

Sequence assembly and mapping with MIRA 5



The Definitive Guide

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Contents

1	Introduction to MIRA	1
1.1	What is MIRA?	1
1.2	What to read in this manual and where to start reading?	1
1.3	The MIRA quick tour	2
1.4	For which data sets to use MIRA and for which not	3
1.4.1	Genome de-novo	3
1.4.2	Genome mapping	4
1.4.3	De-novo ESTs / RNASeq	4
1.4.4	Mapping EST / RNASeq	4
1.5	Any special features I might be interested in?	4
1.5.1	MIRA learns to discern non-perfect repeats, leading to better assemblies	4
1.5.2	MIRA has integrated editors for data from Sanger, 454, IonTorrent sequencing	7
1.5.3	MIRA lets you see why contigs end where they end	11
1.5.4	MIRA tags problematic decisions in hybrid assemblies	13
1.5.5	MIRA for polishing of PacBio or ONT assemblies	14
1.5.6	MIRA allows older finishing programs to cope with amount data in Illumina mapping projects	14
1.5.7	MIRA tags SNPs and other features, outputs result files for biologists	16
1.5.8	MIRA has ... much more	18
1.6	Versions, Licenses, Disclaimer and Copyright	18
1.6.1	Versions	18
1.6.2	License	19
1.6.2.1	MIRA	19
1.6.2.2	Documentation	19
1.6.3	Copyright	19
1.6.4	External libraries	19
1.7	Getting help / Mailing lists / Reporting bugs	19
1.8	Author	20
1.9	Miscellaneous	20
1.9.1	Citing MIRA	20
1.9.2	Postcards, gold and jewellery	20

2	Installing MIRA	21
2.1	Where to fetch MIRA	21
2.1.1	Package naming and versioning scheme	21
2.1.2	SourceForge: source package and precompiled binaries	21
2.1.3	GitHub: git repository, source, and binary packages	22
2.2	Installing from a precompiled binary package	22
2.3	Integration with third party programs (gap4, consed)	23
2.4	Compiling MIRA yourself	23
2.4.1	Prerequisites	23
2.4.2	Compiling and installing	24
2.4.3	Configure arguments for MIRA	24
2.4.3.1	Configure arguments for the BOOST library	24
2.4.3.2	MIRA specific configure arguments	24
2.5	Installation walkthroughs	25
2.5.1	(K)Ubuntu 18.04	25
2.5.2	openSUSE 12.1	26
2.5.3	Fedora 17	26
2.5.4	Mac OSX	27
2.5.5	Compile everything from scratch	28
2.5.6	Dynamically linked MIRA	28
2.6	Compilation hints for other platforms.	29
2.6.1	NetBSD 5 (i386)	29
2.7	Notes for distribution maintainers / system administrators	29
2.7.1	Additional data files	29
3	Preparing data	30
3.1	Introduction	30
3.2	Sanger	30
3.3	Roche / 454	30
3.4	Illumina	30
3.5	Ion Torrent	31
3.6	Short Read Archive (SRA)	31
4	Cookbook: de-novo genome assemblies	32
4.1	Introduction	32
4.2	General steps	32
4.2.1	Copying and naming the sequence data	33
4.2.2	Writing a simple manifest file	33
4.2.3	Starting the assembly	34

4.3	Manifest files for different use cases	34
4.3.1	Manifest for shotgun data	34
4.3.2	Assembling with multiple sequencing technologies (hybrid assemblies)	34
4.3.3	Manifest for data sets with paired reads	35
4.3.4	De-novo with multiple strains	37
5	Cookbook: Mapping	38
5.1	Introduction	38
5.2	General steps	38
5.2.1	Copying and naming the sequence data	39
5.2.2	Copying and naming the reference sequence	39
5.2.3	Writing a simple manifest file	40
5.2.4	Starting the assembly	40
5.3	Manifest files for different use cases	41
5.3.1	Mapping with shotgun data	41
5.3.2	Manifest for data sets with paired reads	41
5.3.3	Mapping with multiple sequencing technologies (hybrid mapping)	44
5.3.4	Mapping with multiple strains	45
5.3.5	Mapping EST / RNASeq data	46
5.4	Walkthroughs	47
5.4.1	Walkthrough: mapping of E.coli from Lenski lab against E.coli B REL606	47
5.5	Useful things to know about reference sequences	47
5.6	Known bugs / problems	48
6	Cookbook: de-novo EST / RNASeq assemblies	49
6.1	Introduction	49
6.2	Preliminaries: on the difficulties of assembling ESTs /RNASeq	49
6.2.1	Poly-A tails	50
6.2.2	Lowly expressed transcripts	50
6.2.3	Very highly expressed transcripts	50
6.2.4	Chimeras	50
6.3	Preprocessing of ESTs	51
6.4	The difference between <i>assembly</i> and <i>clustering</i>	52
6.4.1	Splitting transcripts into contigs based on SNPs	52
6.4.2	Splitting transcripts into contigs based on larger gaps	53
6.5	A simple step by step pipeline for reliable RNASeq assembly of eukaryotes	53
6.6	Solving common problems of EST assemblies	54

7	Working with the results of MIRA	55
7.1	MIRA output directories and files	55
7.1.1	The *_d_results directory	55
7.1.1.1	Additional 'large contigs' result files for de-novo assemblies of genomes	56
7.1.2	The *_d_info directory	56
7.2	First look: the assembly info	57
7.3	Converting results	59
7.3.1	Converting to and from other formats: miracore	59
7.3.2	Steps for converting data from / to other tools	59
7.3.2.1	Example: converting to and from the Staden package (gap4 / gap5)	60
7.3.2.2	Example: converting to and from SAM (for samtools, tablet etc.)	61
7.4	Filtering results	61
7.5	Places of importance in a de-novo assembly	62
7.5.1	Tags set by MIRA	62
7.5.2	Other places of importance	63
7.5.3	Joining contigs	63
7.5.3.1	Joining contigs at true repetitive sites	64
7.5.3.2	Joining contigs at "wrongly discovered" repetitive sites	65
7.6	Places of interest in a mapping assembly	66
7.6.1	Where are SNPs?	67
7.6.2	Where are insertions, deletions or genome re-arrangements?	68
7.6.3	Other tags of interest	72
7.7	Post-processing mapping assemblies	72
7.7.1	Manual cleanup and validation (optional)	72
7.7.2	Comprehensive SNP analysis spreadsheet tables (for Excel or OOcalc)	73
7.7.3	HTML files depicting SNP positions and deletions	74
7.7.4	WIG files depicting contig coverage or GC content	74
7.7.5	Comprehensive spreadsheet tables for gene expression values / genome deletions & duplications	74
8	Utilities in the MIRA package	75
8.1	miracore	75
8.1.1	Synopsis	75
8.1.2	Description	75
8.1.3	Options	75
8.1.3.1	General options	76
8.1.3.2	Options for input containing contig data	77
8.1.4	Examples	78
8.2	mirabait - a "grep" like tool to select reads with kmers up to 256 bases	78
8.2.1	Synopsis	78

8.2.2	Description	79
8.2.3	Options	79
8.2.3.1	Main options	79
8.2.3.2	File type options	80
8.2.3.3	Other options	81
8.2.4	Usage examples	81
8.2.5	Installing different rRNA databases	83
9	MIRA 5 reference manual	84
9.1	Synopsis	84
9.2	Requirements	84
9.3	Configuring an assembly: files and parameters	85
9.3.1	The manifest file: introduction	85
9.3.2	The manifest file: basics	86
9.3.3	The manifest file: information on the data you have	87
9.3.3.1	Starting a new readgroup	87
9.3.3.2	Defining data files to load	88
9.3.3.3	Setting default quality	90
9.3.3.4	Defining technology used to sequence	90
9.3.3.5	Preventing statistics for technologies with biases	90
9.3.3.6	Setting reference sequence for mapping jobs	90
9.3.3.7	Autopairing: letting MIRA find out pair info by itself	90
9.3.3.8	Setting size of read templates	91
9.3.3.9	Read segment placement	91
9.3.3.10	Read segment naming	93
9.3.3.11	Strain naming	94
9.3.3.12	Data directory for SCF files	94
9.3.3.13	Renaming read name prefixes	94
9.3.4	The manifest file: extended parameters	94
9.3.4.1	Parameter groups	95
9.3.4.2	Technology sections	95
9.3.4.3	Parameter short names	97
9.3.4.4	Order dependent quick switches	97
9.3.4.5	Parameter group: -GENERAL (-GE)	98
9.3.4.6	Parameter group: -ASSEMBLY (-AS)	99
9.3.4.7	Parameter group: -STRAIN/BACKBONE (-SB)	102
9.3.4.8	Parameter group: -DATAPROCESSING (-DP)	103
9.3.4.9	Parameter group: -CLIPPING (-CL)	103
9.3.4.10	Parameter group: -SKIM (-SK)	108

9.3.4.11	Parameter group: -KMERSTATISTICS (-KS)	110
9.3.4.12	Parameter group: -ALIGN (-AL)	112
9.3.4.13	Parameter group: -FINALMAPPING (-FM)	113
9.3.4.14	Parameter group: -CONTIG (-CO)	114
9.3.4.15	Parameter group: -EDIT (-ED)	116
9.3.4.16	Parameter group: -MISC (-MI)	117
9.3.4.17	Parameter group: -NAG_AND_WARN (-NW)	118
9.3.4.18	Parameter group: -DIRECTORY (-DIR, -DI)	119
9.3.4.19	Parameter group: -OUTPUT (-OUT)	119
9.4	Resuming / restarting assemblies	121
9.5	Input / Output	121
9.5.1	Directories	121
9.5.2	Filenames	122
9.5.2.1	Output	122
9.5.2.2	Assembly statistics and information files	122
9.5.3	File formats	123
9.5.4	STDOUT/STDERR	125
9.5.5	SSAHA2 / SMALT ancillary data	125
9.5.6	XML TRACEINFO ancillary data	126
9.5.7	Contig naming	127
9.5.8	Recovering strain specific consensus as FASTA	128
9.6	Tags used in the assembly by MIRA and EdIt	128
9.6.1	Tags read (and used)	128
9.6.2	Tags set (and used)	128
9.7	Where reads end up: contigs, singlets, debris	130
9.8	Detection of bases distinguishing non-perfect repeats and SNP discovery	130
9.9	Data reduction: subsampling vs. lossless digital normalisation	132
9.10	Caveats	133
9.10.1	Using data not from sequencing instruments: artificial / synthetic reads	133
9.10.2	Ploidy and repeats	134
9.10.3	Handling of repeats	134
9.10.3.1	Uniform read distribution	134
9.10.3.2	Keeping 'long' repetitive contigs separate	135
9.10.3.3	Helping finishing by tagging reads with HAF tags	135
9.10.4	Consensus in finishing programs (gap4, consed, ...)	136
9.10.5	Some other things to consider	137
9.11	Things you should not do	137
9.11.1	Do not run MIRA on NFS mounted directories without redirecting the tmp directory	137
9.11.2	Do not assemble without quality values	138

9.12	Useful third party programs	138
9.13	Speed and memory considerations	139
9.13.1	Estimating needed memory for an assembly project	139
9.13.2	Some numbers on speed	141
9.14	Known Problems / Bugs	141
9.15	TODOs	141
9.16	Working principles	141
9.17	See Also	142
10	Assembly of <i>hard</i> genome or EST / RNASeq projects	143
10.1	Getting 'mean' genomes or EST / RNASeq data sets assembled	143
10.1.1	For the impatient	143
10.1.2	Introduction to 'masking'	143
10.1.3	How does 'nasty repeat' masking work?	144
10.1.4	Selecting a "nasty repeat ratio"	144
10.2	How MIRA tags different repeat levels	144
10.3	The readrepeats info file	145
10.4	Pipeline to find worst contaminants or repeats in sequencing data	145
10.5	Examples for kmer statistics	148
10.5.1	Caveat: -SK:kmer_size	148
10.5.2	Sanger sequencing, a simple bacterium	148
10.5.3	454 Sequencing, a somewhat more complex bacterium	148
10.5.4	Solexa sequencing, E.coli MG1655	150
10.5.5	(NEED EXAMPLES FOR EUKARYOTES)	151
10.5.6	(NEED EXAMPLES FOR PATHOLOGICAL CASES)	151
11	Description of sequencing technologies	152
11.1	Introduction	152
11.2	Illumina (formerly Solexa)	152
11.2.1	Caveats for Illumina data	152
11.2.2	Illumina highlights	152
11.2.2.1	Quality	152
11.2.3	Lowlights	153
11.2.3.1	Long homopolymers	153
11.2.3.2	The GGCxG and GGC motifs	153
11.2.3.3	Chimeric reads	156
11.2.3.4	Sample barcode misidentification	157
11.2.3.5	Nextera library prep	157
11.2.3.6	Strong GC bias in some Solexa data (2nd half 2009 until advent of TruSeq kit at end of 2010)	157

11.3	Ion Torrent	159
11.3.1	Homopolymer insertions / deletions	160
11.3.2	Sequencing direction dependent insertions / deletions	161
11.3.3	Coverage variance	162
11.3.4	GC bias	162
11.3.5	Other sources of error	162
11.3.6	Where to find further information	162
11.4	Pacific BioSciences	162
11.4.1	Highlights	163
11.4.1.1	Sequence lengths	163
11.4.1.2	GC bias	163
11.4.1.3	Accuracy of corrected reads	163
11.4.1.4	Assemblies of corrected reads	163
11.4.2	Lowlights	163
11.4.2.1	Naming confusion	163
11.4.2.2	Forward / reverse chimeric sequences	164
11.4.2.3	Accuracy of uncorrected subreads	164
11.4.2.4	Immense need for CPU power	164
11.4.2.5	Increased quality requirements for clean DNA sample prep	164
12	Some advice when going into a sequencing project	165
12.1	Talk to your sequencing provider(s) before sequencing	165
12.2	Choosing a sequencing provider	165
12.2.1	WHAT DO YOU WANT?!	166
12.2.2	WHAT DO YOU NEED?!	167
12.2.3	WHAT WILL IT COST ME?	167
12.2.4	WHERE TO SEQUENCE?	167
12.2.5	Summary of all the above	168
12.3	Specific advice	169
12.3.1	Technologies	169
12.3.1.1	Sanger	169
12.3.1.2	Pacific Biosciences	169
12.3.1.3	Oxford Nanopore Technologies (ONT)	170
12.3.1.4	Illumina	170
12.3.1.5	Ion Torrent	170
12.3.1.6	Roche 454	170
12.3.2	Sequencing de-novo	170
12.3.3	Re-sequencing / mapping	170
12.4	A word or two on coverage ...	171

12.4.1	Low coverage isn't worth it	171
12.4.2	Catch-22: too high coverage	171
12.5	A word of caution regarding your DNA in hybrid sequencing projects	171
12.6	Advice for bacteria	172
12.6.1	Do not sample DNA from bacteria in exponential growth phase!	172
12.6.2	Beware of (high copy number) plasmids!	172
13	Bits and pieces	173
13.1	Using SSAHA2 / SMALT to screen for vector sequence	173
14	Frequently asked questions	175
14.1	Assembly quality	175
14.1.1	What is the effect of uniform read distribution (-AS:urd)?	175
14.1.2	There are too many contig debris when using uniform read distribution, how do I filter for "good" contigs?	176
14.1.3	When finishing, which places should I have a look at?	177
14.2	454 data	178
14.2.1	What do I need SFFs for?	178
14.2.2	What's sff_extract and where do I get it?	178
14.2.3	Do I need the sfftools from the Roche software package?	178
14.2.4	Combining SFFs	178
14.2.5	Adaptors and paired-end linker sequences	178
14.2.6	What do I get in paired-end sequencing?	179
14.2.7	Sequencing protocol	180
14.2.8	Filtering sequences by length and re-assembly	180
14.3	Solexa / Illumina data	180
14.3.1	Can I see deletions?	180
14.3.2	Can I see insertions?	181
14.3.3	De-novo assembly with Solexa data	181
14.4	Hybrid assemblies	181
14.4.1	What are hybrid assemblies?	181
14.4.2	What differences are there in hybrid assembly strategies?	181
14.5	Masking	182
14.5.1	Should I mask?	182
14.5.2	How can I apply custom masking?	182
14.6	Miscellaneous	183
14.6.1	What are megahubs?	183
14.6.2	Passes and loops	184
14.6.3	Debris	184
14.6.4	Log and temporary files: more info on what happened during the assembly	184
14.6.4.1	Sequence clipping after load	184
14.7	Platforms and Compiling	185
14.7.1	Windows	185

15 The MAF format	186
15.1 Introduction: why an own assembly format?	186
15.2 The MAF format	187
15.2.1 Basics	187
15.2.2 Reads	187
15.2.2.1 Simple example	187
15.2.2.2 List of records for reads	188
15.2.2.3 Interpreting clipping values	189
15.2.3 Contigs	189
15.2.3.1 Simple example 2	190
15.2.3.2 List of records for contigs	190
16 Log and temporary files used by MIRA	192
16.1 Introduction	192
16.2 The files	192
16.2.1 mira_error_reads_invalid	192
16.2.2 mira_info_reads_tooshort	193
16.2.3 mira_int_alignnextends_preassembly1.0.txt	193
16.2.4 mira_int_clippings.0.txt	193
16.2.5 mira_int_posmatch_megahubs_pass.X.lst	193
16.2.6 mira_int_posmatch_multicopystat_preassembly.0.txt	193
16.2.7 mira_int_posmatch_rawhashhits_pass.X.lst	193
16.2.8 mira_int_skimmarknastyrepeats_hist_pass.X.lst	193
16.2.9 mira_int_skimmarknastyrepeats_nastyseq_pass.X.lst	193
16.2.10 mira_int_vectorclip_pass.X.txt	194
16.2.11 miratmp.ads_pass.X.forward and miratmp.ads_pass.X.complement	194
16.2.12 miratmp.ads_pass.X.reject	194
16.2.13 miratmp.noqualities	194
16.2.14 miratmp.usedids	194
16.2.15 mira_readpoolinfo.lst	194

List of Figures

1.1	How MIRA learns from misassemblies (1)	5
1.2	How MIRA learns from misassemblies (2)	6
1.3	How MIRA learns from misassemblies (3)	7
1.4	Slides presenting the repeat locator at the GCB 99	7
1.5	Slides presenting the Edit automatic Sanger editor at the GCB 99	8
1.6	Sanger assembly without EdIt automatic editing routines	8
1.7	Sanger assembly with EdIt automatic editing routines	9
1.8	454 assembly without 454 automatic editing routines	10
1.9	454 assembly with 454 automatic editing routines	11
1.10	Coverage of a contig.	11
1.11	Repetitive end of a contig	12
1.12	Non-repetitive end of a contig	13
1.13	MIRA pointing out problems in hybrid assemblies (1)	14
1.14	Coverage equivalent reads (CERs) explained.	15
1.15	Coverage equivalent reads let SNPs become very visible in assembly viewers	16
1.16	SNP tags in a MIRA assembly	17
1.17	Tag pointing out a large deletion in a MIRA mapping assembly	18
7.1	Format conversions with miraconvert	59
7.2	Conversions needed for other tools.	60
7.3	Join at a repetitive site which should not be performed due to missing spanning templates.	64
7.4	Join at a repetitive site which should be performed due to spanning templates being good.	65
7.5	Pseudo-repeat in 454 data due to sequencing artifacts	66
7.6	"SROc" tag showing a SNP position in a Solexa mapping assembly.	67
7.7	"SROc" tag showing a SNP/indel position in a Solexa mapping assembly.	68
7.8	"MCVc" tag (dark red stretch in figure) showing a genome deletion in Solexa mapping assembly.	69
7.9	An IS150 insertion hiding behind a WRMc and a SRMc tags	70
7.10	A 16 base pair deletion leading to a SROc/UNsC xmas-tree	71
7.11	An IS186 insertion leading to a SROc/UNsC xmas-tree	72
11.1	The Solexa GGCxG problem.	153

11.2 The Solexa GGC problem, forward example	154
11.3 The Solexa GGC problem, reverse example	155
11.4 A genuine place of interest almost masked by the GGCxG problem.	156
11.5 Example for no GC coverage bias in 2008 Solexa data.	158
11.6 Example for GC coverage bias starting Q3 2009 in Solexa data.	158
11.7 Example for GC coverage bias, direct comparison 2008 / 2010 data.	159
11.8 Example for good IonTorrent data (100bp reads)	160
11.9 Example for problematic IonTorrent data (100bp reads)	160
11.10 Example for a sequencing direction dependent indel	161

Preface

“How much intelligence does one need to sneak upon lettuce? ”

—Solomon Short

Note

A quick note at the start: I am scaling back the scope I recommend MIRA for. For genome de-novo assemblies or mapping projects, haploid organisms up to 20 to 40 megabases should be the limit.

Do not use MIRA if you have PacBio or Oxford Nanopore reads. Then again, for polishing those assemblies with Illumina data, MIRA is really good.

For mapping projects, do not use MIRA if you expect splicing like, e.g., RNASeq against an eukaryotic genome.

Lastly, Illumina projects with more than 40 to 60 million reads start to be so resource intensive that you might be better served with other assemblers or mapping programs. I know some people use MIRA for de-novo RNASeq with 300 million reads because they think it's worth it, but they wait more than a month then for the calculation to finish.

This "book" is actually the result of an exercise in self-defense. It contains texts from several years of help files, mails, postings, questions, answers etc.pp concerning MIRA and assembly projects one can do with it.

I never really intended to push MIRA. It started out as a PhD thesis and I subsequently continued development when I needed something to be done which other programs couldn't do at the time. But MIRA has always been available as binary on the Internet since 1999 ... and as Open Source since 2007. Somehow, MIRA seems to have caught the attention of more than just a few specialised sequencing labs and over the years I've seen an ever growing number of mails in my inbox and on the MIRA mailing list. Both from people having been "since ever" in the sequencing business as well as from labs or people just getting their feet wet in the area.

The help files -- and through them this book -- sort of reflect this development. Most of the chapters¹ contain both very specialised topics as well as step-by-step walk-throughs intended to help people to get their assembly projects going. Some parts of the documentation are written in a decidedly non-scientific way. Please excuse, time for rewriting mails somewhat lacking, some texts were re-used almost verbatim.

The last few years have seen tremendous change in the sequencing technologies and MIRA 5 reflects that: core data structures and routines had to be thrown overboard and replaced with faster and/or more versatile versions suited for the broad range of technologies and use-cases I am currently running MIRA with.

While other programs are nowadays better suited for many assembly tasks -- think SPADES for larger Illumina data sets or Falcon / Canu for PacBio -- I still regularly come back to MIRA when accuracy is important or catch the above mentioned programs doing something silly.

Nothing is perfect, and both MIRA and this documentation (even if it is rather pompously called *Definitive Guide*) are far from it. If you spot an error either in MIRA or this manual, feel free to report it. Or, even better, correct it if you can. At least with the manual files it should be easy: they're basically just some decorated text files.

I hope that MIRA will be as useful to you as it has been to me. Have a lot of fun with it.

Berlin, Feb 2019

Bastien Chevreux

¹ Avid readers of David Gerrold will certainly recognise the quotes from his books at the beginning of each chapter

Chapter 1

Introduction to MIRA

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Half of being smart is to know what you’re dumb at.”

—Solomon Short

1.1 What is MIRA?

MIRA is a multi-pass DNA sequence data assembler/mapper for whole genome and EST/RNASEq projects. MIRA assembles/maps reads gained by

- electrophoresis sequencing (aka Sanger sequencing)
- Illumina (Solexa) sequencing
- Less so: 454 pyro-sequencing (GS20, FLX or Titanium)
- Less so: Ion Torrent

into contiguous sequences (called *contigs*). One can use the sequences of different sequencing technologies either in a single assembly run (a *true hybrid assembly*) or by mapping one type of data to an assembly of other sequencing type (a *semi-hybrid assembly (or mapping)*) or by mapping a data against consensus sequences of other assemblies (a *simple mapping*).

The MIRA acronym stands for **M**imicking **I**ntelligent **R**ead **A**ssembly and the program pretty well does what its acronym says (well, most of the time anyway). It is the Swiss army knife of sequence assembly that I’ve used and developed during the past 20 years to get assembly jobs I work on done efficiently - and especially accurately. That is, without me actually putting too much manual work into it.

Over time, other labs and sequencing providers have found MIRA useful for assembly of extremely ‘unfriendly’ projects containing lots of repetitive sequences. As always, your mileage may vary.

1.2 What to read in this manual and where to start reading?

At the last count, this manual had almost 200 pages and this might seem a little bit daunting. However, you very probably do not need to read everything.

You should read most of this introductory chapter though: e.g.,

- the part with the MIRA quick tour
-

- the part which gives a quick overview for which data sets to use MIRA and for which not
- the part which showcases different features of MIRA (lots of screen shots!)
- where and how to get help if things don't work out as you expected

After that, reading should depend on the type of data you intend to work with: there are specific chapters for assembly of de-novo, of mapping and of EST / RNASeq projects. They all contain an overview on how to define your data and how to launch MIRA for these data sets. There is also chapter on how to prepare data sets from specific sequencing technologies.

The chapter on working with results of MIRA should again be of general interest to everyone. It describes the structure of output directories and files and gives first pointers on what to find where. Also, converting results into different formats -- with and without filtering for specific needs -- is covered there.

As the previously cited chapters are more introductory in their nature, they do not go into the details of MIRA parametrisation. While MIRA has a comprehensive set of standard settings which should be suited for a majority of assembly tasks, there are more than 150 switches / parameters with which one can fine tune almost every aspect of an assembly. A complete description for each and every parameter and how to correctly set parameters for different use cases and sequencing technologies can be found in the reference chapter.

As not every assembly project is simple, there is also a chapter with tips on how to deal with projects which turn out to be "hard." It certainly helps if you at least skim through it even if you do not expect to have problems with your data ... it contains a couple of tricks on what one can see in result files as well as in temporary and log files which are not explained elsewhere.

MIRA comes with a number of additional utilities which are described in an own chapter. While the purpose of **miraconvert** should be quite clear quite quickly, the versatility of use cases for **mirabait** might surprise more than one. Be sure to check it out.

As from time to time some general questions on sequencing are popping up on the MIRA talk mailing list, I have added a chapter with some general musings on what to consider when going into sequencing projects. This should be in no way a replacement for an exhaustive talk with a sequencing provider, but it can give a couple of hints on what to take care of.

There is also a FAQ chapter with some of the more frequently asked questions which popped up in the past few years.

Finally, there are also chapters covering some more technical aspects of MIRA: the MAF format and structure / content of the tmp directory have own chapters.

Complete walkthroughs ... are lacking at the moment for MIRA 4. In the MIRA 3 manual I had them, but so many things have changed (at all levels: MIRA, the sequencing technologies, data repositories) that I did not have time to update them. I probably will need quite some time to write new ones. Feel free to send me some if you are inclined to help fellow scientists.

1.3 The MIRA quick tour

Input can be in various formats like Staden experiment (EXP), Sanger CAF, FASTA, FASTQ or PHD file. Ancillary data containing additional information helpful to the assembly as is contained in, e.g. NCBI traceinfo XML files or Staden EXP files, is also honoured. If present, base qualities in **phred** style and SCF signal electrophoresis trace files are used to adjudicate between or even correct contradictory stretches of bases in reads by either the integrated automatic EdIt editor (written by Thomas Pfisterer) or the assembler itself.

MIRA was conceived especially with the problem of repeats in genomic data and SNPs in transcript (EST / RNASeq) data in mind. Considerable effort was made to develop a number of strategies -- ranging from standard clone-pair size restrictions to discovery and marking of base positions discriminating the different repeats / SNPs -- to ensure that repetitive elements are correctly resolved and that misassemblies do not occur.

The resulting assembly can be written in different standard formats like CAF, Staden GAP4 directed assembly, ACE, HTML, FASTA, simple text or transposed contig summary (TCS) files. These can easily be imported into numerous finishing tools or further evaluated with simple scripts.

The aim of MIRA is to build the best possible assembly by

1. having a more or less full overview on the whole project at any time of the assembly, i.e. knowledge of almost all possible read-pairs in a project,

2. using high confidence regions (HCRs) of several aligned read-pairs to start contig building at a good anchor point of a contig, extending clipped regions of reads on a 'can be justified' basis.
3. using all available data present at the time of assembly, i.e., instead of relying on sequence and base confidence values only, the assembler will profit from trace files containing electrophoresis signals, tags marking possible special attributes of DNA, information on specific insert sizes of read-pairs etc.
4. having 'intelligent' contig objects accept or refuse reads based on the rate of unexplainable errors introduced into the consensus
5. learning from mistakes by discovering and analysing possible repeats differentiated only by single nucleotide polymorphisms. The important bases for discriminating different repetitive elements are tagged and used as new information.
6. using the possibility given by the integrated automatic editor to correct errors present in contigs (and subsequently) reads by generating and verifying complex error hypotheses through analysis of trace signals in several reads covering the same area of a consensus,
7. iteratively extending reads (and subsequently) contigs based on
 - (a) additional information gained by overlapping read pairs in contigs and
 - (b) corrections made by the automated editor.

MIRA was part of a bigger project that started at the DKFZ (Deutsches Krebsforschungszentrum, German Cancer Research Centre) Heidelberg in 1997: the "Bundesministerium für Bildung, Wissenschaft, Forschung und Technologie" supported the PhD thesis of Thomas and myself by grant number *01 KW 9611*. Beside an assembler to tackle difficult repeats, the grant also supported the automated editor / finisher EdIt package -- written by Thomas Pfisterer. The strength of MIRA and EdIt is the automatic interaction of both packages which produces assemblies with less work for human finishers to be done.

I'd like to thank everybody who reported bugs to me, pointed out problems, sent ideas and suggestions they encountered while using the predecessors. Please continue to do so, the feedback made this third version possible.

1.4 For which data sets to use MIRA and for which not

As a general rule of thumb: if you have an organism with more than 100 to 150 megabases or more than 20 to 40 million reads, you might want to try other assemblers first.

1.4.1 Genome de-novo

For genome assembly, the version 5 series of MIRA have been reported to work on projects with something like a million Sanger reads (~80 to 100 megabases at 10x coverage), five to ten million 454 Titanium reads (~100 megabases at 20x coverage) and 20 to 40 million Illumina reads (enough for de-novo of a bacterium or a small eukaryote with 76mers to 300mers).

Provided you have the memory, MIRA is expected to work in de-novo mode with

- Sanger reads: 5 to 10 million
- 454 reads: 5 to 15 million
- Ion Torrent reads: 5 to 15 million
- Illumina reads: in normal operation, up to 40 million reads. Some people use it on up to 300 million, but you'll need a really big machine and month of computation time ... I do not recommend that.

and "normal" coverages, whereas "normal" would be at no more than 50x to 70x for genome projects. Higher coverages will also work, but may create somewhat larger temporary files without heavy parametrisation. Lower coverages (<4x for Sanger, <10x for 454, < 10x for IonTorrent) also need special attention in the parameter settings.

1.4.2 Genome mapping

As the complexity of mapping is a lot lower than de-novo, one can basically double (perhaps even triple) the number of reads compared to 'de-novo'. The limiting factor will be the amount of RAM though, and MIRA will also need lots of it if you go into eukaryotes.

The main limiting factor regarding time will be the number of reference sequences (backbones) you are using. MIRA being pedantic during the mapping process, it might be a rather long wait if you have more than 40 megabase of reference sequences.

1.4.3 De-novo ESTs / RNASeq

The default values for MIRA should allow it to work with many EST and RNASeq data sets, sometimes even from non-normalised libraries. For extreme coverage cases however (like, something with a lot of cases at and above 10k coverage), one would perhaps want to resort to data reduction routines before feeding the sequences to MIRA.

On the other hand, recent developments of MIRA were targeted at making de-novo RNASeq assembly of non-normalised libraries liveable, and indeed I now regularly use MIRA for data sets with up to 50 million Illumina 250bp paired-end reads.

1.4.4 Mapping EST / RNASeq

Mapping 50 to 60 million sequences against 20k to 30k gene reference sequences is doable, but slow and memory intensive.

Note For mapping against an eukaryotic genome: MIRA does *not* splice reads in mapping mode. While it is able to more or less handle insertion / deletions of up to 15% of a read length, splices larger than that will not be made: MIRA will still map the reads, but call a lot of SNPs in the parts which should have been spliced.

1.5 Any special features I might be interested in?

A few perhaps.

Note

The screen shots in this section show data from assemblies produced with MIRA, but the visualisation itself is done in a finishing program named **gap4**.

Some of the screen shots were edited for showing a special feature of MIRA. E.g., in the screen shots with Illumina data, quite some reads were left out of the view pane as else -- due to the amount of data -- these screen shots would need several pages for a complete printout.

1.5.1 MIRA learns to discern non-perfect repeats, leading to better assemblies

MIRA is an iterative assembler (it works in several passes) and acts a bit like a child when exploring the world: it explores the assembly space and is specifically parameterised to allow a couple of assembly errors during the first passes. But after each pass some routines (the "parents", if you like) check the result, searching for assembly errors and deduce knowledge about specific assemblies MIRA should not have ventured into. MIRA will then prevent these errors to re-occur in subsequent passes.

As an example, consider the following multiple alignment:

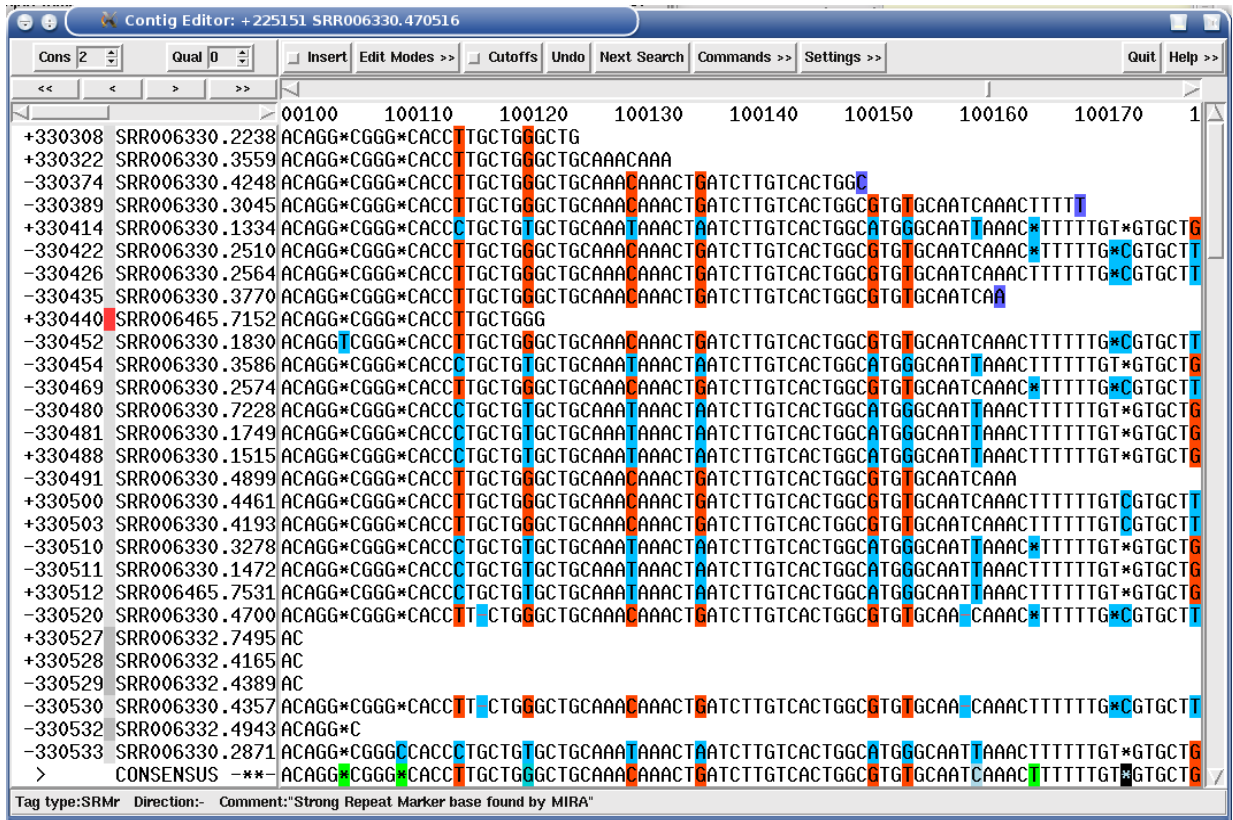


Figure 1.1: How MIRA learns from misassemblies (1). Multiple alignment after 1st pass with an obvious assembly error, notice the clustered columns discrepancies. Two slightly different repeats were assembled together.

These kind of errors can be easily spotted by a human, but are hard to prevent by normal alignment algorithms as sometimes there's only one single base column difference between repeats (and not several as in this example).

MIRA spots these things (even if it's only a single column), tags the base positions in the reads with additional information and then will use that information in subsequent passes. The net effect is shown in the next two figures:

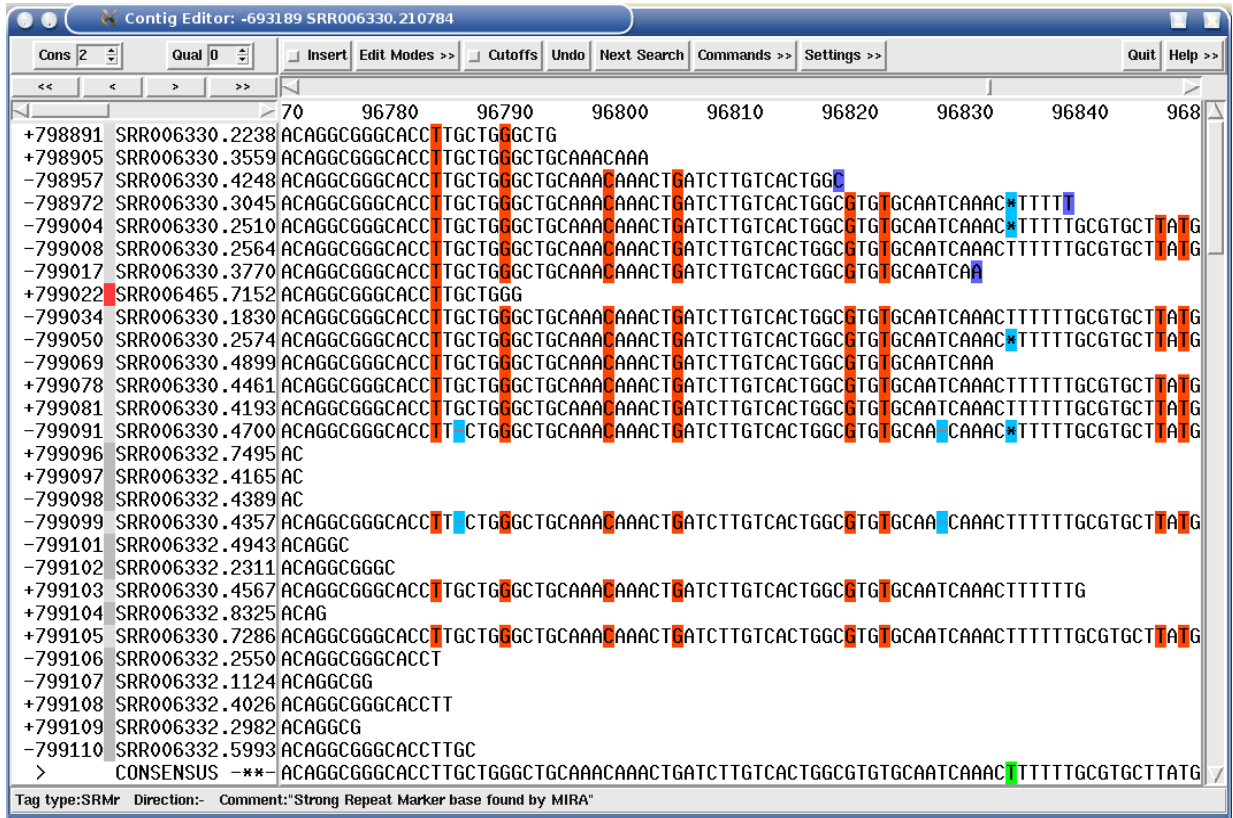


Figure 1.2: Multiple alignment after last pass where assembly errors from previous passes have been resolved (1st repeat site)

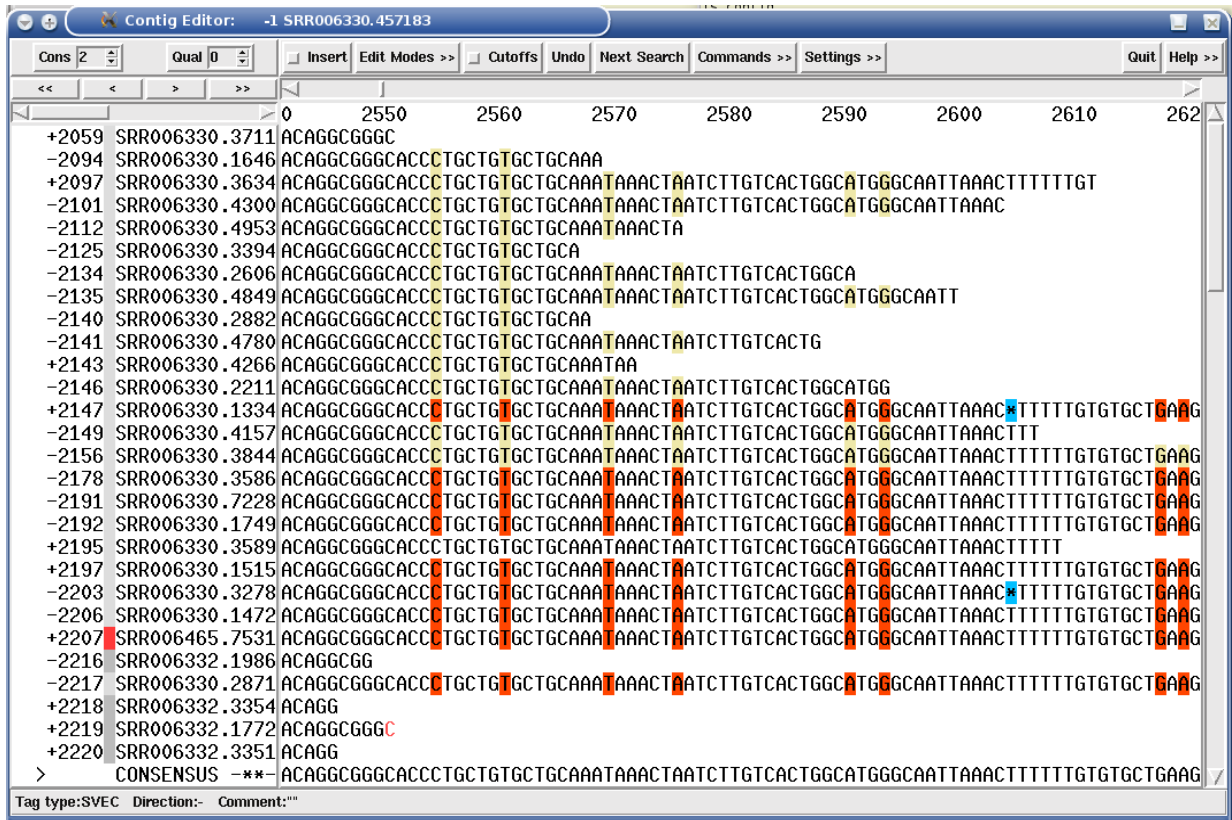


Figure 1.3: Multiple alignment after last pass where assembly errors from previous passes have been resolved (2nd repeat site)

The ability of MIRA to learn and discern non-identical repeats from each other through column discrepancies is nothing new. Here’s the link to a paper from a talk I had at the German Conference on Bioinformatics in 1999: <http://www.bioinfo.de/isb/gcb99/talks/chevreux/>

I’m sure you’ll recognise the basic principle in figures 8 and 9. The slides from the corresponding talk also look very similar to the screen shots above:

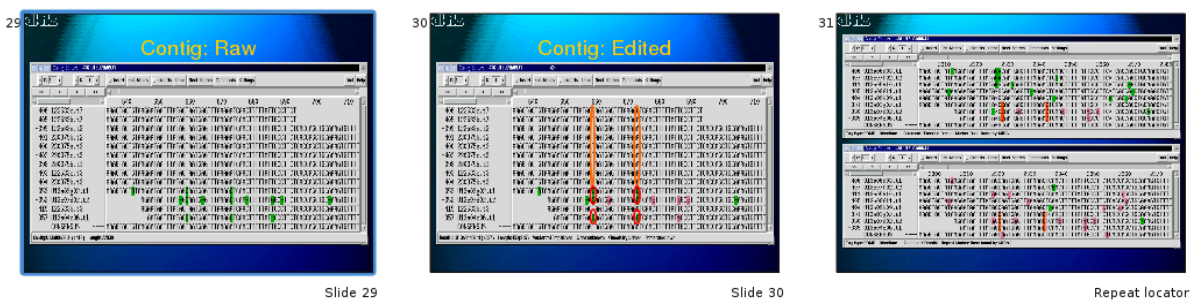
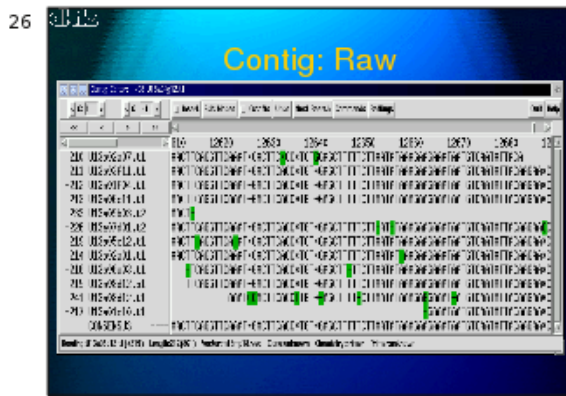


Figure 1.4: Slides presenting the repeat locator at the GCB 99

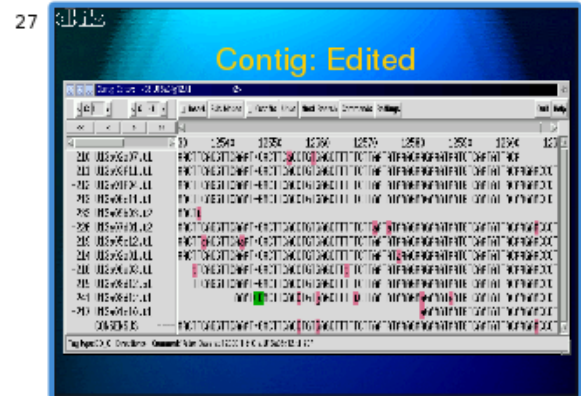
You can get the talk with these slides here: http://chevreux.org/dkfold/gcb99/bachvortrag_gcb99.ppt

1.5.2 MIRA has integrated editors for data from Sanger, 454, IonTorrent sequencing

In the first versions in 1999, the *EdIt* automatic Sanger sequence editor from Thomas Pfisterer was integrated into MIRA.



Contig: Raw



Contig: Edited

Figure 1.5: Slides presenting the Edit automatic Sanger editor at the GCB 99

The routines used a combination of hypothesis generation/testing together with neural networks (trained on ABI and ALF traces) for signal recognition to discern between base calling errors and true multiple alignment differences. They went back to the trace data to resolve potential conflicts and eventually recall bases using the additional information gained in a multiple alignment of reads.



Figure 1.6: Sanger assembly without EdIt automatic editing routines. The bases with blue background are base calling errors.



Figure 1.7: Sanger assembly with EdIt automatic editing routines. Bases with pink background are corrections made by EdIt after assessing the underlying trace files (SCF files in this case). Bases with blue background are base calling errors where the evidence in the trace files did not show enough evidence to allow an editing correction.

With the introduction of 454 and Illumina reads, MIRA also got in 2007 specialised editors to search and correct for typical 454 and Illumina sequencing problems, like the homopolymer run over-/undercalls or the Illumina GGCxG motif problem. These editors are now integrated into MIRA itself and are not part of EdIt anymore.

While not being paramount to the assembly quality, both editors provide additional layers of safety for the MIRA learning algorithm to discern non-perfect repeats even on a single base discrepancy. Furthermore, the multiple alignments generated by these two editors are way more pleasant to look at (or automatically analyse) than the ones containing all kind of gaps, insertions, deletions etc.pp.

```

Contig Editor: +128973 SRR006330.349400
Cons 2  Qual 0  Insert  Edit Modes >>  Cutoffs  Undo  Next Search  Commands >>  Settings >>  Quit  Help >>
117890  117900  117910  117920  117930  117940  117950  117960
+111432 SRR006330.1168 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111442 SRR006330.1122 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111435 SRR006330.4625 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111449 SRR006330.3062 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111430 SRR006330.3381 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111451 SRR006330.2563 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG
-111444 SRR006330.3341 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG
-111440 SRR006330.2625 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111446 SRR006330.1114 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCC
-111439 SRR006330.2904 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111438 SRR006330.2546 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111437 SRR006330.3496 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111431 SRR006330.3417 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111428 SRR006330.3334 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111420 SRR006330.3989 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
+111433 SRR006330.4461 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
+111445 SRR006465.5363 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCAT
-111423 SRR006330.5115 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
+111414 SRR006330.4208 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111419 SRR006330.2371 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111452 SRR006465.3088 GGTGCC*ATTCAACGTCAG
-111425 SRR006330.1341 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111427 SRR006330.2000 GGTGCC*ATTCAACGTCAGCCTTCCCTTACCTAACTCATCTTGGACTATCCATCAGGTAGGGTTTTGGT*CAGTGACGT
-111422 SRR006330.1736 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
+111429 SRR006330.3732 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111415 SRR006330.2221 GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCA**AGG*TTTTGGT*CAGTGACGT
-111426 SRR006330.5007 CTTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
-111403 SRR006330.3026 CTTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
<> CONSENSUS -*- GGTGCC*ATTCAACGTCAGC**TTCC*TT*CCTAACTCATCTTGGACTATCCATCAG**AGG*TTTTGGT*CAGTGACGT
SRR006330.500766(#111426) Clone:unknown Vector:unknown Type:forward universal;unknown primer Tmpl:E

```

Figure 1.8: 454 assembly without 454 automatic editing routines

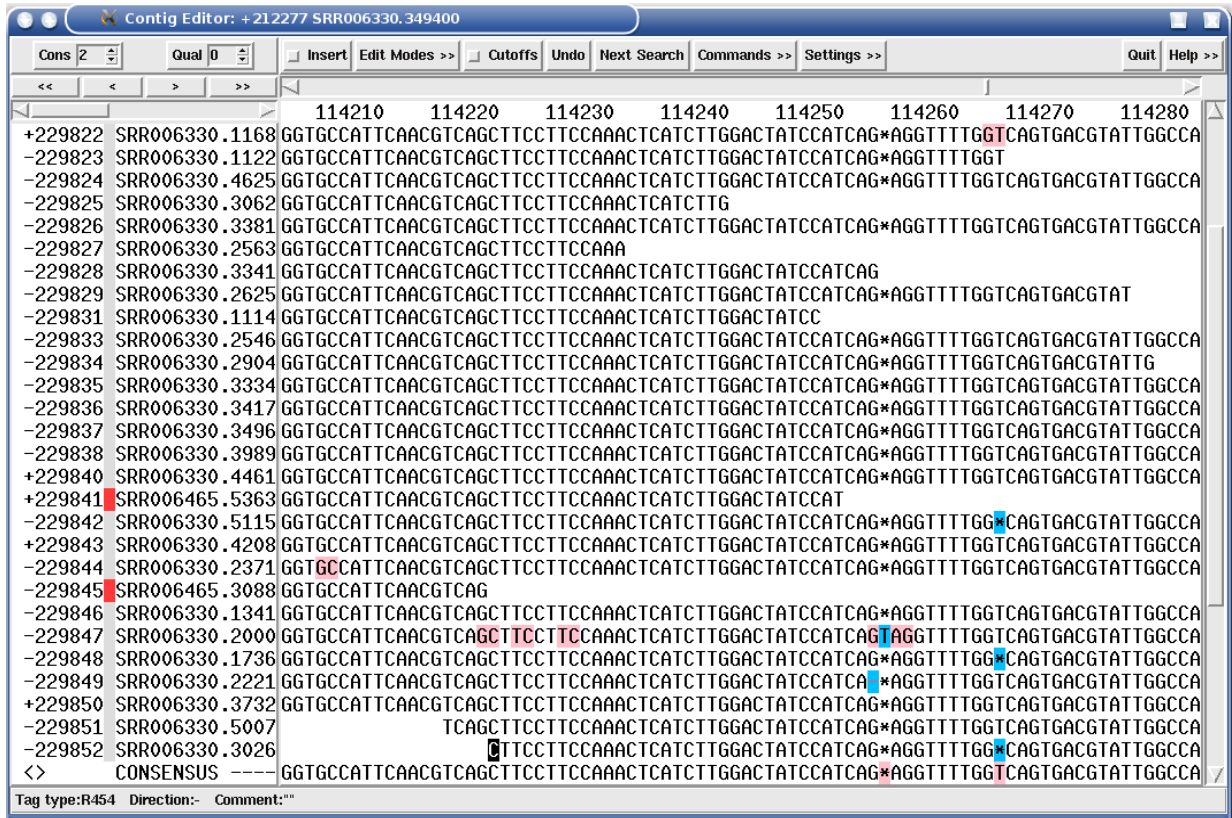


Figure 1.9: 454 assembly with 454 automatic editing routines

1.5.3 MIRA lets you see why contigs end where they end

A very useful feature for finishing are kmer (hash) frequency tags which MIRA sets in the assembly. Provided your finishing editor understands those tags (**gap4**, **gap5** and **consed** are fine but there may be others), they'll give you precious insight where you might want to be cautious when joining to contigs or where you would need to perform some primer walking. MIRA colours the assembly with the hash frequency (HAF) tags to show repetitiveness.

You will need to read about the HAF tags in the reference manual, but in a nutshell: the HAF5, HAF6 and HAF7 tags tell you potentially have repetitive to very repetitive read areas in the genome, while HAF2 tags will tell you that these areas in the genome have not been covered as well as they should have been.

As an example, the following figure shows the coverage of a contig.

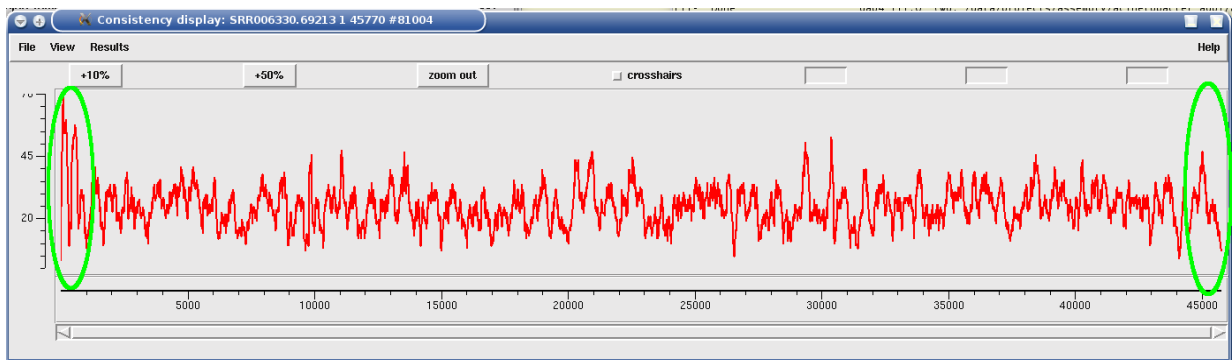


Figure 1.10: Coverage of a contig.

The question is now: why did MIRA stop building this contig on the left end (left oval) and why on the right end (right oval).

Looking at the HAF tags in the contig, the answer becomes quickly clear: the left contig end has HAF5 tags in the reads (shown in bright red in the following figure). This tells you that MIRA stopped because it probably could not unambiguously continue building this contig. Indeed, if you BLAST the sequence at the NCBI, you will find out that this is an rRNA area of a bacterium, of which bacteria normally have several copies in the genome:

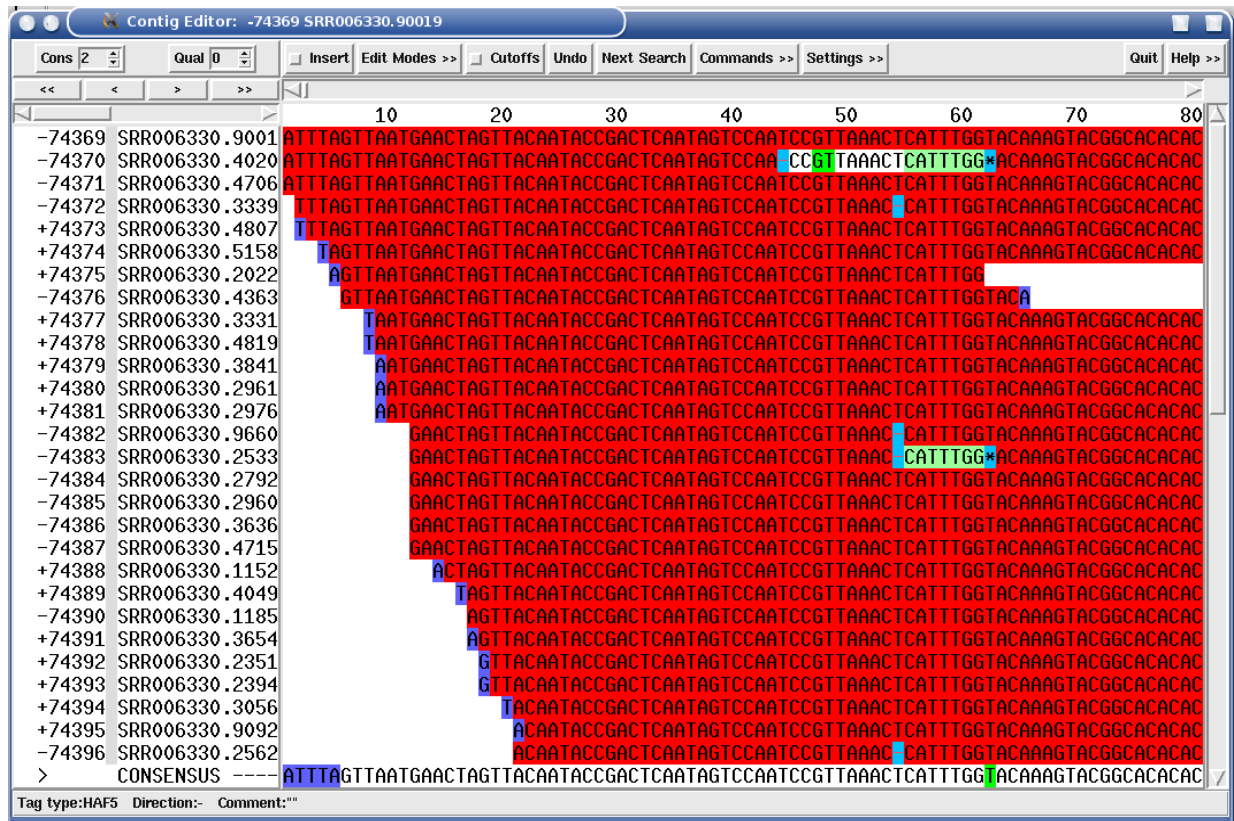


Figure 1.11: HAF5 tags (reads shown with red background) covering a contig end show repetitiveness as reason for stopping a contig build.

The right end of the same contig however ends in HAF3 tags (normal coverage, bright green in the next figure) and even HAF2 tags (below average coverage, pale green in the next image). This tells you MIRA stopped building the contig at this place simply because there were no more reads to continue. This is a perfect target for primer walking if you want to finish a genome.

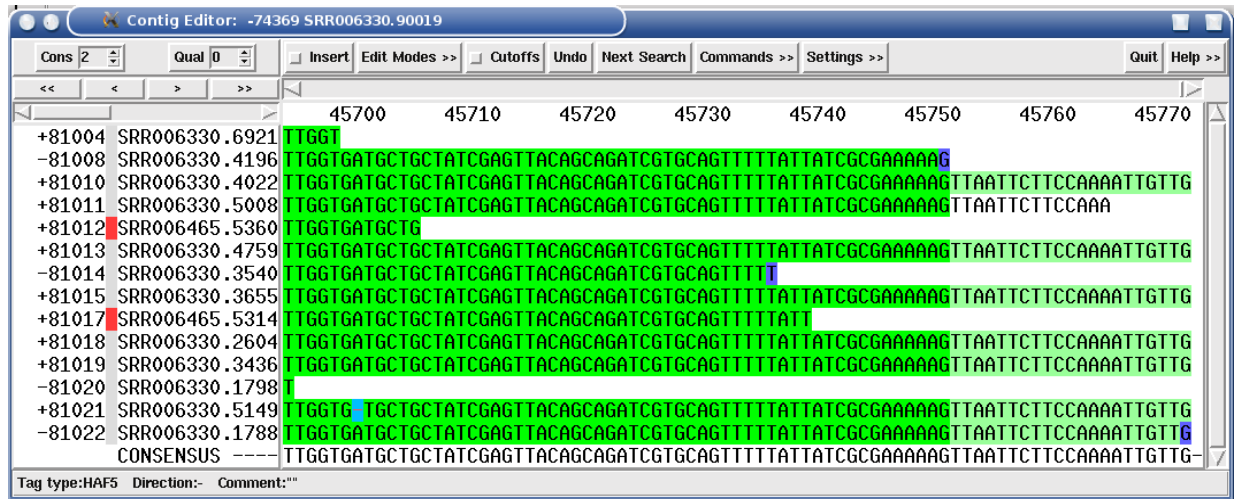


Figure 1.12: HAF2 tags covering a contig end show that no more reads were available for assembly at this position.

1.5.4 MIRA tags problematic decisions in hybrid assemblies

Many people combine Sanger & 454 or 454 & Illumina to improve the sequencing quality of their project through two (or more) sequencing technologies. To reduce time spent in finishing, MIRA automatically tags those bases in a consensus of a hybrid assembly where reads from different sequencing technologies severely contradict each other.

The following example shows a hybrid 454 / Illumina assembly where reads from 454 (highlighted read names in following figure) were not sure whether to have one or two "G" at a certain position. The consensus algorithm would have chosen "two Gs" for 454, obviously a wrong decision as all Illumina reads at the same spot (the reads which are not highlighted) show only one "G" for the given position. While MIRA chose to believe Illumina in this case, it tagged the position anyway in case someone chooses to check these kind of things.

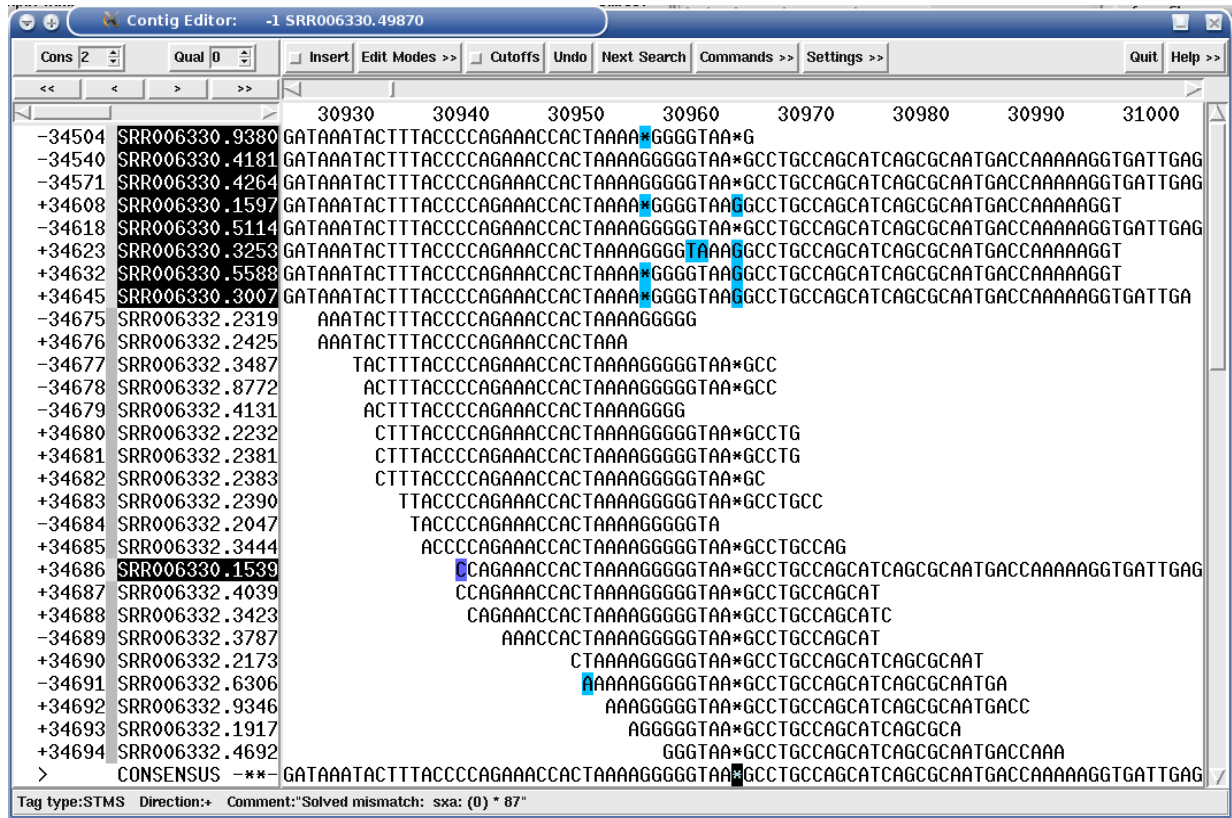


Figure 1.13: A "STMS" tag (Sequencing Technology Mismatch Solved, the black square base in the consensus) showing a potentially difficult decision in a hybrid 454 / Illumina de-novo assembly.

