



Session 4.3 - Annotation

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3 al 10 noviembre 2022 CURSO FORMACIÓN AESAN-CNA





Bacterial genome characteristics

- A bacterial genome is a single "circular" DNA molecule with several million base pairs in size
- Bacteria can contains plasmids (small and circular DNA molecules, that contain (usually) non-essential genes)
- Genomes contain a few thousand genes.
- "Gene density" is much higher than in humans, one million base pairs of bacterial DNA contains about 500 to 1000 genes.
 - bacterial genes have no introns,
 - the average number of codons in bacterial genes is less than in human genes
 - neighboring genes are very close together throughout the genome



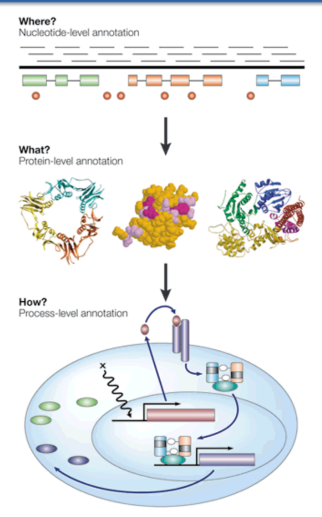


Annotation

Genome annotation is the process of attaching biological (and positional) information to sequences. It consists of three main steps:

- identifying portions of the genome that do not code for proteins
- Identifying coding elements on the genome, a process called gene prediction
- attaching biological information to these elements

https://galaxyproject.github.io/training-material/topics/genome-annotation/tutorials/genome-annotation/tutorial.html







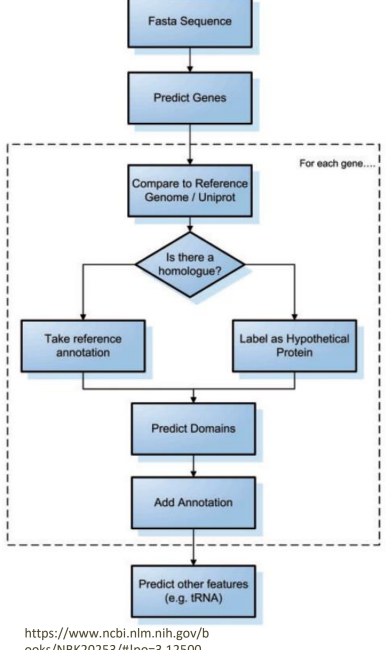
Main categories

- Structural annotation Finding genes and other biologically relevant sites with specific locations but unknown function
 - ORFs
 - Coding sequences(cds)
 - Promoters and regulatory regions
- Functional annotation Elements are used in database searches to attach biologically relevant information to whole sequence and individual objects



Automatic annotation

- Exponential submission of bacterial genomes
- **Databases**
 - Uniprot
 - RefSeq
 - Encyclopedia of DNA elements (ENCODE)
 - Entrez Gene
 - Ensembl
 - GENCODE
 - Gene Ontology Consortium (COGs)
 - GeneRIF
 - KEGG
 - Vertebrate and Genome Annotation Project (Vega)
 - Pfam
 - etc







Automatic annotation

Two strategies for identifying coding genes:

- Sequence alignment to find known protein sequences in the contigs
 - transfer the annotation across
 - will miss proteins not present in your database
 - may miss partial proteins
- Ab initio gene finding o find candidate open reading frames:
 - Build model of ribosome binding sites
 - predict coding regions
 - may choose the incorrect start codon
 - may miss atypical genes, overpredict small genes





Automatic annotation

- tRNA: easy to find and annotate: anti-codon
- rRNA: easy to find and annotate: 5s 16s 23s
- CDS: straightforward to find candidates
 - false positives are often small ORFs
 - wrong start codon o partial genes
 - Pseudogenes
 - assigning function is the bulk of the workload





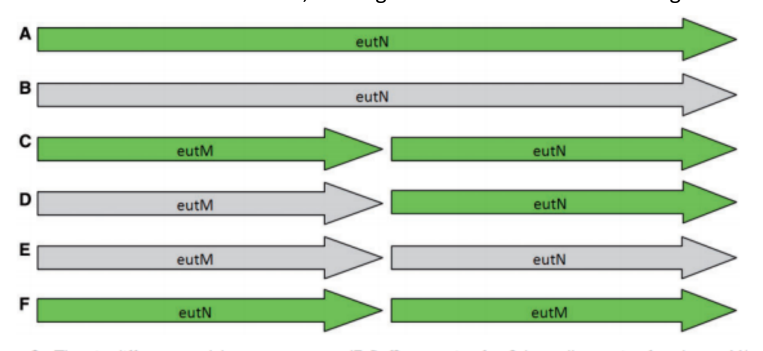
- If sequence homologous are found, may not be functional homologous
- If no homology found- limited information can be inferred
- Incorrect annotation can be propagated when similarity is over part on sequence not used in annotation
 - Multidomain proteins (HMM)
- Inconsistent annotation (Different names, same protein)
- Same gene name, different product name
- Spelling mistakes
- Looking for new genes, not present in DDBB
- Expression experiments / Manual annotation needed

Richardson and Watson. Briefings in Bioinformatics. 2012





Inconsistent annotation, en un gen descrito evento de fusión genica



Salmonella typhi CT18 (NC_003198) and Salmonella typhi Ty2 (NC_004631) there is a single ORF of 690 bp

Figure 2: The six different models present across I7 RefSeq entries for Salmonella species for the eutM/eutN locus. Green indicates normal gene/CDS features, lighter grey indicates gene features annotated as pseudogenes.

