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# Sentieon

## Unique Molecular Identifiers

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## 1 Introduction

This document describes using the Sentieon® tools to process next-generation sequence data while taking advantage of molecular barcode information (also called unique molecular indices or UMIs). Molecular barcodes introduce unique tags on the ends of template DNA molecules prior to sequencing to greatly reduce the impact of PCR duplicates and sequencing errors on the variant calling process.

The Sentieon® tools provide functionality for extracting UMI tags from read data and performing barcode-aware consensus generation. This pipeline expects adapter-free barcoded reads as input. The output of the UMI consensus pipeline is a BAM file containing consensus molecules derived from the barcoded read data. These consensus molecules can be used as input to most variant calling software.

## 2 Sentieon® UMI pipeline

### 2.1 Overall UMI pipeline structure

Sentieon® provides two utilities for UMI NGS data processing:

- `umi extract`: UMI tag extraction from unaligned input reads with adapter already removed
- `umi consensus`: Barcode-aware duplicate removal and consensus calling on aligned input

Sentieon® suggests the following typical UMI processing pipeline (Fig. 2.1):

1. UMI tag extraction from unaligned input reads with `umi extract` utility
2. Alignment to the reference genome with Sentieon® `bwa mem`
3. UMI consensus calling with `umi consensus` utility
4. Alignment and sorting of consensus reads to the reference genome with Sentieon® `bwa mem`.

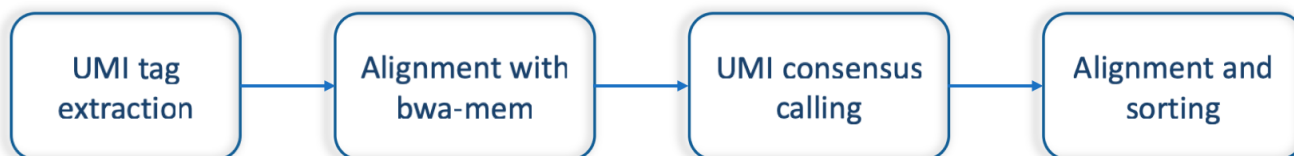


Fig. 2.1: Sentieon® UMI processing pipeline

Below is a code example. Details of each component will be explained in the subsequent sections.

```
sentieon umi extract \  
  8M12S+T,+T \  
  sample_R1.fastq.gz \  
  sample_R2.fastq.gz | \  
sentieon bwa mem \  
  -R "@RG\tID:$GROUP\tSM:$SAMPLE\tLB:$LIBRARY\tPL:$PLATFORM" \  
  -t $NT \  
  -K $BWA_K_SIZE \  
  -p \  
  -C \  
  $REF \  
  - | \  
sentieon umi consensus \  
  -o sample_consensus.fastq.gz  
  
sentieon bwa mem \  
  -R "@RG\tID:$GROUP\tSM:$SAMPLE\tLB:$LIBRARY\tPL:$PLATFORM" \  
  -t $NT \  
  -K $BWA_K_SIZE \  
  -p \  
  -C \  
  $REF \  
  sample_consensus.fastq.gz | \  
sentieon util sort \  
  -i - \  
  -o sample_consensus.bam \  
  --sam2bam --umi_post_process
```

---

## 2.2 Determine the read structure and extract barcode sequences

As a first step, you need to extract the barcode sequences from the input reads. This is performed with Sentieon® `umi extract` command, which extracts the barcode sequence information from the reads and adds it to the read description. As noted earlier, the adapter sequence should be removed from the input reads before running `umi tag` extraction. This can be performed by other third-party tools.

The output of `umi extract` is in FASTQ format with interleaved R1 and R2 reads. By default, the output from the `extract` command will be sent to standard output unless otherwise defined with the option `-o`.

The syntax of the `umi extract` command is as follows:

```
sentieon umi extract [options] read_structure fastq1 [fastq2] [fastq3]
```

Options:

```
-o      Output file (default: stdout)
-d      Turn on duplex mode
--umi_tag    Logic umi tag (default 'XR')
--output_format    Output format FASTQ or SAM (default 'FASTQ')
```

The first argument of `umi extract` command defines the read structure. For paired-end reads, two read structures separated by comma ',' should be specified.

