Adaptive nanopore sequencing on miniature flow cell detects extensive antimicrobial resistance

Abstract

Rapid screening of hospital admissions to detect asymptomatic carriers of resistant bacteria can prevent pathogen outbreaks. However, the resulting isolates rarely have their genome sequenced due to cost constraints and long turn-around times to get and process the data, limiting their usefulness to the practitioner. Here we use real-time, on-device target enrichment ("adaptive") sequencing on a new type of low-cost nanopore flow cell as a highly multiplexed assay covering 1,147 antimicrobial resistance genes. Using this method, we detected three types of carbapenemase in a single isolate of Raoultella ornithinolytica (NDM, KPC, VIM). Further investigation revealed extensive horizontal gene transfer within the underlying microbial consortium, increasing the risk of resistance spreading. From a technical point of view, we identify two important variables that can increase the enrichment of target genes: higher nucleotide identity and shorter read length. Real-time sequencing could thus quickly inform how to monitor this case and its surroundings.

Keywords Nanopore sequencing · Target enrichment · Antimicrobial Resistance · Carbapenemases · Plasmids

Background

Screening patients for multiresistant bacteria on hospital admission can detect asymptomatic colonization early and reduce subsequent complications. However, corresponding isolates rarely have their genome sequenced, which would enable genomic surveillance, and, as a result, source control and reduced spread. Such resistant strains can colonize patients for years, increasing the value of this information. Long-term carriage is surprising in the absence of a selective stimuli such as treatment with antimicrobials. Recently, the underlying microbial consortia in which these strains are embedded have been implicated in resistance maintenance through ongoing horizontal gene transfer of mobile elements. This finding suggests that in special cases, genomic surveillance should be expanded to include metagenomic data.
Here we report on a patient with multiple carbapenem-resistant strains detected in a rectal swab. One of the isolates simultaneously carried four carbapenemases, an unusually high number. To support a timely response, we integrated the results from multiple modalities of real-time nanopore sequencing. First, we reconstructed the genomes of individual isolates and then complemented them with metagenomic data from the swab. In a proof-of-concept, we then applied real-time on-device target enrichment of 1,147 resistance genes on a miniature flow cell to create an ultra-high multiplex assay.

**Results**

During resistance screening of rectal swabs, we found three bacterial species growing on carbapenem agar (*Raoultella ornithinolytica*, *Citrobacter freundii*, and *Citrobacter amalonaticus*). The patient’s history revealed no apparent source, although past occupations included work in waste management and training in agriculture, both of which have increased exposure to antibiotic resistance genes. Surprisingly, we detected multiple carbapenemases in *R. ornithinolytica* using PCR (*NDM*, *KPC*, *VIM*). To identify all resistance genes in the isolates and any putative horizontal transfer between them, we performed real-time nanopore sequencing, both of the isolates individually and of the entire rectal swab, generating in total 3.9 M reads and 23.3 Gb on a standard ("MinION") flow cell.

The *R. ornithinolytica* isolate carried nine plasmids and three carbapenemases: *NDM-1*, *KPC-2*, and *VIM-1* (Figure 1A). All carbapenemases were encoded on one plasmid each, except *VIM*, which was located on the bacterial chromosome. The two *Citrobacter* isolates only carried *VIM-1*. An alignment of the genomic region 10 Kb upstream and downstream of *VIM* across the isolates revealed a transposase-mediated resistance transfer, for which we propose the following gene flow: The genomes of *C. freundii* and *C. amalonaticus* both carry *VIM-1* on an *IncHI2* plasmid (> 95% sequence identity). In *C. freundii*, this transposon then likely copied itself into an *IncN* plasmid with the help of an *ISKpn19* transposase (Figure 1B). The same transposase is found flanking the *VIM* transposon in the *R. ornithinolytica* chromosome, which makes the *IncN* plasmid of *C. freundii* its likely source. A similar transfer pattern was observed for the penicillinase *OXA-1* (data not shown).