- (A) A single intact gene of 690 bp; (B) a single pseudogene of 690 bp; (C) two short intact genes ~300 bp in length;
- (D) one pseudogene and one intact gene, each ~300 bp in length; (E) two pseudogenes, each 300 bp in length; and
- (F) two intact genes with the order reversed.

Richardson and Watson. Briefings in Bioinformatics. 2012





Inconsistent annotation

These two regions are more than 97% identical at the nucleotide level; however, the annotation differs considerably.

While E. coliK12MG1655 contains features with gene names araA, araB and araC, the equivalent features in E. coli 0157:H7 Sakai do not have those gene names and have been assigned uninformative locus tags

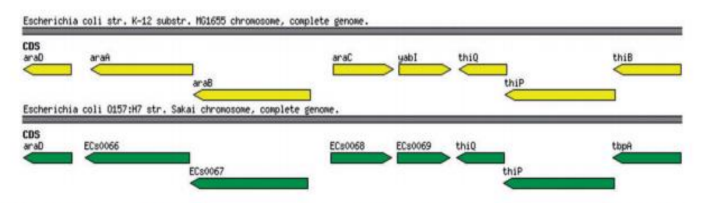


Figure 3: A syntenic block of genes showing inconsistent gene name annotations in E.coli KI2 MGI655 and E. coli 0157:H7 Sakai.





Spelling mistakes

- There are 128 proteins in UniProt that contain the word 'syntase', an incorrect spelling of the word 'synthase'
- If a user was to visit any of these databases and search for 'dihydrofolate synthase' the misspelled entries would be omitted from the search results





- <u>'Same gene name, different product name'</u>
 - The NCBI validation software specifically highlights when this occurs intra-genomically with the description 'Same gene name, different product name'

Table 1: Different product names assigned to features with the gene name 'int' across 17 different RefSeq entries for Salmonella species

| Gene name | Product name | Accession |
|-----------|---|--|
| int | bacteriophage integrase | NC.003198, NC.004631, NC.015761 |
| int | Gifsy-I prophage Int | NC.006905 |
| nt | hypothetical protein | NC.006905 |
| nt | Integrase | NC.003198, NC.004631, NC.006511, NC.012125 |
| nt | integrase (fragment) | NC.003I98 |
| it | phage integrase family site specific recombinase | NC.006905 |
| it | putative cytoplasmic protein | NC.006905 |
| t | Putative integrase | NC.003384 |
| t | putative integrase protein | NC.006905 |
| t | putative P4-type integrase | NC.006905 |
| t | putative phage integrase protein | NC.006905 Richardson and Watson. Brie |
| t | site-specific recombinase, phage integrase family | NC012125 in Bioinformatics. 2012 |





Hypothetical proteins

- These may be real genes with no known function or they may be artifacts of the gene prediction process.
- Often there are features which are only orthologous to other hypothetical features and do not contain any domains. These could either be regions with no functionality, a relic of the feature prediction software or the domains present have not been discovered yet
- Whether or not to include them is often a decision made by the annotation team and varies between groups
- As experimental data becomes more ubiquitous evidence tags should play a larger role in annotation.





<u>Distinguishing orthologs from paralogs</u>

orthologs tend to retain similar functions, whereas paralogs tend to diverge over time to perform different functions

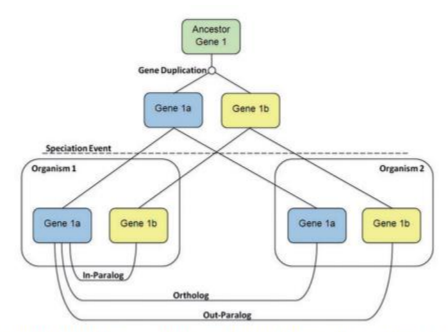


Figure 4: A diagram displaying the processes that can lead to, and define, orthologs and paralogs. Gene duplication and speciation events create complex evolutionary relationships between genes.

Richardson and Watson. Briefings in Bioinformatics. 2012





- RefSeq is one attempt to standardize and improve the quality of genome annotation
 - WP_ prefix. All identical proteins regardless of species
 - Standard classification

```
beta-lactamase (conceptual)
   class A beta-lactamase (HMM:NF033103)
   metallo-beta-lactamase (HMM:NF012229)
      subclass B1 metallo-beta-lactamase (HMM:NF033088)
         NDM family subclass B1 metallo-beta-lactamase (HMM:NF000259)
             subclass B1 metallo-beta-lactamase NDM-1 (allele)
             subclass B1 metallo-beta-lactamase NDM-2 (allele)
             subclass B1 metallo-beta-lactamase NDM-3 (allele)
         VIM family subclass B1 metallo-beta-lactamase (HMM:NF012100)
         SPM family subclass B1 metallo-beta-lactamase (HMM:NF012150)
      subclass B2 metallo-beta-lactamase (HMM:NF033087)
      subclass B3 metallo-beta-lactamase (HMM:NF033105)
   class C beta-lactamase (HMM:NF033085)
   class D beta-lactamase (conceptual)
      class D beta-lactamase (main branch) (HMM:NF012161)
      class D beta-lactamase (other branch) (HMM:NF000270)
```



