



Element
Biosciences

Cloudbreak™ Sequencing

User Guide

FOR USE WITH

AVITI™ System

AVITI™ System LT

AVITI24™ System

AVITI Operating Software v3.4 or later

Cloudbreak, Cloudbreak Freestyle™, and Cloudbreak UltraQ™ Sequencing Kits

ELEMENT BIOSCIENCES

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Overview

The Cloudbreak sequencing workflow uses a Cloudbreak Freestyle, Cloudbreak, or Cloudbreak UltraQ kit to sequence libraries on an AVITI System or an AVITI24 System.

- **Cloudbreak Freestyle**—Provides multiple read lengths and output options to meet a diversity of applications. Cloudbreak Freestyle kits enable direct loading of linear libraries without library conversion, including third-party libraries.
- **Cloudbreak**—Provides the same read length and output options as Cloudbreak Freestyle with potential requirements for library circularization.
- **Cloudbreak UltraQ**—Provides high-quality Q40 and Q50 data for highly sensitive assays.

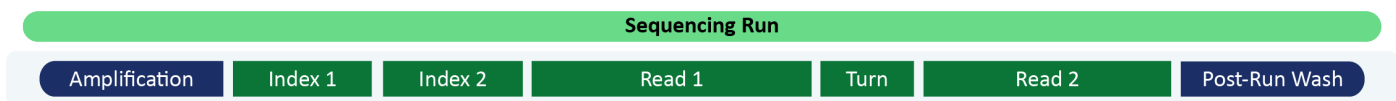
All Cloudbreak kits are designed for minimal waste and easy disposal. Reagent overage supports the extra cycles that index sequences and unique molecular identifiers (UMIs) need to identify samples with high confidence. For a list of kit configurations and catalog numbers, see [Cloudbreak Sequencing Kits on page 27](#).

Cloudbreak kits are compatible with a variety of library preparation workflows. For more information on compatibility, see the [Product Compatibility](#) page on the Element website.

Sequencing Run Stages

AVITI Operating Software (AVITI OS) generates a recipe based on the run parameters entered during run setup. The recipe governs each stage of a run. A run is complete when the recipe is executed and primary analysis is finished. The following stages comprise a sequencing run:

- **Amplification**—Hybridizes the library to the flow cell and performs amplification to form colonies, each containing multiple copies of the same sequence from the library.
- **Sequencing**—Performs each read in the run, including imaging and primary analysis.
- **Post-run wash**—Automatically flushes buffer from the sequencing cartridge through the fluidic system to remove salts and residual library.



Reads in a Sequencing Run

Up to four reads comprise a sequencing run: Index 1, Index 2, Read 1, and Read 2.

- **Index reads**—A run can include one, two, or no index reads.
 - » **Index 1** sequences the Index 1 sequence.
 - » **Index 2** sequences the Index 2 sequence.
 - » A dual-index run sequences Index 1 and Index 2.
- **Read 1 and Read 2**—All runs must have a Read 1.
 - » **Read 1** sequences the forward strand of the DNA insert.
 - » **Read 2** sequences the reverse strand.
 - » A paired-end run sequences Read 1 and Read 2, including a paired-end turn before Read 2 to generate the complementary strand.

Number of Cycles

Read length is the total number of cycles performed in a run. For bioinformatics purposes, adding one extra cycle to each read is recommended. The additional cycle improves the accuracy of the Q score for the 150th cycle. The software and chemistry used for the run prescribe a minimum number of cycles. Read 1 requires at least five cycles and at least 25 cycles to generate all run metrics. The maximum number of cycles depends on the kit:

- A 2 x 75 kit performs up to 184 cycles, supporting one 2 x 76 run with indexing and unique molecular identifiers (UMIs).
- A 2 x 150 kit performs up to 334 cycles, supporting one 2 x 151 run with indexing and UMIs.
- A 2 x 300 kit performs up to 634 cycles, supporting one 2 x 301 run with indexing and UMIs.

Run Times

Kit Output Configuration	Read Length	Run Time (approximate) ¹
High output	2 x 75	24 hours ²
	2 x 150	38 hours ²
	2 x 300	60 hours
Medium output	2 x 75	20 hours ²
	2 x 150	31 hours ²
	2 x 300	51 hours
Low output	2 x 75	17 hours ²
	2 x 150	27 hours ²

¹ Individually addressable lanes and custom recipes can extend run times.

² AVITI24 sequencing runs with AVITI OS v3.4 or later increases sequencing output by ~50%. As a result, run times can slightly increase up to 5%.

Read Counts and Output on AVITI System

Kit Configuration	Kit Size	Target Read Counts	Output (Gb)
Cloudbreak or Cloudbreak Freestyle High Output	2 x 75	1 billion	150
	2 x 150	1 billion	300
	2 x 300	300 million	180
Cloudbreak or Cloudbreak Freestyle Medium Output	2 x 75	500 million	75
	2 x 150	500 million	150
	2 x 300	100 million	60
Cloudbreak or Cloudbreak Freestyle Low Output	2 x 75*	100 million	15
	2 x 150	250 million	75
Cloudbreak UltraQ High Output	2 x 150	800 million	240

* Available as Cloudbreak Freestyle chemistry only

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Read Counts and Output on AVITI24 System

Read counts and output on the AVITI24 System vary depending on the version of AVITI OS you are using. AVITI24 Systems running AVITI OS v3.4 or later delivers up to 50% more output with 2 x 75 or 2 x 150 Cloudbreak or Cloudbreak Freestyle chemistry.

AVITI24 with AVITI OS v3.3 or earlier

Kit Configuration	Kit Size	Target Read Counts	Output (Gb)
Cloudbreak or Cloudbreak Freestyle High Output	2 x 75	1 billion	150
	2 x 150	1 billion	300
	2 x 300 ¹	300 million	180
Cloudbreak or Cloudbreak Freestyle Medium Output	2 x 75	500 million	75
	2 x 150	500 million	150
	2 x 300 ¹	100 million	60
Cloudbreak or Cloudbreak Freestyle Low Output	2 x 75 ²	100 million	15
	2 x 150	250 million	75
Cloudbreak UltraQ High Output	2 x 150 ¹	800 million	240

¹ For the Cloudbreak UltraQ kit and any 2 x 300 cycle kit, target read counts and output are unchanged based on the AVITI OS version.

² Available as Cloudbreak Freestyle chemistry only

AVITI24 with AVITI OS v3.4 or later

Kit Configuration	Kit Size	Target Read Counts	Output (Gb)
Cloudbreak or Cloudbreak Freestyle High Output	2 x 75	1.5 billion	225
	2 x 150	1.5 billion	450
	2 x 300 ¹	300 million	180
Cloudbreak or Cloudbreak Freestyle Medium Output	2 x 75	750 million	112.5
	2 x 150	750 million	225
	2 x 300 ¹	100 million	60
Cloudbreak or Cloudbreak Freestyle Low Output	2 x 75 ²	150 million	22.5
	2 x 150	375 million	112.5
Cloudbreak UltraQ High Output	2 x 150 ¹	800 million	240

¹ For the Cloudbreak UltraQ kit and any 2 x 300 cycle kit, target read counts and output are unchanged based on the AVITI OS version.

² Available as Cloudbreak Freestyle chemistry only

Library Considerations

Some libraries have special considerations for sequencing. Make sure to follow the applicable requirements for your library.

Low-Diversity Amplicon Libraries

For low-diversity, high-multiplex libraries, such as a 16S amplicon library, Element recommends that you enable the Low-Diversity High-Multiplex setting during run setup. This setting requires a library pool that meets the following requirements:

- Adept™ libraries or third-party libraries
- High plexity of ≥ 64 unique dual indexed (UDI) libraries
- A 1–5% spike-in of PhiX Control Library



CAUTION

Exceeding a 5% spike-in can reduce the index diversity of the pool, leading to a reduction in quality.

