

oncoReveal™ 4 Gene Methylation

USER MANUAL



UM-0077 version 1.0

For Research Use Only. Not for use in diagnostic procedures.

Revision History

Version 1.0: Initial Release

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1. Product Introduction

The **oncoRevealTM** 4 Gene Methylation Panel targets specific genetic sequences in DNA that have undergone a bisulfite conversion for methylome analysis. This panel covers CpG sites within $BRCA_1$, $BRCA_2$, RAD_5_1C , and $XRCC_3$. For more information on the panel specifications, please see the oncoRevealTM 4 Gene Methylation Panel product sheet (*Doc. No. MK-0046*).

2. Product Description

The oncoReveal[™] 4 Gene Methylation Panel utilizes our proprietary SLIMamp® (stem-loop inhibition mediated amplification) technology, allowing researchers to amplify regions of interest in a simple, single-tube, multiplex reaction. Pairs of DNA oligos designed for each region of interest are used in the first round of gene-specific PCR (GS-PCR) and the products are subsequently purified via size selection. After purification, a second round of PCR adds index adaptors and P5 & P7 sequences to each library for sample tracking and sequencing. The final libraries are further purified and can be sequenced on the Illumina sequencing platform.

The panel requires an indexing kit, available in low or high throughput options, and the resulting libraries are designed for sequencing on the Illumina platform using a paired-end read length of 150 (2x150). The workflow can be performed and loaded onto the sequencing instrument within one day. Additionally, this protocol also contains numerous stopping points for users who have time limitations.



Figure 1. Overview of the oncoReveal $^{\text{TM}}$ 4 Gene Methylation Panel library preparation.

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3. General Laboratory Guidelines

The following steps are recommended to improve consistency and reduce contamination:

- Work areas: To reduce the risk of contamination from PCR amplicons, supplies should not be moved from one area
 to another. Separate storage areas (refrigerator, freezer) should also be designated for pre- and post-PCR products.
- Lab cleanliness: To further reduce the possibility of contamination, work areas should be cleaned between experiments with laboratory cleaning solution (70% alcohol or freshly-made 10% hypochlorite solution). A periodic cleaning of the floor is also recommended.
- Floor: Items that have fallen to the floor are assumed to be contaminated and should be discarded. If a sample tube or non-consumable item has fallen to the floor and remained sealed, the outer surface of the item should be thoroughly cleaned before use using 70% alcohol or freshly-made 10% hypochlorite solution. Gloves should be changed after handling a contaminated item.
- Aliquot reagents: Frozen reagents should be aliquot into smaller volumes to reduce freeze/thaw cycles. To reduce
 the risk of stock contamination it is recommended to aliquot from the stock and work from the aliquots. In case of
 contamination, the use of aliquots can also help to determine the source more quickly and easily.
- Multichannel pipettes: Multichannel pipettes should be used to maintain consistency and efficiency across numerous samples.
- **Pipette tips:** Tips should be changed between each sample to prevent cross-contamination. Any tips that may have become contaminated due to contact with gloves, the lab bench, tube exteriors, etc., should be discarded.
- Open containers and lids: To prevent possible contamination from the air, tubes should be kept closed when not
 directly in use, and plates, troughs, and similar reservoirs should be covered with seals or lint-free laboratory wipes.
 Additionally, reaching over open containers should be avoided.

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4. Reagents & Equipment

This section describes the necessary equipment, reagents, and consumables needed before performing the protocol.

All reagents should be used in designated pre-PCR or post-PCR areas to prevent amplicon contamination. Each area designated for pre- and post-PCR should have dedicated equipment, reagents, and supplies (including gloves, lab coats, etc.) to prevent contamination.

4.1 Kit Components

oncoReveal™ 4 Gene Methylation Panel Part No.: HDA-HR-1009-24

Reagent	Use	Area Use	Storage
Gene-Specific PCR Master Mix (2x)	Gene-Specific PCR	Pre-PCR	-25° to -15°C
oncoReveal™ 4 Gene Methylation Oligo Pool	Gene-Specific PCR	Pre-PCR	-25° to -15°C
Indexing PCR Master Mix (2x)	Indexing PCR	Pre-PCR	-25° to -15°C

4.2 Indexing Kits

Reagent	Part Number	Use	Area Use	Storage
Pillar Custom Indexing Primers Kit A , indices PI501-8, PI701-4 (32 combinations – 96 reactions)	IDX-PI-1001-96	Indexing PCR	Pre-PCR	-25° to -15°C
Pillar Custom Indexing Primers Kit D , indices PI501-8, PI701-12 (96 combinations – 192 reactions)	IDX-PI-1004-192	Indexing PCR	Pre-PCR	-25° to -15°C

Only one index kit is needed per assay. Multiple options are available to meet a variety of throughput needs.

4.3 User-Supplied Reagents

Reagent	Area Use	Supplier
10N NaOH or 1N NaOH	Post-PCR	General lab supplier
AMPure XP Beads	Post-PCR	Beckman Coulter, Cat# A63881 or A63880
Ethanol, 200 proof for molecular biology	Post-PCR	General lab supplier
Nuclease-free water	Post-PCR	General lab supplier
Qubit dsDNA High Sensitivity Assay kit	Post-PCR	Invitrogen, Cat# Q32851 or Q32854
Agarose gel, 2% (optional) ¹	Post-PCR	General lab supplier
DNA molecular weight markers (optional) ¹	Post-PCR	General lab supplier
TapeStation or equivalent	Post-PCR	Agilent Technologies
10 mM Tris-HCl w/ 0.1% Tween-20, pH 8.5 (optional)	Post-PCR	Teknova, Cat# T7724
PhiX Control v ₃	Post-PCR	Illumina, Cat# FC-110-3001
200 mM Tris-HCl, pH 7.0 (optional) ²	Post-PCR	General lab supplier

¹ The Qubit dsDNA High Sensitivity Assay kit is the primary DNA quantitation assay used throughout this protocol. Additional DNA quantification can optionally be performed using an agarose gel or using TapeStation (or equivalent).

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4.4 Other Consumables

Compatible Sequencing Reagents

Sequencing Reagent Kit	Supplier	Catalog No.
MiniSeq™ Mid Output kit (300 cycles)	Illumina	FC-420-1004
MiniSeq™ High Output kit (300 cycles)	Illumina	FC-420-1003
MiSeq™ Reagent Nano kit v2 (300 cycles) ‡	Illumina	MS-103-1001
MiSeq™ Reagent Micro kit v2 (300 cycles) ‡	Illumina	MS-103-1002
MiSeq™ Reagent kit v2 (300 cycles) ‡	Illumina	MS-102-2002
MiSeq™ Reagent kit v3 (600 cycles) ‡	Illumina	MS-102-3003
NextSeq™ 500/550 Mid Output v2.5 kit (300 cycles)	Illumina	20024905
NextSeq™ 500/550 High Output v2.5 kit (300 cycles)	Illumina	20024908
NextSeq™ 1000/2000 P1 Reagents (300 cycles)	Illumina	20050264
NextSeq™ 1000/2000 P1 Reagents (600 cycles)	Illumina	20075294
NextSeq™ 1000/2000 P2 Reagents v3 (300 cycles)	Illumina	20046813
NextSeq™ 1000/2000 P2 300M Reagents (600 cycles)	Illumina	20075295
NextSeq™ 2000 P3 Reagents (300 cycles)	Illumina	20040561

[‡] Indicates the flowcells the oncoReveal™ 4 Gene Methylation Panel has been validated on. However, libraries generated using this protocol are compatible with all Illumina sequencers.

