Snipper Documentation

Version 1.2

Author: Ryan Welch ([welchr@umich.edu](mailto:welchr@umich.edu?subject=[Snipper]%20YOUR_SUBJECT_HERE))

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**Purpose**

Snipper is a research tool for investigating genes near associated loci from GWAS studies. The user can supply a SNP or list of SNPs, a list of genes, and/or a list of chromosomal regions, after which Snipper will:

* Create a gene list, where genes are added by:
  + For each SNP, find genes nearby up to a certain distance specified by the user, or those genes whose expression is known to be associated with the SNP
  + For each region (ex: chr9:1911-939393), find genes within or overlapping the region
  + Include each gene specified by the user
* Retrieve annotations for each gene from NCBI Entrez Gene, OMIM, and the Michigan Molecular Interactions database (MiMI)
* If the user supplies search terms: search PubMed for each combination of search term and gene
* Search annotation information on each gene for user's search terms
* Create an HTML (or console) report containing all of the available information for each gene (including where search terms matched and how often)

Snipper is designed to handle a modest number of loci (25-50), but has been tested to handle up to 100. Submitting a large number of SNPs beyond this is not recommended, as the program may require extreme amounts of time and memory. We have typically used a handful of search terms (less than 10) - using substantially more than this may also require very large amounts of time for Snipper to finish.

**Installation**

**Installing on Windows (Vista/7)**

To install under Windows, simply run the installer exe downloaded from our website. This will put Snipper under your appropriate \Program Files\ directory, and create a desktop icon to launch the GUI.

If you wish to use the command line instead of the GUI, you will need to:

1. Install Python 2.6 or greater (but **not** the 3.x branch). You can download this from <http://www.python.org/>.
2. Install Snipper from the setup executable downloaded from our website
3. Navigate to where Snipper was installed (usually something like C:\Program Files (x86)\Snipper\, the installer will tell you this)
4. Run the bin/setup\_snipper.py script.
5. Use the command line script bin/snipper.py to run your queries.

**Installing from source (Linux/Unix)**

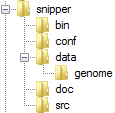
To install Snipper, you will need the following:

* Python version 2.6 or greater (but **not** the 3.x branch!)
* A working internet connection

To use the GUI for Snipper, you will also need the proper Tk package installed on your system. Under Ubuntu Linux, you can install it by using:

sudo apt-get install python-tk

1. Extract Snipper to the location of your choice. Note that you must extract the files into the directory structure given in the archive. It should follow this tree structure:



You can do this by running the following command:

tar zxf snipper\_tarball.tar.gz

1. Next, navigate to snipper/bin, and run the **setup\_snipper.py** script. This script will install dependencies for Snipper, and ensure that they were installed correctly.

python setup\_snipper.py

You should first see dependencies being installed. It takes roughly a minute or so to install. The packages are installed to a virtual python environment under snipper/pyenv, and are not installed globally on your system (therefore, you do not need admin privileges.)

1. Snipper is ready to run, and can be launched by navigating to snipper/bin and running **snipper.py**. For simplicity, you could create a shortcut to this by doing the following:

ln –s snipper/bin/snipper.py /usr/local/bin/snipper

**Using Snipper (command line)**

**Synopsis**

Typical usage of Snipper will be of the form:

|  |
| --- |
| **snipper** --snpfile <file containing SNPs> |

If one had a file containing SNPs, wanted to search 250kb away from each SNP for genes:

|  |
| --- |
| **snipper** --snpfile <file containing SNPs> -d 250kb |

A user will generally want to include search terms with their query, for example:

|  |
| --- |
| **snipper** --snpfile <file containing SNPs> -d 250kb --terms “glucose,insulin” |

Snipper can include genes explicitly requested by the user:

|  |
| --- |
| **snipper** –-gene “TCF7L2,P53,BRCA1” |

Or, the program can be run with chromosomal regions:

|  |
| --- |
| **snipper** –-regions “chr#:start-end” |

All of these can be mixed together, for example:

|  |
| --- |
| **snipper** -s "rs7903146,rs1002227" –-regions “chr3:12393001-12475854” --gene "RB1,PDE8B" |

You can verify that Snipper has installed correctly by running our test example. Simply change directory into the example/ directory, and execute "run\_example.py". This script runs a simple test using a few SNPs, genes, and chromosomal regions. The script will explain what it is doing, as well as showing the command line used to run Snipper. A directory called "example\_results" will be created, containing the HTML output. There is also a "precompiled\_results" directory, which gives the output if the program were to execute successfully. You can compare the two outputs to ensure that the program is working as expected (though we note that as databases and minor revisions to the HTML report format take place, the two may be slightly out of sync.)

**Options**

Snipper supports a wide variety of command line arguments for tailoring what and how much information is retrieved. Please see the table below for a full listing.