Bead-Based Normalization

PCR is required when sequencing a library pool that has undergone the bead-based normalization protocol. Before diluting to the target loading concentration, use amplification and Qubit kits to amplify and quantify the library pool.

- For Cloudbreak chemistry with Adept libraries, use the Adept Rapid PCR-Plus Kit for amplification.
- For Cloudbreak Freestyle chemistry with third-party libraries, use the KAPA HiFi HotStart Library Amplification Kit with Primer Mix. Follow manufacturer instructions.

Short Insert or Long Insert Libraries

Sequencing short insert or long insert libraries require that you specify the preloaded custom recipe during run setup.

Short insert libraries

- If you are using Cloudbreak Freestyle chemistry and libraries with < 100 bp inserts, the short insert recipe is required.
- For libraries with 100–300 bp inserts, the short insert recipe is recommended.
- The short insert recipe is compatible with 2 x 75 and 2 x 150 kits.
- For more information, see [Sequencing Short Insert Libraries with Cloudbreak Freestyle](#).

Long insert libraries

- If you are using Cloudbreak or Cloudbreak Freestyle chemistry and libraries with > 750 bp inserts, the long insert recipe is required.
- The long insert recipe is compatible with 2 x 75 and 2 x 150 kits. Using a 2 x 300 kit accounts for long insert conditions.

Input Recommendations

The recommended input for sequencing is ≥ 1 nM library. The input library is normalized to 1 nM, denatured into single strands, and diluted to the target loading concentration. When starting with a 0.2–1 nM library, the library is denatured and diluted but not normalized. Library pools that start at < 0.2 nM are not supported.

PhiX Control Library Spike-In

For most applications, Element recommends a spike-in of PhiX Control Library. The following recommendations for spike-in percentages optimize the benefits of PhiX Control Library for specific experiments.

Experiment	Spike-In (%)
QC and error rate reporting	> 2
Low-complexity indexing (≤ 2 -plex)	> 2
Libraries that use Low-Diversity High-Multiplex setting	1–5
Other low-diversity libraries*	≥ 5

* For Adept and third-party workflows, the first four cycles of Read 1 require high diversity. Index 1 includes high diversity for Elevate™ workflows.

Custom Primers

You can sequence any combination of I1, I2, R1, and R2 custom primers for third-party libraries with Cloudbreak Freestyle chemistry and Adept libraries with Cloudbreak chemistry. The custom primers must be HPLC-purified.

The custom primers are added to the cartridge using these two applicable methods:

- **Spike-in**—Spike-in custom primers into the applicable wells of the Cloudbreak Freestyle cartridge or the Adept Primer Set Cloudbreak tubes.
- **Replacement**—Replace the primers in the cartridge with buffer tubes from the Custom Primer Set Cloudbreak Freestyle or Adept Custom Primer Set Cloudbreak and add custom primers.

Sequencing Primer Compatibility

- For Cloudbreak Freestyle chemistry, Element oligonucleotides include sequencing primers that are compatible with standard Nextera, TruSeq, TruSeq small RNA, and Elevate libraries.
- For original Cloudbreak chemistry with Adept libraries, replace primers provided in the cartridge with primers provided in the Adept Primer Set for sequencing standard Nextera and TruSeq libraries.
- For original Cloudbreak chemistry with Elevate libraries, the primers provided in the cartridge are compatible with standard Nextera, TruSeq, and Elevate libraries.
- Libraries with sequencing primer binding sites that do not meet these requirements *must* use custom primers.

Custom primers require special consideration and planning. To determine if your library requires custom primers and ensure a run with custom primers meets specifications, contact Element Technical Support early in experiment planning. For more information on Cloudbreak Freestyle custom primer recommendations, see [Cloudbreak Freestyle Compatibility with Third-Party Libraries](#).

Loading Concentration

Use the following recommendations as a starting point to determine your optimal loading concentration. Loading concentration recommendations depend on your library prep workflow, the kit chemistry, output level, and kit size.

- Polony counts increase as the loading concentration increases. Lower polony counts promote higher data quality but lower the amount of data output. Some libraries require a higher or lower concentration than the indicated ranges.
- Average library size is the full length of the library, including DNA insert and adapters. Pooled libraries must contain similar size distributions.
- Sequencing runs on the AVITI24 System running AVITI OS v3.4 or later require a slightly higher loading concentration with 2 x 75 or 2 x 150 Cloudbreak or Cloudbreak Freestyle chemistry.

AVITI or AVITI24 with AVITI OS v3.3 or earlier

Cloudbreak Chemistry, 2 x 75 and 2 x 150 Kits

Average Library Size	Adept v1.1	Adept Rapid PCR-Plus	Elevate PCR-Free	Elevate PCR-Plus
Small (250–450 bp)	4–6 pM	10–14 pM	6–10 pM	8–11 pM
Medium (450–700 bp)	6–10 pM	10–14 pM	7–11 pM	9–12 pM
Large (≥ 700 bp)	10–14 pM	10–14 pM	7–11 pM	9–12 pM

Cloudbreak Freestyle Chemistry, 2 x 75 and 2 x 150 Kits

Average Library Size	Elevate PCR-Free	Elevate PCR-Plus	Third Party PCR-Free	Third Party PCR-Plus
Small (250–450 bp)	5–9 pM	7–10 pM	6–9 pM	7–10 pM
Medium (450–700 bp)	6–10 pM	8–11 pM	7–10 pM	9–12 pM
Large (≥ 700 bp)	6–10 pM	8–11 pM	7–10 pM	9–12 pM

Cloudbreak Chemistry, 2 x 300 Kits ¹

Average Library Size	Adept v1.1	Adept Rapid PCR-Plus	Elevate PCR-Free	Elevate PCR-Plus
Medium (450–700 bp)	4–6 pM	5–8 pM	3–5 pM	4–6 pM
Large (≥ 700 bp)	6–8 pM	5–8 pM	3–5 pM	4–6 pM

Cloudbreak Freestyle Chemistry, 2 x 300 Kits ¹

Average Library Size	Elevate PCR-Free	Elevate PCR-Plus	Third Party PCR-Free	Third Party PCR-Plus
Medium (450–700 bp)	3–5 pM	4–8 pM	4–6 pM	6–9 pM
Large (≥ 700 bp)	3–5 pM	4–8 pM	4–6 pM	8–12 pM

Cloudbreak UltraQ Chemistry, 2 x 150 Kit ¹

Average Library Size	Elevate Libraries
450–550 bp	5–6 pM

¹ For the Cloudbreak UltraQ kit and any 2 x 300 cycle kit, loading concentration is unchanged based on the AVITI OS version.