General Laboratory Consumables

In addition to the consumables listed below other general laboratory supplies needed to carry out the protocol include gloves, prechilled cooler, tube racks, etc.

Consumable	Area Use	Supplier
1.5 mL microcentrifuge tubes	Pre- and Post-PCR	General lab supplier
96-well PCR plates, 0.2 mL	Pre- and Post-PCR	Corning Life Sciences, Cat# 6551 or
90-well 1 Cit plates, 0.2 IIIL	Tie- and Tost-Tote	equivalent
Microplate sealing film	Pre- and Post-PCR	Corning Life Sciences, Cat# PCR-TS or
whereplate scanng min	Tie- and Tost-Tork	equivalent
Conical tubes, 15 mL	Pre- and Post-PCR	General lab supplier
Conical tubes, 50 mL	Post-PCR	General lab supplier
Low retention, aerosol filter pipette tips	Pre- and Post-PCR	General lab supplier
Solution basin (trough or reservoir)	Pre- and Post-PCR	Fisher Scientific, Cat# 13-681-506 or
Solution basin (trough of reservoir)	The and 10st 1 Cit	equivalent
Qubit Assay tubes	Post-PCR	Invitrogen, Cat# Q32856

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² The 200 mM Tris-HCl, pH 7.0 reagent is only required for denaturing libraries for sequencing on the Illumina NextSeq or MiniSeq. If sequencing on the MiSeq this reagent is not needed.

4.5 Equipment

Equipment	Area Use	Supplier
Centrifuge adapted for PCR plates, tabletop	Pre- and Post-PCR	General lab supplier
Gel electrophoresis apparatus (optional)*	Post-PCR	General lab supplier
TapeStation or equivalent*	Post-PCR	Agilent Technologies
Magnetic stand for 96 wells	Post-PCR	Life Technologies, Cat# 12331D or 12027
Microfuge	Pre- and Post-PCR	General lab supplier
Thermal cycler, heated lid capability	Post-PCR	General lab supplier
Pipettes, 0.5-1000 μL capabilities	Pre- and Post-PCR	General lab supplier
Qubit Fluorometer	Post-PCR	Invitrogen, Cat# Q33216/Q33218
Vortex mixer	Pre- and Post-PCR	General lab supplier

^{*} The Qubit dsDNA High Sensitivity kit is the primary DNA quantitation assay used throughout this protocol. Additional DNA quantification can optionally be performed using either an agarose gel or TapeStation (or equivalent).

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5. Workflow

The following chart demonstrates the workflow for performing the oncoReveal $^{\text{\tiny IM}}$ 4 Gene Methylation Panel library preparation.

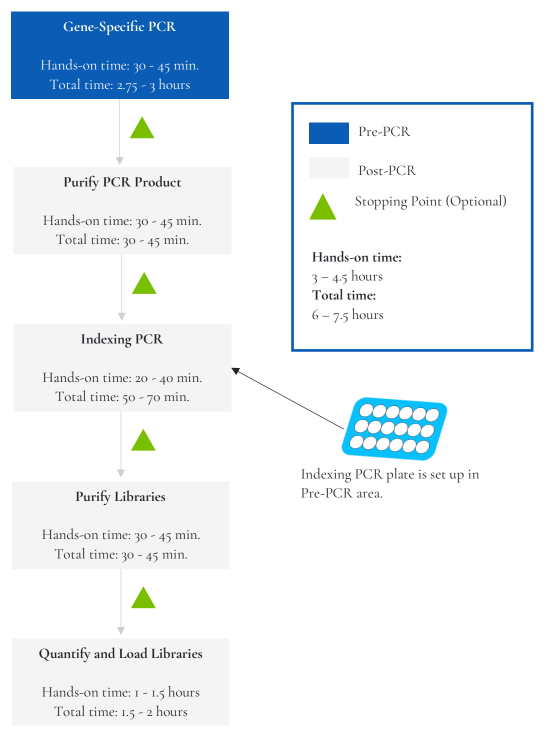


Figure 2. Library preparation workflow for oncoReveal $^{\text{TM}}$ 4 Gene Methylation Panel. The workflow can be completed within one day but contains multiple optional stopping points for users with time constraints.

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6. DNA Input Information

The following protocol includes information for preparing libraries using DNA that has undergone a bisulfite or enzyme conversion.

The recommended DNA input is $\mathbf{5}$ – $\mathbf{20}$ \mathbf{ng} of bisulfite converted DNA per PCR reaction.

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Symbol	Description
•	Information: Information that follows this symbol is important and may require action.
	Optional Stopping Point: A point in the workflow at which work may be safely paused, and samples can be stored appropriately.
Caution: Information that follows this symbol is critical to the workflow. Information this symbol should not be skipped or ignored.	

Symbols used throughout this protocol and their associated meaning.

7. Library Preparation Protocol

Hands-on Time	3 – 4.5 hours	
Total Time	6 – 7.5 hours	

7.1 Gene-Specific PCR (GS-PCR) and Purification

Hands-on Time	30 – 40 min.
Total Time	2.75 – 3 hours

Amplify Genomic DNA Targets

The following steps should be performed in a pre-PCR area.

For this portion of the protocol prepare a prechilled benchtop cooler. Keep the gene-specific PCR Master Mix (GS-PCR MMX) and the custom oligo pool in the cooler until needed.

See recommended DNA input quantities in the section titled "DNA Input Information."

- Prepare Gene-Specific PCR Mix 1: Vortex and centrifuge the GS-PCR MMX and oligo pool before use. For each PCR reaction, the volume of each component is listed below.
- **Important**: The gene-specific PCR master mix reagent is viscous. Ensure the mix is fully homogenized before adding other reaction components. Vortexing is recommended and will not adversely affect enzyme activity.

GS-PCR Mix 1	
Reagent	Volume (μL)
Gene-Specific PCR Master Mix	12.5
oncoReveal™ 4 Gene Methylation Panel Oligo Pool	5.0
Subtotal	17.5

2. **Transfer:** Transfer 17.5 μL of GS-PCR Mix 1 to each sample well in a PCR plate, strip tube, or PCR tube.

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3. Dilute input DNA: Add 7.5 μ L of DNA (diluted if necessary)* to each sample well containing GS-PCR Mix 1. Add 7.5 μ L of nuclease-free water to the no-template control well.

Final GS-PCR Mix		
Reagent	Volume (μL)	
GS-PCR Mix 1	17.5	
DNA (or water)	7.5	
Total	25.0	

^{*} The DNA concentration can be determined using the Qubit dsDNA HS Assay Kit.

- 4. **Seal and mix:** Carefully seal the reactions and vortex for 10 15 seconds.
- 5. **Spin:** Briefly centrifuge the reactions to remove any air bubbles from the bottom of the wells and to spin down droplets from the seal or side walls.

The following steps should be performed in a post-PCR area.

6. Perform GS-PCR: Perform the following program with the heated lid on:

GS-PCR Program		
Temperature	Time	Number of Cycles
95°C	15 min	I
98°C 56°C	1 min	6
56°C	1 min	6
95°C 66°C	30 sec	D. /
66°C	1 min	24
8°C	Hold	I



Do not leave reactions on the thermocycler overnight at 8°C. Once cycling is complete the reactions can be stored at -20°C.



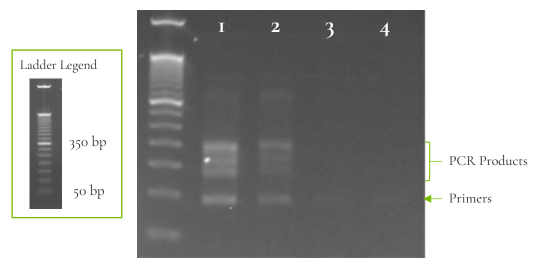


Optional Stopping Point: The gene-specific PCR reactions may be stored at -20°C after cycling is complete.

