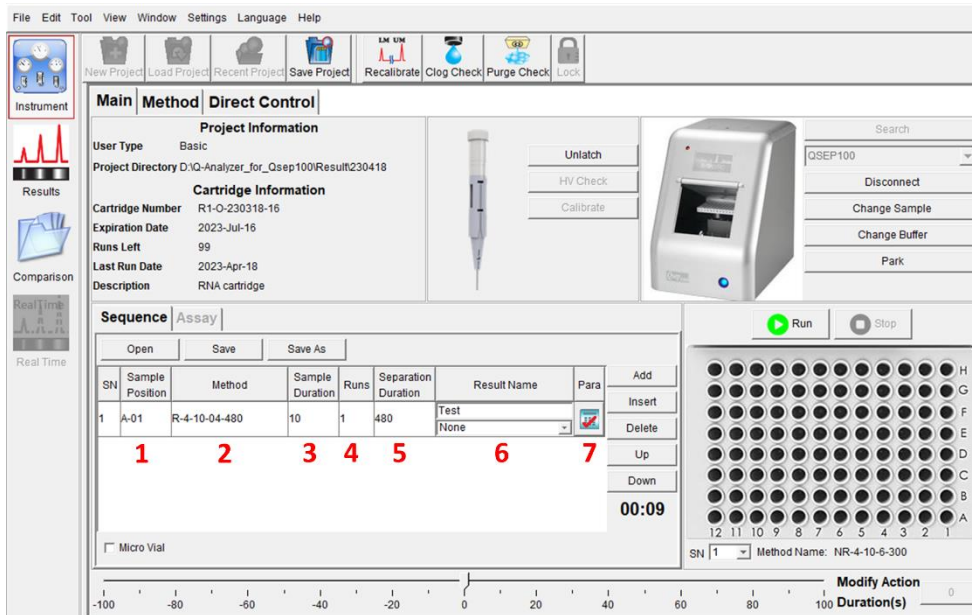
#### Contact Information:

Company Name: Bioptic Inc.  
Address: (23141) 5F., No.6, Ln. 130, Minquan Rd., Xindian District, New Taipei City, Taiwan (R.O.C)  
Tel: +886-2-2218-8726, Fax: +886-2-2218-8727, E-mail: service@bioptic.com.tw

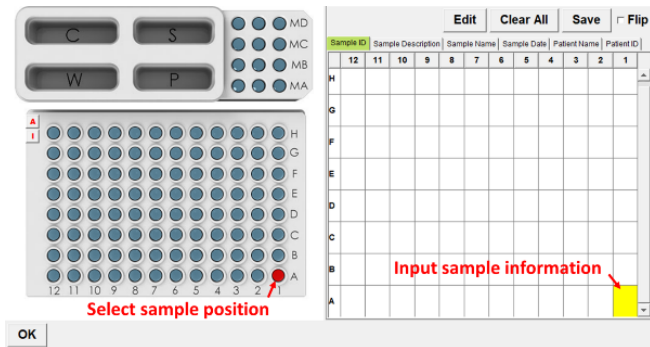
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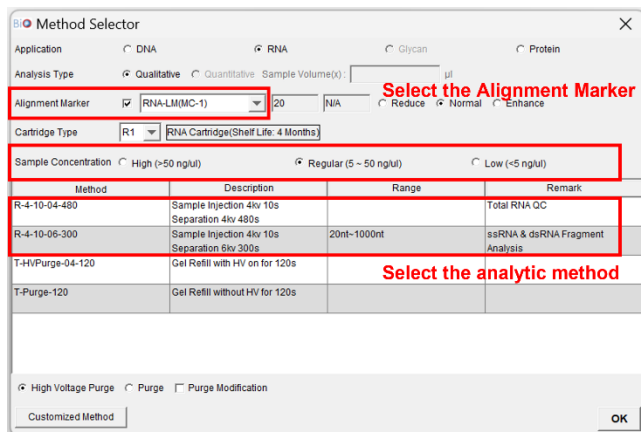
### F. Software Operation



1. Place the sample and select corresponding position, and then input sample information (optional).



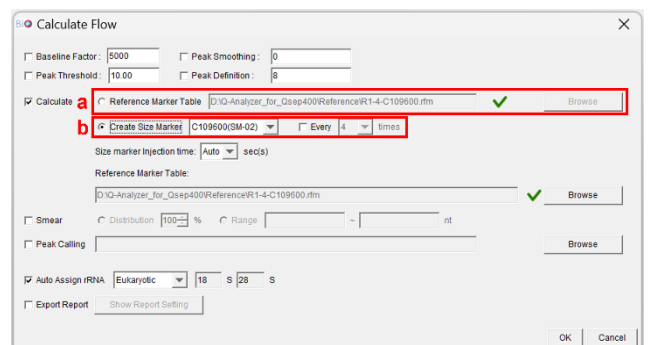
2. Select the alignment marker and the analytic method in Method Selector.