AVITI24 with AVITI OS v3.4 or later

Cloudbreak Chemistry, 2 x 75 and 2 x 150 Kits ²

Average Library Size	Adept v1.1	Adept Rapid PCR-Plus	Elevate PCR-Free	Elevate PCR-Plus
Small (250–450 bp)	5–7.5 pM	12.5–17.5 pM	7.5–12.5 pM	10–13.75 pM
Medium (450–700 bp)	7.5–12.5 pM	12.5–17.5 pM	8.75–13.75 pM	11.25–15 pM
Large (≥ 700 bp)	12.5–17.5 pM	12.5–17.5 pM	8.75–13.75 pM	11.25–15 pM

Cloudbreak Freestyle Chemistry, 2 x 75 and 2 x 150 Kits ²

Average Library Size	Elevate PCR-Free	Elevate PCR-Plus	Third Party PCR-Free	Third Party PCR-Plus
Small (250–450 bp)	6.25–11.25 pM	8.75–12.5 pM	7.5–11.25 pM	8.75–12.5 pM
Medium (450–700 bp)	7.5–12.5 pM	10–13.75 pM	8.75–12.5 pM	11.25–15 pM
Large (≥ 700 bp)	7.5–12.5 pM	10–13.75 pM	8.75–12.5 pM	11.25–15 pM

Cloudbreak Chemistry, 2 x 300 Kits ¹

Average Library Size	Adept v1.1	Adept Rapid PCR-Plus	Elevate PCR-Free	Elevate PCR-Plus
Medium (450–700 bp)	4–6 pM	5–8 pM	3–5 pM	4–6 pM
Large (≥ 700 bp)	6–8 pM	5–8 pM	3–5 pM	4–6 pM

Cloudbreak Freestyle Chemistry, 2 x 300 Kits ¹

Average Library Size	Elevate PCR-Free	Elevate PCR-Plus	Third Party PCR-Free	Third Party PCR-Plus
Medium (450–700 bp)	3–5 pM	4–8 pM	4–6 pM	6–9 pM
Large (≥ 700 bp)	3–5 pM	4–8 pM	4–6 pM	8–12 pM

Cloudbreak UltraQ Chemistry, 2 x 150 Kit ¹

Average Library Size	Elevate Libraries
450–550 bp	5–6 pM

¹ For the Cloudbreak UltraQ kit and any 2 x 300 cycle kit, loading concentration is unchanged based on the AVITI OS version.

² Recommended loading concentration is increased 25% to account for higher polony density on the AVITI24 System.

Cloudbreak Sequencing User Guide

Cloudbreak Workflow Summary

Performing a Cloudbreak sequencing run includes steps to prepare reagents and dilute the library to the appropriate volume and concentration for sequencing. For more information, see [Loading Concentration on page 9](#).

Prepare for the Run

- 1 Thaw the reagent cartridge
- 2 Add primers, if applicable for Adept libraries
- 3 Dilute and denature libraries
- 4 Prepare custom primers (optional)

Set Up the Run

- 5 Define run parameters
- 6 Add custom primers (optional)
- 7 Add library to cartridge
- 8 Load the reagent cartridge and buffer bottle
- 9 Empty waste and prime reagents
- 10 Load the flow cell
- 11 Review, start, and monitor the run

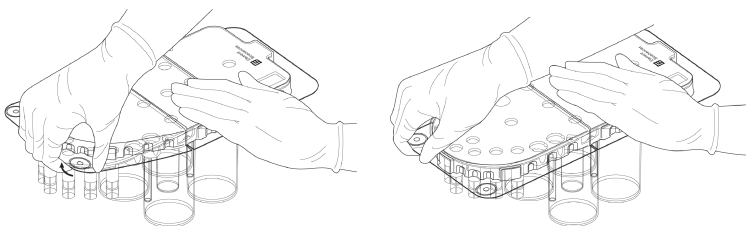
- Off instrument
- On instrument

Run Preparation

Run preparation includes thawing the sequencing cartridge and adding appropriate primers, if applicable. The subsequent library dilution procedure includes the option to store a normalized library. If you intend to store a library, do not prepare the cartridge until you are ready to sequence. Prepare the cartridge within a day of sequencing.

Thaw Reagents

1. If the cartridge includes a shipping cover, remove the shipping cover:
 - a. While supporting the cartridge, lift the removal tab at the left corner until it releases from the cartridge.



- b. Moving across the front edge of the shipping cover, repeatedly lift the edge until the cover is fully released.
 - c. Pull to remove the remainder of the shipping cover from the cartridge.
2. Thaw the sequencing cartridge. Protect the cartridge from light until loading onto the instrument.

Cartridge	Room Temperature Water Bath	Refrigerator
2 x 75	1.5 hours	8 hours
2 x 150, 2 x 300	2.5 hours	24 hours

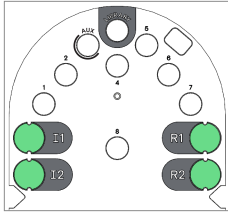
3. Make sure reagents are *fully* thawed. Inspect each well as reagents thaw at varying rates.
4. If any ice remains, continue thawing.
5. Set aside the thawed cartridge at room temperature and initiate the run within 2 hours. If you do not immediately initiate the run, place the thawed cartridge at 2°C to 8°C. You can store overnight at 2°C to 8°C for use the next day.

Add Primer Tubes

1. If you are using a Cloudbreak Freestyle sequencing kit without custom primers *or* you are sequencing Elevate libraries, skip the following steps and proceed to [Dilute Library and Custom Primers on page 13](#).
2. Remove applicable primer set from -25°C to -15°C storage.

Chemistry and Library	Primer Strategy	Primer Set
Cloudbreak, Adept	No custom primers	Adept Primer Set Cloudbreak
	Custom primers (spike-in method)	Adept Primer Set Cloudbreak
	Custom primers (replacement method)	Adept Custom Primer Set Cloudbreak
Cloudbreak Freestyle, Third Party	Custom primers (replacement method)	Custom Primer Set Cloudbreak Freestyle
	Custom primers (spike-in method)	Not applicable

3. To replace the primer tubes in the cartridge with primer set, twist the primer tubes in wells labeled I1, I2, R1, and R2 toward the left to unlock.



4. Remove the primer tubes from the cartridge and discard per the SDS.
If you have trouble removing the tubes, peel the labels off and twist the tubes as you push upwards.
5. Insert the tubes from the primer set into the vacated wells. Match the abbreviation on the tube label to the well label.
6. Twist each tube right until it locks into place.

Dilute Library and Custom Primers

To prepare for sequencing, the input library is normalized to 1 nM, denatured with NaOH, neutralized, and further diluted. The dilution procedures prepare 1.4 ml diluted library at the target loading concentration with an optional spike-in. Custom primers are diluted as applicable.

If you are using the Individually Addressable Lanes add-on, follow the applicable procedures for both libraries. Both libraries use the same denature and dilution methods, resulting in a total volume of 1.4 ml for each library.

Gather the following consumables:

- 0.2 M Tris-HCl buffer, pH 7.0
- 1 N NaOH
- 2 ml DNA LoBind tubes (4–7)
- 10 mM Tris-HCl, pH 8.0 with 0.1 mM EDTA (low TE buffer)
- Nuclease-free water

Prepare the Library

For bead-normalized libraries, perform amplification and quantification before proceeding. See [Bead-Based Normalization on page 7](#).

1. Remove the following components from -25°C to -15°C storage and thaw on ice:
 - » Library Loading Buffer
 - » Experimental library
 - » [Optional] PhiX Control Library
2. Pulse vortex the thawed libraries and briefly centrifuge.
3. If the experimental library is ≥ 1 nM, normalize:
 - a. In a new DNA LoBind tube, use low TE buffer to dilute the library to 1 nM.
 - b. Proceed immediately or cap the tube, store the 1 nM library at -25°C to -15°C.

Calculate Library Loading Concentration

Calculate the experimental library and PhiX Control Library loading concentrations and volumes for sequencing using a total loading volume of 1.4 ml. You will need the target loading concentration, relative amounts of each library, and spike-in percentage of PhiX Control Library to perform the calculations.

NOTE

The experimental and control library concentrations do not need to match.

