

# The Biological Repository (BioR) Quick-start

This is a quick start for those in a hurry. For more detailed instructions, please see the user manual on the following page: <http://bioinformaticstools.mayo.edu/research/bior/>

BioR tools and their uses (shortlist)

Tool Command	Input Format	Output Format	Function
<code>bior_overlap</code>	TJSON	TJSON	Extract from catalog all the attributes based on genomic coordinates
<code>bior_same_variant</code>	TJSON	TJSON	Matches variants based on position, reference, and alternate alleles
<code>bior_lookup</code>	TJSON	TJSON	Matches based on a string/identifier
<code>bior_drill</code>	TJSON	Tab-Delim	Extracts key-value relationships from JSON object
<code>bior_pretty_print</code>	TJSON	JSON	Prints the JSON to the screen in a more readable way
<code>bior_vcf_to_tjson</code>	VCF	TJSON	Converts VCF format to JSON objects
<code>bior_bed_to_json</code>	BED	TJSON	Converts BED format to JSON objects
<code>bior_snpeff</code>	VCF	TJSON	Wraps the SNPEFF tool (if installed)
<code>bior_vep</code>	VCF	TJSON	Wraps the VEP tool (if installed)
<code>bior_annotate</code>	VCF	TJSON	Wrapper to get commonly used variant annotations
<code>bior_index_catalog</code>	TJSON	INDEX	Creates and index on a string/identifier

## Prerequisites:

1) Make sure that you have BioR and its dependencies installed correctly and in your path. This includes Java JDK 1.7, Tabix and BGZIP. Most BioR functions will still work if you don't install SNPEFF/VEP and all of their dependencies, but `bior_annotate` will have limited functionality and `bior_vep` and `bior_snpeff` will not work. The environment variable, `$BIOR_LITE_HOME` represents the location where BioR is installed.

2) To check it is installed and on your path, execute `bior_pretty_print -h` at the command line: it should output a description and the text on how to use `bior_pretty_print`.

```
$ bior_pretty_print -h

NAME
    bior_pretty_print -- prints out a single data row in a readable format

SYNOPSIS
    bior_pretty_print [--row-number <number>] [--log] [--help]

DESCRIPTION
    ...
```

3) Ready to run examples are available in `$BIOR_LITE_HOME/examples`.

## Example 1:

Change directory to `$BIOR_LITE_HOME/examples/quickstart1`. The first example in this guide is to convert gene names into gene positions, which follows this process:

1. Get the gene list
2. Find a catalog that has a gene name in it
  - in this example: `$BIOR_LITE_HOME/examples/quickstart1/data/NCBIGene/GRCh37_p10/genes.tsv.bgz`
3. Look up your gene name in that catalog with `bior_lookup`
4. Extract the key/value pairs that you are interested with `bior_drill`

**STEP1.** The file `geneList.txt` contains a list of genes. Cat the file to see what you are starting with.

```
$ cat geneList.txt
BRCA1
BRCA2
CD28
ZBTB7A
$
```

**STEP2.** Look at the contents of the gene catalog to see what fields are there (NOTE: use *gzcat* on a mac or *zcat* on Linux). For brevity, we are only showing selected portions in this manual, but the complete outputs are given in STEP1.txt, STEP2.txt, etc.

```
$ gzcat data/NCBIGene/GRCh37_p10/genes.tsv.bgz | bior_pretty_print
... {
  "_landmark": "1",
  "_strand": "+",
  "_minBP": 10954,
  "_maxBP": 11507,
  "gene": "LOC100506145"
...
}
$
```

**STEP3.** Find the genes in geneList.txt in the gene field of the genes.tsv.bgz catalog using bior\_lookup:

```
$ cat geneList.txt | bior_lookup -p gene -d data/NCBIGene/GRCh37_p10/genes.tsv.bgz | bior_pretty_print
# COLUMN NAME      COLUMN VALUE
- -----
1 UNKNOWN_1        BRCA1
2 bior.gene37p10 {
...
  "_landmark": "17",
  "_strand": "-",
  "_minBP": 41196312,
  "_maxBP": 41277500,
  "gene": "BRCA1",
...
  "MIM": "113705"
}
$
```

**STEP4.** Create a simple tab-delimited file of the gene and position information using bior\_drill:

```
$ cat geneList.txt | bior_lookup -p gene -d data/NCBIGene/GRCh37_p10/genes.tsv.bgz | bior_drill -p _landmark
-p _strand -p _minBP -p _maxBP > genePos.txt
$ cat genePos.txt
...
#UNKNOWN_1      bior.gene37p10._landmark      bior.gene37p10._strand bior.gene37p10._minBP
      bior.gene37p10._maxBP
BRCA1  17      -      41196312      41277500
BRCA2  13      +      32889617      32973809
CD28   2      +      204571198     204603635
ZBTB7A 19      -      4045216 4066816
$
```

## Example 2:

The second example shows briefly how to annotate a VCF file. In this example, we are interested in finding out if any of my variants are clinically relevant from a subset of the ClinVar database ([http://www.ncbi.nlm.nih.gov/variation/docs/human\\_variation\\_vcf/](http://www.ncbi.nlm.nih.gov/variation/docs/human_variation_vcf/)). From the glossary of the catalog source (<http://www.ncbi.nlm.nih.gov/variation/docs/glossary>), CLNSIG is the annotation we are looking for, since it classifies variants as pathogenic, non-pathogenic, drug-response, etc. For this example, you will need to change directory to \$BIOR\_LITE\_HOME/examples/quickstart2. The file example.vcf contains a VCF file of genetic variants.

To annotate the VCF:

1. VCFs must be converted into a tjson file before searching using `bior_vcf_to_tjson`
2. We need to use `bior_same_variant` to match alternate alleles between our VCF and the ClinVar catalog
3. Drill out the key/value pairs of interest
4. Convert back to VCF for future use or pipe directly into another command

```
$ cat example.vcf | bior_vcf_to_tjson | bior_same_variant -d data/dbSNP/137/clinvarPart.tsv.bgz | bior_drill
-p INFO.CLNSIG | bior_tjson_to_vcf
##fileformat=VCFv4.0
##INFO=<ID=bior.clinvar137.INFO.CLNSIG,Number=.,Type=String,Description="VCF INFO. A string that describes
the variant's clinical significance, where 0 - unknown, 1 - untested, 2 - non-pathogenic, 3 - probable-non-
pathogenic, 4 - probable-pathogenic, 5 - pathogenic, 6 - drug-response, 7 - histocompatibility, 255 -
other.">
#CHROM POS ID REF ALT QUAL FILTER INFO
13 32907403 . T C . . bior.clinvar137.INFO.CLNSIG=1
15 3240324 . N C . . .
$
```

\*Notice the “INFO.CLNSIG” instead of just “CLNSIG”. This is because the CLNSIG key is in the hierarchy of INFO {} (see STEP5.txt).