

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

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## Background

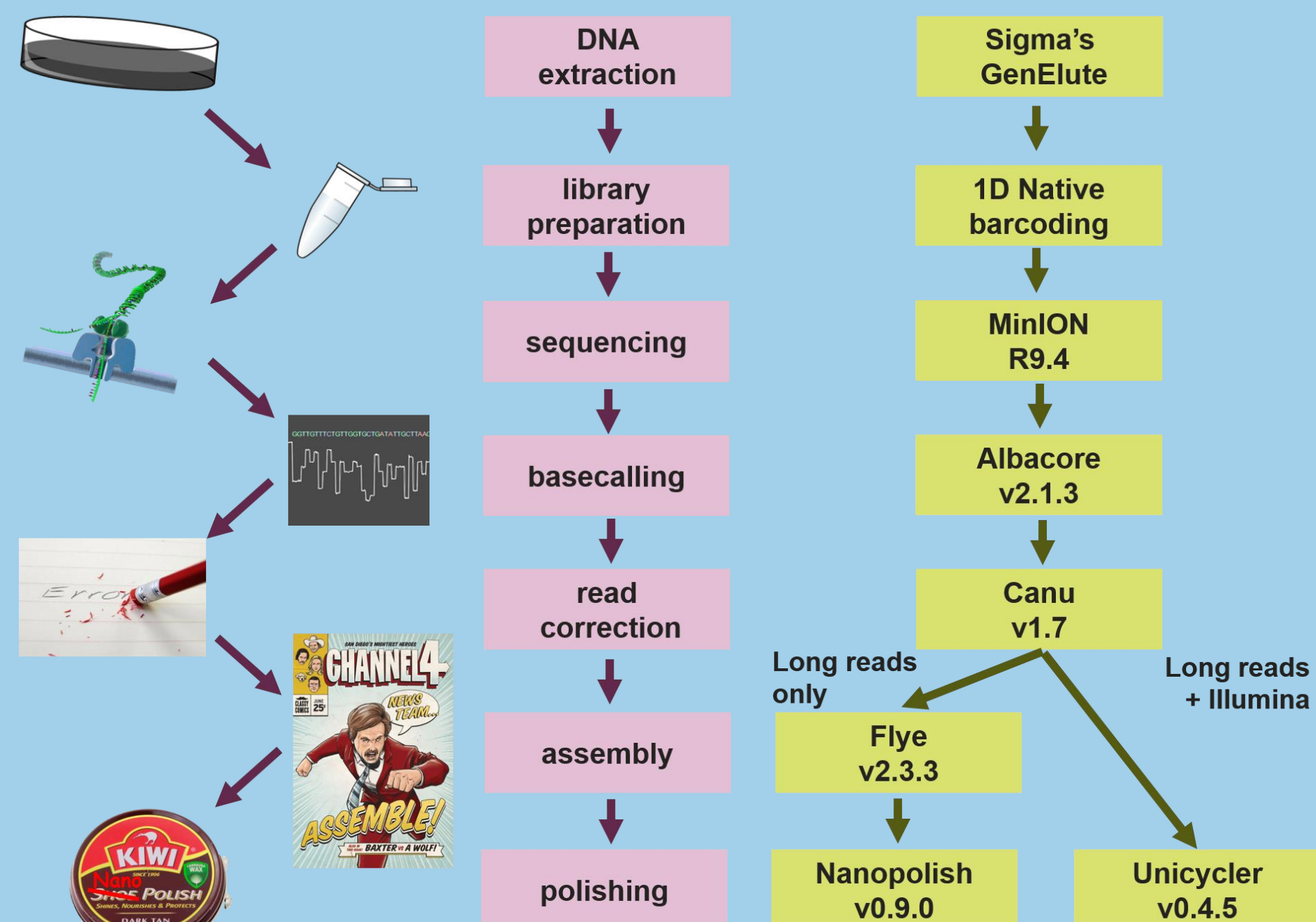
The *B. pertussis* genome is **repetitive**. The average *B. pertussis* genome contains **280 copies of >1,000 bp insertion sequence (IS) elements**, representing **~7% of the 4.1 Mb total genome length**. The many IS copies mean that **closed genome assemblies cannot be produced using short-read sequencing**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