1.5.5 MIRA for polishing of PacBio or ONT assemblies

While MIRA is not really suited to assemble PacBio or Oxford Nanopore reads de-novo, it is very well suited to polish the results of such assemblies with Illumina reads in mapping projects.

TODO: example

1.5.6 MIRA allows older finishing programs to cope with amount data in Illumina mapping projects

Quality control is paramount when you do mutation analysis for biologists: I know they'll be on my doorstep the very next minute they found out one of the SNPs in the resequencing data wasn't a SNP, but a sequencing artefact. And I can understand them: why should they invest -- per SNP -- hours in the wet lab if I can invest a couple of minutes to get them data false negative rates (and false discovery rates) way below 1%? So, finishing and quality control for any mapping project is a must.

Both **gap4** and **consed** start to have a couple of problems when projects have millions of reads: you need lots of RAM and scrolling around the assembly gets a test to your patience. Still, these two assembly finishing programs are amongst the better ones out there, although **gap5** starts to quickly arrive in a state in which it allows itself to substitute to **gap4**.

So, MIRA reduces the number of reads in Illumina mapping projects without sacrificing information on coverage. The principle is pretty simple: for 100% matching reads, MIRA tracks coverage of every reference base and creates long synthetic, coverage equivalent reads (CERs) in exchange for the Illumina reads. Reads that do not match 100% are kept as own entities, so that no information gets lost. The following figure illustrates this:

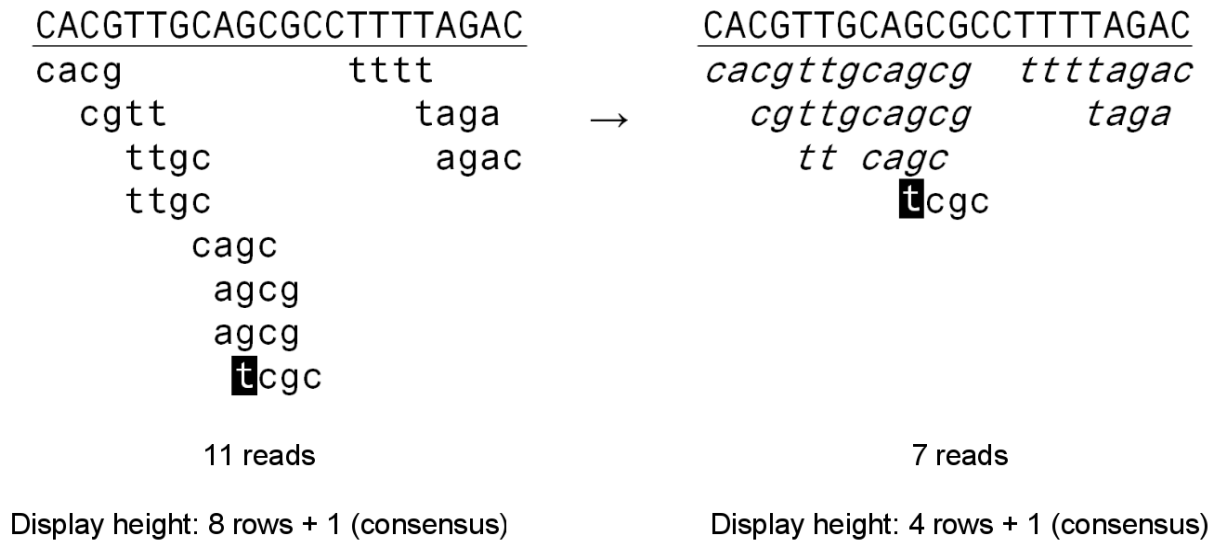


Figure 1.14: Coverage equivalent reads (CERs) explained. Left side of the figure: a conventional mapping with eleven reads of size 4 against a consensus (in uppercase). The inversed base in the lowest read depicts a sequencing error. Right side of the figure: the same situation, but with coverage equivalent reads (CERs). Note that there are less reads, but no information is lost: the coverage of each reference base is equivalent to the left side of the figure and reads with differences to the reference are still present.

This strategy is very effective in reducing the size of a project. As an example, in a mapping project with 9 million Illumina 36mers, MIRA created a project with 1.7m reads: 700k CER reads representing ~8 million 100% matching Illumina reads, and it kept ~950k mapped reads as they had ≥ mismatch (be it sequencing error or true SNP) to the reference. A reduction of 80%, and numbers for mapping projects with Illumina 100bp reads are in a similar range.

Also, mutations of the resequenced strain now really stand out in the assembly viewer as the following figure shows:

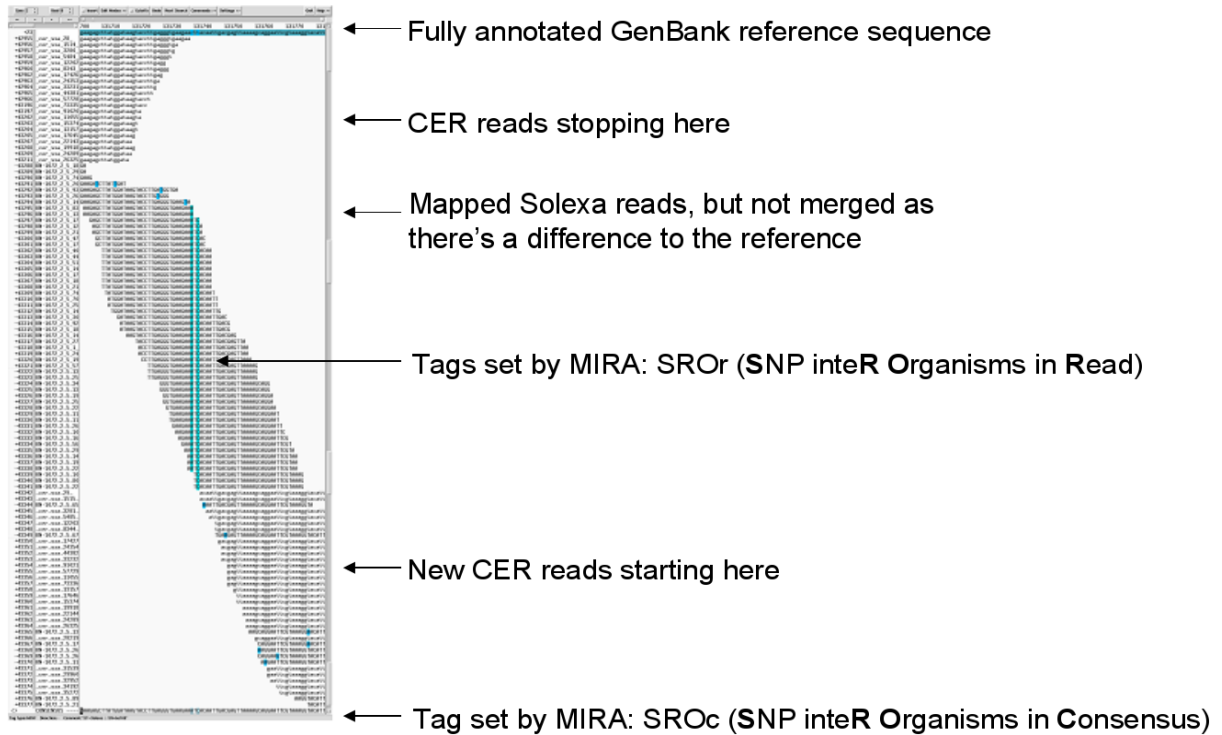


Figure 1.15: Coverage equivalent reads let SNPs become very visible in assembly viewers

1.5.7 MIRA tags SNPs and other features, outputs result files for biologists

Want to assemble two or several very closely related genomes without reference, but finding SNPs or differences between them? Tired of looking at some text output from mapping programs and guessing whether a SNP is really a SNP or just some random junk?

MIRA tags all SNPs (and other features like missing coverage etc.) it finds so that -- when using a finishing viewer like gap4 or consed -- one can quickly jump from tag to tag and perform quality control. This works both in de-novo assembly and in mapping assembly, all MIRA needs is the information which read comes from which strain.

The following figure shows a mapping assembly of Illumina 36mers against a bacterial reference sequence, where a mutant has an indel position in a gene:

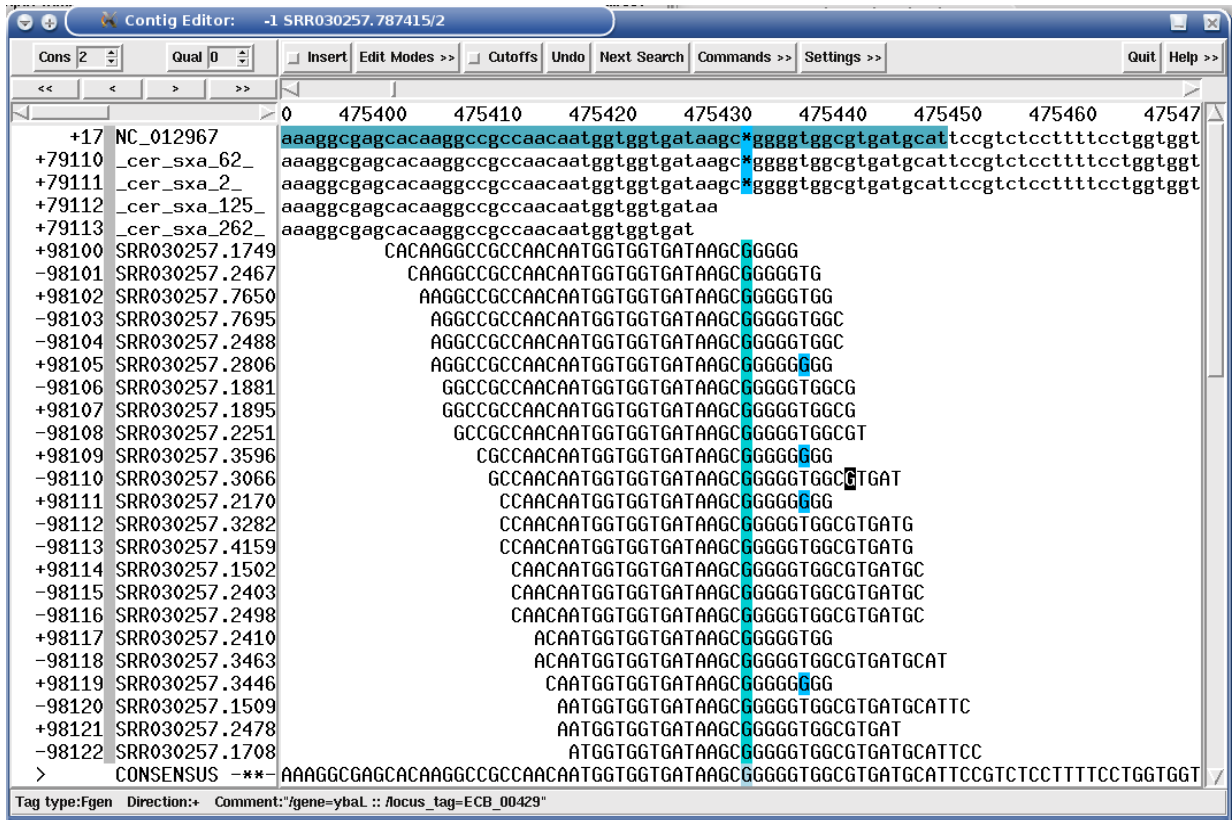


Figure 1.16: "SROc" tag (Snp inteR Organism on Consensus) showing a SNP position in a Illumina mapping assembly.

Other interesting places like deletions of whole genome parts are also directly tagged by MIRA and noted in diverse result files (and searchable in assembly viewers):

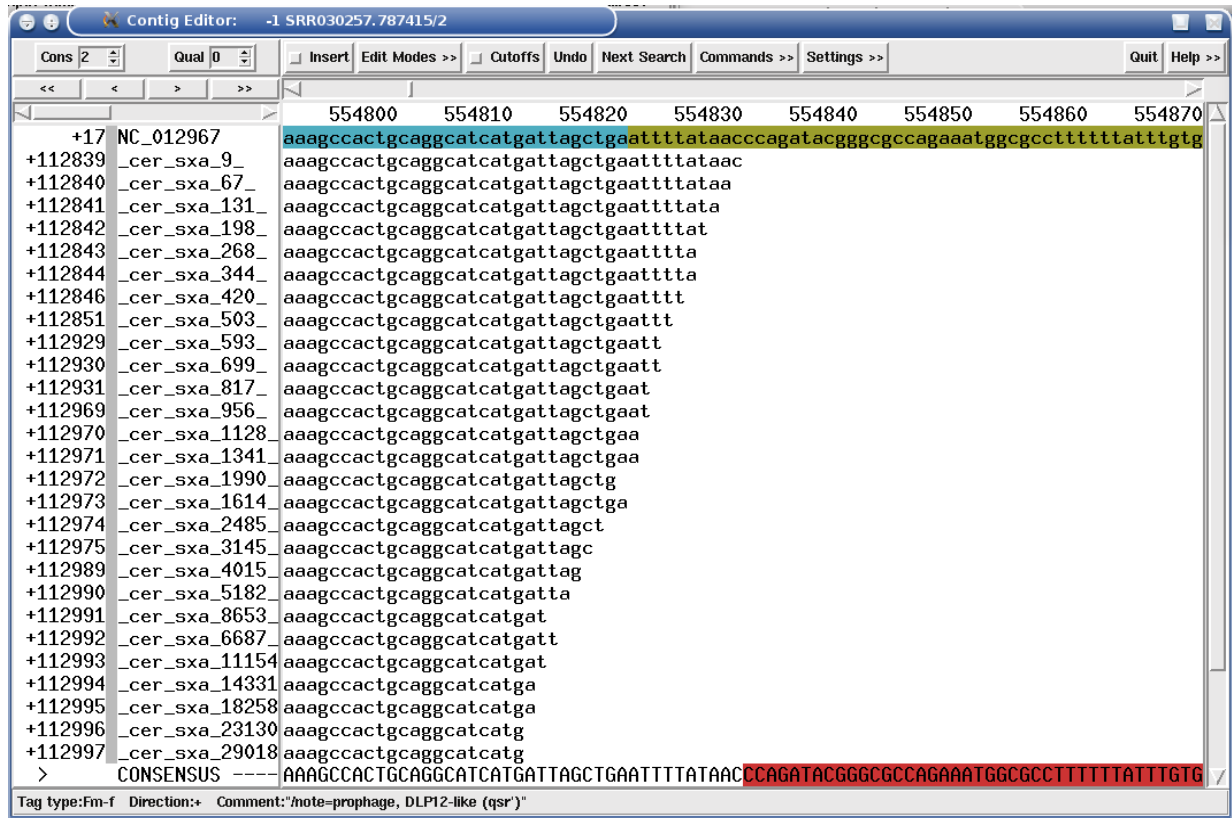


Figure 1.17: "MCVc" tag (Missing CoVerage in Consensus, dark red stretch in figure) showing a genome deletion in Illumina mapping assembly.

Note For bacteria -- and if you use annotated GenBank files as reference sequence -- MIRA will also output some nice lists directly usable (in Excel) by biologists, telling them which gene was affected by what kind of SNP, whether it changes the protein, the original and the mutated protein sequence etc.pp.

1.5.8 MIRA has ... much more

- Extensive possibilities to clip data if needed: by quality, by masked bases, by A/T stretches, by evidence from other reads, ...
- Routines to re-extend reads into clipped parts if multiple alignment allows for it.
- Read in ancillary data in different formats: EXP, NCBI TRACEINFO XML, SSAHA2, SMALT result files and text files.
- Detection of chimeric reads.
- Support for many different of input and output formats (FASTA, EXP, FASTQ, CAF, MAF, ...)
- Automatic memory management (when RAM is tight)
- Over 150 parameters to tune the assembly for a lot of use cases, many of these parameters being tunable individually depending on sequencing technology they apply to.

1.6 Versions, Licenses, Disclaimer and Copyright

1.6.1 Versions

There are two kind of versions for MIRA that can be compiled from source files: production and development.

Production versions are from the stable branch of the source code. These versions are available for download from SourceForge. Development versions are from the development branch of the source tree. These are also made available to the public and should be compiled by users who want to test out new functionality or to track down bugs or errors that might arise at a given location. Release candidates (rc) also fall into the development versions: they are usually the last versions of a given development branch before being folded back into the production branch.

1.6.2 License

1.6.2.1 MIRA

MIRA has been put under the GPL version 2.

This program is free software; you can redistribute it and/or modify it under the terms of the GNU General Public License as published by the Free Software Foundation; either version 2 of the License, or (at your option) any later version.

This program is distributed in the hope that it will be useful, but WITHOUT ANY WARRANTY; without even the implied warranty of MERCHANTABILITY or FITNESS FOR A PARTICULAR PURPOSE. See the GNU General Public License for more details.

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You may also visit <http://www.opensource.org/licenses/gpl-2.0.php> at the Open Source Initiative for a copy of this licence.

1.6.2.2 Documentation

The documentation pertaining to MIRA is licensed under the Creative Commons Attribution-NonCommercial-ShareAlike 3.0 Unported License. To view a copy of this license, visit <http://creativecommons.org/licenses/by-nc-sa/3.0/> or send a letter to Creative Commons, 171 Second Street, Suite 300, San Francisco, California, 94105, USA.

1.6.3 Copyright

© 1997-2000 Deutsches Krebsforschungszentrum Heidelberg -- Dept. of Molecular Biophysics and Bastien Chevreux (for MIRA) and Thomas Pfisterer (for EdIt)

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1.6.4 External libraries

MIRA uses the excellent Expat library to parse XML files. Expat is Copyright © 1998, 1999, 2000 Thai Open Source Software Center Ltd and Clark Cooper as well as Copyright © 2001, 2002 Expat maintainers.

See <http://www.libexpat.org/> and <http://sourceforge.net/projects/expat/> for more information on Expat.

1.7 Getting help / Mailing lists / Reporting bugs

Please try to find an answer to your question by first reading the documents provided with the MIRA package (FAQs, READMEs, usage guide, guides for specific sequencing technologies etc.). It's a lot, but then again, they hopefully should cover 90% of all questions.

If you have a tough nut to crack or simply could not find what you were searching for, you can subscribe to the MIRA talk mailing list and send in your question (or comment, or suggestion), see http://www.chevreux.org/mira_mailinglists.html for more information on that. Now that the number of subscribers has reached a good level, there's a fair chance that someone could answer your question before I have the opportunity or while I'm away from mail for a certain time.

Note

Please very seriously consider using the mailing list before mailing me directly. Every question which can be answered by participants of the list is time I can invest in development and documentation of MIRA. I have a day job as bioinformatician which has nothing to do with MIRA and after work hours are rare enough nowadays.

Furthermore, Google indexes the mailing list and every discussion / question asked on the mailing list helps future users as they show up in Google searches.

Only mail me directly (bach@chevreux.org) if you feel that there's some information you absolutely do not want to share publicly.

Note Subscribing to the list *before sending mails to it* is necessary as messages from non-subscribers will be stopped by the system to keep the spam level low.

To report bugs or ask for new features, please use the GitHub issue tool at: <http://github.com/bachev/mira/issues>. This ensures that requests do not get lost **and** you get the additional benefit to automatically know when a bug has been fixed as I will not send separate emails, that's what bug trackers are there for.

Finally, new or intermediate versions of MIRA will be announced on the separate MIRA announce mailing list. Traffic is very low there as the only one who can post there is me. Subscribe if you want to be informed automatically on new releases of MIRA.

1.8 Author

Bastien Chevreux (mira): bach@chevreux.org

WWW: <http://www.chevreux.org/>

MIRA can use automatic editing routines for Sanger sequences which were written by Thomas Pfisterer (EdIt): t.pfisterer@dkfz-heidelberg.de

1.9 Miscellaneous

1.9.1 Citing MIRA

Please use these citations:

For mira Chevreux, B., Wetter, T. and Suhai, S. (1999): *Genome Sequence Assembly Using Trace Signals and Additional Sequence Information*. Computer Science and Biology: Proceedings of the German Conference on Bioinformatics (GCB) 99, pp. 45-56.

Chevreux, B., Pfisterer, T., Drescher, B., Driesel, A. J., Müller, W. E., Wetter, T. and Suhai, S. (2004): *Using the miraEST Assembler for Reliable and Automated mRNA Transcript Assembly and SNP Detection in Sequenced ESTs*. Genome Research, 14(6)

1.9.2 Postcards, gold and jewellery

If you find this software useful, please send the author a postcard. If postcards are not available, a treasure chest full of Spanish doubloons, gold and jewellery will do nicely, thank you.

Chapter 2

Installing MIRA

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“A problem can be found to almost every solution.”

—Solomon Short

2.1 Where to fetch MIRA

2.1.1 Package naming and versioning scheme

Source packages are usually named

```
mira-miraversion.tar.bz2
```

Note

For *miraversion*, the stable versions of MIRA with the general public as audience usually have a version number in three parts, like 3.0.5, sometimes also followed by some postfix like in 3.2.0rc1 to denote release candidate 1 of the 3.2.0 version of MIRA. On very rare occasions, stable versions of MIRA can have four part like in, e.g., 3.4.0_1: these versions create identical binaries to their parent version (in this case: 3.4.0) and just contains fixes to the source build machinery or documentation.

The version string sometimes can have a different format: *sometext_somenumber+gsomehexnumber* like in, e.g., *ft_fastercontig_10+g4a27c91*. These versions of MIRA are snapshots from the development tree of MIRA and usually contain new functionality which may not be as well tested as the rest of MIRA, hence contains more checks and more debugging output to catch potential errors

Precompiled binary packages are named in the following way:

```
mira_miraversion_OS-and-binarytype.tar.bz2
```

Note

The *OS-and-binarytype* part defines for which operating system and which processor class the package is destined. E.g., *linux-gnu_x86_64_static* contains static binaries for Linux running a X86 64 bit processor (most Intel/AMD nowadays).

2.1.2 SourceForge: source package and precompiled binaries

SourceForge: <https://sourceforge.net/projects/mira-assembler/files/MIRA/>

There you will normally find two directories (stable and development), both with a couple of precompiled binaries -- usually for Linux and Mac OSX -- or the source package for compiling yourself.

Examples for packages at SourceForge:

- `mira_4.9.8_linux-gnu_x86_64_static.tar.bz2`
- `mira_4.9.8_OSX_snowleopard_x86_64_static.tar.bz2`
- `mira-4.9.8.tar.bz2`

2.1.3 GitHub: git repository, source, and binary packages

GitHub main page: <https://github.com/bachev/mira>

GitHub releases (go here for binary packages): <https://github.com/bachev/mira/releases>

GitHub contains the live source of MIRA as well as source packages and binary packages for selected versions. Two main branches are generally interesting to the public: "master" and "development".

To get a clone of the repository on on GitHub, do:

```
git clone https://github.com/bachev/mira.git
cd mira
./bootstrap.sh
```

Once bootstrap has run through (see below for necessary prerequisites), the system is ready for a `./configure` call.

2.2 Installing from a precompiled binary package

The distributable package follows the one-directory-which-contains-everything-which-is-needed philosophy, but after unpacking and moving the package to its final destination, you need to run a script which will create some data files.

1. Download the package, unpack it.
2. Move the directory somewhere to your disk. Either to one of the "standard" places like, e.g., `/opt/mira`, `/usr/local/mira` or somewhere in your home directory
3. Softlink the binaries which are in the 'bin' directory into a directory which is in your shell PATH. Then have the shell reload the location of PATH binaries (either `hash -r` for sh/bash or `rehash` for csh/tcsh).
Alternatively, add the `bin` directory of the MIRA package to your PATH variable.
4. Test whether the binaries are installed ok via `mirabait -v` which should return with the current version you downloaded and installed.
5. Now you need to run a script which will unpack and reformat some data needed by MIRA. That script is located in the `dbdata` directory of the package and should be called with the name of the *SLS* file present in the same directory like this:

```
arcadia:/path/to/mirapkg$ cd dbdata
arcadia:/path/to/mirapkg/dbdata$ ls -l
drwxr-xr-x 3 bach bach    4096 2016-03-18 14:31 mira-createsls
-rwxr-xr-x 1 bach bach    2547 2015-12-14 04:33 mira-install-sls-rrna.sh
-rw-r--r-- 1 bach bach     337 2016-01-01 14:50 README.txt
lrwxrwxrwx 1 bach bach 10421035 2016-03-18 14:28 rfam_rrna-21-12.sls.gz
arcadia:/path/to/mirapkg/dbdata$ ./mira-install-sls-rrna.sh rfam_rrna-21-12.sls.gz
```

This will take a minute or so. Then you're done for MIRA.

Additional scripts for special purposes are in the `scripts` directory. You might or might not want to have them in your \$PATH. Scripts and programs for MIRA from other authors are in the `3rdparty` directory. Here too, you may or may not want to have (some of them) in your \$PATH.

2.3 Integration with third party programs (gap4, consed)

MIRA sets tags in the assemblies that can be read and interpreted by the Staden **gap4** package or **consed**. These tags are extremely useful to efficiently find places of interest in an assembly (be it de-novo or mapping), but both **gap4** and **consed** need to be told about these tags.

Data files for a correct integration are delivered in the `support` directory of the distribution. Please consult the README in that directory for more information on how to integrate this information in either of these packages.

2.4 Compiling MIRA yourself

Note As a general rule of thumb, you do NOT want to compile MIRA yourself. Fetch a precompiled binary if possible.

2.4.1 Prerequisites

Compiling the 5.x series of MIRA needs a C++14 compatible tool chain, i.e., systems starting from 2013/2014 should be OK. The requisites for *compiling* MIRA are:

- gcc \geq 6.1, with libstdc++6. Or clang \geq 3.5.

You really want to use a simple installation package pre-configured for your system, but in case you want or have to install gcc or clang yourself, you will have a lot of fun.

- BOOST library \geq 1.48 (\geq 1.61 on OSX).

You really want to use a simple installation package pre-configured for your system, but in case you want or have to install BOOST yourself, please refer to <http://www.boost.org/> for more information on the BOOST library.



Warning Do NOT use a so called *staged* BOOST library, that will not work.

- zlib. Should your system not have zlib installed or available as simple installation package, please see <http://www.zlib.net/> for more information regarding zlib.
- GNU make. Should your system not have gmake installed or available as simple installation package, please see www.gnu.org/software/make/ for more information regarding GNU make.
- GNU flex \geq 2.6.0. Should your system not have flex installed or available as simple installation package, please see <http://flex.sourceforge.net/> for more information regarding flex.
- Expat library \geq 2.0.1. Should your system not have the Expat library and header files already installed or available as simple installation package, you will need to download and install a yourself. Please see <http://www.libexpat.org/> and <http://sourceforge.net/projects/expat/> for information on how to do this.
- xxd. A small utility from the **vim** package.

For *building the documentation*, additional prerequisites are:

- help2man for manual pages
 - xsltproc + docbook-xsl for HTML output
 - a LaTeX system and dblatex for PDF output
-

When *building from a GitHub checkout*, additional prerequisites are a couple of additional tools:

- current automake and autoconf
- current libtool

2.4.2 Compiling and installing

If you're lucky, after fetching the source (and eventually calling `./bootstrap.sh` if you worked from a GitHub checkout), all you need to do is basically:

```
arcadia:/path/to/mira-5.0.0$ ./configure
arcadia:/path/to/mira-5.0.0$ make
arcadia:/path/to/mira-5.0.0$ make install
```

This should install the following programs:

- **mira**
- **miraconvert**
- **mirabait**
- **miramer**
- **miramem**

Should the `./configure` step fail for some reason or another, you should get a message telling you at which step this happens and either install missing packages or tell **configure** where it should search the packages it did not find. See also next section or the system specific walkthroughs in this chapter.

2.4.3 Configure arguments for MIRA

MIRA understands all standard autoconf configure arguments, the most important one probably being `--prefix=` to tell where you want to have MIRA installed. Please consult `./configure --help` to get a full list of currently supported arguments.

2.4.3.1 Configure arguments for the BOOST library

BOOST is maybe the most tricky library to get right in case it does not come pre-configured for your system. The two main arguments for helping to locate BOOST are probably `--with-boost=[ARG]` and `--with-boost-libdir=LIB_DIR`. Only if those two fail, try using the other `--with-boost-*=` arguments you will see from the `./configure help` text.

2.4.3.2 MIRA specific configure arguments

The configure scripts honours the following MIRA specific arguments:

--with-compiler=gcc/clang Force the compilation to occur with the given compiler. The default is clang on OSX systems and gcc on all other.

Note Using homebrew installed GCC on OSX compiled, but the executable did not run (no idea why). Using clang on Linux is untested atm.

--with-brew Automatically configure the build to be made entirely with and from software (compiler, libraries etc.) installed via Homebrew. Works perfectly on OSX, untested on Linux.

- enable-warnings** Enables compiler warnings, useful only for developers, not for users.
- enable-debug** Lets the MIRA binary contain C/C++ debug symbols.
- enable-mirastatic** Builds static binaries which are easier to distribute. Some platforms (like OpenSolaris) might not like this and you will get an error from the linker.
- enable-optimisations** Instructs the configure script to set optimisation arguments for compiling (on by default). Switching optimisations off (warning, high impact on run-time) might be interesting only for, e.g, debugging with valgrind.
- enable-publicquietmira** Some parts of MIRA can dump additional debug information during assembly, setting this switch to "no" performs this. Warning: MIRA will be a bit chatty, using this is not recommended for public usage.
- enable-developmentversion** Using MIRA with enabled development mode may lead to extra output on stdout as well as some additional data in the results which should not appear in real world data
- enable-boundtracking**
- enable-bugtracking** Both flags above compile in some basic checks into mira that look for sanity within some functions: Leaving this on "yes" (default) is encouraged, impact on run time is minimal

2.5 Installation walkthroughs

2.5.1 (K)Ubuntu 18.04

You will need to install a couple of tools and libraries before compiling MIRA. Here's the recipe:

```
sudo apt-get install gcc make flex
sudo apt-get install libexpat1-dev libboost-all-dev
```

Once this is done, you can unpack and compile MIRA. For a dynamically linked version, use:

```
tar xvjf mira-5.0.0.tar.bz2
cd mira-5.0.0
./configure
make && make install
```

For a statically linked version, just change the configure line from above into:

```
./configure --enable-mirastatic
```

In case you also want to build documentation yourself, you will need to install this in addition:

```
sudo apt-get install xsltproc docbook-xsl dblatex
```

And building the documentation in PDF and HTML format is done like this:

```
make docs
```

Note

People working on git checkouts of the MIRA source code will obviously need some more tools. Get them with this:

```
sudo apt-get install automake libtool xutils-dev
```

2.5.2 openSUSE 12.1

Note This guide for OpenSUSE is outdated, but may give you clues on how to perform this on current systems.

You will need to install a couple of tools and libraries before compiling MIRA. Here's the recipe:

```
sudo zypper install gcc-c++ boost-devel
sudo zypper install flex libexpat-devel zlib-devel
```

Once this is done, you can unpack and compile MIRA. For a dynamically linked version, use:

```
tar xvjf mira-5.0.0.tar.bz2
cd mira-5.0.0
./configure
make && make install
```

In case you also want to build documentation yourself, you will need this in addition:

```
sudo zypper install docbook-xsl-stylesheets dblatex
```

Note

People working on git checkouts of the MIRA source code will obviously need some more tools. Get them with this:

```
sudo zypper install automake libtool xutils-dev
```

2.5.3 Fedora 17

Note This guide for Fedora is outdated, but may give you clues on how to perform this on current systems.

You will need to install a couple of tools and libraries before compiling MIRA. Here's the recipe:

```
sudo yum -y install gcc-c++ boost-devel
sudo yum install flex expat-devel vim-common zlib-devel
```

Once this is done, you can unpack and compile MIRA. For a dynamically linked version, use:

```
tar xvjf mira-5.0.0.tar.bz2
cd mira-5.0.0
./configure
make && make install
```

In case you also want to build documentation yourself, you will need this in addition:

```
sudo yum -y install docbook-xsl dblatex
```

Note

People working on git checkouts of the MIRA source code will obviously need some more tools. Get them with this:

```
sudo yum -y install automake libtool xorg-x1-util-devel
```

2.5.4 Mac OSX

These instructions are for OSX 10.14 (Mojave) and use Homebrew. There are other ways to do this (e.g., see the "compile everything from scratch"), but they are definitely more painful.

First, you will need to install the Apple XCode command line tools. This may be as easy as simply open a terminal and typing:

```
xcode-select --install
```

However, on the machine I have, I got weird compilation errors and found <https://github.com/frida/frida/issues/338#issuecomment-426777849> which told me to do this for things to work (don't ask me why, but this resolved my problems):

```
cd /Library/Developer/CommandLineTools/Packages/  
open macOS_SDK_headers_for_macOS_10.14.pkg
```

Then, if you do not already have it, install Homebrew. See <https://brew.sh/>.

Then go on and install with homebrew the LLVM clang compiler suite and a couple of other things needed:

```
brew install llvm boost flex expat
```

To compile from a git checkout, you will also need these:

```
brew install autoconf automake libtool git
```

To build the HTML documentation, this:

```
brew install docbook-xsl
```

And to build the PDF documentation, we need MacTex (huge download, around 1 GiB, 6 GiB on disk) and the dblatex package:

```
brew cask install mactex  
pip install dblatex
```

Finally, if you want to build small, rudimentary man pages:

```
brew install help2man
```

Now unpack MIRA, configure it and compile:

```
tar xvf mira-5.0.0.tar.bz2  
cd mira-5.0.0  
./configure --with-brew --enable-debug  
make -j 2
```

That's it for the dynamic version.

For building an almost static version, we need some trickery: configure with the `--enable-mirastatic` argument), then ask the build system to create a distribution. This will launch a toolchain at the end of which you will get a bundle which contains everything needed to run, both on OSX and other Unix systems.

```
./configure --with-brew --enable-mirastatic --enable-debug  
make  
make distrib
```

2.5.5 Compile everything from scratch

This lets you build a self-contained static MIRA binary. The only prerequisite here is that you have a working `gcc` with the minimum version described above. Please download all necessary files (expat, flex, etc.pp) and then simply follow the script below. The only things that you will want to change are the path used and, maybe, the name of some packages in case they were bumped up a version or revision.

Contributed by Sven Klages.

```
## whatever path is appropriate
cd /home/gls/SvenTemp/install

## expat
tar zxvf expat-2.0.1.tar.gz
cd expat-2.0.1
./configure --prefix=/home/gls/SvenTemp/expat
make && make install

## flex
cd /home/gls/SvenTemp/install
tar zxvf flex-2.5.35.tar.gz
cd flex-2.5.35
./configure --prefix=/home/gls/SvenTemp/flex
make && make install
cd /home/gls/SvenTemp/flex/bin
ln -s flex flex++
export PATH=/home/gls/SvenTemp/flex/bin:$PATH

## boost
cd /home/gls/SvenTemp/install
tar zxvf boost_1_48_0.tar.gz
cd boost_1_48_0
./bootstrap.sh --prefix=/home/gls/SvenTemp/boost
./b2 install

## mira itself
export CXXFLAGS="-I/home/gls/SvenTemp/flex/include"

cd /home/gls/SvenTemp/install
tar zxvf mira-3.4.0.1.tar.gz
cd mira-3.4.0.1
./configure --prefix=/home/gls/SvenTemp/mira \
--with-boost=/home/gls/SvenTemp/boost \
--with-expat=/home/gls/SvenTemp/expat \
--enable-mirastatic
make && make install
```

2.5.6 Dynamically linked MIRA

In case you do not want a static binary of MIRA, but a dynamically linked version, the following script by Robert Bruccoleri will give you an idea on how to do this.

Note that he, having root rights, puts all additional software in `/usr/local`, and in particular, he keeps updated versions of Boost and Flex there.

```
#!/bin/sh -x

make distclean
oze=`find . -name "*.o" -print`
if [[ -n "$oze" ]]
then
```

```

    echo "Not clean."
    exit 1
fi

export prefix=${BUILD_PREFIX:-/usr/local}
export LDFLAGS="-Wl,-rpath,$prefix/lib"

./configure --prefix=$prefix \
    --enable-debug=yes \
    --enable-mirastatic=no \
    --with-boost-libdir=$prefix/lib \
    --enable-optimisations \
    --enable-boundtracking=yes \
    --enable-bugtracking=yes \
    --enable-extendedbugtracking=no

make
make install

```

2.6 Compilation hints for other platforms.

2.6.1 NetBSD 5 (i386)

Contributed by Thomas Vaughan

The system flex (*/usr/bin/flex*) is too old, but the devel/flex package from a recent pkgsrc works fine. BSD make doesn't like one of the lines in *src/progs/Makefile*, so use GNU make instead (available from pkgsrc as *devel/gmake*). Other relevant pkgsrc packages: *devel/boost-libs*, *devel/boost-headers* and *textproc/expat*. The configure script has to be told about these pkgsrc prerequisites (they are usually rooted at */usr/pkg* but other locations are possible):

```
FLEX=/usr/pkg/bin/flex ./configure --with-expat=/usr/pkg --with-boost=/usr/pkg
```

If attempting to build a pkgsrc package of MIRA, note that the LDFLAGS passed by the pkgsrc mk files don't remove the need for the *--with-boost* option. The configure script complains about flex being too old, but this is harmless because it honours the \$FLEX variable when writing out makefiles.

2.7 Notes for distribution maintainers / system administrators

2.7.1 Additional data files

Depending on options/paramaters, the MIRA/mirabait binary may need to load some additional data during the run. By default this data will always be searched at this location: `LOCATION_OF_BINARY/../../share/mira/...`

That is: If the binary is, e.g., `/opt/mira5/bin/mira` with a softlink pointing from `/usr/local/bin/mira -> /opt/mira5/bin/mira` (because, e.g., `/usr/local/bin` may be by default in your PATH variable), then the additional data will be searched in `/opt/mira5/share/mira/...` and NOT in `/usr/local/share/mira/...`

Note In short: since MIRA 4.9.6, moving the binary is not enough anymore. Take care to have the *share* directory in the right place, i.e., adjacent to the directory the MIRA binary lives in.

Chapter 3

Preparing data

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Rome didn’t fall in a day either.”

—Solomon Short

3.1 Introduction

Most of this chapter and many sections are just stubs at the moment.

3.2 Sanger

Outside MIRA: transform .ab1 to .scf, perform sequencing vector clip (and cloning vector clip if used), basic quality clips.

Recommended program: **gap4** (or rather **pregap4**) from the Staden 4 package.

3.3 Roche / 454

Outside MIRA: convert SFF instrument from Roche to FASTQ, use **sff_extract** for that. In case you used "non-standard" sequencing procedures: clip away MIDs, clip away non-standard sequencing adaptors used in that project.

3.4 Illumina

Outside MIRA: for heavens’ sake: do NOT try to clip or trim by quality yourself. Do NOT try to remove standard sequencing adaptors yourself. Just leave Illumina data alone! (really, I mean it).

MIRA is much, much better at that job than you will probably ever be ... and I dare to say that MIRA is better at that job than 99% of all clipping/trimming software existing out there. Just make sure you use the [-CL:pec] (proposed_end_clip) option of MIRA.

Note The *only* exception to the above is if you (or your sequencing provider) used decidedly non-standard sequencing adaptors. Then it might be worthwhile to perform own adaptor clipping. But this will not be the case for 99% of all sequencing projects out there.

Joining paired-ends: if you want to do this, feel free to use any tool which is out there (TODO: quick list). Just make sure they do not join on very short overlaps. For me, the minimum overlap is at least 17 bases, but I more commonly use at least 30.

3.5 Ion Torrent

Outside MIRA: need to convert BAM to FASTQ. Need to clip away non-standard sequencing adaptors if used in that project. Apart from that: leave the data alone.

3.6 Short Read Archive (SRA)

Outside MIRA: you need to convert SRA format to FASTQ format. This is done using **fastq-dump** from the SRA toolkit from the NCBI. Make sure to have at least version 2.4.x of the toolkit. Last time I looked (March 2015), the software was at <http://www.ncbi.nlm.nih.gov/Traces/sra/?view=software>, the documentation for the whole toolkit was at http://www.ncbi.nlm.nih.gov/Traces/sra/?view=toolkit_doc, and for **fastq-dump** it was http://www.ncbi.nlm.nih.gov/Traces/sra/?view=toolkit_doc&f=fastq-dump

After extraction, proceed with preprocessing as described above, depending on the sequencing technology used.

For extracting Illumina data, use something like this:

```
arcadia:/some/path$ fastq-dump -I --split-files somefile.sra
```

Note

As **fastq-dump** unfortunately uses a pretty wasteful variant of the FASTQ format, you might want to reduce the file size for each FASTQ it produces by doing this:

```
sed -i '3~4 s/^+.*$/+/' file.fastq
```

The above command performs an in-file replacement of unnecessary name and comments on the quality divider lines of the FASTQ. The exact translation of the **sed** is: do an in-file replacement (-i); starting on the third line, then every fourth line (3~4); substitute (s/); a line which starts (^); with a plus (+); and then can have any character (.); repeated any number of times including zero (*); until the end of the line (\$); by just a single plus character (/+).

This alone reduces the file size of a typical Illumina data set with 100mers extracted from the SRA by about 15 to 20%.

Chapter 4

Cookbook: de-novo genome assemblies

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“The universe is full of surprises - most of them nasty.”

—Solomon Short

4.1 Introduction

This guide assumes that you have basic working knowledge of Unix systems, know the basic principles of sequencing (and sequence assembly) and what assemblers do.

While there are step by step instructions on how to setup your data and then perform an assembly, this guide expects you to read at some point in time

- Before the assembly, Chapter 3: “**Preparing data**” (p. 30) to know what to do (or not to do) with the sequencing data before giving it to MIRA.
- After the assembly, Chapter 7: “**Working with the results of MIRA**” (p. 55) to know what to do with the results of the assembly. More specifically, Section 7.1: “**MIRA output directories and files**” (p. 55), Section 7.2: “**First look: the assembly info**” (p. 57), Section 7.3: “**Converting results**” (p. 59), Section 7.4: “**Filtering results**” (p. 61) and Section 7.5: “**Places of importance in a de-novo assembly**” (p. 62).
- And also Chapter 9: “**MIRA 5 reference manual**” (p. 84) to look up how manifest files should be written (Section 9.3.2: “**The manifest file: basics**” (p. 86) and Section 9.3.3: “**The manifest file: information on the data you have**” (p. 87) and Section 9.3.4: “**The manifest file: extended parameters**” (p. 94)), some command line options as well as general information on what tags MIRA uses in assemblies, files it generates etc.pp
- Last but not least, you may be interested in some observations about the different sequencing technologies and the traps they may contain, see Chapter 11: “**Description of sequencing technologies**” (p. 152) for that. For advice on what to pay attention to *before* going into a sequencing project, have a look at Chapter 12: “**Some advice when going into a sequencing project**” (p. 165).

4.2 General steps

This part will introduce you step by step how to get your data together for a simple de-novo assembly. I’ll make up an example using an imaginary bacterium: *Bacillus chocorafoliensis* (or short: *Bchoc*). You collected the strain you want to assemble somewhere in the wild, so you gave the strain the name *Bchoc_wt*.

Just for laughs, let’s assume you sequenced that bug with lots of more or less current sequencing technologies: Sanger, 454, Illumina, Ion Torrent and Pacific Biosciences.

4.2.1 Copying and naming the sequence data

You need to create (or get from your sequencing provider) the sequencing data in any supported file format. Amongst these, FASTQ and FASTA + FASTA-quality will be the most common, although the latter is well on the way out nowadays. The following walkthrough uses what most people nowadays get: FASTQ.

Create a new project directory (e.g. `myProject`) and a subdirectory of this which will hold the sequencing data (e.g. `data`).

```
arcadia:/path/to mkdir myProject
arcadia:/path/to cd myProject
arcadia:/path/to/myProject$ mkdir data
```

Put the FASTQ data into that `data` directory so that it now looks perhaps like this:

```
arcadia:/path/to/myProject$ ls -l data
-rw-r--r-- 1 bach users 263985896 2008-03-28 21:49 bchocwt_lane6.solexa.fastq
```

Note I completely made up the file names above. You can name them anyway you want. And you can have them live anywhere on the hard-disk, you do not need to put them in this `data` directory. It's just the way I do it ... and it's where the example manifest files a bit further down in this chapter will look for the data files.

We're almost finished with the setup. As I like to have things neatly separated, I always create a directory called `assemblies` which will hold my assemblies (or different trials) together. Let's quickly do that:

```
arcadia:/path/to/myProject$ mkdir assemblies
arcadia:/path/to/myProject$ mkdir assemblies/1sttrial
arcadia:/path/to/myProject$ cd assemblies/1sttrial
```

4.2.2 Writing a simple manifest file

A manifest file is a configuration file for MIRA which tells it what type of assembly it should do and which data it should load. In this case we'll make a simple assembly of a genome with unpaired Illumina data

```
# Example for a manifest describing a genome de-novo assembly with
# unpaired Illumina data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome,denovo,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# here comes the unpaired Illumina data

readgroup = SomeUnpairedIlluminaReadsIGotFromTheLab
data = ../../data/bchocwt_lane6.solexa.fastq
technology = solexa
```

Note

Please look up the parameters of the manifest file in the main manual or the example manifest files in the following section. The ones above basically say: make an accurate denovo assembly of unpaired Illumina reads.

4.2.3 Starting the assembly

Starting the assembly is now just a matter of a simple command line:

```
arcadia:/path/to/myProject/assemblies/1sttrial$ mira manifest.conf >&log_assembly.txt
```

For this example - if you followed the walk-through on how to prepare the data - everything you might want to adapt in the first time are the following thing in the manifest file: options:

- project= (for naming your assembly project)

Of course, you are free to change any option via the extended parameters, but this is the topic of another part of this manual.

4.3 Manifest files for different use cases

This section will introduce you to manifest files for different use cases. It should cover the most important uses, but as always you are free to mix and match the parameters and readgroup definitions to suit your specific needs.

Taking into account that there may be *a lot* of combinations of sequencing technologies, sequencing libraries (shotgun, paired-end, mate-pair, etc.) and input file types (FASTQ, FASTA, GenBank, GFF3, etc.pp), the example manifest files just use Illumina and 454 as technologies, GFF3 as input file type for the reference sequence, FASTQ as input type for sequencing data ... and they do not show the multitude of more advanced features like, e.g., using ancillary clipping information in XML files, ancillary masking information in SSAHA2 or SMALT files etc.pp.

I'm sure you will be able to find your way by scanning through the corresponding section on manifest files in the reference chapter :-)

4.3.1 Manifest for shotgun data

Well, we've seen that already in the section above, but here it is again ... but this time with 454 data.

```
# Example for a manifest describing a denovo assembly with
# unpaired 454 data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome, denovo, accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# here's the 454 data

readgroup = SomeUnpaired454ReadsIGotFromTheLab
data = ../../data/some454data.fastq
technology = 454
```

4.3.2 Assembling with multiple sequencing technologies (hybrid assemblies)

Hybrid de-novo assemblies follow the general manifest scheme: tell what you want in the first part, then simply add as separate readgroup the information MIRA needs to know to find the data and off you go. Just for laughs, here's a manifest for 454 shotgun with Illumina shotgun

```

# Example for a manifest describing a denovo assembly with
# shotgun 454 and shotgun Illumina data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome,denovo,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# now the shotgun 454 data
readgroup = DataForShotgun454
data = ../../data/project454data.fastq
technology = 454

# now the shotgun Illumina data

readgroup = DataForShotgunIllumina
data = ../../data/someillumina.fastq
technology = solexa

```

4.3.3 Manifest for data sets with paired reads

When using paired-end data, you should know

1. the orientation of the reads toward each other. This is specific to sequencing technologies and / or the sequencing library preparation.
2. at which distance these reads should be. This is specific to the sequencing library preparation and the sequencing lab should tell you this.

In case you do not know one (or any) of the above, don't panic! MIRA is able to estimate the needed values during the assembly if you tell it to.

The following manifest shows you the most laziest way to define a paired data set by simply adding *autopairing* as keyword to a readgroup (using Illumina just as example):

```

# Example for a lazy manifest describing a denovo assembly with
# one library of paired reads

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome,denovo,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# now the Illumina paired-end data

readgroup = DataIlluminaPairedLib
autopairing
data = ../../data/project_1.fastq ../../data/project_2.fastq
technology = solexa

```

If you know the orientation of the reads and/or the library size, you can tell this MIRA the following way (just showing the readgroup definition here):

```
readgroup = DataIlluminaPairedEnd500Lib
data = ../../data/project_1.fastq ../../data/project_2.fastq
technology = solexa
template_size = 250 750
segment_placement = ----> <----
```

In cases you are not 100% sure about, e.g., the size of the DNA template, you can also give a (generous) expected range and then tell MIRA to automatically refine this range during the assembly based on real, observed distances of read pairs. Do this with *autorefine* modifier like this:

```
template_size = 50 1000 autorefine
```

The following manifest file is an example for assembling with several different libraries from different technologies. Do not forget you can use *autopairing* or *autorefine* :-)

```
# Example for a manifest describing a denovo assembly with
# several kinds of sequencing libraries

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome,denovo,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# now the Illumina paired-end data

readgroup = DataIlluminaForPairedEnd500bpLib
data = ../../data/project500bp-1.fastq ../../data/project500bp-2.fastq
technology = solexa
strain = bchoc_sel1
template_size = 250 750
segment_placement = ----> <----
```

```
# now the Illumina mate-pair data

readgroup = DataIlluminaForMatePair3kbLib
data = ../../data/project3kb-1.fastq ../../data/project3kb-2.fastq
technology = solexa
strain = bchoc_sel1
template_size = 2500 3500
segment_placement = <---- ---->
```

```
# some Sanger data (6kb library)

readgroup = DataForSanger6kbLib
data = ../../data/sangerdata.fastq
technology = sanger
template_size = 5500 6500
segment_placement = ----> <----
```

```
# some 454 data

readgroup = DataFo454Pairs
data = ../../data/454data.fastq
technology = 454
```

```

template_size = 8000 1200
segment_placement = 2----> 1---->

# some Ion Torrent data

readgroup = DataFoIonPairs
data = ../../data/iondata.fastq
technology = iontor
template_size = 1000 300
segment_placement = 2----> 1---->

```

4.3.4 De-novo with multiple strains

MIRA will make use of ancillary information present in the manifest file. One of these is the information to which strain (or organism or cell line etc.pp) the generated data belongs.

You just need to tell in the manifest file which data comes from which strain. Let's assume that in the example from above, the "lane6" data were from a first mutant named *bhoc_se1* and the "lane7" data were from a second mutant named *bhoc_se2*. Here's the manifest file you would write then:

```

# Example for a manifest describing a de-novo assembly with
# unpaired Illumina data, but from multiple strains

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstAssembly
job = genome,denovo,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# now the Illumina data

readgroup = DataForSE1
data = ../../data/bhocse_lane6.solexa.fastq
technology = solexa
strain = bhoc_se1

readgroup = DataForSE2
data = ../../data/bhocse_lane7.solexa.fastq
technology = solexa
strain = bhoc_se2

```

Note While assembling de-novo with multiple strains is possible, the interpretation of results may become a bit daunting in some cases. For many scenarios it might therefore be preferable to successively use the data sets in own assemblies or mappings.

This *strain* information for each readgroup is really the only change you need to perform to tell MIRA everything it needs for handling strains.

Chapter 5

Cookbook: Mapping

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“You have to know what you’re looking for before you can find it.”

—Solomon Short

5.1 Introduction

This guide assumes that you have basic working knowledge of Unix systems, know the basic principles of sequencing (and sequence assembly) and what assemblers do.

While there are step by step instructions on how to setup your data and then perform an assembly, this guide expects you to read at some point in time

- Before the mapping, Chapter 3: “[Preparing data](#)” (p. 30) to know what to do (or not to do) with the sequencing data before giving it to MIRA.
- Generally, the Chapter 7: “[Working with the results of MIRA](#)” (p. 55) to know what to do with the results of the assembly. More specifically, Section 7.3: “[Converting results](#)” (p. 59) Section 7.6: “[Places of interest in a mapping assembly](#)” (p. 66) Section 7.7: “[Post-processing mapping assemblies](#)” (p. 72)
- And also Chapter 9: “[MIRA 5 reference manual](#)” (p. 84) to look up how manifest files should be written (Section 9.3.2: “[The manifest file: basics](#)” (p. 86) and Section 9.3.3: “[The manifest file: information on the data you have](#)” (p. 87) and Section 9.3.4: “[The manifest file: extended parameters](#)” (p. 94)), some command line options as well as general information on what tags MIRA uses in assemblies, files it generates etc.pp
- Last but not least, you may be interested in some observations about the different sequencing technologies and the traps they may contain, see Chapter 11: “[Description of sequencing technologies](#)” (p. 152) for that. For advice on what to pay attention to *before* going into a sequencing project, have a look at Chapter 12: “[Some advice when going into a sequencing project](#)” (p. 165).

5.2 General steps

This part will introduce you step by step how to get your data together for a simple mapping assembly.

I’ll make up an example using an imaginary bacterium: *Bacillus chocorafoliensis* (or short: *Bchoc*).

In this example, we assume you have two strains: a wild type strain of *Bchoc_wt* and a mutant which you perhaps got from mutagenesis or other means. Let’s imagine that this mutant needs more time to eliminate a given amount of chocolate, so we call the mutant *Bchoc_se* ... SE for slow eater

You wanted to know which mutations might be responsible for the observed behaviour. Assume the genome of *Bhoc_wt* is available to you as it was published (or you previously sequenced it), so you resequenced *Bhoc_se* with Solexa to examine mutations.

5.2.1 Copying and naming the sequence data

You need to create (or get from your sequencing provider) the sequencing data in either FASTQ or FASTA + FASTA quality format. The following walkthrough uses what most people nowadays get: FASTQ.

Create a new project directory (e.g. `myProject`) and a subdirectory of this which will hold the sequencing data (e.g. `data`).

```
arcadia:/path/to mkdir myProject
arcadia:/path/to cd myProject
arcadia:/path/to/myProject$ mkdir data
```

Put the FASTQ data into that `data` directory so that it now looks perhaps like this:

```
arcadia:/path/to/myProject$ ls -l data
-rw-r--r-- 1 bach users 263985896 2008-03-28 21:49 bhocse_lane6.solexa.fastq
-rw-r--r-- 1 bach users 264823645 2008-03-28 21:51 bhocse_lane7.solexa.fastq
```

Note I completely made up the file names above. You can name them anyway you want. And you can have them live anywhere on the hard disk, you do not need to put them in this `data` directory. It's just the way I do it ... and it's where the example manifest files a bit further down in this chapter will look for the data files.

5.2.2 Copying and naming the reference sequence

The reference sequence (the backbone) can be in a number of different formats: GFF3, GenBank (`.gb`, `.gbk`, `.gbf`, `.gbff`), MAF, CAF, FASTA. The first three have the advantage of being able to carry additional information like, e.g., annotation. In this example, we will use a GFF3 file like the ones one can download from the NCBI.

Note TODO: Write why GFF3 is better and where to get them at the NCBI.

So, let's assume that our wild type strain is in the following file: `NC_someNCBInumber.gff3`.

You do not need to copy the reference sequence to your directory, but I normally copy also the reference file into the directory with my data as I want to have, at the end of my work, a nice little self-sufficient directory which I can archive away and still be sure that in 10 years time I have all data I need together.

```
arcadia:/path/to/myProject$ cp /somewhere/NC_someNCBInumber.gff3 data
arcadia:/path/to/myProject$ ls -l data
-rw-r--r-- 1 bach users 6543511 2008-04-08 23:53 NC_someNCBInumber.gff3
-rw-r--r-- 1 bach users 263985896 2008-03-28 21:49 bhocse_lane6.solexa.fastq
-rw-r--r-- 1 bach users 264823645 2008-03-28 21:51 bhocse_lane7.solexa.fastq
```

We're almost finished with the setup. As I like to have things neatly separated, I always create a directory called `assemblies` which will hold my assemblies (or different trials) together. Let's quickly do that:

```
arcadia:/path/to/myProject$ mkdir assemblies
arcadia:/path/to/myProject$ mkdir assemblies/1sttrial
arcadia:/path/to/myProject$ cd assemblies/1sttrial
```

5.2.3 Writing a simple manifest file

A manifest file is a configuration file for MIRA which tells it what type of assembly it should do and which data it should load. In this case we have unpaired sequencing data which we want to map to a reference sequence, the manifest file for that is pretty simple:

```
# Example for a manifest describing a mapping assembly with
# unpaired Illumina data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstMapping
job = genome,mapping,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first, the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the Illumina data

readgroup = SomeUnpairedIlluminaReadsIGotFromTheLab
data = ../../data/*fastq
technology = solexa
strain = bchoc_se
```

Note

Please look up the parameters of the manifest file in the main manual or the example manifest files in the following section. The ones above basically say: make an accurate mapping of Solexa reads against a genome; in one pass; the name of the backbone strain is 'bchoc_wt'; the data with the backbone sequence (and maybe annotations) is in a specified GFF3 file; for Solexa data: assign default strain names for reads which have not loaded ancillary data with strain info and that default strain name should be 'bchoc_se'.

5.2.4 Starting the assembly

Starting the assembly is now just a matter of a simple command line:

```
arcadia:/path/to/myProject/assemblies/1sttrial$ mira manifest.conf >&log_assembly.txt
```

For this example - if you followed the walk-through on how to prepare the data - everything you might want to adapt in the first time are the following thing in the manifest file: options:

- `project=` (for naming your assembly project)
- `strain_name=` to give the names of your reference and mapping strain

Of course, you are free to change any option via the extended parameters, but this is the topic of another part of this manual.

5.3 Manifest files for different use cases

This section will introduce you to manifest files for different use cases. It should cover the most important uses, but as always you are free to mix and match the parameters and readgroup definitions to suit your specific needs.

Taking into account that there may be *a lot* of combinations of sequencing technologies, sequencing libraries (shotgun, paired-end, mate-pair, etc.) and input file types (FASTQ, FASTA, GenBank, GFF3, etc.pp), the example manifest files just use Illumina and 454 as technologies, GFF3 as input file type for the reference sequence, FASTQ as input type for sequencing data ... and they do not show the multitude of more advanced features like, e.g., using ancillary clipping information in XML files, ancillary masking information in SSAHA2 or SMALT files etc.pp.

I'm sure you will be able to find your way by scanning through the corresponding section on manifest files in the reference chapter :-)

5.3.1 Mapping with shotgun data

Well, we've seen that already in the section above, but here it is again ... this time with Ion Torrent data though.

```
# Example for a manifest describing a mapping assembly with
# unpaired Ion data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstMapping
job = genome,mapping,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first, the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the Ion Torrent data

readgroup = SomeUnpairedIonReadsIGotFromTheLab
data = ../../data/someiondata.fastq
technology = iontor
strain = bchoc_se
```

5.3.2 Manifest for data sets with paired reads

When using paired-end data in mapping, you must decide whether you want

1. use the MIRA feature to create long 'coverage equivalent reads' (CERs) which saves a lot of memory (both in the assembler and later on in an assembly editor). However, you then *lose information about read pairs!*
2. or whether you want to *keep information about read pairs* at the expense of larger memory requirements both in MIRA and in assembly finishing tools or viewers afterwards.
3. or a mix of the two above

The Illumina pipeline generally normally gives you two files for paired-end data: a `project-1.fastq` and `project-2.fastq`. The first file containing the first read of a read-pair, the second file the second read. Depending on the preprocessing pipeline of your sequencing provider, the names of the reads are either the very same in both files or already have a `/1` or `/2` appended. Also, your sequencing provider may give you one big file where the reads from both ends are present.

Note

MIRA can read all FASTQ variants produced by various Illumina pipelines, be they with or without the `/1` and `/2` already appended to the names. You generally do not need to do any name mangling before feeding the data to MIRA. However, MIRA will shell out a warning if read names are longer than 40 characters.

When using paired-end data, you should know

1. the orientation of the reads toward each other. This is specific to sequencing technologies and / or the sequencing library preparation.
2. at which distance these reads should be. This is specific to the sequencing library preparation and the sequencing lab should tell you this.

In case you do not know one (or any) of the above, don't panic! MIRA is able to estimate the needed values during the assembly if you tell it to.

The following manifest shows you the most laziest way to define a paired data set by simply adding *autopairing* as keyword to a readgroup (using Illumina just as example):

```
# Example for a lazy manifest describing a mapping assembly with
# one library of paired reads

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstMapping
job = genome,mapping,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the Illumina paired-end data

readgroup = DataIlluminaPairedLib
autopairing
data = ../../data/project_1.fastq ../../data/project_2.fastq
technology = solexa
strain = bchoc_sel
```

See? Wasn't hard and it did not hurt, did it? One just needs to tell MIRA it should expect paired reads via the *autopairing* keyword and that is everything you need.

If you are lazy, or want to efficiently include multiple FASTQ files, you can use regular expressions in the line which defines the data to load. E.g., if there were only two FASTQ files in the directory named `_1` and `_2`, you can write this:

```
data = ../../data/project_?.fastq
```

Notice the question mark in the above line? Read more about how to use this and other shortcuts in Section 9.3.3.2: “Defining data files to load” (p. 88)

If you know the orientation of the reads and/or the library size, you can tell this MIRA the following way (just showing the readgroup definition here):

```
readgroup = DataIlluminaPairedEnd500Lib
data = ../../data/project_1.fastq ../../data/project_2.fastq
technology = solexa
template_size = 250 750
segment_placement = ---> <---
```

In cases you are not 100% sure about, e.g., the size of the DNA template, you can also give a (generous) expected range and then tell MIRA to automatically refine this range during the assembly based on real, observed distances of read pairs. Do this with *autorefine* modifier like this:

```
template_size = 50 1000 autorefine
```

The following manifest file is an example for mapping a 500 bp paired-end and a 3kb mate-pair library of a strain called *bhoc_sel* against a GenBank reference file containing a strain called *bhoc_wt*:

```
# Example for a manifest describing a mapping assembly with
# paired Illumina data, not merging reads and therefore keeping
# all pair information

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode
# As special parameter, we want to switch off merging of Solexa reads

project = MyFirstMapping
job = genome,mapping,accurate
parameters = SOLEXA_SETTINGS -CO:msr=no

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first, the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bhoc_wt

# now the Illumina data

readgroup = DataForPairedEnd500bpLib
autopairing
data = ../../data/project500bp-1.fastq ../../data/project500bp-2.fastq
technology = solexa
strain = bhoc_sel

readgroup = DataForMatePair3kbLib
data = ../../data/project3kb-1.fastq ../../data/project3kb-2.fastq
technology = solexa
strain = bhoc_sel
template_size = 2000 4000 autorefine
segment_placement = <--- --->
```

Please look up the parameters used in the main manual. The ones above basically say: make an accurate mapping of Solexa reads against a genome. Additionally do not merge short short Solexa reads to the contig.

For the paired-end library, be lazy and let MIRA find out everything it needs. However, that information should be treated as "information only" by MIRA, i.e., it is not used for deciding whether a pair is well mapped.

For the mate-pair library, assume a DNA template template size of 2000 to 4000 bp (but let MIRA automatically refine this using observed distances) and the segment orientation of the read pairs follows the reverse / forward scheme. That information should be treated as "information only" by MIRA, i.e., it is not used for deciding whether a pair is well mapped.

Comparing this manifest with a manifest for unpaired-data, two parameters were added in the section for Solexa data:

1. `-CO:msr=no` tells MIRA not to merge reads that are 100% identical to the backbone. This also allows to keep the template information (distance and orientation) for the reads.
2. `template_size` tells MIRA at which distance the two reads should normally be placed from each other.
3. `segment_placement` tells MIRA how the different segments (reads) of a DNA template have to be ordered to form a valid representation of the sequenced DNA.

Note

Note that in mapping assemblies, these `template_distance` and `segment_placement` parameters are normally treated as *information only*, i.e., MIRA will map the reads regardless whether the distance and orientation criteria are met or not. This enables post-mapping analysis programs to hunt for genome rearrangements or larger insertions/deletion.



Warning

If template size and segment placement checking were on, the following would happen at, e.g. sites of re-arrangement: MIRA would map the first read of a read-pair without problem. However, it would very probably reject the second read because it would not map at the specified distance or orientation from its partner. Therefore, in mapping assemblies with paired-end data, checking of the template size must be switched off to give post-processing programs a chance to spot re-arrangements.

