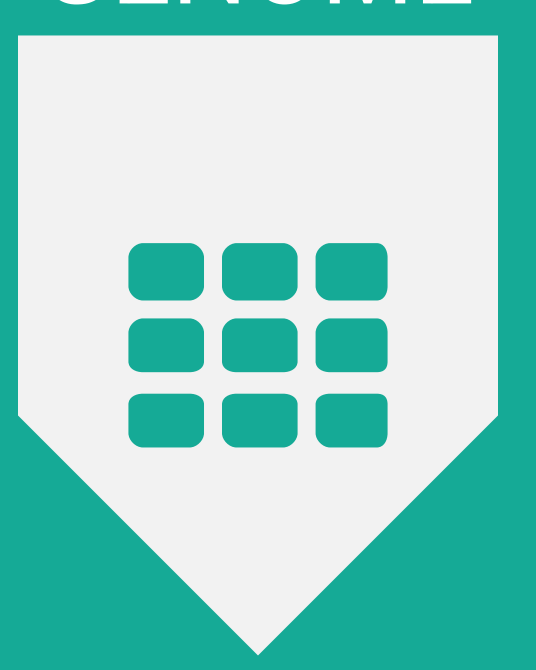
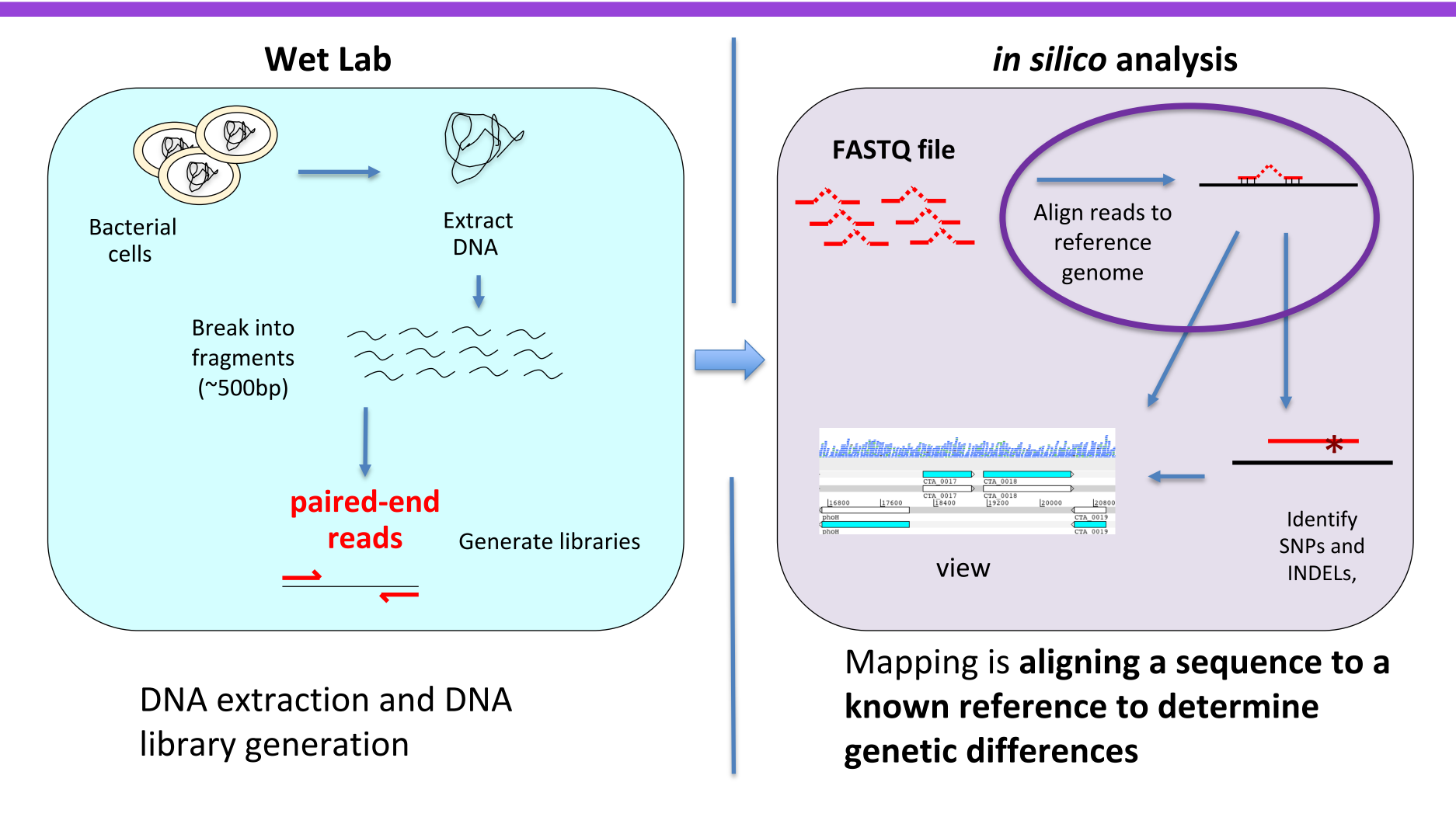
MAPPING READS TO A REFERENCE GENOME

