Velvet Manual - version 0.7

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For impatient people

```
> make
> ./velveth
> ./velvetg
> less sillyDirectory/stats.txt
```

- > ./velveth sillyDirectory 21 -shortPaired data/test_reads.fa
- > ./velvetg sillyDirectory

(Final graph has 16 nodes and n50 of 24184, max 44966, total 100080, using 0/142858 reads)

- > ./velvetg sillyDirectory -cov_cutoff 5 -read_trkg yes -amos_file yes (Final graph has 1 nodes and n50 of 99975, max 99975, total 99975, using 135862/142858 reads > less sillyDirectory/velvet_asm.afg
- > ./velvetg sillyDirectory -cov_cutoff auto (Final graph has 1 nodes and n50 of 99975, max 99975, total 99975, using 0/142858 reads)
- > ./velvetg sillyDirectory -exp_cov 19 -ins_length 100 (Final graph has 12 nodes and n50 of 99975, max 99975, total 100165, using 135855/142858 real
- > ./velvetg sillyDirectory -exp_cov auto (Final graph has 1 nodes and n50 of 99975, max 99975, total 99975, using 135862/142858 reads
- > ./velveth sillyDirectory 21 -short data/test_reads.fa -long data/test_long.fa > ./velvetg sillyDirectory -exp_cov 19 (Final graph has 10 nodes and n50 of 80927, max 80927, total 100122, using 137863/144858 real
- > ./velvetg sillyDirectory -exp_cov auto (Final graph has 1 nodes and n50 of 99975, max 99975, total 99975, using 137862/144858 reads

Installation

Requirements

Velvet should function on any standard 64bit Linux environment with gcc. A good amount of physical memory (12GB to start with, more is no luxury) is recommended.

It can in theory function on a 32bit environment, but such systems have memory limitations which might ultimately be a constraint for assembly.

2.2Compiling instructions

From a GNU environment, simply type:

2.3 Compilation settings

2.3.1 Colorspace Velvet

To produce the colorspace version of Velvet, compile with the instruction:

> make color

All the rest of the manual remains valid, except that the executables are now called velveth_de and velveth_de .

Beware that color- and sequence space are incompatible, hence separate sets of executables. In other words, don't try to hash sequence files with colorspace velvet or vice-versa, under penalty of meaningless results!

2.3.2 CATEGORIES

Because of the use of fixed-length arrays, a number of variables have to be set at compilation time.

One of the is the number of channels, or categories of reads, which can be handled independently. This is for example useful is you want to distinguish reads from different insert libraries, or from different samples altogether.

By default, there are only two short read categories, but this variable can be extended to your needs. For example, to obtain 57 different channels, compile with the parameter:

make 'CATEGORIES=57'

(Note the single quotes and absence of spacing.)

Obviously, the greater the number, the longer the corresponding arrays, the more memory will be required to run Velvet. Adjust this variable according to your needs and your memory requirements.

2.3.3 MAXKMERLENGTH.

Another useful compilation parameter is the MAXKMERLENGTH. As explained in 5.1, the hash length can be crucial to getting optimal assemblies. Depending on the dataset, you might wish to use long hash lengths.

By default, hash-lengths are limited to 31bp, but you can push up this limit by adjusting the MAXKMERLENGTH parameter at compilation time:

make 'MAXKMERLENGTH=57'

(Note the single quotes and absence of spacing.)

By storing longer words, Velvet will be requiring more memory, so adjust this variable according to your needs and memory resources.

3 Running instructions

3.1 Running velveth

Velveth helps you construct the dataset for the following program, velvetg, and indicate to the system what each sequence file represents.

If, on the command line, you forget the syntax, you can print out a short help message:

> ./velveth

Velveth takes in a number of sequence files, produces a hashtable, then outputs two files in an output directory (creating it if necessary), Sequences and Roadmaps, which are necessary to velvetg. The syntax is as follows:

The hash length, also known as k-mer length, corresponds to the length, in base pairs, of the words being hashed. See 5.1 for a detailed explanation of how to choose the hash length.

