Ultra-long reads and ultra-long duplications: What nanopore sequencing is revealing about Bordetella pertussis

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Background

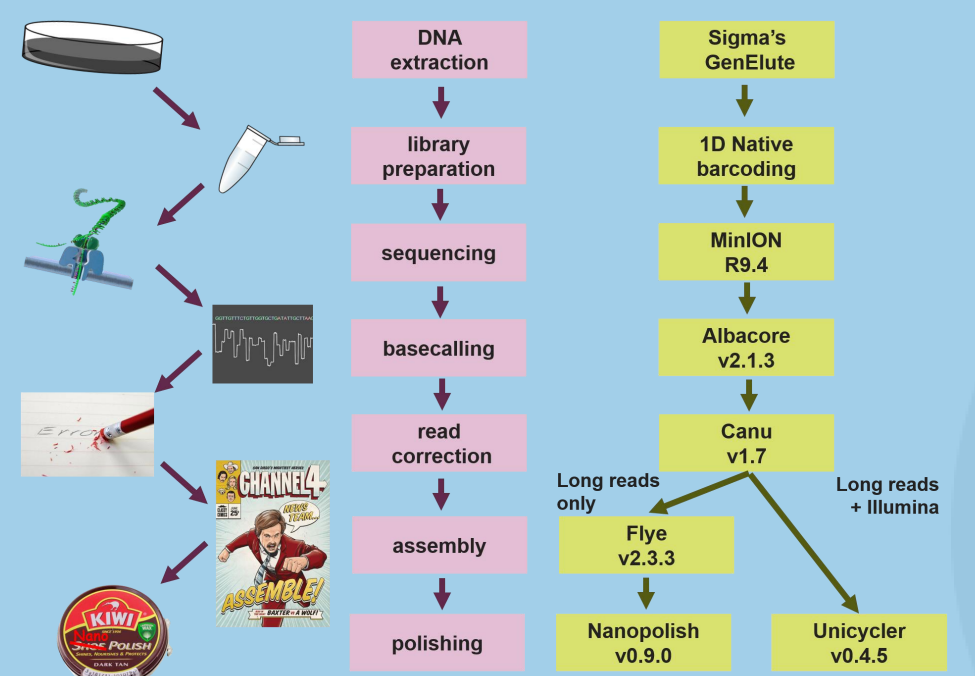
The B. pertussis genome is repetitive. The average B. pertussis genome contains 280 copies of >1,000 bp insertion sequence (IS) elements, representing ~7% of the 4.1 Mb total genome length. The many IS copies mean that closed genome assemblies cannot be produced using short-read sequencing (e.g. Illumina), because each IS element is longer than the short reads

B. pertussis is traditionally described as a monomorphic species: very few base-level differences exist between different strains. The presence of so many mobile IS elements in the genome, however, means that genome-level differences, such as rearrangements, deletions and duplications, are possible We are using long-read sequencing to identity genome-level differences between otherwise highly similar B. pertussis strains

Our nanopore sequencing pipeline*

Through extensive testing and optimisation, we defined a sequencing and data processing pipeline, using Oxford Nanopore Technologies' MinION sequencing, to produce reads longer than 1,000 bp which can be used to assemble closed *B. pertussis* genomes

Using barcodes, up to 12 genomes can be sequenced per flow cell



Our long read genomes

Strain	Contigs	Size Mb	IS 481 copies
UK36	1	4.108	258
UK38	1	4.108	258
UK39	1	4.108	258
UK48	2	4.112	262
UK76	1	4.113	262

We used our nanopore sequencing pipeline to sequence five *B. pertussis* strains, isolated during the 2012 whooping cough outbreak in the UK [1,2]. The genomes of all but one of these strains could be assembled into a closed contig using this pipeline, which produced reads with a mean length >6,000 bp

Two genomes, UK48 and UK76, were longer than the others, and also had more copies of IS 481. On closer inspection, the genomes of these two strains appear to contain the same ultralong duplicated region, ~1.3 Mb into the reference genome

We then used an ultra-long gDNA extraction method [3] to produce 100x coverage of the UK48 genome in reads longer than 100,000 bp, and a maximum read length of 645,000 bp. However, we produced only 30x coverage in reads longer than the duplicated region (~180,000 bp), which was insufficient to produce a closed genome sequence