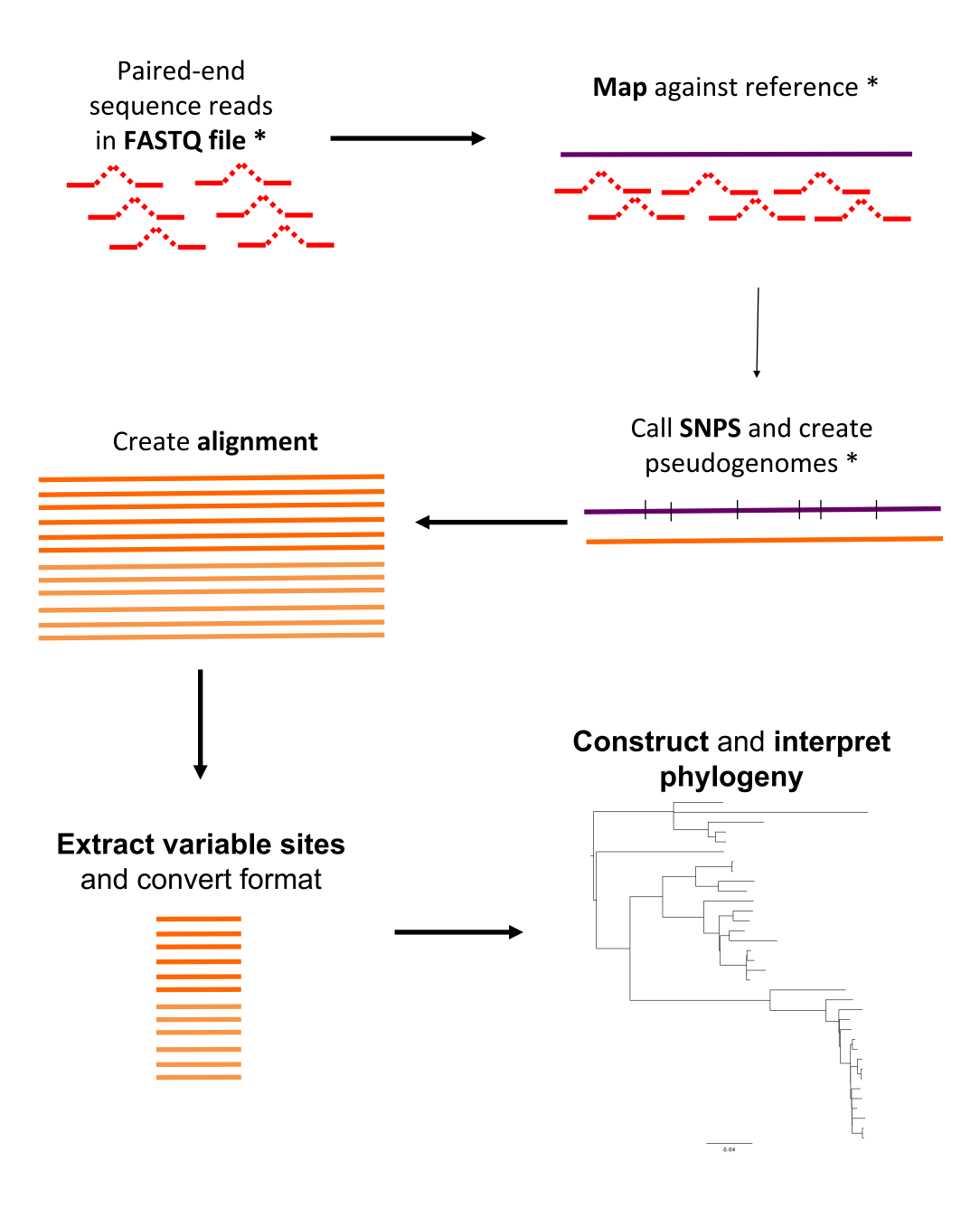


Workflow: generating sequencing reads and *in silico* analysis



Why do we map reads to a reference?

- Identify variation:
 - Single Nucleotide Polymorphisms (SNPs),
 - <u>insertions and deletions (indels)</u>
 - Copy Number Variants (CNVs) between variants of the same bacteria.
 - Presence / absence of genes (AMR)



There are lots of options for mapping!