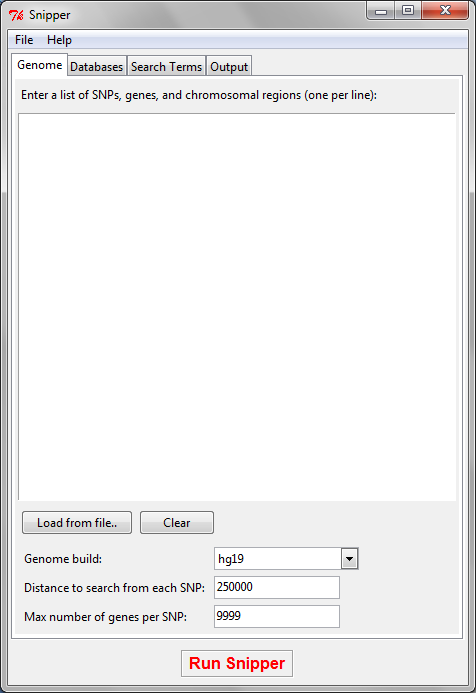
|  |  |
| --- | --- |
| **Argument** | **Description** |
| -o, --out <string> | Specify output directory for report. The output directory will contain directories with the following:   * HTML report (what you’ll likely want to read) * RST files (used to generate the HTML report) * Text output (what would have been printed in plaintext using –console) * A readme file that explains the directories created, along with the command line used to run Snipper |
| --console | Write output to the console instead of creating a report directory. |
| **Options for specifying SNPs, genes, and regions** | |
| -s, --snp <string> | Lookup information for a list of SNPs - these must be separated by commas, surrounded by quotes (whitespace ignored.)  Example:  -s "rs1002227, rs35712349" |
| --snpfile <string> | Provide a list of SNPs to lookup from a file. The file may have \*ANY\* format, provided the file contains plain text. The program will pattern match rs### identifiers from your file. If you have SNPs in the 1000G format (e.g. chr4:9393) they must be specified on the command line using the -s option for now. |
| --build <string> | Select build to use for finding the positions of SNPs and genes. Snipper comes with support for hg19 by default, though other databases can be built. See "[Building your own position database](#Database)" for more information. |
| -g, --gene <string> | Lookup information for a list of gene symbols - these must be separated by commas, surrounded by quotes (whitespace ignored) |
| --genefile <string> | A file of genes to include in the Snipper report. Genes should be the primary HGNC gene symbol. One per line. |
| -r, --regions <string> | Provide a list of chromosomal regions. Genes within these regions will be included in the report.  Example:  --regions "chr4:19141-939393,chrX:9191-939393"  The position numbers themselves should not contain commas. |
| -d <string> | Distance away from SNP to search, default is 1000000. If a distance is specified, the program will return \*ALL\* genes within the distance you specify, not just the default of 1. To specify a new distance, but still only return 1 gene (or arbitrary number of genes), use -n <number>. Distances can be specified using a kb or mb suffix, or as a raw distance. Examples: 500kb, 0.5MB, 1.4MB,834141. |
| -n <int> | Number of genes to return per SNP, default is 1. Note that this works in conjunction with the –d parameter listed above. For example, if you specify a distance of 10MB, but set –n 3, the program will search within 10 megabases of your SNP and return the 3 nearest genes. |
| **Options related to PubMed: search terms, how to search, and how many articles to return** | |
| --terms <string> | Comma-delimited string of terms, enclosed in quotes, to use in searching the literature. This will execute a search, per gene, for any of the search terms.  For example:  Genes: RB1, TCF7L2  Search terms: "glucose,retinoblastoma"  What happens:  -- Search literature for RB1 AND (glucose OR retinoblastoma)  -- Search literature for TCF7L2 AND (glucose OR retinoblastoma)  The information for genes RB1 and TCF7L2 will contain a list of PubMed articles that matched at least 1 of your search terms, all of which will be lumped together.  Searching for terms with spaces is possible, but the entire argument must be enclosed in quotation marks. For example:  --terms "type 2 diabetes, insulinemia, metabolic syndrome" |
| --each-term | When specified, the program will search each gene x search term pair, instead of lumping together search terms. For example:  Genes: RB1, TCF7L2  Search terms: "glucose,retinoblastoma"  What happens:  -- Search literature for RB1 AND glucose  -- Search literature for RB1 AND retinoblastoma  -- Search literature for TCF7L2 AND glucose  -- Search literature for TCF7L2 AND retinoblastoma  The information for genes RB1 and TCF7L2 will have sections of PubMed articles that matched each search term individually. While this makes it more apparent why each PubMed article was returned, it also requires sending more queries to NCBI, and therefore increases the runtime of the program significantly. This is disabled by default. |
| --papernum <int> | Number of recent papers to display, default is 5 |
| **Options for disabling various databases (all are enabled by default)** | |
| --no-generif | Disable GeneRIFs. |
| --no-omim | Disable OMIM. |
| --no-pubmed | Disable PubMed. |
| **Options related to ScanDB (eQTL database)** | |
| --no-scandb | Disables use of ScanDB for finding eQTLs connecting user defined SNPs to genes. |
| --scandb-pval | Change the p-value threshold for calling an eQTL association as “significant.” The default is 1.0E-06. |

**Using Snipper (GUI)**

To launch the GUI:

* Double-click the “Snipper” icon on your desktop (or in the Start Menu, if you chose not to create a desktop icon) (Windows)
* Run bin/snipper.py without any command line arguments (Linux; see Installation to make sure you have the dependencies installed to run the GUI)

When the GUI launches, you should see the following screen:



The main text area is where SNPs, genes, and chromosomal regions can be listed. You can mix these interchangeably, so long as you keep them one per line. For example, you could try:

rs7903146

PDE8B

chrX:70608087-70685854

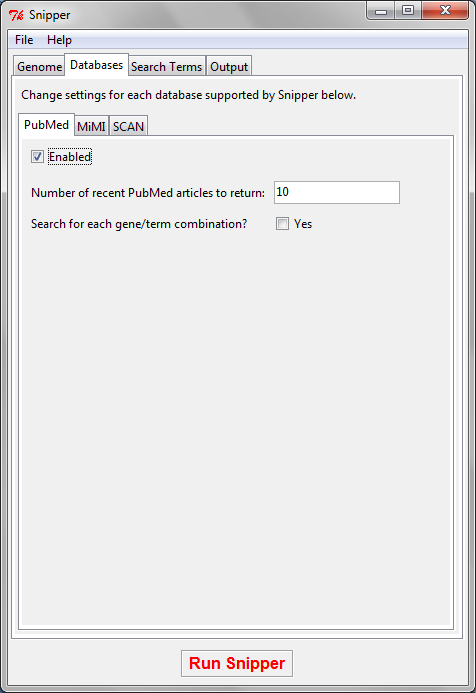
KCNJ11

rs1002227

Alternatively, you could click “**Load from file..**” to read a list of SNPs, genes, and chromosomal regions from a file.

Towards the bottom, there are a few additional options. **Genome build** denotes which human genome build to use when looking the positions of SNPs and genes. **Distance to search from each SNP** specifies the maximal distance that will be searched from each SNP when looking for genes. **Max number of genes per SNP** controls the number of genes that will be returned near a SNP (prioritized by distance, so if there were 5 genes but you desired only the nearest, you would set this to 1.) Leave the value at 9999 to return all genes near a SNP.