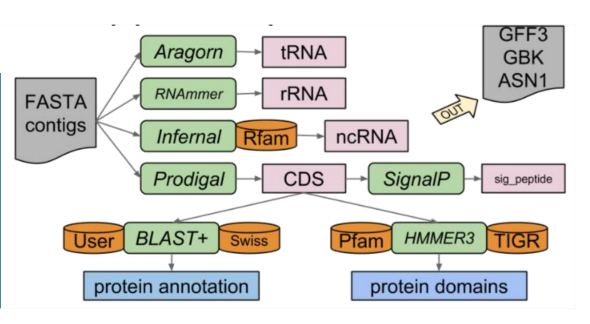
annotation)



Automatic annotation: Prokka (Rapid prokaryotic genome

Seeman, Bioinformatics 2014

| Features predicted | | | | |
|----------------------------|--|--|--|--|
| Coding sequence (CDS) | | | | |
| Ribosomal RNA genes (rRNA) | | | | |
| Transfer RNA genes | | | | |
| Signal leader peptides | | | | |
| Non-coding RNA | | | | |
| | | | | |
| Specific function or name | | | | |
| Personal database | | | | |
| | | | | |



https://galaxyproject.github.io/training-material/topics/genome-annotation/tutorials/annotation-with-prokka/slides.html#8





- Optional user-provided set of annotated proteins
- All bacterial proteins in UniProt
- All proteins from finished bacterial genomes in RefSeq
- Hidden Markov model profile databases, Pfam and TIGRFAMs
- Hypothetical protein

Prokka uses this method, but in a hierarchical manner, starting with a smaller trustworthy database, moving to medium sized but domain-specific databases, and finally to curated models of protein families





Facts

- searching against smaller databases is faster
- searching against similar sequences is faster

• <u>Idea</u>

- start with small set of close proteins
- advance to larger sets of more distant proteins

• Prokka

- your own custom "trusted" set (optional)
- core bacterial proteome (default)
- genus specific proteome (optional)
- whole protein HMMs: PRK clusters, TIGRfams
- protein domain HMMs: Pfam

Prokka uses this method, but in a hierarchical manner, starting with a smaller trustworthy database, moving to medium sized but domain-specific databases, and finally to curated models of protein families





Core Bacterial proteome

- Many bacterial proteins are conserved
 - experimentally validated
 - good annotations
- Prokka provides this database
 - derived from UniProt-Swissprot
 - only bacterial proteins
 - only accept evidence level 1 (aa) or 2 (RNA)
 - reject "Fragment" entries
 - extract /gene /EC_number /product /db_xref ●
- First step gets ~50% of the genes
 - BLAST+ blastp, multi-threading to use all CPUs





Prokka has genus specific databases

- aim to capture "genus specific" naming conventions
- derived from proteins in completed genomes
- proteins are clustered and majority annotation wins
- some annotations are rubbish though

Custom model databases

I took COG/PRK MSAs and made HMMs

Existing model databases

- Pfam, TIGRfams are well curated

And if all else fails

— we always have our friend "hypothetical protein"





Automatic annotation: Prokka output

| Suffix | Description of file contents | |
|--------|---|--|
| .fna | FASTA file of original input contigs (nucleotide) | |
| .faa | FASTA file of translated coding genes (protein) | |
| .ffn | FASTA file of all genomic features (nucleotide) | |
| .fsa | Contig sequences for submission (nucleotide) | |
| .tbl | Feature table for submission | |
| .sqn | Sequin editable file for submission | |
| .gbk | Genbank file containing sequences and annotations | |
| .gff | GFF v3 file containing sequences and annotations | |
| .log | Log file of Prokka processing output | |
| .txt | Annotation summary statistics | |