Comparison of different mappers

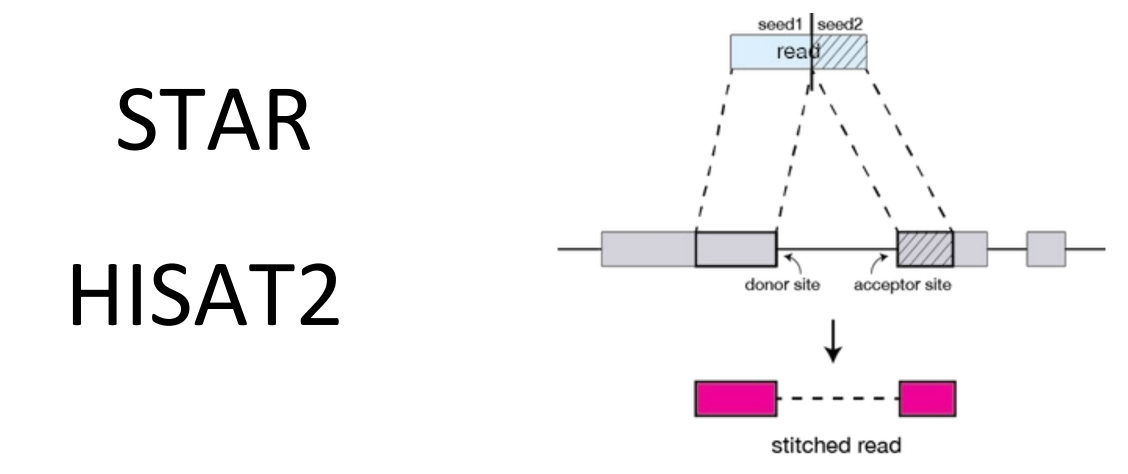
Manner Data A	vellebilit.	Verslen	O.S. N	umbor Citations	Sea Diet	Innest	Output	Min Di	May	DI Mie	matabaa Indola Cana Ali	un Donostod Alic	enment Bereilel OA DE	Culialna Indov
Mapper Data A BatMeth Bisulfite	vailability OS	Version 1.03		umber Citations 34	Seq.Plat.	Input (C)FAST(A/Q)	Output Native	Min. RL		100	matches Indels Gaps Aliq 5 N N	gn. Reported Alig B,U	G NYN	Splicing Index N Reference
Batmis DNA	os	3.0		23	I,So	FASTA/Q	SAM		33	100	10 0 N	A,U,S	NYY	N Reference
BFAST DNA	os	0.7.0		553	I,So,4, Hel	(C)FAST(A/Q)	SAM TSV			*	Y Y Y	B,R,U	G SM N Y	N Reference
Bismark Bisulfite	os	0.7.3		887	1,00,1,110	FASTA/Q	SAM		16 1	10K	Score Score N	_,,U	SM Y Y	N
BLASR DNA	os	1.4			Р	FASTA/Q hdf5	SAM TSV		50 1000		0.2 0.2 Y	A,B,R	GL NYN	De novo Reference
Blat DNA	os	34		6252	N	FASTA	TSV BLAST		11 500		Score Score Y	В	L NNN	De novo Reference
Bowtie DNA	OS	0.12.7	Linux, Mac, Windows	11207	I,So,4,Sa,P	(C)FAST(A/Q)	SAM TSV		4	1K	Score Score N	A,B,R,S	GL SM Y Y	N Reference
Bowtie2 DNA	OS	2.0beta5		8586	I,4,Ion	FASTA/Q	SAM TSV		4 500	00K	Score Score Y	A,B,R,S	GL SM Y Y	N Reference
BRAT Bisulfite	os	1.2.3		60	I	FASTA/Q	TSV				Y 0 N		NNY	N Reference
BRAT-BW Bisulfite	os	2.0.1	Linux	53		FASTA/Q	TSV		32	*	Y 0 N		NNY	N Reference
BS-Seeker Bisulfite	OS		Linux,Mac	193	!	FASTA/Q	SAM		40	000	3 0 N	U	SM Y N	N
BS-Seeker2 Bisulfite	os	2.0.0		107	!	FASTA/Q qseq	SAM BAM			200	Score Score Y	B,U,S	GL SM N Y	N Reference
BSMAP Bisulfite BWA DNA	OS OS	2.73 0.6.2		347 13341	I,So,4,Sa,P	FASTA/Q SAM/BAM FASTA/Q	SAM BAM Native SAM			144 200	15 1 N Y 8 Y	B,R,U R,S	G SM N Y G SM Y Y	N Reference N Reference
BWA-PSSM DNA	OS	0.5.11	Linux	26	1,30,4,3a,F	FASTQ/Q PSSM	SAM BAM			200	30 10 N	B,R,U,S	G SM Y Y	N Reference
BWA-SW DNA	OS	0.6.2		3494	I,4,Sa,Hel,Ion,P	FASTA/Q	SAM			00K	0.1 0.1 Y	R,S	L SM Y N	N Both
BWT-SW DNA		20070916		133	1,4,0a,1101,1011,1 N	FASTA	TSV		7 100	1K	Score Score Y	A	N N N	N Reference
CLC Mapper DNA	Com	4			I,4,So,Sa,Ion,P,Hel	FASTA/Q	SAM BAM						N Y	
CloudBurst DNA	os	1.1	Linux, Mac, Windows	650	N	FASTA	TSV			1K	Y Y Y	A,B	G Cloud N N	N Reads
ContextMap RNA	OS	2.2	Linux, Unix, Mac	22	I,4,So,Sa,Ion,P,Hel	FASTA/Q	SAM		1 50	000	20 10 Y	A,B	G SM N Y	Lib or de novo Reference