Supported file formats are:

```
fasta (default)
fastq
fasta.gz
fastq.gz
sam
bam
eland
gerald
Read categories are:
short (default)
shortPaired
short2 (same as short, but for a separate insert-size library)
shortPaired2 (see above)
long (for Sanger, 454 or even reference sequences)
longPaired
```

For concision, options are stable. In other words, they are true until contradicted by another operator. This allows you to write as many filenames as you wish without having to re-type identical descriptors. For example:

> ./velveth output_directory/ 21 -fasta -short solexa1.fa solexa2.fa solexa3.fa -long capillary.fa

In this example, all the files are considered to be in FASTA format, only the read category changes. However, the default options are "fasta" and "short", so the previous example can also be written as:

> ./velveth output_directory/ 21 solexa*.fa -long capillary.fa

3.1.1 Piping Velvet

If you wish to run velveth directly on the output of another program, without the cost of creating and reading a new sequence file, then you can pipe it. In that case the character '-' is used to represent the standard input within the command line.

Therefore, instead of having:

```
> myprogram > selected_reads.fa
> velveth directory 21 -fastq other_reads.fastq -fasta selected_reads
> velvetg directory (... parameters...)

You can simply do:
> myprogram | velveth directory 21 -fastq other_reads.fastq -fasta -
> velvetg directory (... parameters...)
```

3.1.2 Strand specific sequencing

If you are using a strand specific transcriptome sequencing protocol, you may wish to use this option for better results. Even though it can be used anywhere on the command line, it applies to all the dataset. To toggle it on, type:

> ./velveth output_directory/ 21 (...data files...) -strand_specific

3.2 Running velvetg

Velvetg is the core of Velvet where the de Bruijn graph is built then manipulated. Note that although velvetg saves some files during the process to avoid useless recalculations, the parameters are **not** saved from one run to the next. Therefore:

```
> ./velvetg output_directory/ -cov_cutoff 4
> ./velvetg output_directory/ -min_contig_lgth 100
...is different from:
```

> ./velvetg output_directory/ -cov_cutoff 4 -min_contig_lgth 100

This means you can freely play around with parameters, without re-doing most of the calculations:

```
> ./velvetg output_directory/ -cov_cutoff 4
> ./velvetg output_directory/ -cov_cutoff 3.8
> ./velvetg output_directory/ -cov_cutoff 7
> ./velvetg output_directory/ -cov_cutoff 10
> ./velvetg output_directory/ -cov_cutoff 2
```

On the other hand, within a single velvetg command, the order of parameters is not important.

Finally, if you have any doubt at the command line, you can obtain a short help message by typing:

> ./velvetg

3.2.1 Single reads

Initally, you simply run:

> ./velvetg output_directory/

This will produce a fasta file of contigs and output some statistics. Experience shows that there are many short, low-coverage nodes left over from the intial correction. Determine as you wish a coverage cutoff value (cf. 5.2), say 5.2x, then type:

> ./velvetg output_directory/ -cov_cutoff 5.2

On the other hand, if you want to exclude highly covered data from your assembly (e.g. plasmid, mitochondrial, and chloroplast sequences) you can use a maximum coverage cutoff:

```
> ./velvetg output_directory/ -max_coverage 300 (... other parameters ...)
```

The coverage cutoff can also be set automatically to half the length weighted median contig coverage depth. Although you may wish to optimise this parameter in further iterations, this option allows you to quickly obtain a decent assembly in your first run. To do so, type:

> ./velvetg output_directory/ -cov_cutoff auto

Note that typing $-exp_cov$ auto (cf. next subsection) is equivalent to typing $-exp_cov$ auto $-cov_cutoff$ auto, unless you decide to manually set the coverage cutoff. On the contrary, $-cov_cutoff$ auto does not affect the expected coverage setting.

3.2.2 Adding long reads

Reminder: you must have flagged your long reads as such when running velveth (cf. 3.1).

If you have a sufficient coverage of short reads, and any quantity of long reads (obviously the deeper the coverage and the longer the reads, the better), you can use the long reads to resolve repeats in a greedy fashion.

To do this, Velvet needs to have a reasonable estimate of the expected coverage in short reads of unique sequence (see 5.1 for a definition of k-mer coverage). The simplest way to obtain this value is simply to observe the distribution of contig coverages (as described in 5.3), and see around which value the coverages of nodes seem to cluster (especially the longer nodes in your dataset). Supposing the expected coverage is 19x, them you indicate it with the exp_cov marker:

```
> ./velvetg output_directory/ -exp_cov 19 (... other parameters ...)
```

If you have reasons to believe that the coverage is reasonably uniform over your sample, you can request that Velvet estimate it for you, by typing:

```
> ./velvetg output_directory/ -exp_cov auto (... other parameters ...)
```

This will set exp_cov to the length weighted median conting coverage, and cov_cutoff to half that value (unless you override it with a manual setting).