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Gel Image After Gene-Specific PCR

The following image is an example of samples after gene-specific PCR on a 2% agarose gel.



Lane 1: 20 ng Bisulfite-converted DNA (50% methylated, 50% unmethylated)

Lane 2: 10 ng Bisulfite-converted DNA (50% methylated, 50% unmethylated)

Lane 3: 20 ng unconverted genomic DNA

Lane 4: No Template Control (NTC)

Figure 3. Analysis of GS-PCR product on 2% agarose gel. The doublet in the PCR product is a result of the presence of amplicons of different sizes. PCR product bands may appear faint when using low quantities of DNA.

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Purify the Gene-Specific PCR Product

Hands-on Time	30 – 45 min.
Total Time	30 – 45 min.

The following steps should be performed in a post-PCR area.

Before beginning the purification:

Remove AMPure XP beads from 4°C and incubate at room temperature for at least 30 minutes before use.



Caution: Ensure that the AMPure bead solution reaches room temperature before performing the purification. The temperature of the bead solution can have adverse effects on the purification process.

- If gene-specific PCR products were stored at -20°C, remove from the freezer to thaw at room temperature before purification.
- Prepare 70% ethanol by mixing three parts of water with seven parts of absolute ethanol in a conical tube, invert tube a few times to mix, and dispense sufficient volume to a disposable trough for convenient dispensing using a multichannel pipette.



Caution: fresh 70% ethanol is required for optimal results.

Gene-Specific Product Purification

- I. If condensation has formed or if reactions were stored at -20°C briefly centrifuge the reactions upon thawing. Carefully remove the seal.
- 2. Mix Beads: Vortex AMPure XP beads thoroughly until all beads are well-dispersed.



Caution: Ensure that the AMPure bead solution is homogenous. A non-uniform distribution can have unpredictable effects on the purification process.

- 3. Add water to sample: Add 25 μ L of nuclease-free water to each well containing GS-PCR product to bring the reaction volume to 50 μ L.
- 4. Add beads: Add 75 μL of AMPure beads (equivalent to a 1.5x bead ratio) to each well. Pipette the mixture up and down 10 times. If bubbles form on the bottom of the wells, briefly centrifuge the samples and mix again.
- 5. Bind GS-PCR product to beads: Incubate for 5 minutes at room temperature.
- 6. **Separate beads containing GS-PCR product:** Place the PCR plate on a magnetic rack until the solution appears clear, which can take up to 5 minutes.
- Remove supernatant: Carefully remove the supernatant from each well without disturbing the beads.

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- 8. Wash beads: With the PCR plate still on the magnetic rack add 150 μL of freshly prepared 70% ethanol to each well without disturbing the beads. Incubate at room temperature for 30 seconds, and then remove the supernatant from each well.
- Important: Do not allow the ethanol mixture to remain open to the air. The ethanol concentration will change over time, affecting the washing of the beads. Pour only enough solution for each wash.
- 9. **Second wash:** Repeat step 8 for a second 70% ethanol wash. Remove the supernatant from each well. The unused solution of ethanol can be used to purify the libraries after the indexing PCR is completed.
- 10. **Remove remaining ethanol wash:** Centrifuge for 10–15 seconds, place the PCR plate back on the magnetic rack, and use a 10 or 20 μL tip to remove any trace amounts of ethanol from each well.
- II. Dry beads: Keep the PCR plate on the magnetic rack and let the beads air dry at room temperature for 2-5 minutes.
- Important: To mitigate product loss, do not allow the beads to over-dry. The beads have sufficiently dried when the bead mass has small cracks in the middle. If large cracks have appeared among the entire bead ring or they are flaky, they are over-dried. Beads that are too dry may be difficult to resuspend.
- 12. Resuspend beads: Remove the samples from the magnetic rack and resuspend the dried beads in each well using 64 μL of nuclease-free water. Gently pipette the suspension up and down 10 times. If bubbles form on the bottom of the wells, briefly centrifuge and mix again.
- 13. **Incubate and Elute:** Incubate for 5 minutes to fully elute the product.
 - TIP: After resuspending the beads, cover the reactions and prepare for the indexing PCR step using the Indexing PCR Master Mix. The Indexing PCR Master Mix should be handled in the pre-PCR area. Alternatively, the purified gene-specific PCR product may be stored at -20°C after elution.
- 14. Separate GS-PCR product from beads: Place the reactions on a magnetic rack at room temperature until the solution appears clear. Transfer $62 \mu L$ of clear supernatant from each well to a new PCR plate.
- Optional Stopping Point: The purified GS-PCR products can be stored at -20°C.

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7.2 Index PCR & Purification

Indexing PCR: Amplify the Libraries

Hands-on Time	20 – 40 min.
Total Time	50 – 70 min.

The following steps should be performed in a pre-PCR area.

For this portion of the protocol prepare a prechilled cooler. The Indexing PCR Master Mix should be kept in the cooler until it is needed.

1. Prepare Indexing Primer Mix: In a new PCR plate add $4\,\mu\text{L}$ each of the appropriate forward and reverse indexing primers to each sample well that will be used.

Indexing Primer Mix (per reaction)	
Reagent	Volume (µL)
Pi700 Pillar Index	4.0
Pi500 Pillar Index	4.0
Subtotal 8.0	

2. **Prepare Indexing PCR Mix 1:** Briefly vortex and centrifuge the Indexing PCR Master Mix before use. Prepare Indexing PCR Mix 1 by combining Indexing PCR Master Mix and water with sufficient overage.

Indexing PCR Mix 1 (per reaction)	
Reagent	Volume (μL)
Indexing PCR Master Mix (2x)	25.0
Nuclease-free water	II.O
Subtotal	36.0

3. Add Indexing PCR Mix I to Indexing Primer Mix: Transfer 36 μL of the Indexing PCR Mix I to each sample well of the PCR plate in step I that contains Indexing Primer Mix. To prevent cross-contamination of indices, be sure to change tips between each well.

Indexing PCR Mix 2	
(per reaction)	
Reagent	Volume (μL)
Indexing Primer Mix	8.0
Indexing PCR Mix 1 36.0	
Subtotal	44.0

The following steps should be performed in a post-PCR area. Cover or seal the reactions before transferring from the pre-PCR area to the post-PCR area.

If the purified GS-PCR products were stored at -20° C after the GS-PCR purification, ensure that they have been thawed at room temperature before proceeding.

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 Add purified GS-PCR product: Aliquot 6 μL of purified GS-PCR product into the appropriate wells containing Indexing PCR Mix 2.

Indexing PCR Final Mix (per reaction)	
Reagent	Volume (μL)
Indexing PCR Mix 2	44.0
Purified Gene-Specific PCR product	6.0
Total	50.0

- 5. Seal and mix: Carefully seal the reactions and vortex for 10-15 seconds.
- 6. **Spin:** Briefly centrifuge the reactions to remove any bubbles from the bottom of the wells and to spin down droplets from the seal or side walls.
- 7. **Perform Indexing PCR:** Perform the following program with the heated lid on.

Indexing PCR Program		
Temperature	Time	Number of Cycles
95°C	2 min	I
95°C 66°C	30 sec	
66°C	30 sec	8*
72°C	1 min	
72°C	5 min	I
8°C	Hold	Ι

^{*} Additional Indexing PCR cycles can be performed if final library yield is low or initial DNA input is below recommended minimum.



Optional Stopping Point: The indexed libraries can be stored at -20°C after cycling is complete.