> Our "ultra-long" **UK48** reads:

Read length / bp	Number of reads
1-100,000	35,968
100,001-200,000	2300
200,001-300,000	277
300,001-400,000	45
400,001-500,000	8
500,000+	2

Some B. pertussis genomes contain long duplicated regions

Even using nanopore sequencing reads longer than IS 481, some strains' genomes cannot be assembled into closed contigs

Mapping the raw reads from these strains to the genome of the reference strain, Tohama I, reveals regions of enriched coverage

In UK48 (below), we see a region of ~0.2 Mb, with almost exactly twice as much coverage as the rest of the genome. This suggests that this region of the genome is present twice in UK48

0 0.5 1.0 1.5 2.0 2.5 3.0 3.5 4.0

For more about copy

number variation in

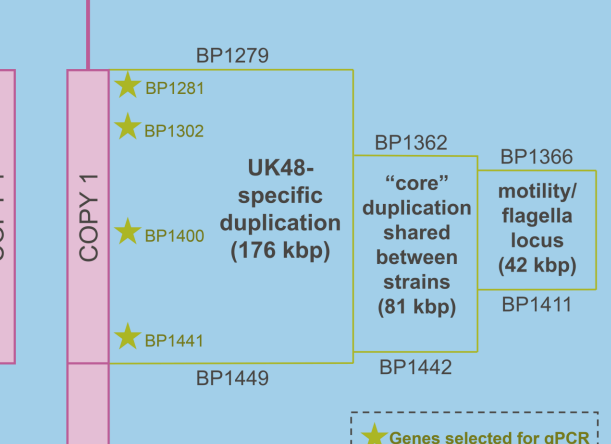
Bordetella pertussis,

visit the poster of

Jonathan Abrahams!



UK71 UK48



Log-phase RNA will be extracted from both strains in Bvg(+) and Bvg(-) conditions, and expression of four genes from the duplicated region will

be measured

Future questions:

duplication

We are using **qPCR** to

expression in UK71, a

does not contain the

in **UK48** with gene

compare gene expression

strain which is highly similar

to UK48 but whose genome

Does the duplication affect growth rate?

Does the duplication affect motility?

Genes (BPxxxx)

N.B. new basecalling tools,

which are more accurate than

Albacore, now exist. A new

pore, R10, was also released in

March 2019

The "motility"

duplication

Over 15 strains sequenced

region which is duplicated in UK48

Although the exact length and genes included in

the duplicated region varies slightly from strain to

strain, the duplication tends to be centred around

globally appear to have a

duplication of the same

and UK76 [4]

*For more information, see Ring et al. 2018 [1]

References

[1] Ring et al. (2018). Resolving the complex Bordetella pertussis genome using barcoded nanopore sequencing. Microbial Genomics 4(11) [2] Sealey et al. (2015). Genomic analysis of isolates from the United Kingdom 2012 pertussis outbreak reveals that vaccine antigen genes are unusually fast evolving. <u>Journal of Infectious Diseases</u> **212**(2) [3] Quick (2018). Ultra-long read sequencing protocol for Rad004. https://www.protocols.io/view/ultra-long-read-sequencing-protocol-for-[4] Abrahams et al. (In preparation). Duplications drive diversity in the

monomorphic pathogen Bordetella pertussis on an underestimated scale.

the same set of "core"

42 of the 77 genes in the

"core" duplication are

related to motility

We are now aiming

to characterise this

duplication further

and/or flagella

synthesis

genes: BP1334 to

BP1492

Tools

https://github.com/bcgsc/abyss https://community.nanoporetech.com/downloads https://github.com/marbl/canu

https://github.com/jts/nanopolish https://github.com/tseemann/prokka

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About the author

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I am a 3nd year PhD student at the University of Bath, researching Bordetella pertussis genomics with an emphasis on sequencing and bioinformatics. I previously worked for 4 years as a bioinformatician at MRC Harwell, and have a PGDip in Science Communication



Scan the QR code to view full methodology, results and data repository

https://github.com/fenderglass/Flye

https://github.com/rrwick/Unicycler