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# **LoopCap™ Viral Target Capture Kit**

ML5120 (96 reactions)  
ML5220 (384 reactions)

## **User Guide**

Molecular Loop Proprietary

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## Product Description

LoopCap™ Viral Target Capture Kits are used to prepare next-generation sequencing libraries for whole genome or targeted sequencing of RNA viruses. This protocol contains instructions for library preparation for sequencing on Illumina platforms, with IL and TN probe designs. Refer to product labeling for the probe design code, which contains either IL or TN in the descriptor, as well as the probe fill size and the design version number.

## Kit Contents

Reagent	Color	Quantity	Storage
RT Mix	Clear	1	-15 to -25 °C
Probe Mix	Blue	1	-15 to -25 °C
Fill-in Mix	Yellow	1	-15 to -25 °C
Cleanup Mix	Red	1	-15 to -25 °C
PCR Mix	Clear	1	-15 to -25 °C

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## Accessories

This protocol requires the use of ML2 series Index Primer Kits, and is not compatible with standard Illumina library amplification primers. Primers are supplied as ready-to-use premixes in 96-well plates. Refer to Index Primer Kit Product Data Sheets for plate layouts and index sequences.

Part #	Product Description	Contents
ML2100	Index Primer Kit, Illumina (96 samples)	1 x 96-well plate
ML2200	Index Primer Kit, Illumina (384 samples)	4 x 96-well plates
ML2100-TN	Index Primer Kit, Illumina (96 samples)	1 x 96-well plate
ML2101-TN	Index Primer Kit B, Illumina (96 samples)	1 x 96-well plate
ML2102-TN	Index Primer Kit C, Illumina (96 samples)	1 x 96-well plate
ML2103-TN	Index Primer Kit D, Illumina (96 samples)	1 x 96-well plate
ML2200-TN	Index Primer Kit, Illumina (384 samples)	4 x 96-well plates

**NOTE:** Index Primers must always be matched to the probe design. Use ML2100/ML2200 for IL probe designs only, and ML2100-TN for TN probe designs only. Amplification primers are not cross-compatible and use of the incorrect primers will result in reaction failure.

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## Reagents, Consumables, and Equipment

The items listed below are recommended when this protocol is performed manually in 96-well plates. Reactions may also be performed in single PCR tubes or strip tubes. Contact support@molecularloop.com for assistance when using liquid handling automation.

### User-supplied reagents

Item	Vendor	Catalog #
KAPA Pure Beads	Roche	KK8002
80% Ethanol (for bead purification)	Various	N/A
Bioanalyzer DNA 1000 Assay	Agilent	5067-1504
1X TE Buffer	Various	N/A
Custom Sequencing Primers (see below)	IDT	N/A

### Custom sequencing primers

Custom sequencing primers are required for **IL probe designs** only. Primers should be HPLC-purified, resuspended to a concentration of 100 µM, and diluted according to Illumina's specified final concentration and volume.

Primer	Sequence
Read 1 Primer	5' - ACGGATACCCACGACATGTAAAACGACGGCCAGT - 3'
Read 2 Primer	5' - CCAGAGGCAAACGACGCTAGTTATTGCTCAGCGG - 3'
Index 1 Primer (i7)	5' - CCGCTGAGCAATAACTAGCGTCGTTTTGCCTCTGG - 3'
Index 2 Primer (i5)*	5' - ACTGGCCGTCGTTTTACATGTCGTGGGTATCCGT - 3'

\* Index 2 Primer is not required for Illumina systems that use the forward strand workflow for dual-indexing, as described in Illumina document 15057455, "Indexed Sequencing Overview Guide".