1. Calculate the loading concentration of each library based on the target loading concentration as follows:

Library	Calculation	Resulting Loading Concentration (pM)
Experimental	Target loading concentration in pM * (100% – spike-in %)	_____
Control	Target loading concentration in pM * spike-in %	_____

2. Calculate the volume for each library based on the calculated loading concentration, starting library concentration, and a 1.4 ml loading volume as follows:

Library	Calculation	Resulting Loading Volume (µl)
Experimental	(Library loading concentration in pM * 1400 µl) / library starting concentration in pM	_____
Control	(Control library loading concentration in pM * 1400 µl) / control library starting concentration in pM	_____

NOTE

If the volume calculated for the control library is < 1 µl, dilute control library in low TE buffer to use a volume ≥ 1 µl for accurate pipetting.

3. Record the total volume of diluted experimental and control library in µl.

Example Calculation

An example calculation using a target loading concentration of 9 pM with a 2% spike-in of PhiX Control Library and a starting experimental library concentration of 1 nM (1000 pM) is shown in the following table. The PhiX Control Library was diluted and used at 100 pM concentration to allow for ≥ 1 µl volume for accurate pipetting.

Library	Loading Concentration (pM)	Loading Volume (µl)
Experimental	9 * 98% = 8.82 pM	(8.82 * 1400)/1000 = 12.3 µl
Control	9 * 2% = 0.18 pM	(0.18 * 1400)/100 = 2.5 µl

Denature the Library with NaOH

This procedure uses equal volumes of library, 0.2 N NaOH, and 0.2 M Tris-HCl buffer, pH 7.0. Therefore, the volume recorded during the calculations step is used in two subsequent steps

1. Combine the following reagents to prepare 0.2 N NaOH. Use 0.2 N NaOH within the day and discard.

Reagent	Volume
1 N NaOH	20 µl
Nuclease-free water	80 µl
Total	100 µl

2. In a new DNA LoBind tube, combine the experimental and control library volumes calculated in step 3 of the calculations step.
3. Add an equal volume of freshly prepared 0.2 N NaOH.
4. Vortex the tube to mix and briefly centrifuge.
5. Incubate the tube at room temperature for 5 minutes to denature the library.
6. Vortex the tube to mix and briefly centrifuge.
7. Add 0.2 M Tris-HCl buffer, pH 7.0 at an equal volume of 0.2 N NaOH to neutralize the reaction.
8. Vortex the tube to mix and briefly centrifuge.

—The library is denatured, neutralized, and at 1/3 the input concentration in 3x input volume.—

9. Add a sufficient volume of Library Loading Buffer to the combined library volume to reach a total volume of 1.4 ml:

$$\text{buffer volume in } \mu\text{l} = 1400 \mu\text{l} - 3 * \text{library volume in } \mu\text{l}$$

10. Vortex the tube to mix and briefly centrifuge.
11. Place the diluted sequencing library on ice. Use within the day.

Prepare Custom Primers

1. If you are not using custom primers, skip the following steps and proceed to [Run Setup on page 16](#).
2. In a new DNA LoBind tube, prepare each applicable custom primer using low TE buffer:

Custom Primer	Volume	Concentration
Index 1	19 μl	100 μM
Index 2	19 μl	100 μM
Read 1	32.4 μl	100 μM
Read 2	19 μl	100 μM

3. Set aside the 100 μM custom primers on ice. Use within the same day.

Run Setup

Run setup for sequencing prompts you to define run parameters, load sequencing consumables, and empty the waste bottle. Before initiating a run, review the overview, software, troubleshooting, and safety information in the user guide for your instrument.

Initiate a Sequencing Run

1. Gather the following materials:
 - » Buffer bottle
 - » Cartridge
 - » Cartridge basket
 - » Towel or wipe
 - » Used flow cell

—A used flow cell might already be present on the instrument.—
2. If applicable, stage run manifests for import:
 - » If setting up the run manually, save the manifest on a USB and connect the USB drive to an instrument USB port.
 - » Alternatively, you can save the manifest to the specified SMB storage connection.
 - » If you planned the run in ElemBio Cloud, upload the manifest to the planned run.
3. On the Home screen, select **New Run**.
4. If AVITI OS prompts that the flow cell is missing, load a **used** flow cell:
 - a. Select **Open Nest**.
 - b. Place the used flow cell onto the nest and close the lid.
 - c. Select **Close Nest**.
5. Select **Sequencing**.
6. Select the side for sequencing:
 - » **Side A**—Set up a run on side A.
 - » **Both**—Set up runs on sides A and B.
 - » **Side B**—Set up a run on side B.
7. For chemistry type, select **Cloudbreak**, and then select **Next**.
8. Proceed as follows:
 - » For a planned run created in ElemBio Cloud, proceed to [Select a Planned Run](#).
 - » For a manual run, proceed to [Define Manual Run Parameters on page 17](#).

Select a Planned Run

1. Select **Planned Run**.

AVITI OS displays a list of compatible planned runs for the instrument and run type. For information on planned run compatibility, see [Run Planning](#) in the [Online Help](#).
2. Select the run you want to use from the list of planned runs.
3. Review the run parameter fields to make sure they are correct.

If you need to edit a planned run, modify it in ElemBio Cloud. See [Run Planning](#) in the [Online Help](#).
4. In the Storage drop-down menu, select the storage connection for the run.

5. Select **Next** to proceed to the Prepare Reagents or the Run Side B screen.
 - » After you proceed, the selected planned run becomes unavailable for other connected instruments.
 - » If you exit run setup before priming, the run returns to the list of available planned runs.
6. If applicable, repeat steps 2–5 to set up a dual start run with a second planned run.
7. Proceed to [Inspect and Mix Reagents on page 18](#).

Define Manual Run Parameters

1. Make sure **Manual Run** is selected for the type of run.
2. In the Run Name field, enter a unique name to identify the run.
 - The field accepts 1–64 alphanumeric characters, hyphens, and underscores.—
3. If applicable, select **Browse** and import the run manifest.
4. [Optional] In the Description field, enter a description that represents the run.
 - The field accepts ≤ 500 alphanumeric characters, hyphens, underscores, spaces, and periods.—
5. In the Storage drop-down menu, select a storage location:
 - » To output run data to the default storage location, leave the default selection.
 - » To override the default storage location for the current run, select a storage connection.
6. Select a Library Type:
 - » **Elevate**—Sequence libraries prepared with Elevate indexes and adapters.
 - » **Adept**—Sequence libraries prepared with the Adept Workflow. Only compatible with Cloudbreak sequencing kits.
 - » **Third Party**—Sequence libraries prepared with a third-party workflow. Only compatible with Cloudbreak Freestyle sequencing kits.
7. If applicable, select a Library Structure:
 - » **Circular**—Sequence libraries that complete circularization before loading.
 - » **Linear**—Sequence libraries prepared for on-instrument circularization.
8. In the Sequencing Kit drop-down menu, select the kit you are using. For information on kit compatibility, see the [Product Compatibility](#) page on the Element website.
 - The kits listed depend on compatibility with the instrument type, and the selected library type and library structure.—
9. If you are using the Adept or Third Party library type, select a Low-Diversity High-Multiplex option.
 - » **Yes**—Sequence low-diversity high-multiplex libraries. This option requires at least 4 cycles for Index 1.
 - » **No**—Sequence other libraries.
10. If you are using the Individually Addressable Lanes add-on and a compatible sequencing kit, select the number of library pools.
11. In the Cycles fields, enter the number of cycles to perform in each read.
 - » Do not exceed the maximum number of cycles for the sequencing kit. See [Number of Cycles on page 5](#).
 - » Add one cycle to the desired number of Read 1 and Read 2 cycles. For example, enter **151** in the Read 1 field to perform 150 cycles in Read 1.
 - » To skip a read, enter **0**.
 - » For Elevate libraries, use the default Index cycle settings and avoid changing the Index cycle settings. Sequencing 12 cycles for Index 1 enables the system to read the full-length 9 bp index barcode and the 3 random bases (N) in front of the barcode.
 - » See the following table for minimum and default cycle values. Aside from the minimum cycle limitations, AVITI OS lets you distribute the available cycles among reads as necessary. However, performance specifications for each kit are based on sequencing Element control libraries using default read length settings specific to the kit.