5.3.3 Mapping with multiple sequencing technologies (hybrid mapping)

I'm sure you'll have picked up the general scheme of manifest files by now. Hybrid mapping assemblies follow the general scheme: simply add as separate readgroup the information MIRA needs to know to find the data and off you go. Just for laughs, here's a manifest for 454 shotgun with Illumina paired-end

```
# Example for a manifest describing a mapping assembly with
# shotgun 454 and paired-end Illumina data, not merging reads and therefore keeping
# all pair information

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode
# As special parameter, we want to switch off merging of Solexa reads

project = MyFirstMapping
job = genome,mapping,accurate
parameters = SOLEXA_SETTINGS -CO:msr=no

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first, the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the shotgun 454 data
readgroup = DataForShotgun454
```

```

data = ../../data/project454data.fastq
technology = 454
strain = bchoc_se1

# now the paired-end Illumina data

readgroup = DataForPairedEnd500bpLib
data = ../../data/project500bp-1.fastq ../../data/project500bp-2.fastq
technology = solexa
strain = bchoc_se1
template_size = 250 750
segment_placement = ---> <---
```

5.3.4 Mapping with multiple strains

MIRA will make use of ancillary information present in the manifest file. One of these is the information to which strain (or organism or cell line etc.pp) the generated data belongs.

You just need to tell in the manifest file which data comes from which strain. Let's assume that in the example from above, the "lane6" data were from a first mutant named *bchoc_se1* and the "lane7" data were from a second mutant named *bchoc_se2*. Here's the manifest file you would write then:

```

# Example for a manifest describing a mapping assembly with
# unpaired Illumina data

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map a genome in accurate mode

project = MyFirstMapping
job = genome,mapping,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first, the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the Illumina data

readgroup = DataForSE1
data = ../../data/bchocse_lane6.solexa.fastq
technology = solexa
strain = bchoc_se1

readgroup = DataForSE2
data = ../../data/bchocse_lane7.solexa.fastq
technology = solexa
strain = bchoc_se2
```

Note While mapping (or even assembling de-novo) with multiple strains is possible, the interpretation of results may become a bit daunting in some cases. For many scenarios it might therefore be preferable to successively use the data sets in own mappings or assemblies.

This *strain* information for each readgroup is really the only change you need to perform to tell MIRA everything it needs for handling strains.

5.3.5 Mapping EST / RNASeq data

For bacteria you can map EST / RNASeq data against any reference sequence(s), be it a genome reference or transcriptome reference.

For eukaryotes, I recommend mapping against transcriptome sequences only as MIRA does not handle splice sites very well. You can get away mapping RNASeq data against 'simple' eukaryotes which have very few and small introns, like *S. cerevisiae* or simple fungi; but just note that the expression level for exons near larger introns will be slightly lower than the rest of the gene.

Note

The standard MIRA parameters are geared towards handling difficult mapping cases where the reference sequence can be quite different to what is actually mapped against it (10% average difference, up to 15% locally).

If the data you are mapping is actually from a strain which is very similar to your reference (give or take a couple of SNPs, genome reorganisations and / or insertions and deletions), you can save a lot of time in the mapping process by relaxing a few parameters like [-SK:kms=(high-value)], [-SB:bnb=no], [-FM:mte=(low-value)].

Here's a very simple, albeit powerful manifest which will allow you to retrieve both SNPs and feature coverage data from the mapping results:

```
# Example for a lazy manifest describing a mapping assembly with
# one library of paired reads

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should map EST / RNASeq data in accurate mode
# Accurate mode may be "too good" for mapping, see next example

project = MyFirstMapping
job = est,mapping,accurate

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups"

# first the reference sequence
readgroup
is_reference
data = ../../data/NC_someNCBInumber.gff3
strain = bchoc_wt

# now the Illumina data

readgroup
data = ../../data/project_1.fastq ../../data/project_2.fastq
technology = solexa
strain = bchoc_sel
```

If you compare the above manifest to a manifest for a simple genome mapping, the only line which really changed is the *job=...* line, which changed from *genome,mapping,accurate* to *est,mapping,accurate*

As already alluded to, the standard MIRA mapping parameters are made to find almost anything that maps, and that takes time. In case you are mapping data which is very closely related to the reference sequence you are using, you can save time by changing the manifest slightly. Start it like this:

```
# Example for a lazy manifest describing a mapping assembly with
# one library of paired reads which is close to your reference sequence

# First part: defining some basic things
```

```
project = MyFirstMapping
job = est,mapping,accurate

# Accurate mode may be "too good" for mapping, therefore these parameters to
# speed up things
parameters = -SK:kms=20 SOLEXA_SETTINGS -SB:bnb=no -FM:mte=6

# remaining manifest as above
# ...
```

5.4 Walkthroughs

5.4.1 Walkthrough: mapping of E.coli from Lenski lab against E.coli B REL606

TODO: Sorry, needs to be re-written for the relatively new SRR format distributed at the NCBI ... and changes in MIRA 3.9.x. Please come back later.

5.5 Useful things to know about reference sequences

There are a few things to consider when using reference sequences:

1. MIRA is not really made to handle a big amount of reference sequences as they currently need inane amounts of memory. Use other programs for mapping against more than, say, 200 megabases.
2. Reference sequences can be as long as needed! They are not subject to normal read length constraints of a maximum of 32k bases. That is, if one wants to load one or several entire chromosomes of a bacterium or lower eukaryote as backbone sequence(s), this is just fine.
3. Reference sequences can be single sequences like provided in, e.g., FASTA, FASTQ, GFF or GenBank files. But reference sequences also can be whole assemblies when they are provided as, e.g., MAF or CAF format. This opens the possibility to perform semi-hybrid assemblies by assembling first reads from one sequencing technology de-novo (e.g. PacBio) and then map reads from another sequencing technology (e.g. Solexa) to the whole PacBio alignment instead of mapping it to the PacBio consensus.

A semi-hybrid assembly will therefore contain, like a hybrid assembly, the reads of both sequencing technologies.

4. Reference sequences will not be reversed! They will always appear in forward direction in the output of the assembly. Please note: if the backbone sequence consists of a MAF or CAF file that contain contigs which contain reversed reads, then the contigs themselves will be in forward direction. But the reads they contain that are in reverse complement direction will of course also stay reverse complement direction.
5. Reference sequences will not not be assembled together! That is, even if a reference sequence has a perfect overlap with another reference sequence, they will still not be merged.
6. Reads are assembled to reference sequences in a first come, first served scattering strategy.
Suppose you have two identical reference sequences and a read which would match both, then the read would be mapped to the first backbone. If you had two identical reads, the first read would go to the first backbone, the second read to the second backbone. With three identical reads, the first backbone would get two reads, the second backbone one read. Etc.pp.
7. Only in references loaded from MAF or CAF files: contigs made out of single reads (singlets) loose their status as reference sequence and will be returned to the normal read pool for the assembly process. That is, these sequences will be assembled to other reference sequences or with each other.

5.6 Known bugs / problems

These are actual for version 4.0 of MIRA and might or might not have been addressed in later version.

Bugs:

1. mapping of paired-end reads with one read being in non-repetitive area and the other in a repeat is not as effective as it should be. The optimal strategy to use would be to map first the non-repetitive read and then the read in the repeat. Unfortunately, this is not yet implemented in MIRA.

Chapter 6

Cookbook: de-novo EST / RNASeq assemblies

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Expect the worst. You’ll never get disappointed.”

—Solomon Short

6.1 Introduction

This document is not complete yet and some sections may be a bit unclear. I’d be happy to receive suggestions for improvements.

Some reading requirements

This guide assumes that you have basic working knowledge of Unix systems, know the basic principles of sequencing (and sequence assembly) and what assemblers do. Basic knowledge on mRNA transcription should also be present.

Please read at some point in time

- Before the assembly, Chapter 3: [“Preparing data”](#) (p. 30) to know what to do (or not to do) with the sequencing data before giving it to MIRA.
- For setting up the assembly, Chapter 4: [“Cookbook: de-novo genome assemblies”](#) (p. 32) to know how to start a denovo assembly (except you obviously will need to change the `--job` setting from *genome* to *est*).
- After the assembly, Chapter 7: [“Working with the results of MIRA”](#) (p. 55) to know what to do with the results of the assembly. More specifically, Section 7.1: [“MIRA output directories and files”](#) (p. 55), Section 7.2: [“First look: the assembly info”](#) (p. 57), Section 7.3: [“Converting results”](#) (p. 59), Section 7.4: [“Filtering results”](#) (p. 61) and Section 7.5: [“Places of importance in a de-novo assembly”](#) (p. 62).
- And also Chapter 9: [“MIRA 5 reference manual”](#) (p. 84) to look up how manifest files should be written (Section 9.3.2: [“The manifest file: basics”](#) (p. 86) and Section 9.3.3: [“The manifest file: information on the data you have”](#) (p. 87) and Section 9.3.4: [“The manifest file: extended parameters”](#) (p. 94)), some command line options as well as general information on what tags MIRA uses in assemblies, files it generates etc.pp

6.2 Preliminaries: on the difficulties of assembling ESTs /RNASeq

Assembling ESTs can be, from an assemblers point of view, pure horror. E.g., it may be that some genes have thousands of transcripts while other genes have just one single transcript in the sequenced data. Furthermore, the presence of 5’ and 3’ UTR, transcription variants, splice variants, homologues, SNPs etc.pp complicates the assembly in some rather interesting ways.

6.2.1 Poly-A tails

Poly-A tails are part of the mRNA and therefore also part of sequenced data. They can occur as poly-A or poly-T, depending from which direction and which part of the mRNA was sequenced. Having poly-A/T tails in the data is a something of a double edged sword. More specifically., if the 3' poly-A tail is kept unmasked in the data, transcripts having this tail will very probably not align with similar transcripts from different splice variants (which is basically good). On the other hand, homopolymers (multiple consecutive bases of the same type) like poly-As are features that are pretty difficult to get correct with today's sequencing technologies, be it Sanger, Solexa or, with even more problems problems, 454. So slight errors in the poly-A tail could lead to wrongly assigned splice sites ... and wrongly split contigs.

This is the reason why many people cut off the poly-A tails. Which in turn may lead to transcripts from different splice variants being assembled together.

Either way, it's not pretty.

6.2.2 Lowly expressed transcripts

Single transcripts (or very lowly expressed transcripts) containing SNPs, splice variants or similar differences to other, more highly expressed transcripts are a problem: it's basically impossible for an assembler to distinguish them from reads containing junky data (e.g. read with a high error rate or chimeras). The standard setting of many EST assemblers and clusterers is therefore to remove these reads from the assembly set. MIRA handles things a bit differently: depending on the settings, single transcripts with sufficiently large differences are either treated as debris or can be saved as *singlet*.

6.2.3 Very highly expressed transcripts

Another interesting problem for de-novo assemblers are non-normalised libraries. In each cell, the number of mRNA copies per gene may differ by several orders of magnitude, from a single transcripts to several tens of thousands. Pre-sequencing normalisation is a wet-lab procedure to approximately equalise those copy numbers. This can however, introduce other artifacts.

If an assembler is fed with non-normalised EST data, it may very well be that an overwhelming number of the reads comes only from a few genes (house-keeping genes). In Sanger sequencing projects this could mean a couple of thousand reads per gene. In 454 sequencing projects, this can mean several tens of thousands of reads per genes. With Solexa data, this number can grow to something close to a million.

Several effects then hit a de-novo assembler, the three most annoying being (in ascending order of annoyance): a) non-random sequencing errors then look like valid SNPs, b) sequencing and library construction artefacts start to look like valid sequences if the data set was not cleaned "enough" and more importantly, c) an explosion in time and memory requirements when attempting to deliver a "good" assembly. While MIRA has methods to deal with this kind of data (e.g. via digital normalisation), a sure sign of the latter are messages from MIRA about *megahubs* in the data set.

Note The guide on how to tackle *hard* projects with MIRA gives an overview on how to hunt down sequences which can lead to the assembler getting confused, be it sequencing artefacts or highly expressed genes.

6.2.4 Chimeras

Chimeras are sequences containing adjacent base stretches which are not occurring in an organism as sequenced, neither as DNA nor as (m)RNA. Chimeras can be created through recombination effects during library construction or sequencing. Chimeras can, and often do, lead to misassemblies of sequence stretches into one contig although they do not belong together. Have a look at the following example where two stretches (denoted by x and o) are joined by a chimeric read *r4* containing both stretches:

```
r1 xxxxxxxxxxxxxxxxxxxx
r2 xxxxxxxxxxxxxxxxxxxx
r3 xxxxxxxxxxxxxxxxxxxx
r4 xxxxxxxxxxxxxxxxxxxx | oooooooooooooooooo
r5                          oooooooooooooo
r6                          oooooooooooooo
r7                          oooooooooooooo
```

The site of the recombination event is denoted by \times | \circ in read $r4$.

MIRA does have a chimera detection -- which works very well in genome assemblies due to high enough coverage -- by searching for sequence stretches which are not covered by overlaps. In the above example, the chimera detection routine will almost certainly flag read $r4$ as chimera and only use a part of it: either the \times or \circ part, depending on which part is longer. There is always a chance that $r4$ is a valid read though, but that's a risk to take.

Now, that strategy would also work totally fine in EST projects if one would not have to account for lowly expressed genes. Imagine the following situation:

```
s1 xxxxxxxxxxxxxxxxxxxx
s2      xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx
s3                                xxxxxxxxxxxxxxxxxxxx
```

Look at read $s2$; from an overlap coverage perspective, $s2$ could also very well be a chimera, leading to a break of an otherwise perfectly valid contig if $s2$ were cut back accordingly. This is why chimera detection is switched off by default in MIRA.



Warning

When starting an EST assembly via the `--job=est, . . .` switch, chimera detection is switched off by default. It is absolutely possible to switch on the SKIM chimera detection afterwards via `[-CL:ascdc]`. However, this will have exactly the effects described above: chimeras in higher coverage contigs will be detected, but perfectly valid low coverage contigs will be torn apart.

It is up to you to decide what you want or need.

6.3 Preprocessing of ESTs

With contributions from Katrina Dlugosch

EST sequences necessarily contain fragments of vectors or primers used to create cDNA libraries from RNA, and may additionally contain primer and adaptor sequences used during amplification-based library normalisation and/or high-throughput sequencing. These contaminant sequences need to be removed prior to assembly. MIRA can trim sequences by taking contaminant location information from a SSAHA2 or SMALT search output, or users can remove contaminants beforehand by trimming sequences themselves or masking unwanted bases with lowercase or other characters (e.g. 'x', as with **cross_match**). Many folks use preprocessing trimming/masking pipelines because it can be very important to try a variety of settings to verify that you've removed all of your contaminants (and fragments thereof) before sending them into an assembly program like MIRA. It can also be good to spend some time seeing what contaminants are in your data, so that you get to know what quality issues are present and how pervasive.

Two features of next generation sequencing can introduce errors into contaminant sequences that make them particularly difficult to remove, arguing for preprocessing: First, most next-generation sequence platforms seem to be sensitive to excess primers present during library preparation, and can produce a small percentage of sequences composed entirely of concatenated primer fragments. These are among the most difficult contaminants to remove, and the program TagDust (<http://genome.gsc.riken.jp/osc/english/dataresource/>) was recently developed specifically to address this problem. Second, 454 EST data sets can show high variability within primer sequences designed to anchor to polyA tails during cDNA synthesis, because 454 has trouble calling the length of the necessary A and T nucleotide repeats with accuracy.

A variety of programs exist for preprocessing. Popular ones include **cross_match** (<http://www.phrap.org/phredphrapconsequence.html>) for primer masking, and SeqClean (<http://compbio.dfci.harvard.edu/tgi/software/>), Lucy (<http://lucy.sourceforge.net/>), and SeqTrim (http://www.scbi.uma.es/cgi-bin/seqtrim/seqtrim_login.cgi) for both primer and polyA/T trimming. The pipeline SnoWhite (<http://evopipes.net>) combines Seqclean and TagDust with custom scripts for aggressive sequence and polyA/T trimming (and is tolerant of data already masked using **cross_match**). In all cases, the user must provide contaminant sequence information and adjust settings for how sensitive the programs should be to possible matches. To find the best settings, it is helpful to look directly at some of the sequences that are being trimmed and inspect them for remaining primer and/or polyA/T fragments after cleaning.



Warning When using **mira** with the the simplest parameter calls (using the "--job=..." quick switches), the default settings used include pretty heavy sequence pre-processing to cope with noisy data. Especially if you have your own pre-processing pipeline, you *must* then switch off different clip algorithms that you might have applied previously yourself. Especially poly-A clips should never be run twice (by your pipeline and by **mira**) as they invariably lead to too many bases being cut away in some sequences,

Note Here too: In some cases MIRA can get confused if something with the pre-processing went wrong because, e.g., unexpected sequencing artefacts like unknown sequencing vectors or adaptors remain in data. The guide on how to tackle *hard* projects with MIRA gives an overview on how to hunt down sequences which can lead to the assembler getting confused, be it sequencing artefacts or highly expressed genes.

6.4 The difference between *assembly* and *clustering*

MIRA in its base settings is an *assembler* and not a *clusterer*, although it can be configured as such. As assembler, it will split up read groups into different contigs if it thinks there is enough evidence that they come from different RNA transcripts.

6.4.1 Splitting transcripts into contigs based on SNPs

Imagine this simple case: a gene has two slightly different alleles and you've sequenced this:

```
A1-1 .....T.....
A1-2 .....T.....
A1-3 .....T.....
A1-4 .....T.....
A1-5 .....T.....
B2-1 .....G.....
B2-2 .....G.....
B2-3 .....G.....
B2-4 .....G.....
```

Depending on base qualities and settings used during the assembly like, e.g., [-CO:mr:mrpg:mnq:mgqrt:emea:amgb] MIRA will recognise that there's enough evidence for a T and also enough evidence for a G at that position and create two contigs, one containing the "T" allele, one the "G". The consensus will be >99% identical, but not 100%.

Things become complicated if one has to account for errors in sequencing. Imagine you sequenced the following case:

```
A1-1 .....T.....
A1-2 .....T.....
A1-3 .....T.....
A1-4 .....T.....
A1-5 .....T.....
B2-1 .....G.....
```

It shows very much the same like the one from above, except that there's only one read with a "G" instead of 4 reads. MIRA will, when using standard settings, treat this as erroneous base and leave all these reads in a contig. It will likewise also not mark it as SNP in the results. However, this could also very well be a lowly expressed transcript with a single base mutation. It's virtually impossible to tell which of the possibilities is right.

Note You can of course force MIRA to mark situations like the one depicted above by, e.g., changing the parameters for [-CO:mrpg:mnq:mgqrt]. But this may have the side-effect that sequencing errors get an increased chance of getting flagged as SNP.

Further complications arise when SNPs and potential sequencing errors meet at the same place. consider the following case:

```

A1-1 .....T.....
A1-2 .....T.....
A1-3 .....T.....
A1-4 .....T.....
B1-5 .....T.....
B2-1 .....G.....
B2-2 .....G.....
B2-3 .....G.....
B2-4 .....G.....
E1-1 .....A.....

```

This example is exactly like the first one, except an additional read E1-1 has made its appearance and has an "A" instead of a "G" or "T". Again it is impossible to tell whether this is a sequencing error or a real SNP. MIRA handles these cases in the following way: it will recognise two valid read groups (one having a "T", the other a "G") and, in assembly mode, split these two groups into different contigs. It will also play safe and define that the single read E1-1 will not be attributed to either one of the contigs but, if it cannot be assembled to other reads, form an own contig ... if need to be even only as single read (a *singleton*).

Note Depending on some settings, singletons may either appear in the regular results or end up in the debris file.

6.4.2 Splitting transcripts into contigs based on larger gaps

Gaps in alignments of transcripts are handled very cautiously by MIRA. The standard settings will lead to the creation of different contigs if three or more consecutive gaps are introduced in an alignment. Consider the following example:

```

A1-1 .....CGA.....
A1-2 .....*GA.....
A1-3 .....**A.....
B2-1 .....***.....
B2-2 .....***.....

```

Under normal circumstances, MIRA will use the reads A1-1, A1-2 and A1-3 to form one contig and put B2-1 and B2-2 into a separate contig. MIRA would do this also if there were only one of the B2 reads.

The reason behind this is that the probability for having gaps of three or more bases only due to sequencing errors is pretty low. MIRA will therefore treat reads with such attributes as coming from different transcripts and not assemble them together, though this can be changed using the [-AL:egp:egpl] parameters of MIRA if wanted.



Problems with homopolymers, especially in 454, Ion Torrent and high coverage Illumina

As 454 and Ion Torrent sequencing has a general problem with homopolymers, this rule of MIRA will sometimes lead to formation of more contigs than expected due to sequencing errors at "long" homopolymer sites ... where long starts at ~6-7 bases. Though MIRA does know about the problem in 454 homopolymers and has some routines which try to mitigate the problem. this is not always successful.

The same applies for Illumina data with long homopolymers (~ 8-9 bp) and high coverage ($\geq 100x$).

6.5 A simple step by step pipeline for reliable RNASeq assembly of eukaryotes

1. Remove rRNA sequences. For that I use **mirabait** like this:

```

arcadia:/path/to/myProject$ mirabait -I -j rrna -p norRNAfile_1.fastq norRNAfile_2. ←
fastq ...

```

2. Clean the data. For this I use **mira**, asking it to perform only a preprocessing of the data from step 1 via a line like this

```
parameters = -AS:ppo=yes
```

in the manifest file. After preprocessing, the results will be present as MAF file in the file `*_assembly/*_d_chkpt/readpool.maf`.

3. As the MAF file contains paired reads together, they need to be separated again. Additionally, I perform a hard cut of the clipped sequence. This is a job for **miraconvert**:

```
arcadia:/path/to/myProject$ miraconvert -C -F -F readpool.maf
```

4. I then use FLASH to merge paired read together, using high overlap and zero allowed errors.

```
arcadia:/path/to/myProject$ ...
```

FLASH will create three file for this: one file with joined pairs, one file with unjoined pairs and one file with orphan reads (i.e., reads which have no mate). I generally continue with just the joined and unjoined files.

5. Reduce the dataset to a reasonable size. Using 3 or 4 gigabases to reconstruct an eukaryotic transcriptome should yield in pretty good transcripts without too much noise and loose all but the rarest transcripts.

Depending on the Illumina read length (100, 125, 150, 250 or 300) I generally go for a 1:1 or 2:1 ratio of joined versus unjoined reads. E.g., if I need to extract 2 gigabases of joined FLASH results and 1 gigabase of unjoined FLASH results I do this:

```
arcadia:/path/to/myProject$ miraconvert -Y 2000000 flashjoined.fastq ↔
reduced2gb_flashjoined.fastq
arcadia:/path/to/myProject$ miraconvert -Y 1000000 flashunjoined.fastq ↔
reduced1gb_flashunjoined.fastq
```

6. Assemble the cleaned, joined and reduced data set. A simple manifest file like this will suffice:

```
project = myRNASEQ
job=est,denovo,accurate

readgroup
technology=solexa
autopairing
data=reduced2gb_flashjoined.fastq reduced1gb_flashunjoined.fastq
```

The result can be annotated and quality controlled. However, this will still contain duplicate genes (due to, e.g., ploidy variants) or gene fragments (due to ploidy variants, splice variants, sequencing errors). To reduce this number I generally do the following:

1. Extract CDS of the annotated sequences. make sure that your pipeline also annotates hypothetical proteins with a length ≥ 300 bp.
2. Cluster the CDS sequences with MIRA, using a high similarity threshold:

```
project = myRNASEQclustering
job=est,clustering,accurate
parameters = --noclipping
parameters = TEXT_SETTINGS -AS:mrs=94

readgroup
technology=text
autopairing
data=fna::CDSfromAnnotation.fasta
```

6.6 Solving common problems of EST assemblies

... continue here ...

Megahubs => track down reason (high expr, seqvec or adaptor: see mira_hard) and eliminate it

Chapter 7

Working with the results of MIRA

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“You have to know what you’re looking for before you can find it.”

—Solomon Short

MIRA makes results available in quite a number of formats: CAF, ACE, FASTA and a few others. The preferred formats are CAF and MAF, as these format can be translated into any other supported format.

7.1 MIRA output directories and files

For the assembly MIRA creates a directory named `projectname_assembly` in which a number of sub-directories will have appeared.

These sub-directories (and files within) contain the results of the assembly itself, general information and statistics on the results and -- if not deleted automatically by MIRA -- a `tmp` directory with log files and temporary data:

- `projectname_d_results`: this directory contains all the output files of the assembly in different formats.
- `projectname_d_info`: this directory contains information files of the final assembly. They provide statistics as well as, e.g., information (easily parsable by scripts) on which read is found in which contig etc.
- `projectname_d_tmp`: this directory contains log files and temporary assembly files. It can be safely removed after an assembly as there may be easily a few GB of data in there that are not normally not needed anymore.

The default settings of MIRA are such that really big files are automatically deleted when they not needed anymore during an assembly.

- `projectname_d_chkpt`: this directory contains checkpoint files needed to resume assemblies that crashed or were stopped.

7.1.1 The `*_d_results` directory

The following files in `projectname_d_results` contain results of the assembly in different formats. Depending on the output options you defined for MIRA, some files may or may not be there. As long as the CAF or MAF format are present, you can translate your assembly later on to about any supported format with the `miraconvert` program supplied with the MIRA distribution:

- `projectname_out.txt`: this file contains in a human readable format the aligned assembly results, where all input sequences are shown in the context of the contig they were assembled into. This file is just meant as a quick way for people to have a look at their assembly without specialised alignment finishing tools.

- `projectname_out.padded.fasta`: this file contains as FASTA sequence the consensus of the contigs that were assembled in the process. Positions in the consensus containing gaps (also called 'pads', denoted by an asterisk) are still present. The computed consensus qualities are in the corresponding `projectname_out.padded.fasta.qual` file.
- `projectname_out.unpadded.fasta`: as above, this file contains as FASTA sequence the consensus of the contigs that were assembled in the process, but positions in the consensus containing gaps were removed. The computed consensus qualities are in the corresponding `projectname_out.unpadded.fasta.qual` file.
- `projectname_out.caf`: this is the result of the assembly in CAF format, which can be further worked on with, e.g., tools from the *caftools* package from the Sanger Centre and later on be imported into, e.g., the Staden gap4 assembly and finishing tool.
- `projectname_out.ace`: this is the result of the assembly in ACE format. This format can be read by viewers like the TIGR clview or by consed from the phred/phrap/consed package.
- `projectname_out.gap4da`: this directory contains the result of the assembly suited for the *direct assembly* import of the Staden gap4 assembly viewer and finishing tool.

7.1.1.1 Additional 'large contigs' result files for de-novo assemblies of genomes

For de-novo assemblies of genomes, MIRA makes a proposal regarding which contigs you probably want to have a look at ... and which ones you can probably forget about.

This proposal relies on the *largecontigs* file in the info directory (see section below) and MIRA automatically extracted these contigs into all the formats you wanted to have your results in.

- The result files for 'large contigs' are all named: `projectname_LargeContigs_out.resulttype`:
- `extractLargeContigs.sh`: this is a small shell script which just contains the call to **miracconvert** with which MIRA extracted the large contigs for you. In case you want to redefine what large contigs are for you, feel free to use this as template.

7.1.2 The *_d_info directory

The following files in `projectname_info` contain statistics and other information files of the assembly:

- `projectname_info_assembly.txt`: This file should be your first stop after an assembly. It will tell you some statistics as well as whether or not problematic areas remain in the result.
- `projectname_info_callparameters.txt`: This file contains the parameters as given on the mira command line when the assembly was started.
- `projectname_info_consensustaglist.txt`: This file contains information about the tags (and their position) that are present in the consensus of a contig.
- `projectname_info_contigreadlist.txt`: This file contains information which reads have been assembled into which contigs (or singlets).
- `projectname_info_contigstats.txt`: This file contains in tabular format statistics about the contigs themselves, their length, average consensus quality, number of reads, maximum and average coverage, average read length, number of A, C, G, T, N, X and gaps in consensus.
- `projectname_info_debrislist.txt`: This file contains the names of all the reads which were not assembled into contigs (or singlets if appropriate MIRA parameters were chosen). The file has two columns: first column is the name of the read, second column is a code showing the reason and stage at which the read was put into the debris category.
- `projectname_info_largecontigs.txt`: This file contains as simple list the names of all the contigs MIRA thinks to be more or less important at the end of the assembly. To be present in this list, a contig needed to reach a certain length (usually 500, but see [-MI:lcs]) and had a coverage of at least 1/3 of the average coverage (per sequencing technology) of the complete project.

Note: only present for de-novo assemblies of genomes.



Warning The default heuristics (500bp length and 1/3 coverage per sequencing technology) generally work well enough for most projects. However, Projects with extremely different coverage numbers per sequencing technology may need to use different numbers. E.g.: a project with 80x Illumina and 6x Sanger would have contigs consisting only of 2 or 3 Sanger sequence but with the average coverage ≥ 2 also in this list although clearly no one would look at these under normal circumstances.

- `projectname_info_groups.txt`: This file contains information about readgroups as determined by MIRA. Most interesting will probably be statistics concerning read-pair sizes.
- `projectname_info_readrepeats`: This file helps to find out which parts of which reads are quite repetitive in a project. Please consult the chapter on how to tackle "hard" sequencing projects to learn how this file can help you in spotting sequencing mistakes and / or difficult parts in a genome or EST / RNASeq project.
- `projectname_info_readstooshort`: A list containing the names of those reads that have been sorted out of the assembly only due to the fact that they were too short, before any processing started.
- `projectname_info_readtaglist.txt`: This file contains information about the tags and their position that are present in each read. The read positions are given relative to the forward direction of the sequence (i.e. as it was entered into the the assembly).
- `projectname_error_reads_invalid`: A list of sequences that have been found to be invalid due to various reasons (given in the output of the assembler).

7.2 First look: the assembly info

Once finished, have a look at the file `*_info_assembly.txt` in the info directory. The assembly information given there is split in three major parts:

1. some general assembly information (number of reads assembled etc.). This part is quite short at the moment, will be expanded in future
2. assembly metrics for 'large' contigs.
3. assembly metrics for all contigs.

The first part for large contigs contains several sections. The first of these shows what MIRA counts as large contig for this particular project. As example, this may look like this:

```
Large contigs:
-----
With      Contig size          >= 500
      AND (Total avg. Cov  >= 19
            OR Cov(san)    >= 0
            OR Cov(454)   >= 8
            OR Cov(pbs)   >= 0
            OR Cov(sxa)   >= 11
            OR Cov(sid)   >= 0
      )
```

The above is for a 454 and Solexa hybrid assembly in which MIRA determined large contigs to be contigs

1. of length of at least 500 bp and
2. having a total average coverage of at least 19x or an average 454 coverage of 8 or an average Solexa coverage of 11

The second section is about length assessment of large contigs:

```

Length assessment:
-----
Number of contigs:      44
Total consensus:      3567224
Largest contig:       404449
N50 contig size:     186785
N90 contig size:     55780
N95 contig size:     34578

```

In the above example, 44 contigs totalling 3.56 megabases were built, the largest contig being 404 kilobases long and the N50/N90 and N95 numbers give the respective lengths.

The next section shows information about the coverage assessment of large contigs. An example:

```

Coverage assessment:
-----
Max coverage (total): 563
Max coverage
  Sanger: 0
  454: 271
  PacBio: 0
  Solexa: 360
  Solid: 0
Avg. total coverage (size >= 5000): 57.38
Avg. coverage (contig size >= 5000)
  Sanger: 0.00
  454: 25.10
  PacBio: 0.00
  Solexa: 32.88
  Solid: 0.00

```

Maximum coverage attained was 563, maximum for 454 alone 271 and for Solexa alone 360. The average total coverage (computed from contigs with a size ≥ 5000 bases is 57.38. The average coverage by sequencing technology (in contigs ≥ 5000) is 25.10 for 454 and 32.88 for Solexa reads.

Note

For genome assemblies, the value for *Avg. total coverage (size ≥ 5000)* is currently always calculated for contigs having 5000 or more consensus bases. While this gives a very effective measure for genome assemblies, assemblies of EST or RNASeq will often have totally irrelevant values here: even if the default of MIRA is to use smaller contig sizes (1000) for EST / RNASeq assemblies, the coverage values for lowly and highly expressed genes can easily span a factor of 10000 or more.

The last section contains some numbers useful for quality assessment. It looks like this:

```

Quality assessment:
-----
Average consensus quality:          90
Consensus bases with IUPAC:        11      (you might want to check these)
Strong unresolved repeat positions (SRMc): 0      (excellent)
Weak unresolved repeat positions (WRMc): 19      (you might want to check these)
Sequencing Type Mismatch Unsolved (STMU): 0      (excellent)
Contigs having only reads wo qual:  0      (excellent)
Contigs with reads wo qual values:  0      (excellent)

```

Beside the average quality of the contigs and whether they contain reads without quality values, MIRA shows the number of different tags in the consensus which might point at problems.

The above mentioned sections (length assessment, coverage assessment and quality assessment) for *large* contigs will then be re-iterated for *all* contigs, this time including also contigs which MIRA did not take into account as large contig.

7.3 Converting results

7.3.1 Converting to and from other formats:miraconvert

miraconvert is tool in the MIRA package which reads and writes a number of formats, ranging from full assembly formats like CAF and MAF to simple output view formats like HTML or plain text.

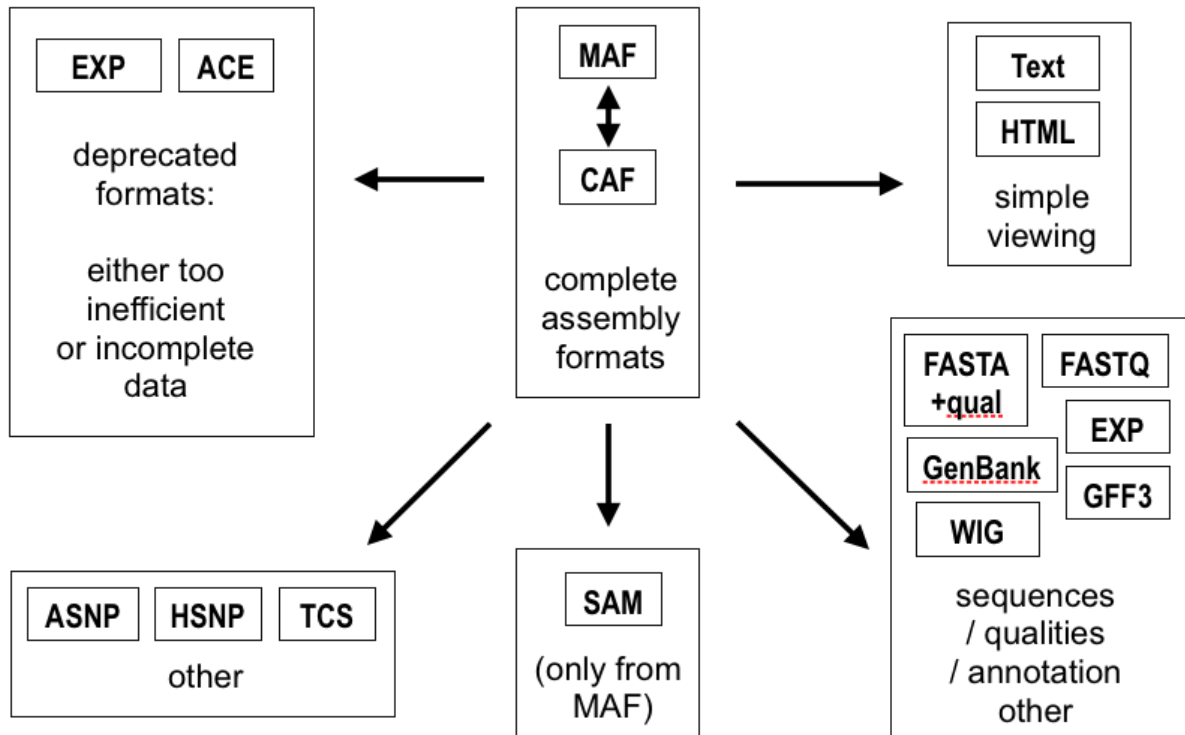


Figure 7.1: **miraconvert** supports a wide range of format conversions to simplify export / import of results to and from other programs

7.3.2 Steps for converting data from / to other tools

The question "How Do I convert to / from other tools?" is complicated by the plethora of file formats and tools available. This section gives an overview on what is needed to reach the most important ones.

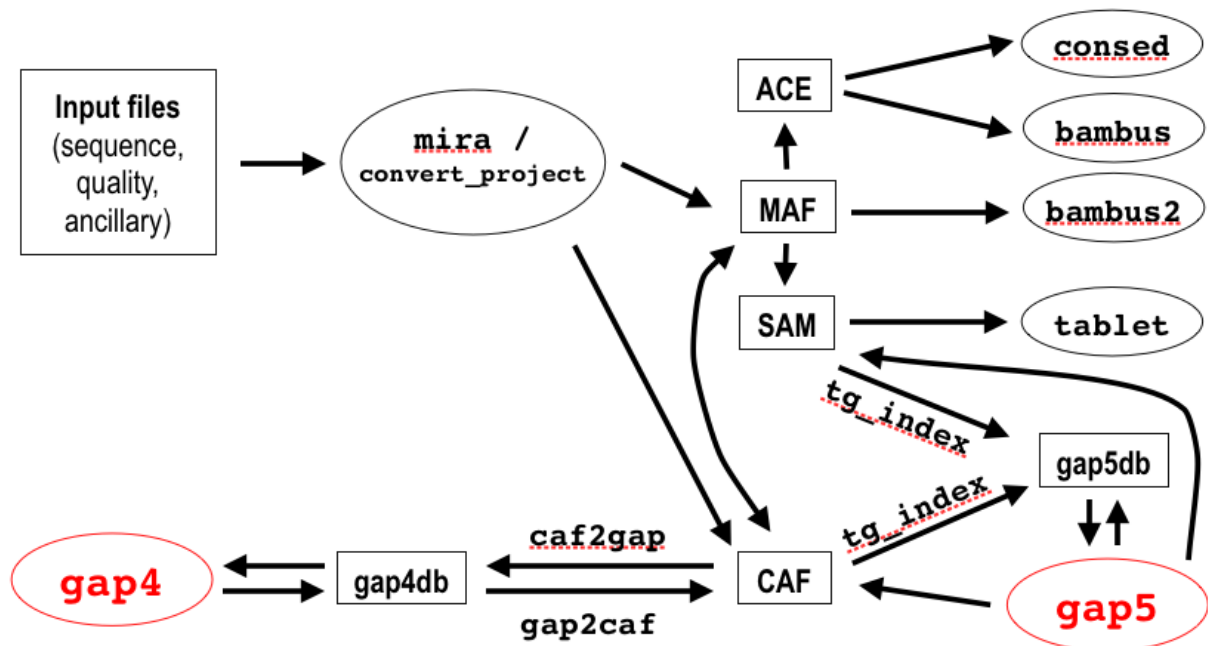


Figure 7.2: Conversion steps, formats and programs needed to reach some tools like assembly viewers, editors or scaffolders.

Please also read the chapter on MIRA utilities in this manual to learn more on **miraconvert** and have a look at `miraconvert -h` which lists all possible formats and other command line options.

7.3.2.1 Example: converting to and from the Staden package (gap4 / gap5)

The **gap4** program (and its successor **gap5** from the Staden package) are pretty useful finishing tools and assembly viewers. They have an own database format which MIRA does not read or write, but there are interconversion possibilities using the CAF format (for gap4) and SAM format (for gap5)

[gap4] You need the **caf2gap** and **gap2caf** utilities for this, which are distributed separately from the Sanger Centre (<http://www.sanger.ac.uk/Software/formats/CAF/>). Conversion is pretty straightforward. From MIRA to gap4, it's like this:

```
$ caf2gap -project YOURGAP4PROJECTNAME -ace mira_result.caf >&/dev/null
```

Note Don't be fooled by the `-ace` parameter of **caf2gap**. It needs a CAF file as input, not an ACE file.

From gap4 to CAF, it's like this:

```
$ gap2caf -project YOURGAP4PROJECTNAME >tmp.caf
$ miraconvert -r c tmp.caf somenewname.caf
```

Note Using **gap2caf**, be careful to use the simple `>` redirection to file and *not* the `>&` redirection.

Note Using first **gap2caf** and then **miraconvert** is needed as gap4 writes an own consensus to the CAF file which is not necessarily the best. Indeed, gap4 does not know about different sequencing technologies like 454 and treats everything as Sanger. Therefore, using **miraconvert** with the `[-r c]` option recalculates a MIRA consensus during the "conversion" from CAF to CAF.

Note If you work with a 32 bit executable of `caf2gap`, it might very well be that the converter needs more memory than can be handled by 32 bit. Only solution: switch to a 64 bit executable of `caf2gap`.



caf2gap bug for sequence annotations in reverse direction

`caf2gap` has currently (as of version 2.0.2) a bug that turns around all features in reverse direction during the conversion from CAF to a `gap4` project. There is a fix available, please contact me for further information (until I find time to describe it here).

[gap5] The `gap5` program is the successor for `gap4`. It comes with on own import utility (`tg_index`) which can import SAM and CAF files, and `gap5` itself has an export function which also writes SAM and CAF. It is suggested to use the SAM format to export data `gap5` as it is more efficient and conveys more information on sequencing technologies used.

Conversion is pretty straightforward. From MIRA to `gap5`, it's like this:

```
$ tg_index INPUT_out.sam
```

This creates a `gap5` database named `INPUT_out.g5d` which can be directly loaded with `gap5` like this:

```
$ gap5 INPUT_out.g5d
```

Exporting back to SAM or CAF is done in `gap5` via the *File->Export Sequences* menu there.

7.3.2.2 Example: converting to and from SAM (for samtools, tablet etc.)

Converting to SAM is done by using `miraconvert` on a MIRA MAF file, like this:

```
$ miraconvert maf -t sam INPUT.maf OUTPUT
```

The above will create a file named `OUTPUT.sam`.

Converting from SAM to a format which either `mira` or `miraconvert` can understand takes a few more steps. As neither tool currently reads SAM natively, you need to go via the `gap5` editor of the Staden package: convert the SAM via `tg_index` to a `gap5` database, load that database in `gap5` and export it there to CAF.

7.4 Filtering results

It is important to remember that, depending on assembly options, MIRA will also include very small contigs (with eventually very low coverage) made out of reads which were rejected from the "good" contigs for quality or other reasons. You probably do not want to have a look at this contig debris when finishing a genome unless you are really, really, really picky.

Many people prefer to just go on with what would be large contigs. Therefore, in de-novo assemblies, MIRA writes out separate files of what it thinks are "good", large contigs. In case you want to extract contigs differently, the `miraconvert` program from the MIRA package can selectively filter CAF or MAF files for contigs with a certain size, average coverage or number of reads.

The file `*_info_assembly.txt` in the info directory at the end of an assembly might give you first hints on what could be suitable filter parameters. As example, for "normal" assemblies (whatever this means), one could want to consider only contigs larger than 500 bases and which have at least one third of the average coverage of the N50 contigs.

Here's an example: In the "Large contigs" section, there's a "Coverage assessment" subsection. It looks a bit like this:

```
...
Coverage assessment:
-----
Max coverage (total): 43
Max coverage
Sanger: 0
454: 43
```

```

Solexa: 0
Solid: 0
Avg. total coverage (size ≥ 5000): 22.30
Avg. coverage (contig size ≥ 5000)
Sanger: 0.00
454: 22.05
Solexa: 0.00
Solid: 0.00
...

```

This project was obviously a 454 only project, and the average coverage for it is ~22. This number was estimated by MIRA by taking only contigs of at least 5kb into account, which for sure left out everything which could be categorised as debris. Normally it's a pretty solid number.

Now, depending on how much time you want to invest performing some manual polishing, you should extract contigs which have at least the following fraction of the average coverage:

- 2/3 if a quick and "good enough" is what you want and you don't want to do some manual polishing. In this example, that would be around 14 or 15.
- 1/2 if you want to have a "quick look" and eventually perform some contig joins. In this example the number would be 11.
- 1/3 if you want quite accurate and for sure not loose any possible repeat. That would be 7 or 8 in this example.

Example (useful with assemblies of Sanger data): extracting only contigs ≥ 1000 bases and with a minimum average coverage of 4 into FASTA format:

```
$ miraconvert -x 1000 -y 4 sourcefile.maf targetfile.fasta
```

Example (useful with assemblies of 454 data): extracting only contigs ≥ 500 bases into FASTA format:

```
$ miraconvert -x 500 sourcefile.maf targetfile.fasta
```

Example (e.g. useful with Sanger/454 hybrid assemblies): extracting only contigs ≥ 500 bases and with an average coverage ≥ 15 reads into CAF format, then converting the reduced CAF into a Staden GAP4 project:

```
$ miraconvert -x 500 -y 15 sourcefile.maf tmp.caf
$ caf2gap -project somename -ace tmp.caf
```

Example (e.g. useful with Sanger/454 hybrid assemblies): extracting only contigs ≥ 1000 bases and with ≥ 10 reads from MAF into CAF format, then converting the reduced CAF into a Staden GAP4 project:

```
$ miraconvert -x 500 -z 10 sourcefile.maf tmp.caf
$ caf2gap -project somename -ace tmp.caf
```

7.5 Places of importance in a de-novo assembly

7.5.1 Tags set by MIRA

MIRA sets a number of different tags in resulting assemblies. They can be set in reads (in which case they mostly end with a *r*) or in the consensus.(then ending with a *c*).

Note

If you use the Staden **gap4**, **gap5** or **consed** assembly editor to tidy up the assembly, you can directly jump to places of interest that MIRA marked for further analysis by using the search functionality of these programs.

However, you need to tell these programs that these tags exist. For that you must change some configuration files. More information on how to do this can be found in the `support/README` file of the MIRA distribution.

You should search for the following "consensus" tags for finding places of importance (in this order).

- IUPc
- UNSc
- SRMc
- WRMc
- STMU (only hybrid assemblies)
- MCVc (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- SROc (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- SAOc (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- SIOc (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- STMS (only hybrid assemblies)

of lesser importance are the "read" versions of the tags above:

- UNSr
- SRMr
- WRMr
- SROr (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- SAOr (only when assembling different strains, i.e., mostly relevant for mapping assemblies)
- SIOr (only when assembling different strains, i.e., mostly relevant for mapping assemblies)

In normal assemblies (only one sequencing technology, just one strain), search for the IUPc, UNSc, SRMc and WRMc tags.

In hybrid assemblies, searching for the IUPc, UNSc, SRMc, WRMc, and STMU tags and correcting only those places will allow you to have a qualitatively good assembly in no time at all.

Columns with SRMr tags (SRM in **R**eads) in an assembly without a SRMc tag at the same consensus position show where mira was able to resolve a repeat during the different passes of the assembly ... you don't need to look at these. SRMc and WRMc tags however mean that there may be unresolved trouble ahead, you should take a look at these.

Especially in mapping assemblies, columns with the MCVc, SROx, SIOx and SAOx tags are extremely helpful in finding places of interest. As they are only set if you gave strain information to MIRA, you should always do that.

For more information on tags set/used by MIRA and what they exactly mean, please look up the according section in the reference chapter.

7.5.2 Other places of importance

The read coverage histogram as well as the template display of gap4 will help you to spot other places of potential interest. Please consult the gap4 documentation.

7.5.3 Joining contigs

I recommend to invest a couple of minutes (in the best case) to a few hours in joining contigs, especially if the uniform read distribution option of MIRA was used (but first filter for large contigs). This way, you will reduce the number of "false repeats" in improve the overall quality of your assembly.

7.5.3.1 Joining contigs at true repetitive sites

Joining contigs at repetitive sites of a genome is always a difficult decision. There are, however, two rules which can help:

1. If the sequencing was done without a paired-end library, don't join.
2. If the sequencing was done with a paired-end library, but no pair (or template) span the join site, don't join.

The following screen shot shows a case where one should not join as the finishing program (in this case **gap4**) warns that no template (read-pair) span the join site:

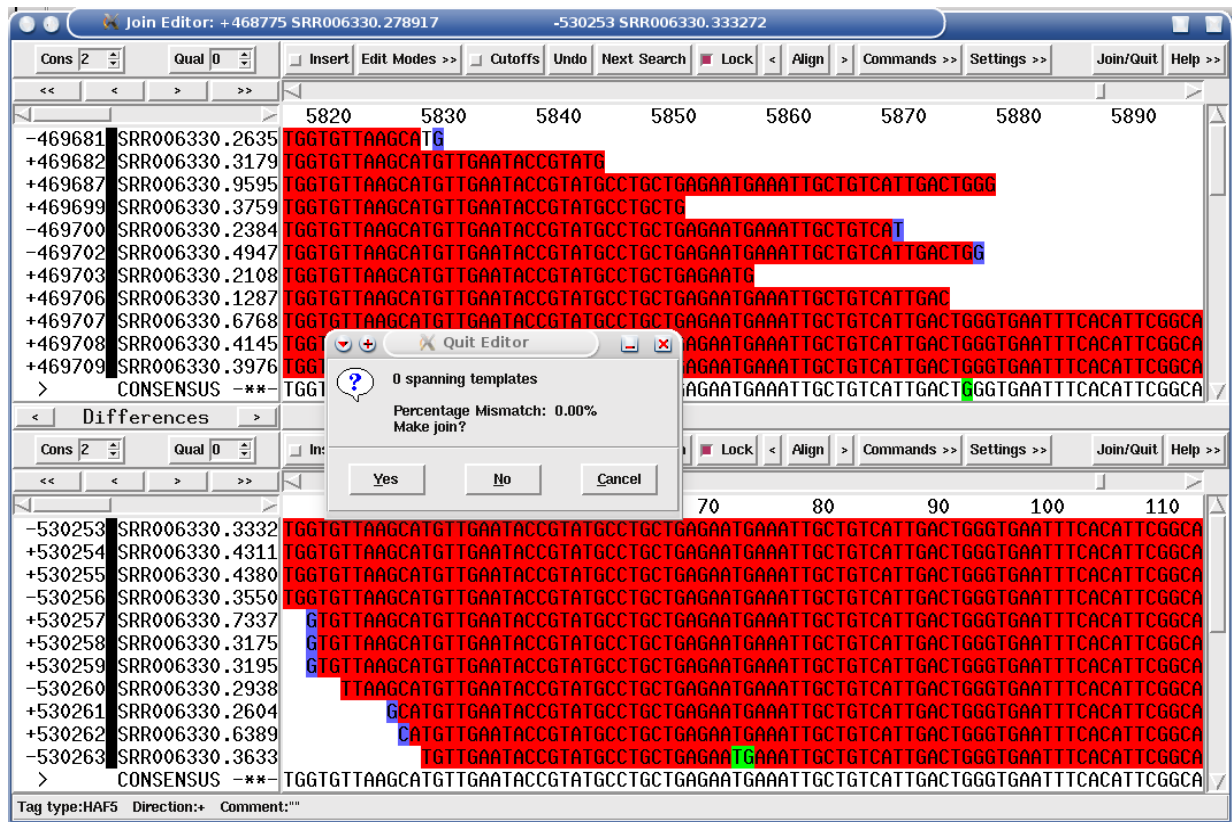


Figure 7.3: Join at a repetitive site which should not be performed due to missing spanning templates.

The next screen shot shows a case where one should join as the finishing program (in this case **gap4**) finds templates spanning the join site and all of them are good:

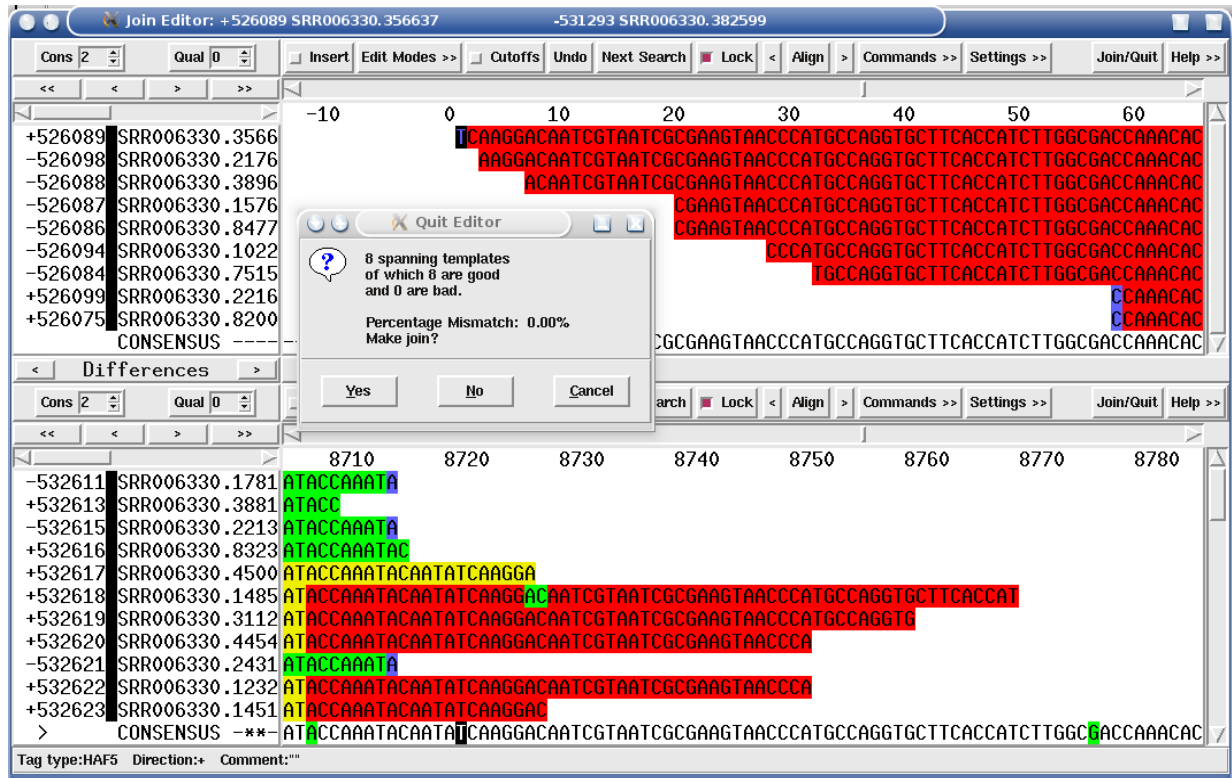


Figure 7.4: Join at a repetitive site which should be performed due to spanning templates being good.

7.5.3.2 Joining contigs at "wrongly discovered" repetitive sites

Remember that MIRA takes a very cautious approach in contig building, and sometimes creates two contigs when it could have created one. Three main reasons can be the cause for this:

1. when using *uniform read distribution*, some non-repetitive areas may have generated so many more reads that they start to look like repeats (so called pseudo-repeats). In this case, reads that are above a given coverage are *shaved off* (see [-AS:urdcn]) and kept in reserve to be used for another copy of that repeat ... which in case of a non-repetitive region will of course never arrive. So at the end of an assembly, these shaved-off reads will form short, low coverage contig debris which can more or less be safely ignored and sorted out via the filtering options ([-x -y -z]) of **miracore**.

Some 454 library construction protocols -- especially, but not exclusively, for paired-end reads -- create pseudo-repeats quite frequently. In this case, the pseudo-repeats are characterised by several reads starting at exact the same position but which can have different lengths. Should MIRA have separated these reads into different contigs, these can be -- most of the time -- safely joined. The following figure shows such a case:

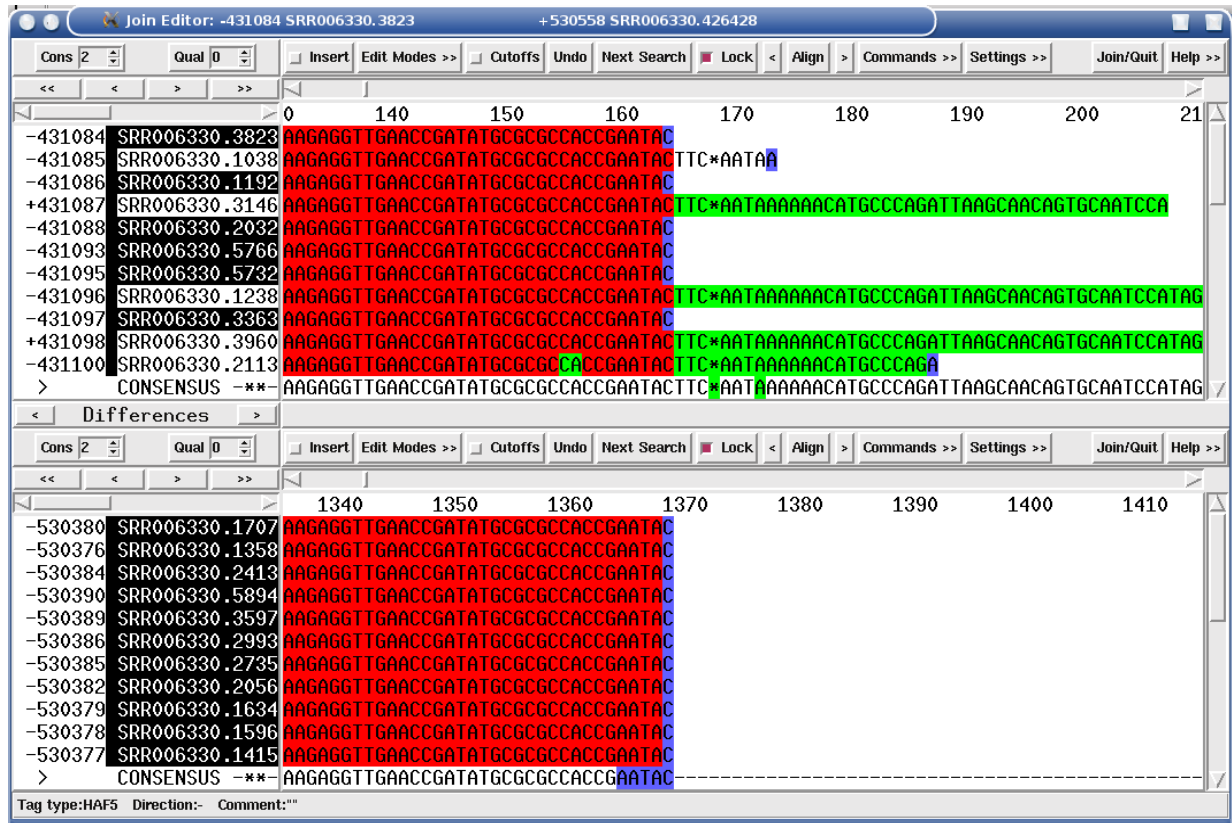


Figure 7.5: Pseudo-repeat in 454 data due to sequencing artifacts

For Solexa data, a non-negligible GC bias has been reported in genome assemblies since late 2009. In genomes with moderate to high GC, this bias actually favours regions with lower GC. Examples were observed where regions with an average GC of 10% less than the rest of the genome had between two and four times more reads than the rest of the genome, leading to false "discovery" of duplicated genome regions.

2. when using unpaired data, the above described possibility of having "too many" reads in a non-repetitive region can also lead to a contig being separated into two contigs in the region of the pseudo-repeat.
3. a number of reads (sometimes even just one) can contain "high quality garbage", that is, nonsense bases which got - for some reason or another - good quality values. This garbage can be distributed on a long stretch in a single read or concern just a single base position across several reads.

While MIRA has some algorithms to deal with the disrupting effects of reads like, the algorithms are not always 100% effective and some might slip through the filters.

7.6 Places of interest in a mapping assembly

This section just give a short overview on the tags you might find interesting. For more information, especially on how to configure gap4 or consed, please consult the *mira usage* document and the *mira* manual.

In file types that allow tags (CAF, MAF, ACE), SNPs and other interesting features will be marked by MIRA with a number of tags. The following sections give a brief overview. For a description of what the tags are (SROc, WRMc etc.), please read up the section "Tags used in the assembly by MIRA and EdIt" in the main manual.

Note Screen shots in this section are taken from the walk-through with Lenski data (see below).

7.6.1 Where are SNPs?

- the **SROc** tag will point to most SNPs. Should you assemble sequences of more than one strain (I cannot really recommend such a strategy), you also might encounter **SIOc** and **SAOc** tags.

```

Contig Editor: -1 SRR030257.787415/2
Cons 2 Qual 0 Insert Edit Modes Cutoffs Undo Next Search Commands Settings Quit Help
161080 161090 161100 161110 161120 161130 161140 161150
+17 NC_012967 gggTaccagaacatggcggcaaacaggaacgCGGGTcAcgcgcataTcgTtatggatacgcgtatcggTattcttCag
+18 _cer_sxa_0_ gggTaccagaacatggcggcaaacaggaacgCGGGT
+19 _cer_sxa_60_ gggTaccagaacatggcggcaaacaggaacgCGGG
+20 _cer_sxa_123_ gggTaccagaacatggcggcaaacaggaacgCGGG
+3663 _cer_sxa_190_ gggTaccagaacatggcggcaaacaggaacgCC
+3664 _cer_sxa_260_ gggTaccagaacatggcggcaaacaggaacgCC
-33401 SRR030257.8883 ACCAGAACATGGCGGCAAACAGGAACGCCGGGTGCA
-33402 SRR030257.2989 CCAGAACATGGCGGCAAACAGGAACGCCGGGTGCAC
+33403 SRR030257.1128 CAGAACATGGCGGCAAACAGGAACGCCGGGTGCACG
+33404 SRR030257.7073 AGAACATGGCGGCAAACAGGAACGCCGGGTGCACGC
-33405 SRR030257.1204 AGAACATGGCGGCAAACAGGAACGCCGGGTGCACGC
-33406 SRR030257.2602 GAACATGGCGGCAAACAGGAACGCCGGGTGCACGCG
-33407 SRR030257.2767 GAACATGGCGGCAAACAGGAACGCCGGGTGCACGCG
-33408 SRR030257.1413 AACATGGCGGCAAACAGGAACGCCGGGTGCACGCGC
+33409 SRR030257.1755 ACATGGCGGCAAACAGGAACGCCGGGTGCACGCGCA
+33410 SRR030257.1463 ACATGGCGGCAAACAGGAACGCCGGGTGCACGCGCA
+33411 SRR030257.1623 CATGGCGGCAAACAGGAACGCCGGGTGCACGCGCAT
+33412 SRR030257.6602 TGGCGGCAAACAGGAACGCCGGTGG
-33413 SRR030257.1821 GGGCGGCAAACAGGAACGCCGGGTGCACGC
-33414 SRR030257.2932 GGGCGGCAAACAGGAACGCCGGG
+33415 SRR030257.5986 GGGCGGCAAACAGGAACGCCGGTGG
-33416 SRR030257.3729 GGGCGGCAAACAGGAACGCCGGTGCACGCGCATATC
+33417 SRR030257.3423 GCGGCAAACAGGAACGCCGGTGCACGCGCATATCG
+33418 SRR030257.2482 GCGGCAAACAGGAACGCCGGTGCACGCGCATATCG
-33419 SRR030257.1401 CGGCAAACAGGAACGCCGGTGCACGCGCATATCG
-33420 SRR030257.3565 CGGCAAACAGGAACGCCGGTGCACGCGCATATCGT
+33421 SRR030257.1125 CGGCAAACAGGAACGCCGGTGCACGCGCATATCGT
-33422 SRR030257.3529 GGC AACAGGAACGCCGGTGCACGCGCATATCGTT
> CONSENSUS -*- GGGTACCAGAACATGGCGGCAAACAGGAACGCCGGTGCACGCGCATATCGTTATGGATACGCGTATCGGTATTCTTCAG /
Tag type: SVEC Direction: - Comment: ''

```

Figure 7.6: "SROc" tag showing a SNP position in a Solexa mapping assembly.

The screenshot shows the 'Contig Editor' window for file '-1_SRR030257.787415/2'. The interface includes a menu bar with options like 'Cons', 'Qual', 'Insert', 'Edit Modes', 'Cutoffs', 'Undo', 'Next Search', 'Commands', 'Settings', 'Quit', and 'Help'. Below the menu is a navigation bar with arrows and a position indicator showing '0 475400 475410 475420 475430 475440 475450 475460 47547'. The main area displays a sequence alignment with a vertical blue highlight at position 475430. The alignment shows a consensus sequence at the bottom: 'CONSENSUS -**-- AAAGGCGAGCACAAGGCCCAACAATGGTGGTGATAAGCGGGGGTGGCGTGATGCATTCCGTCCTTTTCTGGTGGT'. The tag type is 'Fgen', direction is '+', and the comment is '/gene=ybaL :: locus_tag=ECB_00429'.

Figure 7.7: "SROc" tag showing a SNP/indel position in a Solexa mapping assembly.

- the **WRMc** tags might sometimes point SNPs to indels of one or two bases.

7.6.2 Where are insertions, deletions or genome re-arrangements?

- Large deletions: the **MCVc** tags point to deletions in the resequenced data, where no read is covering the reference genome.