Annotation format: gff3

```
##gff-version 3.2.1
Segid - name
                             ##sequence-region ctg123 1 1497228
Source - program
                              ctg123 . gene
                                                                         ID=gene00001;Name=EDEN
                                                     1000 9000
                              ctg123 . TF_binding_site 1000
                                                                         ID=tfbs00001;Parent=gene00001
                                                          1012
Type - term or SOFA
                             ctg123 . mRNA
                                                                         ID=mRNA00001; Parent=gene00001; Name=EDEN.1
sequence ontology
                              ctg123 . mRNA
                                                                         ID=mRNA00002; Parent=gene00001; Name=EDEN. 2
Start
                              ctg123 . mRNA
                                                     1300
                                                                         ID=mRNA00003; Parent=gene00001; Name=EDEN.3
End
                              ctg123 . exon
                                                     1300
                                                                         ID=exon00001; Parent=mRNA00003
                              ctg123 . exon
                                                                         ID=exon00002; Parent=mRNA00001, mRNA00002
Score
                              ctg123 . exon
                                                      3000
                                                                         ID=exon00003; Parent=mRNA00001, mRNA00003
Strand -(+/-)
                              ctg123 . exon
                                                                         ID=exon00004; Parent=mRNA00001, mRNA00002, mRNA00003
                                                      5000
Phase -(0/1/2)
                                                                         ID=exon00005; Parent=mRNA00001, mRNA00002, mRNA00003
                              ctg123 . exon
Attributes
                              ctg123 . CDS
                                                                         ID=cds00001;Parent=mRNA00001;Name=edenprotein.1
                             ctg123 . CDS
                                                                         ID=cds00001;Parent=mRNA00001;Name=edenprotein.1
    Name
                                                                         ID=cds00001;Parent=mRNA00001;Name=edenprotein.1
                              ctg123 . CDS
    Alias
                              ctg123 . CDS
                                                                         ID=cds00001;Parent=mRNA00001;Name=edenprotein.1
                              ctg123 . CDS
                                                                         ID=cds00002;Parent=mRNA00002;Name=edenprotein.2
    Parent
                                                                         ID=cds00002;Parent=mRNA00002;Name=edenprotein.2
                              ctg123 . CDS
    Target
                              ctg123 . CDS
                                                                         ID=cds00002;Parent=mRNA00002;Name=edenprotein.2
    Gap
                              ctg123 . CDS
                                                                        ID=cds00003;Parent=mRNA00003;Name=edenprotein.3
                                                      3301
    Derives from
                                                                . + 1 ID=cds00003;Parent=mRNA00003;Name=edenprotein.3
                              ctg123 . CDS
                              ctg123 . CDS
                                                                . + 1 ID=cds00003;Parent=mRNA00003;Name=edenprotein.3
    Note
                                                                . + 0 ID=cds00004;Parent=mRNA00003;Name=edenprotein.4
                              ctg123 . CDS
                                                      3391
    Dbxref
                             ctg123 . CDS
                                                                . + 1 ID=cds00004;Parent=mRNA00003;Name=edenprotein.4
    Ontology term
                              ctg123 . CDS
                                                                . + 1 ID=cds00004;Parent=mRNA00003;Name=edenprotein.4
```





Annotation format: gbk

- LOCUS Annotated sequence
- DEFINITION
- ACCESION
- FEATURES
 - source
 - gene
 - CDS
 - Locus tag
 - function
 - Product
 - protein_id
 - Translation (sequence)

```
LOCUS
            AF068625
                                     200 bp
                                                       linear
                                                              ROD 06-DEC-1999
DEFINITION Mus musculus DNA cytosine-5 methyltransferase 3A (Dnmt3a) mRNA,
            complete cds.
ACCESSION
            AF068625 REGION: 1..200
VERSION
            AF068625.2 GI:6449467
KEYWORDS
SOURCE
            Mus musculus (house mouse)
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia;
            Sciurognathi; Muroidea; Muridae; Murinae; Mus.
REFERENCE
           1 (bases 1 to 200)
  AUTHORS
            Okano, M., Xie, S. and Li, E.
            Cloning and characterization of a family of novel mammalian DNA
            (cytosine-5) methyltransferases
  JOURNAL
            Nat. Genet. 19 (3), 219-220 (1998)
            9662389
REFERENCE
           2 (bases 1 to 200)
  AUTHORS
           Xie, S., Okano, M. and Li, E.
  TITLE
            Direct Submission
            Submitted (28-MAY-1998) CVRC, Mass. Gen. Hospital, 149 13th Street,
            Charlestown, MA 02129, USA
REFERENCE
           3 (bases 1 to 200)
           Okano, M., Chijiwa, T., Sasaki, H. and Li, E.
  TITLE
            Direct Submission
            Submitted (04-NOV-1999) CVRC, Mass. Gen. Hospital, 149 13th Street,
            Charlestown, MA 02129, USA
            Sequence update by submitter
COMMENT
            On Nov 18, 1999 this sequence version replaced gi:3327977.
FEATURES
                     Location/Qualifiers
     source
                     1..200
                     /organism="Mus musculus"
                     /mol type="mRNA"
                     /db xref="taxon:10090"
                     /chromosome="12"
                     /map="4.0 cM"
     gene
                     1..>200
                     /gene="Dnmt3a"
ORIGIN
       1 gaattccggc ctgctgccgg gccgcccgac ccgccgggcc acacggcaga gccgcctgaa
       61 gcccagcgct gaggctgcac ttttccgagg gcttgacatc agggtctatg tttaagtctt
      121 agctcttgct tacaaagacc acggcaattc cttctctgaa gccctcgcag ccccacagcg
      181 ccctcgcagc cccagcctgc
//
```