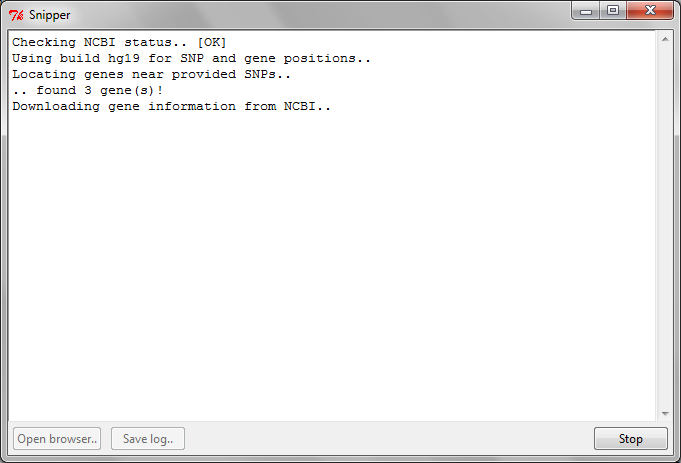
On the “**Databases**” tab there are various options for enabling or disabling databases for querying, as well as changing options that may be specific to a particular one:



The “**Search Terms**” tab is where you can list words or phrases that will be used to search within information returned from OMIM, PubMed, and Entrez Gene. Enter one word or phrase per line.

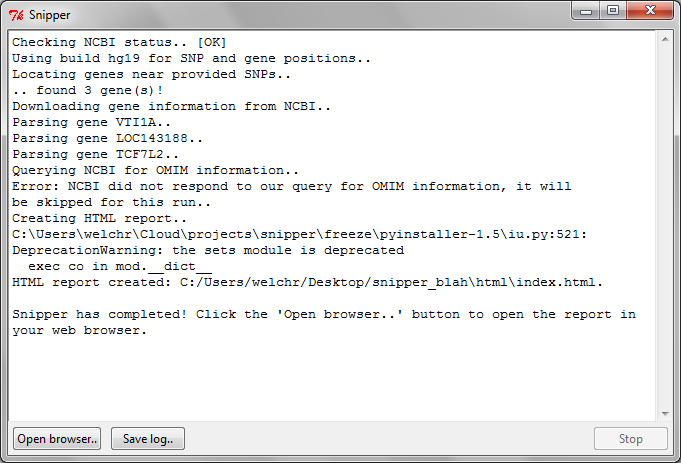
The “**Output**” tab allows the user to change where the snipper report directory will be located. If the directory already exists,

Once you’re satisified with your settings, click the “**Run Snipper**” button at the bottom, or select File 🡪 Run Snipper. You’ll see a console window appear:



This window shows what Snipper is currently doing, and is very similar to what would be seen in command line mode. At the bottom right, the **Stop** button can be used to terminate the current run.

Once the run has completed, you should see the following:



The “**Open browser..**” button will directly open the HTML report in your browser. This is the best way to quickly view the report. You could alternatively navigate to the directory (shown in the log file above) and open the “index.html” file.

If you wish to save the log, you can do so by clicking “**Save log..**” and selecting where the console text will be written.

To run Snipper again, simply close the console, change your settings, and click “**Run Snipper**” again.

**Output**

**Snipper Console Output**

Below, we describe the anatomy of the snipper console output. This mode is less preferable than the HTML output, but can be used for quick inspection via the command line. To activate console output, use the --no-html parameter. In this particular example, we've searched near known type 2 diabetes SNPs, and returned the nearest gene within 250kb for each SNP.

This table gives a listing for each gene identified near a SNP given as input by the user.

#SNPs - number of SNPs given by the user that were near this gene. This can be > 1 when you have 2 SNPs very close by each other.

# Terms - the number of user-defined search terms that were found in the information for this gene.

Total Pubmed - the total number of PubMed articles linked to this gene. This gives the user an idea of how well-researched this particular gene is.

Gene # SNPs # Terms Total Pubmed

---- ------ ------- ------------

CDKAL1 1 4 64

KCNJ11 1 4 198

IGF2BP2 1 4 58

NOTCH2 1 2 76

JAZF1 1 2 42

TCF7L2 1 4 253

KCNQ1 1 4 207

HNF1B 1 4 88

SLC30A8 1 4 83

WFS1 1 4 89

THADA 1 2 20

FTO 1 4 133

HHEX 1 4 89

MTNR1B 1 4 34

ADAMTS9 1 2 26

CDC123 1 2 24

PPARG 1 4 938

TSPAN8 1 2 26

CDKN2B 1 4 165

KIAA1486 1 0 4

SNP Gene/Aliases

--- ------------

For each SNP provided by the user, the table to the left gives the following information:

* The SNP itself
* A row for each gene found near the SNP (this particular example only has 1 gene per SNP, there could be more depending on your settings)
* The primary gene symbol is listed first, followed by the gene's aliases, i.e.:

TCF7L2/TCF4/TCF-4

TCF7L2 is the primary symbol, TCF4 and TCF-4 are aliases.

rs11899863 THADA/GITA/FLJ77530/FLJ44876/FLJ44016/KIAA1767/FLJ21877

rs7903146 TCF7L2/TCF4/TCF-4

rs1387153 MTNR1B/MT2/MEL-1B-R

rs849134 JAZF1/DKFZp761K2222/ZNF802/TIP27

rs4430796 HNF1B/TCF2/FJHN/HNF1beta/HPC11/VHNF1/MODY5/HNF2/LFB3/LF-B3

rs6795735 ADAMTS9/FLJ42955/KIAA1312

rs3802177 SLC30A8/ZNT8/ZnT-8

rs10923931 NOTCH2/AGS2/hN2

rs1801214 WFS1/WFS/FLJ51211/WOLFRAMIN/WFRS

rs163184 KCNQ1/FLJ26167/JLNS1/LQT/KVLQT1/Kv1.9/KCNA9/SQT2/RWS/LQT1/WRS/KCNA8/ATFB3/Kv7.1/ATFB1

rs10965250 CDKN2B/MTS2/TP15/P15/p15INK4b/CDK4I/INK4B

rs11642841 FTO/KIAA1752/MGC5149

rs1470579 IGF2BP2/IMP2/p62/IMP-2/VICKZ2

rs10440833 CDKAL1/FLJ46705/MGC75469/FLJ20342

rs12779790 CDC123/D123/C10orf7/FLJ13863

rs5015480 HHEX/HMPH/HEX/PRH/PRHX/HOX11L-PEN

rs7578326 KIAA1486

rs4760790 TSPAN8/CO-029/TM4SF3

rs5215 KCNJ11/IKATP/TNDM3/PHHI/HHF2/KIR6.2/MGC133230/BIR

rs13081389 PPARG/PPARgamma/GLM1/PPARG2/PPARG1/CIMT1/NR1C3

***From this point forward, Snipper lists genes found near SNPs. It will first list genes known to be associated with SNPs (eQTLs), and then subsequently list the remaining genes in order by distance to SNP. If the user is interested in a particular gene, most text editors can search quickly by using CTRL+F.***

***For this particular example, we list only 1 gene - CDKAL1.***

================================================================================

The gene's full name, primary symbol, and synonyms. Also the UID (Entrez Gene's identifier for the gene), chromosomal location, and type of gene.