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Indexing PCR: Purify the Libraries

Hands-on Time	30 – 45 min.
Total Time	30 – 45 min.

Before beginning the purification:

• Keep AMPure XP beads at room temperature while the Indexing PCR is being performed unless reactions will be stored at -20°C after the program is complete.



Caution: Ensure that the AMPure bead solution reaches room temperature before performing the purification. The temperature of the bead solution can have adverse effects on the purification process.

 If the indexed libraries were stored at -20°C remove them from the freezer to thaw thoroughly to ambient temperature before purification. After samples have thawed, briefly centrifuge to remove any droplets from the side walls.

Library Purification

The following steps should be performed in a post-PCR area.

I. Mix beads: Vortex AMPure XP beads thoroughly until all beads are well dispersed.



Caution: Ensure that the AMPure bead solution is homogenous. A non-uniform distribution can have unpredictable effects on the purification process.

- 2. Add beads: Add $60 \mu L$ beads (equivalent to a 1.2x bead ratio) to each well. Pipette the mixture up and down 10 times. If bubbles form on the bottom of the wells, briefly centrifuge and mix again.
- 3. **Bind libraries to beads:** Incubate for 5 minutes at room temperature.
- 4. **Separate libraries on beads:** Place the PCR plate on a magnetic rack until the solution appears clear, which can take up to 5 minutes.
- 5. Remove supernatant: Carefully remove the supernatant from each well without disturbing the beads.
- 6. Wash beads: With the PCR plate still on the magnetic rack add 150 μL of freshly prepared 70% ethanol to each well without disturbing the beads. Incubate at room temperature for 30 seconds, and then remove the supernatant from each well.



Caution: Do not allow the ethanol mixture to remain open to the air. The ethanol concentration will change over time, affecting the washing of the beads. Pour only enough solution for each wash.

- 7. Second wash: Repeat step 6 for a second 70% ethanol wash. Remove the supernatant from each well.
- 8. **Remove remaining ethanol wash:** Centrifuge for 10 15 seconds, place the PCR plate back on the magnetic rack, and use a 10 or 20 μL tip to remove any trace amounts of ethanol from each well.

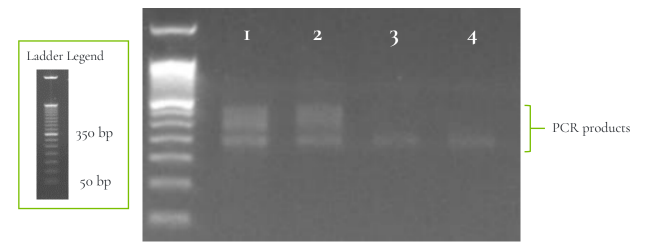
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- 9. Dry beads: Let the beads air dry at room temperature for 2 5 minutes.
- Important: To mitigate product loss, do not allow the beads to over-dry. The beads have sufficiently dried when the bead mass has small cracks in the middle. If large cracks have appeared among the entire bead ring or they are flaky, they are over-dried. Beads that are too dry may be difficult to resuspend.
- 10. Resuspend beads: Remove the PCR plate from the magnetic rack and resuspend the dried beads in each well using $32~\mu\text{L}$ of nuclease-free water. Gently pipette the bead suspension up and down 10 times. If bubbles form on the bottom of the wells briefly centrifuge and mix again.
- 11. Elute libraries: Incubate for 5 minutes at room temperature.
- 12. **Separate libraries from beads:** Place the bead suspensions on the magnetic rack at room temperature until the solution appears clear. Transfer 30 μ L of clear supernatant to a new PCR plate.
 - TIP: During the incubation and magnetic separation of the beads, cover the samples with microplate sealing film and prepare the solutions needed for quantitation performed in the next section.
- 13. Quantification: Analyze an aliquot of each library per the instructions in the next section.
- Optional Stopping Point: The purified libraries can be stored at 4°C for up to 3 days, or at -20°C for longer-term storage.

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Final Library Gel Image

The following image is an example of final libraries after both rounds of PCR and purification on a 2% agarose gel.



Lane 1: 20 ng Bisulfite-converted DNA (50% methylated, 50% unmethylated)

Lane 2: 10 ng Bisulfite-converted DNA (50% methylated, 50% unmethylated)

Lane 3: 20 ng unconverted genomic DNA

Lane 4: No Template Control (NTC)

Figure 4. Gel analysis of final library material. The PCR product doublet stems from differing amplicon sizes.

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7.3 Quantitation and Normalization of Purified Libraries

Prior to sequencing libraries must be quantified, normalized, and then pooled together. The following section describes how to quantify libraries using the Qubit system. Other library quantification methods, such as qPCR quantification or the Bioanalyzer, may be used according to the manufacturer's protocol.

Qubit Quantitation

Hands-on Time	30 – 45 min.
Total Time	30 – 45 min.

The following steps should be performed in a post-PCR area.

 Prepare Qubit working solution: Dilute the Qubit dsDNA HS reagent 1:200 in Qubit dsDNA HS buffer. Vortex briefly to mix the Qubit working solution.

For example, $2000~\mu L$ is sufficient buffer for 10 readings (8 samples + 2 standards). Combine 1990 μL of Qubit dsDNA HS buffer and 10 μL HS reagent.



Important: Fluorescent dyes are sensitive to light. Protect the Qubit working solution from light.

- 2. Label tubes: Set up 0.5 mL Qubit tubes for standards and samples. Label the tube lids.
- 3. Prepare standards: Transfer 190 μ L of Qubit working solution into two tubes for standard 1 and standard 2, and then add 10 μ L of each standard to the corresponding tube.



Caution: New standard dilutions should be prepared with the libraries to be quantified. Do not re-use standard dilutions from previous experiments.

- 4. Prepare samples: Transfer 198 μ L of Qubit working solution to each 0.5 mL tube, and then add 2 μ L of each library to its corresponding Qubit tube (1:100 dilution).
- 5. Mix and spin: Vortex to mix and then centrifuge the tubes briefly.
- 6. Incubate the tubes at room temperature for 2 minutes.
- 7. **Measure concentration**: Measure the concentration of each library on the Qubit 2.0 Fluorometer per the manufacturer's instructions. Use the dsDNA High Sensitivity assay to read standards 1 and 2 followed by the libraries.

If any concentration is above the linear range of the instrument, prepare a new dilution by combining 199 μL Qubit working solution and 1 μL library (1:200 dilution). Repeat steps 5–7.

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8. Calculate concentration: o.83 ng/ μ L of library is equal to 5 nM. Example calculation is below. Adjust dilution factor accordingly.

 $2~\mu L$ of library + 198 μL Qubit working solution:

$$\frac{Qubit \ reading\left(\frac{ng}{mL}\right)}{1,000} \ x \ dilution \ factor \ (100) \ x \ conversion \ factor \ (6) = nM$$



Optional Stopping Point: Once libraries have been quantified either proceed with normalization and pooling or else keep them stored at 4° C for up to 3 days. Store libraries at -20° C for long-term storage.

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Normalization and Pooling

Hands-on Time	30 – 45 min.
Total Time	30 – 45 min.

The following steps should be performed in a post-PCR area.



If sequencing on the MiniSeq, libraries should be normalized to 1 nM prior to pooling. For all other Illumina sequencing platforms, libraries should be normalized to 5 nM prior to pooling.



