Supplementary Information:

CanvasDB: A local database infrastructure for analysis of targeted- and whole genome re-sequencing projects

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Supplementary Tables	2
Supplementary Information	
Implementation of rapid filtering analyses	
CanvasDB file formats	
SNP file format.	6
Indel file format	6
File format for adding new samples to the database	7

Supplementary Tables

Supplementary Table S1. Database structure for the SNP summary table

Column	Туре	Comment
SNP_id	string	Unique id for each SNP in the database
chr	string	Chromosome
pos	integer	SNP coordinate on chromosome
ref	string	Reference allele
alt	string	Alternative allele (SNP allele)
nr_samples ^a	integer	Number of samples in database where SNP is present.
samples ^a	blob	A binary text string consisting of samples where the SNP is present. Each sample is represented by its sample ID number. This binary text string is a comma separated string of sample ID numbers.
snp137 ^b	string	rs id from dbSNP (if SNP is present in dbSNP)
snp137common ^b	string	rs id from dbSNP common, a database of SNPs present in at least 1% frequency
class	string	Classification of SNP based on Annovar (intronic, exonic, nonsynonymous, splicing, synonymous etc)
severity	integer	A number from 1 to 5 representing the putative damaging effect of SNP. Variants that are changing the protein structure have high numbers, while intronic and non-coding variants have low numbers.
gene	string	Name of gene where SNP is located (if any)
details	string	Additional SNP information from Annovar, codon position, amino acid substitution etc.
sift	string	SIFT score
polyphen	string	PolyPhen score
phylop	string	PhyloP score
lrt	string	LRT score
mut_taster	string	MutTaster score
gerp	string	GERP score

^a The 'nr_samples' and 'samples' fields needs to be updated each time a new sample is added to the database.

b In this example the database is using annotations from v137 of dbSNP, this can be updated to later versions.

Supplementary Table S2. Database structure for the indel summary table

Column	Туре	Comment
indel_id	string	Unique id for each indel in the database
chr	string	Chromosome
start	integer	Indel start coordinate on chromosome
end	integer	Indel end coordinate on chromosome
ref	string	Reference allele
alt	string	Alternative allele (indel allele)
type	string	Indel type, i.e. insertion or deletion
size	integer	Size of indel
nr_samples ^a	integer	Number of samples in database where SNP is present.
samples ^a	blob	A binary text string consisting of samples where the SNP is present. Each sample is represented by its sample ID number. This binary text string is a comma separated string of sample ID numbers.
snp137 ^b	string	rs id from dbSNP (if indel is present in dbSNP)
snp137common ^b	string	rs id from dbSNP common, a database of SNPs present in at least 1% frequency
class	string	Classification of indel based on Annovar (intronic, frameshift, splicing, etc)
severity	integer	A number from 1 to 5 representing the putative damaging effect of SNP. Variants that are changing the protein structure have high numbers, while intronic and noncoding variants have low numbers.
gene	string	Name of gene where SNP is located (if any)
details	string	Additional SNP information from Annovar, codon position, amino acid substitution etc.

^a The 'nr_samples' and 'samples' fields needs to be updated each time a new sample is added to the database.

^b In this example the database is using annotations from v137 of dbSNP, this can be updated to later versions.

Supplementary Information

Implementation of rapid filtering analyses

The SNP and indel summary tables (see Supplementary Tables S1 and S2) are crucial for the to the rapid variant filtering in the *canvasDB* system. This section describes the main design of the filtering functionality. The filtering itself is performed by two functions in R, filterSNPs() and filterIndels(), which queries the summary tables. The filtering functions take the following main arguments:

inSamples	Group of samples expected to have a shared
	variant. Corresponds to `in-group' in Figure 3A.
filterSamples	Group of samples where variant should not occur,
	i.e. negative controls. Defaults to all other
	samples outside the 'in-group'. Corresponds to
	`filter-group' in Figure 3A.
discardSamples	Group of samples not used in the filtering.
	Defaults to an empty set. Corresponds to
	'discard-group' in Figure 3A.
minIn	Minimum number of individuals in the 'in-group'
	that are required to have a shared variant.
	Defaults to all samples in `in-group'.
maxOthers	Maximum number of individuals in the 'filter-
	group' that are allowed to carry the variant.
	Defaults to zero.
minSeverity	A score between 1-5 representing the potential
	damaging effect of the variant. Defaults to 3,
	meaning that only variants with a score of at
	least 3 are considered, i.e. amino acid sequence
	altering variants.
dbSNPfilterCommon	A flag indicating whether only rare variants (not
	present in dbSNPCommon) are to be considered.
	Defaults to FALSE.

When the filtering functions are called with the above arguments, the following steps are executed.