[+] GENE: potassium inwardly-rectifying channel, subfamily J, member 11 [KCNJ11]

[+] Entrez Gene UID: 3767

[+] Location: 11p15.1

[+] Type: protein-coding

[+] Synonyms: IKATP TNDM3 PHHI HHF2 KIR6.2 MGC133230 BIR

[+] Search terms matched:

This section shows which search terms provided by the user were found in this gene's information, and also shows where they matched.

-- Location: Gene summary Terms: insulin, diabetes

-- Location: GO Term Terms: insulin, glucose

-- Location: GeneRIF Terms: glucose, insulin, diabetes

-- Location: Pubmed Terms: any

-- Location: Phenotype Terms: diabetes

-- Location: KEGG Pathway Terms: diabetes

-- Location: OMIM Text Terms: glucose, insulin, diabetes

[+] Associated SNPs:

SNP provided by user to search near, and the distance to this SNP.

SNP: rs5215 Distance (bp): 0 Direction: Within

[+] Summary: Potassium channels are present in most mammalian cells, where they

participate in a wide range of physiologic responses. The protein encoded by

Summary of the gene (provided by NCBI Entrez Gene.)

this gene is an integral membrane protein and inward-rectifier type potassium

channel. The encoded protein, which has a greater tendency to allow potassium to

flow into a cell rather than out of a cell, is controlled by G-proteins and is

found associated with the sulfonylurea receptor SUR. Mutations in this gene are

a cause of familial persistent hyperinsulinemic hypoglycemia of infancy (PHHI),

an autosomal recessive disorder characterized by unregulated insulin secretion.

Defects in this gene may also contribute to autosomal dominant non-insulin-

dependent diabetes mellitus type II (NIDDM), transient neonatal diabetes

mellitus type 3 (TNDM3), and permanent neonatal diabetes mellitus (PNDM).

[provided by RefSeq]

Phenotypes associated with the gene (provided by NCBI Entrez Gene.)

[+] Phenotypes: "Adiposity-related heterogeneity in patterns of type 2 diabetes

susceptibility observed in genome wide association data" "Diabetes mellitus,

permanent neonatal, with neurologic features, 606176, {Diabetes mellitus, type

2, susceptibility to" "Diabetes mellitus, type 2, susceptibility to" "Diabetes,

permanent neonatal" "Hyperinsulinemic hypoglycemia, familial, 2" "Meta-analysis

of genome-wide association data and large-scale replication identifies

additional susceptibility loci for type 2 diabetes" "Diabetes mellitus,

transient neonatal, 3" "Replication of genome-wide association signals in UK

samples reveals risk loci for type 2 diabetes" "A genome-wide association study

of type 2 diabetes in Finns detects multiple susceptibility variants" "Diabetes

mellitus, permanent neonatal, with neurologic features" "Genome-wide association

analysis identifies loci for type 2 diabetes and triglyceride levels"

[+] KEGG Pathways:

KEGG pathways.

Type II diabetes mellitus [ http://www.genome.jp/dbget-bin/show\_pathway?hsa04930+3767 ]

Gene ontology terms.

[+] GO Terms: "regulation of membrane potential" "ATP-activated inward rectifier

potassium channel activity" "potassium ion import" "endoplasmic reticulum"

"voltage-gated ion channel activity" "ATP-sensitive potassium channel complex"

"plasma membrane" "microsome" "negative regulation of insulin secretion" "ion

transport" "glucose metabolic process" "protein C-terminus binding" "T-tubule"

"response to drug" "ATP binding" "mitochondrion" "potassium ion binding"

"neurological system process" "response to ATP"

OMIM ID, link to the OMIM article itself, and the OMIM summary for the gene.

[+] OMIM: [600937] Link: http://www.ncbi.nlm.nih.gov/entrez/dispomim.cgi?id=600937

[+] OMIM Text: The KCNJ11 gene encodes a subunit of an inwardly rectifying ATP-

sensitive potassium channel. I(KATP) channels were discovered in cardiac muscle

and later found in pancreatic beta cells, pituitary tissue, skeletal muscle,

brain, and vascular and nonvascular smooth muscle. I(KATP) currents function in

secretion and muscle contraction by coupling metabolic activity to membrane

potential ({13:Inagaki et al., 1995}).

GeneRIFs (Gene References Into Function), provided by NCBI Entrez Gene.

This can be enabled by either --generif or the --all option.

[+] Gene references into function for KCNJ11:

-- Observational study of gene-disease association. (HuGE Navigator) PMID:

19685080

-- Mutations in the pore-forming K(ATP) channel subunit cause neonatal diabetes

&amp; discusses recent advances in understanding of clinical features of

neonatal diabetes, its underlying molecular mechanisms &amp; their impact on

treatment[review] PMID: 18566517

-- mutations in the slide helix of Kir6.2 (V59G) influence the channel kinetics,

providing evidence that this domain is involved in Kir channel gating PMID:

15583126

-- Case of an 18-month-old infant with permanent neonatal diabetes due to an

activating KCNJ11 mutation who successfully transitioned from subcutaneous

insulin therapy to oral sulfonylurea therapy in the outpatient setting. PMID:

18221420

-- the MDR-like core of SUR is linked with the K(IR) pore in KATP channels PMID:

12213829

-- caveolin-3 negatively regulates Kir6.2/SUR2A channel function. PMID: 19481058

-- The prevalent Glu23Lys polymorphism in the potassium inward rectifier 6.2

(KIR6.2) gene is associated with impaired glucagon suppression in response to

hyperglycemia. PMID: 12196481

-- the common E23K genetic variant at the KCNJ11 gene locus was significantly

associated with cardiovascular function PMID: 17720745

Lists the most recent PubMed articles linked to this gene.