ContextMap 2 RNA	OS		Windows, Linux, Unix, Mac	9	I,4,Sa,Ion,P,Hel	FASTA/Q Illumina	SAM BED		20 50	000	0.1 10 Y	В	L No N Y	Lib or de novo Reference
CRAC RNA	OS	2.0.0		41	I,4,Ion,P	(C)FAST(A/Q) RAW	SAM BAM		50	*	score score Y	A,B,U,S	G SM N Y	De novo Both
CUSHAW3 DNA	OS	v3.0.3		33	I,So,4,Ion,P	FASTA/Q	SAM			096	score score Y	A,B,R,U,S	GL SM Y Y	N Reference
DART RNA	OS	1.2.4		0	١	FASTA/Q	SAM		20	000	YYY	ΑU	G SM N Y	De novo Reads
drFAST DNA	os	1.0.0.0		23	So	CFAST(A/Q)	SAM DIVET			200	Score N N	A,B	G NNY	N Reference
DynMap DNA ELAND DNA	OS Com	0.0.20		2 25	IN I	FASTA FASTA	TSV		18 15	8K 150	5 0 N 2 Score N	B B,S	L NNN G NYY	N Reads
ELAND DNA ERNE DNA	OS	1	Linux, Unix, Mac Windows, Linux, Unix, Mac	14	;	FASTA/Q Illumina	SAM BAM Native			600	0.1 5 Y	A,R,U,S	G SM/DM N Y	De novo Reference
Exonerate DNA	OS	2.2		918	Ň	FASTA	TSV		20	*	Score Score Y	B,S	GL NNN	De novo Reference
GEM DNA	Bin	1.x		260	I, So	FASTA/Q	SAM Counts		20	*	YYY	A,S	G SM Y Y	Lib and de novo Reference
GenomeMapper DNA	OS	0.4.3		144	., .,	FASTA/Q	BED TSV		12	2K	10 10 Y	A,B,R	G SM N N	N Reference
GMAP DNA		2012-04-27		868	I,4,Sa,Hel,Ion,P	FASTA/Q	SAM GFF Native		8	*	Y Y Y	В	GL SM N N	De novo Reference
GNUMAP DNA	OS	3.0.2		80	1	FASTA/Q Illumina	SAM TSV		16	1K	Score Score Y	В	G SM/DM Y N	N Reference
GSNAP DNA			Linux,Unix,Mac,Windows	1156	I,4,Sa,Hel,Ion,P	FASTA/Q	SAM Native			250	Y Y Y	A,B,U,S	GL SM N Y	Lib and de novo Reference
HISAT RNA	os		Windows, Linux, Unix, Mac	480	Į.	FASTA/Q	SAM		50	*	0.1 0.1 N	A,B,R,U,S	G SM Y Y	Lib or de novo Reference
HISAT2 DNA	OS		Windows, Linux, Unix, Mac	40	.!	FASTA/Q	SAM		50	000	score score N	В	G SM Y Y	Lib or de novo Reference
Hobbes2 DNA	OS	2.1	Linux	13	N N a d Ca Hallan D	FASTA/Q	SAM			200	0.08 0.08 N	A,U.S	G NNY G NYY	No Reference
hpg-Aligner DNA JAGuaR RNA	OS OS	v2.1.0 2.1	Linux Linux, Unix		I,So,4,Sa,Hel,Ion,P	FASTQ FASTQ	SAM, BAM SAM BAM			000 300	0.3 0.3 Yes	A,B	G NYY G NYY	Lib and de novo Reference Lib Reference
MapReads DNA	OS	2.4.1	Linux,Mac,Windows	15 0	So	FASTA/Q	TSV			120	Score 0 N	8	NYN	N Reference
MapSplice RNA	os	1.15.2		610	I	FASTA/Q	SAM BED		10	120	3 Y	В	SM N Y	De novo
MAQ DNA	OS	0.7.1	Linux,Mac	2592	I,So	(C)FAST(A/Q)	TSV		8	63	Ϋ́ΥΝ̈́		NYY	N Reads
