

DRAGEN RNA for MiSeq i100 v1.1.7

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1 App Highlights

The DRAGEN RNA Pipeline includes the following functional components:

- Decompression of input BCL data
- FASTQ Generation
- FASTQ Compression (DRAGEN ORA or Gzip)
- Mapping & Alignment (includes sorting)
- BAM/CRAM Compression (Optional)
- Gene Fusion Detection (Optional, if the full pipeline is selected)
- Whole Transcriptome Gene Expression (Quantification of Transcripts) (Optional, if the full pipeline is selected)
- Differential Expression (Optional)

The following inputs are required:

- BCL Data generated from the MiSeq i100
- Sample Sheet

The following inputs are optional:

- BedFile (for targeted regions)
- RnaGeneAnnotationFile (see following note)

An RnaGeneAnnotationFile is packaged with each Illumina-provided human reference genome. If an RnaGeneAnnotationFile is provided by the customer on run set-up, it will be used instead of the file packaged with the genome. If not available either with the Illumina-provided genome or by the customer directly, then Gene Fusion and RNA quantification steps will be skipped. An RnaGeneAnnotationFile must be provided if Differential Expression is enabled.

See Pipeline Configuration for Sample Sheet parameters

1.1 DRAGEN core RNA Pipeline info on Gitbook:

<https://help.dragen.illumina.com/product-guide/dragen-v4.4/dragen-rna-pipeline>

2 Pipeline Configuration

DragenRna_Settings

Parameter	Required	Description
SoftwareVersion	Yes	The version of the DRAGEN software used to process the DragenRna pipeline, including conversion to FASTQ, specified using all three integers included in the DRAGEN version name. For example 4.1.5.
AppVersion	Yes	The version of the workflow-specific application (i.e., DRAGEN RNA), using all three integers included in the version name. For example, 1.0.0.
MapAlignOutFormat	Yes	Formatting of the output files. Accepted values are bam, cram, or none. Selecting none produces no map/align output. Selecting none is not allowed if RnaPipelineMode is set to FulPipeline for any sample.
KeepFastQ	Yes	Select whether FASTQs are saved (true) or discarded (false).
DifferentialExpressionEnable	No	Accepted values are true or false. If DifferentialExpressionEnable is true, then RnaPipelineMode must be FullPipeline

DragenRna_Data

Parameter	Required	Description
ReferenceGenomeDir	Yes	Genome name as an alphanumeric string with "_", "-", or ".". Can be placed in either settings or data section. For example hg38-alt_masked.cnv.hla.methylated_combined.rna-10-r4.0-2.
BedFile	No	BED file to be used for targeted regions in text (*.txt) or gzip (*.gz) format. Must include the prefix DragenAmplicon/ before the BED file name. Alphanumeric string with "_", "-", or "." with no spaces allowed.
RnaGeneAnnotationFile	No	Genotype reference file. Alphanumeric string with "_", "-", or "." with no spaces allowed. If DifferentialExpressionEnable is True, the GTF file must be provided by the user or included with the reference genome.
RnaPipelineMode	Yes	Accepted values are MapAlign or FullPipeline. The full pipeline option includes quantification and fusion detection.
DownSampleNumReads	No	Specifies the number of fragments to downsample to. For paired-end sequencing, the number of reads at the down-sampling output will be twice the number of fragments specified. Accepted values are integers. If DownSampleNumReads is placed in the Data section, and downsampling is not desired for that sample, a value of na must be used.
Sample_ID	Yes	See description in the BCL Convert section.

Comparison1	No	Accepted values are control, comparison, or na. Use only if DifferentialExpressionEnable is true. If RNAPipelineMode is MapAlign, this value must be na. Must have at least 2 control and 2 comparison samples. May have a max of 15 control or comparison samples. All control and comparison samples must have FullPipeline for RnaPipelineMode value and have the same ReferenceGenomeDir and RnaGeneAnnotationFile values.
Comparison2	No	
Comparison3	No	
Comparison4	No	
Comparison5	No	

3 Modes of Operations

3.1 Cloud and Local Execution:

DRAGEN can be scheduled to operate in the following modes:

- Cloud:
 - Run planning:
 - Occurs in the Cloud via the Cloud User Interface (UI), or alternatively, sample sheets can be imported into the Cloud UI
 - Sequencing execution:
 - Runs are identified on the sequencer, and are executed locally on the instrument
 - DRAGEN Execution:
 - Occurs in the Cloud automatically after sequencing is complete
- Local:
 - Run planning:
 - Occurs on the instrument via the Instrument UI, or alternatively, sample sheets can be imported into the Instrument UI
 - Sequencing execution:
 - Runs are executed locally on the instrument
 - DRAGEN Execution:
 - Occurs locally on the instrument, automatically after sequencing is complete

3.2 Requeues:

After a sequencing run is complete, it may be desirable to run or re-run DRAGEN with, for example, an updated configuration. Requeues can occur on the instrument or in the Cloud. For on-instrument requeues, the following pre-requisites are necessary in Local Mode:

- The BCL's must be stored on the instrument
- The system must be idle

4 Output Files

4.1 Summary

- Aligned reads - BAM/CRAM format
- Gene counts - BED format
- RNA fusions - CSV/VCF format
- Reports - HTML

4.2 Toplevel DRAGEN

<run_id>/Analysis/<no>/inputs

- <SampleSheet>.csv

<run_id>/Analysis/<no>/Data

- Secondary_Analysis_Complete.txt
- summary
 - <x.y.z> (Note: DRAGEN version)
 - highlevel_summary.json
 - detailed_summary.json
- AggregateReports
 - report.html
 - report_files
 - *Links to workflow level reports*
- Demux
 - AggregateReports
 - *Links to lower-level reports*
 - Demultiplex_Stats.csv
 - *other stats*
- Dragen<Workflow>
 - Data (including FASTQ) for samples configured for the workflow (see below)
- RunInstrumentAnalyticsMetrics
- logs
 - *high-level logs*

4.3 Workflow Level Output

<run_id>/Analysis/<no>/Data/DragenRna

- Dragen<workflow>

- AggregateReports
 - *Links to workflow specific sample level reports*
- fastq (or ora_fastq)
 - `<sample_ID>.S0_Rm_001.fastq.gz` or `*.fastq.ora` (m=1-2)
 - *Additional samples*
 - Reports
 - *Adapter_Metrics.csv*
 - *Quality_Metrics.csv*
 - *Additional metrics files*
- `<sample_ID>`
 - `<workflow>_seq`
 - `<workflow>-specific logs`
 - logs
 - *Sample specific logs*
- logs
 - *Workflow specific logs*

5 Reference Genome Usage Guidelines

DRAGEN supports two types of genome hash tables: linear and pangenome. Pangenome reference genomes extend the reference genomes with alternative variant paths from a sample cohort used to construct the pangenome reference.

5.1 Germline, Enrichment Germline Workflows

- Use pangenome reference genomes.
- Use the hg38 assembly. The recommended reference genome is [Homo sapiens \[1000 Genomes\] hg38 v5 Multigenome](#)
- If not using hg38, the [Homo sapiens \[UCSC\] hg19 v5 Multigenome](#) or [Homo sapiens \[NCBI\] hs37d5 v5 Multigenome](#) are recommended.
- Linear reference is supported, but has reduced accuracy

5.2 Enrichment Somatic, Somatic, and RNA Workflows

- Use linear reference genomes.
- Use the [Homo sapiens \[1000 Genomes\] hg38 v5](#) reference genome or [Homo sapiens \[NCBI\] hs37d5 v5](#) reference genome.

5.3 Reference Genome Version Compatibility with DRAGEN Versions

	HT10	HT11
4.3.13		
4.4.6		

6 Available Resource Files

6.1 Available Reference Genomes:

A reference genome is an input to a DRAGEN pipeline. Reference genomes are available at:

- https://support.illumina.com/sequencing/sequencing_instruments/miseq-i100-plus/product-files.html

and are summarized below.

DRAGEN Reference Genomes Provided with the Instrument (DRAGEN 4.4.6)

Directory Name	Display Name (Species [Source] Assembly)	DN A	RN A	GT F	CN V	Graph	Met h
hg19-alt_masked.cnv.hla.methyl_cg.methylated_combined.ma-11-r5.0-2	Homo sapiens [UCSC] hg19 v5	Yes	Yes	Yes	Yes	No	No
hg38-alt_masked.cnv.hla.methyl_cg.methylated_combined.ma-11-r5.0-2	Homo sapiens [1000 Genomes] hg38 v5	Yes	Yes	Yes	Yes	No	No
mm10-methyl_cg.methylated_combined.ma-11-r5.0-2	Mus musculus [UCSC] mm10 v5	Yes	Yes	Yes	No	No	No
m6-ma-11-r5.0-2	Rattus norvegicus [UCSC] m6 v5	Yes	Yes	Yes	No	No	No
Bacillus_cereus_ATCC_10987-cnv.ma-11-r5.0-2	B. cereus [NCBI] ATCC_10987 v5	Yes	Yes	No	No	No	No
Rhodobacter_sphaeroides_2.4.1_12905-cnv.ma.seed_len19-11-r5.0-2	R.sphaeroides [NCBI] 2.4.1 v5	Yes	Yes	No	No	No	No
eschColi_K12_1-cnv.rna-11-r5.0-2	E. coli [NCBI] K12 MG1655 v5	Yes	Yes	No	No	No	No
phix-cnv.ma-11-r5.0-2	Microvirus Escherichia virus	Yes	Yes	No	No	No	No