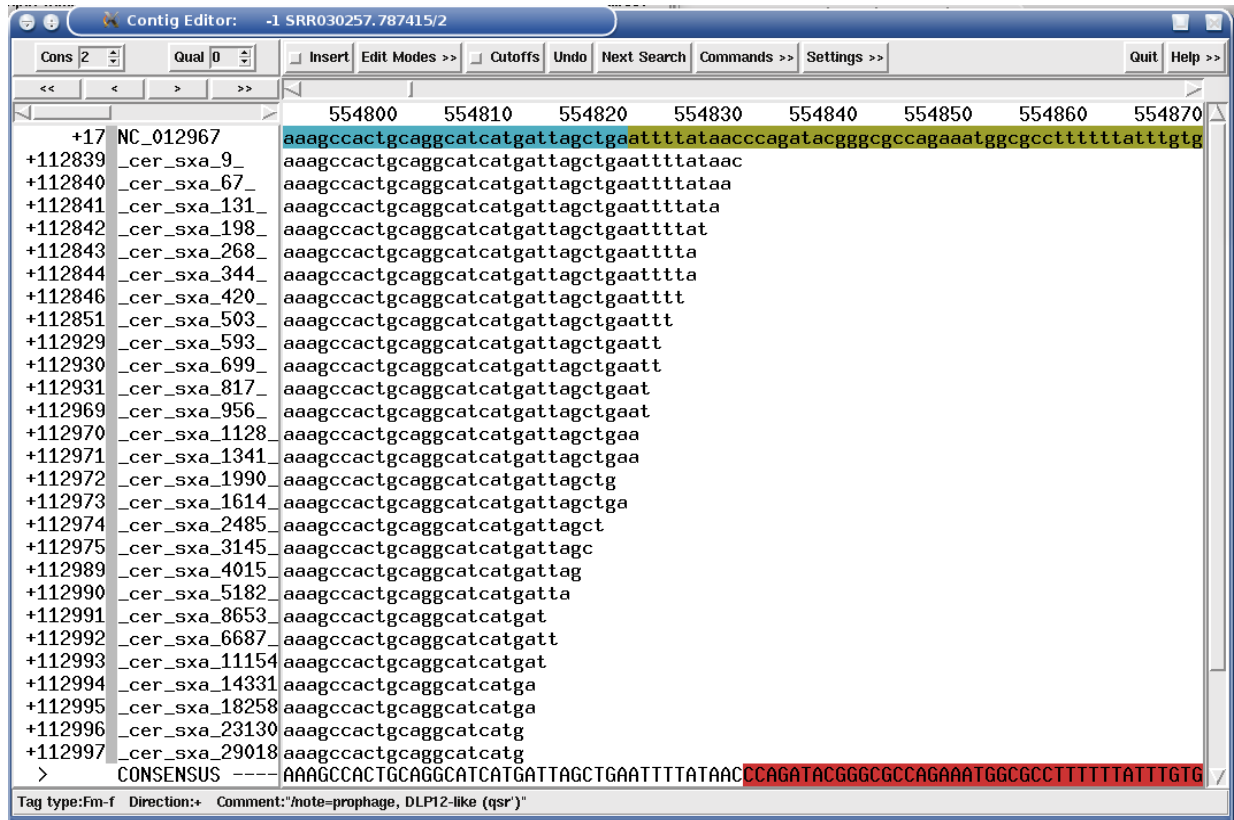


Figure 7.8: "MCVc" tag (dark red stretch in figure) showing a genome deletion in Solexa mapping assembly.

- Insertions, small deletions and re-arrangements: these are harder to spot. In unpaired data sets they can be found looking at clusters of **SROc**, **SRMc**, **WRMc**, and / or **UNSc** tags.

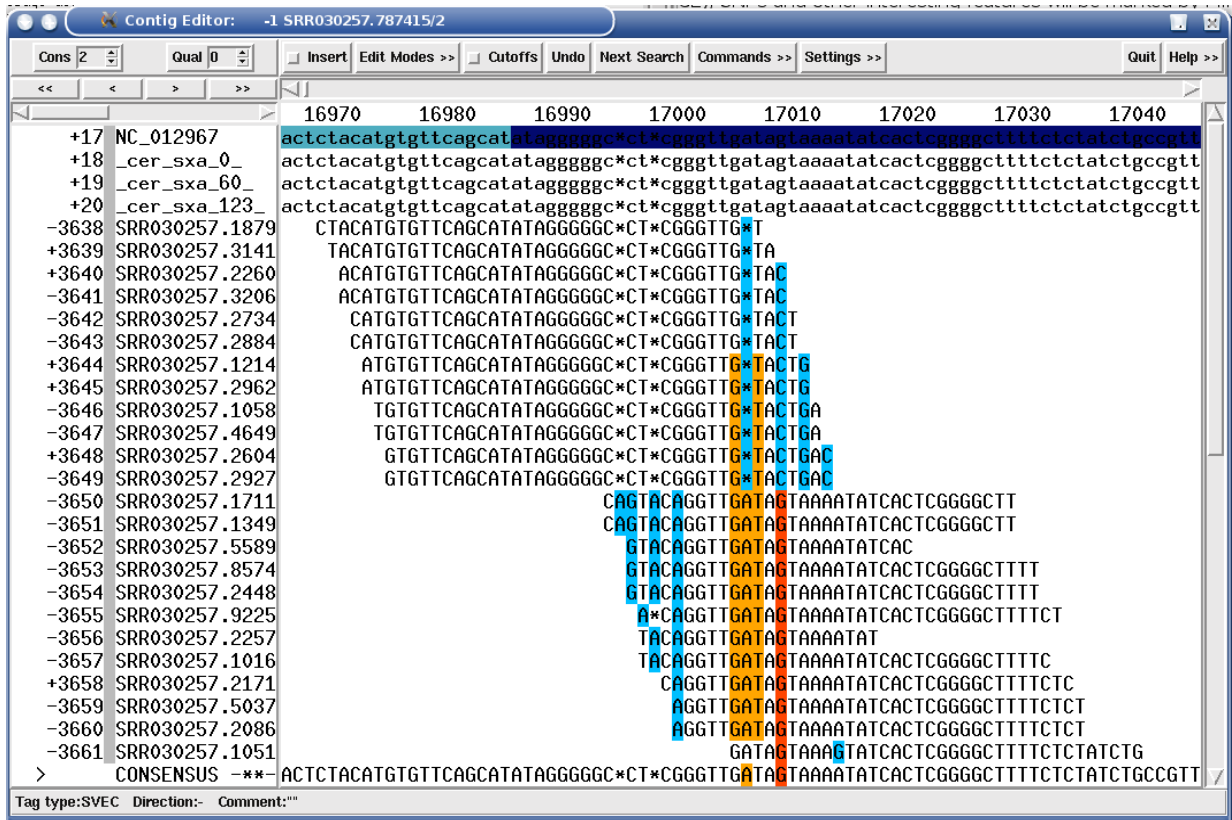


Figure 7.9: An IS150 insertion hiding behind a WRMc and a SRMc tags

more massive occurrences of these tags lead to a rather colourful display in finishing programs, which is why these clusters are also sometimes called Xmas-trees.

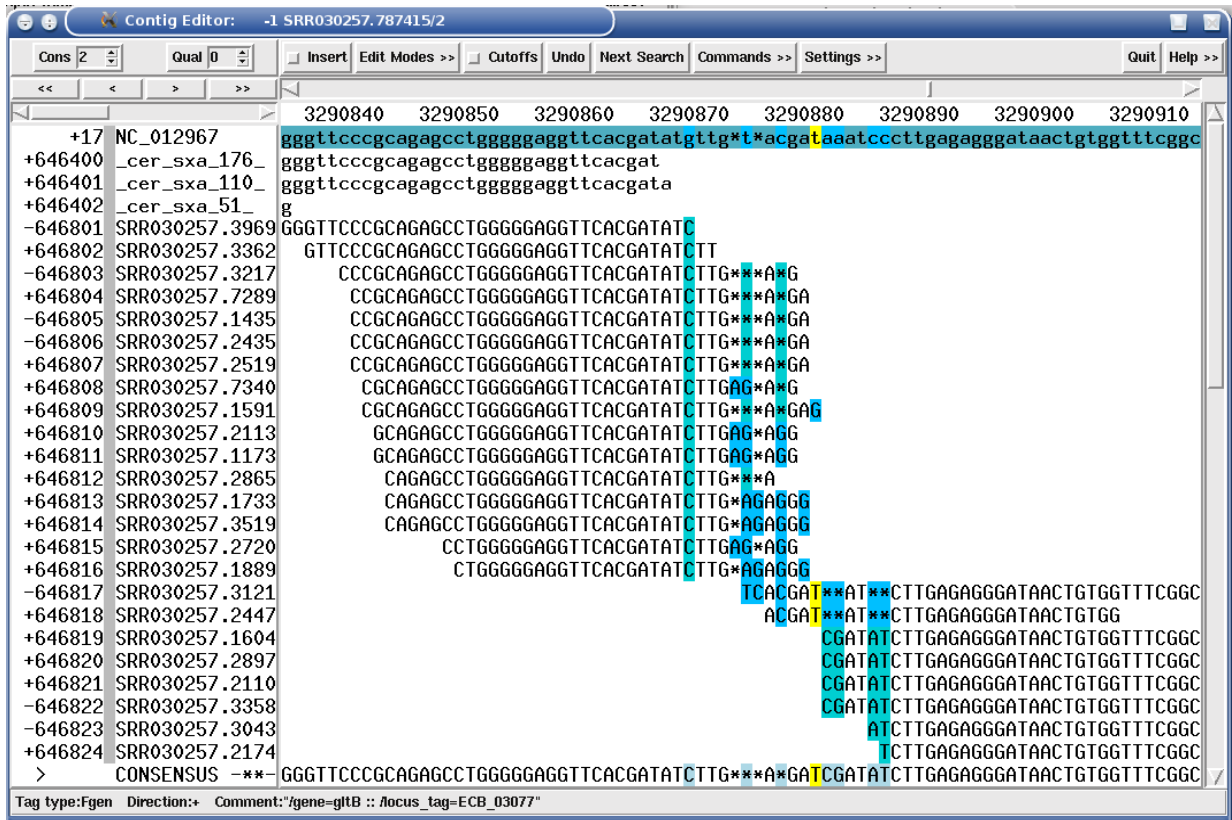


Figure 7.10: A 16 base pair deletion leading to a SROc/UNsC xmas-tree

The screenshot shows the Contig Editor interface with a sequence alignment. The top bar indicates the file path: -1_SRR030257.787415/2. The main window displays a sequence alignment with columns for coordinates (0 to 452588) and rows for individual reads and a consensus sequence. The reads are labeled with IDs like NC_012967, _cer_sxa_59_, etc. The consensus sequence is shown at the bottom, with a tag type of Fgen and a comment: "gene=fimA :: locus_tag=ECB_04183". The sequence contains several tags: SROc (orange) and UNSc (yellow) tags are present in the consensus sequence, indicating problematic regions. The reads are color-coded to match the consensus sequence, with some bases highlighted in blue, orange, or yellow.

Figure 7.11: An IS186 insertion leading to a SROc/UNSc xmas-tree

In sets with paired-end data, post-processing software (or alignment viewers) can use the read-pair information to guide you to these sites (MIRA doesn't set tags at the moment).

7.6.3 Other tags of interest

- the **UNSc** tag points to areas where the consensus algorithm had troubles choosing a base. This happens in low coverage areas, at places of insertions (compared to the reference genome) or sometimes also in places where repeats with a few bases difference are present. Often enough, these tags are in areas with problematic sequences for the Solexa sequencing technology like, e.g., a GGCxG or even GGC motif in the reads.
- the **SRMc** tag points to places where repeats with a few bases difference are present. Here too, sequence problematic for the Solexa technology are likely to have cause base calling errors and subsequently setting of this tag.

7.7 Post-processing mapping assemblies

This section is a bit terse, you should also read the chapter on *working with results* of MIRA.

7.7.1 Manual cleanup and validation (optional)

When working with resequencing data and a mapping assembly, I always load finished projects into an assembly editor and perform a quick cleanup of the results. SNP or small indels normally do not need cleanups, but every mapper will get larger indels mostly wrong, and MIRA is no exception to this.

For close relatives of the reference strain this doesn't take long as MIRA will have set tags (see section earlier in this document) at all sites you should have a look at. For example, very close mutant bacteria with just SNPs or simple deletions and no genome

reorganisation, I usually clean up in 10 to 15 minutes. That gives the last boost to data quality and your users (biologists etc.) will thank you for that as it reduces their work in analysing the data (be it looking at data or performing wet-lab experiments).

The general workflow I use is to convert the CAF file to a gap4 or gap5 database. Then, in gap4 or gap5, I

1. quickly search for the UNSc and WRMc tags and check whether they could be real SNPs that were overseen by MIRA. In that case, I manually set a SROc (or SIOc) tag in gap4 via hotkeys that were defined to set these tags.
2. sometimes also quickly clean up reads that are causing trouble in alignments and lead to wrong base calling. These can be found at sites with UNSc tags, most of the time they have the 5' to 3' GGCxG motif which can cause trouble to Solexa.
3. look at sites with deletions (tagged with MCVc) and look whether I should clean up the borders of the deletion.

After this, I convert the gap4 or gap5 database back to CAF format. But beware: gap4 does not have the same consensus calling routines as MIRA and will have saved it's own consensus in the new CAF. In fact, gap4 performs rather badly in projects with multiple sequencing technologies. So I use `miraconvert` from the MIRA package to recall a good consensus (and save it in MAF as it's more compact and a lot faster in handling than CAF):

And from this MAF file I can then convert with `miraconvert` to any other format I or my users need: CAF, FASTA, ACE, WIG (for coverage analysis), SNP and coverage analysis (see below), HTML etc.pp.

7.7.2 Comprehensive SNP analysis spreadsheet tables (for Excel or OOcalc)

Biologists are not really interested in SNPs coordinates, and why should they? They're more interested where SNPs are, how good they are, which genes or other elements they hit, whether they have an effect on a protein sequence, whether they may be important etc. For organisms without intron/exon structure or splice variants, MIRA can generate pretty comprehensive tables and files if an annotated GenBank file was used as reference and strain information was given to MIRA during the assembly.

Well, MIRA does all that automatically for you if the reference sequence you gave was annotated.

For this, **`miraconvert`** should be used with the `asnp` format as target and a MAF (or CAF) file as input:

```
$ miraconvert -t asnp input.maf output
```

Note that it is strongly suggested to perform a quick manual cleanup of the assembly prior to this: for rare cases (mainly at site of small indels of one or two bases), MIRA will not tag SNPs with a SNP tag (SROc, SAOc or SIOc) but will be fooled into a tag denoting unsure positions (UNSc). This can be quickly corrected manually. See further down in this manual in the section on post-processing.

After conversion, you will have four files in the directory which you can all drag-and-drop into spreadsheet applications like OpenOffice Calc or Excel.

The files should be pretty self-explanatory, here's just a short overview:

1. `output_info_snplist.txt` is a simple list of the SNPs, with their positions compared to the reference sequence (in bases and map degrees on the genome) as well as the GenBank features they hit.
2. `output_info_featureanalysis.txt` is a much extended version of the list above. It puts the SNPs into context of the features (proteins, genes, RNAs etc.) and gives a nice list, SNP by SNP, what might cause bigger changes in proteins.
3. `output_info_featuresummary.txt` looks at the changes (SNPs, indels) from the other way round. It gives an excellent overview which features (genes, proteins, RNAs, intergenic regions) you should investigate.

There's one column (named 'interesting') which pretty much summarises up everything you need into three categories: yes, no, and perhaps. 'Yes' is set if indels were detected, an amino acid changed, start or stop codon changed or for SNPs in intergenic regions and RNAs. 'Perhaps' is set for SNPs in proteins that change a codon, but not an amino acid (silent SNPs). 'No' is set if no SNP is hitting a feature.

4. `output_info_featuresequences.txt` simply gives the sequences of each feature of the reference sequence and the resequenced strain.

7.7.3 HTML files depicting SNP positions and deletions

I've come to realise that people who don't handle data from NextGen sequencing technologies on a regular basis (e.g., many biologists) don't want to be bothered with learning to handle specialised programs to have a look at their resequenced strains. Be it because they don't have time to learn how to use a new program or because their desktop is not strong enough (CPU, memory) to handle the data sets.

Something even biologist know to operate are browsers. Therefore, `miraconvert` has the option to load a MAF (or CAF) file of a mapping assembly at output to HTML those areas which are interesting to biologists. It uses the tags `SROc`, `SAOc`, `SIOc` and `MCVc` and outputs the surrounding alignment of these areas together with a nice overview and links to jump from one position to the previous or next.

This is done with the `'-t hsnp'` option of `miraconvert`:

```
$ miraconvert -t hsnp input.maf output
```

Note I recommend doing this only if the resequenced strain is a very close relative to the reference genome, else the HTML gets pretty big. But for a couple of hundred SNPs it works great.

7.7.4 WIG files depicting contig coverage or GC content

`miraconvert` can also dump a coverage file in WIG format (using `'-t wig'`) or a WIG file for GC content (using `'-t gcwig'`). This comes pretty handy for searching genome deletions or duplications in programs like the Affymetrix Integrated Genome Browser (IGB, see <http://igb.bioviz.org/>) or when looking for foreign sequence in a genome.

7.7.5 Comprehensive spreadsheet tables for gene expression values / genome deletions & duplications

When having data mapped against a reference with annotations (either from GenBank formats or GFF3 formats), `miraconvert` can generate tables depicting either expression values (in RNASeq/EST data mappings) or probable genome multiplication and deletion factors (in genome mappings). For this to work, you must use a MAF or CAF file as input, specify `fcov` as output format and the reference sequence must have had annotations during the mapping with MIRA.

TODO: add example

```
miraconvert -t fcov mira_out.maf myfeaturetable
```

Chapter 8

Utilities in the MIRA package

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Ninety percent of success is just growing up.”

—Solomon Short

8.1 miraconvert

8.1.1 Synopsis

```
miraconvert [options] input_file output_basename
```

8.1.2 Description

miraconvert is a tool to convert, extract and sometimes recalculate all kinds of data related to sequence assembly files.

More specifically, **miraconvert** can

1. convert from multiple alignment files (CAF, MAF) to other multiple alignment files (CAF, MAF, ACE, SAM), and -- if wished -- selecting contigs by different criteria like name, length, coverage etc.
2. extract the consensus from multiple alignments in CAF and MAF format, writing it to any supported output format (FASTA, FASTQ, plain text, HTML, etc.) and -- if wished -- recalculating the consensus using the MIRA consensus engine with MIRA parameters
3. extract read sequences (clipped or unclipped) from multiple alignments and save to any supported format
4. Much more, need to document this.

...

8.1.3 Options

...

8.1.3.1 General options

- f** *caf* / *maf* / *fasta* / *fastq* / *gbf* / *phd* / *fofnexp* “From-type”, the format of the input file. CAF and MAF files can contain full assemblies and/or unassembled (single) sequences while the other formats contain only unassembled sequences.
 - t** *ace* / *asnp* / *caf* / *crlist* / *cstats* / *exp* / *fasta* / *fastq* / *fcov* / *gbf* / *gff3* / *hsnp* / *html* / *ma* “To-type”, the format of the output file. Multiple mentions of [-t] are allowed, in which case **miraconvert** will convert to multiple types.
 - a** Append. Results of conversion are appended to existing files instead of overwriting them.
 - A** Do not adjust sequence case.
When reading formats which define clipping points (like CAF, MAF or EXP), and saving to formats which do not have clipping information, **miraconvert** normally adjusts the case of read sequences: lower case for clipped parts, upper case for unclipped parts of reads. Use -A if you do not want this. See also -C.
-
- Note** Applies only to files/formats which do not contain contigs.
-
- b** Blind data. Replace all bases in all reads / contigs with a 'c'.
 - C** Hard clip reads. When the input is a format which contains clipping points in sequences and the requested output consists of sequences of reads, only the unclipped parts of sequences will be saved as results.
 - d** Delete gap only columns. When output is contigs: delete columns that are entirely gaps (can occur after having deleted reads during editing in gap4, consed or other). When output is reads: delete gaps in reads.
 - F** Filter read groups to different files. Works only for input files containing readgroups, i.e., CAF or MAF. 3 (or 4) files are generated: one or two for paired, one for unpaired and one for debris reads. Reads in paired file are interlaced by default, use -F twice to create separate files.
 - m** Make contigs. Encase single reads as contig singlets into a CAF/MAF file.
 - n** *namefile* Name select. Only contigs or reads are selected for output which name appears in *namefile*. *namefile* is a simple text file having one name entry per line.
 - i** When -n is used, inverts the selection.
 - o** *offset* Offset of quality values in FASTQ files. Only valid if -f is FASTQ.
 - P** *MIRA-PARAMETERSTRING* Additional MIRA parameters. Allows to initialise the underlying MIRA routines with specific parameters. A use case can be, e.g., to recalculate a consensus of an assembly in a slightly different way (see also [-r]) than the one which is stored in assembly files. Example: to tell the consensus algorithm to use a minimum number of reads per group for 454 reads, use: "454_SETTINGS -CO:mrpg=4".
Consult the MIRA reference manual for a full list of MIRA parameters.
 - q** *quality_value* When loading read data from files where sequence and quality are split in several files (e.g. FASTA with .fasta and .fasta.qual files), do not stop if the quality values for a read are missing but set them to be the *quality_value* given.
 - R** *namestring* Rename contigs/singlets/reads with given name string to which a counter is added.
Known bug: will create duplicate names if input (CAF or MAF) contains contigs/singlets as well as free reads, i.e. reads not in contigs nor singlets.
 - S** *namescheme* Naming scheme for renaming reads, important for paired-ends. Only 'solexa' is supported at the moment.
 - Y** *integer* Yield. Defines the maximum number of (clipped/padded) bases to convert. When used on reads: output will contain first reads of file where length of clipped bases totals at least -Y. When used on contigs: output will contain first contigs of file where length of padded contigs totals at least -Y.
-

8.1.3.2 Options for input containing contig data

The following switches will work only if the input file contains contigs (i.e., CAF or MAF with contig data). Though infrequent, note that both CAF and MAF can contain single reads only.

- M** Do not extract contigs (or their consensus), but the reads they are composed of.
- N *namefile*** Name select, sorted. Only contigs/reads are selected for output which name appears in *namefile*. Regardless of the order of contigs/reads in the input, the output is sorted according to the appearance of names in *namefile*. *namefile* is a simple text file having one name entry per line.
Note that for this function to work, all contigs/reads are loaded into memory which may be straining your RAM for larger projects.
- r *c | C | q | f | r*** Recalculate consensus and / or consensus quality values and / or SNP / repeat feature tags of an assembly. This feature is useful in case third party programs create own consensus sequences without handling different sequencing technologies (e.g. the combination of **gap4** and **caf2gap**) or when the CAF/MAF files do not contain consensus sequences at all.
 - c** recalculate consensus & consensus qualities using IUPAC where necessary
 - C** recalculate consensus & consensus qualities forcing ACGT calls and without IUPAC codes
 - q** recalculate consensus quality values only
 - f** recalculate SNP features (SROc, SAOc, SIOc)
 - r** recalculate repeat features (SRMc, WRMc)

Note Only the last of cCq is relevant, 'f' and 'r' work as a switches and can be combined with the others (e.g. "-r Cfr").

Note If the CAF/MAF contains reads from multiple strains, recalculation of consensus & consensus qualities is forced, you can just influence whether IUPACs are used or not. This is due to the fact that CAF/MAF do not provide facilities to store consensus sequences from multiple strains.

- s** Split. Split output into single files, one file per contig. Files are named according to name of contig.
- u** fillUp strain genomes. In assemblies made of multiple strains, holes in the consensus of a strain (bases 'N' or '@') can be filled up with the consensus of the other strains. Takes effect only when '-r' is active.
- Q *quality_value*** Defines minimum quality a consensus base of a strain must have, consensus bases below this will be set to 'N'. Only used when -r is active.
- V *coverage_value*** Defines minimum coverage a consensus base of a strain must have, consensus bases below this coverage will be set to 'N'. Only used when -r is active.
- v** Print version number and exit.
- x *length*** Minimum length a contig (in full assemblies) or read (in single sequence files) must have. All contigs / reads with a length less than this value are discarded. Default: 0 (=switched off).
Note: this is of course not applied to reads in contigs! Contigs passing the [-x] length criterion and stored as complete assembly (CAF, MAF, ACE, etc.) still contain all their reads.
- X *length*** Similar to [-x], but applies only to clipped reads (input file format must have clipping points set to be effective).
- y *contig_coverage*** Minimum average contig coverage. Contigs with an average coverage less than this value are discarded.
- z *min_reads*** Minimum number of reads in contig. Contigs with less reads than this value are discarded.
- l *line_length*** On output of assemblies as text or HTML: number of bases shown in one alignment line. Default: 60.
- c *endgap_character*** On output of assemblies as text or HTML: character used to pad endgaps. Default: ' ' (a blank)

8.1.4 Examples

In the following examples, the CAF and MAF files used are expected to contain full assembly data like the files created by MIRA during an assembly or by the gap2caf program. CAF and MAF could be used interchangeably in these examples, depending on which format currently is available. In general though, MAF is faster to process and smaller on disk.

Simple conversion: a MIRA MAF file to a SAM file

```
miraconvert source.maf destination.sam
```

Note

Previous versions of miraconvert had a slightly different syntax, which however is still supported:

```
miraconvert source.maf destination.sam
```

```
miraconvert source.caf destination.fasta wig ace
```

Simple conversion: the consensus of an assembly to FASTA, at the same time coverage data for contigs to WIG and furthermore

Note

Previous versions of miraconvert had a slightly different syntax, which however is still supported:

```
miraconvert -f caf -t fasta -t wig -t ace source.caf destination
```

```
miraconvert -x 2000 -y 10 source.caf destination.caf
```

Filtering an assembly for contigs of length ≥ 2000 and an average coverage ≥ 10 , while translating from MAF to CAF

```
miraconvert -x 55 -R newname source.fastq destination.fastq
```

Filtering and reordering contigs of an assembly according to external contig name list. This example will fetch the contigs named bchoc_c14, ...3, ...5 and ...13 and save the result in exactly that order to a new file:

```
arcadia:/path/to/myProject$ ls -l
-rw-r--r-- 1 bach users 231698898 2007-10-21 15:16 bchoc_out.caf
-rw-r--r-- 1 bach users      38 2007-10-21 15:16 contigs.lst
arcadia:/path/to/myProject$ cat contigs.lst
bchoc_c14
bchoc_c3
bchoc_c5
bchoc_c13
arcadia:/path/to/myProject$ miraconvert -N contigs.lst bchoc_out.caf myfilteredresult.caf
caf
[...]
arcadia:/path/to/myProject$ ls -l
-rw-r--r-- 1 bach users 231698898 2007-10-21 15:16 bchoc_out.caf
-rw-r--r-- 1 bach users      38 2007-10-21 15:16 contigs.lst
-rw-r--r-- 1 bach users 828726 2007-10-21 15:24 myfilteredresult.caf
```

8.2 mirabait - a "grep" like tool to select reads with kmers up to 256 bases

8.2.1 Synopsis

```
mirabait [options] -b baitfile [-b ...] | -B file [-p file1 file2 | -P file3]* [file4 ...]
```

Note The above command line, especially with mandatory [-b] format appeared only in MIRA 4.9.0 and represents a major change to 4.0.x!

8.2.2 Description

mirabait selects reads from a read collection which are partly similar or equal to sequences defined as target baits. Similarity is defined by finding a user-adjustable number of common k-mers (sequences of k consecutive bases) which are the same in the bait sequences and the screened sequences to be selected, either in forward or reverse complement direction.

When used on paired files (-p or -P), selects read pairs where at least one read matches.

One can use **mirabait** to do targeted assembly by fishing out reads belonging to a gene and just assemble these; or to clean out rRNA sequences from data sets; or to fish out and iteratively reconstruct mitochondria from metagenomic data; or, or, or ... whenever one has to take in or take out subsets of reads based on kmer equality, this tool should come in quite handy.

Note The search performed is exact, that is, sequences selected are guaranteed to have the required number of matching k-mers to the bait sequences while sequences not selected are guaranteed not have these.

8.2.3 Options

8.2.3.1 Main options

-b file A file containing sequences to be used as bait. The file can be in any of the following types: FASTQ, FASTA, GenBank (.gbf, .gbk, .gbff), CAF, MAF or Staden EXP.

If the the file extension is non-standard (e.g. `file.dat`), you can force a file type by using the double colon type specification like in EMBOSS. E.g.: `fastq::file.dat`

Using multiple -b for loading bait sequences from multiple files is allowed.

-B file Load bait from an existing kmer statistics file, not from sequence files. Only one -B allowed, cannot be combined with -b. See -K on how to create such a file.

This option comes in handy when you always want to bait against a given set of sequences, e.g., rRNA sequences or the human genome, and where the statistics computation itself may be quite time and resource intensive. Once computed and saved via [-K], a baiting process loading the statistics via [-B] can start much faster.

-j job Set option for predefined job from supplied MIRA library. Currently available jobs:

- 'rrna': Bait rRNA/rDNA sequences. Locked options: [-b, -B, -k, -K, -n]. In the current version mirabait will use a hash statistics file with 21mers derived from a subset of the RFAM 12 rRNA database to bait rRNA/rDNA reads. The supplied subset should catch all but the most uncommon rRNA data, if needed one could albeit increase the sensitivity by decreasing [-n].

Note that [-j] will hardwire a number of options to be optimal for the chosen job. Note that it is not advisable to change the 'locked' options as this either breaks the functionality or worse, it could lead to undefined results.

-p file_1 file_2 Instructs to load sequences to be baited from files `file_1` and `file_2`. The sequences are treated as pairs, where a read in one file is paired with a read in the second file. The files can be in any of the following types: FASTQ, FASTA, GenBank (.gbf, .gbk, .gbff), CAF, MAF or Staden EXP.

If the the file extension is non-standard (e.g. `file.dat`), you can force a file type by using the double colon type specification like in EMBOSS. E.g.: `fastq::file.dat`

Using multiple -p for baiting sequences from multiple file pairs is allowed.

-P file Instructs to load sequences to be baited from file `file`. The sequences are treated as pairs, where a read in the file is immediately followed by its paired read. The file can be in any of the following types: FASTQ, FASTA, GenBank (.gbf, .gbk, .gbff), CAF, MAF or Staden EXP.

If the the file extension is non-standard (e.g. `file.dat`), you can force a file type by using the double colon type specification like in EMBOSS. E.g.: `fastq::file.dat`

Using multiple -P for baiting sequences from multiple files is allowed.

- k *kmer-length*** k-mer, length of bait in bases (≤ 256 , default=31)
- n *integer*** Default value: 1.
 If the integer given is > 0 : minimum number of kmers needed for a sequence to be selected.
 If the integer given is ≤ 0 : maximum number of missed kmers allowed over sequence length for a sequence to be selected.
- d** Do not use kmers with microrepeats (DUST-like). Standard length for microrepeats is 67% of kmer length, see [-D] to change this.
 Microrepeats are defined as repeats of a 1, 2, 3 or 4 base motif at either end (not in the middle) of a kmer. E.g.: a kmer of 17 will have a microrepeat length of 12 bases, so that, all kmers having 12 A, C, G, T at either end will be filtered away. E.g.: AAAAAAAAAAAnnnnn as well as nnnnnAAAAAAAAAAAAA will be filtered.
 E.g. for repeats of 2 bases: AGAGAGAGAGAGnnnnn or CACACACACACAnnnnn.
 E.g. for repeats of 3 bases: ACGACGACGACGnnnnn.
 E.g. for repeats of 4 bases: ACGTACGTACGTnnnnn.
 Microrepeat motifs will truncate at the end of allocated microrepeat length. E.g. kmer length 20 with microrepeat length of 13 and 4 base repeat: ACGTACGTACGTAnnnnnn.

Note

When saving the kmer statistics via [-K], having [-d] will save kmer statistics where kmers with microrepeats have already been removed. Use this when you always want to have microrepeats removed from a given bait data as [-d] will not be needed when using via [-B] that set in later loads (which saves time).
 If you want to be able to bait from precomputed kmer statistics both with and without microrepeats, use [-d] only when loading the statistics file with [-B], not when creating it with [-K].

- D *integer*** Set length of microrepeats in kmers to discard from bait.
 int > 0 : microrepeat length in percentage of kmer length. E.g.: -k 17 -D 67 --> 67% of 17 bases = 11.39 bases --> 12 bases.
 int < 0 : microrepeat length in bases.
 int $\neq 0$ implies -d, int=0 turns DUST filter off
- i** Inverse selection: selects only sequence that do not meet the -k and -n criteria.
- I** Filters and writes sequences which hit to one file and sequences which do not hit to a second file.
- r** Does not check for hits in reverse complement direction.
- t *integer*** Number of threads to use. The default value of 0 is configured to automatically use up to 4 CPU cores (if present). Numbers higher than 4 (or maybe 8) will probably not make much sense because of diminishing returns.

8.2.3.2 File type options

Normally, mirabait writes separate result files (named `bait_match_*` and `bait_miss_*`) for each input to the current directory. For changing this behaviour, and others relating to output, use these options:

- c *character*** Normally, mirabait will change the case of the sequences it loads to denote kmers which hit a bait in upper case and kmers which did not hit a bait in lower case. Using this option, one can instead mask those bases with the given character.
 Use a blank (-c ' ') to neither mask nor lowercase hits.
- l *integer*** Set length of sequence line in FASTA output.
- K *filename*** Save kmer statistics (for baits loaded via [-b]) to *filename*.
 As the calculation of kmers can take quite some time for larger sequences (e.g., human genome), this option is extremely useful if you want to perform the same baiting operation more than once. Once calculated, the kmer statistics is saved and can be reloaded for a later baiting operation via [-B].
-

- N *name*** Change the file prefix `bait` to `name`. Has no effect if `-o/-O` is used and targets are not directories.
- o *path*** Save sequences matching a bait to `path`. If `path` is a directory, write separate files into this directory. If not, combine all matching sequences from the input file(s) into a single file specified by the path.
- O *path*** Like `-o`, but for sequences not matching.

8.2.3.3 Other options

- T *dir*** Use `dir` as directory for temporary files instead of the current working directory.
- m *integer*** Default is `75`. Defines the memory MIRA can use to compute kmer statistics. Therefore does not apply when using `[-B]`.
 A value of ≥ 100 is interpreted as absolute value in megabyte. E.g., `16384` = 16384 megabyte = 16 gigabyte.
 A value of $0 \leq x \leq 100$ is interpreted as relative value of free memory at the time of computation. E.g.: for a value of `75%` and 10 gigabyte of free memory, it will use 7.5 gigabyte.

Note The minimum amount of memory this algorithm will use is 512 MiB on 32 bit systems and 2 GiB on 64 bit systems.

8.2.4 Usage examples

Note The examples below, together with the manual above, should be enough to get you going. If there's a typical use case you are missing, feel free to suggest it on the MIRA talk mailing list.

Baiting unpaired sequences, bait sequences in FASTA, sequences in FASTQ:

```
mirabait -b b.fasta file.fastq
```

Same as above, but baits in two files (FASTA and GenBank):

```
mirabait -b b1.fasta -b b2.gbk file.fastq
```

Baiting paired sequences, read pairs are in two files:

```
mirabait -b b.fasta -p file_1.fastq file_2.fastq
```

Baiting paired sequences, pairs are interleaved in one file:

```
mirabait -b b.fasta -P file.fastq
```

Like above, but selecting sequences which do not match the baits:

```
mirabait -i -b b.fasta -P file.fastq
```

Baiting paired sequences (`file_1.fastq`, `file_2.fastq` and `file3.fastq`) and unpaired sequences (`file4.fastq`), all at once and different file types:

```
mirabait -b b.fasta -p file_1.fastq file_2.fastq -P file3.fasta file4.caf
```

Like above, but writing sequences matching baits and sequences not matching baits to different files:

```
mirabait -I -b b.fasta -p file_1.fastq file_2.fastq -P file3.fasta file4.caf
```

Change bait criterion to need 10 kmers of size 27:

```
mirabait -k 27 -n 10 -b b.fasta file.fastq
```

Change bait criterion to baiting only reads which have all kmers present in the bait:

```
mirabait -n 0 -b b.fasta file.fastq
```

Change bait criterion to baiting all reads having almost all kmers present in the bait, but allowing for up to 40 kmers not in the bait:

```
mirabait -n -40 -b b.fasta file.fastq
```

Force bait sequences to load as FASTA, force sequences to be baited to be loaded as FASTQ:

```
mirabait -b fasta::b.dat fastq::file.dat
```

Write result files to directory /dev/shm/:

```
mirabait -o /dev/shm/ -b b.fasta -p file_1.fastq file_2.fastq
```

Merge all result files containing sequences hitting baits to file /dev/shm/match:

```
mirabait -o /dev/shm/match -b b.fasta -p file_1.fastq file_2.fastq
```

Like above, but also merge all result files containing sequences not hitting baits to file /dev/shm/nomatch:

```
mirabait -o /dev/shm/match -O /dev/shm/nomatch -b b.fasta -p file_1.fastq file_2.fastq
```

Fetch all reads having rRNA motifs in a paired FASTQ files:

```
mirabait -j rrna -p file1.fastq file2.fastq
```

Fetch all reads not having rRNA motifs in a paired FASTQ files:

```
mirabait -j rrna -i -p file1.fastq file2.fastq
```

Split a paired FASTQ file into two sets of files (4 files total), one containing rRNA reads and one containing non-rRNA reads:

```
mirabait -j rrna -I -p file1.fastq file2.fastq
```

Assuming the file `human_genome.fasta` contains the human genome: bait all read pairs matching the human genome. Also, save the compute kmer statistics for later re-use to file `HG_kmerstats.mhs.gz`:

```
mirabait -b human_genome.fasta -K HG_kmerstats.mhs.gz -p file1.fastq file2.fastq
```

The same as above, but just precompute and save the kmer statistics, no actual baiting done.

```
mirabait -b human_genome.fasta -K HG_kmerstats.mhs.gz
```

Using the precomputed kmer statistics from the command above: bait files with read pairs for human reads:

```
mirabait -B HG_kmerstats.mhs.gz -p file_1.fastq file_2.fastq
```

8.2.5 Installing different rRNA databases

The standard database for rRNA baiting supplied with the MIRA source code and binary packages is called `rfam_rrna-21-12.sls.gz` which will get installed as *MHS* (MiraHashStatistics) file into `$BINDIR/./share/mira/mhs/rfam_rrna-21-12.mhs.gz` (where `$BINDIR` is the directory where the `mira/mirabait` binary resides) and a soft link pointing from `filter_default_rrna.mhs.gz` to `rfam_rrna-21-12.mhs.gz` like so:

```
arcadia:~$ which mira
/usr/local/bin/mira
arcadia:~$ ls -l /usr/local/share/mira/mhs
lrwxrwxrwx 1 bach bach          22 Mar 24 23:58 filter_default_rrna.mhs.gz -> rfam_rrna
-21-12.mhs.gz
-rw-rw-r-- 1 bach bach 148985059 Mar 24 23:58 rfam_rrna-21-12.mhs.gz
```

The file naming scheme for the database is as following: `dbidentifier-kmerlength-kmerfreqcutoff`. The standard database is therefore: `rfam_rrna` as identifier for the RFAM rRNA sequences (currently RFAM 12), then 21 defining a kmer length of 21 and finally a kmer cutoff frequency of 12, meaning that kmers must have been seen at least 12 times in the RFAM database to be stored in the subset.

Note The value of 12 as frequency cutoff for the standard mirabait rRNA database was chosen as a compromise between sensitivity and database size.

Although rRNA are pretty well conserved overall, the cutoff frequency also implies that kmers from rare rRNA variants will not be present in the database, eventually losing some sensitivity for rRNA from rarely sequenced organisms. It follows that more sensitive versions of the rRNA database can be installed by downloading a file from the MIRA repository at SourceForge and calling a script provided by MIRA. To install a version with a kmer size of 21 and a cutoff frequency of, e.g., 3, download `rfam_rrna-21-3.sls.gz` and install it like this:

```
arcadia:~/tmp$ ls
arcadia:~/tmp$ wget https://sourceforge.net/projects/mira-assembler/files/MIRA/slsfiles/ ←
    rfam_rrna-21-3.sls.gz
...
```

TODO: continue docs here.

Chapter 9

MIRA 5 reference manual

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“The manual only makes sense after you learn the program.”

—Solomon Short

9.1 Synopsis

```
mira [-chmMrtv] manifest-file [manifest-file ...]
```

The command line parameters in short:

[-c / --cwd=*directory*] Change working directory.

[-h / --help] Print a short help and exit.

[-m / --mcheck] Only check the manifest file, then exit.

[-M / --mdcheck] Only check the manifest file and presence of data files, then exit.

[-r / --resume] Resume / restart an interrupted assembly. Works only for de-novo assemblies at the moment.

[-t / --thread=*integer* ≥ 0] Force number of threads (overrides equivalent [-GE:not] manifest entry).

[-v / --version] Print version and exit.

9.2 Requirements

To use MIRA itself, one doesn't need very much:

- Sequence data in EXP, CAF, PHD, FASTA or FASTQ format
 - Optionally: ancillary information in NCBI traceinfo XML format; ancillary information about strains in tab delimited format, vector screen information generated with **ssaha2** or **smalt**.
 - Some memory and disk space. Actually lots of both if you are venturing into 454 or Illumina.
-

9.3 Configuring an assembly: files and parameters

MIRA can be used for a number of tasks: you can use it to do de-novo assembly of high-throughput sequencing data for genome or RNA transcript reconstruction. Or you can use the same data and map it to reference sequences to search for mutations, extract coverage or expression information etc.pp. Since 4.9.x, MIRA also has two new modes: you can use for assembly of smaller fragments (BAC size) without MIRA performing heavy preprocessing; or you can use it to cluster sequences (like transcript contigs).

All the configuration needed for an assembly is done in one (or several) configuration file(s): the *manifest* files. This encompasses things like what kind of assembly you want to perform (genome or EST / RNASeq, mapping or de-novo etc.pp) or which data files contain the sequences you want to assemble (and in which format these are).

9.3.1 The manifest file: introduction

A *manifest* file can be seen as a two part configuration file for an assembly: the first part contains some general information and tell MIRA how exactly you want the data to be assembled while the second part contains information about the sequencing data to be loaded. Examples being always easier to follow than long texts, here's an example for a de-novo assembly with single-end (also called shotgun) 454 data:

```
# Example for a manifest describing a simple 454 de-novo assembly

# A manifest file can contain comment lines, these start with the #-character

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should assemble a genome de-novo in accurate mode
# As special parameter, we want to use 4 threads in parallel (where possible)

project = MyFirstAssembly
job = genome,denovo,accurate
parameters = -GE:not=4

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups": this reflects the
# ... that read sequences ...

readgroup = SomeUnpaired454ReadsIGotFromTheLab
data = TCMFS456ZH345.fastq TQF92GT7H34.fastq
technology = 454
```

To make things a bit more interesting, here's an example using a couple more technologies and showing some more options of the manifest file like wild cards in file names, different paired-end/mate-pair libraries and how to let MIRA refine pairing information (or even find out everything by itself):

```
# Example for a manifest describing a de-novo assembly with
# unpaired 454, paired-end Illumina, a mate-pair Illumina
# and a paired Ion Torrent

# First part: defining some basic things
# In this example, we just give a name to the assembly
# and tell MIRA it should assemble a genome de-novo in accurate mode
# As special parameter, we want to use 4 passes with kmer sizes of
# 17, 31, 63 and 127 nucleotides. Obviously, read lengths of the
# libraries should be greater than 127 bp.
# Note: usually MIRA will choose sensible options for number of
# passes and kmer sizes to be used by itself.

project = MyFirstAssembly
job = genome,denovo,accurate
```

```

parameters = -AS:kms=17, 31, 63, 127

# The second part defines the sequencing data MIRA should load and assemble
# The data is logically divided into "readgroups": this reflects the
# ... that read sequences ...

# defining the shotgun (i.e. unpaired) 454 reads
readgroup = SomeUnpaired454ReadsIGotFromTheLab
data = TCMFS456ZH345.fastq TQF92GT7H34.fastq
technology = 454

# defining the paired-end Illumina reads, fixing all needed pair information
readgroup = SomePairedEndIlluminaReadsIGotFromTheLab
data = datape*.fastq
technology = solexa
template_size = 100 300
segment_placement = ---> <---
segment_naming = solexa

# defining the mate-pair Illumina reads, fixing most needed pair information
# but letting MIRA refine the template_size via "autorefine"
readgroup = SomeMatePairIlluminaReadsIGotFromTheLab
data = datamp*.fastq
technology = solexa
template_size = 2000 4000 autorefine
segment_placement = <--- --->
segment_naming = solexa

# defining paired Ion Torrent reads
# example to show how lazy one can be and simply let MIRA estimate by itself
# all needed pairing information via "autopairing"
# Hint: it usually does a better job at it than we do ;- )
readgroup = SomePairedIonReadsIGotFromTheLab
autopairing
data = dataion*.fastq
technology = iontor

```

9.3.2 The manifest file: basics

The first part of an assembly *manifest* contains the very basic information the assembler needs to have to know what you want it to do. This part consists of exactly three entries:

1. **project** = [= *project name*] tells the assembler the name you wish to give to the whole assembly project. MIRA will use that name throughout the whole assembly for naming directories, files and a couple of other things.

You can name the assembly anyway you want, you should however restrain yourself and use only alphanumeric characters and perhaps the characters plus, minus and underscore. Using slashes or backslashes here is a recipe for catastrophe.

2. **job** = [*denovo*|*mapping*], [*genome*|*est*|*fragments*|*clustering*], [*draft*|*accurate*] tells the assembler what kind of data it should expect and how it should assemble it.

You need to make your choice mainly in three steps and in the end concatenate your choices to the [job=] entry of the manifest:

- (a) are you building an assembly from scratch (choose: *denovo*) or are you mapping reads to an existing reference sequence (choose: *mapping*)? Pick one. Leaving this out automatically chooses *denovo* as default.
- (b) are the data you are assembling forming a larger contiguous sequence (choose: *genome*), are you assembling EST or mRNA libraries (choose: *est*), single genes or small plasmids (choose: *fragments*) or do you cluster assembled sequences (choose: *clustering*)? Pick one. Leaving this out automatically chooses *genome* as default.

Since version 4.9.4, a new mode *fragments* is available. This mode is essentially similar to the *EST* mode, but has all safety features switched off which reduce data sizes. Use this mode for assembly of comparatively small EST/mRNA or small plasmid or single gene projects where you want to have highest accuracy and minimal filtering. Warning: contigs with coverages going into the 1000s will lead to really slow assemblies.

Since version 4.9.6, a new mode *clustering* is available. This mode is essentially for clustering assembled contigs like they are created in mRNA or EST assemblies. Basic parameters are: single pass, no clipping, no editing. When in "normal" mode: ~7.5% differences between sequences allowed, gaps ≥ 13 bases disallowed. When in "accurate" mode: ~2% differences between sequences allowed, gaps completely disallowed. Single occurrence of disagreeing base leads to SNP tagging. Warning: do not use that with any type of real sequencing data ... you probably would regret this.

- (c) For de-novo assemblies and mappings: do you want a quick and dirty assembly for first insights (choose: *draft*) or an assembly that should be able to tackle even most nasty cases (choose: *accurate*)?

Pick one. Leaving this out automatically chooses *accurate* as default.

Once you're done with your choices, concatenate everything with commas and you're done. E.g.: `'--job=mapping, genome,` will give you a mapping assembly of a genome in draft quality.

Note For de-novo assembly of genomes, these switches are optimised for 'decent' coverages that are commonly seen to get you something useful, i.e., $\geq 7x$ for Sanger, $\geq 18x$ for 454 FLX or Titanium, $\geq 25x$ for 454 GS20 and $\geq 30x$ for Solexa. Should you venture into lower coverage or extremely high coverage (say, $\geq 60x$ for 454), you will need to adapt a few parameters via extensive switches.

3. **parameters =** is used in case you want to change one of the 150+ extended parameters MIRA has to offer to control almost every aspect of an assembly. This is described in more detail in a separate section below.

9.3.3 The manifest file: information on the data you have

The second part of an assembly *manifest* tells MIRA which files it needs to load, which sequencing technology generated the data, whether there are DNA template constraints it can use during the assembly process and a couple of other things.

9.3.3.1 Starting a new readgroup

readgroup [= *group name*] is the keyword which tells MIRA that you are going to define a new read group. You can optionally name that group.

Understanding readgroups and DNA templates

When you send away your DNA for sequencing, it is going to be prepared for sequencing according to your wishes. Sequencing providers call this "constructing a library" and regardless whether you sequence with Sanger, 454, Illumina, Ion Torrent, Pacific Biosciences or other technologies, the "library prep" is always there.

With most library preps, your DNA is first amplified and then cut into small pieces. These pieces are called *templates* and their length can be anywhere between a few dozen bases, a few hundred bases or even a couple of dozen or even hundred kilobases. The important thing is that these templates can be much bigger in size than the actual read length. While this is a wet lab step, protocols and providers have gotten pretty good at constructing libraries where the DNA templates are all in a given range of bases like, e.g., having a library with template size 500bp (+/- 100bp) and another library with template size around 7kb (+/- 500bp).

Depending on the technology and sequencing strategy used, the DNA templates are used to create either one single read or - and that's important - two or more reads.

Libraries with "single reads" are often called "single read libraries" or "shotgun libraries". They can be found for every sequencing technology and are most of the time easy to construct (therefore cheap) and are often used to provide a decent amount of bases as basic coverage for your project.

Libraries with two reads per DNA template are often called "mate-pair" or "paired-end" libraries. They are harder to construct and sometime have less yield, therefore they are often more expensive. But the sequencing approach using several reads per DNA template allows assembly and scaffolding algorithms to resolve repetitive regions of a genome which are longer than the average read length. Note that Pacific Biosciences has a sequencing mode called "strobed sequencing" which is different from "paired-end/mate-pair" but also creates multiple reads per DNA template.

Long story short: an assembler must know afterwards what kind of reads it has to expect: the sequencing technology, library preparation strategy etc. For this, the notion of *read groups* has emerged: reads coming from the same technology and same library preparation are pooled together in a read group to tell the assembler: in the assembly, if you see two reads coming from a same DNA template, you should expect them to be at a certain distance from each other and they should be oriented in a certain way.

Note The above was a **very** simplified view on the whole area of DNA templates, readgroups, shotgun and paired end sequencing. Enough to hopefully understand the concepts, but you might want to read more about it.

9.3.3.2 Defining data files to load

`data = filepath [filepath ...]` defines the file paths from which sequences should be loaded. A file path can contain just the name of one (or several) files or it can contain the *path*, i.e., the directory (absolute or relative) including the file name.

MIRA automatically recognises what type the sequence data is by looking at the postfix of files. For postfixes not adhering widely used naming schemes for file types, there's additionally a way of explicitly defining the type (see further down at the end of this item on how this is done). Currently allowed file types are:

- `.fasta` for sequences formatted in FASTA format where there exists an additional `.fasta.qual` file which contains quality data. If the file with quality data is missing, this is interpreted as error and MIRA will abort.
- `.fna` and `.fa` also for sequences formatted in FASTA format. The difference to `.fasta` lies in the way MIRA treats a missing quality file (called `.fna.qual` or `.fa.qual`): it does not see that as critical error and continues.
- `.fastq` or `.fq` for files in FASTQ format
- `.gff3` or `.gff` for files in GFF3 format. Note that MIRA will load all sequences and annotations contained in this file.
- `.gbk`, `.gbf`, `.gbff` or `.gb` for files formatted in GenBank format. Note that the MIRA GenBank loader does not understand intron/exon or other multiple-locus structures in this format, use GFF3 instead!
- `.caf` for files in the CAF format (from Sanger Centre)

- `.maf` for files in the MIRA MAF format
- `.exp` for files in the Staden EXP format.
- `.fofnexp` for a *file of EXP filenames* which all point to files in the Staden EXP format.
- `.xml`, `.ssaha2` and `.smalt` for ancillary data in NCBI TRACEINFO, SSAHA2 or SMALT format respectively.

Multiple 'data' lines and multiple entries per line (even different formats) are allowed, as in, e.g.,

```
data = file1.fastq file2.fastq file3.fasta file4.gbk
data = myscreenings.smalt
```

You can also use wildcards and/or directory names. E.g., loading all file types MIRA understand from a given directory `mydir`:

```
data = mydir
```

or loading all files starting with `mydata` and ending with `fastq`:

```
data = mydata*fastq
```

or loading all files in directory `mydir` starting with `mydata` and ending with `fastq`:

```
data = mydir/mydata*fastq
```

or loading all FASTQ files in all directories starting with `mydir`:

```
data = mydir*/*fastq
```

or ... well, you get the gist.

Note Giving a directory like in `mydir` is equivalent to `mydir/*` (saying: give me all files in the directory `mydir`), however the first version should be preferred when the directory contains thousands of files.

Note

GenBank and GFF3 files may or may not contain embedded sequences. If annotations are present in these files for which no sequence is present in the same file, MIRA will look for reads of the same name which it already loaded in this or previously defined read groups and add the annotations there.

As security measure, annotations in GenBank and GFF3 files for which absolutely no sequence or read has been defined are treated as error.

Explicit definition of file types. It is possible to explicitly tell MIRA the type of a file even if said file does not have a 'standard' naming scheme. For this, the EMBOSS double-colon notation has been adapted to work also for MIRA, i.e., you prepend the type of a file and separate it from the file name by a double colon. E.g., the `.dat` postfix is not anything MIRA will recognise, but you can define it should be loaded as FASTQ file like this:

```
data = fastq::myfile.dat
```

Another frequent usage is forcing MIRA to load FASTA files named `.fasta` without complaining in case quality files (which MIRA wants you to provide) are not present:

```
data = fna::myfile.fasta
```

This does (of course) work also with directories or wildcard characters. In the following example, the first line will load all files from `mydirectory` as FASTQ while the second line loads just `.dat` files in a given path as FASTA:

```
data = fastq::mydirectory
data = fasta::/path/to/somewhere/*.dat
```

It is entirely possible (although not really sensible), to give contradicting information to MIRA by using a different explicit file type than one would guess from the standard postfix. In this case, the explicit type takes precedence over the automatic type. E.g.: to force MIRA to load a file as FASTA although it is named `.fastq`, one could use this:

```
data = fasta::file.fastq
```

Note that the above does not make any kind of file conversion, `file.fastq` needs to be already in FASTA format or else MIRA will fail loading that data.

9.3.3.3 Setting default quality

default_qual= *quality_value* is meant to be used as default fall-back quality value for sequences where the data files given above do not contain quality values. E.g., GFF3 or GenBank formats, eventually also FASTA files where quality data files is missing.

9.3.3.4 Defining technology used to sequence

technology= *technology* which names the technology with which the sequences were produced. Allowed technologies are: *sanger*, *454*, *solexa*, *iontor*, *pcbiolq*, *pcbiohq*, *text*.

The *text* technology is not a technology per se, but should be used for sequences which are not coming from sequencing machines like, e.g., database entries, consensus sequences, artificial reads (which do not comply to normal behaviour of 'normal' sequencing data), etc.pp

9.3.3.5 Preventing statistics for technologies with biases

nostatistics used as keyword will prevent MIRA to calculate coverage estimates from reads of the given readgroup.

This keyword should be used in denovo genome assemblies for reads from libraries which produce very uneven coverage (e.g.: old Illumina mate-pair protocols) or have a bias in the randomness of DNA fragmentations (e.g.: Nextera protocol from Illumina).

9.3.3.6 Setting reference sequence for mapping jobs

as_reference This keyword indicates to MIRA that the sequences in this readgroup should not be assembled, but should be used as reference backbone for a mapping assembly. That is, sequencing reads are then placed/mapped onto these reference reads.

9.3.3.7 Autopairing: letting MIRA find out pair info by itself

autopairing This keyword is used to tell MIRA it should estimate values for *template_size* and *segment_placement* (see below).

This is basically the lazy way to tell MIRA that the data in the corresponding readgroup consists of paired reads and that you trust it will find out the correct values.

Note *autopairing* usually works quite well for small and mid-sized libraries (up to, say, 10 kb). For larger libraries it might be a good thing to tell MIRA some rough boundaries via *template_size* / *segment_placement* and let MIRA refine the values for the template size via *autorefine* (see below).

Note *autopairing* is a feature new to MIRA 4.0rc5, it may contain bugs for some corner cases. Feedback appreciated.

9.3.3.8 Setting size of read templates

template_size = *min_size* *max_size* [*infoonly*|*exclusion_criterion*] [*autorefine*]. Defines the minimum and maximum size of "good" DNA templates in the library prep for this read group. This defines at which distance the two reads of a pair are to be expected in a contig, a very useful information for an assembler to resolve repeats in a genome or different splice variants in transcriptome data.

If the term *infoonly* is present, then MIRA will pass the information on template sizes in result files, but will not use it for any decision making during de-novo or mapping assembly. The term *exclusion_criterion* makes MIRA use the information for decision making.

If *infoonly* or *exclusion_criterion* are missing, then MIRA assumes *exclusion_criterion* for de-novo assemblies and *infoonly* for mapping assemblies.

If the term *autorefine* is present, MIRA will start the assembly with the given size information but switch to refined value computed from observed distances in an assembly. However, please note that the size values can *never* be expanded, only shrunk. It is therefore advisable to use generous bounds when using the autorefine feature.

Note The *template_size* line in the manifest file replaces the parameters -GE:uti:tismin:tismax of earlier versions of MIRA (3.4.x and below).

Note The minimum or the maximum size (or both) can be set to a negative value for "don't care and don't check". This allows constructs like `template_size= 500 -1 exclusion_criterion` which would check only the minimum distance but not the maximum distance.

Note

For *mapping* assemblies with MIRA, you usually will want to use *infoonly* as else - in case of genome re-arrangements, larger deletions or insertions - MIRA would probably reject one read of every read pair in the corresponding areas as it would not be at the expected distance and/or orientation ... and you would not be able to simply find the re-arrangement in downstream analysis.

For *de-novo* assemblies however you *should not* use *infoonly* except in very rare cases where you know what you do.

Understanding the size of DNA templates

When using a *paired-end* or *mate-pair* sequencing strategy, two sequences are generated for the ends of each DNA template (see sidebar above: "understanding readgroups and DNA templates"). That is, if one has a library with 6kb fragments, one knows that the outer ends of the two reads will be approximately 6kb apart, like so:

```
DNA template      #####
read 1            .....
read 2            .....
<----- ~6 kb ----->
```

Sequencing labs will try their best to get these two sequences from DNA templates which comply to a given length specification. But as this is chemistry and wet lab, things must be seen with a certain uncertainty and therefore the DNA templates generated are not exactly of the specified size (e.g. 6kb), but the size distribution will vary in a given range, e.g., 5.5kb to 6.5 kb.

9.3.3.9 Read segment placement

Note You do not need to use this when using 'autopairing' (see above).

segment_placement = *placementcode* [*infoonly*|*exclusion_criterion*]. Allowed placement codes are:

- **?** or **unknown** which are place-holders for "well, in the end: don't care." Segments of a template can be reads in any direction and in any relationship to each other.
This is typically used for unpaired libraries (sometimes called *shotgun libraries*), but may be also useful for, e.g., primer walking with Sanger.
- **---> <---** or **FR** or **INNIES**. The *forward / reverse* scheme as used in traditional Sanger sequencing as well as Illumina paired-end sequencing,
This is the usual placement code for Sanger paired-end protocols as well as Illumina paired-end. Less frequently used in IonTorrent paired-end sequencing.
- **<--- --->** or **RF** or **OUTIES**. The *reverse / forward* scheme as used in Illumina mate-pair sequencing.
This is the usual placement code for Illumina mate-pair protocols.
- **1---> 2--->** or **samedir forward** or **SF** or **LEFTIES**. The *forward / forward* scheme. Segments of a template are all placed in the same direction, the segment order in the contig follows segment ordering of the reads.
- **2---> 1--->** **samedir backward** or **SB** or **RIGHTIES**. Segments of a template are all placed in the same direction, the segment order in the contig is reversed compared to segment ordering of the reads.
This is the usual placement code for 454 "paired-end" and IonTorrent long-mate protocols.
- **samedir** Segments of a template are all placed in the same direction, the spatial relationship however is not cared of.
- **>>>** (reserved for sequencing of several equidistant fragments per template like in PacBio strobe sequencing, not implemented yet)

If the term *infoonly* is present, then MIRA will pass the information on segment placement in result files, but will not use it for any decision making during de-novo assembly or mapping assembly. The term *exclusion_criterion* makes MIRA use the information for decision making.

If *infoonly* or *exclusion_criterion* are missing, then MIRA assumes *exclusion_criterion* for de-novo assemblies and *infoonly* for mapping assemblies.

Note

For *mapping* assemblies with MIRA, you usually will want to use *infoonly* as else - in case of genome re-arrangements, larger deletions or insertions - MIRA would probably reject one read of every read pair (as it would not be at the expected distance and/or orientation) and you would not be able to simply find the re-arrangement in downstream analysis.

For *de-novo* assemblies however you *should not* use *infoonly* except in very rare cases where you know what you do.

Note As soon as you tell MIRA that a readgroup contains paired reads (via one of the other typical readgroup parameters like *template_size*, *segment_naming* etc.), the *segment_placement* line becomes mandatory in the manifest. This is because different sequencing technologies and/or library preparations result in different read orientations. E.g., Illumina libraries come in paired-end flavour which have FR (forward/reverse) placements, but there are also mate-pair libraries which have reverse/-forward (RF) placements.

Understanding read segment placement on DNA templates

bla

9.3.3.10 Read segment naming

segment_naming = *naming_scheme* [*rollcomment*]. Defines the naming scheme reads are following to indicate the DNA template they belong to. Allowed naming schemes are: *sanger*, *stlouis*, *tigr*, *FR*, *solexa*, *sra*.

If not defined, the defaults are *sanger* for Sanger sequencing data, while *solexa* for Solexa, 454 and Ion Torrent.

For FASTQ files, the modifier *rollcomment* can be used to let MIRA take the first token in the comment as name of a read instead of the original name. E.g.: for a read

```
@DRR014327.1.1 HWUSI-EAS547_0013:1:1:1106:4597.1 length=91
TTAGAAGGAGATCTGGAGAACATTTTAAACCGGATTGAACAACGCGGCCGTGAGATGGAGCTTCAGACAAGCCGGTCTTATTGGGACGAAC
+
bbb`bbbbabbR`\_bb_bba`b`bb_bb_`^\^Y^\Zb^b`]]\S^a`]a`'bbbb_bbbb]bbb\`^^^]\aaY`\`^\^aa__aB
```

the *rollcomment* modifier will lead to the read being named `HWUSI-EAS547_0013:1:1:1106:4597.1` (which is almost the original instrument read name) instead of `DRR014327.1.1` (which is the SRA read name).



Warning For data from the short read archive (SRA), one will usually need to explicitly specify the 'sra' naming scheme or use the 'rollcomment' modifier in FASTQ files.

Note This has changed with MIRA 3.9.1 and **sff_extract** 0.3.0. Before that, 454 and Ion Torrent were given *fr* as naming scheme.

Understanding read naming schemes

Read naming is a long story with lots of historical gotchas: it needs to be clear and simple, but still people sometimes wanted to convey additional meta-information with it. Unsurprisingly, several "standards" emerged over time. In short: it's a mess. See also XKCD entry on [proliferating standards](#).

How to choose: please read the documentation available at the different centres or ask your sequence provider. In a nutshell (and probably over-simplified):

Sanger scheme "somename.*[pqsfwr][12][bckdeflmnpt][ab|cl]..*" (e.g. U13a08f10.p1ca), but the length of the postfix must be at least 4 characters, i.e., ".p" alone will not be recognised.

Usually, ".p" + 3 characters or ".f" + 3 characters are used for forwards reads, while reverse complement reads take either ".q" or ".r" (+ 3 characters in both cases).

TIGR scheme "somename*TF*|TR*|TA**" (e.g. GCPBN02TF or GCPDL68TABRPT103A58B),

Forward reads take "TF*", reverse reads "TR*".

St. Louis scheme "somename.*[sfrxzyingtpedca]**"

Forward/Reverse scheme "somename.*[fr]**" (e.g. E0K6C4E01DIGEW.f or E0K6C4E01BNDXN.r2nd),

".f*" for forward, ".r*" for reverse.

Solexa scheme Even simpler than the forward/reverse scheme, it allows only for one two reads per template: "somename/*[12]*"

SRA scheme The Short Read Archive (SRA) finally settled on a naming scheme and renames each and every read within its database. When you download sequences from the archive, all reads will be named `XXX000000.Y[Z]` (where X's are characters A-Z, 0 are digits from 0 to 9, Y is a counter and Z is a number denoting the segment (usually 1,2 or 3)). This naming scheme is applied to reads from all technologies, therefore the MIRA technology dependent defaults will not apply and one must specify the 'sra' naming scheme in the command line.

Any wildcard in the forward/reverse suffix must be consistent for a read pair, and is treated as part of the template name. This is to allow multiple sequencing of a fragment, particularly common with Sanger capillary data (e.g. given `somename.f` and `somename.r`, resequenced as `somename.f2` and `somename.r2`, this would be treated as two pairs, with template names `somename` and `somename_2` respectively).

9.3.3.11 Strain naming

strain_name = *string*. Defines the strain / organism-code the reads of this read group are from. If not set, MIRA will assign "StrainX" to normal readgroups and "ReferenceStrain" to readgroups with reference sequences.

Restrictions: in de-novo assemblies you can have 255 strain. In mapping assemblies, you can have at most 8 strains.

Understanding how MIRA uses strain information

bla

9.3.3.12 Data directory for SCF files

datadir_scf = *directory*

For SANGER data only: tells MIRA in which directory it can find SCF data belonging to reads of this read group.

9.3.3.13 Renaming read name prefixes

rename_prefix= *prefix replacement*. Allows to rename reads on the fly while loading data by searching each read name for a given *prefix* string and, if found, replace it with a given *replacement* string.