This information is enabled by either --pubmed or --all.

The number of articles returned is dependent on the --papernum option.

[+] Top Pubmed articles linked to gene KCNJ11, by date:

-- Zhao J et al. "Examination of type 2 diabetes loci implicates CDKAL1 as a

birth weight gene." Diabetes. 2009 Oct;58(10):2414-8. PMID: 19592620

-- Salanti G et al. "Underlying genetic models of inheritance in established

type 2 diabetes associations." Am J Epidemiol. 2009 Sep 1;170(5):537-45.

PMID: 19602701

-- Schulze MB et al. "Use of Multiple Metabolic and Genetic Markers to Improve

the Prediction of Type 2 Diabetes: the European Prospective Investigation

into Cancer and Nutrition (EPIC)-Potsdam study." Diabetes Care. 2009 Aug 31;.

PMID: 19720844

-- Reyes S et al. "K(ATP) channel Kir6.2 E23K variant overrepresented in human

heart failure is associated with impaired exercise stress response." Hum

Genet. 2009 Aug 14;. PMID: 19685080

-- Yoshida T et al. "Association of genetic variants with chronic kidney disease

in individuals with different lipid profiles." Int J Mol Med. 2009

Aug;24(2):233-46. PMID: 19578796

Lists PubMed articles that matched the user's search terms AND are linked to the gene listed here.

If multiple search terms are provided, this section will show papers that matched ANY of the search terms AND the gene.

For more in-depth searching, use --each-term. A section for each search term will then appear, showing the PubMed articles (listed by most recent first) linked to the gene and only that particular search term.

[+] Top Pubmed articles linked to gene KCNJ11 matching any search term:

-- Gach A et al. "Neonatal diabetes in a child positive for islet cell

antibodies at onset and Kir6.2 activating mutation." Diabetes Res Clin Pract.

2009 Nov;86(2):e25-e27. PMID: 19692135

-- 't Hart LM et al. "A Combined Risk Allele Score of Eight Type 2 Diabetes

Genes Is Associated With Reduced First Phase Glucose Stimulated Insulin

Secretion During Hyperglycemic Clamps." Diabetes. 2009 Oct 6;. PMID: 19808892

-- StancÃ¡kovÃ¡ A et al. "Association of 18 confirmed susceptibility loci for

type 2 diabetes with indices of insulin release, proinsulin conversion, and

insulin sensitivity in 5,327 nondiabetic Finnish men." Diabetes. 2009

Sep;58(9):2129-36. PMID: 19502414

-- Salanti G et al. "Underlying genetic models of inheritance in established

type 2 diabetes associations." Am J Epidemiol. 2009 Sep 1;170(5):537-45.

PMID: 19602701

-- Nikolac N et al. "Metabolic control in type 2 diabetes is associated with

sulfonylurea receptor-1 (SUR-1) but not with KCNJ11 polymorphisms." Arch Med

Res. 2009 Jul;40(5):387-92. PMID: 19766903

-- Ting WH et al. "Improved diabetic control during oral sulfonylurea treatment

in two children with permanent neonatal diabetes mellitus." J Pediatr

Endocrinol Metab. 2009 Jul;22(7):661-7. PMID: 19774848

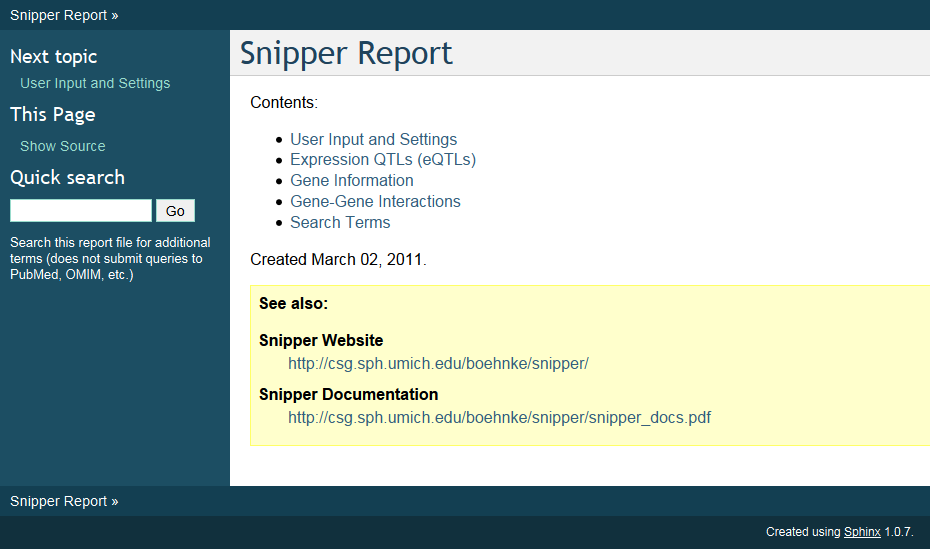
**Snipper HTML output**

This mode is the preferred method of running Snipper, and is also the default. Snipper will produce a formatted HTML report file in a directory of your choosing by doing the following:

|  |
| --- |
| **snipper** --snpfile yoursnps.txt –o my\_html\_directory |

If no directory is given using –o, a directory called “snipper\_report” is created by default.