Annotation format: gbk

- LOCUS Annotated sequence
- DEFINITION
- ACCESION
- FEATURES
 - source
 - gene
 - CDS
 - Locus tag
 - function
 - Product
 - protein_id
 - Translation (sequence)

```
FEATURES
                     Location/Qualifiers
                     /organism="Klebsiella pneumoniae subsp. pneumoniae SA1"
                     /mol_type="genomic DNA"
                     /strain="SA1"
                     /sub species="pneumoniae"
                     /db_xref="taxon: 1379688"
                     /note="contig LPSB1_2557_Contig_49"
                     415..1536
    gene
                     /locus_tag="KPST86_490001"
                     415..1536
                     /locus tag="KPST86 490001"
                     /inference="ab initio prediction:AMIGene:2.0"
                     /note="Evidence 4:Homologs of previously reported genes of
                     unknown function"
                     /codon start=1
                     /transl_table=11
                     /product="conserved hypothetical protein"
                     /protein id="CDI25656.1"
                     /translation="MAYQLNINWPEFLEKYWQKQPVVLKNAFPDFVDPITPDELAGLA
                     MEPEVDSRLVSLKNGKWOASNGPFEHFDGLGETGWSLLAOAVNHWHMPAAELVRPFRV
                     LPDWRLDDLMISFSVPGGGVGPHIDQYDVFIIQGMGSRRWRVGDKLPMRQFCPHPALL
                     HVDPFPPIIDEDLQPGDILYIPPGFPHDGITHETALNYSVGFRGPNGRDLISSFADYV
                     LENDLGDEHYSDPDLTCREHPGRVEEYELERLRTMMIDMIRQPEDFKQWFGSFVTTPR
                     HELDIAPAEPPYEEEEVLDALLGGEKLSRLSGLRVLHIGDSFFVHSEOLDTTDAEALD
                     ALCRYTSLGQEELGSGLQNPAFVSELTRLINQGYWYFEE"
                     complement(1584..2117)
                     /locus_tag="KPST86_490002"
                     complement(1584..2117)
                     /locus tag="KPST86 490002"
                     /inference="ab initio prediction:AMIGene:2.0"
                     /note="Evidence 4:Homologs of previously reported genes of
                     unknown function"
                     /codon_start=1
                     /transl table=11
                     /product="conserved hypothetical protein"
                     /protein id="CDI25658.1"
                     /translation="MEQQLTIEMIADAFSYDITGFDCGEEALNTFLKEHLKRQHDGQI
                     LRGYALVSGDTVPRLLGYYTLSGSCFERGMLPSKTQQKKIPYQNAPSVTLGRLAIDKS
                     VQGQGWGEMLVAHAMRVVWGASKAVGIYGLFVEALNEKAKAFYLRLGFIQLVDENSNL
                     LFYPTKSIEOLFTDDES"
                     complement(2128..2394)
     gene
                     /locus tag="KPST86 490003"
                     complement(2128..2394)
     CDS
                     /locus tag="KPST86 490003"
                     /inference="ab initio prediction:AMIGene:2.0"
                     /note="Evidence 4:Homologs of previously reported genes of
                     unknown function'
```





Resistance prediction using WGS

Hendrisken et al. Frontiers in Microbiology. 2019.

Concordance between phenotypic susceptibility testing and WGS based predicted antimicrobial resistance

| | Pathogen | No. of pathogens | AST method | No. of antimicrobials | Bioinformatic tool | Sequencing data | Concordance | Sensitivity | Specificity | Comment | References |
|------|------------------------|------------------|-------------------|-----------------------|-------------------------------------|---------------------------|-------------|-------------|-------------|--|------------|
| 2013 | S. Typhimurium | 49 | MIC | 17 | ResFinder | Assembled, Velvet | 99.74% | | | Disagreement: 7 isolates | (7) |
| | E. coli | 48 | | | | | | | | including 6 E. coli resistent to | |
| | E. faecalis | 50 | | 14 | | | | | | Spec | |
| | E. faecium | 50 | | | | | | | | | |
| 2013 | E. coli (ESBL) | 74 | DD | 7 | BLASTn, selected panel | Assembled, Velvet | | 96% | 97% | VM rate: 1.2%/M rate: 2.1% | (8) |
| | K. pneumonia (ESBL) | 69 | | | | | | | | | |
| 2014 | S. aureus | 501 | DD/MIC (Vitek) | 12 | BLASTn, selected panel | Assembled, Velvet | | 97% | 99% | VM rate: 0.5%/M rate: 0.7% | (9) |
| 2016 | C. jejuni | 32 | MIC | 9 | BLASTx | Assembled, | 99.2% | | | Lower concordance to | (10) |
| | C. coli | 82 | | | | CLC-bio | | | | Gen, Azi, Clin, Tel | |
| 2016 | S. enterica | 104 | MIC | 14 | ResFinder/ ARG-ANNOT/ | Assembled, CLC-bio | 99.0% | 99.2% | 99.3% | Lower concordance to | (11) |
| | | 536 | | | CARD/BLAST | | | 97.6% | 98.0% | aminoglycosides/β-lactams | |
| 2017 | E. coli | 31 | MIC | 4 | Custom DB based on | | | 87% | 98% | Neg. predictive value: 97% | (12) |
| | K. pneumonia | 24 | | | ARDB/CARD/β- lactamase | | | | | Pos. Predictive value: 91% | |
| | P. aeruginosa | 22 | | | allelles | | | | | | |
| | E. cloacae | 13 | | | | | | | | | |
| 2017 | S. enterica | 50 | MIC | 4 | ResFinder/ PointFinder | Assembled, SPAdes | 98.4% | | | Disagreement: | (13) |
| | E. coli | 50 | | 6 | Pointringer | | | | | 2/2 C.jejuni to FQ/ERY | |
| 0010 | C. jejuni | 50 | | 4 | 5 5 1 1100 | | 00.50/ | | | 5 E. coli to COL (pmrB) | |
| 2018 | E. faecalis | 97 | MIC | 11 | ResFinder/NCBI Pathogen DB/BLAST | Assembled, CLC-bio | 96.5% | | | | (14) |
| 0040 | E. faecium | 100 | DD # #0 | 10 | GeneFinder/ | | 00.00/ | | | D: | (45) |
| 2018 | S. aureus | 501 491 | DD/MIC | 12 | Mykrobe/ | FASTQ/assembled, BLAST | 98.3% | | | Disagreements: 0.7% predicted resistant | (15) |
| | | 397 | MIC | | Typewriter | | | | | 0.6% predicted susceptible | |
| 2018 | M. tuberculosis | 10,209 | MGIT 960 | 4 | Cortex | Assembled | 89.5% | | | 97.1%/99.0% predicted R/S 97.5%/98.8% predicted R/S | (16) |
| | | | | 4 | | | | | | 94.6%/93.6% predicted R/S | |
| | | | | 4 | | | | | | 91.3%/96.8% predicted R/S | |
| 2019 | H. pylori | 140 | MIC (E-test) | 5 | ARIBA | FASTQ | 99% | | | Phenotype issues to metronidazole | (17) |