I. Normalize libraries to 5 nM: Dilute an aliquot (e.g., 4 μL) of each sample library to 5 nM using nuclease-free water or 10 mM Tris-HCl with 0.1% Tween-20, pH 8.5. An example calculation is as follows:

$$\frac{\textit{Library concentration (nM) x 4 \mu L library}}{5 \textit{nM}} = \textit{final volume of library}$$

Final volume of library $-4 \mu L$ library = volume of diluent

- 2. Mix and spin: Mix the 5 nM libraries thoroughly by vortexing and then centrifuge briefly.
- 3. **Prepare library pool:** Label a new 1.5 mL microtube for the library pool. Prepare an equimolar 5 nM mixture of libraries by combining each library at equal volume (e.g., mixing 5 µL of each 5 nM library). Gently pipette the entire solution up and down 10 times to mix thoroughly. Quickly vortex the pool and then briefly centrifuge.
- 4. **Quantify library pool (recommended):** The libraries prepared using the oncoReveal[™] 4-Gene Methylation Panel cluster very efficiently on the MiSeq.

It is recommended that the library pool be quantified using Qubit or another library quantitation method (qPCR) to ensure the pool is at 5 nM (\pm 10%) to prevent poor sequencing performance. If the final dilution is not 5 nM (\pm 10%), adjust the dilution for loading the sequencer accordingly to obtain the desired concentration.



Optional Stopping Point: The normalized libraries can be stored at 4°C overnight for sequencing the next day. For longer storage, the normalized libraries can be stored at -20°C.

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8. Sequencing

Prepare Diluted Libraries for Sequencing

Hands-on Time	30 – 40 min.
Total Time	30 – 40 min.

The libraries generated using this protocol can be multiplexed and sequenced on Illumina® sequencers. Table 1 outlines the sequencing parameters and the recommended per-sample sequencing coverage.

The number of samples that can be multiplexed together is dependent on several factors, among them are the estimated throughput of the flowcell and sequencing platform, the desired sequencing depth, as well as the number of unique index combinations available.

The estimated **maximum** number of samples that can be multiplexed on a single flowcell using each kit is displayed in Table 2. Select the appropriate sequencing kit based on the number of samples to be sequenced.

Sequencing Configuration	Index	Recommended Coverage	Recommended Reads Per Sample
2 X 150	Dual (8 bp each)	Mean: 2000x Min: 400x	48,000 PE reads

Table 1. Sequencing recommendations.



Libraries generated with this protocol require 33% PhiX DNA to be added prior to sequencing.

Sequencing Flowcell	Estimated Instrument Output (million PE reads)	Estimated Maximum Sequencing Batch Size
MiSeq Nano	2	25
MiSeq Micro	8	100
MiSeq v2	30	376
MiSeq v3	50	>600

Table 2. Multiplexing recommendations for Illumina sequencers. Calculations assume \geq 90% effective on-target rate after read mapping and minimum segment coverage > 20% of the mean coverage. Multiplexing values account for the PhiX DNA required for sequencing.

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The following steps should be performed in a post-PCR area.

For this portion of the protocol prepare a prechilled cooler.

Sequencing Using Illumina MiSeq[™] (v2 or v3) Reagents

MiSeq recommended final library pool loading concentration: 15 pM MiSeq recommended PhiX spike-in concentration: 12.5 pM

The following steps can be found in greater detail in Illumina's "MiSeq System: Denature & Dilute Libraries Guide" (Doc# 15039740).

- I. **Normalize:** Dilute libraries to 5 nM, as demonstrated in the previous section "Quantification and Normalization of Purified Libraries."
- 2. **Prepare 0.2 N NaOH:** Label a new 1.5 mL microtube for 0.2 N NaOH. Prepare the NaOH by combining **800 \muL** of nuclease-free water with **200 \muL** of 1 N NaOH. Vortex the solution to mix.
 - **NOTE:** Alternatively, prepare a 1 N NaOH solution by combining $500 \mu L$ of 10 N NaOH with 4.5 mL of nuclease-free water. Vortex the solution to mix. If 1 N NaOH has not been prepared within the last week from a 10 N solution, prepare a new 1 N NaOH solution.
- 3. Denature the library pool: Label a new 1.5 mL microtube for the denatured 25 pM library pool.
 - Denature the 5 nM library pool by combining 5 μ L of the library pool and 5 μ L of the freshly prepared 0.2 N NaOH.
 - b. Vortex thoroughly for 10 seconds and centrifuge for 1 minute.
 - c. Let the solution stand at room temperature for 5 minutes.
 - d. Add 990 µL of Illumina's HT1 hybridization buffer to the denatured library pool.
 - e. Invert the mixture several times, centrifuge briefly, and place in prechilled cooler.
- 4. Dilute 25pM library mix to 15 pM and add PhiX control: Label a new 1.5 mL microtube for the 15 pM library mix. Combine 240 μL of the 25 pM library mix (step 2) with 120 μL of Illumina's HT1 solution and 240uL 12.5pM PhiX. Adjust the volumes as needed for libraries that are over or under 25 pM. Invert the mixture several times, spin briefly, and place on ice.
- 5. **Load MiSeq cartridge:** Using a clean 1000 μL tip puncture the foil cap above the sample loading well on the MiSeq cartridge. Load the **600 μL** library and PhiX mixture into the cartridge, ensuring that the solution has reached the bottom of the well.
- 6. **Run the MiSeq:** Sequence the libraries on the MiSeq per the manufacturer's instructions using a paired-end read length of 150 (2x150) and two indexing reads of 8 cycles each. See "MiSeq System User Guide" (part #15027617).
- 7. Store diluted libraries and mixtures at -20°C for long-term storage.

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Preparing a Sample Sheet for Sequencing

TIP: Prepare the sample sheet prior to loading the sequencing reagent cartridge. If an error has been made during the indexing PCR where samples have the same indices, it can be remedied before loading the samples on the sequencer.

The available Pillar indexing primers and their barcode sequences are listed in Appendix A. For the i5 indexing primers, indexing on the NextSeq requires the reverse complement of the barcode sequence. The correct barcode sequences for sequencing on the MiSeq and the NextSeq are provided in Appendix A. Additionally, the Pillar sample sheet generator will automatically populate the correct barcode sequence when the indexing primer is selected.

In Appendix A, note that indexing primers highlighted in yellow have the same barcode sequences as Illumina TruSeq Custom Amplicon (TSCA) indices.

In the Pillar sample sheet generator, prepare a sample sheet that contains the information for the samples that are being loaded. Ensure that the appropriate sample sheet is being made for the sequencer used.

- 1. Open the Pillar sample sheet generator and enter user input in the shaded cells. Cells that are shaded blue are required and cells that are shaded grey are optional.
- 2. Enter the "Sample_ID" for each sample. Each Sample_ID must be unique and contain only alphanumeric characters, dashes (-), and underscores (_). All other characters are not allowed. To check that the Sample_ID meets all requirements click "Reset Sample_ID color" and then click "Check Sample_ID".
- 3. If text is green, the Sample_ID is acceptable. If text is red, Sample_ID is not acceptable. Change Sample_ID accordingly and repeat step 2 until all text is green.
- 4. Next, enter indices into appropriate fields. Index sequences will be populated once the index_ID is entered.
- 5. Check that the index combination for each sample is unique. If "Check_index_uniqueness" column is green, then all index combinations are unique. If the column is red for a sample, then the index combination is not unique. Do not load samples together in the same run that have the same index combination.
- 6. Once all requirements for the sample sheet are met, export the sample sheet as a comma-separated values (.csv) file by clicking "Export".