Library Type	Kit Size	Minimum Values				Default Values			
		Index 1	Index 2	Read 1	Read 2	Index 1	Index 2	Read 1	Read 2
Adept or Third Party	2 x 75	0	0	5	0	Blank	Blank	76	76
	2 x 150	0	0	5	0	Blank	Blank	151	151
	2 x 300	0	0	5	0	Blank	Blank	301	301
Elevate	2 x 75	4	0	5	0	12	9	76	76
	2 x 150	4	0	5	0	12	9	151	151
	2 x 300	4	0	5	0	12	9	301	301

- If you are using the Advanced Run Settings, select **Advanced Settings** and proceed to [Configure Advanced Run Settings](#).
- Select **Next** to proceed to the Run Side B or Prepare Reagents screen.
- If applicable, repeat steps **2–13** to set up a dual start run.

Configure Advanced Run Settings

Use Advanced Run Settings to modify primary analysis and recipe configurations for a run. Available settings depend on kit compatibility. Some settings require the activation of an add-on. For more information, see the Advanced Run Settings and Add-On information in the user guide for your instrument.

- If you are using the Polony Density setting, select a Polony Density option.
 - » **Standard**—Uses the standard read output.
 - » **High Density**—Increases the read output.
- If you are using the Filter Mask setting, enter a base mask to use for filtering.
 - » Use the base mask format. For more information, see [Base Masks](#) in the [Online Help](#).
 - » If you do not use the Filter Mask setting, the default filter mask is R1 : Y15N*–R2 : Y15N*.
- If you are using the Custom Recipes setting, import the custom recipe file from preloaded recipes or a USB drive:
 - Select **Browse**.
 - Select **Element Recipes** for preloaded recipes or **USB** to upload from a connected USB drive.
 - Select the recipe file, and then select **Open**.
- If you are using the PMG Shift setting, enter the number of cycles to skip.

You can skip up to 20 cycles. Any skipped cycles reduces the maximum cycles AVITI OS allows for a run.
- Select **Next** to proceed.

Inspect and Mix Reagents

- Inspect each cartridge well to make sure reagents are fully thawed.
- Make sure the cartridge contains the appropriate primers.
- Make sure the tubes in the I1, I2, R1, and R2 wells are secure. If necessary, twist each tube to the right.
- Gently invert the cartridge **10 times** to mix reagents.



CAUTION

Inadequately mixed reagents can cause run failure.

- Tap the cartridge base on the benchtop to remove any large droplets from the tube tops.
- Inspect the small tubes to make sure reagents are settled at the bottom.
- Place the cartridge into a clean cartridge basket and lock the clips. Wipe any excess moisture.

Add Custom Primers to the Cartridge

- If you are not using custom primers, skip the following steps and proceed to [Add Library to the Cartridge](#).
- Using a new 1 ml pipette tip, pierce the center of the applicable I1, I2, R1, and R2 wells to create one hole. Push the foil to the edges.
- Discard the pipette tip.
- Add the applicable volume of 100 μM custom primer to each pierced well.

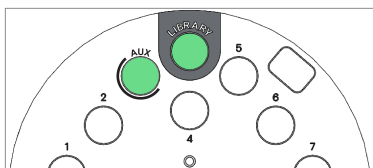
Custom Primer	Volume	Well
Index 1	19 μl	I1
Index 2	19 μl	I2
Read 1	32.4 μl	R1
Read 2	19 μl	R2

—The final concentration of each custom primer is 1 μM .—

- Pipette the content of each tube 15 times to mix. Avoid losing existing primer volume.

Add Library to the Cartridge

- Using a new 1 ml pipette tip, pierce the center of the Library well to create one hole. Push the foil to the edges.



- Discard the pipette tip.
- Briefly centrifuge the diluted sequencing library to remove bubbles and foam from the tube lid.
- Transfer the entire volume of diluted sequencing library to the Library well, dispensing along the well wall.
 - » Avoid aspirating any foam or dispensing air.
 - » Do not allow the library to contact the foil.
 - » Make sure the tube contains ≥ 1.3 ml diluted sequencing library.
- If you are using the Individually Addressable Lanes add-on, repeat steps 1–4 with the AUX well and the second library.

—The library for the AUX well contains the samples for Lane 2 in the run manifest.—



CAUTION

Transferring a library to the AUX well of an incompatible cartridge damages the library and the cartridge. For more information on Individually Addressable Lanes add-on compatibility, see the user guide for your instrument.

- Inspect the Library well through the window at the front of the basket.
 - » Make sure the library is free of foam and that bubbles are minimal.
 - » If an air gap appears below the surface, use a new pipette tip to remove it.
- If the cartridge include shipping locks, twist each shipping lock left to unlock and remove them from the cartridge lid.


Confirm Reagent Preparation

1. If you selected Adept, select the **Swap primer tubes** checkbox to confirm that the I1, I2, R1, and R2 wells contain Adept primers or custom primers.
2. Select the **Invert cartridge** checkbox to confirm that reagents are mixed.
3. Select the **Insert into basket** checkbox to confirm that the cartridge is in the cartridge basket.
4. Select any load library checkboxes to confirm that the cartridge contains diluted library.
5. Select **Next** to proceed to the Load Reagents screen.

Load Reagents and Buffer

1. Open the reagent bay door.
2. Remove any materials from the reagent bay and set aside.
3. Slide the basket containing the thawed cartridge into the reagent bay until it stops.
4. Support the buffer bottle with both hands and slide it into the reagent bay until it stops.
5. Close the reagent bay door, and then select **Next** to proceed.

Empty Waste

1. Open the waste bay door.
2. Unscrew the transport cap from the cap holder above the waste bay.
3. Remove the waste bottle from the waste bay and close the transport cap.
 **CAUTION** Waste bottle contents are considered hazardous. Dispose of waste according to local, state, and regional laws and regulations.
4. [Optional] Insert a funnel into a waste receptacle. Make sure the funnel is secure.
5. Open the transport cap and the vent cap.
6. Support the waste bottle with both hands and empty the waste:
 - a. Position the bottle over the funnel or waste receptacle.
 - If you inserted a funnel, align the handle to the inner edge of the funnel.
 - If you did not insert a funnel, center the handle over the waste receptacle.
 - b. Tip the bottle forward and drain. Invert the bottle and shake to expel all droplets.
 - c. If necessary, wipe liquid off the bottle.
7. Close the vent cap and return the empty waste bottle to the waste bay.
8. Screw the transport cap onto the cap holder and close the waste bay door.

Prime Reagents

1. Bring a new Cloudbreak flow cell to room temperature:
 - a. Remove a flow cell pouch from 2°C to 8°C storage. **Do not open the pouch.**
 - b. Set aside the pouch for at least 5 minutes.

NOTE

Before priming, if you need to discard the run setup, remove the cartridge, correct the run setup as needed, and reload the same cartridge. Proceed to the priming step.

2. Select **Next** to *automatically* start priming.
Priming takes ~ 5 minutes or up to 8 minutes at high elevations.
3. When priming is complete, select **Next** to proceed to the Load Flow Cell screen.
AVITI OS moves the nest forward and opens the nest bay door. A brief delay is normal.

NOTE

After priming and prior to loading a new flow cell, if you need to discard the run setup, remove the cartridge, correct the run setup as needed, and reload the same cartridge. Proceed to loading the flow cell.