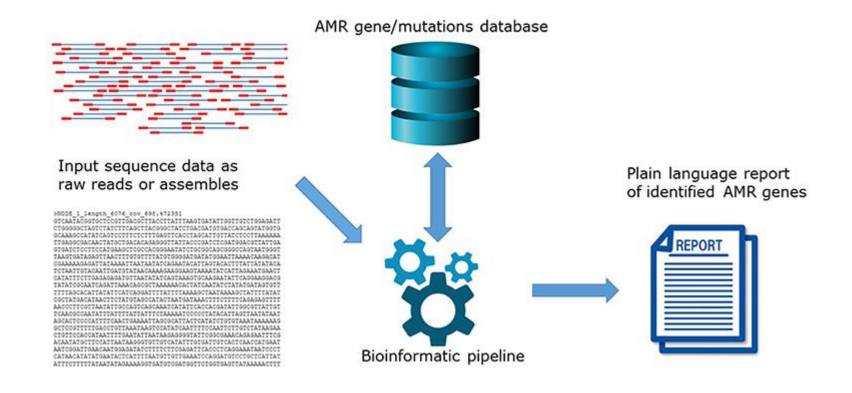
¹⁾ ESBL: Extended Spectrum Beta-Lactamase, 2) MIC: Minimum Inhibitory Concentration, 3) DD: Disk diffusion, 4) VM: Very Major, 5) M: Major, 6) R/S: Resistant/Susceptible, 7) SPEC: Spectinomycin, 8) GEN: Gentamicin, 9) AZI: Azithromycin, 10) CLIN: Clindamycin, 11) TEL: Telithromycin, 12) FQ: Fluoroquinolone, 13) ERY: Erythromycin, 14) COL: colistin.





Resistance prediction using WGS

Hendrisken et al. Frontiers in Microbiology. 2019.







Resistance prediction using WGS

Hendrisken et al. Frontiers in Microbiology. 2019.

Huge list here: https://www.frontiersin.org/files/Articles/478239/fpubh-07-00242 HTML/image m/fpubh-07-00242-t002.jpg

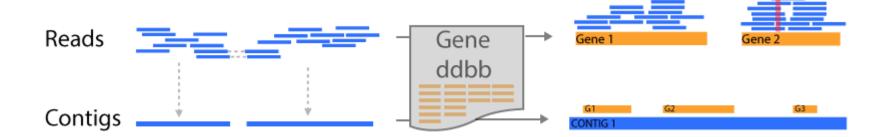
| Software | Туре |
|-----------|--------------------|
| SRST2 | Mapping |
| Ariba | Mapping + assembly |
| ABRICATE | Assembly |
| ResFinder | Assembly |





Mapping vs Assembly

- Functional annotation based on mapping (srst2)
 - Pro: more resolutive / high quality ddbb
 - Con: Unable to locate genes / no ab initio annotation
- Functional annotation based on assembly (Resfinder)
 - Pro: genes are located / related
 - Depend on assembly (close to repetitive regions)







