

# Getting Started with the Genome Analysis Toolkit (GATK)

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## 1 Build Prerequisites

GATK requires JDK 1.6 and Ant 1.7.1 to compile.

## 2 Getting and Building the Source

GATK is located in the Sting svn repository, and compiles using a build.xml in the root directory.

Download and build the source as follows:

```
svn co https://svnrepos/Sting/trunk Sting
cd Sting
ant
```

## 3 Getting Started

The core concept behind GATK is the walker, a class that implements the three core operations: filtering, mapping, and reducing.

**filter** reduces the size of the dataset by applying a predicate.

**map** Applies a function to each individual element in a dataset, effectively 'mapping' it to a new element.

**reduce** Inductively combines the elements of a list. The base case is supplied by the reduceInit() function, and the inductive step is performed by the reduce() function.

Users of the GATK will provide a walker to run their analyses. The engine will produce a result by first filtering the dataset, running a map operation, and finally reducing the map operation to a single result.

## 4 Creating a Walker

To be loaded by GATK, the walker must satisfy the following properties:

1. It must be a loose class, not packaged into a jar file.
2. It must be in the unnamed package (in other words, the source should not start with a package declaration).
3. It must subclass one of the basic walkers in the `org.broadinstitute.sting.gatk.walkers` package: `BasicReadWalker` or `BasicLociWalker`.
4. It must live in the directory `$STING_HOME/dist/walkers`.

## 5 Example

This walker will print output for each read it sees, eventually computing the total number of reads by mapping every read to 1 and summing all the 1s to realize the total number of reads.

Copy the following text into the file \$STING\_HOME/dist/walkers/HelloWalker.java:

```
import net.sf.samtools.SAMRecord;

import org.broadinstitute.sting.gatk.LocusContext;
import org.broadinstitute.sting.gatk.walkers.BasicReadWalker;

/**
 * Define a class extending from BasicReadWalker with types
 * <MapType,ReduceType>.
 */
public class HelloWalker extends BasicReadWalker<Integer,Long> {
    private Long currentRead = 0L;

    // Maps each read to the value 1.
    public Integer map(LocusContext context, SAMRecord read) {
        System.out.printf("Hello read %d%n", ++currentRead );
        return 1;
    }

    // Provides an initial value for the reduce function.
    public Long reduceInit() { return 0L; }

    // Defines how to compute the reduction given a value in the list.
    public Long reduce(Integer value, Long sum) {
        return sum + value;
    }
}
```

To compile the walker:

```
setenv CLASSPATH $STING_HOME/dist/GenomeAnalysisTK.jar:$STING_HOME/dist/sam-1.0.jar
javac HelloWalker.java
```

To run the walker:

```
mkdir $STING_HOME/dist/walkers
java -Xmx4096m -jar dist/GenomeAnalysisTK.jar \
    -I /broad/1KG/legacy_data/trio/na12878.bam -T Hello \
    -L chr1:10000000-10000100 -l INFO
```

This command will run the walker across a subsection of chromosome 1, operating on reads which align to that subsection.