Isolate sequencing captured 79.7% of resistance genes detected in the underlying microbial consortium through metagenomic sequencing (total yield 10 Gb, Figure 1C). Of the remainder, few genes were clinically relevant, such as several efflux pumps. Other resistance genes were associated with Gram-positive bacteria, which we did not screen for with culture (Figure 1C). Surprisingly, metagenomics did not detect five resistance gene types (6.8%), including *KPC*, two out of three *OXA* copies, and two out of four *VIM* copies. This omission likely occurs because the metagenome was dominated by *Proteus vulgaris* (44.6% of reads), leaving fewer reads (depth) for the carbapenemase-carrying strains (*C. freundii* 19.7%, *R. ornithinolytica* 1.8%, *C. amalonaticus* 0.01%). Selective culture enriched these low-abundant species.

We also observed substantial horizontal gene transfer between our isolate members of the *Enterobacteriaceae* (Figure 1D). For example, *C. freundii* and *R. ornithinolytica* share 15 loci. A region was labeled as a putative transfer if its length exceeded one kilobase with 99.9% sequence identity between any two genomes. No additional transfer was found in two uncultured, metagenome-assembled genomes (MAGs), namely *Enterococcus faecium* and *Serratia*
Figure 1: Real-time sequencing reveals extensive resistance load and horizontal gene transfer. (A) Genome reconstruction of a strain of *R. ornithinolytica* carrying nine plasmids and four carbapenemase genes. Color-coded coverage from 90x (black, e.g., chromosome) to 250x (red, e.g., plasmid carrying *OXA-1*). (B) Gene transfer of *VIM-1* across three strains and four loci. The carbapenemase is flanked by multiple transposases (see annotation), which likely mediate its mobilization. Vertical lines indicate 100% sequence identity between corresponding genes. (C) Comparison of shared resistance genes between the enrichment sequencing run ("Flongle"), the *R. ornithinolytica* isolate, all four "isolates" combined and the "metagenome" assembly. Of all resistance genes identified in the metagenome, 79.7% were found in the isolates. Surprisingly, several resistance genes were not identified in the metagenome, among them several carbapenemase copies. In the *R. ornithinolytica* isolate genome, about two-thirds of resistance genes were also found using on-device target enrichment. All plasmid-encoded genes among them were detected, including all carbapenemases. (D) Pairwise shared sequences between isolates and metagenome-assembled genomes. Putative transfers were defined as loci with a minimum length of one kilobase and 99.9% sequence identity between each pair of loci. Extensive sequence transfer is observed between the three isolate genomes (and their corresponding bins from the metagenomic assembly). (E) Miniature, low-cost flow cell used for on-device target enrichment ("Flongle", Oxford Nanopore Technologies), with a one-cent coin placed on top as scale.

*ureilytica*. None of the remaining metagenomic contigs showed putative transfers. Again, metagenomics did not add important information beyond the culture isolates.

The sensitivity of metagenomic sequencing can be increased with depth, but the associated cost limits the applicability in the routine laboratory. Therefore, going in the opposite direction, a recently introduced miniature nanopore flow cell ("Flongle") aims to reduce per-run costs through reduced sequencing yield. Because the yield is reduced, however, targeted sequencing of relevant genes or loci is desirable. Such target enrichment can be performed "on-device", i.e., during the sequencing run in real-time and without any changes in the sample preparation, using a method also known as nanopore "adaptive sequencing" [10,12]. Here, reads are rejected from the pore when the read fragment that already passed through it does not match any sequence in a target database. The nanopore is then free to sequence another molecule.
Adaptive sequencing can be used to enrich or deplete either entire organisms from a sample of DNA or to target specific genes. Here we aimed to enrich 1,147 representative antimicrobial resistance genes (ARGs, see methods), which to our knowledge, is the first time that adaptive sequencing has been used to target a microbial gene panel. We define "enrichment" as the difference in total read count over a corresponding ARG between standard and adaptive sequencing. In the latter condition and unless stated otherwise, we exclude rejected reads, i.e., those for which the adaptive selection algorithm has not recognized a target in the first bases of the read (Figure S1).

Read count over target genes is a commonly used metric in RNA-Seq experiments, with two caveats: First, most RNA-Seq studies use fixed-sized short reads, i.e., the read length distribution is the same for all conditions. Second, the read count is normalized by target length and the overall number of mapped reads, which allows the comparison of targets within a condition ("relative expression"), although their size might differ. Here, we only use raw read count to measure enrichment without further adjustments. We justify this because, first, a single library was used in both conditions (standard, adaptive), and the resulting read length distributions for "standard" and "unrejected adaptive" reads are near-identical (Figure S1). Second, we only compare each target across conditions, and so there is no need to normalize by target length.