INPUT

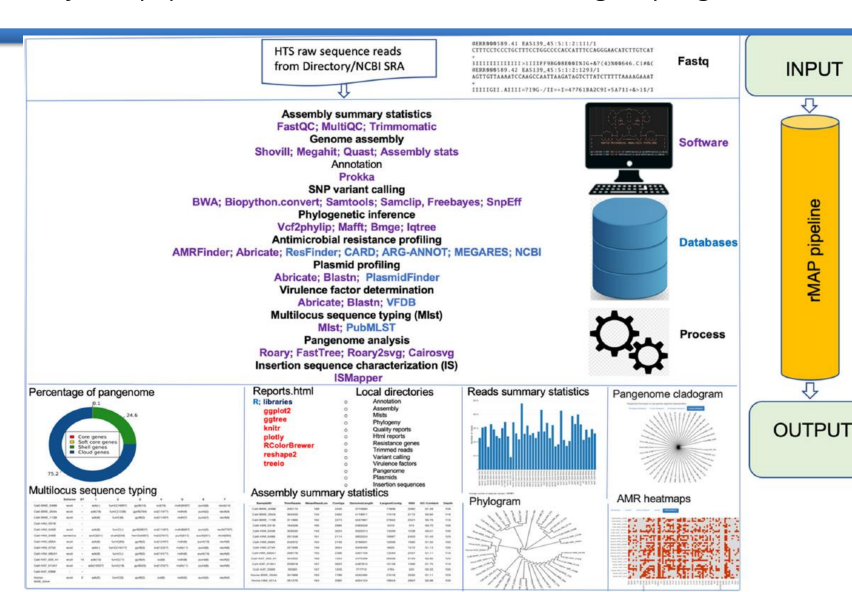
pipeline

MAP

rMAP - rapid microbial analysis pipeline- for ESKAPE bacterial group wgs data

The resistomes of ESKAPE pathogens (Enterococcus faecium, Staphylococcus aureus, Klebsiella pneumoniae, Acinetobacter baumannii, Pseudomonas aeruginosa and Enterobacter species)

Sserwadda & Mboowa, Microbiology, 2021

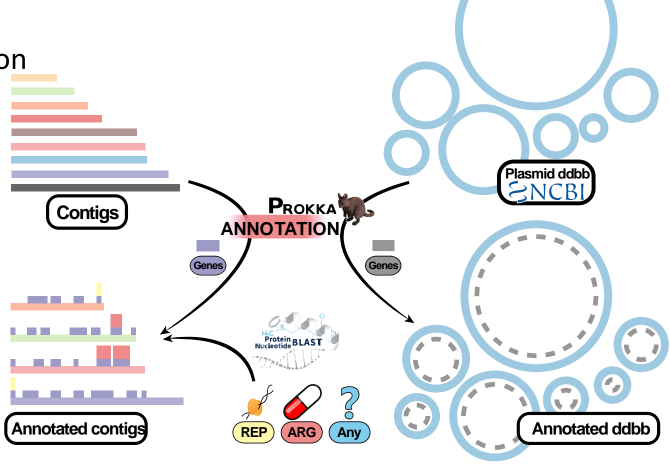






Annotation visualization using PlasmidID

- Automatic annotation
 - Prokka
 - DDBB plasmid
 - Contigs
 - Gff to bed
- Specific annotation
 - BLAST+
 - ABR & REP
 - User input FASTA





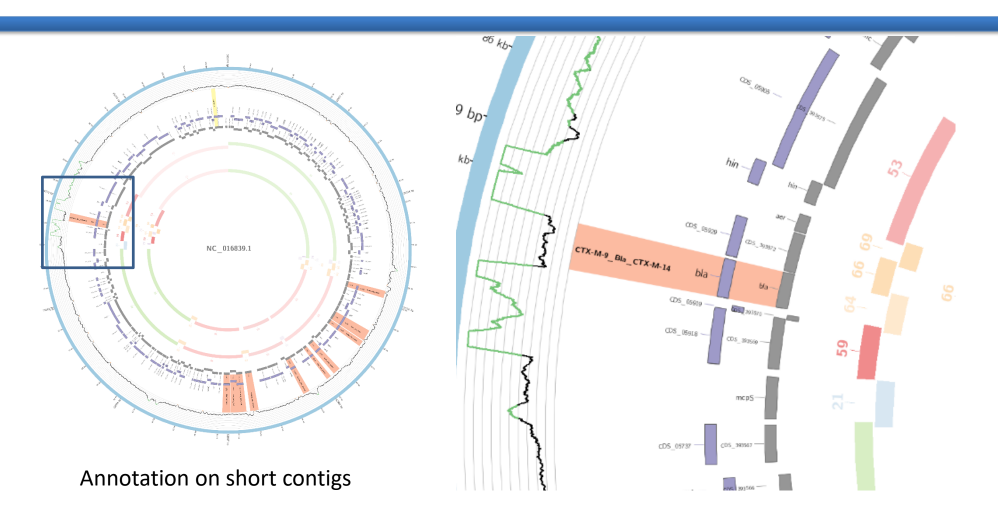


Annotation using PlasmidID Aligned contigs **Mapped plasmids** circos. Annotated contigs Annotated ddbb





Annotation using PlasmidID







Plasmid Track 0000 100 80% **PLASMID NAME** - mapped more than [8]0% - representative of similar plasmids in the ddbb