N.B. new basecalling tools, which are more accurate than Albacore, now exist. A new pore, R10, was also released in March 2019

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

## 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 ultra-long 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**

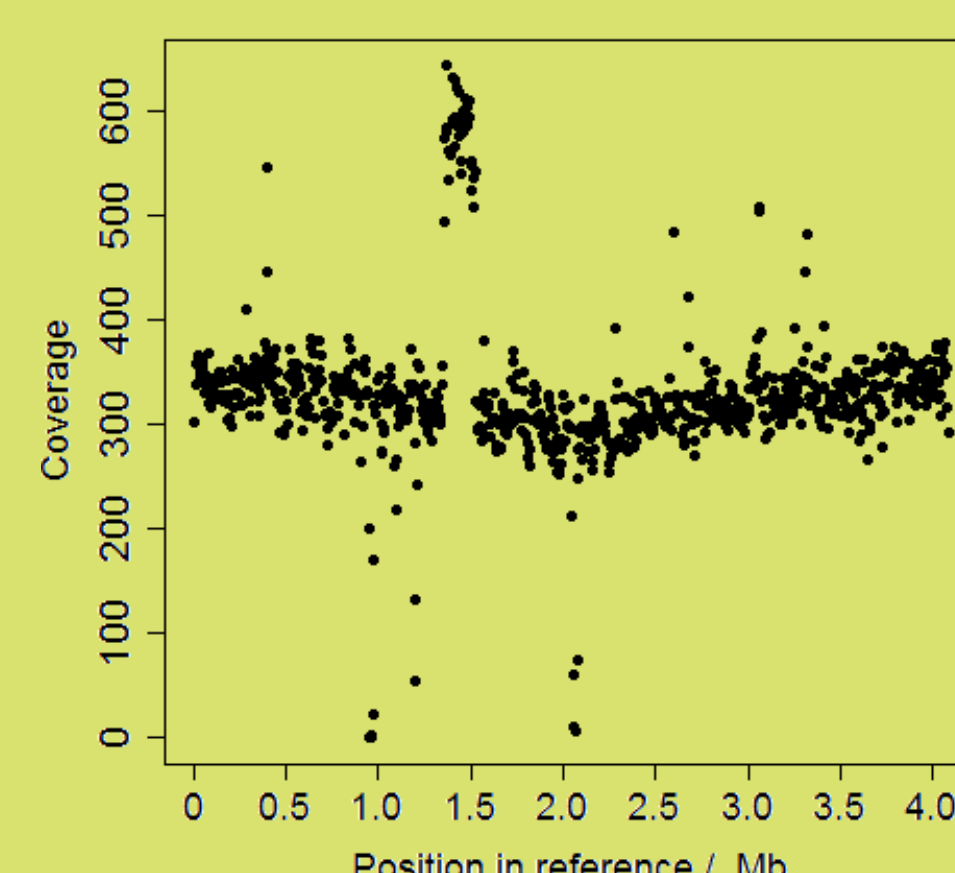
Our “ultra-long” UK48 reads:

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

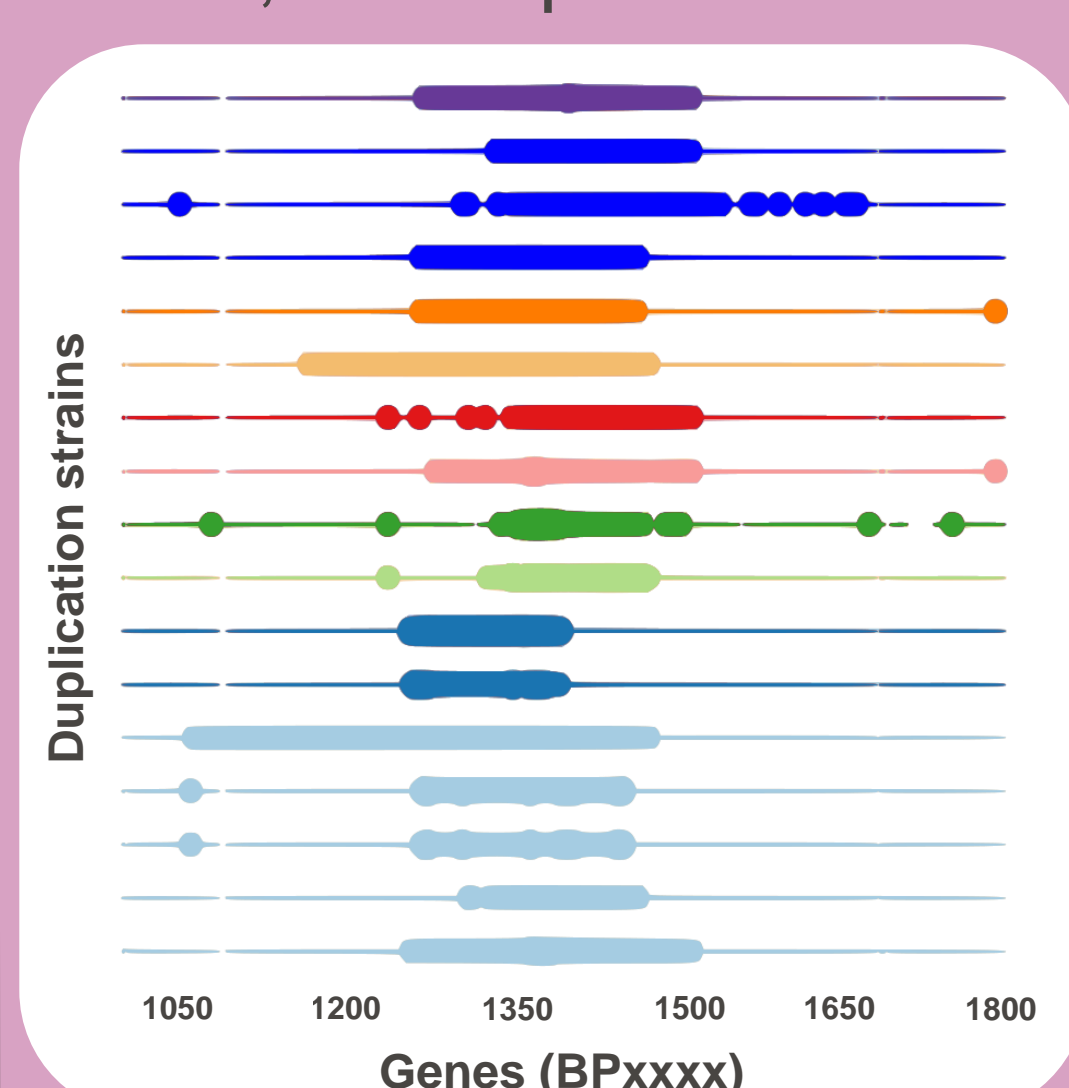


For more about copy number variation in *Bordetella pertussis*, visit the poster of Jonathan Abrahams!

## The “motility” duplication

Over **15 strains** sequenced globally appear to have a **duplication of the same region** which is duplicated in UK48 and UK76 [4]

Although the exact length and genes included in the duplicated region varies slightly from strain to strain, the duplication tends to be **centred around**