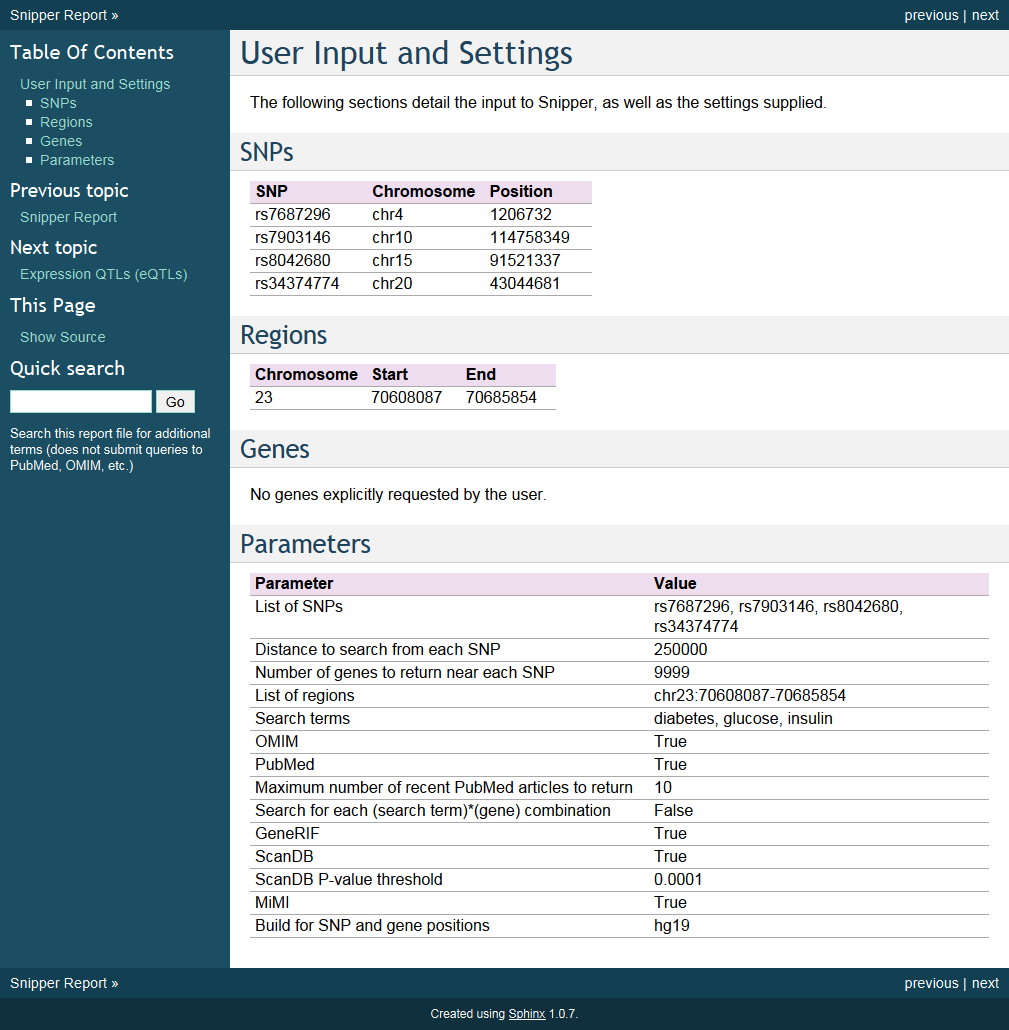
The HTML report begins with “index.html”, which has the table of contents. An image of this table (cropped to remove whitespace) is shown below.



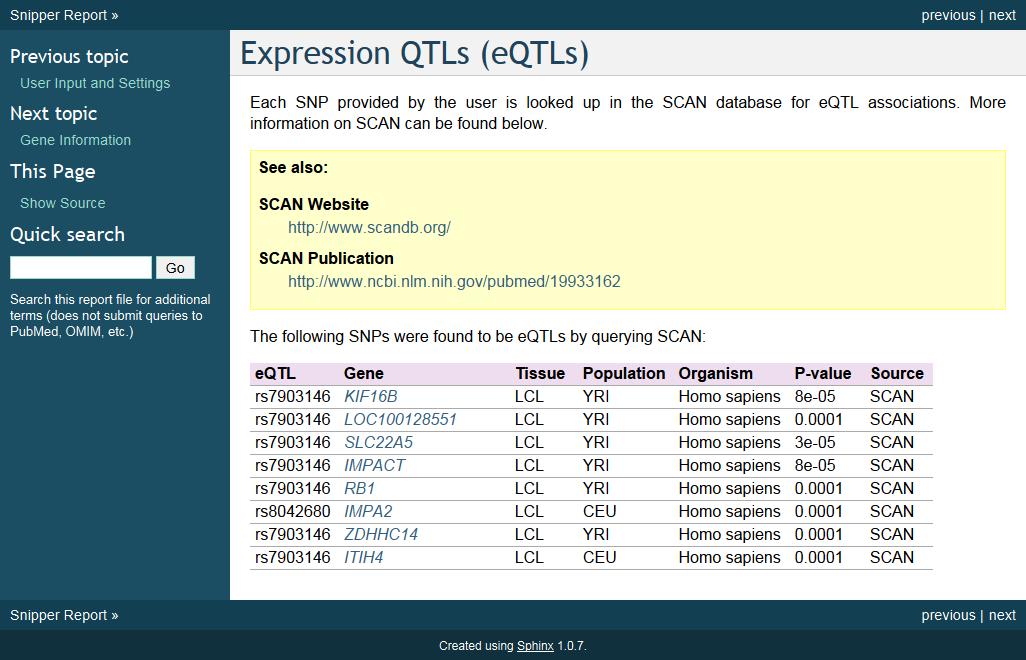
The report contains 5 main sections:

* User Input and Settings: lists all of the input SNPs and regions as given by the user, as well as all command line options that were specified
* Expression QTLs: contains information on genes whose expression is associated with SNPs given by the user
* Gene Information: information on each gene found near the input SNPs/regions
* Gene-Gene Interactions: lists direct interactions between all genes found near input SNPs/regions
* Search Terms: lists each search term given by the user, and where they matched within each gene’s information