Load the Flow Cell

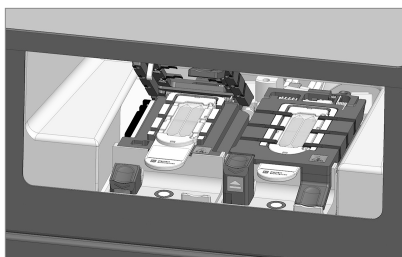
1. Make sure the nest status light is blue.
2. To open the lid, fully press down on the button to the left of the nest until it reaches the bottom. Failure to fully press down on the button can cause errors when closing the lid or aligning the flow cell.
3. Remove the used flow cell from the nest. Discard or store at room temperature for use with priming or washes.
4. Unpackage the Cloudbreak flow cell before loading onto the nest. Ensure the flow cell is used within 2 hours.
Handle the flow cell by the gripper only. If any debris contacts the flow cell surface during or after package opening, gently wipe the flow cell surface with an ethanol wipe and then a lens wipe.



CAUTION

Touching the glass can introduce debris, smudges, and scratches, compromising data quality.

5. Face the label up and place the flow cell over the three registration pins on the nest.



6. Lower the tab on the right side of the lid until the lid snaps into place.
7. Select **Close Nest** to close the nest bay door and retract the stage.
8. Select **Next** to proceed to the Run Summary screen.

Review and Start the Run

1. On the Details page, review the run parameters:

Parameter	Description
Library	The workflow that prepared the libraries and the library type
Sequencing Kit	The size and version of the sequencing kit
Storage	The location where sequencing output is stored
Manifest	The file name of the uploaded run manifest, if applicable
Cycles	The number of cycles in each read
Description	An optional description of the run
Advanced	If applicable, the advanced run settings for the run

NOTE

If you notice any errors in run set up, refer to the guidance in [Knowledge Base](#) to discard the run, skip the priming step, and restart the run setup process.

- Review the flow cell, cartridge, and buffer bottle information:

Field	Description
Lot Number	The number assigned to the batch the consumable was manufactured with
Expires on	The year, month, and date that the consumable expires
Serial Number	The unique identifier or all zeros indicating an unscanned barcode
Part Number	The Element-assigned identifier for the consumable

- Select **Run** to start sequencing.
- [Optional] If you imported run manifests from a USB drive, disconnect the USB drive:
 - In the taskbar, select **USB Drive**, and then select **Eject**.
 - Detach the USB drive from the instrument.
- Process the materials removed from the reagent bay:
 - » For a used cartridge and buffer bottle, follow the instructions in [Discard the Cartridge and Bottle on page 24](#).
 - » For a wash tray, follow the guidelines in the user guide for your instrument. Residual liquid in the wash tray is normal.

Monitor Run Metrics

- If necessary, select **Details** to open run details.
- Monitor run metrics as they appear onscreen. AVITI OS indicates the expected cycle that metrics appear. —The expected cycles are approximate, and all metrics are estimates. Bases2Fastq generates the final metrics.—
- Continue monitoring the run as AVITI OS refreshes the metrics.
 - » Each cycle refreshes the Q scores, error rates, base compositions, and index metrics.
 - » If you are using the Individually Addressable Lanes add-on, AVITI OS displays metrics for each library pool.
 - » AVITI OS refreshes the yield and reads metrics after cycle 15 of Read 2:
 - If Read 2 contains no cycles, metrics refresh after cycle 15 of Read 1.
 - If Read 1 or Read 2 contain fewer than 15 cycles, metrics refresh when the last cycle begins.
- When the run is complete, leave all materials on the instrument.
 - » To return to the Details view, select **Overview**.
 - » To access run data, go to your storage location.

Initiate Flexible Start

Flexible start provides the option to start a run or recovery wash while another run is in progress. AVITI OS safely pauses the run on the adjacent side.

- On the Home screen, select **New Run**.
- When prompted to request flexible start and pause the active run, select **New Run**.

Step	Estimated Wait Time to Pause*
Amplification	2 hours
Index 1, Index 2, Read 1, or Read 2	A few minutes
Turn	30 minutes
Wash	1 hour

* Estimates are for 2 x 75 and 2 x 150 runs with Cloudbreak chemistry. For 2 x 300 runs, the wait time for a pause at the amplification step can exceed 2 hours.

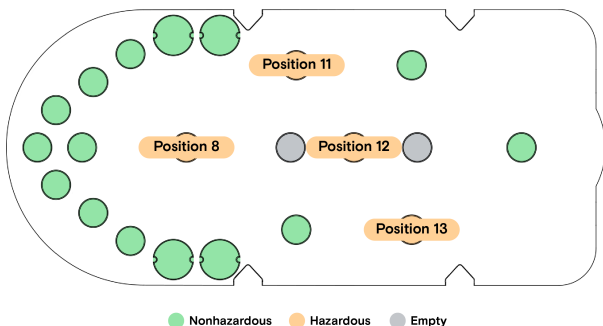
- Wait for the run to pause.

- » To cancel flexible start while waiting, select **Cancel Request**.
 - » Contact Element Technical Support if the wait time exceeds 5 hours at the amplification step or 1.5 hours at any other step.
4. When the run pauses, proceed through run setup and start the second run or recovery wash.
 - » For run setup instructions, proceed to [Initiate a Sequencing Run on page 16](#).
 - » For recovery wash instructions, see the user guide for your instrument.
 5. To cancel setup of the second run or recovery wash, select **Back** to return to the Home screen, and then select **Resume**.

Discard the Cartridge and Bottle

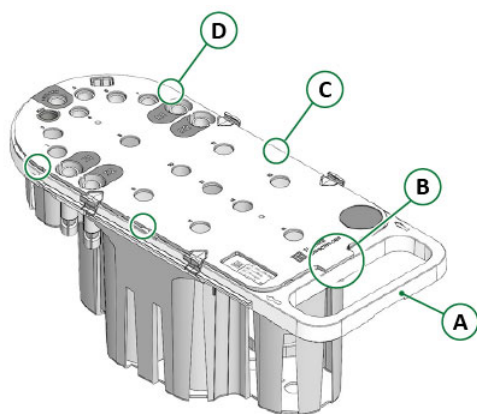
The cartridge and buffer bottle contain reagents with region-specific disposal requirements, which are described in the Safety Data Sheets (SDS) at [elementbiosciences.com/resources](https://www.elementbiosciences.com/resources). The amount of reagent remaining in each well after a run depends on how many cycles the run performed.

The following wells contain hazardous reagents. The position numbers in the figure align with the position numbers in the SDS.



Dispose of Reagents

1. Keep the cartridge in the basket with the clips locked.
2. Hold the basket handle with one hand and lift the cartridge lid tab with the other. Expect resistance at three points.



- A Basket handle
- B Cartridge lid tab and first point of resistance
- C Second point of resistance on both sides of the cartridge
- D Final point of resistance on both sides of the cartridge

3. Remove the hazardous wells from the cartridge.
—The volume remaining in each well depends on the number of cycles performed.—
4. Using a pipette tip or a similar tool, enlarge the hole in each foil seal to form a triangle.



5. Empty each well into hazardous waste or other appropriate container per the SDS.
6. Unlock the clips and remove the cartridge from the basket.
7. Remove the remaining wells from the cartridge and enlarge the hole in each foil seal.
8. Empty each well into the appropriate container per the SDS.
9. Discard the cartridge and buffer bottle per the SDS.
10. Rinse the basket with nuclease-free water and dry upside down.