To compare target read abundance between adaptive and "standard" nanopore sequencing, we sequenced three isolates in technical duplicates on a single MinION flow cell, periodically alternating between adaptive and standard sequencing in 16 one-hour intervals (Figure 2A). Overall, adaptive sequencing could roughly double the abundance of many targets, while others were hardly detected at all (Figure 2B). We subsequently identified two factors that substantially affected target abundance between the two conditions: nucleotide identity and read length.

First, high nucleotide identity between an isolate’s gene and the corresponding member of the target gene panel resulted in a higher on-target read count (Figure 2B). Surprisingly, most similar and thus most enriched genes were located on plasmids. This likely reflects a bias in the database composition, where common resistance plasmids are well annotated while strain-specific, chromosomal gene isoforms are undersampled. As expected, target sequence similarity did not affect abundance in the standard condition, which did not use a target database. To quantify this effect, we performed Bayesian regression and modeled the effects of variables for which a contribution to abundance seemed plausible, namely sequence similarity, coverage, read length and whether the target was located on a plasmid (including interaction effects, see methods). The largest effect was observed for similarity conditional on whether adaptive sequencing was turned on ($\beta = 11.98, 95\% CI \pm 0.15$). However, it can be hard to interpret any single coefficient in an interaction model in isolation; it is more informative to plot samples from the posterior distribution for any variable of interest. All else being equal, adaptive enrichment only outperforms standard sequencing when an isolate’s gene has a nucleotide identity of at least 95% to a record in the target database, and up to two-fold for near-identical targets (Figure 2C). Several targets are enriched four times over the standard baseline. Other studies reported a similar enrichment of two- to four-fold for bacterial genomes, albeit partly using different real-time matching algorithms.

Second, we observed that reads from plasmids were shorter than chromosomal ones (Figure 2D). Furthermore, the length of chromosomal reads is shorter for adaptive sequencing than the standard because if a target is not identified on a given read, sequencing is terminated prematurely. Our statistical model takes this conditionality into account. The model indicates that adaptive sequencing outperforms the standard only for reads smaller than 3 kb, all else being equal.
This bias contributes to the higher target abundance for plasmid-encoded genes than chromosomal ones within the adaptive sequencing condition.

It seems counterintuitive that short reads are enriched more than longer ones. However, this is due to how adaptive sequencing rejects reads. The algorithm scans the first part of each read for target sequences, and if none is found after several hundred bases, the read is rejected (median 415 bases, Figure S1). To further investigate the relationship between read length, target length, and false-negative rate (FNR), we simulated combinations of, e.g., long reads and short targets and long reads and long targets (see methods). A fixed-length target occupies a smaller fraction in long reads than in shorter ones. We find that the smaller this fraction, the higher the false-negative rate, i.e., the more reads which contain the target are falsely rejected (Figure S3). The intuition behind this result is that a target has many potential starting points on a read. The longer the read and the shorter the target, the more likely the target starts at a position after the read interval that the selection algorithm uses for its decision.

The relative abundance of reads outside of target regions should not change during adaptive sequencing. We checked this by comparing the coverage of "standard" and "rejected adaptive" reads across assembly contigs, which did not differ substantially (Figure S2). Note that we normalize coverage to that of the chromosome for both read sets because the rejected reads from the adaptive condition are much shorter and more numerous (Figure S1, median read length 3,194 vs. 415 bases).

We then tested the enrichment on a new type of miniature, low-cost flow cell and generated 5.4 Mb within four hours from the carbapenemase-rich R. ornithinolytica isolate (Figure 1E). 97.2 % of reads were rejected; of those, 0.2 % (n=43) were false negative. Correspondingly, 2.8 % of reads were accepted, of which 20.4 % (n=104) were true positive, i.e., could be found in the target database. A positive database hit was defined as a read with at least 100 bp mapped to a target with a minimum of 50 % matching positions. 57.9 % of the resistance genes found in the high-quality genome reconstruction were found using adaptive sampling, too, including all four carbapenemases (Figure 2C). As expected from the adaptive-standard state switching experiment, the probability of detection was determined by genomic location: All un-detected genes were located on the chromosome, and all plasmid-encoded resistance genes were detected (odds ratio 26.7, p < 0.001). While plasmids are present in higher copy numbers relative to the chromosome (Figure 1A), our regression model did not assign a large effect to this variable. Since many resistance determinants are located on plasmids, we argue that enrichment sequencing is a promising approach for antimicrobial gene detection in routine settings.