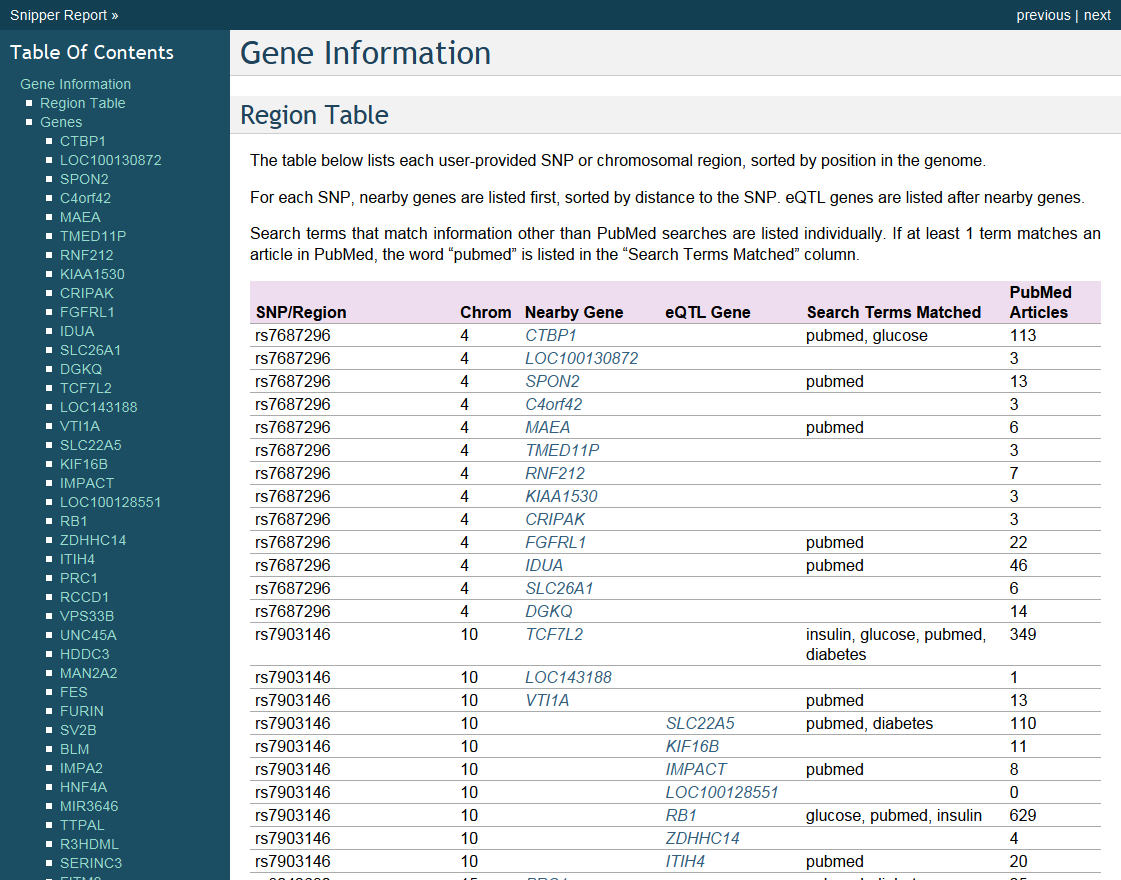
An example of the input section:

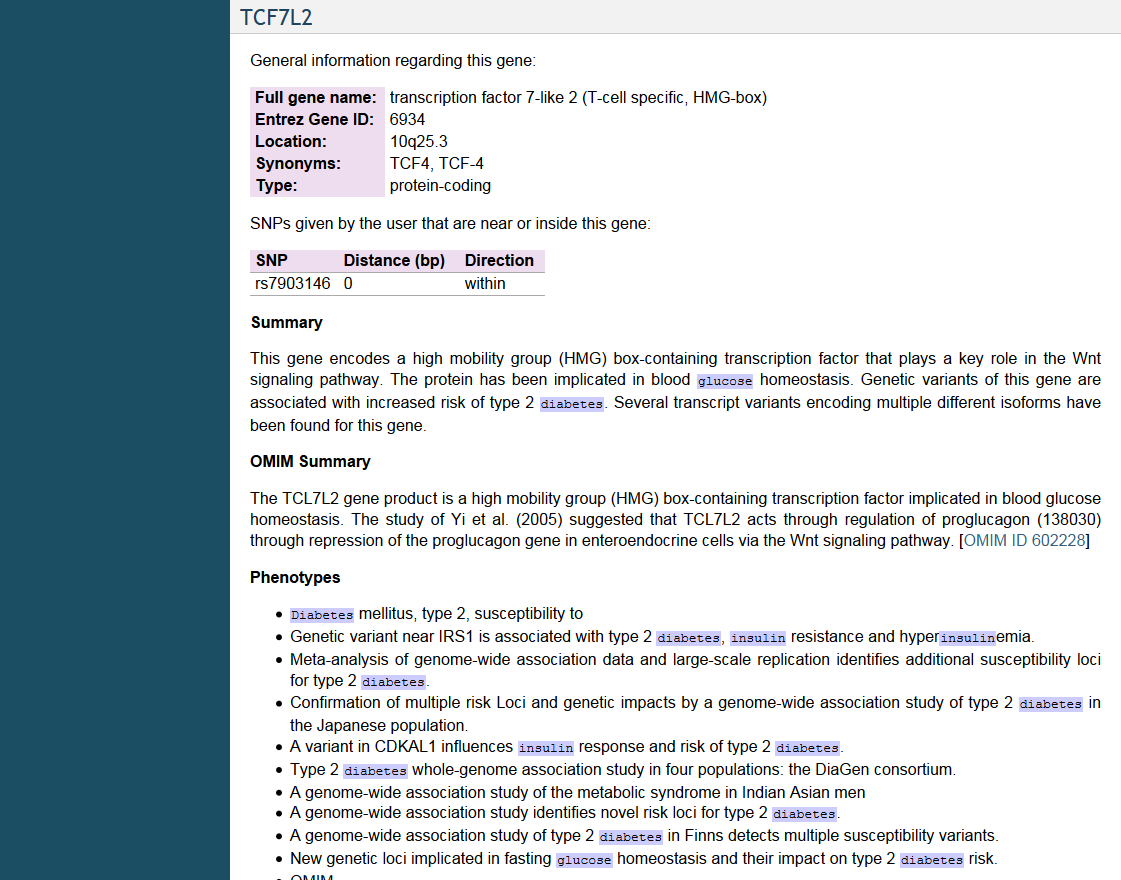


The eQTL report contains a list of genes that were associated with user input SNPs. Each gene is listed along with the information for the association as retrieved from the SCAN database.