**\*NOTE: Based on sample concentration to adjust injection condition.**

Sample concentration	High (2kV, 10s)	Regular (4kV, 10s)	Low (8kV, 10s)
RNA	> 50 ng/μl	5~50 ng/μl	2~5 ng/μl

3. Sample Duration: adjust the sample injection time to increase/decrease injection amount.  
**\*NOTE: Do not set the injection time over 20 sec.**
4. Runs: repetition time.
5. Separation Duration: adjust the duration to extend/reduce the separation time.  
**\*NOTE: Step 3-5 are optional.**
6. Input the result name for result file.
7. Click "Para" . Choose (a) reference or (b) create size marker to do the calculations.



**\*NOTE: When using function "Create Size Marker", select C109600 to pair with the Lower Marker.**

8. Click "Run" to start the analysis.

#### Contact Information:

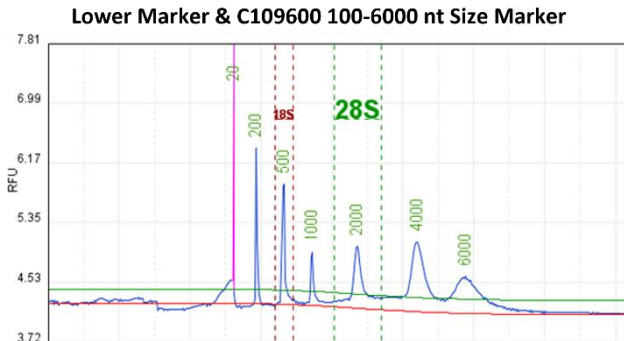
Company Name: BiOptic Inc.  
Address: (23141) 5F., No.6, Ln. 130, Minquan Rd., Xindian District, New Taipei City, Taiwan (R.O.C)  
Tel: +886-2-2218-8726, Fax: +886-2-2218-8727, E-mail: service@bioptic.com.tw

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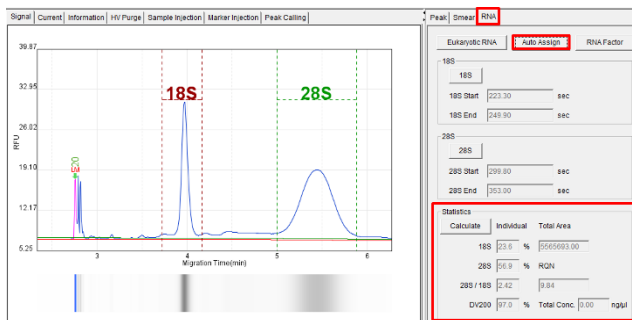
#### G. Results

##### • Lower Marker & Size Marker



##### • RNA Quality Number (RQN)

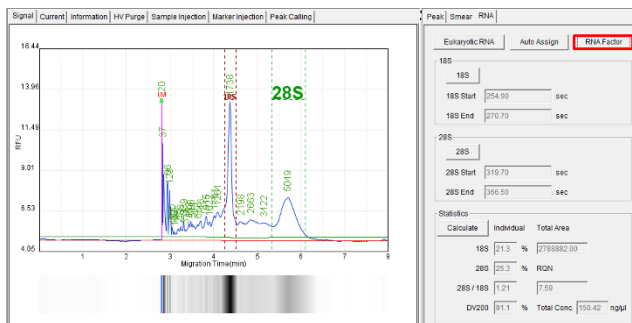
The software will identify 18S and 28S automatically. Then, the 28S/18S ratio and RQN value from 1 to 10 will be provided. If the software does not assign 18S and 28S regions, click "Auto Assign".



##### • 18S/28S identification

If the software cannot identify 18S and 28S region correctly, please follow the instructions below.

1. Adjust the 18S and 28S regions by manually dragging the red line (18S) and the green line (28S).
2. Adjust the "RNA factor" to identify 18S/28S region for total RNA from same species.



3. Enter the lower marker and 18S/28S start and end times according to the result. Click "Save" and "Apply".

Item	Time	nt
LM	195.60	20
18Ss	327.56	1700
18Se	346.28	2100
28Ss	389.21	4200
28Se	439.28	5200

#### H. Troubleshooting

Please ensure the whole system is working well and the operation is following the instructions.

Sometimes, there are some unknown substances in PCR reagent buffer or other kit buffer. These substances may cause unstable current in sample injection or separation steps. Here is a list of solutions to help fix the occurrence.

1. Use dilution buffer to dilute the sample.
2. Centrifuge the sample for a while to make the residues accumulate at the bottom of the tube.
3. Insert a "T-Purge-120" method between sample runs. E.g., Insert a "T-Purge-120" every 5-10 sample runs.

SN	Sample Position	Method	Sample Duration	Runs	Separation Duration	Result Name	Para	Add
1	A-01A-0	M-4-10-06-300	10	1	300	Test	None	Insert
2		T-Purge-120	0	1	0	Test	None	Delete
3	A-05A-0	M-4-10-06-300	10	1	300	Test	None	Down

e.g. Insert 1 run "T-Purge-120" every 5-10 sample runs

#### I. Cartridge Discard

Please wear gloves before discarding cartridge.

##### Gel reservoir



1. Bend the cartridge tip.
2. Open the cap on gel reservoir and remove the inner cap.
3. Pour the gel into the chemical waste container.
4. Cartridge can be thrown into the bin.

Cartridge tip

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