Masai DNA	os	0.4	*	1	I, lon	FASTA/Q	SAM		20 320	678	32 32 N	A, B, U	G NNY	N Both
MicroRazerS miRNA	os	0.1	Linux	40	N	FASTA	SAM TSV		10	*	Score 0 N	S	G NNN	N Reference
MIRA DNA	OS	3	Linux,Unix		I,4,Sa,Ion,P		SAM GFF Counts CAF		25 190	000	Score Score Y	B,R	L SM Y Y	N Both
MOM DNA	Bin	0.6		48	1,4	FASTA	TSV		. –		Y 0 N	Α	L SM N Y	N Either
MOSAIK DNA	OS	2.1	Linux,Unix,Mac,Windows		I,So,4,Sa,Hel,Ion,P	(C)FAST(A/Q)	BAM			000	YYY	A,B	G SM Y Y	N Reference
mrFAST DNA	os	2.5.0.1	Linux,Unix	602	!	FASTA/Q	SAM DIVET			000	Score 4 N	A,B	G NYY	N Reference
mrsFAST DNA mrsFAST-Ultra DNA	OS	2.4.0.4		229	!	FASTA/Q	SAM DIVET			100	Score N N	A	G NYY G SMYY	N Reference
mrsFAST-Ultra DNA Mummer 3 DNA	OS OS	3.3.1 3.23	Linux, Mac Linux,Mac	28 2446	I NI	FASTA/Q FASTA	SAM DIVET TSV		8 ; 10	500	Score N N	A,B,S A,B	G SM Y Y G N N N	N Reference N Reference
NextGenMap DNA	OS	0.4.6		82	I 4 Ion	(C)FAST(A/Q),SAM,BAM	SAM BAM			000	Score Score N	R,S	GL SM N Y	N Reference
Novoalign(CS) DNA	Bin	V2.08.03		0	I,So,4,Hel,Ion		SAM Native			250	Y Y Y	A,B,R,U	G SM/DM Y Y	Lib Reference
OSA RNA	Bin		Windows, Linux, Unix, Mac	54	I,4,lon	FASTA/Q	SAM BAM			000	* * Y	A,B,U	G SM Y Y	Lib and de novo Reference
PASS DNA	Bin	1.62		142	I,So,4	(C)FAST(A/Q)	SAM GFF3 BLAST		23	1K	YYY	A,B	G SM Y Y	De novo Reference
PASS-bis Bisulfite	OS	2.01	Linux	14	I,So,4,Sa	FASTA/Q	SAM GFF Counts			000	Score N N	A, B, U, S	G SM Y Y	N Reference
Passion RNA	OS	1.2.0		28	I,4,Sa,P	FASTA/Q	BED				Y Y Y	U	SM Y Y	De novo
PatMaN miRNA	OS	1.2.2		140	N	FASTA	TSV		1	*	Y Y N	Α	G NNN	N Reads
PerM DNA	OS	0.4.0		113	I,So	(C)FAST(A/Q)	SAM TSV		20	128	9 0 Y	A,U	G DM Y Y	N Reference
ProbeMatch DNA	os	000	Linux,Mac	4	I,4,Sa	FASTA	ELAND		36	50	3 Y N	A,B	N N N	N Reference
QPALMA RNA RazerS DNA	OS	0.9.2		169 165	I,4 I 4	Specific	TSV		11	*	1 1 1	АВИС		Lib and de novo
RazerS DNA RazerS3 DNA	OS OS	1.2 3.1	Linux,Mac,Windows Windows, Linux, Mac	165 81	1,4	FASTQ FASTA/Q	TSV ELAND SAM TSV GFF		11 11	*	Score Score Y 0.5 Y N	A,B,U,S A,B,U,S	G NNY G SMNY	N Reference N Reads
REAL DNA	OS	0.0.28		32	i	FASTA/Q FASTA/Q	TSV		4	*	Score N N	А,Б,О,З В,U	G SM Y N	N Reference
NEAE DIVA	00	0.0.20	Linux	10		TAOTA Q	104		AA 400	222	17	5,0	0 011 1 11	N Telefolio