## User-supplied consumables

Item	Vendor	Catalog #
96-well PCR plates, white/clear	Bio-Rad	HSP9601
Microseal B film	Bio-Rad	MSB1001
1.5 ml low-bind microcentrifuge tubes	Eppendorf	22431021
Pipette tips, filtered (various sizes)	Various	Various

## Equipment

Item	Vendor	Catalog #
Centrifuge (5810 or similar)	Eppendorf	5811000017
Centrifuge (5418 or similar)	Eppendorf	5418000017
Mastercycler X50s	Eppendorf	6311000010
Microfuge	VWR	C1413V-115V
Pipettes (P10, P20, P200, P1000)	Eppendorf / Gilson	Various
Film sealing roller for PCR plates	Bio-Rad	MSR0001
PCR workstation, AC600 series	Airclean	AC624LFUV
DynaMag-2 tube magnet	Thermo Fisher	12321D
Bioanalyzer 2100 (or similar)	Agilent	G2939A

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## RNA Input Recommendations

This protocol has been optimized for viral RNA applications. Depending on the application, variable amounts of RNA may be appropriate. For simple detection of RNA, single-copy sensitivity may be possible but best results will be achieved if reactions contain at least 100 copies of RNA. For whole RNA sequencing, best results will be achieved if reactions contain at least 10,000 copies of RNA.

**NOTE:** *The relationship between input copies and RT-qPCR Ct value may vary, and must be empirically determined by the user. Higher amounts of input RNA (lower Ct values) will generally produce superior results.*

Purified RNA should be resuspended in RNase-free water or TE with a pH no greater than 7.5. Contaminants including ethanol, sodium azide, sodium acetate, and guanidine salts may affect performance. DNase treatment is not recommended as trace amounts of residual DNase could negatively impact performance. The presence of small amounts of host DNA should not affect performance.

If RNA is quantified, a method that is specific for RNA is recommended (e.g. Qubit RNA BR Assay Kit or RT-qPCR), rather than one that will also detect DNA. To reduce inter-sample performance variability, all samples in a batch should be quantified using the same method and normalized to the same concentration.

## Thermal Cyclers and Programs

This protocol is sensitive to evaporation, particularly during the hybridization step. It is therefore important to use thermal cyclers with proper lid engagement that form a tight seal against the reaction plate. In addition, the cycler lid should be heated to 105 °C for all steps except the 4 °C hold at the end of the CleanupPCR program, where it may be omitted. Note that some thermal cyclers may not allow the use of a heated lid for lower block temperatures such as the 25 °C step during Reverse Transcription. Should this be the case, use a heated lid for all steps where it is permissible by the instrument to do so.

The thermal cycler programs listed below are used in this protocol.

Program and Steps	Temperature and Time (hh:mm:ss)
<b>RTHybFill</b> Reverse Transcription Denature Hybridize and Fill Hold	<b>Heated lid 105 °C</b> 25 °C 00:10:00, 50 °C 00:50:00 95 °C 00:01:00 55 °C 24:00:00 55 °C ∞
<b>CleanupPCR</b> Cleanup Denature PCR Hold	<b>Heated lid 105 °C</b> 45 °C 00:30:00 95 °C 00:03:00 N x [98 °C 00:00:15, 60 °C 00:00:15, 72 °C 00:01:00] 4 °C ∞

The number of PCR cycles to run (N) should be adjusted based on the amount of input RNA, the number of probes, the hybridization time, and the sensitivity of the quantification method. On-target yield (an amplicon of ≥250 bp) will change approximately linearly with respect to both input RNA and number of probes. A nonspecific amplicon of ~170-200 bp may also be observed, especially in reactions with lower RNA input. Recommended cycle numbers are provided on the next page.

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## PCR Cycle Number Recommendations

The recommended PCR cycle numbers in the table below are based on a 16 hour hybridization with quantification using an Agilent Bioanalyzer 2100 DNA 1000 assay. More sensitive quantitation methods may allow for fewer PCR cycles.

The relationship between number of input copies and RT-qPCR Ct value may vary, and must be empirically determined by the user. If RNA samples are unnormalized with highly variable Cts or unknown copy numbers, assume a copy number of at least 10,000 and employ a high sensitivity quantitation method (with proportionally fewer PCR cycles) to limit overamplification of higher input samples. In general, minimizing PCR cycles will produce more on-target and uniform coverage, so run as few cycles as necessary to achieve precise and reproducible quantitation.