A read structure is defined by `<number><operator>` pairs. The number can be any digit or '+' to indicate the end of the read. Possible operators include:

- **T** The template sequence.
- **B** The cell barcode sequence.
- **M** The molecular barcode sequence.
- **S** A sequence of bases that should be ignored.

The `-d` option is used to extract duplex UMI and label the strand it originates. Duplex UMI extraction requires an identical read structure for both strands.

As an example, the following command demonstrates a single-ended UMI extraction on pair-ended reads. In this case, the first read in the pair contains an 8bp molecular barcode followed by a 12bp spacer and then the template sequence. The second read contains only the template sequence. The paired reads will be interleaved in the output file. Please note that in this example the output is piped to `gzip` to generate compressed FASTQ file. In general, we recommend piping the output directly to the next step (Sentieon® `bwa mem`).

```
sentieon umi extract "8M12S+T,+T" \  
sample_R1.fastq.gz \  
sample_R2.fastq.gz | \  
gzip -c \  
> sample_extracted_pair.fastq.gz
```

The command below demonstrates duplex UMI extraction where both reads contain 4bp molecular barcode followed by template sequence.

```
sentieon umi extract \  
-d \  
"4M+T,4M+T" \  
sample_R1.fastq.gz \  
sample_R2.fastq.gz | \  
gzip -c \  
> sample_extracted_pair.fastq.gz
```

Below is a use case when the UMI sequence is already in a separate FASTQ file `sample_I1.fastq.gz`. When running in this mode, only one additional UMI index read is allowed. The UMI index read should contain no template sequences.

This mode does not allow duplex UMI extraction.

```
sentieon umi extract \  
  "+M,+T,+T" \  
  sample_I1.fastq.gz \  
  sample_R1.fastq.gz \  
  sample_R2.fastq.gz | \  
  gzip -c \  
  > sample_extracted_pair.fastq.gz
```

The command below produces a SAM format output. This format is useful when using the RNA-seq aligner STAR. In this example, the first read in the pair contains a 16bp cell barcode and a 10bp molecular barcode. The second read contains only the template sequence. The output is single-end reads in the SAM format.

```
sentieon umi extract \  
  --output_format SAM \  
  "16B10M+S,+T" \  
  sample_R1.fastq.gz \  
  sample_R2.fastq.gz \  
  > sample_extracted.sam
```

The output of `umi extract` contains additional tags. By default, the output contains CR tag for cell or sample barcode, and XR tag for UMI sequence to be used by `umi consensus`.

Table 2.1: Additional tags generated by `umi extract`

Tags	Meaning
RX	Extracted UMI sequence bases.
XR	UMI tag for grouping in <i>umi consensus</i> .
CR	Cell barcode.

## 2.3 Alignment to the reference genome

The interleaved fastq can be aligned to the reference genome using `bwa mem`. The `-p` option is necessary to specify that the input file is an interleaved fastq while the `-C` option will cause the barcode tags in the fastq description to be appended to the read's SAM record.

```
sentieon bwa mem \  
  -R "@RG\tID:$GROUP\tSM:$SAMPLE\tLB:$LIBRARY\tPL:$PLATFORM" \  
  -t $NT \  
  -K $BWA_K_SIZE \  
  -p \  
  -C \  
  $REF \  
  sample_extracted_pair.fastq.gz > sample_aligned.sam
```

## 2.4 Consensus molecule creation

The next stage of the pipeline is creating consensus molecules from aligned barcode-tagged reads using `Sentieon® umi consensus`.