Caveat emptor: if this option is used on datasets with very heterogeneous coverage (e.g. transcriptomes, ChIP-seq, etc) then the results are simply unpredictable, and mis-assemblies are to be expected. This option is intended mainly for standard genomic sequencing.

3.2.3 Paired-ends reads

Reminder: you must have flagged your reads as being paired-ends when running velveth (cf. 3.1).

To activate the use of read pairs, you must specify two parameters: the expected (i.e. average) insert length (or at least a rough estimate), and the expected short-read k-mer coverage (see 5.1 for more information). The insert length is understood to be the length of the sequenced fragment, i.e. it includes the length of the reads themselves. If you expect your insert length to be around 400bp, and your coverage to be around 21.3x, you would type:

If you happen to have hashed paired long reads and you ordered them as explained in 4.1 you can also tell Velvet to use this information for scaffolding by indicating the corresponding insert length (remember that you still need to indicate the short-read k-mer coverage):

```
> ./velvetg output_directory/ -exp_cov 21 -ins_length_long 40000
(... other parameters ...)
```

As with long reads, if you have reasons to believe that the coverage is reasonably uniform over your sample, you can request that Velvet estimate it for you, by typing:

```
> ./velvetg output_directory/ -exp_cov auto (... other parameters ...)
```

This will set exp_cov to the length weighted median contig coverage, and cov_cutoff to half that value (unless you override it with a manual setting).

Caveat emptor: if this option is used on datasets with very heterogeneous coverage (e.g. transcriptomes, ChIP-seq, etc) then the results are simply unpredictable, and mis-assemblies are to be expected. This option is intended mainly for standard genomic sequencing.

Automatically measuring the insert length

If the insert length of a library is unspecified, Velvet will attempt to measure it for you, based on the read-pairs which happen to map onto a common node. The corresponding $ins_length*$ and $ins_length*_sd$ parameters are thus set for you. This option is designed as a convenience if your assembly is initially good enough to provide long contigs to bootstrap the process. If that is not the case, the library insert length will be determined from a small sample size, and is liable to produce a poor estimate. You should check the Velvet console output to make sure the insert length estimates are not absurd.

Scaffolding By default, Velvet will try to scaffold contigs that it cannot quite connect. This results in sequences of Ns in the contigs.fa file, which correspond to the estimated distance between two neighbouring contigs. If you do not want this scaffolding, you can turn it off with the following switch:

```
> ./velvetg output_directory/ -exp_cov 21 -ins_length_long 200
-scaffolding no (... other parameters ...)
```

Standard deviations This is a more subtle point which you can ignore if you have only one dataset of paired-end reads or if the standard deviation (SD) of the insert lengths is roughly proportional to the expected length (e.g. if the insert-lengths are described as $length \pm p\%$).

Velvet does not use the absolute values of the insert-length SDs, but their relative values. Therefore, you do not need to spend too much time on the estimation of the SDs, as long as you are consistent. You can then enter your own *a priori* SD's. To do so simply indicate them as follows:

```
> ./velvetg output_directory/ -exp_cov 21
-ins_length 200 -ins_length_sd 20
-ins_length2 20000 -ins_length2_sd 5000
-ins_length_long 40000 -ins_length_long_sd 1000
(... other parameters ...)
```

3.2.4 Controlling Velvet's output

Selecting contigs for output By default, Velvet will print out as many contigs as possible. This has the drawback of potentially flooding the output

with lots of unwanted very short contigs, which are hardly useable in a significant way. If you wish, you can request that the contigs in the contigs fa file be longer than a certain length, say 100bp:

```
> ./velvetg -min_contig_lgth 100 (... other parameters ...)
```

Using read tracking Velvet's read tracking can be turned on with the read-tracking option. This will cost slightly more memory and calculation time, but will have the advantage of producing in the end a more detailed description of the assembly:

```
> ./velvetg output_directory/ -read_trkg yes (... other parameters ...)
```

Producing an .afg file If you turn on the read tracking, you might also want to have all the assembly information in one datastructure. For this purpose Velvet can produce AMOS files (cf 4.2.3). Because the .afg files tend to be very large, they are only produced on demand:

```
> ./velvetg output_directory/ -amos_file yes (... other parameters ...)
```

Using multiple categories You can be interested in keeping several kinds of short read sets separate. For example, if you have two paired-end experiments, with different insert lengths, mixing the two together would be a loss of information. This is why Velvet allows for the use of 2 short read channels (plus the long reads, which are yet another category).