1. Based on the number of samples in the inSamples, filterSamples and discardSamples groups, as well as the arguments minIn and maxOthers, the two values minSamples and maxSamples are calculated, representing the minimal and maximal number of samples in which candidate variant should be found. Variants present in fewer samples than minSamples or more than maxSamples cannot fulfill the filtering criteria.

- 2. From the SNP or indel summary table only variants where nr_samples is in the range between minSamples and maxSamples are returned. This is done by a MySQL query that can be rapidly executed since the relevant columns of the summary tables are indexed. In the same query, filtering on minSeverity and dbSNPfilterCommon are performed (if those arguments were set in the function call). The resulting variants from the MySQL query are returned into R. This step can reduce the number of candidate variants dramatically.
- 3. The variants returned into R from step 2) are present in a number of samples that is consistent with the original filtering function call. For these remaining variants, the sample information is present in the binary text string samples. R immediately converts the binary samples string into a vector object, resulting in a numerical vector with sample IDs for each of the candidate variants. These vectors are then compared to the samples in the inSamples group using built-in vector matching functions in R. Those that fulfill the original filtering criteria having at least minIn samples present in the inSamples group represent the final set of candidate variants.
- 4. Once the candidate variants have been detected from the summary table (in steps 1-3 above) the actual data for each of the samples (read counts, quality scores etc) are fetched from the individual SNP and indel tables for each of the samples, after which the final results are formatted as a table that is returned to the user.

As described above, the filtering is essentially performed by one single MySQL query on indexed columns in a summary table and subsequent analyses of vector objects in R. This allows for very rapid execution for most types of filtering analyses.

CanvasDB file formats

When importing SNP and indel variant calls into *canvasDB*, the files are parsed into a specific format that can be imported into the database. These file formats for SNPs and indels are described in this section. In the system there are pre-defined functions for parsing some common variant call file formats (e.g. VCF) into the cdb files. However, it is also possible to parse other formats into cdb files. One advantage of parsing the files outside of the system is that it makes the import into the database faster.

SNP file format

A separate SNP file needs to be created for each individual sample. The *canvasDB* SNP file format is a tab delimited text file. Each row corresponds to a SNP and the following columns are required:

SNP_id	A unique ID for each SNP in the database. On the format
	`chr pos ref alt', e.g. chr1 879482 G C
Chr	Chromosome, e.g. chr1
Pos	SNP position on chromosome, e.g. 879482
Ref	Reference allele, e.g. G
Alt	Alternative allele, e.g. C
Cov	Read coverage
ref_reads	Number of reference reads
ref_starts	Number of reference reads with unique starting points
ref_qual	Quality score for reference allele
alt_reads	Number of alternative reads
alt_starts	Number of alternative reads with unique starting points
alt_qual	Quality score for alternative allele
Het	Heterozygosity flag. 0=homozygous, 1=heterozygous

Indel file format

The file format for indels is a tab delimited text file, similar to the SNP file format described above. Each row corresponds to an indel and the following columns are required:

indel id	A unique ID for each indel in the database. On the format
_	`chr start end ref alt', e.g. chr1 1276972 1276973 - CACA
chr	Chromosome, e.g. chr1
start	Indel start position on chromosome, e.g. 1276972
end	Indel end position on chromosome, e.g. 1276973
ref	Reference allele, e.g
alt	Alternative allele, e.g. CACA (insertion)
cov	Read coverage
ref_reads	Number of reference reads
alt_reads	Number of alternative reads
het	Heterozygosity flag. 0=homozygous, 1=heterozygous

File format for adding new samples to the database

The easiest way to add new samples into the database is to construct a text file holding information about multiple samples and send this file as a parameter to the the batchImport() function in R. This section describes the structure of the sample text file.

The sample file is a text file delimited by the character '|'. This implies that the '|' character may not be used in sample names, descriptions or any other fields included in the file. The file should include a header with the following fields:

canvasId	Sample identifier used within the canvasDB
	system. Each sequenced sample must have a
	unique canvasId. These ids are used in
	filtering analysis functions etc.
sampleName	Alternative name of the sequenced samples.
seq.platform	Sequencing technology used for the experiments
library.type	The library type used for the experiments, for
	example WholeExome or WholeGenome
read.type	Type of read and read length, for example
	Frag75, PE100x100 or similar
capture.method	Method used for capturing genomic target region
date	Date of the sequencing experiment
principal.investigator	Name/initials of principal investigator
instrument.name	Name of sequencing facility/instrument
gender	Gender of sequenced individual
comments	Any comments about experiment
geographic.location	Geographic origin of sample
phenotypes	Text string describing the phenotye
SNP.file	Path to SNP data file
indel.file	Path to indel data file
file.format	File format for SNP/indel files. Currently the
	following options are supported: CanvasDB, GATK
	(VCF), Lifescope, TorrentSuite (VCF)
reads.total	Total number of reads for the sample
reads.on.target	Number of reads mapping to target region