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9. Methylation Calling with PiVAT®

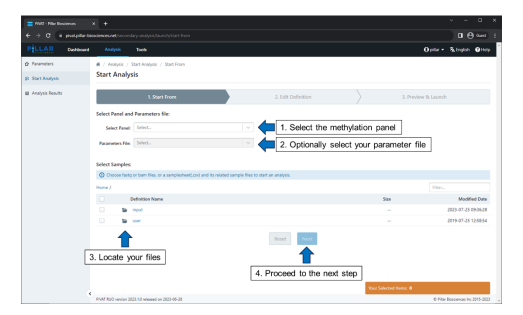
The oncoRevealTM 4 Gene Methylation Panel data analysis is performed using the methylation module in Pillar's secondary analysis pipeline, PiVAT®. This module accurately identifies methylation levels at targeted CpG sites covered by the oncoRevealTM 4 Gene Methylation Panel. The output of the methylation analysis is provided in a Microsoft Excel file format, allowing flexibility in post-analysis data reporting. For example, final methylation values can be adjusted in the Excel output file based on the expected tumor content of the sequenced samples.

The following section provides a walkthrough explaining how to initialize the analysis and how to understand the output format generated by PiVAT.

9.1 Starting your PiVAT Run

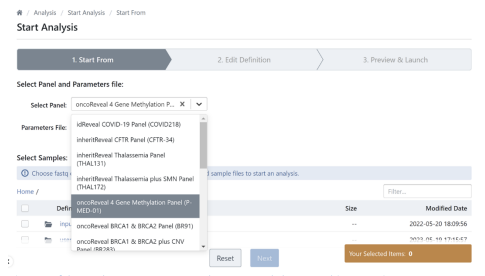
Identify negative control samples in PiVAT using the following steps:

1. Set up the analysis run by selecting the correct panel and FASTQ output.



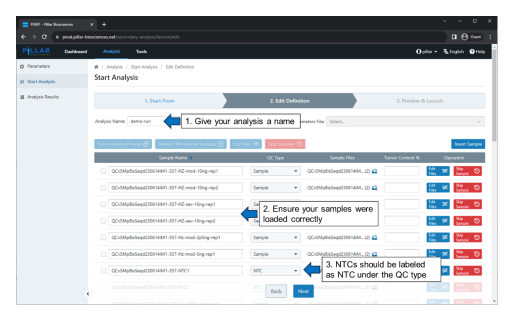
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oncoReveal™ 4 Gene Methylation Panel User Manual



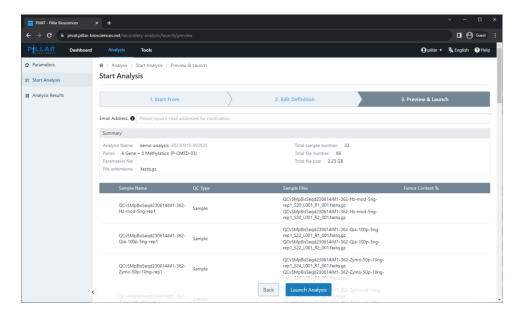
The name of this panel in PiVAT is "oncoReveal™ 4 Gene Methylation Panel (P-MED-01)"

2. Input an appropriate name for the analysis run and verify that the samples have been input correctly.



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 Verify the run setup. Remember to scroll all the way down to confirm that the negative control samples are reflected appropriately.



4. Once the run setup is confirmed, launch the analysis.

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9.2 Output Format in PiVAT

PiVAT reports methylation levels as percentages. It is recommended that users make inferences about differences in methylation levels based on existing literature and on their own prior testing and controls.

The output from PiVAT will be exported in a Microsoft Excel file containing the following information:

- 1. The methylation levels, computed at 3 levels: per-site, per-amplicon, and per-gene
- 2. Bisulfite conversion rates
- 3. Overall sequencing and mapping statistics
- 4. Run-specific parameters

The information containing in the Excel file is organized in multiple sheets:

Sheet Name in Excel Output File	Description
Mathylation Summary By Sample	High-level summary sheet that contains samples in each row and methylation summary by gene and amplicon as columns.
Methylation Summary By Sample	The methylation summary consists of mean, standard deviation, peramplicon coverage, and coefficient of variation as a percentage.
Methylation Site Type	Methylation and bisulfite conversion levels at each of the targeted positions across each sample.
Methylation Summary By Site	A variation of the "Methylation Site Type" sheet, wherein the table is transformed to display CpG sites as rows and samples as columns, to make it easy to compare per-site methylation levels between samples.
Bisulfite Conversion Rates	A two-column table with sample and its corresponding bisulfite conversion values (expressed as a percentage).
	Overall sequencing and mapping statistics.
Overall Stats	For a more detailed description of this sheet, see the <i>PiVAT® User Manual</i> (Doc. No. UM-0073).
Segment Coverage	Per-amplicon coverage (rows) across all samples (columns). Mean, standard deviation, and minimum coverages are reported.
Run Parameters	Parameters used within the PiVAT run, recorded for audit trail purposes.

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9.3 Example PiVAT Output

Methylation Summary By Sample

The example below is focused on a single amplicon and gene.

A	В	c	D	E	F	G	Н	1	J	к
Sample_ID	Sample Ty pe	Methylati on_Mean :BRCA1		Coverage _Mean:Bi sSeq04.B RCA1-1		Methylati on_Mean :BisSeq04 .BRCA1-1		STDEV:Bi sSeq04.B RCA1-1		%CV:BisS eq04.BRC A1-1
Sample1	Sample	0.66		635.29		0.67		0.23		
Sample2	Sample	93.95		883.71		96.88		1.55		0.02
Sample3	Sample	0.5		469		0.48		0.55		

NOTE: In the provided example, the data in some columns have been removed for brevity and for visual clarity. Actual output will contain data for all genes and amplicons covered in this panel.

Column Label	Description							
Sample_ID	Sample name							
SampleType	Sample type (e.g., sample, positive control, or no-template control)							
	Mean methylation level							
Methylation_Mean	Gene-specific methylation mean is represented in Column C in the example above.							
	Amplicon-specific methylation mean is represented in Column G in the example above.							
Coverage_Mean	Mean amplicon coverage							
STDEV	Standard deviation of the mean							
%CV	Coefficient of specification for a specific amplicon This column will be empty if the mean methylation level is below 5%.							

Methylation Site Type

This provides site-by-site methylation information for each sample.