To do so, you simply need to use the appropriate options when hashing the reads (see 3.1). Put the shorter inserts in the first category. Supposing your first readset has an insert length around 400bp and the second one a insert length around 10,000bp, you should type:

```
> ./velvetg output_directory/ -ins_length 400 -ins_length2 10000
(... other parameters ...)
```

Note: Increasing the amount of categories is possible. It's simply a bit more expensive memory-wise.

Note: In the stats.txt file, you will find all three categories (long, short1 and short2) treated separately.

Obtaining reads unused in the assembly If you wish to obtain the reads which do not belong in the assembly, you should type:

```
> ./velvetg output_directory/ -unused_reads yes (... other parameters ...)

The reads will be copied into a FastA file called UnusedReads.fa .
```

3.3 Advanced parameters: Tour Bus

Caveat Emptor

The following parameters are probably best left untouched. If set unwisely, Velvet's behaviour may be unpredictable.

Nonetheless, some users are curious to control the way in which Tour Bus (cf. 7) decides whether to merge polymorphisms or not.

Before we go into the actual details, it is worth discussing the pros and cons of bubble smoothing. The original idea is that a few SNPs, in the case of diploid assembly, should not prevent the construction of an overall contig. Detecting them post assembly is just a matter of scanning the assembly files and detecting discrepancies between the consensus sequence and the reads.

On the other hand, if you have two copies of a repeat in a haploid genome, you want to reduce the merging to a minimum, so that later analysis with paired-end reads or long reads may allow you to retrieve both individual copies, instead of just one artificial "consensus" sequence.

Hopefully, these issues will eventually be resolved by further thought and experiment. In the mean time, Velvet allows direct access to these parameters for those who want to play around, or maybe tailor Velvet to specific needs (e.g. multi-strain sequencing).

Maximum branch length Partly for engineering issues and partly to avoid aberrant transformations, there is a limit as to how long two paths must before simplification. By default, it is set a 100bp. This means that Velvet will not merge together two sequences which are sufficiently divergent so as not to have any common k-mer over 100bp. If you want to allow greater simplifications, then you can set this length to, say, 200bp:

```
> ./velvetg output_directory/ -max_branch_length 200 (...other parameters...)
```

Maximum divergence rate After aligning the two sequences with a standard dynamic alignment, Velvet compares the number of aligned pairs of nucleotides to the length of the longest of the two sequences. By default, Velvet will not simplify two sequences if they are more than 20% diverged. If you want to change that limit to 33%:

```
> ./velvetg output_directory/ -max_divergence 0.33 (...other parameters...)
```

Maximum gap count After aligning the two sequences with a standard dynamic alignment, Velvet compares the number of aligned pairs of nucleotides to the length of the longest of the two sequences. By default, Velvet will not simplify to sequences if more than 3bp of the longest sequence are unaligned.

```
> ./velvetg output_directory/ -max_gap_count 5 (...other parameters...)
```

3.4 Advanced parameters: Rock Band

Minimum long read connection cutoff When using long reads to connect and complete contigs, Velvet applies a simple multiplicity cutoff (by default 2) to reduce the amount of noise. In other words, at least two long reads are required to validate a connection. Nonetheless, you may want to lower this cutoff to 0 if the long sequences you are using are in fact high confidence non-overlapping sequences, such as contigs. On the contrary, if your long read coverage is very high, you might consider raising this cutoff.

3.5 Advanced parameters: Pebble

Minimum read-pair validation Velvet will by default assume that paired end reads are perfectly placed. With experimental data this assumption can be contradicted by occasional mis-pairings or by incorrect mappings of the reads because of errors. To avoid being misled by random noise, and therefore avoid missassemblies, Velvet requires that a connection between two contigs be corroborated by at least 10 mate pairs. If you want to change this cutoff to, say, 20, simply type:

> ./velvetg output_directory/ -min_pair_count 20 (...other parameters...)