Troubleshooting

The following troubleshooting information addresses problems that can occur during run setup and sequencing with a Cloudbreak, Cloudbreak Freestyle, or Cloudbreak UltraQ kit. If a problem persists, contact Element Technical Support. For more information on troubleshooting, see the user guide for your instrument.

Run Setup Problems

Problem	Resolution
The flow cell is cracked, scratched, or otherwise damaged.	Contact Element Technical Support.
Small particulates are visible in the flow cell lane.	See Cloudbreak Flow Cell Variations on page 32 .
The lid does not engage when a flow cell is on the nest.	Remove the flow cell and wipe the nest. Inspect the flow cell for large debris and wipe with an alcohol pad if necessary. Reload the flow cell.
AVITI OS cannot detect a loaded cartridge or waste bottle.	Follow the onscreen prompt to reload the cartridge or waste bottle. Make sure the applicable bay, reagent or waste, is unobstructed, and that the cartridge is contained within a cartridge basket.
The system cannot scan or detect a barcode on the cartridge, buffer bottle, or flow cell.	Follow the onscreen prompt to reload the consumable or continue by manually entering consumable information.
The flow cell version is incompatible with the cartridge.	Load a flow cell that is the same version as the cartridge.

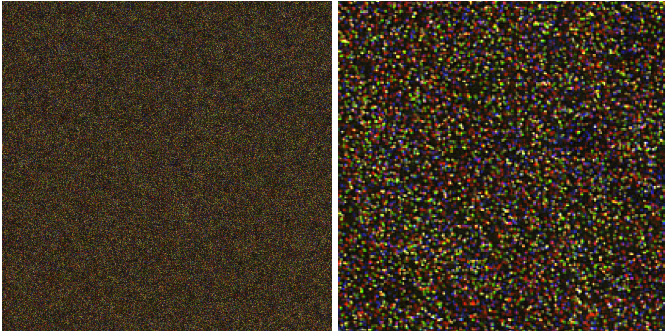
Sequencing Problems

Problem	Resolution
Polony density is lower or higher than expected.	Contact Element Technical Support or stop the run. For instructions on stopping a run, see the troubleshooting section of the user guide for your instrument.
The assigned or perfect match metrics are lower than expected.	Make sure that the index sequences recorded in the run manifest are correct.
The samples with low representation metric is higher than expected.	Select Sample Details to view the samples with low representation. Make sure that the index sequences recorded in the run manifest and the pooling concentration are correct.
The Q30 percentage is lower than expected.	Contact Element Technical Support.
The PhiX error rate is higher than expected.	
The flow cell contains very few polonies or no polonies.	
The user interface is frozen.	

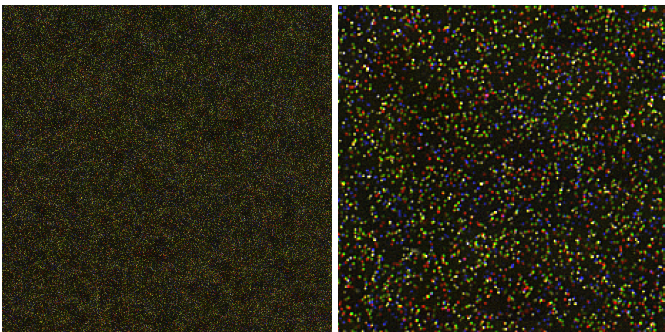
Thumbnail Image Troubleshooting

The following figures show example thumbnail images for a standard flow cell, an underloaded flow cell, and an overloaded flow cell. If the thumbnail image for a run indicates an underloaded flow cell, increase the loading concentration. For an overloaded flow cell, reduce the loading concentration. If problems persist, contact Element Technical Support.

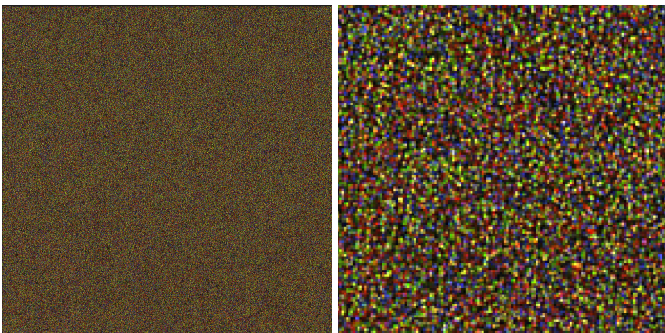
Example expected thumbnail image, full-size and zoomed



Example thumbnail image with underloading, full-size and zoomed



Example thumbnail image with overloading, full-size and zoomed



Cloudbreak Consumables

Cloudbreak consumables include a sequencing kit and optional controls and custom primers. The workflow also requires user-supplied consumables. For a list of required equipment, see the site prep guide for your instrument.

Cloudbreak Sequencing Kits

The following tables list the kit contents and storage requirements. Kits contain one of each part listed. The Library Loading Buffer pouch includes two tubes. For SDS information, see [elementbiosciences.com/resources](https://www.elementbiosciences.com/resources).

AVITI 2x150 Sequencing Kit Cloudbreak UltraQ (860-00018)

Part #	Component	Shipping	Storage
820-00026	AVITI 2x150 Cartridge Cloudbreak UltraQ	-25°C to -15°C	-25°C to -15°C
810-00008	AVITI Flow Cell Cloudbreak UltraQ	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x75 Sequencing Kit Cloudbreak Freestyle High Output (860-00015)

Part #	Component	Shipping	Storage
820-00022	AVITI 2x75 Cartridge Cloudbreak Freestyle High Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x75 Sequencing Kit Cloudbreak Freestyle Medium Output (860-00014)

Part #	Component	Shipping	Storage
820-00021	AVITI 2x75 Cartridge Cloudbreak Freestyle Medium Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x75 Sequencing Kit Cloudbreak Freestyle Low Output (860-00034)

Part #	Component	Shipping	Storage
820-00032	AVITI 2x75 Cartridge Cloudbreak Freestyle Low Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak Freestyle High Output (860-00013)

Part #	Component	Shipping	Storage
820-00020	AVITI 2x150 Cartridge Cloudbreak Freestyle High Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak Freestyle Medium Output (860-00012)

Part #	Component	Shipping	Storage
820-00019	AVITI 2x150 Cartridge Cloudbreak Freestyle Medium Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak Freestyle Low Output (860-00011)

Part #	Component	Shipping	Storage
820-00018	AVITI 2x150 Cartridge Cloudbreak Freestyle Low Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x300 Sequencing Kit Cloudbreak Freestyle High Output (860-00017)

Part #	Component	Shipping	Storage
820-00024	AVITI 2x300 Cartridge Cloudbreak Freestyle High Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x300 Sequencing Kit Cloudbreak Freestyle Medium Output (860-00016)

Part #	Component	Shipping	Storage
820-00023	AVITI 2x300 Cartridge Cloudbreak Freestyle Medium Output	-25°C to -15°C	-25°C to -15°C
810-00003	AVITI Flow Cell Cloudbreak Freestyle	Room temperature	2°C to 8°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x75 Sequencing Kit Cloudbreak High Output (860-00004)

Part #	Component	Shipping	Storage
820-00015	AVITI 2x75 Cartridge Cloudbreak High Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x75 Sequencing Kit Cloudbreak Medium Output (860-00007)

Part #	Component	Shipping	Storage
820-00014	AVITI 2x75 Cartridge Cloudbreak Medium Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak High Output (860-00003)

Part #	Component	Shipping	Storage
820-00013	AVITI 2x150 Cartridge Cloudbreak High Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak Medium Output (860-00006)

Part #	Component	Shipping	Storage
820-00012	AVITI 2x150 Cartridge Cloudbreak Medium Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x150 Sequencing Kit Cloudbreak Low Output (860-00005)