This is most useful for systems like Illumina or PacBio which generate quite long read names which, in the end, are either utterly useless for an end user or are even breaking older programs which have a length restriction on read names. E.g.:

```
rename_prefix = DQT9AAQ4:436:H371HABMM: Sample1_
```

will rename reads like `DQT9AAQ4:436:H371HABMM:5:1101:9154:3062` into `Sample1_5:1101:9154:3062`

Note `rename_prefix` entries are valid per readgroup. I.e., an entry for a readgroup will not rename reads of another readgroup.

Note

Multiple `rename_prefix` entries are allowed per readgroup. E.g.:

```
rename_prefix = DQT9AAQ4:436:H371HABMM: S1sxa_
rename_prefix = m140328_002546_42149_c100624422550000001823118308061414_s1_ S1pb_
```

will rename a read called `DQT9AAQ4:436:H371HABMM:1:1101:3099:2186` into `S1sxa_1:1101:3099:2186` while renaming another read called `m140328_002546_42149_c100624422550000001823118308061414_s1_p0/100084/10792_20790/0_9573` into `S1pb_p0/100084/10792_20790/0_9573`

9.3.4 The manifest file: extended parameters

The **parameters=** line in the manifest file opens up the full panoply of possibilities the MIRA assembler offers. This ranges from fine-tuning assemblies to setting parameters in a way so that MIRA is suited also for very special assembly cases.

9.3.4.1 Parameter groups

Some parameters one can set in MIRA somehow belong together. Example given: when specifying an overlap in an alignment of two sequences, one could tell the assembler it should look at overlaps only if they have a certain similarity and a certain length. On the other hand, specifying how many processors / threads the assembler should use or whether the results of an assembly should be written out as SAM format does not seem to relate to alignments.

MIRA uses *parameter groups* to keep parameters together which somehow belong together. Example given:

```
parameters = -GENERAL:number_of_threads=4 \  
             -ALIGN:min_relative_score=70 -ASSEMBLY:minimum_read_length=150 \  
             -OUTPUT:output_result_caf=no
```

The parameters of the different parameter groups are described in detail a bit later in this manual.

9.3.4.2 Technology sections

With the introduction of new sequencing technologies, MIRA also had to be able to set values that allow technology specific behaviour of algorithms. One simple example for this could be the minimum length a read must have to be used in the assembly. For Sanger sequences, having this value to be 150 (meaning a read should have at least 150 unclipped bases) would be a very valid, albeit conservative choice. For 454 reads and especially Solexa and ABI SOLiD reads however, this value would be ridiculously high.

To allow very fine grained behaviour, especially in hybrid assemblies, and to prevent the explosion of parameter names, MIRA knows two categories of parameters:

1. **technology independent parameters** which control general behaviour of MIRA like, e.g., the number of assembly passes or file names etc.
2. **technology dependent parameters** which control behaviour of algorithms where the sequencing technology plays a role. Example for this would be the minimum length of a read (like 200 for Sanger reads and 120 for 454 FLX reads).

More on this a bit further down in this documentation.

As example, a manifest using technology dependent and independent parameters could look like this:

```
parameters = COMMON_SETTINGS -GENERAL:number_of_threads=4 \  
             SANGER_SETTINGS -ALIGN:min_relative_score=70 -ASSEMBLY:minimum_read_length ↔  
             =150 \  
             454_SETTINGS -ALIGN:min_relative_score=75 -ASSEMBLY:minimum_read_length=100 \  
             SANGER_SETTINGS -ALIGN:min_relative_score=90 -ASSEMBLY:minimum_read_length=75
```

Now, assume the following read group descriptions in a manifest:

```
...  
  
readgroup  
technology=454  
...  
  
readgroup  
technology=solexa  
...
```

For MIRA, this means a number of parameters should apply to the assembly as whole, while others apply to the sequencing data itself ... and some parameters might need to be different depending on the technology they apply to. MIRA dumps the parameters it is running with at the beginning of an assembly and it makes it clear there which parameters are "global" and which parameters apply to single technologies.

Here is as example a part of the output of used parameters that MIRA will show when started with 454 and Illumina (Solexa) data:

```

...
Assembly options (-AS):
  Number of passes (nop)           : 1
  Skim each pass (sep)            : yes
  Maximum number of RMB break loops (rbl) : 1
  Spoiler detection (sd)          : no
  Last pass only (sdlpo)         : yes

  Minimum read length (mrl)       : [454] 40
                                   [sxa] 20
  Enforce presence of qualities (epoq) : [454] no
                                   [sxa] yes
...

```

You can see the two different kind of settings that MIRA uses: *common settings* (like [-AS:nop]) which allows only one value and *technology dependent settings* (like [-AS:mrl]), where for each sequencing technology used in the project, the setting can be different.

How would one set a minimum read length of 40 and not enforce presence of base qualities for Sanger reads, but for 454 reads a minimum read length of 30 and enforce base qualities? The answer:

```

job=denovo,genome,draft
parameters= SANGER_SETTINGS -AS:mrl=40:epoq=no 454_SETTINGS -AS:mrl=40:epoq=yes

```

Notice the `..._SETTINGS` section in the command line (or parameter file): these tell MIRA that all the following parameters until the advent of another switch are to be set specifically for the said technology.

Note

For improved readability, you can distribute parameters across several lines either by pre-fixing every line with `parameter=`, like so:

```

job=denovo,genome,draft
parameter= SANGER_SETTINGS -AS:mrl=80:epoq=no
parameter= 454_SETTINGS -AS:mrl=30:epoq=yes

```

Alternatively you can use a backslash at the end of a parameter line to indicate that the next line is a continuing line, like so:

```

job=denovo,genome,draft
parameter= SANGER_SETTINGS -AS:mrl=80:epoq=no \
          454_SETTINGS -AS:mrl=30:epoq=yes

```

Note that the very last line of the parameters settings **MUST NOT** end with a backslash.

Beside `COMMON_SETTINGS` there are currently 6 technology settings available:

1. `SANGER_SETTINGS`
 2. `454_SETTINGS`
 3. `IONTOR_SETTINGS`
 4. `PCBIOLQ_SETTINGS` (currently not supported)
 5. `PCBIOHQ_SETTINGS`
 6. `SOLEXA_SETTINGS`
 7. `TEXT_SETTINGS`
-

Some settings of MIRA are influencing global behaviour and are not related to a specific sequencing technology, these must be set in the `COMMON_SETTINGS` environment. For example, it would not make sense to try and set different number of assembly passes for each technology like in

```
parameters= 454_SETTINGS -AS:nop=4 SOLEXA_SETTINGS -AS:nop=3
```

Beside being contradictory, this makes not really sense. MIRA will complain about cases like these. Simply set those common settings in an area prefixed with the `COMMON_SETTINGS` switch like in

```
parameters= COMMON_SETTINGS -AS:nop=4 454_SETTINGS ... SOLEXA_SETTINGS ...
```

Since MIRA 3rc3, the parameter parser will help you by checking whether parameters are correctly defined as `COMMON_SETTINGS` or technology dependent setting.

9.3.4.3 Parameter short names

Writing the verbose form of parameters can be quite a long task. Here a short example:

```
parameters = COMMON_SETTINGS -GENERAL:number_of_threads=4 \
             SANGER_SETTINGS -ALIGN:min_relative_score=70 -ASSEMBLY:minimum_read_length ↔
             =150 \
             454_SETTINGS -ALIGN:min_relative_score=75 -ASSEMBLY:minimum_read_length=100 \
             SOLEXA_SETTINGS -ALIGN:min_relative_score=90 -ASSEMBLY:minimum_read_length=75
```

However, every parameter has a shortened form. The above could be written like this:

```
parameters = COMMON_SETTINGS -GE:not=4 \
             SANGER_SETTINGS -AL:mrs=70 -AS:mrl=150 \
             454_SETTINGS -AL:mrs=75 -AS:mrl=100 \
             SOLEXA_SETTINGS -AL:mrs=90 -AS:mrl=75
```

Please note that it is also perfectly legal to decompose the switches so that they can be used more easily in scripted environments (notice the multiple `-AL` in some sections of the following example):

```
parameters = COMMON_SETTINGS -GE:not=4 \
             SANGER_SETTINGS \
             -AL:mrs=70 \
             -AL:mrl=150 \
             454_SETTINGS -AL:mrs=75:mrl=100 \
             SOLEXA_SETTINGS \
             -AL:mrs=90 \
             -AL:mrl=75
```

9.3.4.4 Order dependent quick switches

For some parameters, the order of appearance in the parameter lines of the manifest is important. This is because the *quick parameters* are realised internally as a collection of extended parameters that will overwrite any previously manually set extended parameters. It is generally a good idea to place quick parameters in the order as described in this documentation, that is: first the order dependent quick parameters, then other quick parameters, then all the other extended parameters.

[--hirep_best], [--hirep_good], [--hirep_something] These are modifier switches for genome data that is deemed to be highly repetitive. With *hirep_good* and *hirep_best*, the assemblies will run slower due to more iterative cycles and slightly different default parameter sets that give MIRA a chance to resolve many nasty repeats. The *hirep_something* switch goes the other way round and resolves repeats less well than a normal assembly, but allows MIRA to finish even on more complex data.

Usage recommendations bacteria: starting MIRA without any hirep switches yields good enough result in most cases. Under normal circumstances one can use *hirep_good* or even *hirep_best* without remorse as data sets and genome complexities are small enough to run within a couple of hours at most.

Usage recommendations for 'simple' lower eukaryotes: starting MIRA without any hirep switches yields good enough result in most cases. If the genomes are not too complex, using *hirep_good* can be a possibility.

Usage recommendations for lower eukaryotes with complex repeats: starting MIRA without any hirep switches might already take too long or create temporary data files which are too big. For these cases, using *hirep_something* makes MIRA use a parameter set which is targeted as resolving the non-repetitive areas of a genome and additionally all repeats which occur less than 10 times in the genome. Repeats occurring more often will not be resolved, but using the debris information one can recover affected reads and use these with harsh data reduction algorithms (e.g. digital normalisation) to get a glimpse into these.

Note These switches replace the '--highlyrepetitive' switch from earlier versions.

[--noclipping=...] Switching off clipping options. If used as `--noclipping` or `--noclipping=all`, this switches off really everything, both technology dependent and independent switches. Clipping options for technology dependent options be switched off via entries being *sanger*, *454*, *iontor*, *solexa* or *solid*. Multiple entries separated by comma are allowed.

Examples:

1. Switch off 454 and Solexa, but keep technology independent clippings and all clippings for other technologies, (like, e.g., Sanger) `--noclipping=454,solexa`
2. Switch off really everything: `--noclipping` or `--noclipping=all`

Note

Switching off technology independent clippings (`[-CL:pec]`, `[-CL:gbcdc]`, `[-CL:kjd]`) via this switch has been implemented for consistency in MIRA 4.9.6. Prior to this they were kept active, which created a good deal of confusion with a number of users.

As soon as you have any kind of 'real' sequencing data, you really should use at least `[-CL:pec]` and `[-CL:gbcdc]`.

9.3.4.5 Parameter group: -GENERAL (-GE)

General options control the type of assembly to be performed and other switches not belonging anywhere else.

[number_of_threads(not)=0 ≤ integer ≤ 256] Default is 0. Master switch to set the number of threads used in different parts of MIRA.

A value of 0 tells MIRA to set this to the number of available physical cores on the machine it runs on. That is, hyper-threaded "cores" are not counted in as using these would cause a tremendous slowdown in the heavy duty computation parts. E.g., a machine with 2 processors having 4 cores each will have this value set to 8.

In case MIRA cannot find out the number of cores, the fall-back value is 2.

Note: when running the SKIM algorithm in parallel threads, MIRA can give different results when started with the same data and same arguments. While the effect could be averted for SKIM, the memory cost for doing so would be an additional 50% for one of the large tables, so this has not been implemented at the moment. Besides, at the latest when the Smith-Watermans run in parallel, this could not be easily avoided at all.

[automatic_memory_management(amm)=only[es]|t[rue], off|n[o]|f[alse]] Default is Yes. Whether MIRA tries to optimise run time of certain algorithms in a space/time trade-off memory usage, increasing or reducing some internal tables as memory permits.

Note 1: This functionality currently relies on the `/proc` file system giving information on the system memory ("MemTotal" in `/proc/meminfo`) and the memory usage of the current process ("VmSize" in `/proc/self/status`). If this is not available, the functionality is switched off.

Note 2: The automatic memory management can only work if there actually is unused system memory. It's not a wonder switch which reduces memory consumption. In tight memory situations, memory management has no effect and the algorithms fall back to minimum table sizes. This means that the effective size in memory can grow larger than given in the memory management parameters, but then MIRA will try to keep the additional memory requirements to a minimum.

[max_process_size(mps)=0 ≤ integer] Default is 0. If automatic memory management is used (see above), this number is the size in gigabytes that the MIRA process will use as maximum target size when looking for space/time trade-offs. A value of 0 means that MIRA does not try keep a fixed upper limit.

Note: when in competition to [-GE:kpmf] (see below), the smaller of both sizes is taken as target. Example: if your machine has 64 GiB but you limit the use to 32 GiB, then the MIRA process will try to stay within these 32 GiB.

[keep_percent_memory_free(kpmf)=0 ≤ integer] Default is 10. If automatic memory management is used (see above), this number works a bit like [-GE:mps] but the other way round: it tries to keep x percent of the memory free.

Note: when in competition to [-GE:mps] (see above), the argument leaving the most memory free is taken as target. Example: if your machine has 64 GiB and you limit the use to 42 GiB via [-GE:mps] but have a [-GE:kpmf] of 50, then the MIRA process will try to stay within $64 - (64 * 50\%) = 32$ GiB.

[preprocess_only(ppo)=on/y[es]/t[rue], off/n[o]/f[alse]] Default is no. As a special use case, MIRA will just run the following tasks: loading and clipping of reads as well as calculating kmer frequencies and read repeat information. The resulting reads can then be found as MAF file in the checkpoint directory; the read repeat information in the info directory.

No assembly is performed.

[est_snp_pipeline_step(esps)=1 ≤ integer ≤ 4] Default is 1. Controls the starting step of the SNP search in EST pipeline and is therefore only useful in miraSearchESTSNPs.

EST assembly is a three step process, each with different settings to the assembly engine, with the result of each step being saved to disk. If results of previous steps are present in a directory, one can easily "play around" with different setting for subsequent steps by reusing the results of the previous steps and directly starting with step two or three.

[print_date(pd)=on/y[es]/t[rue], off/n[o]/f[alse]] Default is yes. Controls whether date and time are printed out during the assembly. Suppressing it is not useful in normal operation, only when debugging or benchmarking.

[bang_on_throw(bot)=on/y[es]/t[rue], off/n[o]/f[alse]] Default is no. For debugging purposes only. Controls whether MIRA raises a signal when detecting an error which triggers a running debugger like gdb.

9.3.4.6 Parameter group: -ASSEMBLY (-AS)

General options for controlling the assembly.

[num_of_passes(nop)=integer ≥ 0] Default is 0. Defines how many iterations of the whole assembly process are done.

The default of 0 will let MIRA choose automatically the number of passes and the kmer sizes used in each pass (see also [-AS:kms] below).

Early termination: if the number of passes was chosen too high, one can simply create a file `projectname_assembly/projectname_chkpt/terminate`. At the beginning of a new pass, MIRA checks for the existence of that file and, if it finds it, acknowledges by renaming it to `terminate_acknowledged` and then run 2 more passes (with special "last pass routines") before finishing the assembly.

Note As a rule of thumb, *de-novo* assemblies should always have at least two passes, while *mapping* assemblies should work with only one pass. Not doing this will lead to results unexpected by users. The reason is that the MIRA the learning routines either have no chance to learn enough about the assembly (for *de-novo* with one pass) or learn "too much" (*mapping* with more than one pass).



Warning MIRA versions ≤ 4.0.2 were interpreting the value of '0' in a different way and only performed pre-processing of reads. MIRA can still do this, but this is controlled by the new parameter [-GE:ppo].

[kmer_series(kms)=comma separated list of integers ≥ 0 and ≤ 256] Default is an empty value. If set, overrides [-AS:nop] and [-SK:kms].

If set, this parameter provides a one-stop-shop for defining the number of passes and the kmer size used in each pass. E.g.: -AS:kms=17, 31, 63, 127 defines an assembly with 4 passes which uses a kmer size of 17 in pass 1, 31 in pass 2, 63 in pass 3 and 127 in pass 4.

Note that it is perfectly valid to use the same kmer size more than once, e.g.: 17, 31, 63, 63 will perform a 4 pass assembly, using a kmer size of 63 in passes 3 and 4. It also makes sense to do this, as with default parameters MIRA uses its integrated automatic editor which edits away obvious sequencing errors in each step, thus the second pass with a kmer size of 63 bases can rely on improved reads.

[rmb_break_loops(rbl)=integer > 0] Default is dependent of the sequencing technology and assembly quality level. Defines the maximum number of times a contig can be rebuilt during a main assembly pass (see [-AS:nop] or [-AS:kms]) if misassemblies due to possible repeats are found.

[max_contigs_per_pass(mcpp)=integer] Default is 0. Defines how many contigs are maximally built in each pass. A value of 0 stands for 'unlimited'. Values >0 can be used for special use cases like test assemblies etc.

If in doubt, do not touch this parameter.

[automatic_repeat_detection(ard)=only[es]|t[rue], off|n[o]|f[alse]] Default is currently yes. Tells MIRA to use coverage information accumulated over time to more accurately pinpoint reads that are in repetitive regions.

[coverage_threshold(ardct)=float > 1.0] Default is 2.0 for all sequencing technologies in most assembly cases. This option says this: if MIRA a read has ever been aligned at positions where the total coverage of all reads of the same sequencing technology attained the average coverage times [-AS:ardct] (over a length of [-AS:ardml], see below), then this read is considered to be repetitive.

[min_length(ardml)=integer > 1] Default is dependent of the sequencing technology, currently 400 for Sanger and 200 for 454 and Ion Torrent.

A coverage must be at least this number of bases higher than [-AS:ardct] before being really treated as repeat.

[grace_length(ardgl)=integer > 1] Default is dependent of the sequencing technology.

[uniform_read_distribution(urd)=only[es]|t[rue], off|n[o]|f[alse]] Default is currently always no as these algorithms were supplanted by better ones in MIRA 5.0.

Takes effect only if uniform read distribution ([-AS:urd]) is on.

When set to yes, MIRA will analyse coverage of contigs built at a certain stage of the assembly and estimate an average expected coverage of reads for contigs. This value will be used in subsequent passes of the assembly to ensure that no part of the contig gets significantly more read coverage of reads that were previously identified as repetitive than the estimated average coverage allows for.

This switch is useful to disentangle repeats that are otherwise 100% identical and generally allows to build larger contigs. It is expected to be useful for Sanger and 454 sequences. Usage of this switch with Solexa and Ion Torrent data is currently not recommended.

It is a real improvement to disentangle repeats, but has the side-effect of creating some "contig debris" (small and low coverage contigs, things you normally can safely throw away as they are representing sequence that already has enough coverage).

This switch must be set to no for EST assembly, assembly of transcripts etc. It is recommended to also switch this off for mapping assemblies.

[urd_startinpass(urdsip)=integer > 0] Default is dependent of the sequencing technology and assembly quality level. Recommended values are: 3 for an assembly with 3 to 4 passes ([-AS:nop]). Assemblies with 5 passes or more should set the value to the number of passes minus 2.

Takes effect only if uniform read distribution ([-AS:urd]) is on.

[urd_clipoffmultiplier(urdcn)=float > 1.0] Default is 1.5 for all sequencing technologies in most assembly cases.

This option says this: if MIRA determined that the average coverage is x , then in subsequent passes it will allow coverage for reads determined to be repetitive to be built into the contig only up to a total coverage of $x*urdcn$. Reads that bring the

coverage above the threshold will be rejected from that specific place in the contig (and either be built into another copy of the repeat somewhere else or end up as contig debris).

Please note that the lower [-AS:urdcn] is, the more contig debris you will end up with (contigs with an average coverage less than half of the expected coverage, mostly short contigs with just a couple of reads).

Takes effect only if uniform read distribution (-AS:urd) is on.

[spoiler_detection(sd)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology and assembly quality level. A spoiler can be either a chimeric read or it is a read with long parts of unclipped vector sequence still included (that was too long for the [-CL:pvc] vector leftover clipping routines). A spoiler typically prevents contigs to be joined, MIRA will cut them back so that they represent no more harm to the assembly.

Recommended for assemblies of mid- to high-coverage genomic assemblies, not recommended for assemblies of ESTs as one might lose splice variants with that.

A minimum number of two assembly passes (-AS:nop) must be run for this option to take effect.

[sd_last_pass_only(sdlpo)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Defines whether the spoiler detection algorithms are run only for the last pass or for all passes (-AS:nop).

Takes effect only if spoiler detection (-AS:sd) is on. If in doubt, leave it to 'yes'.

[minimum_read_length(mrl)=integer ≥ 20] Default is dependent of the sequencing technology. Defines the minimum length that reads must have to be considered for the assembly. Shorter sequences will be filtered out at the beginning of the process and won't be present in the final project.

[minimum_reads_per_contig(mrpc)=integer ≥ 1] Default is dependent of the sequencing technology and the [--job] parameter. For genome assemblies it's usually around 2 for Sanger, 5 for 454, 5 for Ion Torrent, 5 for PacBio and 10 for Solexa. In EST assemblies, it's currently 2 for all sequencing technologies.

Defines the minimum number of reads a contig must have before it is built or saved by MIRA. Overlap clusters with less reads than defined will not be assembled into contigs but reads in these clusters will be immediately transferred to debris.

This parameter is useful to considerably reduce assembly time in large projects with millions of reads (like in Solexa projects) where a lot of small "junk" contigs with contamination sequence or otherwise uninteresting data may be created otherwise.

Note Important: a value larger 1 of this parameter interferes with the functioning of [-OUT:sssip] and [-OUT:stsip].

[enforce_presence_of_qualities(epoq)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. When set to yes, MIRA will stop the assembly if any read has no quality values loaded.

Note [-AS:epoq] switches on/off the quality check for a complete sequencing technology. A more fine grained control for switching checks of per readgroup is available via the *default_qual* readgroup parameter in the manifest file.

[use_genomic_pathfinder(ugpf)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. MIRA has two different pathfinder algorithms it chooses from to find its way through the (more or less) complete set of possible sequence overlaps: a genomic and an EST pathfinder. The genomic looks a bit into the future of the assembly and tries to stay on safe grounds using a maximum of information already present in the contig that is being built. The EST version on the contrary will directly jump at the complex cases posed by very similar repetitive sequences and try to solve those first and is willing to fall back to first-come-first-served when really bad cases (like, e.g., coverage with thousands of sequences) are encountered.

Generally, the genomic pathfinder will also work quite well with EST sequences (but might get slowed down a lot in pathological cases), while the EST algorithm does not work so well on genomes. If in doubt, leave on yes for genome projects and set to no for EST projects.

[use_emergency_search_stop(uess)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Another important switch if you plan to assemble non-normalised EST libraries, where some ESTs may reach coverages of several hundreds or thousands of reads. This switch lets MIRA save a lot of computational time when aligning those extremely high coverage areas (but only there), at the expense of some accuracy.

[ess_partnerdepth(esspd)=integer > 0] Default is 500. Defines the number of potential partners a read must have for MIRA switching into emergency search stop mode for that read.

[use_max_contig_buildtime(umcibt)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. Defines whether there is an upper limit of time to be used to build one contig. Set this to yes in EST assemblies where you think that extremely high coverages occur. Less useful for assembly of genomic sequences.

[buildtime_in_seconds(bts)=integer > 0] Default is 3600 for genome assemblies, 720 for EST assemblies with Sanger or 454 and 360 for EST assemblies with Solexa or Ion Torrent. Depending on [-AS:umcibt] above, this number defines the time in seconds allocated to building one contig.

9.3.4.7 Parameter group: -STRAIN/BACKBONE (-SB)

Controlling backbone options in mapping assemblies:

[bootstrap_new_backbone(bnb)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for mapping assemblies with Illumina data, no otherwise.

When set to 'yes', MIRA will use a two stage mapping process which bootstraps an intermediate backbone (reference) sequence and greatly improves mapping accuracy at indel sites.

Note Currently only works with Illumina data, other sequencing technologies will not be affected by this flag.

[startbackboneusage_inpass(sbuip)=0 < integer] Default is dependent on assembly quality level chosen: 0 for 'draft' and [-AS:nop] divided by 2 for 'accurate'.

When assembling against backbones, this parameter defines the pass iteration (see [-AS:nop]) from which on the backbones will be really used. In the passes preceding this number, the non-backbone reads will be assembled together as if no backbones existed. This allows MIRA to correctly spot repetitive stretches that differ by single bases and tag them accordingly. Note that full assemblies are considerably slower than mapping assemblies, so be careful with this when assembling millions of reads.

Rule of thumb: if backbones belong to same strain as reads to assemble, set to 1. If backbones are a different strain, then set [-SB:sbuip] to 1 lower than [-AS:nop] (example: nop=4 and sbuip=3).

[backbone_raillength(brl)=0 ≤ integer ≤ 10000] Default is 0. Parameter for the internal sectioning size of the backbone to compute optimal alignments. Should be set to two times length of longest read in input data + 15%. When set to 0, MIRA will compute optimal values from the data loaded.

[backbone_railoverlap(bro)=0 ≤ integer ≤ 2000] Default is 0. Parameter for the internal sectioning size of the backbone to compute optimal alignments. Should be set to length of the longest read. When set to 0, MIRA will compute optimal values from the data loaded.

[trim_overhanging_reads(tor)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes.

When set to 'yes', MIRA will trim back reads at end of contigs which outgrow the reference sequence so that boundaries of the reference and the mapped reads align perfectly. That is, the mapping does not perform a sequence extension.

Note The trimming is performed via setting low quality cutoffs in the reads, i.e., the trimmed parts are not really gone but just not part of the active contig anymore. They can be uncovered when working on the assembly in finishing programs like, e.g., **gap4** or **gap5**.



Warning Previous versions of MIRA (up to and including 3.9.18) behaved as if this option had been set to 'no'. This is a major change in behaviour, but it is also what probably most people expect from a mapping.

9.3.4.8 Parameter group: -DATAPROCESSING (-DP)

Options for controlling some data processing during the assembly.

[use_read_extension(ure)=only[es]|t[rue], off/n[o]|f[alse]] Default is dependent of the sequencing technology used: yes for Sanger, no for all others. MIRA expects the sequences it is given to be quality clipped. During the assembly though, it will try to extend reads into the clipped region and gain additional coverage by analysing Smith-Waterman alignments between reads that were found to be valid. Only the right clip is extended though, the left clip (most of the time containing sequencing vector) is never touched.

[read_extension_window_length(rewl)=integer > 0] Default is dependent of the sequencing technology used. Only takes effect when [-DP:ure] (see above) is set to yes. The read extension routines use a sliding window approach on Smith-Waterman alignments. This parameter defines the window length.

[read_extension_with_maxerrors(rewme)=integer > 0] Default is dependent of the sequencing technology used. Only takes effect when [-DP:ure] (see above) is set to yes. The read extension routines use a sliding window approach on Smith-Waterman alignments. This parameter defines the number maximum number of errors (=disagreements) between two alignment in the given window.

[first_extension_in_pass(feip)=integer ≥ 0] Default is dependent of the sequencing technology used. Only takes effect when [-DP:ure] (see above) is set to yes. The read extension routines can be called before assembly and/or after each assembly pass (see [-AS:nop]). This parameter defines the first pass in which the read extension routines are called. The default of 0 tells MIRA to extend the reads the first time before the first assembly pass.

[last_extension_in_pass(leip)=integer ≥ 0] Default is dependent of the sequencing technology used. Only takes effect when [-DP:ure] (see above) is set to yes. The read extension routines can be called before assembly and/or after each assembly pass (see [-AS:nop]). This parameter defines the last pass in which the read extension routines are called. The default of 0 tells MIRA to extend the reads the last time before the first assembly pass.

9.3.4.9 Parameter group: -CLIPPING (-CL)

Controls for clipping options: when and how sequences should be clipped.

Every option in this section can be set individually for every sequencing technology, giving a very fine grained control on how reads are clipped for each technology.

[msvs_gap_size(msvsgs)=integer ≥ 0] Default is dependent of the sequencing technology used. Takes effect only when loading data from ancillary SSAHA2 or SMALT files.

While performing the clip of screened vector sequences, MIRA will look if it can merge larger chunks of sequencing vector bases that are a maximum of [-CL:msvsgs] apart.

[msvs_max_front_gap(msvsmfg)=integer ≥ 0] Default is dependent of the sequencing technology used. Takes effect only when loading data from ancillary SSAHA2 or SMALT files.

While performing the clip of screened vector sequences at the start of a sequence, MIRA will allow up to this number of non-vector bases in front of a vector stretch.

[msvs_max_end_gap(msvsmeg)=integer ≥ 0] Default is dependent of the sequencing technology used. Takes effect only when loading data from ancillary SSAHA2 or SMALT files.

While performing the clip of screened vector sequences at the end of a sequence, MIRA will allow up to this number of non-vector bases behind a vector stretch.

[possible_vector_leftover_clip(pvlc)=only[es]|t[rue], off/n[o]|f[alse]] Default is dependent of the sequencing technology used: yes for Sanger, no for any other. MIRA will try to identify possible sequencing vector relics present at the start of a sequence and clip them away. These relics are usually a few bases long and were not correctly removed from the sequence in data preprocessing steps of external programs.

You might want to turn off this option if you know (or think) that your data contains a lot of repeats and the option below to fine tune the clipping behaviour does not give the expected results.

You certainly want to turn off this option in EST assemblies as this will quite certainly cut back (and thus hide) different splice variants. But then make certain that you pre-processing of Sanger data (sequencing vector removal) is good, other sequencing technologies are not affected then.

[pvc_maxlenallowed(pvcmla)=integer ≥ 0] Default is dependent of the sequencing technology used. The clipping of possible vector relics option works quite well. Unfortunately, especially the bounds of repeats or differences in EST splice variants sometimes show the same alignment behaviour than possible sequencing vector relics and could therefore also be clipped.

To refrain the vector clipping from mistakenly clip repetitive regions or EST splice variants, this option puts an upper bound to the number of bases a potential clip is allowed to have. If the number of bases is below or equal to this threshold, the bases are clipped. If the number of bases exceeds the threshold, the clip is **NOT** performed.

Setting the value to 0 turns off the threshold, i.e., clips are then always performed if a potential vector was found.

[quality_clip(qc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is **no**. This will let MIRA perform its own quality clipping before sequences are entered into the assembly. The clip function performed is a sequence end window quality clip with back iteration to get a maximum number of bases as useful sequence. Note that the bases clipped away here can still be used afterwards if there is enough evidence supporting their correctness when the option [-DP:ure] is turned on.

Warning: The windowing algorithm works pretty well for Sanger, but apparently does not like 454 type data. It's advisable to not switch it on for 454. Beside, the 454 quality clipping algorithm performs a pretty decent albeit not perfect job, so for genomic 454 data (not! ESTs), it is currently recommended to use a combination of [-CL:emrc] and [-DP:ure].

[qc_minimum_quality(qcmq)=integer ≥ 15 and ≤ 35] Default is dependent of the sequencing technology used. This is the minimum quality bases in a window require to be accepted. Please be cautious not to take too extreme values here, because then the clipping will be too lax or too harsh. Values below 15 and higher than 30-35 are not recommended.

[qc_window_length(qcwl)=integer ≥ 10] Default is dependent of the sequencing technology used. This is the length of a window in bases for the quality clip.

[bad_stretch_quality_clip(bsqc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is **no**. This option allows to clip reads that were not correctly preprocess and have unclipped bad quality stretches that might prevent a good assembly.

MIRA will search the sequence in forward direction for a stretch of bases that have in average a quality less than a defined threshold and then set the right quality clip of this sequence to cover the given stretch.

[bsqc_minimum_quality(bsqcmq)=integer ≥ 0] Default is dependent of the sequencing technology used. Defines the minimum average quality a given window of bases must have. If this quality is not reached, the sequence will be clipped at this position.

[bsqc_window_length(bsqcwl)=integer ≥ 0] Default is dependent of the sequencing technology used. Defines the length of the window within which the average quality of the bases are computed.

[maskedbases_clip(mbc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used. This will let MIRA perform a 'clipping' of bases that were masked out (replaced with the character X). It is generally not a good idea to use mask bases to remove unwanted portions of a sequence, the EXP file format and the NCBI traceinfo format have excellent possibilities to circumvent this. But because a lot of preprocessing software are built around cross_match, scylla- and phrap-style of base masking, the need arose for MIRA to be able to handle this, too. MIRA will look at the start and end of each sequence to see whether there are masked bases that should be 'clipped'.

[mbc_gap_size(mbcgs)=integer ≥ 0] Default is dependent of the sequencing technology used. While performing the clip of masked bases, MIRA will look if it can merge larger chunks of masked bases that are a maximum of [-CL:mbcgs] apart.

[mbc_max_front_gap(mbcmfg)=integer ≥ 0] Default is dependent of the sequencing technology used. While performing the clip of masked bases at the start of a sequence, MIRA will allow up to this number of unmasked bases in front of a masked stretch.

[mbc_max_end_gap(mbcmeg)=integer ≥ 0] Default is dependent of the sequencing technology used. While performing the clip of masked bases at the end of a sequence, MIRA will allow up to this number of unmasked bases behind a masked stretch.

[lowercase_clip_front(lccf)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used: on for 454 and Ion Torrent data, off for all others. This will let MIRA perform a 'clipping' of bases that are in lowercase at the front end of a sequence, leaving only the uppercase sequence. Useful when handling 454 data that does not have ancillary data in XML format.

[lowercase_clip_back(lccb)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used: on for 454 and Ion Torrent data, off for all others. This will let MIRA perform a 'clipping' of bases that are in lowercase at the back end of a sequence, leaving only the uppercase sequence. Useful when handling 454 data that does not have ancillary data in XML format.

[clip_polyat(cpat)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for all EST/RNASeq assemblies. Poly-A stretches in forward reads and poly-T stretches in reverse reads get either clipped or tagged here (see [-CL:cpkps] below). The assembler will not use these stretches for finding overlaps, but it will use these to discern and disassemble different 3' UTR endings.



Warning Should poly-A / poly-T stretches have been trimmed in pre-processing steps before MIRA got the reads, this option **MUST** be switched off.

[cp_keep_poly_stretch (cpkps)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes but takes effect only if [-CL:cpat] (see above) is also set to yes.

Instead of clipping the poly-A / poly-T sequence away, the stretch in question in the reads is kept and tagged. The tags provide additional information for MIRA to discern between different 3' UTR endings and also a good visual anchor when looking at the assembly with different programs.

Note One side-effect of this option is that the poly-A / poly-T stretch are 'cleaned'. That is, single non-poly A / poly-T bases within the stretch are automatically edited to be conforming to the surrounding stretch. This is necessary as homopolymers are by nature one of the hardest motifs to be sequenced correctly by any sequencing technology and one frequently gets 'dirty' poly-A sequence from sequencing and this interferes heavily with the methods MIRA uses to discern repeats.

Note Keeping the poly-A sequence is a two-edged sword: on one hand it enabled to discern different 3' UTR endings, on the other hand it might be that sequencing problems toward the end of reads create false-positive different endings. If you find that this is the case for your data, just switch off this option: MIRA will then simply build the longest possible 3' UTRs.

[cp_min_sequence_len(cpmsl)=integer > 0] Default is 10. Only takes effect when [-CP:cpat] (see above) is set to yes. Defines the number of 'A' (in forward direction) or 'T' (in reverse direction) must be present to be considered a poly-A sequence stretch.

[cp_max_errors_allowed(cpmea)=integer > 0] Default is 1. Only takes effect when [-CL:cpat] (see above) is set to yes. Defines the maximum number of errors allowed in the potential poly-A sequence stretch. The distribution of these errors is not important.

[cp_max_gap_from_end(cpmgfe)=integer > 0] Default is 9. Only takes effect when [-CL:cpat] (see above) is set to yes. Defines the number of bases from the end of a sequence (if masked: from the end of the masked area) within which a poly-A sequence stretch is looked for.

[clip_3ppolybase (c3pp)=on|y[es]|t[rue], off|n[o]|f[alse]] c3p* options to be described ...

[clip_known_adaptorsright (ckar)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Defines whether MIRA should search and clip known sequencing technology specific sequencing adaptors. MIRA knows adaptors for Illumina best, followed by Ion Torrent and some 454 adaptors.

As the list of known adaptors changes quite frequently, the best place to get a list of known adaptors by MIRA is by looking at the text files in the program sources: src/mira/adaptorsforclip.*.xxd.

[ensure_minimum_left_clip(emlc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used. If on, ensures a minimum left clip on each read according to the parameters in [-CL:mlcr:smc]

[minimum_left_clip_required(mlcr)=integer ≥ 0] Default is dependent of the sequencing technology used. If [-CL:emlc] is on, checks whether there is a left clip which length is at least the one specified here.

[set_minimum_left_clip(smlc)=integer ≥ 0] Default is dependent of the sequencing technology used. If [-CL:emlc] is on and actual left clip is < [-CL:mlcr], set left clip of read to the value given here.

[ensure_minimum_right_clip(emrc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used. If on, ensures a minimum right clip on each read according to the parameters in [-CL:mrcr:smrc]

[minimum_right_clip_required(mrcr)=integer ≥ 0] Default is dependent of the sequencing technology used. If [-CL:emrc] is on, checks whether there is a right clip which length is at least the one specified here.

[set_minimum_right_clip(smrc)=integer ≥ 0] Default is dependent of the sequencing technology used. If [-CL:emrc] is on and actual right clip is < [-CL:mrcr], set the length of the right clip of read to the value given here.

[gb_chimeradetectionclip(gbcdc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for all jobs.

Very safe chimera detection, should have no false positives. For repetitive data, a low number of false negatives is possible.

[kmerjunk_detection(kjd)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is currently yes.

Reads that look "fishy" are marked as potentially chimeric. This mark leads either to a read being completely killed or to a read being included into a contig only if no other possibility remains.

It is currently suggested to leave this parameter switched on and to fine-tune via [-CL:kjck] (see below).

[kmerjunk_completekil(kjck)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is currently no for genome assemblies and yes for EST/RNASEq assemblies.

If set to yes, reads marked as junk (see above) are completely removed from an assembly. If set to no, reads are not removed but included only into a contig as a very last resort.

Having reads killed guarantees assemblies of extremely high quality containing virtually no misassembly due to chimeric sequencing errors. The downside is that, computationally, there is no difference between junk and stretches with correct but very low coverage data (generally < 3x coverage). It's up to you to decide what is more important: total accuracy or longer contigs.

Note

As a rule of thumb: I set this to no for genome assemblies with at least medium average coverage (≥ 20 -30x) as MIRA does a pretty good job to incorporate these reads so late in an assembly that they do not lead to misassemblies. In transcript assemblies I set this to yes as there is a high chance that high coverage transcripts could be extended via chimeric reads.

With this in mind: deciding for metagenome assemblies would be really difficult though. It probably depends on what you need the data for.

[propose_end_clips(pec)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is is dependent on --job settings currently yes for all genome assemblies and for EST/RNASEq assemblies with 454, Illumina or Ion torrent data.

This implements a pretty powerful strategy to ensure a good "high confidence region" (HCR) in reads, basically eliminating 99.9% of all junk at the 5' and 3' ends of reads. Note that one still must ensure that sequencing vectors (Sanger) or adaptor sequences (454, Solexa ion Torrent) are "more or less" clipped prior to assembly.



Warning

Extremely effective, but should NOT be used for very low coverage data, like low coverage genome or rare EST/RNASEq transcripts.

The defaults for EST/RNASEq guarantee high quality transcripts at the expense of losing low coverage / rare transcripts.

[pec_continuous(pec)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is is dependent on --job settings currently yes for all genome assemblies and for EST/RNASeq assemblies with 454, Illumina or Ion torrent data.

This is [-CL:pec] on steroids. The 'simple' -CL:pec is run only twice, with a fixed kmer size, during data preprocessing before performing the assembly. The continuous version additionally run at every pass, effectively taking advantage of larger kmer size and catching more difficult cases.



Warning

Extremely effective, but should NOT be used for very low coverage data, like low coverage genome or rare EST/RNASeq transcripts.

The defaults for EST/RNASeq increase the proportion of high quality transcripts especially with Illumina data as many chimeric sequences are caught which would otherwise remain undetected.

[handle_solexa_ggcxg_problem(pechsgp)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is is dependent yes.

Solexa data has a pretty awful problem with in some reads when a GGCxG motif occurs (read more about it in the chapter on Solexa data). In short: the sequencing errors produced by this problem lead to many false positive SNP discoveries in mapping assemblies or problems in contig building in de-novo assembly.

MIRA knows about this problem and can look for it in Solexa reads during the proposed end clipping and further clip back the reads, greatly minimising the impact of this problem.

[pec_kmer_size(peckms)=10 ≤ integer ≤ 32] Default is is dependent on technology and quality in the --job switch: usually between 17 and 21 for Sanger, higher for 454 (up to 27) and highest for Solexa (31). Ion Torrent has at the moment 17, but this may change in the future to somewhat higher values.

This parameter defines the minimum number of bases at each end of a read that should be free of any sequencing errors.

[pec_minimum_kmer_forward_reverse(pmkfr)=integer ≥ 0] Default is is dependent on technology and quality in the --job switch: usually between 1 and 3 when [-CL:pec=yes].

This parameter defines the minimum number of occurrence of a kmer at each end of a read that should be free of any sequencing errors.

[rare_kmer_mask(rkm)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is is dependent on --job switch: currently it's yes for Solexa data and no otherwise. If this parameter is active, MIRA will completely mask with 'X' those parts of a read which have kmer occurrence (in forward and reverse direction) less than the value specified via [-CL:pmkfr].

This is a quality ensuring move which improves assembly of ultra-high coverage contigs by cleaning out very likely, low frequency sequence dependent sequencing errors which passed all previous filters. The drawback is that very rare transcripts or very lowly covered genome parts with an occurrence less than the given value will also be masked out. However, Illumina gives so much data that this is almost never the case.

Note This works only if [-CL:pec] is active.

[search_phix174(spx174)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is *on* for Illumina data, off otherwise.

PhiX 174 is a small phage of enterobacteria whose DNA is often spiked-in during Illumina sequencing to determine error rates in datasets and to increase complexity in low-complexity samples (amplicon, chipseq etc) to help in cluster identification.

If it remains in the sequenced data, it has to be seen as a contaminant for projects working with organisms which should not contain the PhiX 174 phage.

Note However, PhiX may be part of some genome sequences (enterobacteria). In these cases, the PhiX174 search will report genuine genome data.

[filter_phix174(fpx174)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is *on* for Illumina data in EST (RNASeq) assemblies, off otherwise.

If [-CL:spx174] is on and [-CL:fpx174] also, MIRA will filter out as contaminants all reads which have PhiX174 sequence recognised.

The default value of having the filtering on only for Illumina EST (RNASeq) data is a conservative approach: the overwhelming majority of RNASeq data will indeed not sequence some enterobacteria, so having PhiX174 containing reads thrown out is indeed a valid move. For genomes however, MIRA currently is cautious and will not filter these reads by default.



Warning However, PhiX may be part of some genome sequences (enterobacteria). In these cases, the PhiX174 filter will remove reads from valid genome or expression data.

[filter_rrna(frna)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is *on* for EST (RNASeq) assemblies, off otherwise.

If enabled, MIRA will filter out (and not assemble) all reads (or pairs, see below) it recognises as being rRNA or rDNA. This is useful to reduce computing time on data sets which contain large contamination of rRNA which were not filtered away in wet lab.

[filter_rrna_pairs(frnap)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is *on* for EST (RNASeq) assemblies, off otherwise.

If enabled together with [-CL:frna], MIRA will filter out (and not assemble) all reads pairs where at least one of the reads is recognised as being rRNA or rDNA.

This option is useful to also catch less conserved parts of rRNA transcribed like, e.g. the internal transcribed spacers (ITS) in eukaryotic data.

[filter_rrna_numkmers(frnanck)=integer > 0] Default is 20.

The rRNA recognition in MIRA works with a precompiled set of preserved rRNA kmers, at the time of this writing with 21-mers. To allow for specific recognition, the rRNA filtering process expects to find at least this number of kmers per read before identifying it as rRNA.

To increase sensitivity (and at the same time risk more false positives): reduce this parameter. To increase specificity (and at the same time risk more reads not being recognised): increase this parameter.

The default parameters together with the default database seem to work pretty well and this is expected to work for all but the most exotic rRNA containing organisms.

9.3.4.10 Parameter group: -SKIM (-SK)

Options that control the behaviour of the initial fast all-against-all read comparison algorithm. Matches found here will be confirmed later in the alignment phase. The new SKIM3 algorithm that is in place since version 2.7.4 uses a kmer based algorithm that works similarly to SSAHA (see Ning Z, Cox AJ, Mullikin JC; "SSAHA: a fast search method for large DNA databases."; Genome Res. 2001;11;1725-9).

The major differences of SKIM3 and SSAHA are:

1. the word length *n* of a kmer (hash) in SSAHA2 must be < 15, but can be up to 32 bases in 64 bit versions of MIRA < 4.0.2 and lower, and up to 256 bases for higher versions of MIRA.
2. SKIM3 uses a maximum fixed amount of RAM that is independent of the word size. E.g., SSAHA would need 4 exabyte to work with word length of 30 bases ... SKIM3 just takes a couple of hundred MB.

The parameters for SKIM3:

[number_of_threads(not)=integer ≥ 1] Number of threads used in SKIM, default is 2. A few parts of SKIM are non-threaded, so the speedup is not exactly linear, but it should be very close. E.g., with 2 processors I get a speedup of 180-195%, with 4 between 350 and 395%.

Although the main data structures are shared between the threads, there's some additional memory needed for each thread.

[also_compute_reverse_complements(acrc)=only[es]|t[rue], off[n[o]|f[alse]] Default is on. Defines whether SKIM searches for matches only in forward/forward direction or whether it also looks for forward/reverse direction.

You usually will not want to touch the default, except for very special application cases where you do not want MIRA to use reverse complement sequences at all.

[kmer_size(kms)=10 < integer ≤ 256] Defaults are dependent on "--job" switch and sequencing technologies used.

Controls the number of consecutive bases n which are used as a kmer. The higher the value, the faster the search. The lower the value, the slower the search and the more weak matches are found.

A secondary effect of this parameter is the estimation of MIRA on whether stretches within a read sequence are repetitive or not. Large values of [-SK:kms] allow a better distinction between "almost identical" repeats early in the assembly process and, given enough coverage, generally lead to less and longer contigs.

Note This parameter gets overridden by the one-stop-shop parameter [-AS:kms] which determines number of passes and kmer size to use in each pass.



Warning For de-novo assemblies, values below 15 are not recommended. For mapping assemblies, values below 10 should not be used.

[kmer_save_stepping(kss)=integer ≥ 1] Default is 1. This is a parameter controlling the stepping increment s with which kmers are generated. This allows for more or less fine grained search as matches are found with at least $n+s$ (see [-SK:kms]) equal bases. The higher the value, the faster the search. The lower the value, the more weak matches are found.

[percent_required(pr)=integer ≥ 1] Default is dependent of the sequencing technology used and assembly quality wished. Controls the relative percentage of exact word matches in an approximate overlap that has to be reached to accept this overlap as possible match. Increasing this number will decrease the number of possible alignments that have to be checked by Smith-Waterman later on in the assembly, but it also might lead to the rejection of weaker overlaps (i.e. overlaps that contain a higher number of mismatches).

Note: most of the time it makes sense to keep this parameter in sync with [-AL:mrs].

[maxhits_perread(mhpr)=integer ≥ 1] Default is 2000. Controls the maximum number of possible hits one read can maximally transport to the overlap edge reduction phase. If more potential hits are found, only the best ones are taken.

In the pre-2.9.x series, this was an important option for tackling projects which contain *extreme* assembly conditions. It still is if you run out of memory in the graph edge reduction phase. Try then to lower it to 1000, 500 or even 100.

As the assembly increases in passes ([-AS:nop]), different combinations of possible hits will be checked, always the probably best ones first. So the accuracy of the assembly should only suffer when lowering this number too much.

[filter_megahubs(fmh)=only[es]|t[rue], off[n[o]|f[alse]] Default is yes. Defines whether megahubs (reads with extremely many overlaps to other reads) are filtered. See also [-SK:mhc:mmhr].

[megahub_cap(mhc)=integer > 0] Default is 150000. Defines the number of kmer overlaps a read may have before it is categorised as megahub.

Note You basically don't want to mess with this one. Except for assemblies containing very long reads. Rule of thumb: you might want to multiply the 150k value by n where n is the average read length divided by 2000. Don't overdo, max n at 15 or so.

[max_megahub_ratio(mmhr)=integer ≥ 0] Default is 0. If the number of reads identified as megahubs exceeds the allowed ratio, MIRA will abort.

This is a fail-safe parameter to avoid assemblies where things look fishy. In case you see this, you might want to ask for advice on the `mira_talk` mailing list. In short: bacteria should never have megahubs (90% of all cases reported were contamination of some sort and the 10% were due to incredibly high coverage numbers). Eukaryotes are likely to contain megahubs if filtering is `[-KS:mnr]` not on.

EST project however, especially from non-normalised libraries, will very probably contain megahubs. In this case, you might want to think about masking, see `[-KS:mnr]`.

[sw_check_on_backbones(swcb)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is currently (3.4.0) yes for accurate mapping jobs. Takes effect only in mapping assemblies. Defines whether SKIM hits against a backbone (reference) sequence with less than 100% identity are double checked with Smith-Waterman to improve mapping accuracy.

You will want to set this option to yes whenever your reference sequence contains more complex or numerous repeats and your data has SNPs in those areas.

[max_kmers_in_memory(mkim)=integer ≥ 100000] Default is 15000000. Has no influence on the quality of the assembly, only on the maximum memory size needed during the skimming. The default value is equivalent to approximately 500MB.

Note: reducing the number will increase the run time, the more drastically the bigger the reduction. On the other hand, increasing the default value chosen will not result in speed improvements that are really noticeable. In short: leave this number alone if you are not desperate to save a few MB.

[memcap_hitreduction(mchr)=integer ≥ 10] Default is 1024, 2048 when Solexa sequences are used. Maximum memory used (in MiB) during the reduction of skim hits.

Note: has no influence on the quality of the assembly, reducing the number will increase the runtime, the more drastically the bigger the reduction as hits then must be streamed multiple times from disk.

The default is good enough for assembly of bacterial genomes or small eukaryotes (using Sanger and/or 454 sequences). As soon as assembling something bigger than 20 megabases, you should increase it to 2048 or 4096 (equivalent to 2 or 4 GiB of memory).

9.3.4.11 Parameter group: -KMERSTATISTICS (-KS)

Hash statistics (nowadays called kmer statistics in literature or other software packages) allows to quickly assess reads from a coverage point of view without actually assembling the reads. MIRA uses this as a quick pre-assembly evaluation to find and tag reads which are from repetitive and non-repetitive parts of a project.

The length of the kmer is defined via `[-SK:kms]` or `[-AS:kms]` while the parameters in this section define the boundaries of the different repeat levels.

A more in-depth description on kmer statistics is given in the sections *Introduction to 'masking'* and *How does 'nasty repeat' masking work?* in the chapter dealing with the assembly of hard projects.

[freq_est_minnormal(fenn)=float > 0] During kmer statistics analysis, MIRA will estimate how repetitive parts of reads are. Parts which are occurring less than `[-KS:fenn]` times the average occurrence will be tagged with a HAF2 (less than average) tag.

[freq_est_maxnormal(fexn)=float > 0] During kmer statistics analysis, MIRA will estimate how repetitive parts of reads are. Parts which are occurring more than `[-KS:fenn]` but less than `[-KS:fexn]` times the average occurrence will be tagged with a HAF3 (normal) tag.

[freq_est_repeat(fer)=float > 0] During kmer statistics analysis, MIRA will estimate how repetitive parts of reads are. Parts which are occurring more than `[-KS:fexn]` but less than `[-KS:fer]` times the average occurrence will be tagged with a HAF4 (above average) tag.

[freq_est_heavyrepeat(fehr)=float > 0] During kmer statistics analysis, MIRA will estimate how repetitive parts of reads are. Parts which are occurring more than `[-KS:fer]` but less than `[-KS:fehr]` times the average occurrence will be tagged with a HAF5 (repeat) tag.

[freq_est_crazyrepeat(fecr)=float > 0] During kmer statistics analysis, MIRA will estimate how repetitive parts of reads are. Parts which are occurring more than [-KS:fehr] but less than [-KS:fecr] times the average occurrence will be tagged with a HAF6 (heavy repeat) tag. Parts which are occurring more than [-KS:fecr] but less than [-KS:nrr] times the average occurrence will be tagged with a HAF7 (crazy repeat) tag.

[mask_nasty_repeats(mnr)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent on --job type: yes for de-novo, no for mapping.

Tells MIRA to tag (during the kmer statistics phase) read subsequences of length [-SK:kms] nucleotides that appear more than X times more often than the median occurrence of subsequences would otherwise suggest. The threshold X from which on subsequences are considered nasty is set by [-KS:nrr] or [-KS:nrc], the action MIRA should take when encountering those sequences is defined by [-KS:ldn] (see below).

When not using lossless digital normalisation ([-KS:ldn]), the tag used by MIRA will be "MNRr" which stands for "Mask Nasty Repeat in read". This tag has an active masking function in MIRA and the fast all-against-all overlap searcher (SKIM) will then completely ignore the tagged subsequences of reads. There's one drawback though: the smaller the reads are that you try to assemble, the higher the probability that your reads will not span nasty repeats completely, therefore leading to a abortion of contig building at this site. Reads completely covered by the MNRr tag will therefore land in the debris file as no overlap will be found.

This option is extremely useful for assembly of larger projects (fungi-size) with a high percentage of repeats.

Note Although it is expected that bacteria will not really need this, leaving it turned on will probably not harm except in unusual cases like several copies of (pro-)phages integrated in a genome.

[nasty_repeat_ratio(nrr)=integer ≥ 2] Default is depending on the [--job=...] parameters. Normally it's high (around 100) for genome assemblies, but much lower (20 or less) for EST assemblies.

Sets the ratio from which on subsequences are considered nasty and hidden from the kmer statistics overlapper with a *MNRr* tag. E.g.: A value of 10 means: mask all k-mers of [-SK:kms] length which are occurring more than 10 times more often than the average of the whole project.

[nasty_repeat_coverage(nrc)=integer ≥ 0] Default is depending on the [--job=...] parameters: 0 for genome assemblies, 200 for EST assemblies.

Closely related to the [-KS:nrr] parameter (see above), but while the above works on ratios derived from a calculated average, this parameter allows to set an absolute value. Note that this parameter will take precedence over [-KS:nrr] if the calculated value of nrr is larger than the absolute value given here. A value of 0 de-activates this parameter.

[lossless_digital_normalisation(ldn)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent on --job type: yes for de-novo EST/RNAseq assembly, no otherwise.

Tells MIRA how on whether or not digitally normalising reads containing nasty repeats when [-KS:mnr] is active.

When set to *yes*, MIRA will apply a modified digital normalisation step to the reads, effectively decreasing the coverage of a given repetitive stretch down to a minimum needed to correctly represent one copy of the repeat. However, contrary to the published method, MIRA will keep enough reads of repetitive regions to also correctly reconstruct slightly different variants of the repeats present in the genome or EST / RNASeq data set, even if they differ in only a single base.

The tag used by MIRA to denote stretches which may have contributed to the digital normalisation will be "DGNr". Additionally, contigs which contain reads completely covered by a DGNr tag will get an additional "_dn" as part of their name to show that they contain read representatives for digital normalisation. E.g.: "contig_dn_c1".

This option is extremely useful for non-normalised EST / RNASeq projects, to get at least the sequence of overrepresented transcripts assembled even if the coverage values then cannot be interpreted as expression values anymore.

Note The lossless digital normalisation will be applied as soon as the kmer size of the active pass (see [-AS:kms]) reaches a size of at least 50 or, at the latest, in the second to last pass.

Note Once digital normalisation has been applied, the parameters [-KS:nrr] and [-KS:nrc] do not take effect anymore.



Warning The effect of lossless digital normalisation on genome data has not been studied sufficiently by me to approve it for genomes. Use with care in genome assemblies.

[repeatlevel_in_infofile(rliif)=integer; 0, 5-8] Default is 6. Sets the minimum level of the HAF tags from which on MIRA will report tentatively repetitive sequence in the *_info_readrepeats.lst file of the info directory.

A value of 0 means "switched off". The default value of 6 means all subsequences tagged with *HAF6*, *HAF7* and *MNRr* will be logged. If you, e.g., only wanted *MNRr* logged, you'd use 8 as parameter value.

See also [-KS:fenn:fxn:fer:fehr:mnr:nrr] to set the different levels for the *HAF* and *MNRr* tags.

[memory_to_use(mtu)=integer] Default is 75. Defines the memory MIRA can use to compute kmer statistics.

A value of >100 is interpreted as absolute value in megabyte. E.g., 16384 = 16384 megabyte = 16 gigabyte.

A value of 0 ≤ x ≤ 100 is interpreted as relative value of free memory at the time of computation. E.g.: for a value of 75% and 10 gigabyte of free memory, it will use 7.5 gigabyte.

Note The minimum amount of memory this algorithm will use is 512 Mib on 32 bit systems and 2 Gib on 64 bit systems.

9.3.4.12 Parameter group: -ALIGN (-AL)

The align options control the behaviour of the Smith-Waterman alignment routines. Only read pairs which are confirmed here may be included into contigs. Affects both the checking of possible alignments found by SKIM as well as the phase when reads are integrated into a contig.

Every option in this section can be set individually for every sequencing technology, giving a very fine grained control on how reads are aligned for each technology.

[bandwidth_in_percent(bip)=integer > 0 and ≤100] Default is dependent of the sequencing technology used. The banded Smith-Waterman alignment uses this percentage number to compute the bandwidth it has to use when computing the alignment matrix. E.g., expected overlap is 150 bases, bip=10 -> the banded SW will compute a band of 15 bases to each side of the expected alignment diagonal, thus allowing up to 15 unbalanced inserts / deletes in the alignment. INCREASING AND DECREASING THIS NUMBER: *increase*: will find more non-optimal alignments, but will also increase SW runtime between linear and \Circum2. *decrease*: the other way round, might miss a few bad alignments but gaining speed.

[bandwidth_min(bmin)=integer > 0] Default is dependent of the sequencing technology used. Minimum bandwidth in bases to each side.

[bandwidth_max(bmax)=integer > 0] Default is dependent of the sequencing technology used. Maximum bandwidth in bases to each side.

[min_overlap(mo)=integer > 0] Default is dependent of the sequencing technology used. Minimum number of overlapping bases needed in an alignment of two sequences to be accepted.

[min_score(ms)=integer > 0] Default is dependent of the sequencing technology used. Describes the minimum score of an overlap to be taken into account for assembly. MIRA uses a default scoring scheme for SW align: each match counts 1, a match with an N counts 0, each mismatch with a non-N base -1 and each gap -2. Take a bigger score to weed out a number of chance matches, a lower score to perhaps find the single (short) alignment that might join two contigs together (at the expense of computing time and memory).

[min_relative_score(mrs)=integer > 0 and ≤100] Default is dependent of the sequencing technology used. Describes the min % of matching between two reads to be considered for assembly. Increasing this number will save memory, but one might loose possible alignments. I propose a maximum of 80 here. Decreasing below 55% will make memory and time consumption probably explode.

Note: most of the time it makes sense to keep this parameter in sync with [-SK:pr].

[extra_gap_penalty(egp)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is dependent of the sequencing technology used. Defines whether or not to increase penalties applied to alignments containing long gaps. Setting this to 'yes' might help in projects with frequent repeats. On the other hand, it is definitively disturbing when assembling very long reads containing multiple long indels in the called base sequence ... although this should not happen in the first place and is a sure sign for problems lying ahead.

When in doubt, set it to yes for EST projects and de-novo genome assembly, set it to no for assembly of closely related strains (assembly against a backbone).

When set to no, it is recommended to have [-CO:amgb] and [-CO:amgbemc] both set to yes.

[egp_level(egpl)=comma separated list of integer ≥ 0 and ≤ 100] Default is dependent of the sequencing technology and job used. Has no effect if extra_gap_penalty is off.

...

[egp_level(megpp)=0 \leq integer \leq 100] Default is 100. Has no effect if extra_gap_penalty is off. Defines the maximum extra penalty in percent applied to 'long' gaps.

9.3.4.13 Parameter group: -FINALMAPPING (-FM)

The options control in this section take effect only in mapping mode and control the behaviour of the Smith-Waterman alignment routines only during the mapping stage of reads to the reference.

I.e., parameters from -AL take effect in this stage, but the -FM parameters do not take effect in the -AL stage. This allows for a more fine grained control.

Parameters in this section are currently more of a hack just for Solexa/Illumina data.

Note The former parameter [-AL:shme] has been removed, its functionality replaced and extended by parameters in this section.

[active(act)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Only takes effect when in mapping mode and activates a fine grained mapping mode.

The result is usually a much better mapping in areas with larger discrepancies between reference sequence and mapped data. Note that the final mapping stage takes longer if this mode is active, usually a factor of a bit above 2.

Switching off this functionality may be useful when mapping expression data where coverage values may be more important than the best possible alignment.

[(act)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Defines whether or not to map perfect matches.

Setting this option to no can come in handy when one wants to concentrate a mapping process only on differences between reference and mapped reads.

[maxtotalerrors(mte)=integer ≥ -1] When running in mapping mode, this defines the maximum number of mismatches and gaps a read may have compared to the reference to be allowed to map.

The default value of -1 lets MIRA choose this value automatically. It sets it to 15% of the average Illumina read lengths loaded.

[maxmismatches(mmm)=integer ≥ -1] Like [-FM:mte], but just for mismatches and except that a value of -1 means "don't care" instead of "don't compute automatically."

[maxgaps(mg)=integer ≥ -1] Like [-FM:mte], but just for gaps and except that a value of -1 means "don't care" instead of "don't compute automatically."

[clean_end_distance(ced)=integer ≥ 0] The default value of 0. When looking at overlaps between reads and reference, the ends of reads must have a mismatch/gap-free zone wrt to the reference of at least this length to be accepted.

Setting this parameter to a value > 0 has subtle implications for the results you will get in mapping when the data has larger insertions or deletions.

In short, the value of 0 will allow you to better manually reconstruct in **gap4** what happened at such a place, but the automatically generated FASTA exports of the mapping results will not show the exact gap length.

A value > 0 will give you better exports of the final results, but in **gap4** you will not be able to reconstruct what happened at an insertion or deletion site.

Note

Suggested use: take '4' for jobs in which you use MIRA as a better **pilon** replacement, i.e., you map Illumina data to a sequence to correct and polish the sequence. Take the default '0' for all other cases.

9.3.4.14 Parameter group: -CONTIG (-CO)

The contig options control the behaviour of the contig objects.

[name_prefix(np)=string] Default is <projectname>. Contigs will have this string prepended to their names. Normally, the [project=] line in the manifest will set this.

[reject_on_drop_in_relscore(rodirs)=integer ≥ 0 and ≤ 100] Default is dependent of the sequencing technology used.

When adding reads to a contig, reject the reads if the drop in the minimum relative score of the alignment of the current consensus and the new read is > the expected value calculated during the alignment phase. Lower values mean stricter checking.

This value is doubled should a read be entered that has an assembled template partner (a read pair) at the right distance in the current contig.

[cmin_relative_score(cmrs)=integer ≥ -1 and ≤ 100] Default is -1. Works similarly to [-AL:mrs], but during contig construction phase instead of read vs read alignment phase: describes the min % of matching between a read being added to a contig and the current contig consensus.

If value is set to -1, then the value of [-AL:mrs] is used.

Note: most of the time it makes sense to keep this parameter at -1. Else have it at approximately [-AL:mrs]-10 or switch it completely off via 0.

[mark_repeats(mr)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. One of the most important switches in MIRA: if set to yes, MIRA will try to resolve misassemblies due to repeats by identifying single base stretch differences and tag those critical bases as RMB (Repeat Marker Base, weak or strong). This switch is also needed when MIRA is run in EST mode to identify possible inter-, intra- and intra-and-interorganism SNPs.

[only_in_result(mroir)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. Only takes effect when [-CO:mr] (see above) is set to yes. If set to yes, MIRA will not use the repeat resolving algorithm during build time (and therefore will not be able to take advantage of this), but only before saving results to disk.

This switch is useful in some (rare) cases of mapping assembly.

[assume_snp_instead_repeat(asir)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. Only takes effect when [-CO:mr] (see above) is set to yes, effect is also dependent on the fact whether strain data (see - [-SB:lsd]) is present or not. Usually, MIRA will mark bases that differentiate between repeats when a conflict occurs between reads that belong to one strain. If the conflict occurs between reads belonging to different strains, they are marked as SNP. However, if this switch is set to yes, conflict within a strain are also marked as SNP.

This switch is mainly used in assemblies of ESTs, it should not be set for genomic assembly.