4 File formats

4.1 Input sequence files

Velvet works mainly with fasta and fastq formats.

For paired-end reads, the assumption is that each read is next to its mate read. In other words, if the reads are indexed from 0, then reads 0 and 1 are paired, 2 and 3, 4 and 5, etc.

If for some reason you have forward and reverse reads in two different FASTA files but in corresponding order, the bundled Perl script *shuffleSequences_fasta.pl* will merge the two files into one as appropriate. To use it, type:

> ./shuffleSequences_fasta.pl forward_reads.fa reverse_reads.fa output.fa

There is also a *shuffleSequences_fastq.pl* script for Sanger-style FastQ files.

Concerning read orientation, Velvet expects paired-end reads to come from opposite strands facing each other, as in the traditional Sanger format. If you have paired-end reads produced from circularisation (i.e. from the same strand), it will be necessary to replace the first read in each pair by its reverse complement before running velveth.

4.2 Output files

After running Velvet you will find a number of files in the output directory:

4.2.1 The contigs.fa file

This fasta file contains the sequences of the contigs longer than 2k, where k is the word-length used in velveth. If you have specified a min_contig_lgth threshold, then the contigs shorter than that value are omitted.

Note that the length and coverage information provided in the header of each contig should therefore be understood in k-mers and in k-mer coverage (cf. 5.1) respectively.

The N's in the sequence correspond to gaps between scaffolded contigs. The number of N's corresponds to the estimated length of the gap. For reasons of compatibility with the archives, any gap shorter than 10bp is represented by a sequence of 10 N's.

4.2.2 The stats.txt file

This file is a simple tabbed-delimited description of the nodes. The column names are pretty much self-explanatory. Note however that node lengths are given in k-mers. To obtain the length in nucleotides of each node you simply need to add k-1, where k is the word-length used in velveth.

The in and out columns correspond to the number of arcs on the 5' and 3' ends of the contig respectively.

The coverages in columns *short1_cov*, *short1_Ocov*, *short2_cov*, and *short2_Ocov* are provided in k-mer coverage (5.1).

Also, the difference between *_cov and *_Ocov is the way these values are computed. In the first count, slightly divergent sequences are added to the coverage tally. However, in the second, stricter count, only the sequences which map perfectly onto the consensus sequence are taken into account.

4.2.3 The velvet_asm.afg file

This file is mainly designed to be read by the open-source AMOS genome assembly package. Nonetheless, a number of programs are available to transform this kind of file into other assembly file formats (namely ACE, TIGR, Arachne and Celera). See http://amos.sourceforge.net/ for more information.

The file describes all the contigs contained in the contigs fa file (cf 4.2.1).

If you are overwhelmed by the size of the file, two bundled scripts provided by Simon Gladman can help you out:

- asmbly_splitter.pl breaks down the original .afg file into individual files for each contig,
- snp_view.pl allows you to print out a simple ASCII alignment of reads around a given position on a contig.

4.2.4 The LastGraph file

This file describes in its entirety the graph produced by Velvet, in an idiosyncratic format which evolved with my PhD project. The format of this file is

briefly as follows:

• One header line for the graph:

```
$NUMBER_OF_NODES $NUMBER_OF_SEQUENCES $HASH_LENGTH
```

• One block for each node:

```
NODE $NODE_ID $COV_SHORT1 $0_COV_SHORT1 $COV_SHORT2 $0_COV_SHORT2 $ENDS_OF_KMERS_OF_NODE $ENDS_OF_KMERS_OF_TWIN_NODE
```

Note that the ends of k-mers correspond to the last nucleotides of the k-mers in the node. This means that the two sequences given above are not reverse-complements of each other but reverse complements shifted by k nucleotides. The common length of these sequences is equal to the length of the corresponding contig minus k-1.

See 4.2.2 for an explanation of O_COV values.

• One line for each arc:

```
ARC $START_NODE $END_NODE $MULTIPLICITY
```

Note: this one line implicitly represents an arc from node A to B and another, with same multiplicity, from -B to -A.

• For each long sequence, a block containing its path:

```
SEQ $SEQ_ID
$NODE_ID $OFFSET_FROM_START $START_COORD $END_COORD $OFFSET_FROM_END
$NODE_ID2 etc.
```

The offset variables are distances from the edges of the nodes whereas the start and end coordinates are correspond to coordinates within the read sequence.