Part #	Component	Shipping	Storage
820-00011	AVITI 2x150 Cartridge Cloudbreak Low Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x300 Sequencing Kit Cloudbreak High Output (860-00008)

Part #	Component	Shipping	Storage
820-00016	AVITI 2x300 Cartridge Cloudbreak High Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

AVITI 2x300 Sequencing Kit Cloudbreak Medium Output (860-00009)

Part #	Component	Shipping	Storage
820-00017	AVITI 2x300 Cartridge Cloudbreak Medium Output	-25°C to -15°C	-25°C to -15°C
810-00002	AVITI Flow Cell Cloudbreak	Room temperature	2°C to 8°C
820-00010	Adept Primer Set Cloudbreak	-25°C to -15°C	-25°C to -15°C
820-00002	AVITI Buffer Bottle (Universal Wash Buffer)	Room temperature	Room temperature
820-00004	Library Loading Buffer	-25°C to -15°C	-25°C to -15°C

PhiX Control Library

PhiX Control Library is a color-balanced, ready-to-use library that adds diversity to low-complexity libraries. Each type of PhiX Control Library includes unique index sequences and has a concentration of 1 nM. For a list of sequences, see [Element Index Sequences](#).

Part #	Type	Format	Shipping and Storage
830-00004	PhiX Control Library, Adept	Circular	-25°C to -15°C
830-00017	Cloudbreak PhiX Control Library, Elevate	Linear	-25°C to -15°C
830-00023	Cloudbreak Freestyle PhiX Control, Third Party	Linear	-25°C to -15°C

Adept Primer Set Cloudbreak

Primers for I1, I2, R1, and R2 provided in Cloudbreak cartridges support Elevate libraries. Sequencing Adept libraries requires replacing the prepackaged primers with tubes from the Adept Primer Set Cloudbreak.

Adept Primer Set Cloudbreak (820-00010)

- Index 1—Adept Index 1 (I1) Primer Cloudbreak
- Index 2—Adept Index 2 (I2) Primer Cloudbreak
- Read 1—Adept Read 1 (R1) Primer Cloudbreak
- Read 2—Adept Read 2 (R2) Primer Cloudbreak

The Adept Primer Set Cloudbreak is not compatible with Cloudbreak Freestyle or Cloudbreak UltraQ kits.

Custom Primer Sets

A custom primer set provides read-specific buffers for preparing custom primers for Adept libraries with Cloudbreak chemistry or third-party libraries with Cloudbreak Freestyle chemistry.

Part #	Custom Primer Set	Buffers	Shipping and Storage
820-00009	Adept Custom Primer Set Cloudbreak	<ul style="list-style-type: none"> • Adept Custom Index 1 (I1) Buffer Cloudbreak • Adept Custom Index 2 (I2) Buffer Cloudbreak • Adept Custom Read 1 (R1) Buffer Cloudbreak • Adept Custom Read 2 (R2) Buffer Cloudbreak 	-25°C to -15°C
820-00025	Custom Primer Set Cloudbreak Freestyle	<ul style="list-style-type: none"> • Custom Index 1 (I1) Buffer Cloudbreak Freestyle • Custom Index 2 (I2) Buffer Cloudbreak Freestyle • Custom Read 1 (R1) Buffer Cloudbreak Freestyle • Custom Read 2 (R2) Buffer Cloudbreak Freestyle 	-25°C to -15°C
820-00038	Custom Primer Set Read 1, Cloudbreak Freestyle	<ul style="list-style-type: none"> • Custom Read 1 (R1) Buffer Cloudbreak Freestyle 	-25°C to -15°C

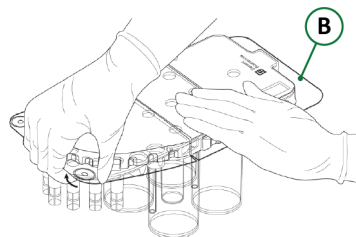
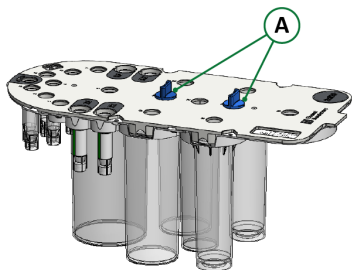
User-Supplied Consumables and Equipment

Consumables	Supplier
DNA LoBind Tubes, 2 ml	Eppendorf, # 022431021
0.2 M Tris-HCl, pH 7.0	General lab supplier
1 N NaOH	
10 mM Tris-HCl, pH 8.0 with 0.1 mM EDTA (Low TE buffer)	
Filtered pipette tips	
Nuclease-free laboratory-grade water	

Cartridge Shipping Configuration

The Cloudbreak sequencing cartridge includes shipping protection in the form of shipping locks or a thermoform shipping cover.

- If your cartridge includes the shipping locks (**A**), remove the shipping locks before loading the cartridge onto the instrument.
- If your cartridge includes the thermoform shipping cover (**B**), remove the cover before thawing reagents.



Cloudbreak Flow Cell Variations

Cloudbreak flow cells might have small particulate within the flow cell lane. These variations are normal and do not impact data quality.



Document History

Revision	Description of Change
February 2026 Document # MA-00058 Rev. H	<ul style="list-style-type: none">• Added a note to use the default index cycle settings for Elevate libraries.• Added a note on wiping flow cell to remove any debris.• Added guidance on discarding a run setup to correct any error during setup.• Converted library loading calculations into tables instead of text for clarity.• Reordered topics to move Thaw Reagents before Add Primer Tubes, and split the Empty Waste and Prime Reagents into separate topics for clarity.• Updated workflow diagram to fix inconsistent numbering.
July 2025 Document # MA-00058 Rev. G	<ul style="list-style-type: none">• Added read counts, output, and loading concentrations for AVITI24 systems running AVITI OS v3.4, or later, with Cloudbreak or Cloudbreak Freestyle chemistry and 2 x 75 or 2 x 150 cycle kits.• Updated steps to dispose reagents to show three main points of resistance when removing the cartridge lid.
May 2025 Document # MA-00058 Rev. F	<ul style="list-style-type: none">• Added Custom Primer Set Read 1 Cloudbreak Freestyle to custom primer sets.• Corrected column order for Cloudbreak Freestyle 2 x 300 loading concentration.
April 2025 Document # MA-00058 Rev. E	<ul style="list-style-type: none">• Updated expected priming time when using AVITI OS v3.3.
March 2025 Document # MA-00058 Rev. D	<ul style="list-style-type: none">• Added run times for each kit configuration.• Added Read Counts and Outputs table to Overview section.• Added Universal Wash Buffer to each mention of AVITI Buffer Bottle.• Removed the term pollination in the Run Stages description.• Updated storage time for thawed cartridges.
January 2025 Document # MA-00058 Rev. C	<ul style="list-style-type: none">• Added recommendations for using short insert and long insert recipes.• Added run specifications for the Cloudbreak Freestyle 2 x 75 low output kit.• Added example of different cartridge shipping configurations, such as shipping locks or shipping cover.• Added statement that Cloudbreak flow cell variations do not impact data quality.• Recommended a pipette tip or similar tool to enlarge hole in foil seal.
December 2024 Document # MA-00058 Rev. B	<ul style="list-style-type: none">• Added 2 x 75 Cloudbreak Freestyle Low Output kit.• Updated name of sequencing basket to cartridge basket.
October 2024 Document # MA-00058 Rev. A	<ul style="list-style-type: none">• Initial release of user guide.

Technical Support

Visit the [Documentation page](#) on the Element Biosciences website for additional guides and the most recent version of this guide. For technical assistance, contact Element Technical Support.

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