# Probes	1,000 copies	10,000 copies	100,000 copies
128	31	28	25
256	30	27	24
512	29	26	23
1,024	28	25	22
2,048	27	24	21
4,096	26	23	20
8,192	25	22	19
16,384	24	21	18
32,768	23	20	17
65,536	22	19	16
131,072	21	18	15

**NOTE:** For pan-viral assays where only a subset of probes are expected to produce signal, cycles should be adjusted based on individual design size, rather than total probe count. Similarly, for single virus assays with ultra-high tiling density, cycle numbers may have to be increased as the relationship between probe count and cycle number may not be linear in these assays.

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## Best Practices

Although not always stated at all steps in this protocol, these best practices should be followed throughout to maximize performance.

## Samples

- RNA samples should be stored at -80 °C until use, and thawed on ice.
- All work surfaces and gloves should be sanitized with RNaseZap (or equivalent) prior to reaction setup.
- For most consistent performance, all samples in a batch, including control samples, should be from the same sample type and extracted with the same RNA extraction procedure.
- A no-RNA control is recommended.
- Upon thawing frozen samples, briefly vortex and spin down prior to use.

## Master Mixes

- Prepare master mixes in a PCR workstation.
- The PCR workstation should be UV-irradiated after each setup. If unsure, UV-irradiate the workstation before setting up a master mix. *Do not turn on the UV light when reagents are in the workstation.*
- Master mixes are prepared in 1.5 ml or 2 ml microfuge tubes. Briefly vortex to mix and spin down.
- All master mixes for the batch should be prepared in the same manner unless specified otherwise in this protocol.
- Master mixes can be warmed to room temperature immediately before and during use but should be stored at -20 °C when not actively in use.

## Reaction Setup

- **Set up RT-hybridization reactions on ice or a 96-well cold block.** Do not allow the RNA samples to warm to room temperature prior to beginning the incubation in the thermal cycler.
- **Set up subsequent reactions (fill-in and cleanup/PCR) at room temperature.** After beginning the RT-hybridization reaction, do not cool the reaction plate below room temperature at any time until a safe stopping point has been reached (after PCR is complete). This includes centrifugation steps - refer to the note in the Reaction Plates section below.

## Reaction Plates

- Always seal plates with microseal B film (clear adhesive) seals. Foil seals are not recommended for any step in this protocol. However, they can be used for plates that will be placed in the freezer for storage.
- Using a roller for microseal B film, apply firm pressure and seal over the tops of all wells. Visually inspect all wells, particularly the edges and corners of the plate, to confirm that the seal is in contact with the plate. If not, apply firm pressure and roll until the film is in contact with the plate.
- When removing plate seals, a heated plate sealer can be used if desired to briefly warm the seal and loosen the adhesive.
- **Pulse centrifuge** in an Eppendorf 5810 fitted with a swinging bucket plate rotor up to maximum rpm for a few seconds. Balance plates with a blank if spinning an odd number of plates.

**NOTE:** Many centrifuges have a cooling setting. Since reactions should not be cooled below room temperature at any point after beginning the RT and Probe Hybridization step, turn the cooling off and centrifuge at room temperature.

- After centrifugation, inspect the bottom of the plate to ensure the expected volume is present in every well.
- It is particularly important to spin down the Index Primer Plate(s) to ensure that the required volume can be easily pipetted from the bottom of the plate, and limit potential cross-contamination between adjacent wells. Wells may appear empty prior to centrifugation.