Good general aligners

bwa bowtie2

Fast, sensitive and easy to use!

Splice-aware aligners for RNA-seq



Why do we map to a reference?

- Identify variation:
 - Single Nucleotide Polymorphisms (SNPs),
 - <u>insertions</u> and <u>deletions</u> (indels)
 - Copy Number Variants (CNVs) between variants of the same bacteria.
 - Presence / absence of genes (AMR)

Single Nucleotide Polymorphisms (SNPs)

Reference CCGTTAGAGTTACAATTCGA

Read 2 TTAGAGTAACAA

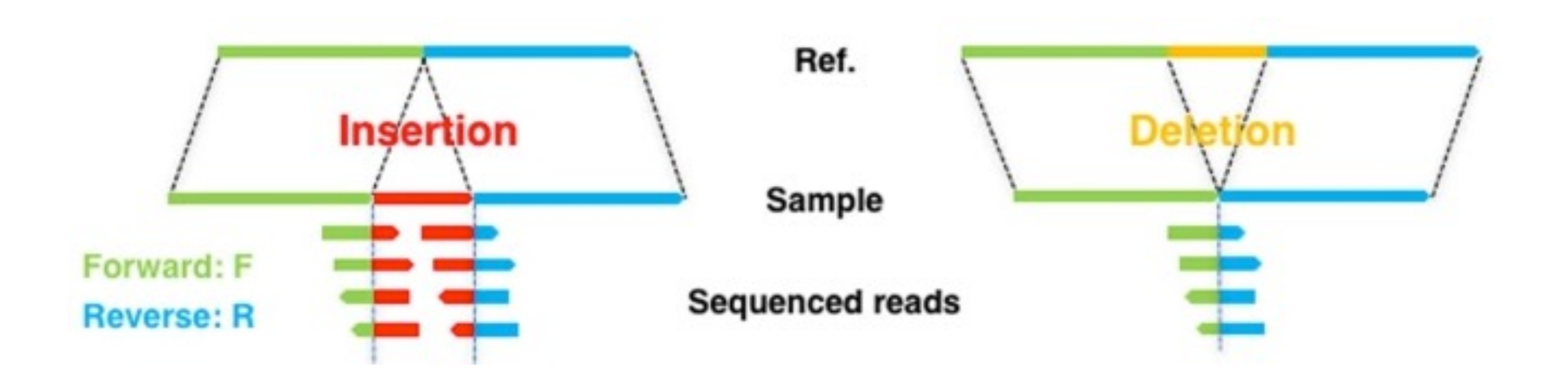
Read 3 CCGTTAGAGTTA

Read 4 TTACAATTCGA

Read 5 GAGTAACAA

Read 6 TTAGAGTAACAAT

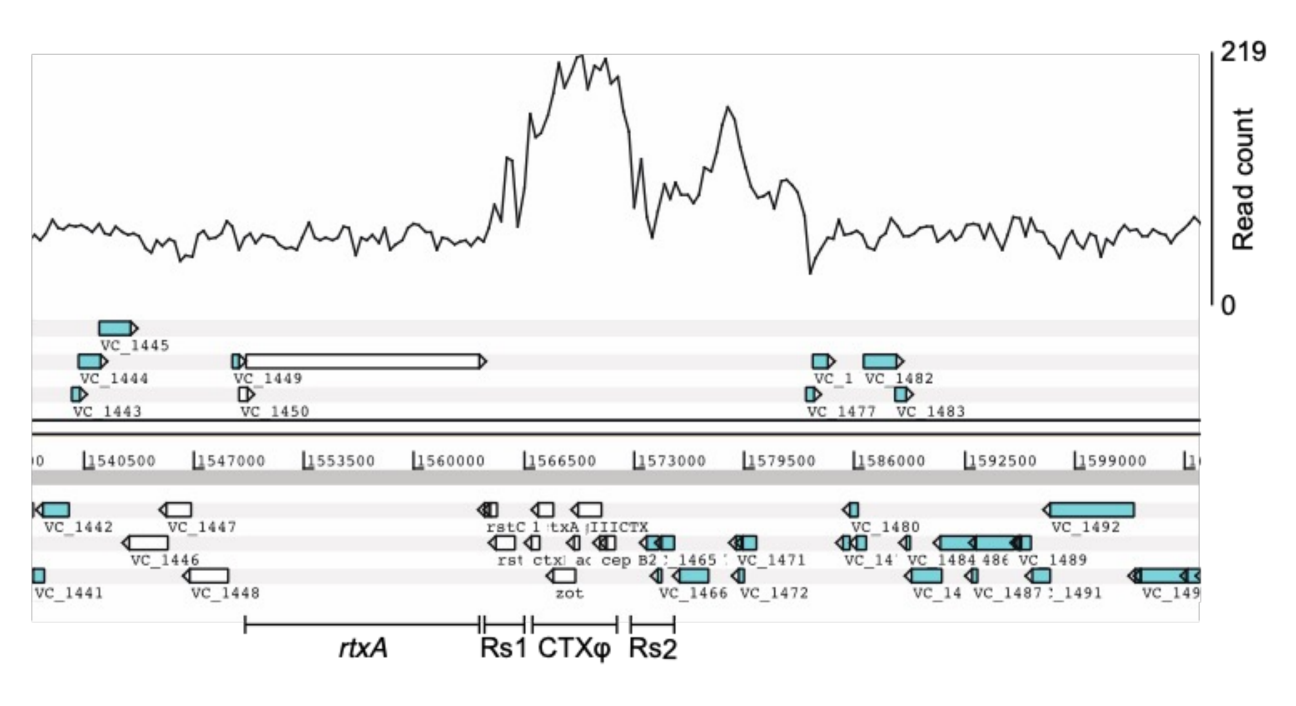
INDELS

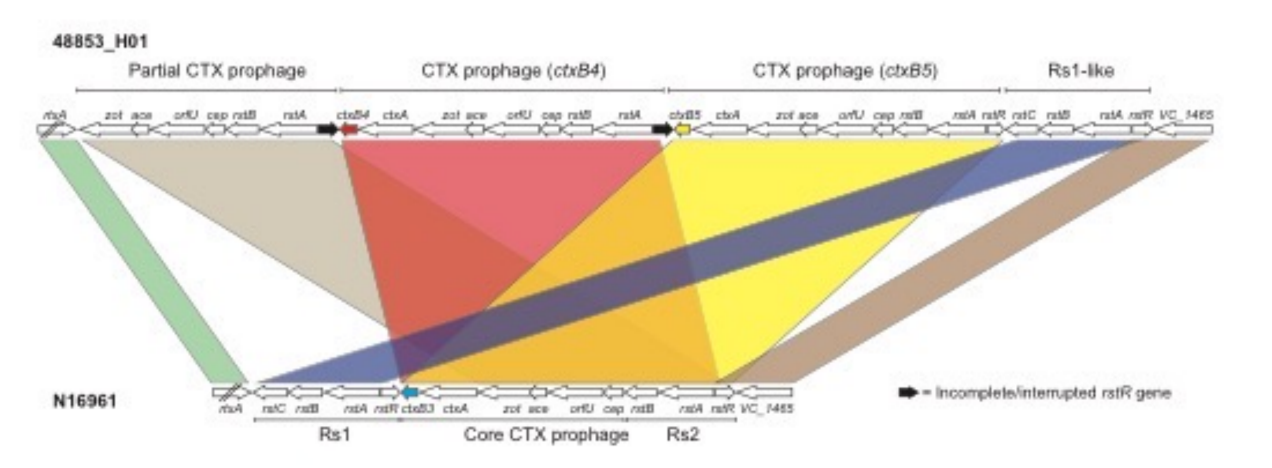


Why do we map to a reference?

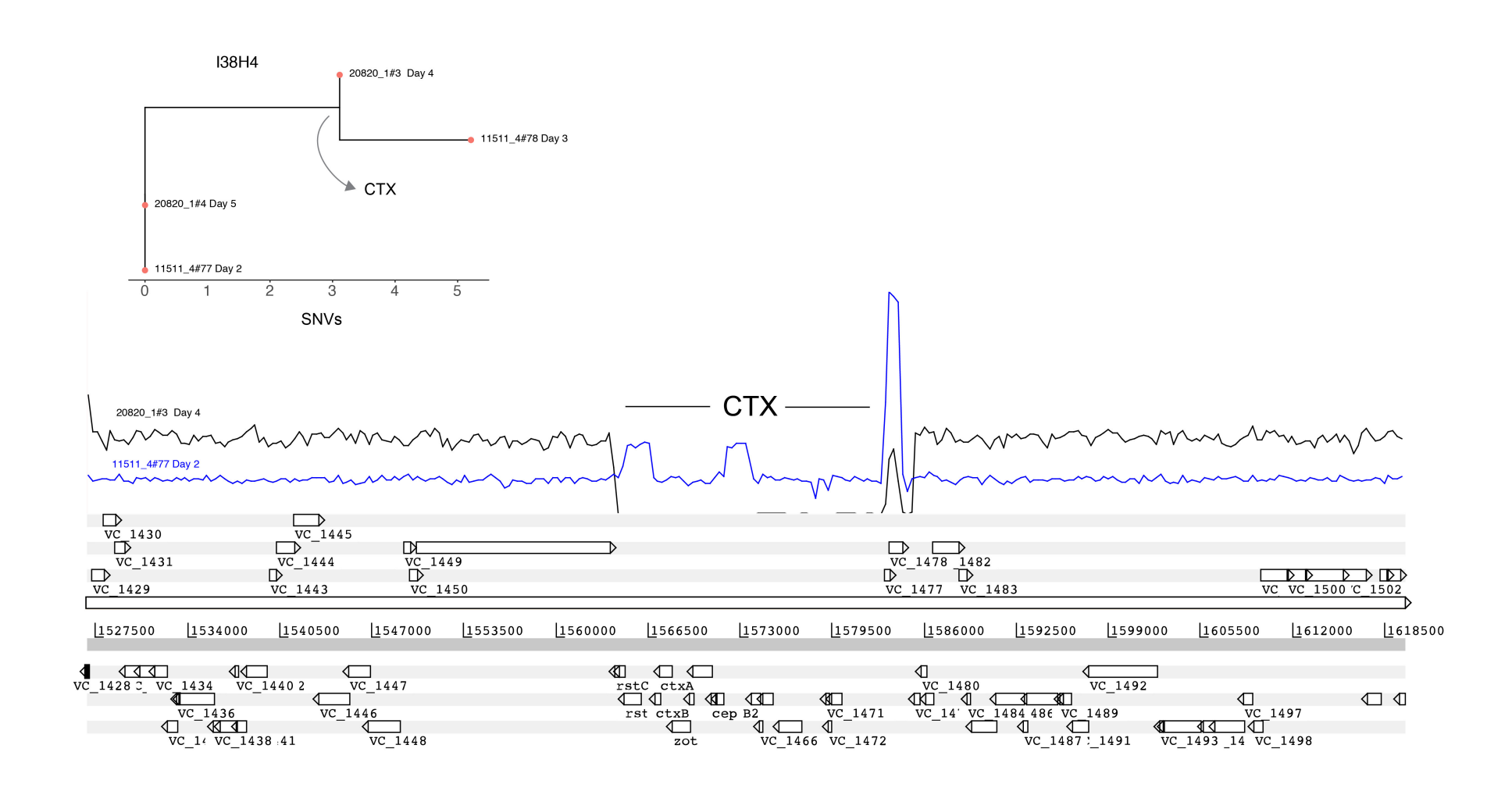
- Identify variation:
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Copy number variation



