1000 Genomes Processing README

This README contains information relating to data associated with the 1000 Genomes resequencing done at New York Genome Center.

Alignment, post-processing and variant calling

Alignment and post-processing are performed exactly as outlined by the Center for Common Disease Genomics project: https://github.com/CCDG/Pipeline-Standardization/blob/master/PipelineStandard.md .

Programs and reference data

The data was aligned to the reference genome using the following programs and reference datasets:

- 1. BWA-MEM bwakit-0.7.15
- 2. Samtools-1.3.1
- 3. Picard-2.4.1
- 4. GATK-3.5-0
- 5. Resource files
 - All the resource files used in the analysis can be obtained here:
 https://console.cloud.google.com/storage/browser/genomics-public-data/resources/broad/hg38/v0/.

Reference genome: GRCh38 with alternative sequences, plus decoys and HLA

The reference genome that the data was aligned to can be obtained here: ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/reference/GRCh38_reference_genome /GRCh38_full_analysis_set_plus_decoy_hla.fa

Command lines

1. Alignment at lane level

```
bwa mem -Y \
-K 100000000 \
-t 16 \
-R $rg_string \
$reference_fasta_file \
$fastq_file(1) \
$fastq_file(2) | samtools view -Shb -o $bam_file -
```

2. Fix mate information in the BAM

```
java $jvm_args -jar picard.jar \
FixMateInformation \
MAX_RECORDS_IN_RAM=2000000 \
VALIDATION STRINGENCY=SILENT \
```

```
ADD_MATE_CIGAR=True \
    ASSUME_SORTED=true \
    I=$bam_file \
    O=$bam file fixedmate
Merging lane-level bam files to Sample level bam files
    java $jvm_args -jar picard.jar \
    MergeSamFiles \
    USE_THREADING=true \
    MAX RECORDS IN RAM=2000000 \
    VALIDATION STRINGENCY=SILENT \
    SORT ORDER=queryname \
    INPUT=$bam1 \
    INPUT=$bam2 \
    OUTPUT=$bam merged
Mark duplicates and coordinate sort BAM
    java $jvm_args -jar picard.jar \
    MarkDuplicates \
    MAX_RECORDS_IN_RAM=2000000 \
    VALIDATION STRINGENCY=SILENT \
    M=$dedup metrics \
    I=$bam_sorted \
    O=$bam_dedup
    java $jvm_args -jar picard.jar \
    SortSam \
    MAX RECORDS IN RAM=2000000 \
    VALIDATION_STRINGENCY=SILENT \
    SORT ORDER=coordinate \
    CREATE_INDEX=true \
    I=$bam_merged \
    O=$bam sorted
Recalibrate base quality scores using known SNPs
    java $jvm_args -jar GenomeAnalysisTK.jar \
    -T BaseRecalibrator \
    -downsample to fraction 0.1 \
    -nct 4 \
    --preserve_qscores_less_than 6 \
    -L $autosomes \
    -R $reference_fasta \
    -o $recal_data.table \
    -I $bam sorted \
```

-knownSites \$known snps from dbSNP138 \

-knownSites \$known indels from mills 1000genomes

-knownSites \$known_indels \

3.

4.