	phiX174 [NCBI] v5						
Cereibacter_sphaeroides_ATCC_17023-rna-11-r5.0-2	C. sphaeroides [ATCC] 2.4.1 v5	Yes	Yes	No	No	No	No
Staphylococcus_aureus_NCTC_8325-rna-11-r5.0-2	S. aureus [NCBI] v5	Yes	Yes	No	No	No	No
Mycobacterium_tuberculosis_H37Rv-rna-11-r5	M. tuberculosis [NCBI] v5	Yes	Yes	No	No	No	No
Streptococcus_pneumoniae_Hu17-rna-11-r5.0-2	S. pneumoniae [NCBI] v5	Yes	Yes	No	No	No	No
Salmonella_enterica_LT2-rna-11-r5.0-2	S. enterica [NCBI] v5	Yes	Yes	No	No	No	No
Shigella_sonnei_SE6-1-rna-11-r5.0-2	S. sonnei [NCBI] v5	Yes	Yes	No	No	No	No
Saccharomyces_cerevisiae_S288C-rna-11-r5.0-2	S. cerevisiae [NCBI] v5	Yes	Yes	No	No	No	No
Human_immunodeficiency_virus_1-rna-11-r5.0-2	HIV 1 [NCBI] v5	Yes	Yes	No	No	No	No
Influenza_A_virus_H1N1-rna-11-r5.0-2	Influenza A H1N1 [NCBI] v5	Yes	Yes	No	No	No	No
Influenza_A_virus_H3N2-rna-11-r5.0-2	Influenza A H3N2 [NCBI] v5	Yes	Yes	No	No	No	No
Influenza_A_virus_H5N1-rna-11-r5.0-2	Influenza A H5N1 [NCBI] v5	Yes	Yes	No	No	No	No
SARS_coronavirus_2-rna-11-r5.0-2	SARS-CoV2 [NCBI] v5	Yes	Yes	No	No	No	No
Drosophila_melanogaster_Fruitfly-rna-11-r5.0-2	D. melanogaster [NCBI] v5	Yes	Yes	No	No	No	No

6.2 Available Non-Reference Genome Resource Files:

DRAGEN requires resource files to function correctly and achieve the optimum performance for certain workflows. All resource files are available for download at the Illumina DRAGEN Product Files support site at:

- https://support.illumina.com/sequencing/sequencing_software/dragen-bio-it-platform/product_files.html

These resources can be imported to the instrument.

7 How To Generate Custom Resource Files

7.1 Reference Genomes:

Custom reference genomes can be imported to the instrument.

- Use the [Reference Builder \(Instruments\) BaseSpace App](#), which accepts a fasta file and generates a .gz file.
- Reference Builder App version 2.0.0 generates reference genomes for DRAGEN versions 4.1.* (v8 hash table format)
- Reference Builder App version 3.0.0 generates reference genomes for DRAGEN versions 4.3.* (v10 hash table format)

8 How To Import Resource Files

You can add and delete reference files on the Reference Files tab in the Resources settings screen. The Reference Files tab displays the reference file name, file type, file description, and the number of related reference genomes.

8.1 Import Reference Files

- Make sure that there are no sequencing runs or on-instrument secondary analyses in progress.
- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select Resources.
- On the Reference Files tab, select Import reference file.
- Navigate to the reference file, and then select Open.
- [Optional] Enter a description for the reference file.
- Select a file type from the drop-down list.

9 How To Install DRAGEN and DRAGEN Applications

9.1 Install DRAGEN Versions:

- When a new DRAGEN version is available, download the DRAGEN installer (*.ires) from the MiSeq i100 Series support page. Save the installer locally or to a network drive.
- Make sure that there are no sequencing runs or on-instrument secondary analysis in progress.
- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select DRAGEN.
- On the Versions tab, select Install version.
- Navigate to the installer, and then select Open.
- Select Install. A message indicates if the installation was successful or failed

9.2 Uninstall DRAGEN Versions:

- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select DRAGEN.
- To uninstall a previous DRAGEN version, do as follows.
 - a. On the Versions tab, select the ellipsis icon in the Actions column.
 - b. Select Uninstall.
 - c. Select Yes, uninstall.
- To uninstall the latest DRAGEN version, do as follows.
 - a. On the Versions tab, select the ellipsis icon in the Actions column.
 - b. Select Uninstall all.
 - c. Select Yes, uninstall all

9.3 Application Installation:

- Download the application (*.iapp) from the MiSeq i100 Series support page. Save the installer to a network drive.
- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select Applications.
- Select Install application.

- Navigate to the application file, and then select Open. After the file uploads, information about the application displays.
- Select Install. After the application installs, you can review the application configuration.

9.4 View Application Settings:

The DRAGEN application provides a default library prep kit, index adapter kit, read information, index information, and permissions. Some applications also provide settings and configuration for secondary analysis.

- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select Applications.
- Select the application to view. After you install an application, the Configuration screen opens automatically.
- Edit any of the following information:
 - Library prep kits
 - Index adapter kits
 - Index reads
 - Read type
 - Index lengths
 - Read length
- Select Save.

9.5 Uninstall Applications

Administrators can uninstall applications.

- Select the instrument icon to open the global navigation menu.
- Select Settings, and then select Applications.
- Select the application to uninstall.
- Select Uninstall.
- Confirm to uninstall the application.

Release History

Revision	Release Reference	Originator	Description of Change
00	1130656	Mark Bilstad	Initial release