Discussion

We detected a highly resistant consortium during hospital admission screening, including a strain that carried three carbapenemases. Nanopore sequencing comprehensively characterized three resistant culture isolates, documenting many resistance genes as well as extensive gene transfer between isolates. Metagenomic sequencing of the corresponding rectal swap added little information and did not detect several important resistance genes. It might be that deeper sequencing would increase sensitivity. Still, because the carbapenemase-carrying strains were low abundant, in practice, this procedure would not be cost-competitive in a routine setting. Cultural screening as a first step reliably identified the strains that carried clinically relevant resistance genes within 24 hours from sample streaking on screening agar plates to detectable growth. From subsequent sequencing, including library preparation to isolate genome assembly, another 24
Figure 2: Effect of two variables during adaptive sequencing on enrichment efficiency compared to a standard nanopore sequencing run. (A) The same three Citrobacter and Raoultella isolates were sequenced with and without enrichment (green and violet, respectively), alternating the conditions periodically on a single flow cell. (B) Each point corresponds to an open reading frame that has been annotated in the final isolate genome assembly as a resistance gene using a dereplicated ARG database (n=1,147). "Read count" is the number of reads from each sequencing condition that map to these genes. As the read passes through the nanopore during enrichment, it is searched in real-time against a database of target genes. If no match is found, the read is ejected prematurely. Reads that are very similar to a database entry (nucleotide identity) pass this filter, while reads with lower sequence identity are falsely rejected. Many highly similar targets reside on plasmids, likely a sampling bias in the resistance database. As expected, similarity has no effect on read count per ORF for standard sequencing because there is no database search involved. (C) Adaptive sequencing outperforms the standard once the nucleotide identity ("similarity") between the target and its match in the panel database surpasses 95 %. For values close to identity, a two-fold enrichment can be expected. We even observed a four-fold enrichment of several targets over the standard baseline. When we fit a Bayesian multivariate regression model, the increase of target abundance with similarity becomes clear (2.5 and 97.5 % quantiles displayed). (D) Reads derived from plasmids are shorter than chromosomal ones. In turn, chromosomal reads from adaptive are shorter than those from standard sequencing because they are more frequently ejected from the nanopore before the read has been sequenced fully for lack of any match. These factors need to be accounted for in a regression model to estimate the effect of read length on target abundance accurately. (E) Adaptive sequencing outperforms the standard for shorter read lengths of 3 kb and less, all else being equal. Short reads of about 1 kb can potentially double target abundance. Therefore, library preparation protocols for adaptive sequencing could add a step to shear extracted DNA to improve the enrichment.
of the sequencing platform. In addition, more computation is required for the final assembly, binning, and validation of
the mixed sample.

We then evaluated a new approach for on-device, real-time target enrichment called "adaptive sequencing". It en-
compassed 1,147 representative antimicrobial resistance genes in an ultra-high multiplex assay. In the enrichment
sequencing data, reads were detected for all resistance genes known to be present in the isolate assemblies from
an independent, previous sequencing run (see methods). However, while some genes could be enriched up to four
times over the baseline, others were hardly captured. To explain the disparity, we found that two variables influence
enrichment substantially. First, the higher the nucleotide similarity of a read to its corresponding entry in the target
database, the more reads were selected. Adaptive outperformed standard sequencing above 95 % identity. Optimization
of the target database to reflect the expected targets as closely as possible is thus crucial. Future studies will have to
determine the influence of database size and redundancy on target abundance. Second, we showed that fragments shorter
than 3 kb are beneficial to target abundance. Counterintuitively for nanopore sequencing, deliberate shearing of DNA
fragments during library preparation should help to enrich targets. From an economics perspective, the fold-enrichment
is inversely proportional to sequencing cost. So an enrichment by a factor of two would translate into 50 % reduced
sequencing costs (excluding library preparation).