⊿ A	В	C	D	E	F	G	н	1	J	K	L	М	N	0	P	Q	R	S	Т	U	V	w	x
Sample_I	SampleTy	Chromos	Position	REF_(Un	ALT	Location	Amplicon	Gene_Sy	Methylati	Methylat	Methylat	Methylat	Methylat	Conseque	Gene_ID	Feature	All_Featu	HGVSC	Exon	Intron	Strand	Repeat	Methylati
D	pe	ome		methylat			_ID	mbol	on_Rate_	ed_Base_	ed+Unme	ed_Base_	ed_Base_	nce			res						on_Site_T
				ed)					(%)	Coverage	thylated_	Quality	Direction										уре
											Total_Cou		_Ratio										
											nt												
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis04.tr	BRCA2	1.05184	14	1331	33	0.04F/R	upstream_	675	NM_0000	NM_0000	c452			1	4G+1	CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis04.tr	BRCA2	1.57068	21	1337	37	0.0F/R	upstream_	675	NM_0000	NM_0000	c410			1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis04.tr	BRCA2	0.81967	11	1342	36	0.0F/R	upstream_	675	NM_0000	NM_0000	c383			1	5T	CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis04.tr	BRCA2	0.9687	13	1342	38	0.0F/R	upstream_	675	NM_0000	NM_0000	c374			1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis04.tr	BRCA2	0.59791	8	1338	37	O.OF/R	upstream_	675	NM_0000	NM_0000	c360			1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis05-D	BRCA2	0.37688	3	796	38	0.0F/R	5_prime_U	675	NM_0000	NM_0000	c95	1/27		1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis05-D	BRCA2	1.00756	8	794	37	O.OF/R	5_prime_U	675	NM_0000	NM_0000	c92	1/27		1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis05-D	BRCA2	0.50314	4	795	36	0.0F/R	5_prime_U	675	NM_0000	NM_0000	c90	1/27		1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis05-D	BRCA2	0.75567	6	794	39	0.0F/R	5_prime_U	675	NM_0000	NM_0000	c70	1/27		1	4T	CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis06.tr	BRCA2	1.48515	9	606	37	0.0F/R	intron_var	675	NM_0000	NM_0000	c41+41		1/26	1	6T	CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis06.tr	BRCA2	1.32013	8	606	36	0.0F/R	intron_var	675	NM_0000	NM_0000	c41+57		1/26	1		CpG Site
Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis06.tr	BRCA2	0.66116	4	605	36	O.OF/R	intron_var	675	NM_0000	NM_0000	c41+62		1/26	1	5G-1	CpG Site
4 Sample1	Sample	chr13	3.3E+07	T	C	chr13:328	N-Bis06.tr	BRCA2	0.99174	6	605	37	0.0F/R	intron_var	675	NM_0000	NM_0000	c41+93		1/26	1		CpG Site
Sample1	Sample	chr14	1E+08	T	C	chr14:104	XRCC3.inti	XRCC3	96.7773	991	1024	35	O.OF/R	intron var	7517	NM 0011	NM 0011	c.406+139	95	6/9	-1		CpG Site
Sample1	Sample	chr14	1E+08	T	C	chr14:104	XRCC3.int	XRCC3	93.0528	951	1022	34	O.OF/R	intron_var	7517	NM_0011	NM_0011	c.406+138	38	6/9	-1		CpG Site
7 Sample1	Sample	chr14	1E+08	T	C	chr14:104	XRCC3.inti	XRCC3	93.8656	964	1027	34	0.0F/R	intron var	7517	NM 0011	NM 0011	c.406+136	51	6/9	-1		CpG Site
8 Sample1	Sample	chr14	1E+08	Т	C	chr14:104	XRCC3.Inti	XRCC3	97.0968	301	310	36	O.OF/R	intron_var	7517	NM_0011	NM_0011	c.406+169	90	6/9	-1		CpG Site
9 Sample1	Sample	chr14	1E+08	Т	C	chr14:104	XRCC3.Inti	XRCC3	97.1154	303	312	36	0.0F/R	intron_var	7517	NM_0011	NM_0011	c.406+167	74	6/9	-1		CpG Site
0 Sample1	Sample	chr14	1E+08	Т	С	chr14:104	XRCC3.Inti	XRCC3	96.4744	301	312	36	0.0F/R	intron var	7517	NM 0011	NM 0011	c.406+166	53	6/9	-1		CpG Site

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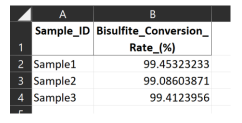
Methylation Summary By Site

This provides an alternate view of the data also shown in the previous sheet, Methylation Site Type.

Α	В	С	D	E	F	G	н	- 1	J	K	L	M	N	0	Р	Q	R	S
Chromos	Position	REF_(Un	ALT	Location	Amplicon	Gene_Sy	Consequ	Gene_ID	Feature	All_Featu	HGVSC	Exon	Intron	Strand	Repeat	Sample1	Sample2	Sample3
ome		methylat ed)			_ID	mbol	ence			res								
chr13	3.3E+07	Т	С	chr13:328	N-Bis04.tr	BRCA2	upstream_	675	NM_0000	NM_0000	c452			1	4G+1	0.69159	2.10809	2.35294
chr13	3.3E+07	Т	С	chr13:328	N-Bis04.tr	BRCA2	upstream_	675	NM_0000	NM_0000	c410			1		0.21622	1.07383	0.2851
chr13	3.3E+07	Т	С	chr13:328	N-Bis04.tr	BRCA2	upstream	675	NM_0000	NM_0000	c383			1	5T	0.64823	0.74727	0.17813
chr13	3.3E+07	Т	С	chr13:328	N-Bis04.tr	BRCA2	upstream	675	NM_0000	NM_0000	c374			1		1.14545	1.26558	0.10688
chr13	3.3E+07	Т	С	chr13:328	N-Bis04.tr	BRCA2	upstream_	675	NM_0000	NM_0000	c360			1		0.3027	0.17255	0.14255
chr13	3.3E+07	Т	С	chr13:328	N-Bis05-D	BRCA2	5_prime_l	675	NM_0000	NM_0000	c95	1/27		1		0.10111	0.52061	0.9848
chr13	3.3E+07	Т	С	chr13:328	N-Bis05-D	BRCA2	5_prime_l	675	NM_0000	NM_0000	c92	1/27		1		0.77519	0.91067	1.79756
chr13	3.3E+07	Т	С	chr13:328	N-Bis05-D	BRCA2	5_prime_l	675	NM_0000	NM_0000	c90	1/27		1		0.30334	0.9974	0.81353
chr13	3.3E+07	Т	С	chr13:328	N-Bis05-D	BRCA2	5_prime_l	675	NM_0000	NM_0000	c70	1/27		1	4T	0.37087	0.43384	1.99058
chr13	3.3E+07	Т	С	chr13:328	N-Bis06.tr	BRCA2	intron_va	675	NM_0000	NM_0000	c41+41		1/26	1	6T	2.80095	0.2313	0.41797
chr13	3.3E+07	Т	С	chr13:328	N-Bis06.tr	BRCA2	intron_va	675	NM_0000	NM_0000	c41+57		1/26	1		1.90817	0.69391	0.31348
chr13	3.3E+07	Т	С	chr13:328	N-Bis06.tr	BRCA2	intron_va	675	NM_0000	NM_0000	c41+62		1/26	1	5G-1	0.35842	0.15456	4.49791
chr13	3.3E+07	Т	С	chr13:328	N-Bis06.tr	BRCA2	intron_va	675	NM_0000	NM_0000	c41+93		1/26	1		0.65672	0.38551	0.10449
chr14	1E+08	Т	С	chr14:104	XRCC3.int	XRCC3	intron_va	7517	NM_0011	NM_0011	c.406+13	95	6/9	-1		95.6714	96.3998	94.2472
chr14	1E+08	Т	С	chr14:104	XRCC3.int	XRCC3	intron_vai	7517	NM_0011	NM_0011	c.406+13	88	6/9	-1		90.8316	93.4458	88.0229

Bisulfite Conversion Rates

This provides the bisulfite conversion rate as a percentage. The example shown is the bisulfite conversion rate across three samples.



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Overall Stats

This provides detailed sequencing and mapping results from the run analyzed.

A	В	С	D
1 Stat	Sample1	Sample2	Sample3
2 Total Reads	42124	38994	26810
3 Overall:Q=30	89.04	84.47	87.08
4 Overall:Q=20	90.53	87.43	88.75
5 Properly Paired Reads	41900	38656	26718
6 Properly Paired Read (%)	99.47	99.13	99.66
7 Mapped Reads	41968	38824	26754
8 Mapping Rate (%)	99.63	99.56	99.79
9 On Target Reads	35605	36874	22882
10 On Target Rate (%)	84.84	94.98	85.53
11 Insert Size Mean	109	109	110
12 Insert Size Median	117	117	117
13 Insert Size Std Dev	31	38	31
14			
15 Stat	Sample1	Sample2	Sample3
16 Coverage_Mean	1348	1285	881
17 STDEV	876	588	884
18 Coverage_Median	1310	952	929
19 Coverage_Max	3770	2569	3394
20 Coverage_Min	308	500	174
Total_Number_Of_Reads	35624	36932	22886
Total_Valid_Reads	33294	30850	21216
23 On_Target_Ratio	0.93459	0.83532	0.92703

Segment Coverage

This provides the amplicon mean coverage, standard deviation, and minimum coverage across all samples.