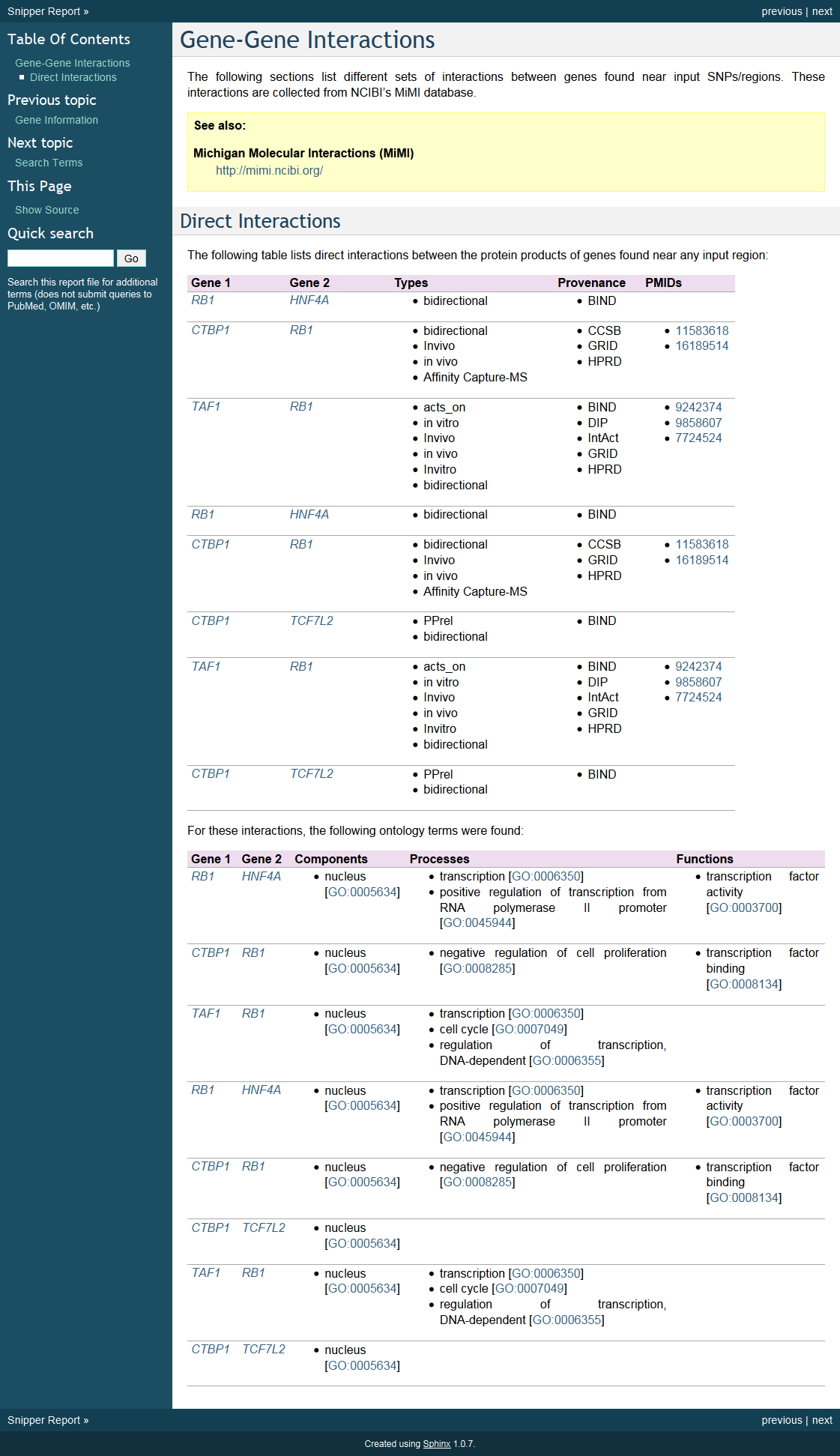
The gene report contains two sections: a table listing each gene found near SNPs, or those found in regions given by the user, and finally an individual section for each gene. An example:





The example is abbreviated and only includes a single gene, TCF7L2. Within this gene, every occurrence of the words “glucose”, “insulin”, and “diabetes” are highlighted (these are search terms supplied with --terms.)

The next section displays a list of direct interactions between all genes (those both near SNPs and within regions), along with their details as downloaded from MiMI. An example:



And finally, a section listing each search term provided by the user, and where each term matched within the information for each gene:



The “any” term is used when search terms are globbed together as 1 query, that is: “term1 OR term2 OR term3”. This makes searching PubMed much faster, but also not quite as specific. The user can supply the --each-term parameter, which forces Snipper to submit independent queries to PubMed for each search term + gene pair.

**Examples**

We list below a few examples of running Snipper by giving both a word description of what the program is doing, along with the command line parameters.

* Using Snipper on a single SNP (-s)
* Search 500 kb away (-d 500kb) from the SNP for genes
* List more PubMed articles than the default (--papernum 10)

|  |
| --- |
| **snipper** -s "rs1002227" -d 500kb --papernum 10 |

* Use Snipper on a file that contains SNP (rs#) names. The file can be of arbitrary format.
* Search 1 MB from each SNP in the file for genes, but return only the 3 nearest genes (-n) for each SNP
* Add search terms "diabetes","insulin","glucose" (--terms)

|  |
| --- |
| **snipper** --snpfile file\_with\_snps.txt -d 1MB -n 3 --terms “diabetes,glucose,insulin” |

**Building your own position database**

Snipper comes pre-loaded with a database file giving the positions of SNPs and genes for human genome build hg19 (UCSC).

To build your own database, a script has been provided in the bin/ directory called "build\_db.py". You simply need to execute this script with the human genome build for which you wish to build a database. For example:

|  |
| --- |
| **bin/build\_db.py --build hg18** |

This will create a database file called "hg18.db" in the data/genome/ directory, and add information about this newly created database to the conf file (conf/snipper.conf). To use this new database, you can use the --build parameter when running Snipper, like so:

|  |
| --- |
| **snipper** -s "rs7903146" --build hg18 |

The build\_db.py script always downloads the latest snp and refFlat tables for a given human genome build.

**Warning**: building the database can take many hours (2-3 at minimum.) You should allow ample time for the script to complete!

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