The degree to which DNA should be sheared for an enrichment experiment depends on the underlying choice of target
length. For the enrichment of antimicrobial resistance genes, as demonstrated in this study and with a mean length of
about one kilobase, we recommend matching this with an equal median read length.

We then performed on-device target enrichment of the most resistant culture isolate on a new type of miniature flow cell
and were able to identify all plasmid-encoded resistance genes and nearly two-thirds of all resistance genes known to be
present.

Conclusions

As a proof of concept, we show that on-device target enrichment on low-cost flow cells could be a valuable complement
to routine microbiology. It takes us closer to an effective point-of-care resistance screening, especially given the
continued rapid improvements in the underlying technology. However, given the variable sequencing yield of this new
flow cell type, further controlled experiments that compare multiple runs with and without enrichment are warranted, as
are studies that optimize sample preparation and target database composition.

Methods

Culture and DNA extraction

All samples were streaked on carbapenemase chromogenic agar plates (CHROMagar, Paris, France). Carbapenemase
carriage was confirmed using PCR and phenotypically using microdilution MIC testing. DNA was extracted from
culture isolates and rectal swabs using the ZymoBIOMICS DNA Miniprep extraction kit according to the manufacturer’s
instructions. The cell disruption was conducted three times for five minutes with the Speedmill Plus (Analytik Jena,
Germany).
Library preparation

DNA quantification steps were performed using the dsDNA HS assay for Qubit (Invitrogen, US). DNA was size-selected by cleaning up with 0.45x volume of Ampure XP buffer (Beckman Coulter, Brea, CA, USA) and eluted in 60 l EB buffer (Qiagen, Hilden, Germany). The libraries were prepared from 1.5 g input DNA. For multiple samples we used the SQK-LSK109 kit (Oxford Nanopore Technologies, Oxford, UK) and the Native Barcoding Expansion-Kit (EXP-NBD104), according to the manufacturer’s protocol. For the Flongle run we used the SQK-RBK004 kit from the same manufacturer.

Nanopore sequencing and on-device target enrichment

All DNA was sequenced on the GridION using a FLO-MIN106D (MinION) and FLO-FGL001 (Flongle) flow cells (MinKNOW software v4.1.2), all from Oxford Nanopore Technologies. Three sequencing runs were performed: For the first run (MinION flow cell) we multiplexed three culture isolates (A2, B1, B2) and a metagenomic sample (3.9 M reads and 23.3 Gb in 48 h, about 10 Gb were metagenomic). The second run (MinION flow cell) was an experiment comparing "adaptive" and "standard" sequencing. On a single flow cell, we periodically alternated between both states by manually turning the sequencing run off and then back on in the other state in one-hour intervals for a total of 16 hours. Toggling between states did not harm sequencing (e.g., through pore blockages; 3.44 M reads and 4.47 Gb in 16 hours). The sequencing yield for all barcodes from three isolates (two Citrobacter and one Raoultella) with two technical replicates each was about equal. Adaptive sequencing groups the sequence data into "rejected", i.e., reads that do not contain a target, and "unrejected". The latter comprises reads with a target found and reads with a pending decision. We used the unrejected reads pooled across isolates and replicates for all further analyses unless stated otherwise. Pooled and per-isolate results did not differ substantially (compare Figure 2B and S4). In the third sequencing run (Flongle flow cell), only isolate A2 was included, and adaptive sampling was applied throughout (18,646 reads and 5.36 Mb in 4h).

As target database, we created a dereplicated version of the CARD database of resistance genes (v3.1.3) using mmseqs2 easy-cluster (v13.45111) using a minimum sequence identity of 0.95 and minimum coverage of 0.8 in coverage mode 1. We thereby reduced the database from 2,979 to 1,147 representative genes. We performed this step to reduce the search space that the adaptive sequencing algorithm has to map against. The total length of all genes in the database was 1.16 Mb. The reduction halves the database size because many resistance genes such as CTX have many documented isoforms, which would lead to uninformative multi-mappings. Reads were basecalled using the guppy GPU basecaller (high accuracy model, v4.2.2, Oxford Nanopore Technologies). For isolate genomes, reads were assigned to their respective barcodes only if matching adapters were detected on both ends of the read to avoid cross-contamination.