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## Workflow

	16 hr Hybridization	
	Hands-on Time	Turnaround Time
Reverse Transcription Probe Hybridization	15 min	17 hr 15 min
↓		
Fill-In	15 min	1 hr 15 min
↓		
Enzymatic Cleanup PCR	15 min	1 hr 15 min
↓		
Library Pooling Library Purification	30 min	30 min
	<b>75 min</b>	<b>20 hr 15 min</b>

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## Procedure

### 1. Reverse Transcription and Probe Hybridization

1.1. Set up the RTHybFill program on a thermal cycler:

Program and Steps	Temperature and Time (hh:mm:ss)
<b>RTHybFill</b>	<b>Heated lid 105 °C</b>
Reverse Transcription	25 °C 00:10:00, 50 °C 00:50:00
Denature	95 °C 00:01:00
Hybridize and Fill	55 °C 24:00:00
Hold	55 °C ∞

1.2. Prepare labware and reagents

- Label one 96-well PCR plate to be used for the capture reactions and place it on ice or on a cold block.
- Retrieve RNA samples from frozen storage.
- Retrieve RT Mix and Probe Mix from frozen storage.

1.3. Add sample RNA to reaction plate

- Per reaction, pipette **6 µl** of RNA into a well of the reaction plate. Adhere to guidelines described under RNA Input Recommendations.

1.4. Prepare RT-Hybridization master mix

- Once thawed, vortex and spin down the RT Mix and Probe Mix.
- Prepare the RT-Hybridization master mix as follows:

Component	Per reaction	96 reactions	384 reactions
Probe Mix	1.6 µl	176 µl	704 µl
RT Mix	0.4 µl	44 µl	176 µl
<b>TOTAL</b>	<b>2.0 µl</b>	<b>220 µl</b>	<b>880 µl</b>

**NOTE:** The 96 and 384 reaction volumes already include the required overage. The RT Mix is viscous; exercise care when pipetting.

- Vortex to mix, and spin down the tube.

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1.5. Add RT-Hybridization master mix to reactions

- Pipette **2 µl** of RT-Hybridization master mix into each sample well. The master mix is viscous; exercise care when pipetting.
- **Mix the contents of each well** by pipetting up and down 3 times, then seal the plate as described in Best Practices. Alternatively, seal and then pulse vortex the plate. Pulse centrifuge the plate briefly.
- The reactions should now be a homogenous **pale blue** color.

**NOTE:** *The plate will be incubated at a high temperature for an extended period of time. Improper sealing may result in significant volume loss due to evaporation.*

1.6. Perform Reverse Transcription and Probe Hybridization

- Place the reaction plate in the thermal cycler and run the RTHybFill program.
- Make a note of the thermal cycler start time.
- Allow the program to run for the desired hybridization time. An overnight hybridization (12 - 20 hours) is recommended.

## 2. Fill-In

Start this procedure roughly 20 minutes prior to the end of the hybridization to ensure that the Fill-in Mix is fully thawed and at room temperature when hybridization is completed.

**NOTE:** Do not remove the reaction plate from the thermal cycler until the Fill-in Mix is fully thawed. Correct timing is important to maximize result quality.

2.1. Retrieve Fill-in Mix from frozen storage. Allow to thaw and equilibrate to room temperature, then vortex and spin down.

2.2. Add Fill-in Mix to reactions

- Remove the reaction plate from the thermal cycler, but leave the RTHybFill program running. Close the lid of the thermal cycler so the temperature of the block does not fluctuate.

**NOTE:** Do not end the thermal cycler program. The thermal cycler should remain at 55 °C for the Fill-in reaction.

- Pulse centrifuge the plate briefly and carefully remove and discard the seal.
- Add **2 µl** of Fill-in Mix to each reaction well in the reaction plate.
- **Mix the contents of each well** by pipetting up and down 3 times, then seal the plate as described in Best Practices. Alternatively, seal and then pulse vortex the plate. Pulse centrifuge the plate briefly.
- The reactions should now be a homogenous **pale green** color.

2.3. Place the reaction plate back in the thermal cycler and allow the RTHybFill program to continue for 60 minutes.

**NOTE:** Record the time the reaction plate was returned to the thermal cycler; correct timing at this step is important to maximize result quality.

### 3. Cleanup and PCR

Start this procedure roughly 20 minutes prior to the end of the fill-in reaction.

**NOTE:** Correct timing is critical to maximize result quality. Ensure that master mixes and index primers are fully thawed before use.

