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SNV+CNV
 dragen -f -r $\{1\} \setminus
    -1 ${2} -2 ${3} --output-directory ${4} --output-file-prefix ${5} \
    --RGID Illumina_RGID --RGSM ${5} \
    --enable-map-align true \
    --enable-map-align-output true \
    --output-format BAM \
    --enable-duplicate-marking true \
    --enable-sort true \
    --enable-variant-caller true \
    --vc-enable-vcf-output true \
    --enable-vcf-compression true \
    --enable-cnv true \
    --cnv-enable-self-normalization true \
SV
--enable-sv true
repeat
--repeat-genotype-enable true
--repeat-genotype-specs /opt/edico/repeat-specs/hg19_expanded/
gvcf
--vc-emit-ref-confidence GVCF
DRAGEN-ML:typically removes 30-50% of SNP FPs, with smaller gains on INDELS. FN counts are reduced by 10% or more.
--vc-ml-dir=/opt/edico/resources/ml_model/hg19 --vc-ml-enable-recalibration=true
Star Allele Caller 星号等位基因:用于药物基因组
--enable-star-allele true
Targeted callers: 输出.targeted.vcf.gz and .targeted.json
--enable-targeted=true
High Sensitivity Mode
--vc-enable-high-sensitivity-mode=true
CNV 分析参数
```

```
--enable-cnv true
--cnv-filter-copy-ratio 0.2  # The default value is 0.2, leading to calls less than CR=0.8 or
--greater than CR=1.2.
--cnv-filter-length 10000  # Specifies the minimum event length in bases at which a reported
--greater than CR=1.2.
--cnv-filter-length 10000  # Specifies the minimum event length in bases at which a reported
--cnv-filter-qual 10  # PASS in the output VCF file
```

## CNV 解析度

WGS_Coverag_per_Sample	Recommended_Resolution(bp)
5X	10000
10X	5000
>=30X	1000

-cnv-interval-width 用来控制解析度,WES 默认是 500,WGS 默认是 1000 该参数在分析是需要设置,如果设置变小会增加分析时间