# Pablo González de la Rosa

Bioinformatician | Oxford Nanopore Technologies Website: pgonzale60.github.io • Email: pgonzale60@gmail.com • GitHub: github.com/pgonzale60 • Remote-friendly (Oxford, UK)

#### PROFESSIONAL SUMMARY

Bioinformatics workflow engineer with 8+ years building reproducible genomic analysis pipelines. Proficient in Nextflow (DSL2) with a focus on debugging, module composition, and automation. Experienced with containerisation and GitLab CI/CD in ISO-aligned quality settings. Built the `wf-pgx` research workflow from a template to a usable pipeline (available upon request).

I use Nextflow as the backbone of bioinformatics automation, supported by Python and R for analysis and glue. I also prototype LLM-assisted helpers to streamline text-heavy tasks when appropriate.

#### **CORE COMPETENCIES**

- Nextflow Automation: DSL2, nf-core patterns, schema-driven parameters, channel design, and debugging.
- Reproducible Environments: Docker/OCI, Singularity/Apptainer.
- **Collaboration & Traceability**: GitLab merge requests, code review, changelog management; JIRA ticketing linked to MRs.
- **CI/CD & Quality**: GitLab CI/CD, smoke tests, automated documentation; quality assurance in ISO-aligned settings.
- HPC & Cloud: Slurm; working familiarity with AWS (S3, EC2 basics).
- Analysis & Glue: Python (pandas, NumPy), R (tidyverse, Shiny), Bash; Rust/PHP (familiar).
- Reporting: Coverage metrics (mosdepth), VCF stats, HTML report generation.

#### PROFESSIONAL EXPERIENCE

## Oxford Nanopore Technologies — Bioinformatician

Oxford, UK • Nov 2023 - Present

- Built and maintained Nextflow pipelines for targeted sequencing (schema-driven parameters, containerised modules, CI smoke tests).
- Assembled and evolved the `wf-pgx` research workflow from a barebones template to a usable pipeline (available upon request); added representative datasets and documentation.
- Co-developed team workflows aimed toward regulatory compliance: `wf-vax-qc`, `wf-plasmid-qc`, and `wf-installation-qualification`.
- Built Python utilities for variant calling and analysis across human, cattle, and bacterial datasets; integrated into pipelines and CI.
- Established practices around GitLab CI/CD, JIRA → MR traceability, and code review quality.
- Presented journal club sessions on pipeline debugging and workflow patterns.

## Wellcome Sanger Institute & University of Cambridge — Doctoral Researcher

Cambridge, UK • 2020 – 2023

- Architected workflows for large-scale genomic and metagenomic analyses in evolutionary studies.
- Automated genome assembly quality assessment using orthology and k-mer metrics; integrated into multiple research projects to reduce manual validation.
- Delivered invited and contributed talks at conferences and seminars on genome assembly and chromosome evolution.
- Mentored MSc/PhD students in bioinformatics best practices and open-source tooling; co-authored peer-reviewed publications.

#### Winter Genomics — Bioinformatics Intern

Mexico City, Mexico • Aug 2013 - Nov 2015

- Designed and deployed a MySQL-backed variant database with a web interface to support collaborators' queries and curation workflows.
- Developed R/Bash automation for microarray QC; standardised gene expression analyses and reduced turnaround time
- Contributed to host–pathogen comparative analysis workflows; supported publication-grade datasets and figures.

#### **SELECTED PROJECTS**

- wf-pgx (ONT): Assembled and evolved research-use pharmacogenomics workflow from a template; prepared a public release note of accompanying data and results representative datasets and docs for internal/research users.
- **Biopharma QC Workflows (team support)**: Contributed to workflows aimed toward regulatory compliance: `wf-vax-qc`, `wf-plasmid-qc`, and `wf-installation-qualification`.
- Chromosome-scale Genome Annotation (MSc): Nextflow workflow to annotate multi-megabase animal genomes; integrated gene prediction, repeat masking, and evidence-driven refinement for midscale HPC.
- Assembly Completeness Assessment (PhD): Automated pipeline to assess completeness of newly assembled multi-megabase animal genomes using orthology- and k-mer-based metrics.
- **GEO Differential Expression Automation (BSc thesis)**: Automated microarray-based differential expression across dozens of GEO experiments using GNU Make; downloaded datasets and Brainarray annotations, inferred experimental designs, executed group-vs-group contrasts, and summarised results in a single table.
- **Regulatory Gene Visualisation (BSc training)**: Automated gene visualisation for a gene regulation website using PHP and available regulatory annotations.
- Al-assisted Preference Matching (personal R&D): Prototyped LLM-assisted workflows to match user preferences against free-text (e.g., adverts) to reduce manual screening.

#### Context and outcomes (selected):

- wf-pgx: assembled and evolved the `wf-pgx` research workflow from a barebones template to a usable pipeline.
- Biopharma QC: supported team delivery of QC workflows aligned with regulatory expectations; clearer installation/qualification steps.

- Verification Suite: standardised release checks and metrics diffs; faster validation cycles.
- Al matching: accelerated manual screening tasks using LLM-assisted triage and ranking prototypes.

# **PUBLICATIONS (SELECTED)**

- Stevens, L., et al. (2024). The genome of Litomosoides sigmodontis illuminates the origins of Y chromosomes in filarial nematodes. PLoS Genetics, 20(1), e1011116.
- Stevens, L., Martínez-Ugalde, I., et al. (2023). Ancient diversity in host-parasite interaction genes in a model parasitic nematode. Nature Communications, 14(1), 7776.
- Gonzalez de la Rosa, P. M., et al. (2021). A telomere-to-telomere assembly of Oscheius tipulae and the evolution of rhabditid nematode chromosomes. G3, 11(1), jkaa020.

Full list of publications on Google Scholar

#### **EDUCATION**

• PhD Evolutionary Biology, University of Cambridge & Wellcome Sanger Institute (2020 – 2023)

Chromosome evolution and programmed DNA elimination in Rhabditina nematodes.

- Developed an automated pipeline to assess completeness of newly assembled multi-megabase animal genomes (orthology/k-mer metrics, reproducible runs).
- MSc Integrative Biology, LANGEBIO-CINVESTAV, Mexico (2016 2018)

Host-specific gene expression in monarch butterflies.

- Built a Nextflow workflow to annotate multi-megabase animal genomes (repeat/structural annotation, evidence integration, HPC scheduling).
- BSc Genomic Sciences, Center for Genomic Sciences, UNAM, Mexico (2011 2015)
- Thesis: Automated GEO differential expression pipeline with GNU Make (data/annotation retrieval, design inference, batch contrasts, unified summary table).

## **TECHNICAL TOOLKIT**

- Languages: Python (advanced), R (advanced), Bash (proficient), SQL (proficient); Rust/PHP (familiar)
- **Workflow Tools**: Nextflow DSL2 (nf-core patterns), schema-driven parameters; containers with Docker/Singularity
- CI/CD & Quality: GitLab CI/CD, smoke tests, docs automation; JIRA for planning and MR traceability;
  ISO-aligned quality familiarity
- HPC & Cloud: Slurm; AWS familiarity (S3, EC2 basics)
- Al/LLM Prototyping: Prompting and LLM-assisted extraction for text triage (early-stage prototypes)
- Reporting: matplotlib, Plotly, Shiny, custom HTML reports

#### **REFERENCES**

Available upon request.