3.1. Set up the CleanupPCR program on a thermal cycler:

Program and Steps	Temperature and Time (hh:mm:ss)
<b>CleanupPCR</b>	<b>Heated lid 105 °C</b>
Cleanup	45 °C 00:30:00
Denature	95 °C 00:03:00
PCR	N x [98 °C 00:00:15, 60 °C 00:00:15, 72 °C 00:01:00]
Hold	4 °C ∞

The number of PCR cycles to run (N) should be adjusted based on the amount of input DNA, the number of probes, the hybridization time, and the sensitivity of the quantification method. On-target yield (an amplicon of ≥250 bp) will change approximately linearly with respect to both input DNA and number of probes. A nonspecific amplicon of ~170-200 bp may also be observed, especially in reactions with lower DNA input. Refer to page 10 for PCR cycle recommendations.

3.2. Retrieve the following reagents from frozen storage:

- Cleanup Mix
- PCR Mix
- Index Primer Plate

**NOTE:** Always ensure that the correct amplification primers are being used, i.e. ML2100/ML2200 for IL probe designs, and ML2100-TN for TN probe designs. Using the incorrect primers will result in reaction failure.

3.3. Allow reagents to thaw and equilibrate to room temperature, then vortex and spin down.

**NOTE:** It is particularly important to spin down the Index Primer Plate(s) to ensure that the required volume can be easily pipetted from the bottom of the plate, and limit potential cross-contamination between adjacent wells. Wells may appear empty prior to centrifugation.

- 3.4. Prepare the **Cleanup and PCR master mix** in a microcentrifuge tube by combining kit components as follows:

Component	Per reaction	96 reactions	384 reactions
Cleanup Mix	2 µl	204 µl	816 µl
PCR Mix	12 µl	1224 µl	4896 µl
<b>TOTAL</b>	<b>14 µl</b>	<b>1428 µl</b>	<b>5712 µl</b>

**NOTE:** The 96 and 384 reaction volumes already include the required overage.

- Vortex to mix, and spin down the tube.
- 3.5. End the RTHybFill program and remove the reaction plate from the thermal cycler.
- 3.6. Pulse centrifuge the plate briefly and carefully remove and discard the seal. The volume of the capture reaction product at this step, net of evaporation, should be approximately **~7.6 µl**.
- 3.7. Set up Cleanup and PCR reactions:
- Add **14 µl** of **Cleanup and PCR master mix** to each reaction
  - Add **2.4 µl** of the appropriate **index primer** to each reaction.

**NOTE:** The total reaction volume in each well should be approximately 24 µl.

- **Mix the contents of each well** by pipetting up and down 3 times, then seal the plate as described in Best Practices. Alternatively, seal and then pulse vortex the plate. Pulse centrifuge the plate briefly.
  - The reactions should now be a homogenous **magenta color**.
- 3.8. Perform Cleanup and PCR by placing the plate in the thermal cycler, and run program CleanupPCR.
- NOTE:** This program may be run overnight, but the reactions should remain at 4 °C until no later than the following day.
- 3.9. After the CleanupPCR program completes, store the reaction plate at -20 °C or proceed directly to Step 4, Library Pooling.

## 4. Library Pooling

4.1. Remove the reaction plate from the thermal cycler (or -20 °C storage). If frozen, allow the plate to thaw and equilibrate to room temperature.

4.2. Spin down the reaction plate and carefully remove and discard the seal.

**CAUTION:** *This plate contains highly concentrated DNA that could contaminate subsequent experiments. Be very careful when handling this plate to avoid spilling or splashing.*

4.3. Prepare the library pool by transferring an equal volume of each reaction into a single microcentrifuge tube. A minimum of 10 µl per reaction is recommended, and the total pool volume should be at least 120 µl (e.g. transfer ≥15 µl per reaction for 8 reactions). Pooling fewer than 8 reactions is not recommended.