• If short reads are tracked, for every node a block of read identifiers:

```
NR $NODE_ID $NUMBER_OF_SHORT_READS
$READ_ID $OFFSET_FROM_START_OF_NODE $START_COORD
$READ_ID2 etc.
```

5 Practical considerations / Frequently asked questions

5.1 Choice of hash length k

The hash length is the length of the k-mers being entered in the hash table. Firstly, you must observe three technical constraints:

- it must be an odd number, to avoid palindromes. If you put in an even number, Velvet will just decrement it and proceed.
- it must be below or equal to MAXKMERHASH length (cf. 2.3.3, by default 31bp), because it is stored on 64 bits
- it must be strictly inferior to read length, otherwise you simply will not observe any overlaps between reads, for obvious reasons.

Now you still have quite a lot of possibilities. As is often the case, it's a tradeoff between specificity and sensitivity. Longer kmers bring you more specificity (i.e. less spurious overlaps) but lowers coverage (cf. below)...so there's a sweet spot to be found with time and experience.

We like to think in terms of "k-mer coverage", i.e. how many times has a k-mer been seen among the reads. The relation between k-mer coverage C_k and standard (nucleotide-wise) coverage C is $C_k = C*(L-k+1)/L$ where k is your hash length, and L you read length.

Experience shows that this kmer coverage should be above 10 to start getting decent results. If C_k is above 20, you might be "wasting" coverage. Experience also shows that empirical tests with different values for k are not that costly to run!

5.2 Choice of a coverage cutoff

Velvet was designed to be explicitly cautious when correcting the assembly, to lose as little information as possible. This consequently will leave some obvious errors lying behind after the Tour Bus algorithm (cf. 7) was run. To detect them, you can plot out the distribution of k-mer coverages (5.1), using plotting software (I use R).

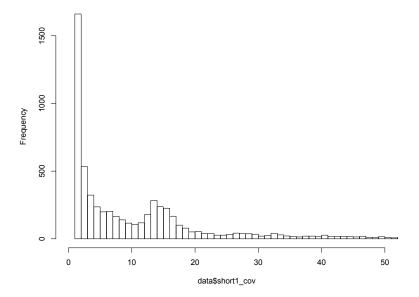
The examples below are produced using the S. suis P1/7 data available from the Sanger Institute (www.sanger.ac.uk/Projects/S_suis/) (Note: a simple script is necessary to convert the sequence files to FastA). I used a k-mer length of 21 and no cutoff.

With the R instruction:

- (R) > data = read.table("stats.txt", header=TRUE)
- (R) > hist(data\$short1_cov, xlim=range(0,50), breaks=1000000)

... vou can obtain:

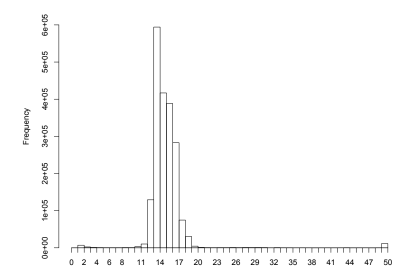
Histogram of data\$short1_cov



However, if you weight the results with the node lengths (you need to install the plotrix package for R to do this):

- (R) > library(plotrix)
- (R) > weighted.hist(data\$short1_cov, data\$lgth, breaks=0:50)

...you obtain:



The comparison of these two plots should convince you that below 7 or 8x you find mainly short, low coverage nodes, which are likely to be errors. Set the exact cutoff at your discretion.

However beware that there is such a thing as an over-aggressive cutoff, which could create mis-assemblies, and destroy lots of useful data.

If you have read-pair information, or long reads, it may be profitable to set a low coverage cutoff and to use the supplementary information resolve the more ambiguous cases.

5.3 Determining the expected coverage

From the previous weighted histogram it must be pretty clear that the expected coverage of contigs is near 14x.

5.4 Visualising contigs and assemblies

This section will be quite vague, as there are a number of solutions currently available, and presumably new ones under development. The following indications are just hints, as I have not done any exhaustive shopping nor benchmarking.

Most assembly viewers require an assembly format, which come in a variety of shapes and colours: ACE, AMOS, CELERA, TIGR, etc. Velvet only ouputs AMOS .afg files, but these can easily be converted with open-source software (amos.sourceforge.net).