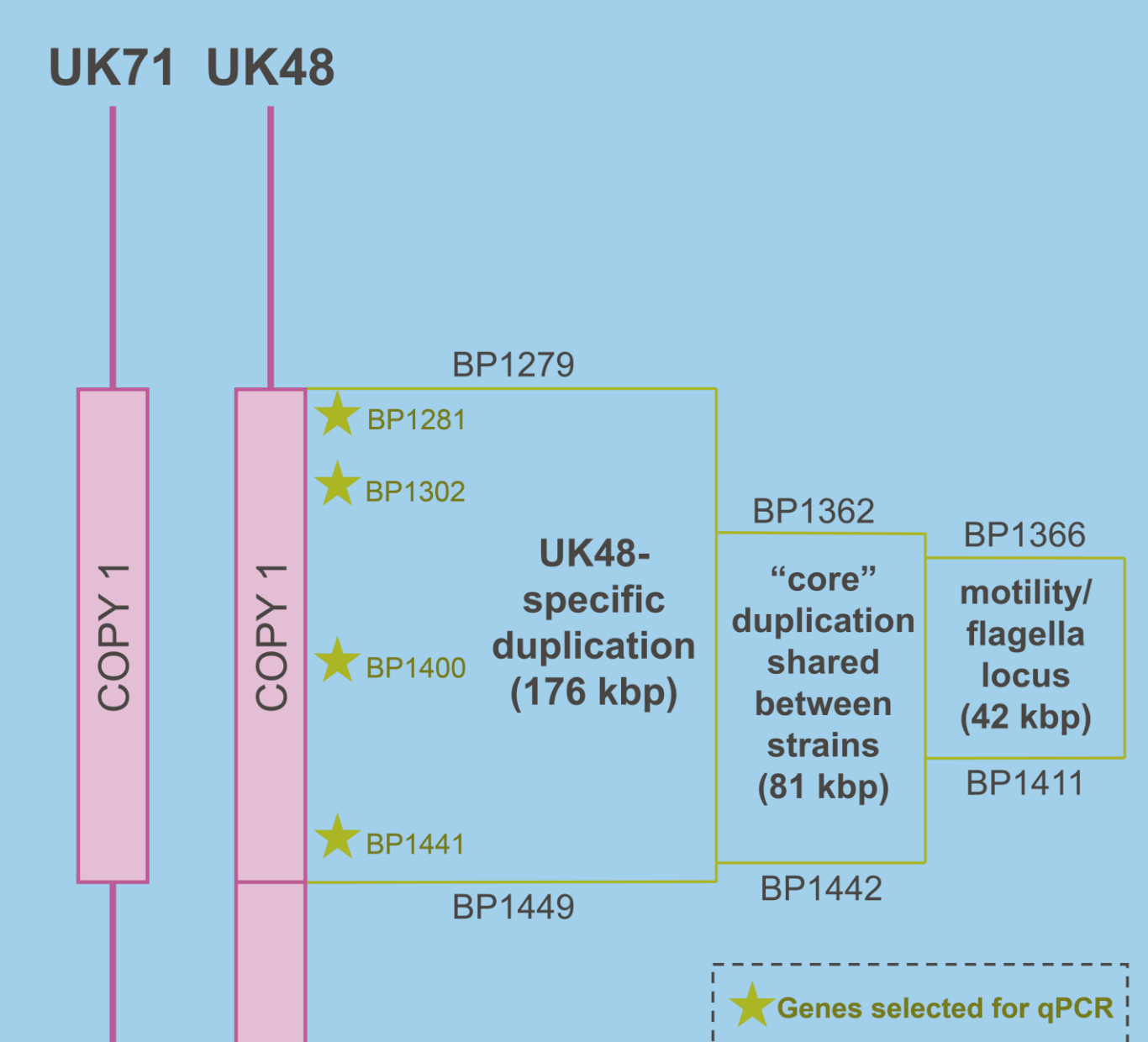


**the same set of “core” genes: BP1334 to BP1492**

42 of the 77 genes in the “core” duplication are related to **motility and/or flagella synthesis**

We are now aiming to characterise this duplication further

## Does the duplication affect gene expression?



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

We are using **qPCR** to compare **gene expression** in **UK48** with gene expression in **UK71**, a strain which is highly similar to UK48 but whose genome **does not contain the duplication**

**Future questions:** Does the duplication affect **growth rate**? Does the duplication affect **motility**?



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

## 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. *Journal of Infectious Diseases* 212(2)
- [3] Quick (2018). Ultra-long read sequencing protocol for Rad004. <https://www.protocols.io/view/ultra-long-read-sequencing-protocol-for-rad004-mrx57n>
- [4] Abrahams et al. (In preparation). Duplications drive diversity in the monomorphic pathogen *Bordetella pertussis* on an underestimated scale.

## Tools

**ABYSS:** <https://github.com/bcgsc/abyss>  
**Albacore:** <https://community.nanoporetech.com/downloads>  
**Canu:** <https://github.com/marbl/canu>  
**Flye:** <https://github.com/fenderglass/Flye>  
**Nanopolish:** <https://github.com/jts/nanopolish>  
**Prokka:** <https://github.com/seemann/prokka>  
**Unicycler:** <https://github.com/rnw/Unicycler>

## Acknowledgements

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## About the author

I am a 3<sup>rd</sup> 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

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