5.

```
java $jvm_args -jar GenomeAnalysisTK.jar \
        -T PrintReads \
        -nct 4 \
        --disable indel quals \
        --preserve_qscores_less_than 6 \
        -SQQ 10 \
        -SQQ 20 \
        -SQQ 30 \
        -rf BadCigar \
        -R $reference fasta \
        -o $recalibrated bam \
        -I $bam sorted \
        -BQSR $recal_data.table
6.
   Creating CRAM files
        samtools view \
        -C \
        -T $reference_fasta \
        -o $cram \
        $recalibrated_bam
        samtools index $cram
    Raw variant calls using HaplotypeCaller on single sample
7.
        java $jvm_args -jar GenomeAnalysisTK.jar \
        -T HaplotypeCaller \
        --genotyping_mode DISCOVERY \
        -A AlleleBalanceBySample \
        -A DepthPerAlleleBySample \
        -A DepthPerSampleHC \
        -A InbreedingCoeff \
        -A MappingQualityZeroBySample \
        -A StrandBiasBySample \
        -A Coverage \
        -A FisherStrand \
        -A HaplotypeScore \
        -A MappingQualityRankSumTest \
        -A MappingQualityZero \
        -A QualByDepth \
        -A RMSMappingQuality \
        -A ReadPosRankSumTest \
        -A VariantType \
        -1 INFO \
        --emitRefConfidence GVCF \
        -rf BadCigar \
        --variant_index_parameter 128000 \
        --variant_index_type LINEAR \
        -R $reference_fasta \
        -nct 1 \
```

```
-I $recalibrated bam \
        -o $gvcf
  Jointly recalibrate Genotype Quality score of all samples
        java $jvm args -jar GenomeAnalysisTK.jar \
        -T GenotypeGVCFs \
        -R $reference fasta \
        -nt 5 \
        --disable_auto_index_creation_and_locking_when_reading_rods \
        --variant $gvcf \
        -o $recalibrated vcf
9. Variant Quality Score Recalibration (VQSR) to assign FILTER status
        java $jvm_args -jar GenomeAnalysisTK.jar /
    -T VariantRecalibrator /
    -R $reference fasta /
    -nt 5 /
    -input $recalibrated vcf /
    -mode SNP /
    -recalFile $vqsr_snp.recal /
    -tranchesFile $vqsr snp.tranches /
    -rscriptFile $vqsr_snp_plots.R /
    -resource:hapmap,known=false,training=true,truth=true,prior=15.0 $hapmap
/
    -resource:omni,known=false,training=true,truth=true,prior=12.0 $kg omni /
    -resource:1000G, known=false, training=true, truth=false, prior=10.0 $kg snps
    -resource:dbsnp,known=true,training=false,truth=false,prior=2.0 $dbsnp /
    -an QD /
    -an MQ /
    -an FS /
    -an MQRankSum /
    -an ReadPosRankSum /
    -an SOR /
    -an DP /
    -tranche 100.0 /
    -tranche 99.8 /
    -tranche 99.6 /
    -tranche 99.4 /
    -tranche 99.2 /
    -tranche 99.0 /
    -tranche 95.0 /
    -tranche 90.0
        java $jvm_args -jar GenomeAnalysisTK.jar /
    -T VariantRecalibrator /
    -R $reference_fasta /
    -nt 5 /
    -input $recalibrated vcf /
    -mode INDEL /
```

```
-recalFile $recalibrate indel.recal /
    -tranchesFile $recalibrate_indel.tranches /
    -rscriptFile $recalibrate_indel_plots.R /
    -resource:mills,known=true,training=true,truth=true,prior=12.0 $kg mills
/
    -resource:dbsnp,known=true,training=false,truth=false,prior=2.0 $dbsnp /
    -an QD /
    -an FS /
    -an ReadPosRankSum /
    -an MQRankSum /
    -an SOR /
    -an DP /
    -tranche 100.0 /
    -tranche 99.0 /
    -tranche 95.0 /
    -tranche 92.0 /
    -tranche 90.0 /
    --maxGaussians 4
        java $jvm_args -jar GenomeAnalysisTK.jar /
    -T ApplyRecalibration /
    -R $reference_fasta /
    -nt 5 /
    -input $recalibrated_vcf /
    -mode SNP /
    --ts filter level 99.80 /
    -recalFile $recalibrate SNP.recal /
    -tranchesFile $recalibrate SNP.tranches /
    -o $vqsr_snp_vcf
        java $jvm_args -jar GenomeAnalysisTK.jar /
    -T ApplyRecalibration /
    -R $reference fasta /
    -nt 5 /
    -input $vqsr_snp_vcf /
    -mode INDEL /
    --ts filter level 99.0 /
    -recalFile $recalibrate_INDEL.recal /
    -tranchesFile $recalibrate INDEL.tranches /
    -o $vqsr_snp_indel_vcf
```