The syntax of `umi consensus` is as follows:

```
umi consensus [-i input] [options] -o output
```

Options for umi\_consensus:

```
-i, --input          Input file (default: stdin SAM)
-o, --output         Output file
  --input_format     SAM/BAM/CRAM
  --umi_tag          Logic UMI tag (default: 'XR')
  --copy_tags        List of tags to be copied (default: XR,RX,MI,BI,BD,XZ)
  --read_name_prefix Read name prefix (default: 'UMI-')
```

By default, umi consensus will read input from the standard input with SAM format. This can be overridden with --input option that defines the input file, and --input\_format option for the file format other than SAM. Output from umi consensus is an interleaved fastq containing consensus molecules, which can be re-mapped by Sentieon® bwa mem. Below is example:

```
cat sample_aligned.sam | \
  sentieon umi consensus \
  -o sample_consensus.fastq.gz
```

The output of umi consensus generates the following additional tags.

Table 2.2: Output fastq tags from umi consensus

Tags	Meaning
BI/BD	Indel quality scores
MI	A unique label for UMI group where the consensus is derived.
XZ	The number of raw reads within the UMI group where the consensus is derived. For duplex UMI this will contain the number of raw reads from each strand.

## Report consensus reads without the BI/BD tags

By default, umi consensus recalibrates the INDEL error rates and store such information in the BI/BD tags. This modeling step can be turned off by removing the BI/BD tags from the --copy\_tags option.

```
cat sample_aligned.sam | \
  sentieon umi consensus \
  --copy_tags XR,RX,MI,XZ \
  -o sample_consensus.fastq.gz
```

## 2.5 Alignment of consensus reads to the reference genome

The interleaved fastq from umi consensus can be aligned to the reference genome using Sentieon® bwa mem. Similar to the previous alignment, the -p option and the -C option are necessary. Piping the output to Sentieon® util sort creates the output BAM file ready for variant calling. The --umi\_post\_process option is used to instruct the tool to perform the necessary post-processing of the consensus reads.

```
sentieon bwa mem \
  -R "@RG\tID:$GROUP\tSM:$SAMPLE\tLB:$LIBRARY\tPL:$PLATFORM" \
  -t $NT \
  -K $BWA_K_SIZE \
  -p \
  -C \
```

(continues on next page)

```
$REF \
sample_consensus.fastq.gz | \
sentieon util sort \
-i - \
-o sample_consensus.bam \
--sam2bam --umi_post_process
```

## 2.6 Variant calling from consensus reads

The BAM file from UMI consensus pipeline is analysis ready for variant calling. Additional steps of duplicate marking or base quality adjustments should not be performed because the UMI consensus step is essentially a combination of both PCR duplicate removal and base quality recalibration.

Although any somatic caller can be used with the consensus reads, we recommend TNScope® because of its high sensitivity for low-frequency variant detection.

## 2.7 Logs from UMI consensus

The log printed by `umi consensus` contains statistical information on the input reads that can be useful for quality control. The two pieces of information currently available are:

- Group size histogram: Group size is the number of supporting raw reads within a group that call one consensus read. The histogram shows the number of UMI consensus reads that have a specific group size and can be used to detect issues:
  - If the proportion of singletons (UMI consensus with group size = 1) is too high, it will be difficult to accurately model the PCR error rates and improve the quality scores.
  - Conversely, if the average reads per UMI group is too high, it could be an indication that the amount of input DNA in the library preparation was too low, which in turn will lead to low coverage of consensus reads.
- Duplex stat: With duplex UMI tags, `umi consensus` is able to recognize reads originating from each strand of DNA by comparing the UMI tags from both R1 and R2 reads, and identify errors raised during sample preparation. The numbers reported in the log represent the number of consensus reads that are either single strand or duplex, grouped by the following codes:
  - Group code 1: Single-strand consensus with no reads from the complementary UMI group.
  - Group code 2: Duplex consensus with reads from two complementary UMI groups.
  - Group code 3: UMI sequences from R1 and R2 reads are the same. In this case, `umi consensus` relies on the strand of inserts to determine whether the input reads are from the same strand. This group is further divided into:
    - \* Group code 31: Single-strand consensus with all reads coming from the same strand.
    - \* Group code 32: Duplex consensus with reads from both strands of DNA.