[min_reads_per_group(mrpg)=integer ≥ 2] Default is dependent of the sequencing technology used. Only takes effect when [-CO:mr] (see above) is set to yes. This defines the minimum number of reads in a group that are needed for the RMB (Repeat Marker Bases) or SNP detection routines to be triggered. A group is defined by the reads carrying the same nucleotide for a given position, i.e., an assembly with mrpg=2 will need at least two times two reads with the same nucleotide (having at least a quality as defined in [-CO:mgqrt]) to be recognised as repeat marker or a SNP. Setting this to a low number increases sensitivity, but might produce a few false positives, resulting in reads being thrown out of contigs because of falsely identified possible repeat markers (or wrongly recognised as SNP).

[min_neighbour_qual(mnq)=integer ≥ 10] Default is dependent of the sequencing technology used. Takes only effect when [-CO:mr] is set to yes. This defines the minimum quality of neighbouring bases that a base must have for being taken into consideration during the decision whether column base mismatches are relevant or not.

[min_groupqual_for_rmb_tagging(mgqrt)=integer ≥ 25] Default is dependent of the sequencing technology used. Takes only effect when [-CO:mr] is set to yes. This defines the minimum quality of a group of bases to be taken into account as potential repeat marker. The lower the number, the more sensitive you get, but lowering below 25 is not recommended as a lot of wrongly called bases can have a quality approaching this value and you'd end up with a lot of false positives. The higher the overall coverage of your project, the better, and the higher you can set this number. A value of 35 will probably remove most false positives, a value of 40 will probably never show false positives ... but will generate a sizable number of false negatives.

[min_coverage_percentage(mcp)=0 < integer ≤ 100] Default is currently 10. Used to reduce the number of IUPAC bases due to non-random PCR artefacts or sequencing errors in very high coverage areas (e.g. Illumina ≥ 80). Once the most probable base has been determined, [-CO:mcp] defines the minimum percentage (calculated from the most probable base) the coverage of alternative bases must have to be considered for consensus. E.g.: with mcp=10 and the most probable base having a coverage of 200x, other bases must have a coverage of 20x.

Drawback is that valid low frequency variants will not show up anymore as IUPAC in the FASTA.

[endread_mark_exclusion_area(emea)=integer ≥ 0] Default is dependent of the sequencing technology used. Takes only effect when [-CO:mr] is set to yes. Using the end of sequences of Sanger type shotgun sequencing is always a bit risky, as wrongly called bases tend to crowd there or some sequencing vector relics hang around. It is even more risky to use these stretches for detecting possible repeats, so one can define an exclusion area where the bases are not used when determining whether a mismatch is due to repeats or not.

[emea_set1_on_clipping_pec(emeas1clpec)=on/y[es]|t[rue], off/n[o]|f[alse]] Default is yes. When [-CL:pec] is set, the end-read exclusion area can be considerably reduced. Setting this parameter will automatically do this.

Note Although the parameter is named "set to 1", it may be that the exclusion area is actually a bit larger (2 to 4), depending on what users will report back as "best" option.

[also_mark_gap_bases(amgb)=on/y[es]|t[rue], off/n[o]|f[alse]] Default is dependent of the sequencing technology used. Determines whether columns containing gap bases (indels) are also tagged.

Note: it is strongly recommended to not set this to 'yes' for 454 type data.

[also_mark_gap_bases_even_multicolumn(amgbemc)=on/y[es]|t[rue], off/n[o]|f[alse]] Default is yes. Takes effect only when [-CO:amgb] is set to yes. Determines whether multiple columns containing gap bases (indels) are also tagged.

[also_mark_gap_bases_need_both_strands(amgbnbs)=on/y[es]|t[rue], off/n[o]|f[alse]] Default is yes. Takes effect only when [-CO:amgb] is set to yes. Determines whether both for tagging columns containing gap bases, both strands need to have a gap. Setting this to no is not recommended except when working in desperately low coverage situations.

[force_nonIUPACconsensus_perseqtype(fnic)=on/y[es]|t[rue], off/n[o]|f[alse]] Default is no for de-novo genome assemblies, yes for all others. If set to yes, MIRA will be forced to make a choice for a consensus base (A,C,G,T or gap) even in unclear cases where it would normally put a IUPAC base. All other things being equal (like quality of the possible consensus base and other things), MIRA will choose a base by either looking for a majority vote or, if that also is not clear, by preferring gaps over T over G over C over finally A.

MIRA makes a considerable effort to deduce the right base at each position of an assembly. Only when cases begin to be borderline it will use a IUPAC code to make you aware of potential problems. It is **suggested** to leave this option to no as IUPAC bases in the consensus are a sign that - if you need 100% reliability - you really should have a look at this particular place to resolve potential problems. You might want to set this parameter to yes in the following cases: 1) when your tools that use assembly result cannot handle IUPAC bases and you don't care about being absolutely perfect in your data (by looking over them manually). 2) when you assemble data without any quality values (which you should not do anyway), then this method will allow you to get a result without IUPAC bases that is "good enough" with respect to the fact that you did not have quality values.

[merge_short_reads(msr)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for all Solexas when in a mapping assembly, else it's no. Can only be used in mapping assemblies. If set to yes, MIRA will merge all perfectly mapping Solexa reads into longer reads (Coverage Equivalent Reads, CERs) while keeping quality and coverage information intact.

This feature hugely reduces the number of Solexa reads and makes assembly results with Solexa data small enough to be handled by current finishing programs (gap4, consed, others) on normal workstations.

[msr_keepcontigendsunmerged(msrme)=integer ≥ 0] Default is 0 for all Solexas when in a mapping assembly. Takes only effect in mapping assemblies if [-CO:msr=yes].

Defines how many "errors" (i.e. differences) a read may have to be merged into a coverage equivalent read. Useful only when one does not need SNP information from an assembly but wants to concentrate either on coverage data or on paired-end information at contig ends.



Warning This feature allows to merge non-perfect reads, which makes most SNP information simply disappear from the alignment. Use with care!

[msr_keepcontigendsunmerged(msrkceu)=-1, integer > 0] Default is -1 for all Solexas when in a mapping assembly. Takes only effect in mapping assemblies if [-CO:msr=yes] and for reads which have a paired-end / mate-pair partner actively used in the assembly.

If set to a value > 0, MIRA will not merge paired-end / mate-pair reads if they map within the given distance of a contig end of the original reference sequence (backbone). Instead of a fixed value, one can also use -1. MIRA will then automatically not merge reads if the distance from the contig end is within the maximum size of the template insert size of the sequencing library for that read (either given via [-GE:tismax] or via XML TRACEINFO for the given read).

This feature allows to use the data reduction from [-CO:msr] while enabling the result of such a mapping to be useful in subsequent scaffolding programs to order contigs.

9.3.4.15 Parameter group: -EDIT (-ED)

General options for controlling the integrated automatic editor. The editors generally make a good job cleaning up alignments from typical sequencing errors like (like base overcalls etc.). However, they may prove tricky in certain situations:

- in EST assemblies, they may edit rare transcripts toward almost identical, more abundant transcripts. Usage must be carefully weighed.
- the editors will not only change bases, but also sometimes delete or insert non-gap bases as needed to improve an alignment when facts (trace signals or other) show that this is what should have been the sequence. However, this can make post processing of assembly results pretty difficult with some formats like ACE, where the format itself contains no way to specify certain edits like deletion. There's nothing one can do about it and the only way to get around this problem is to use file formats with more complete specifications like CAF, MAF (and BAF once supported by MIRA).

The following edit parameters are supported:

[_mira_automatic_contig_editing(mace)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. When set to yes, MIRA will use built-in versions of own automatic contig editors (see parameters below) to improve alignments.

[edit_kmer_singlets(eks)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for all sequencing technologies, but only takes effect if [-ED:mace] is on (see above).

When set to yes, MIRA uses the alignment information of a complete contig at places with sequencing errors which lead to unique kmers and correct the error according to the alignment.

This is an extremely conservative yet very effective editing strategy and can therefore be kept always activated.

[edit_homopolymer_overcalls(ehpo)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes for 454 and Ion Torrent, but only takes effect if [-ED:mace] is on (see above).

When set to yes, MIRA use the alignment information of a complete contig at places with potential homopolymer sequencing errors and correct the error according to the alignment.

This editor should be switched on only for sequencing technologies with known homopolymer sequencing problems. That is: currently only 454 and Ion.

[edit_automatic_contig_editing(eace)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. When set to yes, MIRA will use built-in versions of the "EdIt" automatic contig editor (see parameters below) to correct sequencing errors in Sanger reads.

EdIt will try to resolve discrepancies in the contig by performing trace signal analysis and correct even hard to resolve errors.

Note The current development version has a memory leak in this editor, therefore the option cannot be turned on.

[strict_editing_mode(sem)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Only for Sanger data. If set to yes, the automatic editor will not take error hypotheses with a low probability into account, even if all the requirements to make an edit are fulfilled.

[confirmation_threshold(ct)=integer, 0 < x ≤ 100] Default is 50. Only for Sanger data. The higher this value, the more strict the automatic editor will apply its internal rule set. Going below 40 is not recommended.

9.3.4.16 Parameter group: -MISC (-MI)

Options which would not fit elsewhere.

[knowwhatido(ikwid)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. This switch tells MIRA that you know what you do in some situations and force it not to stop when it thinks something is really wrong, but simply continue.



Warning You generally should not to set this flag except in cases where MIRA stopped and the warning / error message told you to get around that very specific problem by setting this flag.

[large_contig_size(lcs)=integer < 0] Default is 500. This parameter has absolutely no influence whatsoever on the assembly process of MIRA. But is used in the reporting within the *_assembly_info.txt file after the assembly where MIRA reports statistics on *large* contigs and *all* contigs. [-MI:lcs] is the threshold value for dividing the contigs into these two categories.

[large_contig_size_for_stats(lcs4s)=integer < 0] Default is 5000 for [--job=genome] and 1000 for [--job=est].

This parameter is used for internal statistics calculations and has a subtle influence when being in a [--job=genome] assembly mode.

MIRA uses coverage information of an assembly project to find out about potentially repetitive areas in reads (and thus, a genome). To calculate statistics which are reflecting the approximate truth regarding the average coverage of a genome, the "large contig size for stats" value of [-MI:lcs4s] is used as a cutoff threshold: contigs smaller than this value do not contribute to the calculation of average coverage while contigs larger or equal to this value do.

This reflects two facts: on the one hand - especially with short read sequencing technologies and in projects without read pair libraries - contigs containing predominantly repetitive sequences are of a relatively small size. On the other hand, reads which could not be placed into contigs (maybe due to a sequencing technology dependent motif error) often enough form small contigs with extremely low coverage.

It should be clear that one does not want any of the above when calculating average coverage statistics and having this cutoff discards small contigs which tend to muddy the picture. If in doubt, don't touch this parameter.

9.3.4.17 Parameter group: -NAG_AND_WARN (-NW)

Parameters which let MIRA warn you about unusual things or potential problems. The flags in this parameter section come in three flavours: *stop*, *warn* and *no* which let MIRA either stop, give a warning or do nothing if a specific problem is detected.

[check_nfs(cnfs)=stop/warn/no] Default is stop. MIRA will check whether the tmp directory is running on a NFS mount.

**Warning**

You should never ever at all run MIRA on a NFS mounted directory ... or face the the fact that the assembly process may very well take 5 to 10 times longer (or more) than normal. You have been warned. The reason for the slowdown is the same as why one should never run a BLAST search on a big database being located on a NFS volume: access via network is terribly slow when compared to local disks, at least if you have not invested a lot of money into specialised solutions.

[check_duplicate_readnames(cdrn)=stop/warn/no] Default is stop. MIRA will check for duplicate read names after loading.

**Warning**

Duplicate read names usually hint to a serious problem with your input and should really, really be fixed. You can choose to ignore this error by switching off this flag, but this will almost certainly lead to problems with result files (ACE and CAF for sure, maybe also SAM) and probably to other unexpected effects.

[check_template_problems(ctp)=stop/warn/no] Default is stop. MIRA will check read template naming after loading.

**Warning**

Problems in read template naming point to problems with read names or to broken template information. You should try to find the cause of the problem instead of ignoring this error message.

[check_illuminajunkiness(cijie)=stop/warn/no] Default is stop.

MIRA will stop the assembly in EST/RNASeq assemblies with Illumina if either [-CL:kjd=no] or [-CL:kjck=no].

While in de-novo assemblies MIRA will not stop if these two parameters are off, it still is a good idea to keep them on. This warning parameter is there as a foolguard.

[check_proposedendclip(cpec)=stop/warn/no] Default is stop.

MIRA will stop the assembly in EST/RNASeq assemblies with Illumina if [-CL:pec=no].

[check_maxreadnamelength(cmrrl)=stop/warn/no] Default is stop. MIRA will check whether the length of the names of your reads surpass the given number of characters (see [-NW:mrnl]).

While MIRA and many other programs have no problem with long read names, some older programs have restrictions concerning the length of the read name. Example given: the pipeline CAF -> caf2gap -> gap2caf will stop working at the **gap2caf** stage if there are read names having > 40 characters where the names differ only at >40 characters.

This should be a warning only, but as a couple of people were bitten by this, the default behaviour of MIRA is to stop when it sees that potential problem. You might want to rename your reads to have ≤ 40 characters.

On the other hand, you also can ignore this potential problem and force MIRA to continue by using the parameter: [-NW:cmrrl=warn] or [-NW:cmrrl=no]

[maxreadnamelength(cmrrl)=integer ≥ 0] Default is 40. This defines the effective check length for [-NW:cmrrl].

[check_illuminajunkiness(cijie)=stop/warn/no] Default is stop. MIRA will check whether you switched off [-CL:kjd] or [-CL:kjck] when doing a de-novo assembly of Illumina data in EST/RNASeq mode.

[check_proposedendclip(cpec)=stop/warn/no] Default is stop. MIRA will check whether you switched off [-CL:cpec] when using any kind of reads from a 'real' sequencing technology (i.e., anything except TEXT).

[check_average_coverage(cac)=stop/warn/no] Default is stop. In genome de-novo assemblies, MIRA will perform checks early in the assembly process whether the average coverage to be expected exceeds a given value (see [-NW:acv]).

With today's sequencing technologies (especially Illumina, but also Ion Torrent and 454), many people simply take everything they get and throw it into an assembly. Which, in the case of Illumina and Ion, can mean they try to assemble their organism with a coverage of 100x, 200x and more (I've seen trials with more than 1000x).

This is not good. Not. At. All! For two reasons (well, three to be precise).

The first reason is that, usually, one does not sequence a single cell but a population of cells. If this population is not clonal (i.e., it contains subpopulations with genomic differences with each other), assemblers will be able to pick up these differences in the DNA once a certain sequence count is reached and they will try reconstruct a genome containing all clonal variations, treating these variations as potential repeats with slightly different sequences. Which, of course, will be wrong and I am pretty sure you do not want that.

The second and way more important reason is that none of the current sequencing technologies is completely error free. Even more problematic, they contain both random and non-random sequencing errors. Especially the latter can become a big hurdle if these non-random errors are so prevalent that they suddenly appear to be valid sequence to an assembler. This in turn leads to false repeat detection, hence possibly contig breaks or even wrong consensus sequence. You don't want that, do you?

The last reason is that overlap based assemblers (like MIRA is) need *exponentially* more time and memory when the coverage increases. So keeping the coverage comparatively low helps you there.

[average_coverage_value(acv)=integer ≥ 0] Default is 80 for de-novo assemblies, in mapping assemblies it is 120 for Ion Torrent and 160 for Illumina data (might change in future). This defines the effective coverage to check for in [-NW:cac].

[x174_contig_prefix(x174cp)=string] Default is WarnILMNPhiX174_. Setting this string to something else than an empty string will make MIRA check the consensus of every contig it builds against the PhiX174 phage. If more than 30% of the contig length matches PhiX174, the contig gets the prefix given in this option.

Illumina uses PhiX174 in sequencing as spike-in control, and some sequences make it to the final user data. MIRA has the ability to search and or filter against this, but when switched off, this parameter is the last possibility for users to check quickly whether these often unwanted sequences made it to their results.

9.3.4.18 Parameter group: -DIRECTORY (-DIR, -DI)

General options for controlling where to find or where to write data.

[tmp_redirected_to(trt)=<directoryname>] Default is an empty string. When set to a non-empty string, MIRA will create the MIRA-temporary directory at the given location instead of using the current working directory.

This option is particularly useful for systems which have solid state disks (SSDs) and some very fast disk subsystems which can be used for temporary files. Or in projects where the input and output files reside on a NFS mounted directory (current working dir), to put the tmp directory somewhere outside the NFS (see also: Things you should not do).

In both cases above, and for larger projects, MIRA then runs a lot faster.

Note Prior to MIRA 4.0rc2, users had to make sure themselves that the target directory did not already exist. MIRA now handles this automatically by creating directory names with a random substring attached.

9.3.4.19 Parameter group: -OUTPUT (-OUT)

Options for controlling which results to write to which type of files. Additionally, a few options allow output customisation of textual alignments (in text and HTML files).

There are 3 types of results: result, temporary results and extra temporary results. One probably needs only the results. Temporary and extra temporary results are written while building different stages of a contig and are given as convenience for trying to find out why MIRA set some RBMs or disassembled some contigs.

Output can be generated in these formats: CAF, Gap4 Directed Assembly, FASTA, ACE, TCS, WIG, HTML and simple text.

Naming conventions of the files follow the rules described in section **Input / Output**, subsection **Filenames**.

[savesimplesingletsinproject(sssip)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. Controls whether 'unimportant' singlets are written to the result files.

Note Note that a value larger 1 of the [-AS:mrpc] parameter will disable the function of this parameter.

[savetaggedsingletsinproject(stsip)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Controls whether singlets which have certain tags (see below) are written to the result files, even if [-OUT:sssip] (see above) is set.

If one of the (SRMr, CRMr, WRMr, SROr, SAOr, SIOr) tags appears in a singlet, MIRA will see that the singlets had been part of a larger alignment in earlier passes and even was part of a potentially 'important' decision. To give the possibility to human finishers to trace back the decision, these singlets can be written to result files.

Note Note that a value larger 1 of the [-AS:mrpc] parameter will disable the function of this parameter.

[remove_rollover_tmpr(rrt)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes. Removes log and temporary files once they should not be needed anymore during the assembly process.

[remove_tmp_directory(rtd)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no. Removes the complete tmp directory at the end of the assembly process. Some logs and temporary files contain useful information that you may want to analyse though, therefore the default of MIRA is not to delete it.

[output_result_caf(orc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes.

[output_result_maf(orm)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes.

[output_result_gap4da(org)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

Note If set to yes, MIRA will automatically switch back to no (and cannot be forced to 'yes') when 454 or Solexa reads are present in the project as this ensure that the file system does not get flooded with millions of files.

[output_result_fasta(orf)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes.

[output_result_ace(ora)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.



Warning ACE is the least suited file format for NGS data. Use it only when absolutely necessary.

[output_result_txt(ort)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[output_result_tcs(ors)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is yes.

[output_result_html(orb)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[output_tmpresult_caf(otc)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[output_tmpresult_maf(otm)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[output_tmpresult_gap4da(otg)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[output_tmpresult_fasta(otf)=on|y[es]|t[rue], off|n[o]|f[alse]] Default is no.

[**output_tmpresult_ace(ota)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_tmpresult_txt(ott)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_result_tcs(ots)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_tmpresult_html(oth)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_caf(oetc)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_gap4da(oetg)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_fasta(oetf)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_ace(oeta)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_txt(oett)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**output_exttmpresult_html(oeth)**=*on|y[es]|t[rue]*, *off|n[o]|f[alse]*] Default is no.

[**text_chars_per_line(tcpl)**=*integer > 0*] Default is 60. When producing an output in text format ([-OUT:ortlottloett]), this parameter defines how many bases each line of an alignment should contain.

[**html_chars_per_line(tcpl)**=*integer > 0*] Default is 60. When producing an output in HTML format, ([-OUT:orhlothloeth]), this parameter defines how many bases each line of an alignment should contain.

[**text_endgap_fillchar(tegfc)**=*<single character>*] Default is _ (a blank). When producing an output in text format ([-OUT:ortlottloett]), endgaps are filled up with this character.

[**html_endgap_fillchar(hegfc)**=*<single character>*] Default is _ (a blank). When producing an output in HTML format ([-OUT:orhlothloeth]), end-gaps are filled up with this character.

9.4 Resuming / restarting assemblies

It may happen that a MIRA run is interrupted - sometimes rather harshly - due to events more or less outside your control like, e.g., power failures, machine shutdowns for maintenance, missing disk space, run-time quotas etc. This may be less of a problem when assembling or mapping small data sets with run times between a couple of minutes up to a few hours, but becomes a nuisance for larger data sets like in small eukaryotes or RNASeq samples where the run time is measured in days.

If this happens in de-novo assemblies, MIRA has a *resume* functionality: at predefined points in the assembly process, MIRA writes out special files to disk which enables it to resume the assembly at the point where these files were written. Starting MIRA in resume mode is pretty easy: simply add the resume flag [-r] on a command line like this:

```
$ mira -r ...
```

where the ellipsis ("...") above stands for the rest of the command line you would have used to start a new assembly.

9.5 Input / Output

9.5.1 Directories

Since version 3.0.0, MIRA now puts all files and directories it generates into one sub-directory which is named *projectname_assembly*. This directory contains up to four sub-directories:

- *projectname_d_results*: this directory contains all the output files of the assembly in different formats.
- *projectname_d_info*: this directory contains information files of the final assembly. They provide statistics as well as, e.g., information (easily parsable by scripts) on which read is found in which contig etc.

- *projectname_d_tmp*: this directory contains tmp files and temporary assembly files. It can be safely removed after an assembly as there may be easily a few GB of data in there that are not normally not needed anymore.

In case of problems: please do not delete. I will get in touch with you for additional information that might possibly be present in the tmp directory.

- *projectname_d_chkpt*: this directory contains checkpoint files needed to resume assemblies that crashed or were stopped.

9.5.2 Filenames

9.5.2.1 Output

These result output files and sub-directories are placed in in the *projectname_results* directory after a run of MIRA.

projectname_out.<type> Assembled project written in type = (*maf / gap4da / caf / ace / fasta / html / tcs / wig / text*) format by MIRA, final result.

Type *gap4da* is a directory containing experiment files and a file of filenames (called 'fofn'), all other types are files. *gap4da*, *caf*, *ace* contain the complete assembly information suitable for import into different post-processing tools (*gap4*, *consed* and others). *html* and *text* contain visual representations of the assembly suited for viewing in browsers or as simple text file. *tcs* is a summary of a contig suited for "quick" analysis from command-line tools or even visual inspection. *wig* is a file containing coverage information (useful for mapping assemblies) which can be loaded and shown by different genome browsers (IGB, GMOD, USCS and probably many more).

fasta contains the contig consensus sequences (and *.fasta.qual* the consensus qualities). Please note that they come in two flavours: padded and unpadded. The padded versions may contain stars (*) denoting gap base positions where there was some minor evidence for additional bases, but not strong enough to be considered as a real base. Unpadded versions have these gaps removed. Padded versions have an additional postfix *.padded*, while unpadded versions *.unpadded*.

projectname_LargeContigs_out.<type> These files are only written when MIRA runs in *de-novo* mode. They usually contain a subset of contigs deemed 'large' from the whole project. More details are given in the chapter "working with results of MIRA."

9.5.2.2 Assembly statistics and information files

These information files are placed in in the *projectname_info* directory after a run of MIRA.

projectname_info_assembly.txt This file contains basic information about the assembly. MIRA will split the information in two parts: information about *large* contigs and information about all contigs.

For more information on how to interpret this file, please consult the chapter on "Results" of the MIRA documentation manual.

Note In contrast to other information files, this file always appears in the "info" directory, even when just intermediate results are reported.

projectname_info_contigreadlist.txt This file contains information which reads have been assembled into which contigs (or singlets).

projectname_info_contigstats.txt This file contains statistics about the contigs themselves, their length, average consensus quality, number of reads, maximum and average coverage, average read length, number of A, C, G, T, N, X and gaps in consensus.



Warning For contigs containing digitally normalised reads, the coverage numbers may sometimes seem strange. E.g.: a contig may contain only one read, but have an average coverage of 3. This means that the read was a representative for 3 reads. The coverage numbers are computed as if all 3 reads had been assembled instead of the representative. In EST/RNaseq projects, these numbers thus represent the (more or less) true expression coverage.

projectname_info_consensustaglist.txt This file contains information about the tags (and their position) that are present in the consensus of a contig.

projectname_info_largecontigs.txt For de-novo assemblies, this file contains the name of the contigs which pass the (adaptable) 'large contig' criterion.

projectname_info_readrepeats.lst Tab delimited file with three columns: read name, repeat level tag, sequence.

This file permits a quick analysis of the repetitiveness of different parts of reads in a project. See [-SK:rliif] to control from which repetitive level on subsequences of reads are written to this file,

Note Reads can have more than one entry in this file. E.g., with standard settings (-SK:rliif=6) if the start of a read is covered by MNRr, followed by a HAF3 region and finally the read ends with HAF6, then there will be two lines in the file: one for the subsequence covered by MNRr, one for HAF6.

projectname_info_readstooshort A list containing the names of those reads that have been sorted out of the assembly before any processing started only due to the fact that they were too short.

projectname_info_readtaglist.txt This file contains information about the tags and their position that are present in each read. The read positions are given relative to the forward direction of the sequence (i.e. as it was entered into the the assembly).

projectname_info_WARNINGS_*.txt These files collect warning messages MIRA dumped out throughout the assembly process. These warnings cover a wide area of things monitored by MIRA and can - together with the output written to STDOUT - give an insight as to why an assembly does not behave as expected. There are three warning files representing different levels of criticality: *critical*, *medium* and *minor*. These files may be empty, meaning that no warning of the corresponding level was printed. It is strongly suggested to have a look at least at critical warnings during and after an assembly run.

Note These files are quite new to MIRA and not all warning messages appear there yet. This will come over time.

projectname_error_reads_invalid A list of sequences that have been found to be invalid due to various reasons (given in the output of the assembler).

9.5.3 File formats

MIRA can write almost all of the following formats and can read most of them.

ACE This old assembly file format used mainly by phrap and consed. Support for .ace output is currently only in test status in MIRA as documentation on that format is ... sparse and I currently don't have access to consed to verify my assumptions. Using consed, you will need to load projects with -nophd to view them. Tags /in reads and consensus) are fully supported. The only hitch: consed has a bug which prevents it to read consensus tags which are located throughout the whole file (as MIRA writes per default). The solution to that is easy: filter the CAF file through the fixACE4consed.tcl script which is provided in older MIRA distributions (V4.9.5 and before), then all should be well.

If you don't have consed, you might want to try clview (<http://www.tigr.org/tdb/tgi/software/>) from TIGR to look at .ace files.

BAM The binary cousin of the SAM format. MIRA neither reads nor writes BAM, but BAMs can be created out of SAMs (which can be created via **miraconvert**).

CAF Common Assembly Format (CAF) developed by the Sanger Centre. <http://www.sanger.ac.uk/resources/software/caf.html> provides a description of the format and some software documentation as well as the source for compiling caf2gap and gap2caf (thanks to Rob Davies for this).

EXP Standard experiment files used in genome sequencing. Correct EXP files are expected. Especially the ID record (containing the id of the reading) and the LN record (containing the name of the corresponding trace file) should be correctly set. See <http://www.sourceforge.net/projects/staden/> for links to online format description.

FASTA A simple format for sequence data, see <http://www.ncbi.nlm.nih.gov/BLAST/fasta.html>. An often used extension of that format is used to also store quality values in a similar fashion, these files have a .fasta.qual ending.

MIRA writes two kinds of FASTA files for results: *padded* and *unpadded*. The difference is that the padded version still contains the gap (pad) character (an asterisk) at positions in the consensus where some of the reads apparently had some more bases than others but where the consensus routines decided that to treat them as artifacts. The *unpadded* version has the gaps removed.

GBF, **GBK** GenBank file format as used at the NCBI to describe sequences. MIRA is able to read and write this format (but only for viruses or bacteria) for using sequences as backbones in an assembly. Features of the GenBank format are also transferred automatically to Staden compatible tags.

If possible, use GFF3 instead (see below).

GFF3 General feature format used to describe sequences and features on these sequences. MIRA is able to read and write this format.

HTML Hypertext Markup Language. Projects written in HTML format can be viewed directly with any table capable browser. Display is even better if the browser knows style sheets (CSS).

MAF MIRA Assembly Format (MAF). A faster and more compact form than EXP, CAF or ACE. See documentation in separate file.

PHD This file type originates from the phred base caller and contains basically -- along with some other status information -- the base sequence, the base quality values and the peak indices, but not the sequence traces itself.

SAM The Sequence Alignment/Map Format. MIRA does not write SAM directly, but **miracore** can be used for converting a MAF (or CAF) file to SAM.

MIRA cannot read SAM though.

SCF The Staden trace file format that has established itself as compact standard replacement for the much bigger ABI files. See <http://www.sourceforge.net/projects/staden/> for links to online format description.

The SCF files should be V2-8bit, V2-16bit, V3-8bit or V3-16bit and can be packed with compress or gzip.

traceinfo.XML XML based file with information relating to traces. Used at the NCBI and ENSEMBL trace archive to store additional information (like clippings, insert sizes etc.) for projects. See further down for a description of the fields used and <http://www.ncbi.nlm.nih.gov/Traces/trace.cgi?cmd=show&f=rfc&m=main&s=rfc> for a full description of all fields.

TCS Transpose Contig Summary. A text file as written by MIRA which gives a summary of a contig in tabular fashion, one line per base. Nicely suited for "quick" analysis from command line tools, scripts, or even visual inspection in file viewers or spreadsheet programs.

In the current file version (TCS 1.0), each column is separated by at least one space from the next. Vertical bars are inserted as visual delimiter to help inspection by eye. The following columns are written into the file:

1. contig name (width 20)
 2. padded position in contigs (width 3)
 3. unpadded position in contigs (width 3)
 4. separator (a vertical bar)
 5. called consensus base
 6. quality of called consensus base (0-100), but MIRA itself caps at 90.
 7. separator (a vertical bar)
 8. total coverage in number of reads. This number can be higher than the sum of the next five columns if Ns or IUPAC bases are present in the sequence of reads.
-

9. coverage of reads having an "A"
10. coverage of reads having an "C"
11. coverage of reads having an "G"
12. coverage of reads having an "T"
13. coverage of reads having an "*" (a gap)
14. separator (a vertical bar)
15. quality of "A" or "--" if none
16. quality of "C" or "--" if none
17. quality of "G" or "--" if none
18. quality of "T" or "--" if none
19. quality of "*" (gap) or "--" if none
20. separator (a vertical bar)
21. Status. This field sums up the evaluation of MIRA whether you should have a look at this base or not. The content can be one of the following:
 - everything OK: a colon (:)
 - unclear base calling (IUPAC base): a "!M"
 - potentially problematic base calling involving a gap or low quality: a "!m"
 - consensus tag(s) of MIRA that hint to problems: a "!\$". Currently, the following tags will lead to this marker: SRMc, WRMc, DGPc, UNSc, IUPc.
22. list of a consensus tags at that position, tags are delimited by a space. E.g.: "DGPc H454"

9.5.4 STDOUT/STDERR

The actual stage of the assembly is written to `STDOUT`, giving status messages on what MIRA is actually doing. Dumping to `STDERR` is almost not used anymore by MIRA, remnants will disappear over time.

Some debugging information might also be written to `STDOUT` if MIRA generates error messages.

On errors, MIRA will dump these also to `STDOUT`. Basically, three error classes exist:

1. **WARNING:** Messages in this error class do not stop the assembly but are meant as an information to the user. In some rare cases these errors are due to (an always possible) error in the I/O routines of MIRA, but nowadays they are mostly due to unexpected (read: wrong) input data and can be traced back to errors in the preprocessing stages. If these errors arise, you definitely **DO** want to check how and why these errors came into those files in the first place.
Frequent cause for warnings include missing SCF files, SCF files containing known quirks, EXP files containing known quirks etc.
2. **FATAL:** Messages in this error class actually stop the assembly. These are mostly due to missing files that MIRA needs or to very garbled (wrong) input data.
Frequent causes include naming an experiment file in the 'file of filenames' that could not be found on the disk, same experiment file twice in the project, suspected errors in the EXP files, etc.
3. **INTERNAL:** These are true programming errors that were caught by internal checks. Should this happen, please mail the output of `STDOUT` and `STDERR` to the author.

9.5.5 SSAHA2 / SMALT ancillary data

The **ssaha2** or **smalt** programs - both from the Sanger Centre - can be used to detect possible vector sequence stretches in the input data for the assembly. MIRA can load the result files of a **ssaha2** or **smalt** run and interpret the results to tag the possible vector sequences at the ends of reads.

Note that this also uses the parameters `[-CL:msvsgs:msvsmfg:msvsmeg]` (see below).

ssaha2 must be called like this "ssaha2 <ssaha2options> vector.fasta sequences.fasta" to generate an output that can be parsed by MIRA. In the above example, replace `vector.fasta` by the name of the file with your vector sequences and `sequences.fasta` by the name of the file containing your sequencing data.

smalt must be called like this: "smalt map -f ssaha <ssaha2options> hash_index sequences.fasta"

This makes you basically independent from any other commercial or license-requiring vector screening software. For Sanger reads, a combination of **lucy** and **ssaha2** or **smalt** together with this parameter should do the trick. For reads coming from 454 pyro-sequencing, **ssaha2** or **smalt** and this parameter will also work very well. See the usage manual for a walkthrough example on how to use SSAHA2 / SMALT screening data.

Note The output format of SSAHA2 must be the native output format (`-output ssaha2`). For SMALT, the output option `-f ssaha` must be used. Other formats cannot be parsed by MIRA.

Note I currently use the following SSAHA2 options: `-kmer 8 -skip 1 -seeds 1 -score 12 -cmatch 9 -ckmer 6`

Note Anyone contributing SMALT parameters?

Note The sequence vector clippings generated from SSAHA2 / SMALT data do not replace sequence vector clippings loaded via the EXP, CAF or XML files, they rather extend them.

9.5.6 XML TRACEINFO ancillary data

MIRA extracts the following data from the TRACEINFO files:

- `trace_name` (required)
 - `trace_file` (recommended)
 - `trace_type_code` (recommended)
 - `trace_end` (recommended)
 - `clip_quality_left` (recommended)
 - `clip_quality_right` (recommended)
 - `clip_vector_left` (recommended)
 - `clip_vector_right` (recommended)
 - `strain` (recommended)
 - `template_id` (recommended for paired end)
 - `insert_size` (recommended for paired end)
 - `insert_stdev` (recommended for paired end)
 - `machine_type` (optional)
 - `program_id` (optional)
-

Other data types are also read, but the info is not used.

Here's the example for a TRACEINFO file with ancillary info:

```
<?xml version="1.0"?>
<trace_volume>
<trace>
  <trace_name>GCJAA15TF</trace_name>
  <program_id>PHRED (0.990722.G) AND TTUNER (1.1)</program_id>
  <template_id>GCJAA15</template_id>
  <trace_direction>FORWARD</trace_direction>
  <trace_end>F</trace_end>
  <clip_quality_left>3</clip_quality_left>
  <clip_quality_right>622</clip_quality_right>
  <clip_vector_left>1</clip_vector_left>
  <clip_vector_right>944</clip_vector_right>
  <insert_stdev>600</insert_stdev>
  <insert_size>2000</insert_size>
</trace>
<trace>
  ...
</trace>
...
</trace_volume>
```

See <http://www.ncbi.nlm.nih.gov/Traces/trace.cgi?cmd=show&f=rfc&m=main&s=rfc> for a full description of all fields and more info on the TRACEINFO XML format.

9.5.7 Contig naming

MIRA names contigs the following way: `<projectname>_<contigtype><number>`. While `<projectname>` is dictated by the `[--project=]` parameter and `<number>` should be clear, the `<contigtype>` might need additional explaining. There are currently three contig types existing:

1. `_c`: these are "normal" contigs
2. `_rep_c`: only for genome assembly mode. These are contigs containing only repetitive areas. These contigs had `_lrc` as type in previous version of MIRA, this was changed to the `_rep_c` to make things clearer.
3. `_s`: these are singlet-contigs. Technically: "contigs" with a single read.
4. `_dn`: these is an additional contig type which can occur when MIRA ran a digital normalisation step during the assembly. Contigs which contain reads completely covered by a DGNr tag will get an additional `"_dn"` as part of their name to show that they contain read representatives for digital normalisation. E.g.: "contig_dn_c1".

Reads covered only partly by the DGNr tag do not trigger the `_dn` naming.

Important side note Due to the digital normalisation step, the coverage numbers in the info file regarding contig statistics will not represent the number of reads in the contig, but they will show an approximation of the true coverage or expression value as if there had not been a digital normalisation step performed. The approximation may be around 10 to 20% below the true value.

Basically, for genome assemblies MIRA starts to build contigs in areas which seem "rock solid", i.e., not a repetitive region (main decision point) and nice coverage of good reads. Contigs which started like this get a `_c` name. If during the assembly MIRA reaches a point where it cannot start building a contig in a non-repetitive region, it will name the contig `_rep_c` instead of `_c`. This is why "`_rep_c`" contigs occur late in a genome assembly.

Note MIRA has a different understanding of "rock solid" when in EST/RNASEq assembly: here, MIRA will try to reconstruct a full length gene sequence, starting with the most abundant genes.

Note Depending on the settings of [-AS:mrpc], your project may or may not contain `_s` singlet-contigs. Also note that reads landing in the debris file will not get assigned to singlet-contigs and hence not get `_s` names.

9.5.8 Recovering strain specific consensus as FASTA

In case you used strain information in an assembly, you can recover the consensus for just any given strain by using **miraconvert** and convert from a full assembly format (e.g. MAF or CAF) which also carries strain information to FASTA. MIRA will automatically detect the strain information and create one FASTA file per strain encountered.

It will also create a blend of all strains encountered and conveniently add "AllStrains" to the name of these files. Note that this blend may or may not be something you need, but in some cases I found it to be useful.

9.6 Tags used in the assembly by MIRA and EdIt

MIRA uses and sets a couple of tags during the assembly process. That is, if information is known before the assembly, it can be stored in tags (in the EXP and CAF formats) and will be used in the assembly.

9.6.1 Tags read (and used)

This section lists "foreign" tags, i.e., tags that whose definition was made by other software packages than MIRA.

- **ALUS, REPT:** Sequence stretches tagged as ALUS (ALU Sequence) or REPT (general repetitive sequence) will be handled with extreme care during the assembly process. The allowed error rate after automatic contig editing within these stretches is normally far below the general allowed error rate, leading to much higher stringency during the assembly process and subsequently to a better repeat resolving in many cases.
- **Fpas:** GenBank feature for a poly-A sequence. Used in EST, cDNA or transcript assembly. Either read in the input files or set when using [-CL:cpat]. This allows to keep the poly-A sequence in the reads during assembly without them interfering as massive repeats or as mismatches.
- **FCDS, Fgen:** GenBank features as described in GBF/GBK files or set in the Staden package are used to make some SNP impact analysis on genes.
- **other.** All other tags in reads will be read and passed through the assembly without being changed and they currently do not influence the assembly process.

9.6.2 Tags set (and used)

This section lists tags which MIRA sets (and reads of course), but that other software packages might not know about.

- **UNSR, UNSc:** **UN**Sure in **R**ead respectively **C**ontig. These tags denote positions in an assembly with conflicts that could not be resolved automatically by MIRA. These positions should be looked at during the finishing process.

For assemblies using good sequences and enough coverage, something 0.01% of the consensus positions have such a tag. (e.g. ~300 UNSc tags for a genome of 3 megabases).

- **SRMr, WRMc: Strong Repeat Marker and Weak Repeat Marker.** These tags are set in two flavours: as **SRMr** and **WRMr** when set in reads, and as **SRMc** and **WRMc** when set in the consensus. These tags are used on an individual per base basis for each read. They denote bases that have been identified as crucial for resolving repeats, often denoting a single SNP within several hundreds or thousands of bases. While a SRM is quite certain, the WRM really is either weak (there wasn't enough comforting information in the vicinity to be really sure) or involves gap columns (which is always a bit tricky).

MIRA will automatically set these tags when it encounters repeats and will tag exactly those bases that can be used to discern the differences.

Seeing such a tag in the consensus means that MIRA was not able to finish the disentanglement of that special repeat stretch or that it found a new one in one of the last passes without having the opportunity to resolve the problem.

- **DGPc: Dubious Gap Position in Consensus.** Set whenever the gap to base ratio in a column of 454 reads is between 40% and 60%.
- **SAO, SRO, SIO: SNP intra Organism, SNP R Organism, SNP Intra and inter Organism.** As for SRM and WRM, these tags have a **r** appended when set in reads and a **c** appended when set in the consensus. These tags denote SNP positions.

MIRA will automatically set these tags when it encounters SNPs and will tag exactly those bases that can be used to discern the differences. They denote SNPs as they occur within an organism (SAO), between two or more organisms (SRO) or within and between organisms (SIO).

Seeing such a tag in the consensus means that MIRA set this as a valid SNP in the assembly pass. Seeing such tags only in reads (but not in the consensus) shows that in a previous pass, MIRA thought these bases to be SNPs but that in later passes, this SNP does not appear anymore (perhaps due to resolved misassemblies).

- **STMS: (only hybrid assemblies).** The **Sequencing Type Mismatch Solved** is tagged to positions in the assembly where the consensus of different sequencing technologies (Sanger, 454, Ion Torrent, Solexa, PacBio, SOLiD) reads differ, but MIRA thinks it found out the correct solution. Often this is due to low coverage of one of the types and an additional base calling error.

Sometimes this depicts real differences where possible explanation might include: slightly different bugs were sequenced or a mutation occurred during library preparation.

- **STMU: (only hybrid assemblies).** The **Sequencing Type Mismatch Unresolved** is tagged to positions in the assembly where the consensus of different sequencing technologies (Sanger, 454, Ion Torrent, Solexa, SOLiD) reads differ, but MIRA could not find a good resolution. Often this is due to low coverage of one of the types and an additional base calling error.

Sometimes this depicts real differences where possible explanation might include: slightly different bugs were sequenced or a mutation occurred during library preparation.

- **MCVc: The Missing Co{V}erage in Consensus.** Set in assemblies with more than one strain. If a strain has no coverage at a certain position, the consensus gets tagged with this tag (and the name of the strain which misses this position is put in the comment). Additionally, the sequence in the result files for this strain will have an @ character.

- **MNRr: (only with [-KS:mnr] active).** The **Masked Nasty Repeat** tags are set over those parts of a read that have been detected as being many more times present than the average sub-sequence. MIRA will hide these parts during the initial all-against-all overlap finding routine (SKIM3) but will otherwise happily use these sequences for consensus generation during contig building.

- **FpAS:** See "Tags read (and used)" above.

- **ED_C, ED_I, ED_D:** EDit Change, EDit Insertion, EDit Deletion. These tags are set by the integrated automatic editor EdIt and show which edit actions have been performed.

- **HAF2, HAF3, HAF4, HAF5, HAF6, HAF7.** These are **HAsh Frequency** tags which show the status of read parts in comparison to the whole project. Only set if [-AS:ard] is active (default for genome assemblies).

More info on how to use the information conveyed by HAF tags in the section dealing with repeats and HAF tags in finishing programs further down in this manual.

HAF2 coverage below average (standard setting at < 0.5 times average)

HAF3 coverage is at average (standard setting at ≥ 0.5 times average and ≤ 1.5 times average)

HAF4 coverage above average (standard setting at > 1.5 times average and < 2 times average)

HAF5 probably repeat (standard setting at ≥ 2 times average and < 5 times average)

HAF6 'heavy' repeat (standard setting at > 8 times average)

HAF7 'crazy' repeat (standard setting at > 20 times average)

9.7 Where reads end up: contigs, singlets, debris

At the start, things are simple: a read either aligns with other reads or it does not. Reads which align with other reads form contigs, and these MIRA will save in the results with a contig name of `_c`.

However, not all reads can be placed in an assembly. This can have several reasons and these reads may end up at two different places in the result files: either in the *debris* file, then just as a name entry, or as singlet (a "contig" with just one read) in the regular results.

1. reads are too short and get filtered out (before or after the MIRA clipping stages). These invariably land in the debris file.
2. reads are real singlets: they contain genuine sequence but have no overlap with any other read. These get either caught by the `[-CL:pec]` clipping filter or during the SKIM phase
3. reads contain mostly or completely junk.
4. reads contain chimeric sequence (therefore: they're also junk)

MIRA filters out these reads in different stages: before and after read clipping, during the SKIM stage, during the Smith-Waterman overlap checking stage or during contig building. The exact place where these single reads land is dependent on why they do not align with other reads. Reads landing in the debris file will have the reason and stage attached to the decision.

9.8 Detection of bases distinguishing non-perfect repeats and SNP discovery

MIRA is able to find and tag SNPs in any kind of data -- be it genomic or EST -- in both de-novo and mapping assemblies ... provided it knows which read in an assembly is coming from which strain, cell line or organism.

The SNP detection routines are based on the same routines as the routines for detecting non-perfect repeats. In fact, MIRA can even distinguish between bases marking a misassembled repeat from bases marking a SNP within the same project.

All you need to do to enable this feature is to set `[-CO:mr=yes]` (which is standard in all `--job=...` incantations of **mira** and in some steps of **miraSearchESTSNPs**). Furthermore, you will need to provide *strain information*, either in the manifest file or in ancillary NCBI TRACEINFO XML files.

The effect of using strain names attached to reads can be described briefly like this. Assume that you have 6 reads (called R1 to R6), three of them having an A at a given position, the other three a C.

```
R1    .....A.....
R2    .....A.....
R3    .....A.....
R4    .....C.....
R5    .....C.....
R6    .....C.....
```

Note This example is just that: an example. It uses just 6 reads, with two times three reads as read groups for demonstration purposes and without looking at qualities. For MIRA to recognise SNPs, a few things must come together (e.g. for many sequencing technologies it wants forward and backward reads when in de-novo assembly) and a couple of parameters can be set to adjust the sensitivity. Read more about the parameters: `[-CO:mrpg:mnq:mgqrt:emea:amgb:amgbemc:amgbnbs]`

Now, assume you did not give any strain information. MIRA will most probably recognise a problem and, having no strain information, assume it made an error by assembling two different repeats of the same organism. It will tag the bases in the reads with repeat marker tags (SRMr) and the base in the consensus with a SROc tag (to point at an unresolved problem). In a subsequent pass, MIRA will then not assemble these six reads together again, but create two contigs like this:

```
Contig1:
R1  .....A.....
R2  .....A.....
R3  .....A.....

Contig2:
R4  .....C.....
R5  .....C.....
R6  .....C.....
```

The bases in the repeats will keep their SROr tags, but the consensus base of each contig will not get SROc as there is no conflict anymore.

Now, assume you gave reads R1, R2 and R3 the strain information "human", and read R4, R5 and R6 "chimpanzee". MIRA will then create this:

```
R1 (hum)  .....A.....
R2 (hum)  .....A.....
R3 (hum)  .....A.....
R4 (chi)  .....C.....
R5 (chi)  .....C.....
R6 (chi)  .....C.....
```

Instead of creating two contigs, it will create again one contig ... but it will tag the bases in the reads with a SROr tag and the position in the contig with a SROc tag. The SRO tags (SNP inte**R** Organisms) tell you: there's a SNP between those two (or multiple) strains/organisms/whatever.

Changing the above example a little, assume you have this assembly early on during the MIRA process:

```
R1 (hum)  .....A.....
R2 (hum)  .....A.....
R3 (hum)  .....A.....
R4 (chi)  .....A.....
R5 (chi)  .....A.....
R6 (chi)  .....A.....
R7 (chi)  .....C.....
R8 (chi)  .....C.....
R9 (chi)  .....C.....
```

Because "chimp" has a SNP within itself (A versus C) and there's a SNP between "human" and "chimp" (also A versus C), MIRA will see a problem and set a tag, this time a SIOr tag: SNP Intra- and inter **O**rganism.

MIRA does not like conflicts occurring within an organism and will try to resolve these cleanly. After setting the SIOr tags, MIRA will re-assemble in subsequent passes this:

```
Contig1:
R1 (hum)  .....A.....
R2 (hum)  .....A.....
R3 (hum)  .....A.....
R4 (chi)  .....A.....
R5 (chi)  .....A.....
R6 (chi)  .....A.....

Contig2:
R7 (chi)  .....C.....
R8 (chi)  .....C.....
R9 (chi)  .....C.....
```

The reads in Contig1 (hum+chi) and Contig2 (chi) will keep their SIOr tags, the consensus will have no SIOc tag as the "problem" was resolved.

When presented to conflicting information regarding SNPs and possible repeat markers or SNPs within an organism, MIRA will always first try to resolve the repeats marker. Assume the following situation:

```
R1 (hum)  .....A...T.....
R2 (hum)  .....A...G.....
R3 (hum)  .....A...T.....
R4 (chi)  .....C...G.....
R5 (chi)  .....C...T.....
R6 (chi)  .....C...G.....
```

While the first discrepancy column can be "explained away" by a SNP between organisms (it will get a SROr/SROc tag), the second column cannot and will get a SIOr/SIOc tag. After that, MIRA opts to get the SIO conflict resolved:

```
Contig1:
R1 (hum)  .....A...T.....
R3 (hum)  .....A...T.....
R5 (chi)  .....C...T.....

Contig2:
R2 (hum)  .....A...G.....
R4 (chi)  .....C...G.....
R6 (chi)  .....C...G.....
```

9.9 Data reduction: subsampling vs. lossless digital normalisation

Some data sets have way too much data. Sometimes it is simply more than needed like, e.g., performing a de-novo genome assembly with reads enough for 300x coverage is like taking a sledgehammer for cracking a nut. Sometimes it is even more than is good for an assembly (see also: motif dependent sequencing errors).

MIRA being an overlap-based assembler, reducing a data set helps to keep time and memory requirements low. There are basically two ways to perform this: reduction by subsampling and reduction by digital normalisation. Both methods have their pros and cons and can be used effectively in different scenarios.

- *Subsampling* is a process to create a smaller, hopefully representative set from a larger data set.

In sequencing, various ways exist to perform subsampling. As sequencing data sets from current sequencing technologies can be seen as essentially randomised when coming fresh from the machine, the selection step can be as easy as selecting the first n reads. When the input data set is not random (e.g. in SAM/BAM files with mapped data), one must resort to random selection of reads.

Subsampling must be done by the user prior to assembly with MIRA.

On the upside, subsampling preserves the exact copy number structure of the input data set: a repeat with n copies in a genome will always be represented by reads forming n copies of the repeat in the reduced data set. Furthermore, subsampling is comparatively insensitive to motif dependent sequencing errors. On the downside, subsampling will more probably lose rare events of the data set (e.g., rare SNPs of a cell population or rare transcripts in EST/RNASeq). Also, in EST/RNASeq projects, subsampling will not be able to reduce extraordinary coverage events to a level which make the assembly not painfully slow. Examples for the later being rRNA genes or highly expressed house-keeping genes where today's Illumina data sets sometimes contains enough data to reach coverage numbers $\geq 100,000x$ or even a million x .

Subsampling should therefore be used for single genome de-novo assemblies; or for EST/RNASeq assemblies which need reliable coverage numbers for transcript expression data but where at least all rDNA has been filtered out prior to assembly.

- *Digital normalisation* is a process to perform a reduction of sequencing data redundancy. It was made known to a wider audience by the paper "*A Reference-Free Algorithm for Computational Normalization of Shotgun Sequencing Data*" by Brown et al. (see <http://arxiv.org/abs/1203.4802>).

The normalisation process works by progressively going through the sequencing data and selecting reads which bring new, previously unseen information to the assembly and discarding those which describe nothing new. For single genome assemblies, this has the effect that repeats with n copies in the genome are afterwards present often with just enough reads to reconstruct only a single copy of the repeat. In EST/RNASeq assemblies, this leads to reconstructed transcripts having all the more or less same coverage.

The normalisation process as described in the paper allows for a certain lossiness during the data reduction as it was developed to cope with billions of reads. E.g., it will often lose borders in genome reorganisation events or SNP information from ploidies, from closely related genes copies or from closely related species.

MIRA implements a variant of the algorithm: the *lossless digital normalisation*. Here, normalised data has copy numbers reduced like in the original algorithm, but all variants (SNPs, borders of reorganisation events etc.) present in the original data set are retained in the reduced data set. Furthermore, the normalisation is parameterised to take place only for excessively repetitive parts of a data set which would lead to overly increased run-time and memory consumption. This gives the assembler the opportunity to correctly evaluate and work with repeats which do not occur "too often" in a data set while still being able to reconstruct at least one copy of the really nasty repeats.

Digital normalisation should not be done prior to an assembly with MIRA, rather the MIRA parameter to perform a digital normalisation on the complete data set should be used.

The lossless digital normalisation of MIRA should be used for EST/RNASeq assemblies containing highly repetitive data. Metagenome assemblies may also profit from this feature.

Note

MIRA keeps track of the approximate coverage represented by the reads chosen in the digital normalisation process. That is, MIRA is able to give approximate coverage numbers as if digital normalisation had never happened. The approximation may be around 10 to 20% below the true value. Contigs affected by this coverage approximation are denoted with an additional "_dn" in their name.

Due to the digital normalisation step, the coverage numbers in the info file regarding contig statistics will not represent the number of reads in the contig, but they will show an approximation of the true coverage or expression value as if there had not been a digital normalisation step performed.

9.10 Caveats

9.10.1 Using data not from sequencing instruments: artificial / synthetic reads

The default parameters for MIRA assemblies work best when given real sequencing data and they even expect the data to behave like real sequencing data. But some assembly strategies work in multiple rounds, using so called "artificial" or "synthetic" reads in later rounds, i.e., data which was not generated through sequencing machines but might be something like the consensus of previous assemblies.

If one doesn't take utter care to make these artificial reads at least behave a little bit like real sequencing data, a number of quality insurance algorithms of MIRA might spot that they "look funny" and trim back these artificial reads ... sometimes even removing them completely.

Summary tips for creating artificial reads for MIRA assemblies

The following should lead to the least amount of surprises for most assembly use cases when calling MIRA only with the most basic switches `--project=...` `--job=...`

1. **Length:** between 50 and 20000 bp
 2. **Quality values:** give your artificial reads quality values. Using 30 as quality value for your bases should be OK for most applications.
 3. **Orientation:** for every read you create, create a read with the same data (bases and quality values) in reverse complement direction.
-

The following list gives all the gory details on how synthetic reads should look like or which MIRA algorithms to switch off in certain cases:

- Forward and reverse complement directions: most sequencing technologies and strategies yield a mixture of reads with both forward and reverse complement direction to the DNA sequenced. In fact, having both directions allows for a much better quality control of an alignment as sequencing technology dependent sequencing errors will often affect only one direction at a given place and not both (the exception being homopolymers and 454).

The MIRA *proposed end clipping* algorithm [-CL:pec] uses this knowledge to initially trim back ends of reads to an area without sequencing errors. However, if reads covering a given area of DNA are present in only one direction, then these reads will be completely eliminated.

If you use only artificial reads in an assembly, then switch off the *proposed end clipping* [-CL:pec=no].

If you mix artificial reads with "normal" reads, make sure that every part of an artificial read is covered by some other read in reverse complement direction (be it a normal or artificial read). The easiest way to do that is to add a reverse complement for every artificial read yourself, though if you use an overlapping strategy with artificial reads, you can calculate the overlaps and reverse complements of reads so that every second artificial read is in reverse complement to save time and memory afterwards during the computation.

- Sequencing type/technology: MIRA currently knows Sanger, 454, Ion Torrent, Solexa, PacBioHQ/LQ and "Text" as sequencing technologies, every read entered in an assembly must be one of those.

Artificial reads should be classified depending on the data they were created from, that is, Sanger for consensus of Sanger reads, 454 for consensus of 454 reads etc. However, should reads created from Illumina consensus be much longer than, say, 200 or 300 bases, you should treat them as Sanger reads.

- Quality values: be careful to assign decent quality values to your artificial reads as several quality clipping or consensus calling algorithms make extensive use of qualities. Pay attention to values of [-CL:qc:bsqc] as well as to [-CO:mrpg:mnq:mgqrt].
- Read lengths: current maximum read length for MIRA is around ~30kb. However, to account for some safety, MIRA currently allows only 20kb reads as maximum length.

9.10.2 Ploidy and repeats

MIRA treats ploidy differences as repeats and will therefore build a separate contigs for the reads of a ploidy that has a difference to the other ploidy/ploidies.

There is simply no other way to handle ploidy while retaining the ability to separate repeats based on differences of only a single base. Everything else would be guesswork. I thought for some time about doing a coverage analysis around the potential repeat/ploidy site, but came to the conclusion that due to the stochastic nature of sequencing data, this would very probably take wrong decisions in too many cases to be acceptable.

If someone has a good idea, I'll be happy to hear it.

9.10.3 Handling of repeats

9.10.3.1 Uniform read distribution

Under the assumption that reads in a project are uniformly distributed across the genome, MIRA will enforce an average coverage and temporarily reject reads from a contig when this average coverage multiplied by a safety factor is reached at a given site. This strategy reduces over-compression of repeats during the contig building phase and keeps reads in reserve for other copies of that repeat.

It's generally a very useful tool disentangle repeats, but has some slight secondary effects: rejection of otherwise perfectly good reads. The assumption of read distribution uniformity is the big problem we have here: of course it's not really valid. You sometimes have less, and sometimes more than "the average" coverage. Furthermore, the new sequencing technologies - 454 perhaps but certainly the ones from Solexa - show that you also have a skew towards the site of replication origin.

Warning: Solexa data from late 2009 and 2010 show a high GC content bias. This bias can reach 200 or 300%, i.e., sequence part for with low GC

One example: let's assume the average coverage of a project is 8 and by chance at one place there 17 (non-repetitive) reads, then the following happens:

(Note: p is the parameter [-AS:urdsip])

Pass 1 to $p-1$: MIRA happily assembles everything together and calculates a number of different things, amongst them an average coverage of ~ 8 . At the end of pass $p-1$, it will announce this average coverage as first estimate to the assembly process.

Pass p : MIRA has still assembled everything together, but at the end of each pass the contig self-checking algorithms now include an "average coverage check". They'll invariably find the 17 reads stacked and decide (looking at the [-AS:ardct] parameter which is assumed to be 2 for this example) that 17 is larger than $2 \cdot 8$ and that this very well may be a repeat. The reads get flagged as possible repeats.

Pass $p+1$ to end: the "possibly repetitive" reads get a much tougher treatment in MIRA. Amongst other things, when building the contig, the contig now looks that "possibly repetitive" reads do not over-stack by an average coverage multiplied by a safety value ([-AS:urdcml]) which we'll assume now to be 1.5 in this example. So, at a certain point, say when read 14 or 15 of that possible repeat want to be aligned to the contig at this given place, the contig will just flatly refuse and tell the assembler to please find another place for them, be it in this contig that is built or any other that will follow. Of course, if the assembler cannot comply, the reads 14 to 17 will end up as contiglet (contig debris, if you want) or if it was only one read that got rejected like this, it will end up as singlet or in the debris file.

Tough luck. I do have ideas on how to re-integrate those reads at the end of an assembly, but I have deferred doing this as in every case I had looked up, adding those reads to the contigs wouldn't have changed anything ... there's already enough coverage.

What should be done in those cases is simply filter away the contiglets (defined as being of small size and having an average coverage below the average coverage of the project divided 3 (or 2.5)) from a project.

9.10.3.2 Keeping 'long' repetitive contigs separate

MIRA had since 2.9.36 a feature to keep long repeats in separate contigs. Due to algorithm changes, this feature is now standard. The effect of this is that contigs with non-repetitive sequence will stop at a 'long repeat' border which cannot be crossed by a single read or by paired reads, including only the first few bases of the repeat. Long repeats will be kept as separate contigs.

This has been implemented to get a clean overview on which parts of an assembly are 'safe' and which parts will be 'difficult'. For this, the naming of the contigs has been extended: contigs named with a '_c' at the end are contigs which contain mostly 'normal' coverage. Contigs with "rep_c" are contigs which contain mostly sequence classified as repetitive and which could not be assembled together with a 'c' contig.

The question remains: what are 'long' repeats? MIRA defines these as repeats that are not spanned by any read that has non-repetitive parts at the end. Basically -for shotgun assemblies - the mean length of the reads that go into the assembly defines the minimum length of 'long' repeats that have to be kept in separate contigs.

It has to be noted that when using paired-end (or template) sequencing, 'long' repeats which can be spanned by read-pairs (or templates) are frequently integrated into 'normal' contigs as MIRA can correctly place them most of the time.

9.10.3.3 Helping finishing by tagging reads with HAF tags

HAF tags (HAsH Frequency) are set by MIRA when the option to colour reads by kmer frequency ([-GE:crkf]), on by default in most --job combinations) is on. These tags show the status of k-mers (stretch of bases of given length k) in read sequences: whether MIRA recognised them as being present in sub-average, average, above average or repetitive numbers.

When using a finishing programs which can display tags in reads (and using the proposed tag colour schemes for gap4 or consed, the assembly will light up in colours ranging from light green to dark red, indicating whether a certain part of the assembly is deemed non-repetitive to extremely repetitive.

One of the biggest advantages of the HAF tags is the implicit information they convey on why the assembler stopped building a contig at an end.

- if the read parts composing a contig end are mostly covered with HAF2 tags (below average frequency, coloured light-green), then one very probably has a hole in the contig due to coverage problems which means there are no or not enough reads covering a part of the sequence.

- if the read parts composing a contig end are mostly covered with HAF3 tags (average frequency, coloured green), then you have an unusual situation as this should only very rarely occur. The reason is that MIRA saw that there are enough sequences which look the same as the one from your contig end, but that these could not be joined. Likely reasons for this scenario include non-random sequencing artifacts (seen in 454 data) or also non-random chimeric reads (seen in Sanger and 454 data).
- if the read parts composing a contig end are mostly covered with HAF4 tags (above average frequency, coloured yellow), then the assembler stopped at grey zone of the coverage not being normal anymore, but not quite repetitive yet. This can happen in cases where the read coverage is very unevenly distributed across the project. The contig end in question might be a repeat occurring two times in the sequence, but having less reads than expected. Or it may be non-repetitive coverage with an unusual excess of reads.
- if the read parts composing a contig end are mostly covered with HAF5 (repeat, coloured red), HAF6 (heavy repeat, coloured darker red) and HAF7 tags (crazy repeat, coloured very dark red), then there is a repetitive area in the sequence which could not be uniquely bridged by the reads present in the assembly.

This information can be especially helpful when joining reads by hand in a finishing program. The following list gives you a short guide to cases which are most likely to occur and what you should do.

- the proposed join involves contig ends mostly covered by HAF2 tags. Joining these contigs is probably a safe bet. The assembly may have missed this join because of too many errors in the read ends or because sequence having been clipped away which could be useful to join contigs. Just check whether the join seems sensible, then join.
- the proposed join involves contig ends mostly covered by HAF3 tags. Joining these contigs is probably a safe bet. The assembly may have missed this join because of several similar chimeric reads or reads with similar, severe sequencing errors covering the same spot. Just check whether the join seems sensible, then join.
- the proposed join involves contig ends mostly covered by HAF4 tags. Joining these contigs should be done with some caution, it may be a repeat occurring twice in the sequence. Check whether the contig ends in question align with ends of several other contigs. If not, joining is probably the way to go. If potential joins exist with several other contigs, then it's a repeat (see below).
- the proposed join involves contig ends mostly covered by HAF5, HAF6 or HAF7 tags. Joining these contigs should be done with utmost caution, you are almost certainly (HAF5) and very certainly (HAF6 and HAF7) in a repetitive area of your sequence. You will probably need additional information like paired-end or template info in order join your contigs.

9.10.4 Consensus in finishing programs (gap4, consed, ...)

MIRA goes a long way to calculate a consensus which is as correct as possible. Unfortunately, communication with finishing programs is a bit problematic as there currently is no standard way to say which reads are from which sequencing technology.

It is therefore often the case that finishing programs calculate an own consensus when loading a project assembled with MIRA. This is the case for at least, e.g., gap4. This consensus may then not be optimal.

The recommended way to deal with this problem is: import the results from MIRA into your finishing program like you always do. Then finish the genome there, export the project from the finishing program as CAF and finally use `miraconvert` (from the MIRA package) with the `-r` option to recalculate the optimal consensus of your finished project.

E.g., assuming you have just finished editing the gap4 database `DEMO . 3`, do the following. First, export the gap4 database back to CAF:

```
$ gap2caf -project DEMO -version 3 >demo3.caf
```

Then, use `miraconvert` with option `'-r'` to convert it into any other format that you need. Example for converting to a CAF and a FASTA format with correct consensus:

```
$ miraconvert -t caf -t fasta -r c demo3.caf final_result
```


9.10.5 Some other things to consider

- MIRA cannot work with EXP files resulting from GAP4 that already have been edited. If you want to reassemble an edited GAP4 project, convert it to CAF format and use the [-caf] option to load.
- As also explained earlier, MIRA relies on sequencing vector being recognised in preprocessing steps by other programs. Sometimes, when a whole stretch of bases is not correctly marked as sequencing vector, the reads might not be aligned into a contig although they might otherwise match quite perfectly. You can use [-CL:pvc] and [-CO:emea] to address problem with incomplete clipping of sequencing vectors. Also having the assembler work with less strict parameters may help out of this.
- MIRA has been developed to assemble shotgun sequencing or EST sequencing data. There are no explicit limitations concerning length or number of sequences. However, there are a few implicit assumptions that were made while writing portions of the code:
 1. Problems which might arise with 'unnatural' long sequence reads: my implementation of the Smith-Waterman alignment routines. I use a banded version with linear running time (linear to the bandwidth) but quadratic space usage. So, comparing two 'reads' of length 5000 will result in memory usage of 95 MiB, two reads with 50000 bases will need 9.5 GiB.