	Α	В	С	D	Е	F	G	Н	ı	J	K	L	M
Tar	rget_N	Region	Segment	GC_Cont	Coverage	Coverage	Coverage	STDEV:Sa	STDEV:Sa	STDEV:Sa	Coverage	Coverage	Coverage
a	ame		_Size	ent_(%)	_Mean:S	_Mean:S	_Mean:S	mple1	mple2	mple3	_Min:Sa	_Min:Sa	_Min:Sa
1					ample1	ample2	ample3				mple1	mple2	mple3
2 N-B	3is04.tr	chr13:328	96	31.46853	1341	2035	1250	2	12	2	1332	1985	1234
3 N -B	3is05-D	chr13:328	44	32.94118	796	936	591	1	1	1	792	933	588
4 N-B	3is06.tr	chr13:328	99	34.69388	605	893	179	1	4	1	600	866	174
5 XRC	CC3.Inti	chr14:104	56	43.29897	312	744	268	1	2	0	308	734	242
6 XRC	CC3.intr	chr14:104	42	27.08333	1031	1435	351	3	4	0	1022	1423	348
7 XRC	CC3.5U	chr14:104	74	34.61538	1150	1538	203	2	2	0	1141	1533	198
8 N-B	3is02.tr	chr14:104	50	28.125	2767	1337	1852	6	1	5	2735	1332	1827
9 N-B	3is01.tr	chr14:104	39	20	1451	952	964	1	1	1	1449	950	961
10 Biss	Seq04.E	chr17:412	83	26.31579	644	887	477	1	2	1	640	878	473
11 Biss	Seq05.E	chr17:412	66	25.64103	3762	2564	3386	13	7	11	3702	2540	3332
12 Biss	Seq02.F	chr17:567	112	33.11688	1311	516	972	1	8	1	1308	500	968
13 Biss	Seq03.F	chr17:567	103	38.51351	1468	1604	930	2	5	2	1460	1589	921

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10. Troubleshooting Library Preparation

Low Yield of Gene-Specific Product

Potential Cause	Solution
DNA quantity or quality	The recommended input for the assay is 5 – 20 ng of bisulfite converted DNA. Higher quantities may be necessary for low- or poor-quality samples.
Improper cycling	Check that the cycling protocol performed is the appropriate protocol for gene- specific amplification.

Low Indexing Efficiency

Potential Cause	Solution
	Incomplete AMPure purification or loss of gene-specific product will affect the indexing PCR reaction. The purified gene-specific product can be checked on an agarose gel to ensure the product was not lost, and that the clean-up was sufficient to remove excess primers.
Improper AMPure purification	The AMPure bead ratio and ethanol concentration affect the PCR clean-up. Ensure that the correct AMPure bead concentration was used for clean-up, and fresh 70% ethanol was used for the wash.
	Leftover ethanol from the wash steps can hinder the PCR reaction. Remove as much of the ethanol during the final wash step with a pipette and dry the beads to ensure the residual ethanol has evaporated.

Low Yield of Final Library

Potential Cause	Solution
DNA quantity or quality	The recommended input for the assay is 5 – 20 ng of bisulfite converted DNA. Higher quantities may be necessary for low- or poor-quality samples.
	Run the product from the gene-specific PCR on an agarose gel to check the yield.
Improper AMPure purification	Incomplete AMPure purification or loss of product will affect the final yield. The purified product can be checked on an agarose gel to ensure that the product was not lost during PCR clean-up.

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The AMPure bead ratio and ethanol concentration affect the PCR clean-up. Ensure that the correct AMPure bead concentration was used for clean-up, and fresh 70% ethanol was used for the wash.

Amplicon Contamination in No-Template Control

Potential Cause	Solution	
	Make sure to change tips between samples and avoid waving tips over tubes or plates.	
	Poor sealing or residual liquid in tips can cause contamination of nearby samples. If possible, leave adjacent wells empty between samples.	
Cross-contamination	Workspaces and equipment for pre-PCR and post-PCR should be separated to prevent amplicon contamination.	
	Periodically clean the workspace, floor, equipment, and instrumentation with a laboratory cleaning solution to break down amplicons on surfaces.	
	Recommended cleaning solutions are 10% bleach, 70% isopropanol, or 70% ethanol.	

Sequencing Performance

Potential Cause	Solution	
Improper normalization and pooling of libraries	Confirm that the appropriate loading concentration was used for the applicable sequencing instrument.	
	Check the 5 nM (or 1 nM if sequencing on the MiniSeq) library mix using Qubit or RT-PCR. Dilute the denatured library mix as needed to adjust for the difference in concentration.	
Improper library quantitation	Improper library quantitation may result in artificially high or low yields, which affects downstream normalization. Re-quantitate the final libraries and/or the normalized libraries to check for the expected values.	

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Improper AMPure purification	Changing the ratio of AMPure beads affects the purification of the products. Notably, the presence of primer dimers can cause an underestimation of total quantity, causing over-clustering. The final libraries can be checked on an agarose gel for the proper product size and presence of primer dimers.
	The AMPure bead ratio and ethanol concentration affect the PCR clean-up. Ensure that the correct AMPure bead concentration was used for clean-up, and fresh 70% ethanol was used for the wash.

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11. Appendix A: Index Sequences

Indexing primers highlighted in yellow have the same barcode sequences as Illumina TruSeq Custom Amplicon (TSCA) indices.

i7 Index Sequences		
Index ID	Index Barcode Sequence	
Рі701	ATTACTCG	
Pi ₇₀₂	TCCGGAGA	
Pi ₇ 03	CGCTCATT	
Pi ₇₀₄	GAGATTCC	
Pi ₇ 05	ATTCAGAA	
Pi ₇ 06	GAATTCGT	
Pi ₇ 0 ₇	CTGAAGCT	
Pi ₇ 08	TAATGCGC	
Pi ₇₀₉	ATCACGAC	
Pi ₇ 10	ACAGTGGT	
Pi ₇₁₁	CAGATCCA	
Pi ₇₁₂	ACAAACGG	
Pi ₇₁₃	GAAACCCA	
Pi ₇₁₄	TGTGACCA	
Pi ₇₁₅	AGGGTCAA	
Pi ₇ 16	AGGAGTGG	

i5 Index Sequences				
Index ID	Index Barcode Sequence for MiSeq	Index Barcode Sequence for NextSeq		
Pi501	TATAGCCT	AGGCTATA		
Pi502	ATAGAGGC	GCCTCTAT		
Pi503	CCTATCCT	AGGATAGG		
Pi504	GGCTCTGA	TCAGAGCC		
Pi505	AGGCGAAG	CTTCGCCT		
Pi506	TAATCTTA	TAAGATTA		
Pi507	CAGGACGT	ACGTCCTG		
Pi508	GTACTGAC	GTCAGTAC		
Pi509	TGAACCTT	AAGGTTCA		
Pi510	TGCTAAGT	ACTTAGCA		
Pi511	TGTTCTCT	AGAGAACA		
Pi512	TAAGACAC	GTGTCTTA		
Pi513	CTAATCGA	TCGATTAG		
Pi514	CTAGAACA	TGTTCTAG		
Pi515	TAAGTTCC	GGAACTTA		
Pi516	TAGACCTA	TAGGTCTA		

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