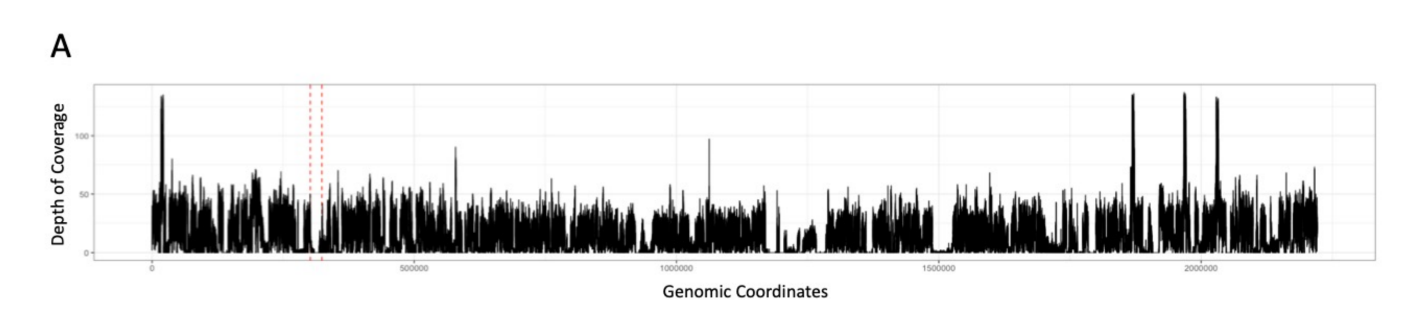


Gene presence / absence

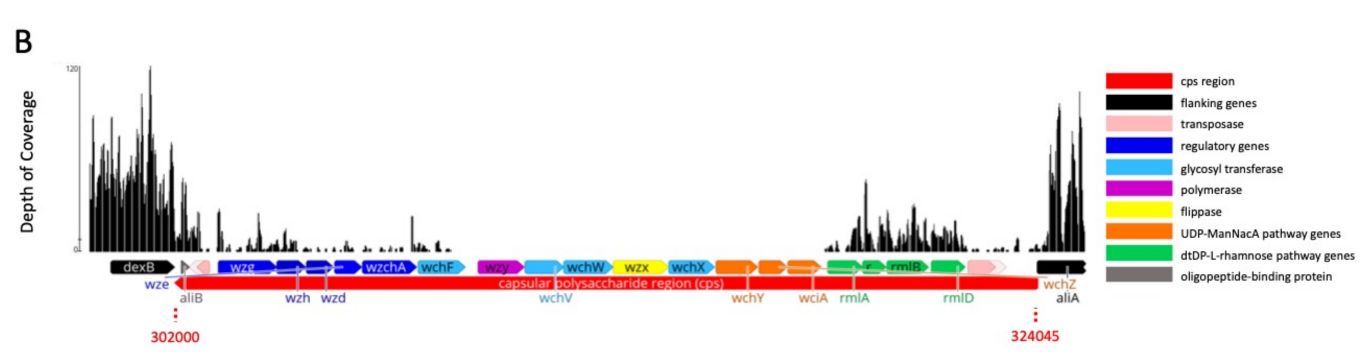


Gene presence/absence: AMR

- Absence/Deletions is easier to spot
- To identify insertions is a little tricky







Today's Agenda







Questions?

Mapping Module