This problem has become acute now with PacBio, I'm working on it. In the mean time, current usable sequence length of PacBio are more in the 3 to 4 kilobase range, with only a few reads attaining or surpassing 20 kb. So Today's machines should still be able to handle the problem more or less effortlessly.
 2. 32 bit versions of MIRA are not supported anymore.
 3. to reduce memory overhead, the following assumptions have been made:
 4. MIRA is not fully multi-threaded (yet), though most bottlenecks are now in code areas which cannot be multi-threaded by algorithm design.
- a project does not contain sequences from more than 255 different:
 - sequencing machine types
 - primers
 - strains (in mapping mode: 7)
 - base callers
 - dyes
 - process status
- a project does not contain sequences from more than 65535 different
 - clone vectors
 - sequencing vectors

9.11 Things you should not do

9.11.1 Do not run MIRA on NFS mounted directories without redirecting the tmp directory

Of course one can run MIRA atop a NFS mount (a "disk" mounted over a network using the NFS protocol), but the performance will go down the drain as the NFS server respectively the network will not be able to cope with the amount of data MIRA needs to shift to and from disk (writes/reads to the tmp directory). Slowdowns of a factor of 10 and more have been observed. In case you have no other possibility, you can force MIRA to run atop a NFS using [-NW:cnfs=warn] ([-NW:cnfs=no]), but you have been warned.

In case you want to keep input and output files on NFS, you can use [-DI:trt] to redirect the tmp directory to a local filesystem. Then MIRA will run at almost full speed.

9.11.2 Do not assemble without quality values

Assembling sequences without quality values is like ... like ... like driving a car downhill a sinuous mountain road with no rails at 200 km/h without brakes, airbags and no steering wheel. With a ravine on one side and a rock face on the other. Did I mention the missing seat-belts? You *might* get down safely, but experience tells the result will rather be a bloody mess.

Well, assembling without quality values is a bit like above, but bloodier. And the worst: you (or the people using the results of such an assembly) will notice the gore only until it is way too late and money has been sunk in follow-up experiments based on wrong data.

All MIRA routines internally are geared toward quality values guiding decisions. No one should ever assembly anything without quality values. Never. Ever. Even if quality values are sometimes inaccurate, they do help.

Now, there are **very rare occasions** where getting quality values is not possible. If you absolutely cannot get them, and I mean only in this case, use the following switch: `--noqualities [=SEQUENCINGTECHNOLOGY]` and additionally give a default quality for reads of a readgroup. E.g.:

```
parameters= --noqualities=454
```

```
readgroup
technology=454
data=...
default_qual=30
```

This tells MIRA not to complain about missing quality values and to fake a quality value of 30 for all reads (of a readgroup) having no qualities, allowing some MIRA routines (in standard parameter settings) to start disentangling your repeats.



Warning Doing the above has some severe side-effects. You will be, e.g., at the mercy of non-random sequencing errors. I suggest combining the above with a `[-CO:mrpg=4]` or higher. You also may want to tune the default quality parameter together with `[-CO:mnq]` and `[-CO:mgqrt]` in cases where you mix sequences with and without quality values.

9.12 Useful third party programs

Viewing the results of a MIRA assembly or preprocessing the sequences for an assembly can be done with a number of different programs. The following ones are just examples, there are a lot more packages available:

HTML browser If you have really nothing else as viewer, a browser who understands tables is needed to view the HTML output. A browser knowing style sheets (CSS) is recommended, as different tags will be highlighted. Konqueror, Opera, Mozilla, Netscape and Internet Explorer all do fine, lynx is not really ... optimal.

Assembly viewer / finishing / preprocessing You'll want GAP4 or its successor GAP5 (generally speaking: the Staden package) to preprocess the sequences, visualise and eventually rework the results when using gap4da output. The Staden package comes with a fully featured sequence preparing and annotating engine (pregap4) that is very useful to preprocess your Sanger data (conversion between file types, quality clipping, tagging etc.).

See <http://www.sourceforge.net/projects/staden/> for further information and also a possibility to download precompiled binaries for different platforms.

Vector screening Reading result files from **ssaha2** or **smalt** from the Sanger Centre is supported directly by MIRA to perform a fast and efficient tagging of sequencing vector stretches. This makes you basically independent from any other commercial or license-requiring vector screening software. For Sanger reads, a combination of **lucy** (see below), **ssaha2** or **smalt** together with the MIRA parameters for SSAHA2 / SMALT support (see all `[-CL:mvs*]` parameters) and quality clipping (`[-CL:qc]`) should do the trick. For reads coming from 454 pyro-sequencing, **ssaha2** or **smalt** and the SSAHA2 / SMALT support also work pretty well.

See <http://www.sanger.ac.uk/resources/software/ssaha2/> and / or <http://www.sanger.ac.uk/resources/software/smalt/> for further information and also a possibility to download the source or pre-compiled binaries for different platforms.

Preprocessing lucy from TIGR (now JCVI) is another useful sequence preprocessing program for Sanger data. Lucy is a utility that prepares raw DNA sequence fragments for sequence assembly. The cleanup process includes quality assessment, confidence reassurance, vector trimming and vector removal.

There's a small script in the MIRA 3rd party package which converts the clipping data from the lucy format into something MIRA can understand (NCBI Traceinfo).

See <ftp://ftp.tigr.org/pub/software/Lucy/> to download the source code of lucy.

Assembly viewer Viewing .ace file output without consed can be done with clview from TIGR. See <http://www.tigr.org/tdb/tgi/software/>.

A better alternative is Tablet <http://bioinf.scri.ac.uk/tablet/> which also reads SAM format.

Assembly coverage analysis The Integrated Genome Browser (IGB) of the GenoViz project at SourceForge (<http://sourceforge.net/projects/genoviz/>) is just perfect for loading a genome and looking at mapping coverage (provided by the wiggle result files of MIRA).

Preprocessing (base calling) TraceTuner (<http://sourceforge.net/projects/tracetuner/>) is a tool for base and quality calling of trace files from DNA sequencing instruments. Originally developed by Paracel, this code base was released as open source in 2006 by Celera.

Preprocessing / viewing phred (basecaller) - cross_match (sequence comparison and filtering) - phrap (assembler) - consed (assembly viewer and editor). This is another package that can be used for this type of job, but requires more programming work. The fact that sequence stretches are masked out (overwritten with the character X) if they shouldn't be used in an assembly doesn't really help and is considered harmful (but it works).

Note the bug of consed when reading ACE files, see more about this in the section on file types (above) in the entry for ACE.

See <http://www.phrap.org/> for further information.

text viewer A text viewer for the different textual output files.

As always, most of the time a combination of several different packages is possible. My currently preferred combo for genome projects is **ssaha2** or **smalt** and or **lucy** (vector screening), MIRA (assembly, of course) and gap4 (assembly viewing and finishing).

For re-assembling projects that were edited in gap4, one will also need the gap2caf converter. The source for this is available at <http://www.sanger.ac.uk/resources/software/caf.html>.

9.13 Speed and memory considerations

9.13.1 Estimating needed memory for an assembly project

Since the V2.9.24x3 version of MIRA, there is **miramem** as program call. When called from the command line, it will ask a number of questions and then print out an estimate of the amount of RAM needed to assemble the project. Take this estimate with a grain of salt, depending on the sequences properties, variations in the estimate can be +/- 30% for bacteria and 'simple' eukaryotes. The higher the number of repeats is, the more likely you will need to restrict memory usage in some way or another.

Here's the transcript of a session with miramem:

```
This is MIRA V3.2.0rc1 (development version).
```

```
Please cite: Chevreur, B., Wetter, T. and Suhai, S. (1999), Genome Sequence Assembly Using Trace Signals and Additional Sequence Information. Computer Science and Biology: Proceedings of the German Conference on Bioinformatics (GCB) 99, pp. 45-56.
```

```
To (un-)subscribe the MIRA mailing lists, see:  
http://www.chevreur.org/mira\_mailinglists.html
```

After subscribing, mail general questions to the MIRA talk mailing list:
 mira_talk@freelists.org

To report bugs or ask for features, please use the SourceForge ticketing system at:

<http://sourceforge.net/p/mira-assembler/tickets/>

This ensures that requests do not get lost.

[...]

miraMEM helps you to estimate the memory needed to assemble a project. Please answer the questions below.

Defaults are give in square brackets and chosen if you just press return. Hint: you can add k/m/g modifiers to your numbers to say kilo, mega or giga.

Is it a genome or transcript (EST/tag/etc.) project? (g/e/) [g]

g

Size of genome? [4.5m] **9.8m**

9800000

Size of largest chromosome? [9800000]

9800000

Is it a denovo or mapping assembly? (d/m/) [d]

d

Number of Sanger reads? [0]

0

Are there 454 reads? (y/n/) [n] **y**

y

Number of 454 GS20 reads? [0]

0

Number of 454 FLX reads? [0]

0

Number of 454 Titanium reads? [0] **750k**

750000

Are there PacBio reads? (y/n/) [n]

n

Are there Solexa reads? (y/n/) [n]

n

***** Estimates *****

The contigs will have an average coverage of ~ 30.6 (+/- 10%)

RAM estimates:

reads+contigs (unavoidable): 7.0 GiB

large tables (tunable): 688. MiB

 total (peak): 7.7 GiB

add if using -CL:pvlc=yes : 2.6 GiB

Estimates may be way off for pathological cases.

Note that some algorithms might try to grab more memory if the need arises and the system has enough RAM. The options for automatic memory management control this:

-AS:amm, -AS:kpmf, -AS:mps

Further switches that might reduce RAM (at cost of run time or accuracy):

-SK:mkim, -SK:mchr (both runtime); -SK:mhpr (accuracy)

If your RAM is not large enough, you can still assemble projects by using disk swap. Up to 20% of the needed memory can be provided by swap without the speed penalty getting too large. Going above 20% is not recommended though, above 30% the machine will be almost permanently swapping at some point or another.

9.13.2 Some numbers on speed

To be rewritten for MIRA4.

9.14 Known Problems / Bugs

File Input / Output:

1. MIRA can only read unedited EXP files.
2. There sometimes is a (rather important) memory leak occurring while using the assembly integrated Sanger read editor. I have not been able to trace the reason yet.

Assembly process:

1. The routines for determining *Repeat Marker Bases* (SRMr) are sometimes too sensitive, which sometimes leads to excessive base tagging and preventing right assemblies in subsequent assembly processes. The parameters you should look at for this problem are [-CO:mrc:nrz:mgqrt:mgqwpc]. Also look at [-CL:pvc] and [-CO:emea] if you have a lot of sequencing vector relics at the end of the sequences.

9.15 TODOs

These are some of the topics on my TODO list for the next revisions to come:

1. Making Smith-Waterman parts of the process multi-threaded or use SIMD (currently stopped due to other priorities like PacBio etc.)

9.16 Working principles

Note: description is old and needs to be adapted to the current 4.x line of MIRA.

To avoid the "garbage-in, garbage-out" problematic, MIRA uses a 'high quality alignments first' contig building strategy. This means that the assembler will start with those regions of sequences that have been marked as good quality (high confidence region - HCR) with low error probabilities (the clipping must have been done by the base caller or other preprocessing programs, e.g. pregap4) and then gradually extends the alignments as errors in different reads are resolved through error hypothesis verification and signal analysis.

This assembly approach relies on some of the automatic editing functionality provided by the EdIt package which has been integrated in parts within MIRA.

This is an approximate overview on the steps that are executed while assembling:

1. All the experiment / phd / fasta sequences that act as input are loaded (or the CAF project). Qualities for the bases are loaded from the FASTA or SCF if needed.
2. the ends of the reads are cleaned ensure they have a minimum stretch of bases without sequencing errors

3. The high confidence region (HCR) of each read is compared with a quick algorithm to the HCR of every other read to see if it could match and have overlapping parts (this is the 'SKIM' filter).
4. All the reads which could match are being checked with an adapted Smith-Waterman alignment algorithm (banded version). Obvious mismatches are rejected, the accepted alignments form one or several alignment graphs.
5. Optional pre-assembly read extension step: MIRA tries to extend HCR of reads by analysing the read pairs from the previous alignment. This is a bit shaky as reads in this step have not been edited yet, but it can help. Go back to step 2.
6. A contig gets made by building a preliminary partial path through the alignment graph (through in-depth analysis up to a given level) and then adding the most probable overlap candidates to a given contig. Contigs may reject reads if these introduce too many errors in the existing consensus. Errors in regions known as dangerous (for the time being only ALUS and REPT) get additional attention by performing simple signal analysis when alignment discrepancies occur.
7. Optional: the contig can be analysed and corrected by the automatic editor ("EdIt" for Sanger reads, or the new MIRA editor for 454 reads).
8. Long repeats are searched for, bases in reads of different repeats that have been assembled together but differ sufficiently (for EdIT so that they didn't get edited and by phred quality value) get tagged with special tags (SRMr and WRMr).
9. Go back to step 5 if there are reads present that have not been assembled into contigs.
10. Optional: Detection of spoiler reads that prevent joining of contigs. Remedy by shortening them.
11. Optional: Write out a checkpoint assembly file and go back to step 2.
12. The resulting project is written out to different output files and directories.

9.17 See Also

The other MIRA manuals and walkthroughs as well as **EdIt**, **gap4**, **pregap4**, **gap5**, **clview**, **caf2gap**, **gap2caf**, **ssaha2**, **smalt**, **compress** and **gzip**, **cap3**, **ttuner**, **phred**, **phrap**, **cross_match**, **consed**, **pilon**.

Chapter 10

Assembly of *hard* genome or EST / RNASeq projects

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“If it were easy, it would have been done already.”

—Solomon Short

10.1 Getting 'mean' genomes or EST / RNASeq data sets assembled

For some EST data sets you might want to assemble, MIRA will take too long or the available memory will not be sufficient. For genomes this can be the case for eukaryotes, plants, but also for some bacteria which contain high number of (pro-)phages, plasmids or engineered operons. For EST data sets, this concerns all projects with non-normalised libraries.

This guide is intended to get you through these problematic genomes. It is (cannot be) exhaustive, but it should get you going.

10.1.1 For the impatient

For bacteria with nasty repeats, try first [--hirep_something]. This will increase runtime and memory requirements, but helps to get this sorted out. If the data for lower eukaryotes leads to runtime and memory explosion, try either [--hirep_good] or, for desperate cases, [--hirep_something].

10.1.2 Introduction to 'masking'

The SKIM phase (all-against-all comparison) will report almost every potential hit to be checked with Smith-Waterman further downstream in the MIRA assembly process. While this is absolutely no problem for most bacteria, some genomes (eukaryotes, plants, some bacteria) have so many closely related sequences (repeats) that the data structures needed to take up all information might get much larger than your available memory. In those cases, your only chance to still get an assembly is to tell the assembler it should disregard extremely repetitive features of your genome.

There is, in most cases, one problem: one doesn't know beforehand which parts of the genome are extremely repetitive. But MIRA can help you here as it produces most of the needed information during assembly and you just need to choose a threshold from where on MIRA won't care about repetitive matches.

The key to this are the three fail-safe command line parameters which will mask "nasty" repeats from the quick overlap finder (SKIM): [-KS:mnr] and [-KS:nrr] respectively [-KS:nrc]. I'll come back to [-SK:kms] later as it also plays a role in this.

10.1.3 How does 'nasty repeat' masking work?

If switched on [-KS:mnr=yes], MIRA will use k-mer statistics to find repetitive stretches. K-mers are nucleotide stretches of length k. In a perfectly sequenced genome without any sequencing error and without sequencing bias, the k-mer frequency can be used to assess how many times a given nucleotide stretch is present in the genome: if a specific k-mer is present as many times as the average frequency of all k-mers, it is a reasonable assumption to estimate that the specific k-mer is not part of a repeat (at least not in this genome).

Following the same path of thinking, if a specific k-mer frequency is now two times higher than the average of all k-mers, one would assume that this specific k-mer is part of a repeat which occurs exactly two times in the genome. For 3x k-mer frequency, a repeat is present three times. Etc.pp. MIRA will merge information on single k-mers frequency into larger 'repeat' stretches and tag these stretches accordingly.

Of course, low-complexity nucleotide stretches (like poly-A in eukaryotes), sequencing errors in reads and non-uniform distribution of reads in a sequencing project will weaken the initial assumption that a k-mer frequency is representative for repeat status. But even then the k-mer frequency model works quite well and will give a pretty good overall picture: most repeats will be tagged as such.

Note that the parts of reads tagged as "nasty repeat" will not get masked per se, the sequence will still be present. The stretches dubbed repetitive will get the "MNRr" tag. They will still be used in Smith-Waterman overlaps and will generate a correct consensus if included in an alignment, but they will not be used as seed.

Some reads will invariably end up being completely repetitive. These will not be assembled into contigs as MIRA will not see overlaps as they'll be completely masked away. These reads will end up as debris. However, note that MIRA is pretty good at discerning 100% matching repeats from repeats which are not 100% matching: if there's a single base with which repeats can be discerned from each other, MIRA will find this base and use the k-mers covering that base to find overlaps.

10.1.4 Selecting a "nasty repeat ratio"

The ratio from which on the MIRA kmer statistics algorithm won't report matches is set via [-KS:nrr]. E.g., using [-KS:nrr=10] will hide all k-mers which occur at a frequency 10 times (or more) higher than the median of all k-mers.

The nastiness of a repeat is difficult to judge, but starting with 10 copies in a genome, things can get complicated. At 20 copies, you'll have some troubles for sure.

The standard values of 10 for the [-KS:nrr] parameter is a pretty good 'standard' value which can be tried for an assembly before trying to optimise it via studying the kmer statistics calculated by MIRA. For the later, please read the section 'Examples for kmer statistics' further down in this guide.

10.2 How MIRA tags different repeat levels

During SKIM phase, MIRA will assign frequency information to each and every k-mer in all reads of a sequencing project, giving them different status. Additionally, tags are set in the reads so that one can assess reads in assembly editors that understand tags (like gap4, gap5, consed etc.). The following tags are used:

HAF2 coverage below average (default: < 0.5 times average)

HAF3 coverage is at average (default: ≥ 0.5 times average and ≤ 1.5 times average)

HAF4 coverage above average (default: > 1.5 times average and < 2 times average)

HAF5 probably repeat (default: ≥ 2 times average and < 5 times average)

HAF6 'crazy' repeat (default: > 5 times average)

MNRr stretches which were masked away by [-KS:mnr=yes] being more repetitive than deduced by [-KS:nrr=...] or given via [-KS:nrc=...].

10.3 The readrepeats info file

If `[-KS:mnr=yes]` is used, MIRA will write an additional file into the info directory: `<projectname>_info_readrepeats.lst`

The "readrepeats" file makes it possible to try and find out what makes sequencing data nasty. It's a key-value-value file with the name of the sequence as "key" and then the type of repeat (HAF2 - HAF7 and MNRr) and the repeat sequence as "values". "Nasty" in this case means *everything which was masked via [-KS:mnr=yes]*.

The file looks like this:

```
read1      HAF5      GCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCT ...
read2      HAF7      CCGAAGCCGAAGCCGAAGCCGAAGCCGAAGCCGAAGCCGAAGCCGAAGCCGAAGCCGAAGC ...
read2      MNRr      AAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA ...
read3      HAF6      GCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCTTCGGCT ...
...
etc.
```

That is, each line consists of the read name where a stretch of repetitive sequences was found, then the MIRA repeat categorisation level (HAF2 to HAF7 and MNRr) and then the stretch of bases which is seen to be repetitive.

Note that reads can have several disjunct repeat stretches in a single read, hence they can occur more than one time in the file as shown with *read2* in the example above.

One will need to search some databases with the "nasty" sequences and find vector sequences, adaptor sequences or even human sequences in bacterial or plant genomes ... or vice versa as this type of contamination happens quite easily with data from new sequencing technologies. After a while one gets a feeling what constitutes the largest part of the problem and one can start to think of taking countermeasures like filtering, clipping, masking etc.

10.4 Pipeline to find worst contaminants or repeats in sequencing data

Note

In case you are not familiar with UNIX pipes, now would be a good time to read an introductory text on how this wonderful system works. You might want to start with a short introductory article at Wikipedia: http://en.wikipedia.org/wiki/Pipeline_%28Unix%29

In a nutshell: instead of output to files, a pipe directs the output of one program as input to another program.

There's one very simple trick to find out whether your data contains some kind of sequencing vector or adaptor contamination which I use. it makes use of the read repeat file discussed above.

The following example shows this exemplarily on a 454 data where the sequencing provider used some special adaptor in the wet lab but somehow forgot to tell the Roche pre-processing software about it, so that a very large fraction of reads in the SFF file had unclipped adaptor sequence in it (which of course wreaks havoc with assembly programs):

```
arcadia:$ grep MNRr badproject_info_readrepeats.lst | cut -f 3 | sort | uniq -c | sort -g -r <-
| head -15
504 ACCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
501 CAACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
489 GGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
483 GCCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
475 AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
442 GATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
429 CGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
424 TTGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
393 ACTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
379 CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
363 ATTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
343 CATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
```

```
334 GTTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
328 AACACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
324 GGTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
```

You probably see a sequence pattern `CTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC` in the above screens hot. Before going into details of what you are actually seeing, here's the explanation how this pipeline works:

grep MNRr badproject_info_readrepeats.lst From the file with the information on repeats, grab all the lines containing repetitive sequence which MIRA categorised as 'nasty' via the 'MNRr' tag. The result looks a bit like this (first 15 lines shown):

```
C6E3C7T12GKN35 MNRr GCGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12JLIBM MNRr TTCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12HQOM1 MNRr CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12G52II MNRr CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12JRMPO MNRr TCTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12H1A8V MNRr GCGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12H34Z7 MNRr AAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12H4HGC MNRr GGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12FNA1N MNRr AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12F074V MNRr CTTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12I1GYO MNRr CAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12I53C8 MNRr CACACTCGTATAGTGACACGCAACAGGGG
C6E3C7T12I4V6V MNRr ATCACTCGTATAGTGACACGCAACAGGGG
C6E3C7T12H5R00 MNRr TCTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
C6E3C7T12IBA5E MNRr AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
...
```

cut -f 3 We're just interested in the sequence now, which is in the third column. The above 'cut' command takes care of this. The resulting output may look like this (only first 15 lines shown):

```
GCGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
TTCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
TCTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
GCGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
AAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
GGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
CTTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
CAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
CACACTCGTATAGTGACACGCAACAGGGG
ATCACTCGTATAGTGACACGCAACAGGGG
TCTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
...
```

sort Simply sort all sequences. The output may look like this now (only first 15 line shown):

```
AAACCTCGTATAGTGACACGCA
AAACCTCGTATAGTGACACGCAACAGG
AAACCTCGTATAGTGACACGCAACAGGG
AAACCTCGTATAGTGACACGCAACAGGGG
AAACCTCGTATAGTGACACGCAACAGGGG
AAACCTCGTATAGTGACACGCAACAGGGG
AAACCTCGTATAGTGACACGCAACAGGGG
AAACCTCGTATAGTGACACGCAACAGGGG
AAACCTCGTATAGTGACACGCAACAGGGGAT
AAACCTCGTATAGTGACACGCAACAGGGGATA
AAACCTCGTATAGTGACACGCAACAGGGGATA
```

```
AAACTCGTATAGTGACACGCAACAGGGGATA
AAACTCGTATAGTGACACGCAACAGGGGATA
AAACTCGTATAGTGACACGCAACAGGGGATA
AAACTCGTATAGTGACACGCAACAGGGGATA
...
```

uniq -c This command counts how often a line repeats itself in a file. As we previously sorted the whole file by sequence, it effectively counts how often a certain sequence has been tagged as MNRr. The output consists of a tab delimited format in two columns: the first column contains the number of times a given line (sequence in our case) was seen, the second column contains the line (sequence) itself. An exemplarily output looks like this (only first 15 lines shown):

```
1 AAACCTCGTATAGTGACACGCA
1 AAACCTCGTATAGTGACACGCAACAGG
1 AAACCTCGTATAGTGACACGCAACAGGG
5 AAACCTCGTATAGTGACACGCAACAGGGG
1 AAACCTCGTATAGTGACACGCAACAGGGGAT
13 AAACCTCGTATAGTGACACGCAACAGGGGATA
6 AAACCTCGTATAGTGACACGCAACAGGGGATAGAC
4 AAACCTCGTATAGTGACACGCAACAGGGGATAGACAA
9 AAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGC
3 AAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCA
257 AAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
1 AACACTCGTATAGTGACACGCAAC
2 AACACTCGTATAGTGACACGCAACAGGG
23 AACACTCGTATAGTGACACGCAACAGGGG
6 AACACTCGTATAGTGACACGCAACAGGGGATA
...
```

sort -g -r We now sort the output of the previous `uniq`-counting command by asking 'sort' to perform a numerical sort (via '-g') and additionally sort in reverse order (via '-r') so that we get the sequences encountered most often at the top of the output. And that one looks exactly like shown previously:

```
504 ACCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
501 CAACCTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
489 GGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
483 GCCACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
475 AATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
442 GATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
429 CGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
424 TTGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
393 ACTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
379 CAGACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
363 ATTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
343 CATACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
334 GTTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
328 AACACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
324 GGTACTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC
...
```

So, what is this ominous `CTCGTATAGTGACACGCAACAGGGGATAGACAAGGCAC` you are seeing? To make it short: a modified 454 B-adaptor with an additional MID sequence.

Note

These adaptor sequences have absolutely no reason to exist in your data, none! Go back to your sequencing provider and ask them to have a look at their pipeline as they should have had it set up in a way that you do not see these things anymore. Yes, due to sequencing errors, sometimes some adaptor or sequencing vectors remnants will stay in your sequencing data, but that is no problem as MIRA is capable of handling that very well.

But having much more than 0.1% to 0.5% of your sequence containing these is a sure sign that someone goofed somewhere ... and it's very probably not your fault.

10.5 Examples for kmer statistics

Selecting the right ratio so that an assembly fits into your memory is not straight forward. But MIRA can help you a bit: during assembly, some frequency statistics are printed out (they'll probably end up in some info file in later releases). Search for the term "Kmer statistics" in the information printed out by MIRA (this happens quite early in the process)

10.5.1 Caveat: -SK:kmer_size

Some explanation how kmer size affects the statistics and why it should be chosen ≥ 16 for [-KS:mnr]

10.5.2 Sanger sequencing, a simple bacterium

This example is taken from a pretty standard bacterium where Sanger sequencing was used:

```
Kmer statistics:
=====
Measured avg. coverage: 15

Deduced thresholds:
-----
Min normal cov: 7
Max normal cov: 23
Repeat cov: 29
Crazy cov: 120
Mask cov: 150

Repeat ratio histogram:
-----
0      475191
1      5832419
2      181994
3       6052
4       4454
5        972
6         4
7         8
14        2
16       10
=====
```

The above can be interpreted like this: the expected coverage of the genome is 15x. Starting with an estimated kmer frequency of 29, MIRA will treat a k-mer as 'repetitive'. As shown in the histogram, the overall picture of this project is pretty healthy:

- only a small fraction of k-mers have a repeat level of '0' (these would be k-mers in regions with quite low coverage or k-mers containing sequencing errors)
- the vast majority of k-mers have a repeat level of 1 (so that's non-repetitive coverage)
- there is a small fraction of k-mers with repeat level of 2-10
- there are almost no k-mers with a repeat level >10

10.5.3 454 Sequencing, a somewhat more complex bacterium

Here's in comparison a profile for a more complicated bacterium (454 sequencing):

Kmer statistics:

```
=====
Measured avg. coverage: 20
```

Deduced thresholds:

```
-----
Min normal cov: 10
Max normal cov: 30
Repeat cov: 38
Crazy cov: 160
Mask cov: 0
```

Repeat ratio histogram:

```
-----
0      8292273
1      6178063
2      692642
3      55390
4      10471
5      6326
6      5568
7      3850
8      2472
9      708
10     464
11     270
12     140
13     136
14     116
15     64
16     54
17     54
18     52
19     50
20     58
21     36
22     40
23     26
24     46
25     42
26     44
27     32
28     38
29     44
30     42
31     62
32     116
33     76
34     80
35     82
36     142
37     100
38     120
39     94
40     196
41     172
42     228
43     226
44     214
45     164
46     168
```

```

47      122
48      116
49      98
50      38
51      56
52      22
53      14
54      8
55      2
56      2
57      4
87      2
89      6
90      2
92      2
93      2
1177    2
1181    2
=====

```

The difference to the first bacterium shown is pretty striking:

- first, the k-mers in repeat level 0 (below average) is higher than the k-mers of level 1! This points to a higher number of sequencing errors in the 454 reads than in the Sanger project shown previously. Or at a more uneven distribution of reads (but not in this special case).
- second, the repeat level histogram does not trail off at a repeat frequency of 10 or 15, but it has a long tail up to the fifties, even having a local maximum at 42. This points to a small part of the genome being heavily repetitive ... or to (a) plasmid(s) in high copy numbers.

Should MIRA ever have problems with this genome, switch on the nasty repeat masking and use a level of 15 as cutoff. In this case, 15 is OK to start with as a) it's a bacterium, it can't be that hard and b) the frequencies above level 5 are in the low thousands and not in the tens of thousands.

10.5.4 Solexa sequencing, E.coli MG1655

```

Kmer statistics:
=====
Measured avg. coverage: 23

Deduced thresholds:
-----
Min normal cov: 11
Max normal cov: 35
Repeat cov: 44
Crazy cov: 184
Mask cov: 0

Repeat ratio histogram:
-----
0      1365693
1      8627974
2      157220
3      11086
4      4990
5      3512
6      3922
7      4904
8      3100

```

9	1106
10	868
11	788
12	400
13	186
14	28
15	10
16	12
17	4
18	4
19	2
20	14
21	8
25	2
26	8
27	2
28	4
30	2
31	2
36	4
37	6
39	4
40	2
45	2
46	8
47	14
48	8
49	4
50	2
53	2
56	6
59	4
62	2
63	2
67	2
68	2
70	2
73	4
75	2
77	4

=====

This kmer statistics shows that MG1655 is pretty boring (from a repetitive point of view). One might expect a few repeats but nothing fancy: The repeats are actually the rRNA and sRNA stretches in the genome plus some intergenic regions.

- the k-mers number in repeat level 0 (below average) is considerably lower than the level 1, so the Solexa sequencing quality is pretty good respectively there shouldn't be too many low coverage areas.
- the histogram tail shows some faint traces of possibly highly repetitive k-mers, but these are false positive matches due to some standard Solexa base-calling weaknesses of earlier pipelines like, e.g., adding poly-A, poly-T or sometimes poly-C and poly-G tails to reads when spots in the images were faint and the base calls of bad quality

10.5.5 (NEED EXAMPLES FOR EUKARYOTES)

10.5.6 (NEED EXAMPLES FOR PATHOLOGICAL CASES)

Vector contamination etc.

Chapter 11

Description of sequencing technologies

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Opinions are like chili powder - best used in moderation.”

—Solomon Short

11.1 Introduction

Note: This section contains things I’ve seen in the past and simply jotted down. These may be fundamentally correct or correct only under circumstances or not correct at all with your data. You may have different observations.

...

11.2 Illumina (formerly Solexa)

11.2.1 Caveats for Illumina data

Note

Even if you can get bacteria sequenced with ridiculously high coverage like 500x or 1000x, this amount of data is simply not needed. Even more important - though counterintuitive - is the fact that due to non-random sequence dependent sequencing errors, a too high coverage may even make the assembly worse.

Another rule of thumb: when having more than enough data, reduce the data set so as to have an average coverage of approximately 100x. In some rare cases (high GC content), perhaps 120x to 150x, but certainly not more.



Warning When reducing a data set, do **NOT**, under no circumstances not, try fancy selection of reads by some arbitrary quality or length criteria. This will introduce a terrible bias in your assembly due to non-random sequence-dependent sequencing errors and non-random sequence dependent base quality assignment. More on this in the next section.

11.2.2 Illumina highlights

11.2.2.1 Quality

For current HiSeq 100bp reads I get - after MIRA clipping - about 90 to 95% reads matching to a reference without a single error. MiSeq 250bp reads contain a couple more errors, but nothing to be alarmed off.

In short: Illumina is currently *the* technology to use if you want high quality reads.

11.2.3 Lowlights

11.2.3.1 Long homopolymers

Long homopolymers (stretches of identical bases in reads) can be a slight problem for Solexa. However, it must be noted that this is a problem of all sequencing technologies on the market so far (Sanger, Solexa, 454). Furthermore, the problem is much less pronounced in Solexa than in 454 data: in Solexa, first problem appears may appear in stretches of 9 to 10 bases, in Ion Torrent a stretch of 3 to 4 bases may already start being problematic in some cases.

11.2.3.2 The GGCxG and GGC motifs

GGCxG or even GGC motif in the 5' to 3' direction of reads. This one is particularly annoying and it took me quite a while to circumvent in MIRA the problems it causes.

Simply put: at some places in a genome, base calling after a GGCxG or GGC motif is particularly error prone, the number of reads without errors declines markedly. Repeated GGC motifs worsen the situation. The following screen shots of a mapping assembly illustrate this.

The first example is the GGCxG motif (in form of a GGCTG) occurring in approximately one third of the reads at the shown position. Note that all but one read with this problem are in the same (plus) direction.

The screenshot shows a sequence alignment in the Contig Editor. The reference sequence is GGAATAGAACAGATTTTGGCAAAAATATCCCACAGCGGCTGTGGGATAATTGAGACAGTTCAGACACGTTAAGTCTCCT. A red box highlights a region of the reads where the motif GGCTG is present, and a red arrow points to the G at the end of this motif in the consensus sequence. The reads are numbered on the left, ranging from +54656 to +54731. The x-axis at the top shows positions from 230 to 260. The bottom status bar indicates 'Base confidence:1 (Probability 0.205672) Position 260355'.

Figure 11.1: The Solexa GGCxG problem.

The next two screen shots show the GGC, once for forward direction and one with reverse direction reads:

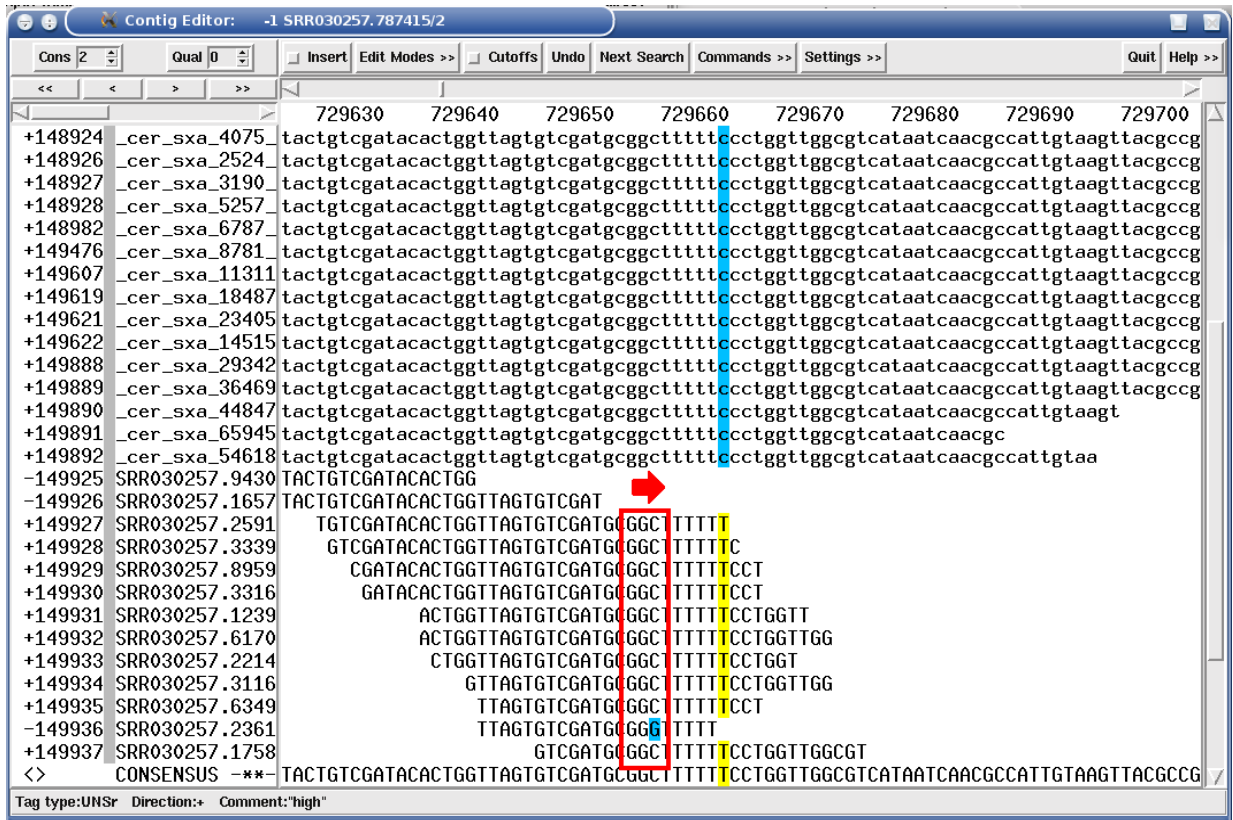


Figure 11.2: The Solexa GGC problem, forward example

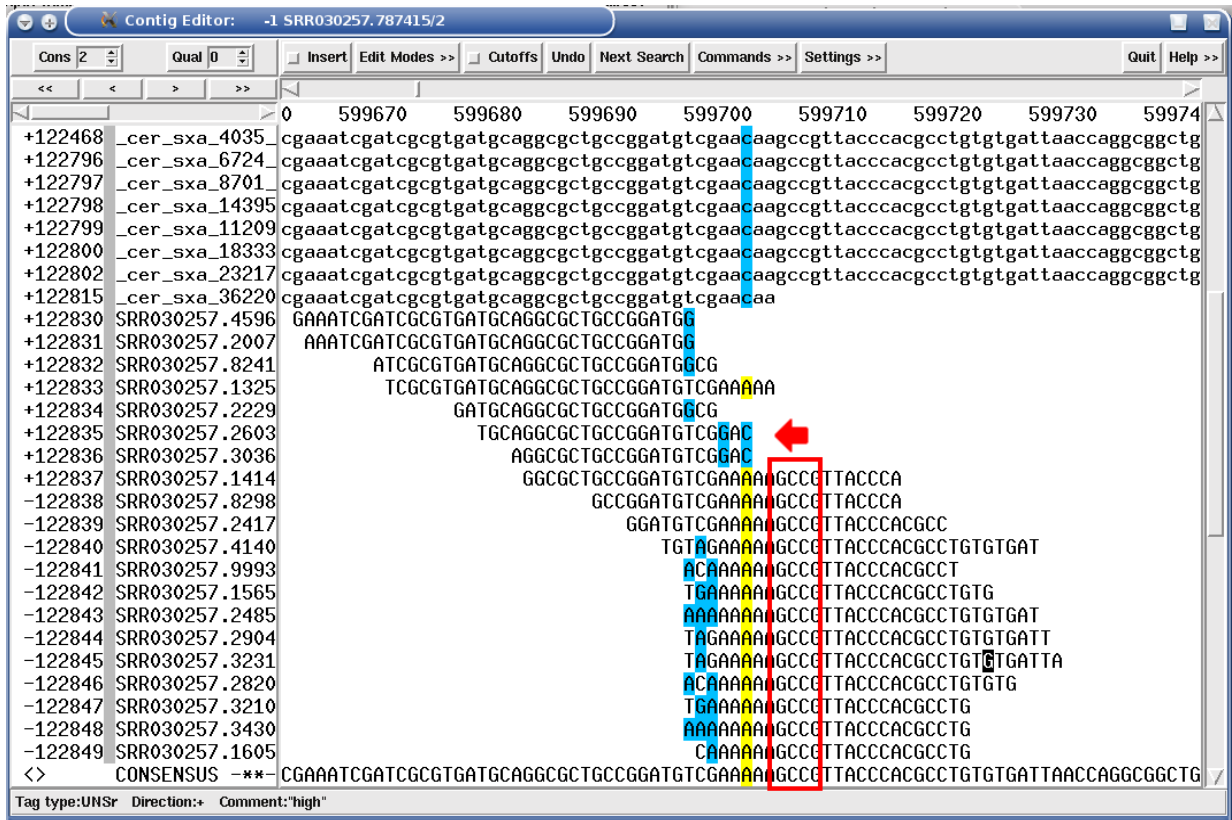


Figure 11.3: The Solexa GGC problem, reverse example

Places in the genome that have GGCGGC GCCGCC (a motif, perhaps even repeated, then some bases and then an inverted motif) almost always have very, very low number of good reads. Especially when the motif is GGCxG.

Things get especially difficult when these motifs occur at sites where users may have a genuine interest. The following example is a screen shot from the Lenski data (see walk-through below) where a simple mapping reveals an anomaly which -- in reality -- is an IS insertion (see http://www.nature.com/nature/journal/v461/n7268/fig_tab/nature08480_F1.html) but could also look like a GGCxG motif in forward direction (GGCCG) and at the same time a GGC motif in reverse direction:

Note that projects using reads ≤ 100 bp assembled fine with MIRA 4.9.3 and before as the default algorithms for proposed-end-clip (`[-CL:pec]`) implicitly caught chimeras occurring near the read ends and the remaining chimeras were caught by the algorithms for low level chimeras.



Warning MIRA 4.9.4 and higher eliminate all chimeras in Illumina reads of any length, you do not need to take any precautionary steps here. But if you use other assemblers and in light of the above, I highly recommend to apply very stringent filters to Illumina data. Especially for applications like metagenomics or RNA de-novo assembly where low coverage may be expected for parts of the results! Indeed, I now treat any assembly result with consensus data generated from a coverage of less than 3 Illumina reads as potentially junk data.

11.2.3.4 Sample barcode misidentification

Long story short: data from multiplexed samples contains "low" amounts of foreign samples from the same lane. Probably not a problem for high coverage assemblies, but can become a problem in multiplexed RNASeq or projects looking for "rare" variants.

In essence, the barcoding used for multiplexing several samples into a single lane is not a 100% foolproof process. I found one paper quantifying this effect to 0.3% of misidentified reads: Kircher et al., *Double indexing overcomes inaccuracies in multiplex sequencing on the Illumina platform*; Nucleic Acids Res. Jan 2012; 40(1): e3. <http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3245947/>

For example, I got some genome sequencing data for a bacterium where closer inspection of some small contigs coming out of the assembly process turned out to be highly expressed genes from a plant. The sequencing provider had multiplexed our bacterial sample with a RNASeq project of that plant.

Another example involved RNASeq of two genomes where one of the organisms had been modified to contain additional genes under a strong promoter. In the data set we suddenly saw those inserted genes pop-up in the samples of the wild type organism. Which, clearly, could not be.

11.2.3.5 Nextera library prep

Opinions seem to be divided about Nextera: some people don't like it as it introduces sometimes terrible coverage bias in the data, other people say they're happy with the data.

Someone told me (or wrote, I do not remember) that this divide may be due to the fact that some people use their sequencing data for de-novo assemblies, while others just do mappings and hunt for SNPs. In fact, this would explain a lot: for de-novo assemblies, I would never use Nextera. When on a hunt for SNPs, they may be OK.

11.2.3.6 Strong GC bias in some Solexa data (2nd half 2009 until advent of TruSeq kit at end of 2010)

I'm recycling a few slides from a couple of talks I held in 2010.

Things used to be so nice and easy with the early Solexa data I worked with (36 and 44mers) in late 2007 / early 2008. When sample taking was done right -- e.g. for bacteria: in stationary phase -- and the sequencing lab did a good job, the read coverage of the genome was almost even. I did see a few papers claiming to see non-trivial GC bias back then, but after having analysed the data I worked with I dismissed them as "not relevant for my use cases." Have a look at the following figure showing exemplarily the coverage of a 45% GC bacterium in 2008:

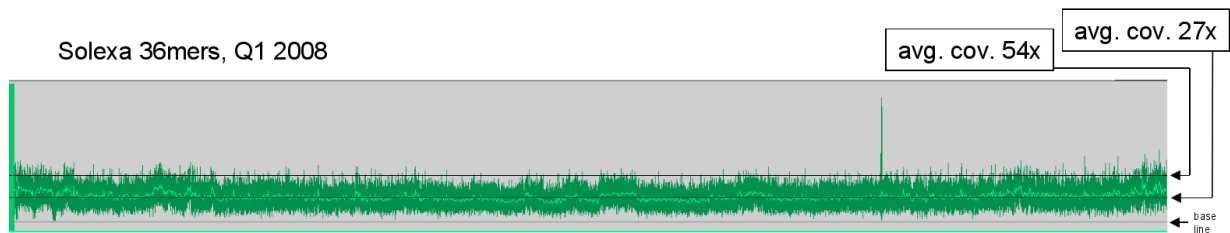


Figure 11.5: Example for no GC coverage bias in 2008 Solexa data. Apart from a slight *smile shape* of the coverage -- indicating the sample taking was not 100% in stationary phase of the bacterial culture -- everything looks pretty nice: the average coverage is at 27x, and when looking at potential genome duplications at twice the coverage (54x), there's nothing apart a single peak (which turned out to be a problem in a rRNA region).

Things changed starting sometime in Q3 2009, at least that's when I got some data which made me notice a problem. Have a look at the following figure which shows exactly the same organism as in the figure above (bacterium, 45% GC):

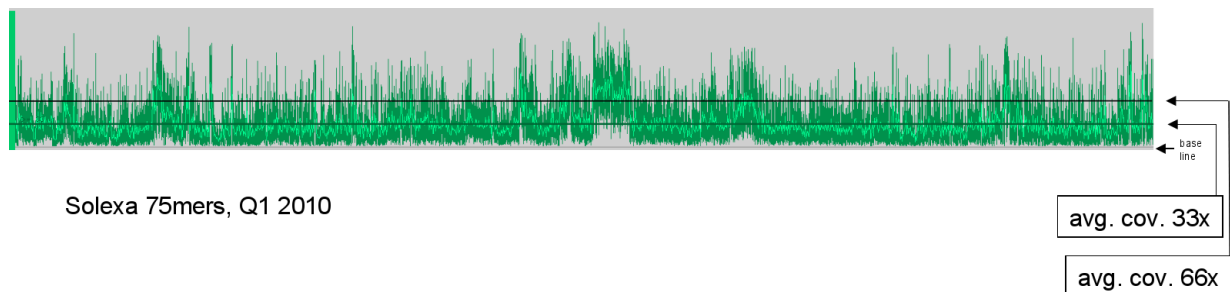


Figure 11.6: Example for GC coverage bias starting Q3 2009 in Solexa data. There's no *smile shape* anymore -- the people in the lab learned to pay attention to sample in 100% stationary phase -- but something else is extremely disconcerting: the average coverage is at 33x, and when looking at potential genome duplications at twice the coverage (66x), there are several dozen peaks crossing the 66x threshold over a several kilobases (in one case over 200 Kb) all over the genome. As if several small genome duplications happened.

By the way, the figures above are just examples: I saw over a dozen sequencing projects in 2008 without GC bias and several dozen in 2009 / 2010 with GC bias.

Checking the potential genome duplication sites, they all looked "clean", i.e., the typical genome insertion markers are missing. Poking around at possible explanations, I looked at GC content of those parts in the genome ... and there was the explanation:

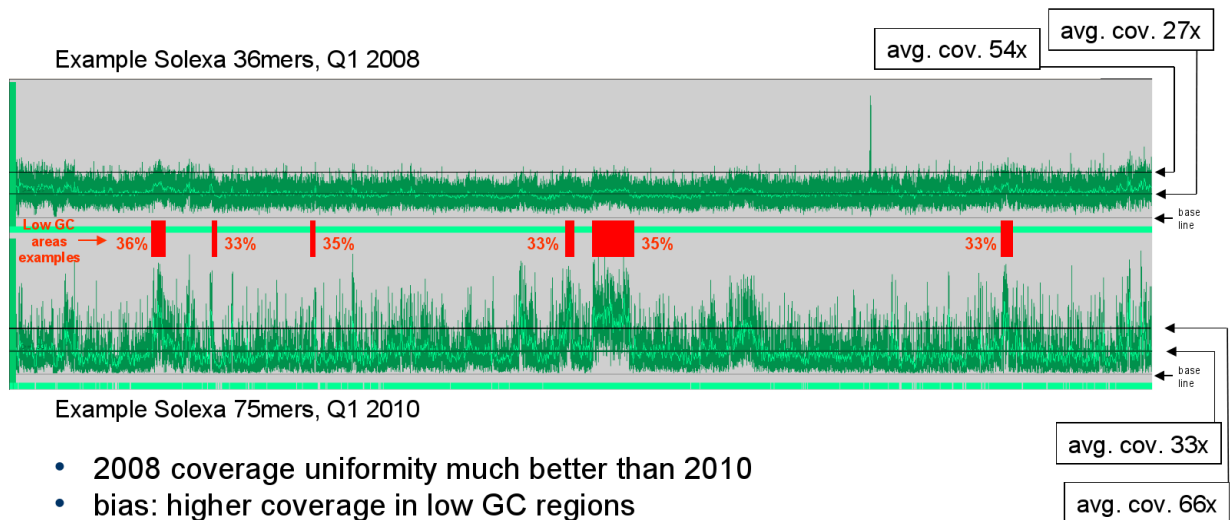


Figure 11.7: Example for GC coverage bias, direct comparison 2008 / 2010 data. The bug has 45% average GC, areas with above average read coverage in 2010 data turn out to be lower GC: around 33 to 36%. The effect is also noticeable in the 2008 data, but barely so.

Now as to actually *why* the GC bias suddenly became so strong is unknown to me. The people in the lab use the same protocol since several years to extract the DNA and the sequencing providers claim to always use the Illumina standard protocols.

But obviously something must have changed.

It took Illumina some 18 months to resolve that problem for the broader public: since data I work on were done with the TruSeq kit, this problem has vanished.

However, if you based some conclusions or wrote a paper with Illumina data which might be affected by the GC bias (Q3 2009 to Q4 2010), I suggest you rethink all the conclusion drawn. This should be especially the case for transcriptomics experiments where a difference in expression of 2x to 3x starts to get highly significant!

11.3 Ion Torrent

As of January 2014, I would say Ion Torrent reads behave very much like late data from the 454 technology (FLX / Titanium chemistry): reads are on average are > 300bp and the homopolymer problem is much less pronounced than 2 years ago. The following figure shows what you can get out of 100bp reads if you're lucky:

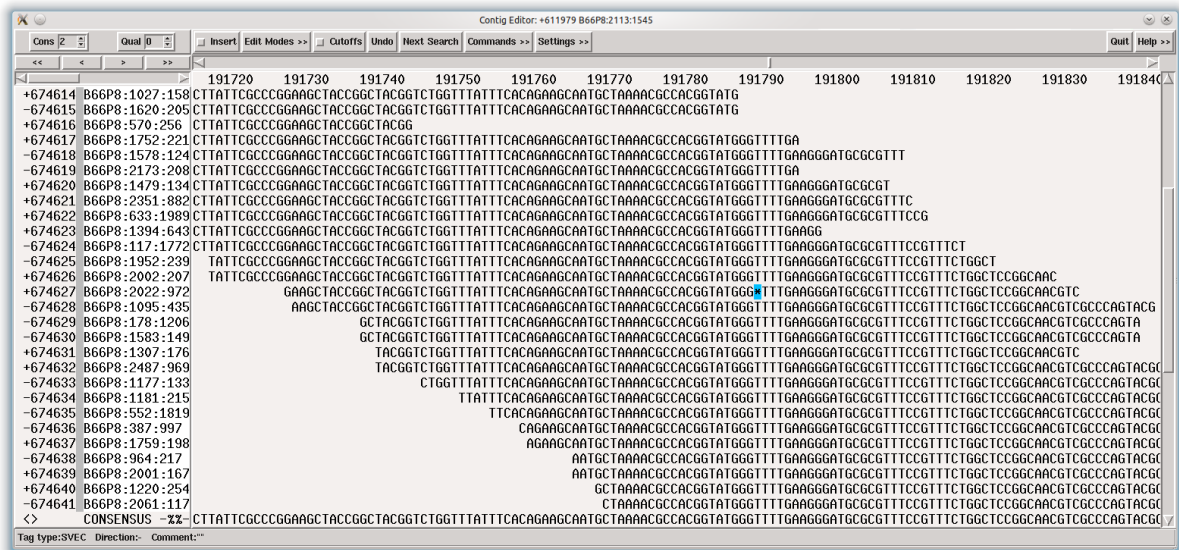


Figure 11.8: Example for good IonTorrent data (100bp reads). Note that only a single sequencing error - shown by blue background - can be seen. Except this, all homopolymers of size 3 and 4 in the area shown are good.

The "if you're lucky" part in the preceding sentence is not there by accident: having so many clean reads is more of an exception rather a rule. On the other hand, most sequencing errors in current IonTorrent data are unproblematic ... if it were not for indels, which is going to be explained on the next sections.

11.3.1 Homopolymer insertions / deletions

The main source of error in your data will be insertions / deletions (indels) especially in homopolymer regions (but not only there, see also next section). Starting with a base run of 4 to 6 bases, there is a distinct tendency to have an increased occurrence of indel errors.

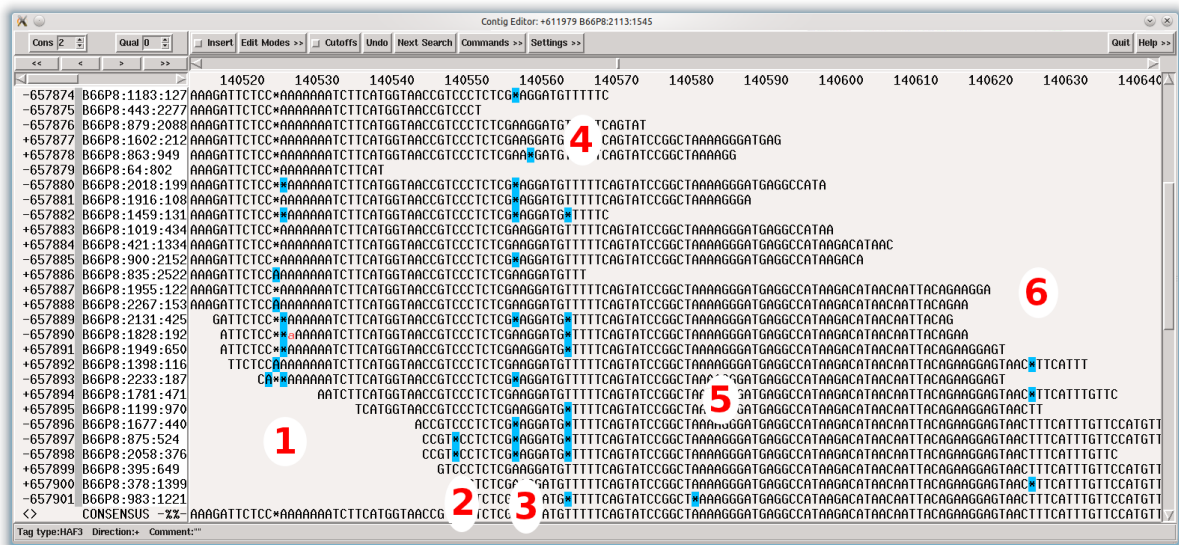


Figure 11.9: Example for problematic IonTorrent data (100bp reads).

The above figure contains a couple of particularly nasty indel problems. While areas 2 (C-homopolymer length 3), 5 (A-homopolymer length 4) and 6 (T-homopolymer length 3) are not a big problem as most of the reads got the length right, the areas 1, 3 and 4 are nasty.

Area 1 is an A-homopolymer of length 7 and while many reads get that length right (enough to tell MIRA what the true length is), it also contains reads with a length of 6 and and others with a length of 8.

Area 2 is a "A-homopolymer" of length 2 where approximately half of the reads get the length right, the other half not. See also the following section.

Area 4 is a T-homopolymer of length 5 which also has approximately half the reads with a wrong length of 4.

11.3.2 Sequencing direction dependent insertions / deletions

In the previous section, the screen shot showing indels had an indel at a homopolymer of 2, which is something quite curious. Upon closer investigation, one might notice a pattern in the gap/nogap distribution: it is almost identical to the orientation of build direction of reads!

I looked for other examples of this behaviour and found quite a number of them, the following figure shows a very clear case of that error behaviour:

```

Cons 2  Qual 0  Insert Edit Modes >> Cutoffs Undo Next Search Commands >> Settings >> Quit Help >>
187210 187220 187230 187240 187250 187260 187270 187280
+673190 B66P8:392:884 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCA
+673191 B66P8:1989:246 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCA
-673192 B66P8:344:2149 TGAATGCCCACTCATGTTGCGCCCT
-673193 B66P8:142:877 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGT
-673194 B66P8:2344:147 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673195 B66P8:1300:214 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGC
-673196 B66P8:1487:243 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673197 B66P8:1920:104 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673198 B66P8:1832:901 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673199 B66P8:535:1520 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCG
+673200 B66P8:2018:193 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673201 B66P8:233:1244 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATA
+673202 B66P8:1542:453 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673203 B66P8:1852:297 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673204 B66P8:1299:970 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673205 B66P8:717:1575 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673206 B66P8:1801:188 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673207 B66P8:1727:142 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673208 B66P8:1450:637 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATA
-673209 B66P8:1791:208 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673210 B66P8:156:477 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673211 B66P8:1331:209 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
-673212 B66P8:1950:190 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673213 B66P8:14:764 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGT
+673214 B66P8:757:2238 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673215 B66P8:1413:154 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
+673216 B66P8:2014:251 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATAT
+673217 B66P8:1503:168 TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
<> CONSENSUS -%- TGAATGCCCACTCATGTTGCGCCCTGTGCTGATTGCC*TCGCGGTGGGATTTGCAGTCGGTATTGCGCAGTATATGCCA
B66P8:1989:2469(#673191) Clone:unknown Vector:unknown Type:forward universal:unknown primer Tmpl:d

```

Figure 11.10: Example for a sequencing direction dependent indel. Note how all but one of the reads in '+' direction miss a base while all reads built in '-' direction have the correct number of bases.

This is quite astonishing: the problem occurs at a site without real homopolymer (calling a 2-bases run a 'homopolymer' starts stretching the definition a bit) and there are no major problematic homopolymer sites near. In fact, this was more or less the case for all sites I had a look at.

Neither did the cases which were investigated show common base patterns, so unlike the Solexa GGCxG motif it does not look like that error of IonTorrent is bound to a particular motif.

While I cannot prove the following statement, I somehow suspect that there must be some kind of secondary structure forming which leads to that kind of sequencing error. If anyone has a good explanation I'd be happy to hear it: feel free to contact me at bach@chevreux.org.

11.3.3 Coverage variance

The coverage variance with the old ~100bp reads was a bit on the bad side for low coverage projects (10x to 15x): it varied wildly, sometimes dropping to nearly zero, sometimes reaching approximately double the coverage.

This has now improved and I have not seen pronounced coverage variance in the data sets I have worked on.

11.3.4 GC bias

The GC bias seems to be small to non-existent, at least I could not immediately make a correlation between GC content and coverage.

11.3.5 Other sources of error

You will want to keep an eye on the clipping of the data in the SFF files from IonTorrent: while it is generally good enough, some data sets of IonTorrent show that - for some error patterns - the clipping is too lax and strange artefacts appear. MIRA will take care of these - or at least of those it knows - but you should be aware of this potential problem.

11.3.6 Where to find further information

IonTorrent being pretty new, getting as much information on that technology is quite important. So here are a couple of links I found to be helpful:

- There is, of course, the TorrentDev site (http://lifetech-it.hosted.jivesoftware.com/community/torrent_dev) at Life Technologies which will be helpful to get a couple of questions answered.
Just be aware that some of the documents over there are sometimes painting an - how should I say it diplomatically? - overly optimistic view on the performance of the technology. On the other hand, so do documents released by the main competitors like 454/Roche, Illumina, PacBio etc. ... so no harm done there.
- I found Nick Loman's blog [Pathogens: Genes and Genomes](#) to be my currently most valuable source of information on IonTorrent. While the group he works for won a sequencer from IonTorrent, he makes that fact very clear and still unsparingly dissects the data he gets from that machine.
His posts got me going in getting MIRA grok IonTorrent.
- The blog of Lex Nederbragt [In between lines of code](#) is playing in the same league: very down to earth and he knows a bluff when he sees it ... and is not afraid to call it (be it from IonTorrent, PacBio or 454).
The analysis he did on a couple of Ion data sets have saved me quite some time.
- Last, but not least, the board with [IonTorrent-related-stuff](#) over at [SeqAnswers](#), the first and foremost one-stop-shop ... erm ... discussion board for everything related to sequencing nowadays.

11.4 Pacific BioSciences

As of January 2014, PacBio should be seen as *the* technology to go to for de-novo sequencing of bacteria and lower eukaryotes. Period. Complement it with a bit of Illumina to get rid of the last remaining errors and you'll have - for a couple of thousand Euros - the best genome sequences money can buy.

11.4.1 Highlights

11.4.1.1 Sequence lengths

Just one word: huge. At least compared to other currently existing technologies. It is not unusual to get average - usable - read lengths of more than 3 to 4 kb, some chemistries doubling that number (at the expense of accuracy). The largest - usable - reads I have seen were > 25kb, though one needs to keep in mind that these are quite rare and one does not see many of them in a project.

11.4.1.2 GC bias

I have seen none in my projects so far, neither have I in public data. But these were certainly not as many projects as Sanger, 454, Illumina and Ion, so take this with a grain of salt.

11.4.1.3 Accuracy of corrected reads

Once the raw PacBio data has been corrected (HGAP pipeline), the resulting reads have a pretty good accuracy. There still are occasional homopolymer errors remaining at non-random locations, but they are a minor problem.

11.4.1.4 Assemblies of corrected reads

The assemblies coming out of the HGAP pipeline are already astoundingly good. Of course you get long contigs, but also the number of miscalled consensus bases is not too bad: 1 error per 20 kb. Once the program **Quiver** went through the assembly to do its magic in polishing, the quality improves further to into the range of 1 error per 50kb to 1 error per 250kb.

In my hands, I get even better assemblies with MIRA (longer contigs which span repeats unresolved by HGAP). When combining this with some low coverage Illumina data (say, 50x) to do cheap polishing, the error rates I get are lower than 1 error in 4 megabases.

Note Take the above with a grain of salt as at the time of this writing, I analysed in-depth only on a couple of bacteria. For ploidal organisms I have just played a bit around with public data without really doing an in depth analysis there.

11.4.2 Lowlights

11.4.2.1 Naming confusion

With PacBio, there are quite a number of read types being thrown around and which do confuse people: *polymerase reads*, *quality clipped reads*, *subreads*, *corrected reads* and maybe some more I currently forgot. Here's the total unofficial guide on how to keep those things apart:

- **polymerase reads** are the rawest and most unedited stuff you may come into contact. You can see it as "data fresh from the machine" and the number of megabases there is usually the one sequencing providers sell to you.

The sequencing technology PacBio employs uses special hairpin adaptors they have named SMRTBell, and these adaptors will be present in the polymerase reads together with the fragments of your DNA.

In terms of regular expression look-alike, the data in polymerase reads has the following form:

```
(Adaptor + (forward fragment sequence + (Adaptor + (fragment sequence in reverse ←
complement)))) *
```

E.g., some of your *polymerase reads* will contain just the adaptor and (part of) a fragment sequence: Adap+FwdSeq. Others might contain: Adap+FwdSeq+Adap+RevSeq. And still others might contain: multiple copies of Adap+FwdSeq+Adap+RevSeq.

- **quality clipped reads** are simply *polymerase reads* where some sort of first quality clipping has been done.
-

- **subreads** are *quality clipped reads* where the adaptors have been removed and the read split into forward fragment sequences and reverse fragment sequences. Hence, one quality clipped polymerase read can yield several subreads.
- **corrected (sub)reads** are subreads where through the magic of lots of computational power and a very high coverage of subreads, the errors have been almost completely removed from the subreads.

This is usually done only on a part of the subreads as it takes already long enough (several hundred hours CPU for a simple bacterium).

11.4.2.2 Forward / reverse chimeric sequences

The splitting of polymerase reads into subreads (see above) needs the SMRTBell adaptor to be recognised by motif searching programs. Unfortunately, it looks like as if some "low percentage" of reads have a self-looped end instead of an adaptor. Which in turn means that the subread splitting will not split those reads and you end up with a chimeric sequence.

11.4.2.3 Accuracy of uncorrected subreads

You need to be brave now: the accuracy of the the unclipped polymerase reads is usually only about 50%. That is: on average every second base is wrong. And I have seen a project where this accuracy was only 14% (6 out of 7 bases are wrong).