5.5 What's long and what's short?

Velvet was pretty much designed with micro-reads (e.g. Illumina) as short and short to long reads (e.g. 454 and capillary) as long. Reference sequences can also be thrown in as long.

That being said, there is no necessary distinction between the types of reads. The only constraint is that a short read be shorter than 32kb. The real difference is the amount of data Velvet keeps on each read. Short reads are presumably too short to resolve many repeats, so only a minimal amount of information is kept. On the contrary, long reads are tracked in detail through the graph.

This means that whatever you call your reads, you should be able to obtain the same initial assembly. The differences will appear as you are trying to resolve repeats, as long reads can be followed through the graph. On the other hand, long reads cost more memory. It is therefore perfectly fine to store Sanger reads as "short" if necessary.

6 Contributed software

The Velvet package is bundled with programs developed by other programers which could be useful to Velvet users:

afg_handling by Simon Gladman (Simon.Gladman@csiro.au)

These two scripts allow you to examine the (generally) large .afg files which can be produced by Velvet:

asmbly_splitter allows you to choose a specific scaffold from the assembly and produce a self-standing .afg file for that scaffold.

snp_view produces an ASCII pileup display of the reads above a given locus

layout by Paul Harrison (pfh@logarithmic.net)

This script converts a (Last)Graph file into a .dot file, which can then be converted into an image by GraphViz (www.graphviz.org). This allows you to directly observe the topology of a graph.

VelvetOptimiser by Simon Gladman (Simon.Gladman@csiro.au)

This script automatically finds the optimal parameter settings for Velvet.

estimate-exp_cov by Torsten Seeman (torsten.seemann@infotech.monash.edu.au)

This script automatically determines the expected coverage value as described in the manual, and displays an ASCII histogram, thus obviating the need to start R for each Velvet run.

fasta2agp by David Studholme (david.studholme@tsl.ac.uk)

This script converts a Velvet assembly in FastA format with N's in the gaps into a AGP file which can be submitted to the EMBL or the NCBI.

extractContigReads by Daniel Zerbino (zerbino@ebi.ac.uk), suggested by Jasper Rees

This script scans the Graph2 file produced by Velvet and produces a FastA file of all the reads which belong to a given contig.

observed-insert-lengths by Daniel Zerbino

This scripts scans the Graph2 file produced by velvetg then computes and displays the insert length distribution of a chosen short read library in the assembly.

shuffleSequences by Eric Cabot (ecabot@wisc.edu) Peter (peter@maubp.freeserve.co.uk) and Daniel Zerbino (zerbino@ebi.ac.uk)

Alternative ways to efficiently shuffle your reads produced in the language of your choice: C, BioPython, Perl or Bash.

show_repeata by Ken Doig (kdd@doig.org)

Plots out the length of the larger repeated contigs in the assembly.

7 For more information

Publication: For more information on the theory behind Velvet, you can turn to:

D.R. Zerbino and E. Birney. 2008. Velvet: algorithms for de novo short read assembly using de Bruijn graphs. *Genome Research*, **18**: 821-829

Please use the above reference when citing Velvet.

Webpage: For general information and FAQ, you can first take a look at www.ebi.ac.uk/~zerbino/velvet.

Mailing list: For questions/requests/etc. you can subscribe to the users' mailing list: velvet-users@ebi.ac.uk.

To do so, see listserver.ebi.ac.uk/mailman/listinfo/velvet-users .

Contact emails: For specific questions/requests you can contact us at the following addresses:

- Daniel Zerbino < zerbino@ebi.ac.uk >
- Ewan Birney:

 birney@ebi.ac.uk>

Reporting bugs: We are very grateful to all the people who send us bugs. However, to speed up the process and avoid useless delays, please:

- 1. ensure that you have the very last version of Velvet, to the last digit, as displayed on the website.
- 2. attach to your e-mail the Log file from within the Velvet directory.
- 3. if the program crashed and created a core dump file could you please:
 - (a) destroy the core.* file
 - (b) recompile Velvet with the instruction "make debug"
 - (c) re-run Velvet and let it crash (therefore creating a new core file)
 - (d) launch the GNU debugger with the instructions:
 - > gdb ./velvetg core.*
 - (e) within gdb, request a backtrace:
 - (gdb) bt
 - (f) send the listing with the entire gdb session.