**NOTE:** *If input quantity and/or quality differs significantly between samples, bin samples with similar input into pools, or vary the volume of each library in a pool to be representative of the expected library yield. Failure to do so may result in significant over- or under-representation of libraries in the final sequencing results.*

4.4. Proceed directly to Step 5, Library Purification, or store the tube at -20 °C.

**NOTE:** *If library QC will include individual reactions and be performed immediately after purification, the reaction plate may be stored at room temperature temporarily. If not, seal the plate and store at -20 °C.*

## 5. Library Purification

- 5.1. Perform library purification using KAPA Pure Beads (or an equivalent product) on the pooled library, with a 0.75X bead ratio (i.e. 75 µl of beads per 100 µl of pooled library).

**NOTE:** The bead ratio may be adjusted depending on the amplicon size and the desired purification stringency and yield. A higher ratio (e.g. 1X) may increase overall yield but slightly reduce the fraction of on-target amplicons. Lower ratios may produce a higher fraction of on-target amplicons but reduce overall yield and sensitivity to deletion-containing amplicons that are smaller than expected - adding an additional cycle of PCR is recommended when using the minimum bead ratios specified below.

**Do not go below the minimum bead ratios recommended in the table below.**

Insert Size	Library Size	Standard Bead Ratio	Minimum Bead Ratio
75 bp	~260 bp	0.75X	0.75X
150 bp	~335 bp		0.67X
225 bp	~410 bp		0.60X

- 5.2. Elute the pooled, purified library DNA in 1X TE Buffer, using an elution volume 10-fold smaller than the original pooled library volume (i.e. for 100 µl of pooled library, elute in 10 µl of 1X TE Buffer).
- 5.3. The purified library is ready for quantification, dilution, and sequencing according to the sequencer manufacturer's instructions, but should be stored at -20 °C until use.

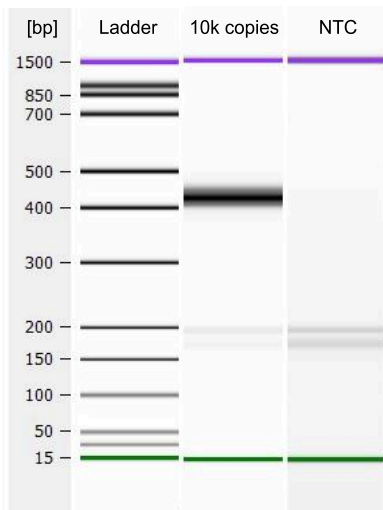
**NOTE:** Be sure to use the custom sequencing primers specified on page 5 when sequencing **IL design libraries** prepared using the Molecular Loop protocol.

## 6. Library Quantification/QC (Optional)

Quantification can be performed using electrophoresis- or fluorescence-based methods. An example of a purified library pool and no-template control analyzed using the Agilent Bioanalyzer 2100 DNA 1000 Assay is shown in Figure 1 below.

**NOTE:** When loading the chip, only go to the first stop of the pipette to prevent air bubbles from being introduced into the DNA 1000 chip. This achieves better accuracy. Add the DNA/ladder directly after adding the matrix.

One or more small peaks of ~170-200 bp may or may not be present; these likely represent self-ligated, unfilled circles. The true target peak should be  $\geq 250$  bp. If a no-RNA reaction is run individually, the self-circle peak(s) may be quite strong, whereas a true target peak should not be present.



**Figure 1.** Agilent Bioanalyzer 2100 DNA 1000 Assay results. Lane 1 contains the library pool; lane 2 contains a no-template control. The strong peak at ~410 bp represents the desired product for a 225 bp fill design. One or more small peaks of ~170-200 bp may or may not be present; these likely represent self-ligated, unfilled circles. The true target peak should be  $\geq 250$  bp. If the no-RNA reaction is run, the self-circle peak(s) may be quite strong, and a true target peak should not be present.

## Data Analysis

Targets are captured by probes containing synthetic sequences at their 5' and 3' ends. These synthetic sequences will appear at the 5' (and sometimes also the 3') end of each read generated by the NGS system. These synthetic sequences should be trimmed from the reads prior to alignment or genotype calling.

For detailed instructions, refer to the **Data Analysis User Guide**.