%05





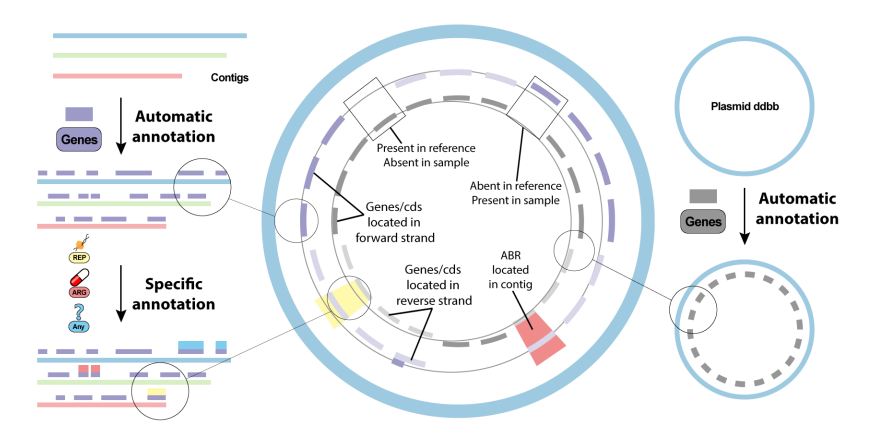
Coverage Track





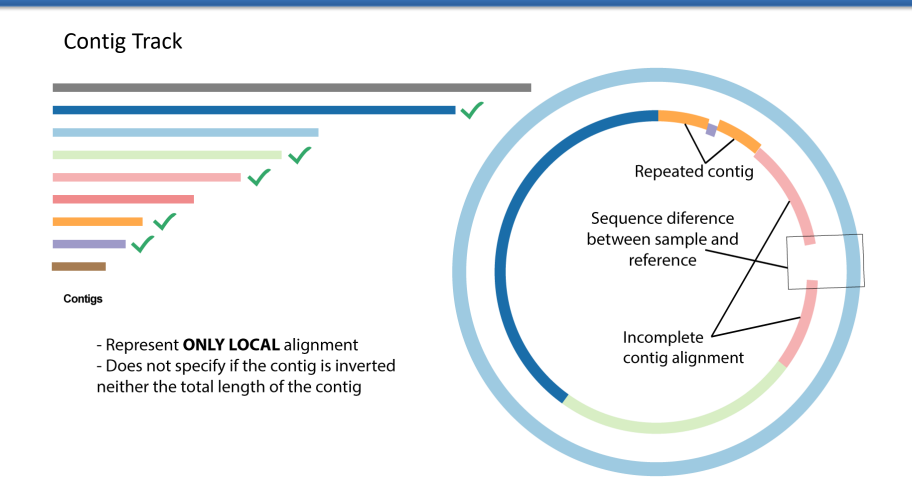


Annotation Track











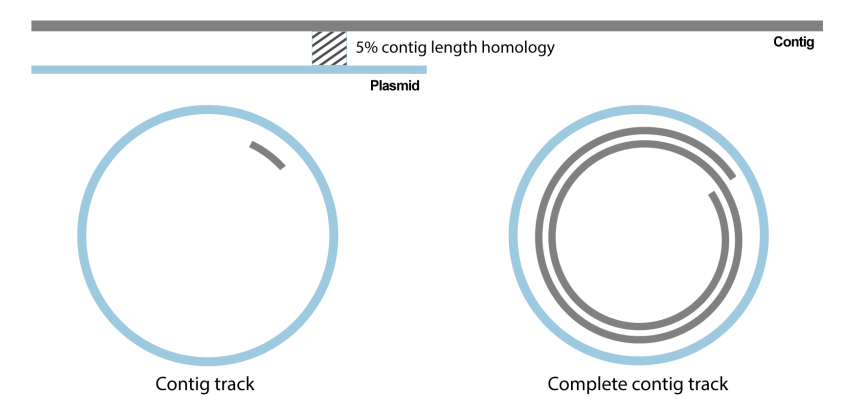


Complete contig Track Alignment is less than 20% of contig size Repeated contigs are absent This region is larger in the This region is difsample so the ferent but the size contig squggle is similar





Complete contig Track

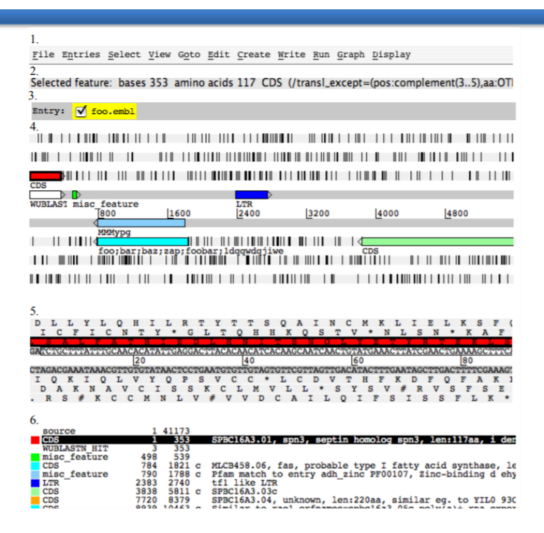






Manual annotation: Artemis

Artemis is a DNA sequence viewer and annotation tool that allows visualisation of sequence features and the results of analyses within the context of the sequence, and its six-frame translation.







Thanks for your attention!