After clipping, the average accuracy of the polymerase reads should be anywhere between 80% and 85% (this depends a little bit on the chemistry used), which translates to: every 5th to every 7th base is wrong. The vast majority of errors being insertions or deletions, not base substitutions.

80% to 85% accuracy with indels as primary error is unfortunately something assemblers cannot use very well. Read: not at all if you want good assemblies (at least I know no program which does that). Therefore, one needs to apply some sort of correction ... which needs quite a deal of CPU, see below.

11.4.2.4 Immense need for CPU power

The above mentioned accuracies of 80% to 85% are too low for any existing assembler I know to be correctly assembled. Therefore, people came up with the idea of doing error correction on subreads to improve their quality.

There are two major approaches: 1) correcting PacBio subreads with other technologies with shorter reads and 2) correcting long PacBio subreads with shorter PacBio subreads. Both approaches have been shown to work, though there seems to be a preference nowadays to use the second option as the "shorter" PacBio reads provide the benefit of being still longer than read from other technologies and hence provide a better repeat resolution.

Anyway, the amount of CPU power needed for any method above is something to keep for: bacteria with 3 to 5 megabases at a 100x polymerase read coverage can take several hundred hours of CPU for the correction step.

11.4.2.5 Increased quality requirements for clean DNA sample prep

This is a problem which cannot be really attributed to PacBio: one absolutely needs to check whether the protocols used "since ever" for DNA extraction yield results which are clean and long enough for PacBio. Often they are not.

The reason for this being a problem is simple: PacBio can sequence really long fragments, but if your DNA extraction protocol smashed the DNA into small pieces, then no sequencing technology in this universe will be able to give you long reads for small fragments.

Chapter 12

Some advice when going into a sequencing project

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Reliable information lets you say ‘I don’t know’ with real confidence.”

—Solomon Short

12.1 Talk to your sequencing provider(s) before sequencing

Well, duh! But it’s interesting what kind of mails I sometimes get. Like in:

“We’ve sequenced a one gigabase, diploid eukaryote with Solexa 36bp paired-end with 200bp insert size at 25x coverage. Could you please tell us how to assemble this data set de-novo to get a finished genome?”

A situation like the above should have never happened. Good sequencing providers are interested in keeping customers long term and will therefore try to find out what exactly your needs are. These folks generally know their stuff (they’re making a living out of it) and most of the time propose you a strategy that fulfills your needs for a near minimum amount of money.

Listen to them.

If you think they try to rip you off or are overselling their competences (which most providers I know won’t even think of trying, but there are some), ask a quote from a couple of other providers. You’ll see pretty quickly if there are some things not being right.

Note As a matter of fact, a rule which has saved me time and again for finding sequencing providers is not to go for the cheapest provider, especially if their price is far below quotes from other providers. They’re cutting corners somewhere others don’t cut for a reason.

12.2 Choosing a sequencing provider

Note This is a slightly reworked version of a post I made on the MIRA talk mailing list. The question *“Could you please recommend me a sequencing provider?”* arrives every now and then in my private inbox, often enough for me decide to make a collage of the responses I gave in the past and post it to MIRA talk.

This response got, errrr, a little longer, but allow me to note that I will not give you names. The reasons are manifold:

- once upon a time I worked for a sequencing company
- the company I am currently employed with is not in the sequencing provider business, but the company uses more than one sequencing provider on a regular base and I get to see quite some data
- due to my development on MIRA in my free time, I'm getting insight into a number of highs and lows of sequencing technologies at different sequencing providers which I would not get if I were to expose them publicly ... I do not want to jeopardise these relationships.

That being said, there are a number of general considerations which could help you. Excuse me in case the detours I am going to make are obvious to you, but I'm writing this also for future references. Also, please bear with me if I look at "sequencing" a bit differently than you might be accustomed to from academia, but I have worked for quite some time now in industry ... and there cost-effectiveness respectively "probability of success" of a project as whole is paramount to everything else. I'll come back to that further down.

There's one -- and only one -- question which you, as sequencing customer, need to be able to answer ... if necessary in every excruciating detail, but you must know the answer. The question is:

12.2.1 WHAT DO YOU WANT?!

Detour - Sequencing -

For me, every "sequencing project", be it genomic or transcriptomic, really consists of four major phases:

1. **data generation:** This can be broadly seen as everything to get the DNA/RNA ready to be sent off to sequencing (usually something the client does), the library prep at the sequencing provider and finally the sequencing itself (including base calling). An area of thousand pitfalls where each step (and the communication) is crucial and even one slight inadvertence can make the difference between a "simple" project and a "hard" project. E.g.: taking DNA from growing cells (especially bacteria in exponential growing phase) might not be a good idea ... it makes assembly more difficult. Some DNA extraction methods generate more junk than good fragments etc.pp

The reason I am emphasizing this is simple: nowadays, the "sequencing" itself is not the most expensive part of a sequencing project, the next two steps are (most of the time anyway).

2. **assembly & finishing:** Still a hard problem. Even a "simple" bacterium can present weeks of effort to get right if its riddled with phages, prophages, transposon elements, genetically engineered repeats etc.pp. And starting with eukaryotes the real fun starts: ploidy, retrotransposons etc. make for an unbelievable genome plasticity and almost always have their own surprises. I've seen "simple" *Saccharomyces cerevisiae* - where biologist swore to high heaven they were "close to the publicly sequenced strains" - being *very* different from what they were expected to be, both on the DNA level and the genome organisation level.

Getting eukaryotes right "down to the last base" might cost quite some money, especially when looping back to step 1 (data generation) to tackle difficult areas.

3. **annotation:** Something many people forget: give the sequence a meaning. Here too, things can get quite costly if done "right", i.e., with hand curation. Especially on organism which are not part of the more commonly sequenced species or are generally more complex.

Annotation of a de-novo transcriptome assembly is also not for the faint of heart, especially if done on short, unpaired read assemblies.

4. **using the sequencing data:** ... for whatever it was generated for.

The above makes it clear that, depending on what you are really interested in within your project and what you expect to be able to do with the sequencing data, one can cut corners and reduce cost here and there (but not everywhere). And therefore, the above question "What do you want?" is one which - after the initial chit-chat of "hi, hello, nice to meet you, a pleasure to be here, etc." - every good representative of respectable sequencing providers I have met so far will ask as very first question. Usually in the form of "what do you want to sequence and what will you want to use the data for (and what not)?"

12.2.2 WHAT DO YOU NEED?!

... difference between "want" and "need" ...

Every other question - like where to sequence, which sequencing technology to use, how to process the sequencing data afterwards - is incidental and subordinated to your answer(s) to the question of "what do you want?!" But often sequencing customers get their priorities wrong by putting forward another question:

12.2.3 WHAT WILL IT COST ME?

And its inevitable companion question "Can you make it cheaper?"

Detour - Putting things into perspective -

Come to think of it, people sometimes have very interesting ideas regarding costs. Interesting as in "outright silly." It may be because they do not really know what they want or feel unsure on a terrain unbeknownst to them, and often instead focus their energy on single aspects of a wider project because they feel more at home there. And suddenly the focus lies on haggling and bartering for some prices because, after all, this is something everyone knows how to do, right?

As I hinted earlier, the pure sequencing costs are nowadays probably not the biggest factor in any sequencing project: 454, Illumina, IonTorrent and other technology providers have seen to that. E.g., in 2003/2004 it still cost somewhere between 150 - 200 k€ to get an 8x Sanger coverage of a moderately sized bacterium (4 to 5 mb). Nowadays, for the same organism, you get coverages in the hundreds (PacBio) or even thousands (going with Illumina) for a few hundred Euro.

Cost for assembly, finishing and annotation have not followed the same decrease. Yes, advances in algorithms have made things easier in some parts, but not really on the same scale. Furthermore, the "short read" technologies have more than made up for algorithmical complexity when compared to the old Sanger reads. Maybe that "(ultra)long read" technologies will alleviate the problem, but I would not hold my breath for them to really work well.

One thing however has almost not changed at all: your costs of actually doing followup experiments and data interpretation! Remember that sequencing in itself is most of the time not the ultimate goal, you actually want to gain something out of it. Be it abstract knowledge for a paper or concrete hints for producing some compounds or whatever, chances are that you will actually devote a substantial amount of your resources (time, manpower, mental health) into followup activities (lab experiments, genetic engineering, writing papers) to turn the abstract act of sequencing into something tangible, be it papers, fame, new products, money, or whatever you want to achieve.

And this is the place where it pays to stop and think: "what do I want? what are my strengths and where are my weaknesses? where are my priorities?" The English have a nice saying: "Being penny-wise and pound-foolish is not wise." I may add: Especially not if you are basing man months / years of lab work and your career on the outcome of something like sequencing. Maybe I'm spoiled because I have left academia for quite some time now, but in sequencing I always prefer to throw a bit more money at the sequencing process itself to minimise risks of the later stages.

12.2.4 WHERE TO SEQUENCE?

There's one last detour I'd like to make, and that is the question of "where to sequence?"

Detour - Public or private, old-timers or young-timers ? -

Choosing a sequencing provider is highly dependent on your answer to "what do you want?" In case you want to keep the sequencing data (or the very act of sequencing) secret (even only for some time) will probably lead you to commercial sequencing companies. There you more or less have complete control on the data. Paranoid people might perhaps argue that you can have that only with own sequencing equipment and personnel, but I have the feeling that only a minority is able to cough-up the necessary money for purchasing sequencing equipment for a small one-time project.

Instead of companies you could however also look whether one of the existing sequencing centers in the world might be a good cooperation candidate. Especially if you are doing this project within the scope of your university. Note however that there might be a number of gotchas lurking there, beside the obvious "the data is not really secret anymore": sometimes the raw sequencing data needs to be publicly released, maybe earlier than you would like; or the sequencing center imposes that each and every paper you publish with that data as basis has them as (co-)first author.

A related problem is "whom do I trust to deliver good work?" Intuition says that institutes with a long sequencing history have amassed quite some knowledge in this field, making them experts in all three aspects (data generation, assembly & finishing, annotation) of a sequencing project ... and intuition probably isn't wrong there. The same thing is probably true for sequencing companies which have existed for more than just a couple of years, though from what I have seen so far is that - due to size - sequencing companies sometimes really focus on the data generation and rely on partner companies for "assembly" and "annotation". This is not to say that younger companies are bad. Incidentally, it is my belief that in this field, people are still more important than technology ... and every once in a while good people split off a well known institute (or company) to try their luck in an own company. Always look for references there.

The following statement is a personal opinion (and you can call me biased for that): Personally, I am however quite wary of sequencing done at locations where a sequencer exists because someone got a grant to buy one (because it was chic & en-vogue to get a shiny new toy) but where the instrument then slowly starts to collect dust after the initial flurry ... and because people often do not calculate chemistry costs which arise in case they'd really thought of using the machine 24/7. I want to know that technicians actually work with those things every day, that they know the ins and outs of the work, the protocols, the chemistry, the moods of the machine (even an instrument can have a bad day). I honestly do not believe that one can build up enough expertise when operating these things "every once in a while".

12.2.5 Summary of all the above

All of the above means that depending on what I need the data for, I have the freedom choose among different providers. In case I just need masses of raw data and potential savings are substantial, I might go with the cheapest whom I know to generate good data. If I want good service and second round of data in case I am not 110% satisfied with the first round (somehow people have stopped questioning me there), this is usually not the cheapest provider ... but the additional costs are not really high. If I wanted my data really really quick, I'd search for a provider with Ion Torrent, or MiSeq (I am actually looking for one with a MiSeq, so if anyone knows a good one, preferably in Europe -> mail me). Though I already did transcriptomics on eukaryotes, in case I needed larger eukaryotes assembled de-novo & also annotated, I would probably look for the help of a larger sequencing center as this starts to get dangerously near the fringe of my field of expertise.

In closing this part, here are a couple of guidelines which have not failed me so far for choosing sequencing providers:

- Building a good relationship helps. In case your institute / university already has good (or OK) experience with a provider, ask there first.
- It is a lot easier to build a good relationship with someone who speaks your language ... or a good(!) English.
- I will not haggle for a couple of hundred Euros in a single project, I'll certainly reconsider this when savings are in the tens of thousands.
- Managing expectations: some sequencing projects are high risk from the start, for lots of possible reasons (underfunded, bad starting material, unclear organism). This is *sometimes* (!) OK as long as everyone involved knows and acknowledges this. However, you should always have a clear target ("what am I looking for?") and preferably know in advance how to treat the data to get there.

- Errors occur, stay friendly at first. In case the expectations were clear (see above), the material and organism are not at fault but the data quality somehow is bad, it is not too difficult to have the sequencing provider acknowledge this and get additional sequencing for no added cost.

Regarding the technologies you can use ... it really depends on what you want to do :-). And note that I base my answers on technologies available today without bigger problems: PacBio, Illumina, with IonTorrent as Joker for quick projects. Since 2018, 454 can be considered to be dead as the Newbler assembler is not available anymore and I am not even sure sequencing kits are manufactured. Oxford Nanopore has become a game changer with reads of several hundred Kb and even over a megabase, but still requires quite some tinkering both in protocols and on the bioinformatics side.

12.3 Specific advice

Here's how I see things as of now (early 2019), which might not necessarily be how others see them.

12.3.1 Technologies

12.3.1.1 Sanger

Use for: checking assemblies; closing gaps by PCR; checking for a couple of genes with known sequence (i.e., where you can design oligos for).

Do not use for: anything else. In particular, if you find yourself designing oligos for a 96 well plate destined for Sanger sequencing of a single bacterial DNA sample, you (probably) are doing something wrong.

12.3.1.2 Pacific Biosciences

Use for: de-novo of bacteria and lower eukaryotes (or higher eukaryotes if you have the money). PacBio should be seen as *the* technology to use when getting the best good quality sequence with least number of contigs is important to you. Also, resequencing of variants of known organisms with lots of genomic reorganisation flexibility due to high numbers of transposons (where short reads will not help in getting the chromosome assembled/mapped correctly).

Do not use for: resequencing of "dull" organisms (where the only differences will be simple SNPs or simple insertion/deletions or simple contig reorganisations at non-repetitive places). Illumina will do a much better and cost effective job there.

Note

As of Fall 2018: aim for at least 100x coverage of raw data, better 130x to 150x as pre-processing (quality clip, removal of adapters and other sequencing artefacts) will take its toll and reduce the data by up to 1/3. After that, the error correction/self-correction of raw reads into corrected reads will again reduce the data considerably.

It's really a numbers game: the more data you have, the more likely you will also get many of those really long reads in the 5 to 30 Kb range which are extremely useful to get over those nasty repeats.

Note MIRA should not be used to assemble PacBio. Use it to polish the results of Canu or Falcon/Quiver/Arrow with Illumina data.



Warning For non-haploid organisms, you might need more coverage to get enough data at ploidy sites to get the reads correctly out of error correction. Aim for at least 80x to 100x per ploidy.



Warning Preparation of your DNA sample is not trivial as many methods will break your DNA into "small" chunks which are good enough for Sanger, 454, Illumina or Ion Torrents, but not for PacBio.

12.3.1.3 Oxford Nanopore Technologies (ONT)

I currently have no hands-on experience with ONT, but from what I see it gets you much longer reads than PacBio. At the moment I would go for it for de-novo of difficult eukaryotes or a quick and dirty first look bacteria with lots of repeats.

12.3.1.4 Illumina

Use for: general resequencing jobs (finding SNPs, indel locations of any size, copy number variations etc.); gene expression analysis; cheap test sequencing of unknown organisms to assess complexity; de-novo sequencing if you are OK with getting hundreds / thousands of contigs (depending on organism, some bacteria get only a few dozen).



Warning Careful with high GC organisms, starting with 60% to 65% GC Illumina reads contain more errors: SNP detection may be less reliable if extreme care is not taken to perform good read clipping. Especially the dreaded GGCxG motif often leads to problems in Illumina reads.



Warning For de-novo assemblies, do *NOT* (never ever at all and under no circumstances) use the Nextera kit, take TruSeq. The non-random fragmentation behaviour of Nextera leads to all sorts of problems for assemblers (not only MIRA) which try to use kmer frequencies as a criterion for repetitiveness of a given sequence.

12.3.1.5 Ion Torrent

Use for: like Illumina. With three notable exceptions: 1) SNP detection is not as good as with Illumina (more false positives and false negatives) 2) de-novo assemblies will contain more single-base indels and 3) Ion having problems with homopolymers, that technology is not as well suited as complimentary hybrid technology for PacBio as is Illumina (except for high-GC perhaps).

Ion has a speed advantage on Illumina: if you have your own machine, getting from your sample to data takes less time than with Illumina.

Also, it looks like as if Ion has less problems with GC content or sequence motifs than Illumina.

12.3.1.6 Roche 454

That technology is dead. Move on, nothing to see here anymore.

12.3.2 Sequencing de-novo

- On a cheap gene fishing expedition? Probably Illumina HiSeq, at least 100bp, 150 to 250bp or 300bp if your provider supports it well. Paired-end definitely a plus. As alternative: Ion Torrent for small organism (maybe up to 100Mb) and when you need results quickly without caring for possible frameshifts.
- Want some larger contigs? PacBio. Add in cheap Illumina 100bp paired-end (150 to 300bp if provider supports it) to get rid of those last frameshifts which may remain.
- Even longer reads to resolve difficult areas? Some tinkering with Oxford Nanopore + polishing with Illumina should do wonders.

12.3.3 Re-sequencing / mapping

There is a reason why Illumina currently dominates the market as it does: a cheap Illumina run (preferably paired-end) will answer most of your questions in 95% of the cases. Things will get difficult for organisms with high numbers of repeats and/or frequent genome re-arrangements. Then using longer read technologies and/or Illumina mate-pair may be required.

12.4 A word or two on coverage ...

12.4.1 Low coverage isn't worth it

There's one thing to be said about coverage and de-novo assembly: especially for bacteria, getting more than 'decent' coverage is *cheap* with any current day technology. Every assembler I know will be happy to assemble de-novo genomes with coverages of 25x, 30x, 40x ... and the number of contigs will still drop dramatically between a 15x Ion Torrent and a 30x Ion Torrent project.

In any case, do some calculations: if the coverage you expect to get reaches 50x (e.g. 200MB raw sequence for a 4MB genome), then you (respectively the assembler) can still throw away the worst 20% of the sequence (with lots of sequencing errors) and concentrate on the really, really good parts of the sequences to get you nice contigs.

Other example: the price for 1 gigabase Illumina paired-end of a single DNA prep is way, way below USD 1000, even with commercial providers. Then you just need to do the math: is it worth to invest 10, 20, 30 or more days of wet lab work, designing primers, doing PCR sequencing etc. and trying to close remaining gaps or hunt down sequencing errors when you went for a 'low' coverage or a non-hybrid sequencing strategy? Or do you invest a few bucks more to get some additional coverage and considerably reduce the uncertainties and gaps which remain?

Remember, you probably want to do research on your bug and not research on how to best assemble and close genomes. So even if you put (PhD) students on the job, it's costing you time and money if you wanted to save money earlier in the sequencing. Penny-wise and pound-foolish is almost never a good strategy :-)

I do agree that with eukaryotes, things start to get a bit more interesting from the financial point of view.

12.4.2 Catch-22: too high coverage

There is, however, a catch-22 situation with coverage: too much coverage isn't good either. Without going into details: sequencing errors sometimes interfere heavily when coverage exceeds ~60x to 80x for 454 & IonTorrent and approximately 150x to 200x for Solexa/Illumina.

In those cases, do yourself a favour: there's more than enough data for your project ... just cut it down to some reasonable amount: 40x to 50x for 454 & IonTorrent, 100x for Solexa/Illumina.

12.5 A word of caution regarding your DNA in hybrid sequencing projects

So, you have decided that sequencing your bug with PacBio and Illumina (or PacBio and Ion Torrent or whatever) may be a viable way to get the best bang for your buck. Then please follow this advice: prepare enough DNA *in one go* for the sequencing provider so that they can sequence it with all the technologies you chose without you having to prepare another batch ... or even grow another culture!

The reason for that is that as soon as you do that, the probability that there is a mutation somewhere that your first batch did not have is not negligible. And if there is a mutation, even if it is only one base, there is a >95% chance that MIRA will find it and thinks it is some repetitive sequence (like a duplicated gene with a mutation in it) and splits contigs at those places.

Now, there are times when you cannot completely be sure that different sequencing runs did not use slightly different batches (or even strains).

One example: the SFF files for SRA000156 and SRA001028 from the NCBI short trace archive should both contain E.coli K12 MG-16650 (two unpaired half plates and a paired-end plate). However, they contain DNA from different cultures. Furthermore, the DNA was prepared by different labs. The net effect is that the sequences in the paired-end library contain a few distinct mutations from the sequences in the two unpaired half-plates. Furthermore, the paired-end sequences contain sequences from phages that are not present in the unpaired sequences.

In those cases, provide strain information to the reads so that MIRA can discern possible repeats from possible SNPs.

12.6 Advice for bacteria

12.6.1 Do not sample DNA from bacteria in exponential growth phase!

The reason is simple: some bacteria grow so fast that they start replicating themselves even before having finished the first replication cycle. This leads to more DNA around the origin of replication being present in cells, which in turn fools assemblers and mappers into believing that those areas are either repeats or that there are copy number changes.

Sample. In. Stationary. Phase!

For de-novo assemblies, MIRA will warn you if it detects data which points at exponential phase. In mapping assemblies, look at the coverage profile of your genome: if you see a smile shape (or V shape), you have a problem.

12.6.2 Beware of (high copy number) plasmids!

This is a source of interesting problems and furthermore gets people wondering why MIRA sometimes creates more contigs than other assemblers when it usually creates less.

Here's the short story: there are data sets which include one or several high-copy plasmid(s). Here's a particularly ugly example: SRA001028 from the NCBI short read archive which contains a plate of paired-end reads for Ecoli K12 MG1655-G (<ftp://ftp.ncbi.nih.gov/pub/TraceDB/ShortRead/SRA001028/>).

The genome is sequenced at ~10x coverage, but during the assembly, three intermediate contigs with ~2kb attain a silly maximum coverage of ~1800x each. This means that there were ~540 copies of this plasmid (or these plasmids) in the sequencing.

When using the uniform read distribution algorithm - which is switched on by default when using "--job=" and the quality level of 'accurate' - MIRA will find out about the average coverage of the genome to be at ~10x. Subsequently this leads MIRA to dutifully create ~500 additional contigs (plus a number of contig debris) with various incarnations of that plasmid at an average of ~10x, because it thought that these were repetitive sites within the genome that needed to be disentangled.

Things get even more interesting when some of the plasmid / phage copies are slightly different from each other. These too will be split apart and when looking through the results later on and trying to join the copies back into one contig, one will see that this should not be done because there are real differences.

DON'T PANIC!

The only effect this has on your assembly is that the number of contigs goes up. This in turn leads to a number of questions in my mailbox why MIRA is sometimes producing more contigs than Newbler (or other assemblers), but that is another story (hint: Newbler either collapses repeats or leaves them completely out of the picture by not assembling repetitive reads).

What you can do is the following:

1. either you assemble everything together and then join the plasmid contigs manually after assembly, e.g. in gap4 (drawback: on really high copy numbers, MIRA will work quite a bit longer ... and you will have a lot of fun joining the contigs afterwards)
2. or, after you found out about the plasmid(s) and know the sequence, you filter out reads in the input data which contain this sequence (you can use **mirabait** for this) and assemble the remaining reads.

Chapter 13

Bits and pieces

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Just when you think it’s finally settled, it isn’t.”

—Solomon Short



Warning

The documentation of MIRA 3.9.x has not completely caught up yet with the changes introduced by MIRA now using manifest files. Quite a number of recipes still show the old command-line style, e.g.:

```
mira --project=... --job=... ...
```

For those cases, please refer to chapter 3 (the reference) for how to write manifest files.

13.1 Using SSAHA2 / SMALT to screen for vector sequence

If your sequencing provider gave you data which was NOT pre-clipped for vector sequence, you can do this yourself in a pretty robust manner using SSAHA2 -- or the successor, SMALT -- from the Sanger Centre. You just need to know which sequencing vector the provider used and have its sequence in FASTA format (ask your provider).

Note This screening is a valid method for any type of Sanger sequencing vectors, 454 adaptors, Illumina adaptors and paired-end adaptors etc. However, you probably want to use it only for Sanger type data as MIRA already knows all standard 454, Ion Torrent and Illumina adaptors.

Note SSAHA2 and SMALT need their input data to be in FASTA format, so for these to run you will need them also in FASTA format. For MIRA however you can load your original data in whatever format it was present.

For SSAHA2 follow these steps (most are the same as in the example above):

```
$ ssaha2 -output ssaha2
-kmer 8 -skip 1 -seeds 1 -score 12 -cmatch 9 -ckmer 6
/path/where/the/vector/data/resides/vector.fasta
yourinputsequences.fasta > screendataforyoursequences.ssaha2
```

Then, in your manifest file, add the following line in the readgroup which contains the sequences you screened:

```
readgroup
...
data = yourinputsequences_inwhateverformat_thisexamplehasfastq.fastq
data = screendataforyoursequences.ssaha2
...
```

For SMALT, the only difference is that you use SMALT for generating the vector-screen file and ask SMALT to generate it in SSAHA2 format. As SMALT works in two steps (indexing and then mapping), you also need to perform it in two steps and then call MIRA. E.g.:

```
$ smalt index -k 7 -s 1 smaltidxdb /path/where/the/vector/data/resides/vector.fasta
$ smalt map -f ssaha -d -1 -m 7 smaltidxdb yourinputsequences.fasta > <
  screendataforyoursequences.smalt
```

Note Please note that, due to subtle differences between output of SSAHA2 (in ssaha2 format) and SMALT (in ssaha2 format), MIRA identifies the source of the screening (and the parsing method it needs) by the name of the screen file. Therefore, screens done with SSAHA2 need to have the postfix `.ssaha2` in the file name and screens done with SMALT need `*.smalt`.

Chapter 14

Frequently asked questions

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Every question defines its own answer. Except perhaps ‘Why a duck?’ ”

—Solomon Short



Warning

The documentation of MIRA 3.9.x has not completely caught up yet with the changes introduced by MIRA now using manifest files. Quite a number of recipes still show the old command-line style, e.g.:

```
mira --project=... --job=... ...
```

For those cases, please refer to chapter 3 (the reference) for how to write manifest files.

This list is a collection of frequently asked questions and answers regarding different aspects of the MIRA assembler.

Note This document needs to be overhauled.

14.1 Assembly quality

1. Test question 1

Test answer 1

2. Test question 2

Test answer 2

14.1.1 What is the effect of uniform read distribution (-AS:urd)?

```
I have a project which I once started quite normally via
"--job=denovo,genome,accurate,454"
and once with explicitly switching off the uniform read distribution
"--job=denovo,genome,accurate,454 -AS:urd=no"
I get less contigs in the second case and I wonder if that is not better.
Can you please explain?
```

Since 2.9.24x1, MIRA has a feature called "uniform read distribution" which is normally switched on. This feature reduces over-compression of repeats during the contig building phase and makes sure that, e.g., a rRNA stretch which is present 10 times in a bacterium will also be present approximately 10 times in your result files.

It works a bit like this: under the assumption that reads in a project are uniformly distributed across the genome, MIRA will enforce an average coverage and temporarily reject reads from a contig when this average coverage multiplied by a safety factor is reached at a given site.

It's generally a very useful tool disentangle repeats, but has some slight secondary effects: rejection of otherwise perfectly good reads. The assumption of read distribution uniformity is the big problem we have here: of course it's not really valid. You sometimes have less, and sometimes more than "the average" coverage. Furthermore, the new sequencing technologies - 454 perhaps but especially the microreads from Solexa & probably also SOLiD - show that you also have a skew towards the site of replication origin.

One example: let's assume the average coverage of your project is 8 and by chance at one place you have 17 (non-repetitive) reads, then the following happens:

`p` = parameter of `-AS:urdsip`

Pass 1 to `$p-1$`: MIRA happily assembles everything together and calculates a number of different things, amongst them an average coverage of `~8`. At the end of pass '`$p-1$`', it will announce this average coverage as first estimate to the assembly process.

Pass `p`: MIRA has still assembled everything together, but at the end of each pass the contig self-checking algorithms now include an "average coverage check". They'll invariably find the 17 reads stacked and decide (looking at the `-AS:urdct` parameter which I now assume to be 2) that 17 is larger than $2 * 8$ and that this very well may be a repeat. The reads get flagged as possible repeats.

Pass `$p+1$` to end: the "possibly repetitive" reads get a much tougher treatment in MIRA. Amongst other things, when building the contig, the contig now looks that "possibly repetitive" reads do not over-stack by an average coverage multiplied by a safety value (`-AS:urdcn`) which I'll assume in this example to be 1.5. So, at a certain point, say when read 14 or 15 of that possible repeat want to be aligned to the contig at this given place, the contig will just flatly refuse and tell the assembler to please find another place for them, be it in this contig that is built or any other that will follow. Of course, if the assembler cannot comply, the reads 14 to 17 will end up as contiglet (contig debris, if you want) or if it was only one read that got rejected like this, it will end up as singlet or in the debris file.

Tough luck. I do have ideas on how to re-integrate those reads at the end of an assembly, but I had deferred doing this as in every case I had looked up, adding those reads to the contigs wouldn't have changed anything ... there's already enough coverage. What I do in those cases is simply filter away the contiglets (defined as being of small size and having an average coverage below the average coverage of the project / 3 (or 2.5)) from a project.

14.1.2 There are too many contig debris when using uniform read distribution, how do I filter for "good" contigs?

```
When using uniform read distribution there are too many contig with low
coverage which I don't want to integrate by hand in the finishing process. How
do I filter for "good" contigs?
```

OK, let's get rid of the cruft. It's easy, really: you just need to look up one number, take two decisions and then launch a command.

The first decision you need to take is on the minimum average coverage the contigs you want to keep should have. Have a look at the file `*_info_assembly.txt` which is in the info directory after assembly. In the "Large contigs" section, there's a "Coverage assessment" subsection. It looks a bit like this:

```
...
Coverage assessment:
-----
Max coverage (total): 43
Max coverage
Sanger: 0
```



```
454:      43
Solexa:  0
Solid:   0
Avg. total coverage (size ≥ 5000): 22.30
Avg. coverage (contig size ≥ 5000)
Sanger:  0.00
454:     22.05
Solexa:  0.00
Solid:   0.00
...
```

This project was obviously a 454 only project, and the average coverage for it is ~22. This number was estimated by MIRA by taking only contigs of at least 5Kb into account, which for sure left out everything which could be categorised as debris. It's a pretty solid number.

Now, depending on how much time you want to invest performing some manual polishing, you should extract contigs which have at least the following fraction of the average coverage:

- 2/3 if a quick and "good enough" is what you want and you don't want to do some manual polishing. In this example, that would be around 14 or 15.
- 1/2 if you want to have a "quick look" and eventually perform some contig joins. In this example the number would be 11.
- 1/3 if you want quite accurate and for sure not loose any possible repeat. That would be 7 or 8 in this example.

The second decision you need to take is on the minimum length your contigs should have. This decision is a bit dependent on the sequencing technology you used (the read length). The following are some rules of thumb:

- Sanger: 1000 to 2000
- 454 GS20: 500
- 454 FLX: 1000
- 454 Titanium: 1500

Let's assume we decide for an average coverage of 11 and a minimum length of 1000 bases. Now you can filter your project with `miraconvert`

```
miraconvert -x 1000 -y 11 sourcefile.caf filtered.caf
```

14.1.3 When finishing, which places should I have a look at?

I would like to find those places where MIRA wasn't sure and give it a quick shot. Where do I need to search?

Search for the following tags in `gap4` or any other finishing program for finding places of importance (in this order).

- IUPc
 - UNSc
 - SRMc
 - WRMc
 - STMU (only hybrid assemblies)
 - STMS (only hybrid assemblies)
-

14.2 454 data

1. *What are little boys made of?*
Snips and snails and puppy dog tails.
2. *What are little girls made of?*
Sugar and spice and everything nice.

14.2.1 What do I need SFFs for?

```
I need the .sff files for MIRA to load ...
```

Nope, you don't, but it's a common misconception. MIRA does not load SFF files, it loads FASTA, FASTA qualities, FASTQ, XML, CAF, EXP and PHD. The reason why one should start from the SFF is: those files can be used to create a XML file in TRACEINFO format. This XML contains the absolutely vital information regarding clipping information of the 454 adaptors (the sequencing vector of 454, if you want).

For 454 projects, MIRA will then load the FASTA, FASTA quality and the corresponding XML. Or from CAF, if you have your data in CAF format.

14.2.2 What's `sff_extract` and where do I get it?

```
How do I extract the sequence, quality and other values from SFFs?
```

Use the `sff_extract` script from Jose Blanca at the University of Valencia to extract everything you need from the SFF files (sequence, qualities and ancillary information). The home of `sff_extract` is: http://bioinf.comav.upv.es/sff_extract/index.html but I am thankful to Jose for giving permission to distribute the script in the MIRA 3rd party package (separate download).

14.2.3 Do I need the `sfftools` from the Roche software package?

No, not anymore. Use the `sff_extract` script to extract your reads. Though the Roche `sfftools` package contains a few additional utilities which could be useful.

14.2.4 Combining SFFs

```
I am trying to use MIRA to assemble reads obtained with the 454 technology but I can't combine my sff files since I have two files obtained with GS20 system and 2 others obtained with the GS-FLX system. Since they use different cycles (42 and 100) I can't use the sfffile to combine both.
```

You do not need to combine SFFs before translating them into something MIRA (or other software tools) understands. Use `sff_extract` which extracts data from the SFF files and combines this into input files.

14.2.5 Adaptors and paired-end linker sequences

```
I have no idea about the adaptor and the linker sequences, could you send me the sequences please?
```

Here are the sequences as filed by 454 in their patent application:

```
>AdaptorA
CTGAGACAGGGAGGGAACAGATGGGACACGCAGGGATGAGATGG
>AdaptorB
CTGAGACACGCAACAGGGGATAGGCAAGGCACACAGGGGATAGG
```

However, looking through some earlier project data I had, I also retrieved the following (by simply making a consensus of sequences that did not match the target genome anymore):

```
>5prime454adaptor???
GCCTCCCTCGCGCCATCAGATCGTAGGCACCTGAAA
>3prime454adaptor???
GCCTTGCCAGCCCGCTCAGATTGATGGTGCCTACAG
```

Go figure, I have absolutely no idea where these come from as they also do not comply to the "tcag" ending the adaptors should have.

I currently know one linker sequence (454/Roche also calls it *spacer* for GS20 and FLX paired-end sequencing):

```
>flxlinker
GTTGGAACCGAAAGGGTTGAATTCAAACCCTTTCGGTTCCAAC
```

For Titanium data using standard Roche protocol, you need to screen for two linker sequences:

```
>titlinker1
TCGTATAACTTCGTATAATGTATGCTATACGAAGTTATTACG
>titlinker2
CGTAATAACTTCGTATAGCATAACATTATACGAAGTTATACGA
```



Warning Some sequencing labs modify the adaptor sequences for tagging and similar things. Ask your sequencing provider for the exact adaptor and/or linker sequences.

14.2.6 What do I get in paired-end sequencing?

```
Another question I have is does the read pair sequences have further
adaptors/vectors in the forward and reverse strands?
```

Like for normal 454 reads - the normal A and B adaptors can be present in paired-end reads. That theory this could look like this:

A-Adaptor - DNA1 - Linker - DNA2 - B-Adaptor.

It's possible that one of the two DNA fragments is *very* short or is missing completely, then one has something like this:

A-Adaptor - DNA1 - Linker - B-Adaptor

or

A-Adaptor - Linker - DNA2 - B-Adaptor

And then there are all intermediate possibilities with the read not having one of the two adaptors (or both). Though it appears that the majority of reads will contain the following:

DNA1 - Linker - DNA2

There is one caveat: according to current paired-end protocols, the sequences will **NOT** have the direction

```
----> Linker <----
```

as one might expect when being used to Sanger Sequencing, but rather in this direction

```
<---- Linker ---->
```

14.2.7 Sequencing protocol

Is there a way I can find out which protocol was used?

Yes. The best thing to do is obviously to ask your sequencing provider.

If this is - for whatever reason - not possible, this list might help.

Are the sequences ~100-110 bases long? It's GS20.

Are the sequences ~220-250 bases long? It's FLX.

Are the sequences ~350-450 bases long? It's Titanium.

Do the sequences contain a linker (GTTGGAACCGAAAGGGTTTGAATTCAAACCCCTTTCGGTTCCAAC)? It's a paired end protocol.

If the sequences left and right of the linker are ~29bp, it's the old short paired end (SPET, also it's most probably from a GS20).

If longer, it's long paired-end (LPET, from a FLX).

14.2.8 Filtering sequences by length and re-assembly

I have two datasets of ~500K sequences each and the sequencing company already did an assembly (using MIRA) on the basecalled and fully processed reads (using of course the accompanying *qual file). Do you suggest that I should redo the assembly after filtering out sequences being shorter than a certain length (e.g. those that are <200bp)? In other words, am I taking into account low quality sequences if I do the assembly the way the sequencing company did it (fully processed reads + quality files)?

I don't think that filtering out "shorter" reads will bring much positive improvement. If the sequencing company used the standard Roche/454 pipeline, the cut-offs for quality are already quite good, remaining sequences should be, even when being < 200bp, not of bad quality, simply a bit shorter.

Worse, you might even introduce a bias when filtering out short sequences: chemistry and library construction being what they are (rather imprecise and sometimes problematic), some parts of DNA/RNA yield smaller sequences per se ... and filtering those out might not be the best move.

You might consider doing an assembly if the company used a rather old version of MIRA (<3.0.0 for sure, perhaps also <3.0.5).

14.3 Solexa / Illumina data

14.3.1 Can I see deletions?

Suppose you ran the genome of a strain that had one or more large deletions. Would it be clear from the data that a deletion had occurred?

In the question above, I assume you'd compare your strain *X* to a strain *Ref* and that *X* had deletions compared to *Ref*. Furthermore, I base my answer on data sets I have seen, which presently were 36 and 76 mers, paired and unpaired.

Yes, this would be clear. And it's a piece of cake with MIRA.

Short deletions (1 to 10 bases): they'll be tagged SROc or WRMc. General rule: deletions of up to 10 to 12% of the length of your read should be found and tagged without problem by MIRA, above that it may or may not, depending a bit on coverage, indel distribution and luck.

Long deletions (longer than read length): they'll be tagged with MCVc tag by MIRA ins the consensus. Additionally, when looking at the FASTA files when running the CAF result through miraconvert: long stretches of sequences without coverage (the @ sign in the FASTAs) of *X* show missing genomic DNA.

14.3.2 Can I see insertions?

Suppose you ran the genome of a strain X that had a plasmid missing from the reference sequence. Alternatively, suppose you ran a strain that had picked up a prophage or mobile element lacking in the reference. Would that situation be clear from the data?

Short insertions (1 to 10 bases): they'll be tagged SROc or WRMc. General rule: deletions of up to 10 to 12% of the length of your read should be found and tagged without problem by MIRA, above that it may or may not, depending a bit on coverage, indel distribution and luck.

Long insertions: it's a bit more work than for deletions. But if you ran a de-novo assembly on all reads not mapped against your reference sequence, chances are good you'd get good chunks of the additional DNA put together

Once the Solexa paired-end protocol is completely rolled out and used on a regular base, you would even be able to place the additional element into the genome (approximately).

14.3.3 De-novo assembly with Solexa data

Any chance you could assemble de-novo the sequence of a from just the Solexa data?



Warning Highly opinionated answer ahead, your mileage may vary.

Allow me to make a clear statement on this: maybe.

But the result would probably be nothing I would call a good assembly. If you used anything below 76mers, I'm highly sceptical towards the idea of de-novo assembly with Solexa (or ABI SOLiD) reads that are in the 30 to 50bp range. They're really too short for that, even paired end won't help you much (especially if you have library sizes of just 200 or 500bp). Yes, there are papers describing different draft assemblers (SHARCGS, EDENA, Velvet, Euler and others), but at the moment the results are less than thrilling to me.

If a sequencing provider came to me with N50 numbers for an *assembled genome* in the 5-8 Kb range, I'd laugh him in the face. Or weep. I wouldn't dare to call this even 'draft'. I'd just call it junk.

On the other hand, this could be enough for some purposes like, e.g., getting a quick overview on the genetic baggage of a bug. Just don't expect a finished genome.

14.4 Hybrid assemblies

14.4.1 What are hybrid assemblies?

Hybrid assemblies are assemblies where one used more than one sequencing technology. E.g.: Sanger and 454, or 454 and Solexa, or Sanger and Solexa etc.pp

14.4.2 What differences are there in hybrid assembly strategies?

Basically, one can choose two routes: multi-step or all-in-one-go.

Multi-steps means: to assemble reads from one sequencing technology (ideally the one from the shorter tech like, e.g., Solexa), fragment the resulting contigs into pseudo-reads of the longer tech and assemble these with the real reads from the longer tech

(like, e.g., 454). The advantage of this approach is that it will be probably quite faster than the all-in-one-go approach. The disadvantage is that you loose a lot of information when using only consensus sequence of the shorter read technology for the final assembly.

All-in-one-go means: use all reads in one single assembly. The advantage of this is that the resulting alignment will be made of true reads with a maximum of information contained to allow a really good finishing. The disadvantage is that the assembly will take longer and will need more RAM.

14.5 Masking

14.5.1 Should I mask?

In EST projects, do you think that the highly repetitive option will get rid of the repetitive sequences without going to the step of repeat masking?

For eukaryotes, yes. Please also consult the [-KS:mnr] option.

Remember: you still **MUST** have sequencing vectors and adaptors clipped! In EST sequences the poly-A tails should be also clipped (or let mira do it).

For prokaryotes, I'm a big fan of having a first look at unmasked data. Just try to start MIRA without masking the data. After something like 30 minutes, the all-vs-all comparison algorithm should be through with a first comparison round. grep the log for the term "megahub" ... if it doesn't appear, you probably don't need to mask repeats

14.5.2 How can I apply custom masking?

I want to mask away some sequences in my input. How do I do that?

First, if you want to have Sanger sequencing vectors (or 454 adaptor sequences) "masked", please note that you should rather use ancillary data files (CAF, XML or EXP) and use the sequencing or quality clip options there.

Second, please make sure you have read and understood the documentation for all -CL parameters in the main manual, but especially -CL:mbc:mbcgs:mbcmfg:mbcmeg as you might want to switch it on or off or set different values depending on your pipeline and on your sequencing technology.

You can without problem mix your normal repeat masking pipeline with the FASTA or EXP input for MIRA, as long as you **mask** and not **clip** the sequence.

An example:

```
>E09238ARF0
tcag GTGTCAGTGTGACTGTAAAAAAAAAGTACGTATGGACTGCATGTGCATGTCATGGTACGTGTCA
GTCAGTACAAAAAAAAAAAAAAAAAGTACGT tgctgacgcacatgatcgtagc
```

(spaces inserted just as visual helper in the example sequence, they would not occur in the real stuff)

The XML will contain the following clippings: left clip = 4 (clipping away the "tcag" which are the last four bases of the adaptor used by Roche) right clip= ~90 (clipping away the "tgctgac..." lower case sequence on the right side of the sequence above).

Now, on the FASTA file that was generated with reads_sff.py or with the Roche sff* tools, you can let run, e.g., a repeat masker. The result could look like this:

```
>E09238ARF0
tcag XXXXXXXXXX TTGACTGTAAAAAAAAAGTACGTATGGACTGCATGTGCATGTCATGGTACGTGTCA
GTCAGTACAAAAAAAAAAAAAAAAAGTACGT tgctgacgcacatgatcgtagc
```

The part with the Xs was masked away by your repeat masker. Now, when MIRA loads the FASTA, it will first apply the clippings from the XML file (they're still the same). Then, if the option to clip away masked areas of a read (-CL:mbc, which is normally on for EST projects), it will search for the stretches of X and internally also put clips to the sequence. In the example above, only the following sequence would remain as "working sequence" (the clipped parts would still be present, but not used for any computation).

```
>E09238ARF0
.....TTGACTGTAAAAAAAAAAGTACGTATGGACTGCATGTGCATGTCATGGTACGTGTCA
GTCAGTACAAAAAAAAAAAAAAAAAAGTACGT.....
```

Here you can also see the reason why your filters should **mask** and not clip the sequence. If you change the length of the sequence, the clips in the XML would not be correct anymore, wrong clippings would be made, wrong sequence reconstructed, chaos ensues and the world would ultimately end. Or something.

IMPORTANT! It might be that you do not want MIRA to merge the masked part of your sequence with a left or right clip, but that you want to keep it something like DNA - masked part - DNA. In this case, consult the manual for the `-CL:mbc` switch, either switch it off or set adequate options for the boundaries and gap sizes.

Now, if you look at the sequence above, you will see two possible poly-A tails ... at least the real poly-A tail should be masked else you will get megahubs with all the other reads having the poly-A tail.

You have two possibilities: you mask yourself with an own program or you let MIRA do the job (`-CL:cpat`, which should normally be on for EST projects but I forgot to set the correct switch in the versions prior to 2.9.26x3, so you need to set it manually for 454 EST projects there).

IMPORTANT! Never ever at all use two poly-A tail masker (an own and the one from MIRA): you would risk to mask too much. Example: assume the above read you masked with a poly-A masker. The result would very probably look like this:

```
>E09238ARF0
tcag XXXXXXXXXX TTGACTGTAAAAAAAAAAGTACGTATGGACTGCATGTGCATGTCATGGTACGTGTCA
GTCAGTAC XXXXXXXXXXXXXXXXXXXXXXXX GTACGT tgetgacgcacatgatcgtagc
```

And MIRA would internally make the following out of it after loading:

```
>E09238ARF0
.....TTGACTGTAAAAAAAAAAGTACGTATGGACTGCATGTGCATGTCATGGTACGTGTCA
GTCAGTAC.....
```

and then apply the internal poly-A tail masker:

```
>E09238ARF0
.....TTGACTGT.....
.....
```

You'd be left with ... well, a fragment of your sequence.

14.6 Miscellaneous

14.6.1 What are megahubs?

```
I looked in the log file and that term "megahub" you told me about appears
pretty much everywhere. First of all, what does it mean?
```

Megahub is the internal term for MIRA that the read is massively repetitive with respect to the other reads of the projects, i.e., a read that is a megahub connects to an insane number of other reads.

This is a clear sign that something is wrong. Or that you have a quite repetitive eukaryote. But most of the time it's sequencing vectors (Sanger), A and B adaptors or paired-end linkers (454), unmasked poly-A signals (EST) or non-normalised EST libraries which contain high amounts of housekeeping genes (always the same or nearly the same).

Countermeasures to take are:

- set clips for the sequencing vectors (Sanger) or Adaptors (454) either in the XML or EXP files
- for ESTs, mask poly-A in your input data (or let MIRA do it with the `-CL:cpat` parameter)

- only after the above steps have been made, use the [-KS:mnr] switch to let mira automatically mask nasty repeats, adjust the threshold with [-SK:rt].
- if everything else fails, filter out or mask sequences yourself in the input data that come from housekeeping genes or nasty repeats.

14.6.2 Passes and loops

```
While processing some contigs with repeats i get
"Accepting probably misassembled contig because of too many iterations."
What is this?
```

That's quite normal in the first few passes of an assembly. During each pass (-AS:nop), contigs get built one by one. After a contig has been finished, it checks itself whether it can find misassemblies due to repeats (and marks these internally). If no misassembly, perfect, build next contig. But if yes, the contig requests immediate re-assembly of itself.

But this can happen only a limited number of times (governed by -AS:rbl). If there are still misassemblies, the contig is stored away anyway ... chances are good that in the next full pass of the assembler, enough knowledge has been gained to correctly place the reads.

So, you need to worry only if these messages still appear during the last pass. The positions that cause this are marked with "SRMc" tags in the assemblies (CAF, ACE in the result dir; and some files in the info dir).

14.6.3 Debris

```
What are the debris composed of?
```

- sequences too short (after trimming)
- megahubs
- sequences almost completely masked by the nasty repeat masker ([-KS:mnr])
- singlets, i.e., reads that after an assembly pass did not align into any contig (or were rejected from every contig).
- sequences that form a contig with less reads than defined by [-AS:mrpc]

14.6.4 Log and temporary files: more info on what happened during the assembly

```
I do not understand why ... happened. Is there a way to find out?
```

Yes. The tmp directory contains, beside temporary data, a number of log files with more or less readable information. While development versions of MIRA keep this directory after finishing, production versions normally delete this directory after an assembly. To keep the logs and temporary files also in production versions, use "-OUT:rtd=no".

As MIRA also tries to save as much disk space as possible, some logs and temporary files are rotated (which means that old logs and tmps get deleted). To switch off this behaviour, use "-OUT:rrot=no". Beware, the size of the tmp directory will increase, sometimes dramatically so.

14.6.4.1 Sequence clipping after load

How MIRA clipped the reads after loading them can be found in the file `mira_int_clippings.0.txt`. The entries look like this:

```
load: minleft. U13a01d05.t1      Left: 11      -> 30
```


Interpret this as: after loading, the read "U13a01d05.t1" had a left clipping of eleven. The "minleft" clipping option of MIRA did not like it and set it to 30.

```
load: bad seq. gnl|ti|1133527649      Shortened by 89 New right: 484
```

Interpret this as: after loading, the read "gnl|ti|1133527649" was checked with the "bad sequence search" clipping algorithm which determined that there apparently is something dubious, so it shortened the read by 89 bases, setting the new right clip to position 484.

14.7 Platforms and Compiling

14.7.1 Windows

```
Also, is MIRA be available on a windows platform?
```

As a matter of fact: it was and may be again. While I haven't done it myself, according to reports I got compiling MIRA 2.9.3* in a Cygwin environment was actually painless. But since then BOOST and multi-threading has been included and I am not sure whether it is still as easy.

I'd be thankful for reports :-)

Chapter 15

The MAF format

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“Design flaws travel in herds. ”

—Solomon Short



Warning

The documentation of MIRA 3.9.x has not completely caught up yet with the changes introduced by MIRA now using manifest files. Quite a number of recipes still show the old command-line style, e.g.:

```
mira --project=... --job=... ...
```

For those cases, please refer to chapter 3 (the reference) for how to write manifest files.

This documents describes purpose and format of the MAF format, version 1. Which has been superceeded by version 2 but is not described here (yet). But as v1 and v2 are very similar only the notion of readgroups is a big change, I'll let this description live until I have time to update this section.

15.1 Introduction: why an own assembly format?

I had been on the hunt for some time for a file format that allow MIRA to quickly save and load reads and full assemblies. There are currently a number of alignment format files on the market and MIRA can read and/or write most of them. Why not take one of these? It turned out that all (well, the ones I know: ACE, BAF, CAF, CALF, EXP, FRG) have some kind of no-go 'feature' (or problem or bug) that makes one life pretty difficult if one wants to write or parse that given file format.

What I needed for MIRA was a format that:

1. is easy to parse
2. is quick to parse
3. contains all needed information of an assembly that MIRA and many finishing programs use: reads (with sequence and qualities) and contigs, tags etc.pp

MAF is not a format with the smallest possible footprint though it fares quite well in comparison to ACE, CAF and EXP), but as it's meant as interchange format, it'll do. It can be easily indexed and does not need string lookups during parsing.

I took the liberty to combine many good ideas from EXP, BAF, CAF and FASTQ while defining the format and if anything is badly designed, it's all my fault.

15.2 The MAF format

This describes version 1 of the MAF format. If the need arises, enhancements like meta-data about total number of contigs and reads will be implemented in the next version.

15.2.1 Basics

MAF ...

1. ... has for each record a keyword at the beginning of the line, followed by exactly one blank (a space or a tab), then followed by the values for this record. At the moment keywords are two character keywords, but keywords with other lengths might appear in the future
2. ... is strictly line oriented. Each record is terminated by a newline, no record spans across lines.

All coordinates start at 1, i.e., there is no 0 value for coordinates.

15.2.2 Reads

15.2.2.1 Simple example

Here's an example for a simple read, just the read name and the sequence:

```
RD      U13a05e07.t1
RS      CTTGCATGCCTGCAGGTCGACTCTAGAAGGACCCCGATCA
ER
```

Reads start with RD and end with ER, the RD keyword is always followed by the name of the read, ER stands on its own. Reads also should contain a sequence (RS). Everything else is optional. In the following example, the read has additional quality values (RQ), template definitions (name in TN, minimum and maximum insert size in TF and TT), a pointer to the file with the raw data (SF), a left clip which covers sequencing vector or adaptor sequence (SL), a left clip covering low quality (QL), a right clip covering low quality (QR), a right clip covering sequencing vector or adaptor sequence (SR), alignment to original sequence (AO), a tag (RT) and the sequencing technology it was generated with (ST).

```
RD      U13a05e07.t1
RS      CTTGCATGCCTGCAGGTCGACTCTAGAAGGACCCCGATCA
RQ      , -+*, 1-+/, 36; :6≤3327<7A1/, ,) . ('..7=@E8:
TN      U13a05e07
DI      F
TF      1200
TT      1800
SF      U13a05e07.t1.scf
SL      4
QL      7
QR      30
SR      32
AO      1 40 1 40
RT      ALUS 10 15 Some comment to this read tag.
ST      Sanger
ER
```

15.2.2.2 List of records for reads

- *RD string: readname*
RD followed by the read name starts a read.
 - *LR integer: read length*
The length of the read can be given optionally in LR. This is meant to help the parser perform sanity checks and eventually pre-allocate memory for sequence and quality.
MIRA at the moment only writes LR lines for reads with more than 2000 bases.
 - *RS string: DNA sequence*
Sequence of a read is stored in RS.
 - *RQ string: qualities*
Qualities are stored in FASTQ format, i.e., each quality value + 33 is written as single as ASCII character.
 - *SV string: sequencing vector*
Name of the sequencing vector or adaptor used in this read.
 - *TN string: template name*
Template name. This defines the DNA template a sequence comes from. In it's simplest form, a DNA template is sequenced only once. In paired-end sequencing, a DNA template is sequenced once in forward and once in reverse direction (Sanger, 454, Solexa). In Sanger sequencing, several forward and/or reverse reads can be sequenced from a DNA template. In PacBio sequencing, a DNA template can be sequenced in several "strokes", leading to multiple reads on a DNA template.
 - *DI character: F or R*
Direction of the read with respect to the template. F for forward, R for reverse.
 - *TF integer: template size from*
Minimum estimated size of a sequencing template. In paired-end sequencing, this is the minimum distance of the read pair.
 - *TT integer: template size to*
Maximum estimated size of a sequencing template. In paired-end sequencing, this is the maximum distance of the read pair.
 - *SF string: sequencing file*
Name of the sequencing file which contains raw data for this read.
 - *SL integer: seqvec left*
Clip left due to sequencing vector. Assumed to be 1 if not present. Note that left clip values are excluding, e.g.: a value of '7' clips off the left 6 bases.
 - *QL integer: qual left*
Clip left due to low quality. Assumed to be 1 if not present. Note that left clip values are excluding, e.g.: a value off '7' clips of the left 6 bases.
 - *CL integer: clip left*
Clip left (any reason). Assumed to be 1 if not present. Note that left clip values are excluding, e.g.: a value of '7' clips off the left 6 bases.
 - *SR integer: seqvec right*
Clip right due to sequencing vector. Assumed to be the length of the sequence if not present. Note that right clip values are including, e.g., a value of '10' leaves the bases 1 to 9 and clips at and including base 10 and higher.
 - *QR integer: qual right*
Clip right due to low quality. Assumed to be the length of the sequence if not present. Note that right clip values are including, e.g., a value of '10' leaves the bases 1 to 9 and clips at and including base 10 and higher.
-

- *CR integer: clip right*
Clip right (any reason). Assumed to be the length of the sequence if not present. Note that right clip values are including, e.g., a value of '10' leaves the bases 1 to 9 and clips at and including base 10 and higher.
- *AO four integers: x1 y1 x2 y2*
AO stands for "Align to Original". The interval [x1 y1] in the read as stored in the MAF file aligns with [x2 y2] in the original, unedited read sequence. This allows to model insertions and deletions in the read and still be able to find the correct position in the original, base-called sequence data.
A read can have several AO lines which together define all the edits performed to this read.
Assumed to be "1 x 1 x" if not present, where 'x' is the length of the unclipped sequence.
- *RT string + 2 integers + optional string: type x1 y1 comment*
Read tags are given by naming the tag type, which positions in the read the tag spans in the interval [x1 y1] and afterwards optionally a comment. As MAF is strictly line oriented, newline characters in the comment are encoded as `\n`.
If $x1 > y1$, the tag is in reverse direction.
The tag type can be a free form string, though MIRA will recognise and work with tag types used by the Staden gap4 package (and of course the MIRA tags as described in the main documentation of MIRA).
- *ST string: sequencing technology*
The current technologies can be defined: Sanger, 454, Solexa, SOLiD.
- *SN string: strain name*
Strain name of the sample that was sequenced, this is a free form string.
- *MT string: machine type*
Machine type which generated the data, this is a free form string.
- *BC string: base caller*
Base calling program used to call bases
- *IB boolean (0 or 1): is backbone*
Whether the read is a backbone. Reads used as reference (backbones) in mapping assemblies get this attribute.
- *IC boolean (0 or 1)*
Whether the read is a coverage equivalent read (e.g. from mapping Solexa). This is internal to MIRA.
- *IR boolean (0 or 1)*
Whether the read is a rail. This also is internal to MIRA.
- ER
This ends a read and is mandatory.

15.2.2.3 Interpreting clipping values

Every left and right clipping pair (SL & SR, QL & QR, CL & CR) forms a clear range in the interval [left right] in the sequence of a read. E.g. a read with SL=4 and SR=10 has the bases 1,2,3 clipped away on the left side, the bases 4,5,6,7,8,9 as clear range and the bases 10 and following clipped away on the right side.

The left clip of a read is determined as $\max(\text{SL}, \text{QL}, \text{CL})$ (the rightmost left clip) whereas the right clip is $\min(\text{SR}, \text{QR}, \text{CR})$.

15.2.3 Contigs

Contigs are not much more than containers containing reads with some additional information. Contrary to CAF or ACE, MAF does not first store all reads in single containers and then define the contigs. In MAF, contigs are defined as outer container and within those, the reads are stored like normal reads.

15.2.3.1 Simple example 2

The above example for a read can be encased in a contig like this (with two consensus tags gratuitously added in):

```
CO      contigname_s1
NR      1
LC      24
CS      TGCCTGCAGGTCGACTCTAGAAGG
CQ      -+/,36;:6≤3327<7A1/,,).
CT      COMM 5 8 Some comment to this consensus tag.
CT      COMM 7 12 Another comment to this consensus tag.
\\
RD      U13a05e07.t1
RS      CTTGCATGCCTGCAGGTCGACTCTAGAAGGACCCCGATCA
RQ      , -+*,1-+/,36;:6≤3327<7A1/,,).('..7=@E8:
TN      U13a05e07
TF      1200
TT      1800
SF      U13a05e07.t1.scf
SL      4
SR      32
QL      7
QR      30
AO      1 40 1 40
RT      ALUS 10 15 Some comment to this read tag.
ST      Sanger
ER
AT      1 24 7 30
//
EC
```

Note that the read shown previously (and now encased in a contig) is absolutely unchanged. It has just been complemented with a bit of data which describes the contig as well as with a one liner which places the read into the contig.

15.2.3.2 List of records for contigs

- *CO string: contig name*
CO starts a contig, the contig name behind is mandatory but can be any string, including numbers.
- *NR integer: num reads in contig*
This is optional but highly recommended.
- *LC integer: contig length*
Note that this length defines the length of the 'clear range' of the consensus. It is 100% equal to the length of the CS (sequence) and CQ (quality) strings below.
- *CT string + 2 integers + optional string: identifier x1 y1 comment*
Consensus tags are defined like read tags but apply to the consensus. Here too, the interval [x1 y1] is including and if x1 > y1, the tag is in reverse direction.
- *CS string: consensus sequence*
Sequence of a consensus is stored in RS.
- *CQ string: qualities*
Consensus Qualities are stored in FASTQ format, i.e., each quality value + 33 is written as single as ASCII character.
- **
This marks the start of read data of this contig. After this, all reads are stored one after the other, just separated by an "AT" line (see below).

- AT *Four integers: x1 y1 x2 y2*

The AT (Assemble_To) line defines the placement of the read in the contig and follows immediately the closing "ER" of a read so that parsers do not need to perform time consuming string lookups. Every read in a contig has exactly one AT line.

The interval [x2 y2] of the read (i.e., the unclipped data, also called the 'clear range') aligns with the interval [x1 y1] of the contig. If $x1 > y1$ (the contig positions), then the reverse complement of the read is aligned to the contig. For the read positions, $x2$ is always $< y2$.

- //

This marks the end of read data

- EC

This ends a contig and is mandatory

Chapter 16

Log and temporary files used by MIRA

MIRA Version V5rc1 Bastien Chevreux 2018Bastien Chevreux

“The amount of entropy in the universe is constant - except when it increases. ”

—Solomon Short

Warning



The documentation of MIRA 3.9.x has not completely caught up yet with the changes introduced by MIRA now using manifest files. Quite a number of recipes still show the old command-line style, e.g.:

```
mira --project=... --job=... ...
```

For those cases, please refer to chapter 3 (the reference) for how to write manifest files.

16.1 Introduction

The tmp directory used by mira (usually <projectname>_d_tmp) may contain a number of files with information which could be interesting for other uses than the pure assembly. This guide gives a short overview.

Note This guide is probably the least complete and most out-of-date as it is updated only very infrequently. If in doubt, ask on the MIRA talk mailing list.



Warning Please note that the format of these files may change over time, although I try very hard to keep changes reduced to a minimum.

Note Remember that mira has two options that control whether log and temporary files get deleted: while [-OUT:rtd] removes the complete tmp directory after an assembly, [-OUT:rrot] removes only those log and temporary files which are not needed anymore for the continuation of the assembly. Setting both options to no will keep all log and temporary files.

16.2 The files

16.2.1 mira_error_reads_invalid

A simple list of those reads that were invalid (no sequence or similar problems).

16.2.2 mira_info_reads_tooshort

A simple list of those reads that were sorted out because the unclipped sequence was too short as defined by [-AS:mrl].

16.2.3 mira_int_alignnextends_preassembly1.0.txt

If read extension is used ([-DP:ure]), this file contains the read name and the number of bases by which the right clipping was extended.

16.2.4 mira_int_clippings.0.txt

If any of the [-CL:] options leads to the clipping of a read, this file will tell when, which clipping, which read and by how much (or to where) the clippings were set.

16.2.5 mira_int_posmatch_megahubs_pass.X.lst

Note: replace the *X* by the pass of mira. Should any read be categorised as megahub during the all-against-all search (SKIM3), this file will tell you which.

16.2.6 mira_int_posmatch_multicopystat_preassembly.0.txt

After the initial all-against-all search (SKIM3), this file tells you to how many other reads each read has overlaps. Furthermore, reads that have more overlaps than expected are tagged with ``mc'' (multicopy).

16.2.7 mira_int_posmatch_rawhashhits_pass.X.lst

Note: replace the *X* by the pass of mira. Similar to mira_int_posmatch_multicopystat_preassembly.0.txt, this counts the kmer hits of each read to other reads. This time however per pass.

16.2.8 mira_int_skimmarknastyrepeats_hist_pass.X.lst

Note: replace the *X* by the pass of mira. Only written if [-KS:mnr] is set to yes. This file contains a histogram of kmer occurrences encountered by SKIM3.

16.2.9 mira_int_skimmarknastyrepeats_nastyseq_pass.X.lst

Note: replace the *X* by the pass of mira. Only written if [-KS:mnr] is set to yes. One of the more interesting files if you want to know the repetitive sequences cause the assembly to be really difficult: for each masked part of a read, the masked sequences is shown here.

E.g.

```
U13a04h11.t1    TATATATATATATATATATATATATA
U13a05b01.t1    TATATATATATATATATATATATATA
U13a05c07.t1    AAAAAAAAAAAAAAA
U13a05e12.t1    CTCTCTCTCTCTCTCTCTCTCTCTC
```

Simple repeats like the ones shown above will certainly pop-up there, but a few other sequences (like e.g. rDNA/rRNA or SINEs, LINEs in eukaryotes) will also appear.

Nifty thing to try out if you want to have a more compressed overview: sort and unify by the second column.

```
sort -k 2 -u mira_int_skimmarknastyrepeats_nastyseq_pass.X.lst
```

16.2.10 mira_int_vectorclip_pass.X.txt

Note: replace the *X* by the pass of mira. Only written if [-CL:pvlc] is set to yes. Tells you where possible sequencing vector (or adaptor) leftovers were found and clipped (or not clipped).

16.2.11 miratmp.ads_pass.X.forward and miratmp.ads_pass.X.complement

Note: replace the *X* by the pass of mira. Which read aligns with Smith-Waterman against which other read, 'forward-forward' and 'forward-complement'.

16.2.12 miratmp.ads_pass.X.reject

Note: replace the *X* by the pass of mira. Which possible read overlaps failed the Smith-Waterman alignment check.

16.2.13 miratmp.noqualities

Which reads went completely without qualities into the assembly.

16.2.14 miratmp.usedids

Which reads effectively went into the assembly (after clipping etc.).

16.2.15 mira_readpoolinfo.lst