For the experiment comparing "adaptive" and "standard" sequencing on a single flow cell, we periodically alternated between both states by manually turning the sequencing run off and then back on in the other state in one-hour intervals for a total of 16 hours. This protocol did not have a negative effect on total sequencing yield over time (e.g., through pore blockages). The sequencing yield for all barcodes from three isolates (two Citrobacter and one Raoultella) with two technical replicates each was about equal. Adaptive sequencing groups the sequence data into "rejected", i.e., reads
that do not contain a target, and "unrejected". The latter comprises reads where a target has been found and reads with a pending decision. We used the unrejected read fraction for all further analyses.

Data analysis

Isolate data were assembled using flye (v2.9)\cite{flye} and consensus sequences corrected using three rounds of polishing with racon (v1.4.3)\cite{racon} followed by medaka (v1.4.3, unpublished, github.com/nanoporetech/medaka). Read mapping was performed using minimap2 (v2.22-r1101)\cite{minimap2}. Genome quality was confirmed using checkm (v1.1.3)\cite{checkm}. All isolate genome assemblies were > 99\% complete and < 1\% "contaminated" (duplicate single-copy marker genes), which counts as high quality by community standards. Resistance gene annotation was performed using abricate (v1.0.1, unpublished, github.com/tseemann/abricate) against the CARD database (see above). Taxonomic assignments were performed using single-copy marker genes\cite{msas} as well as k-mers using sourmash (v4.2)\cite{sourmash}.

For the long read-only metagenomic assembly we used flye with the --meta flag. We then mapped all reads to the assembly using minimap2. We then used racon to perform three rounds of long read-only polishing of the assembly using the alignment. Last, we used medaka to generate the final consensus assembly. Binning and annotation were then performed as described elsewhere\cite{medaka} by feeding the consensus assembly into the corresponding workflow modules using default settings. Pairwise similarity between genes was calculated using mmseqs2 easy-search (v13.45111)\cite{mmseqs2}. The amount of putative horizontal gene transfer between isolate genomes and MAGs was estimated by counting the number of shared genes for each pair. First, we performed pairwise genome alignment using the nucmer command from mummer (v4.0.0rc1)\cite{mummer}. We then searched for "shared genes", defined as such if the alignment was 1 kb or longer and if the pairwise nucleotide identity between genes was > 99.9\%.

To model the influence of several predictor variables on our outcome variable "target abundance" (total on-target read count), including plausible interactions, we fit a Bayesian regression model using brms (v2.13)\cite{brms}. The outcome variable was modeled as a Poisson distribution. We conditioned the effect of nucleotide similarity on sequencing state, i.e., whether adaptive sequencing was turned on or off, by introducing an interaction term. Also, we conditioned read length on sequencing state and whether a read was derived from a plasmid or the chromosome. Finally, we included a term to model the effect of contig coverage, calculated from mapping the reads back to the isolate assemblies.

\[
read\_count \sim \text{Normal}(\mu_i, \sigma)\\
\mu_i = \alpha + \gamma_i\text{adaptive}_i + \delta_i\text{plasmid}_i + \delta_i\text{adaptive}_i + \beta_i\text{coverage}\\
\gamma_i = \beta_2 + \beta_3\text{similarity}\\
\delta_i = \beta_4 + \beta_5\log(\text{read\_length})
\]

Sampling was performed with four chains, each with 2,000 iterations, of which the first half were discarded as warmup, for a total of 4,000 post-warmup samples. Samples were drawn using the NUTS algorithm. All chains converged ($\hat{R} = 1.00$). Coefficient estimates including confidence intervals are available in Table S1. Refer to the project repository for code on how the model was specified.
For the simulation experiment, we first determined that a log-normal distribution can adequately model the different length distributions (Figure S1). We then selected parameters to model combinations of reads and targets of varying length distributions, from long (mean 8,103 bases, or log 9, variance 1.5) to short (mean 665 bases, or log 6.5, variance 0.25). We could thus, for example, assess the effects of combining long reads with short targets and short reads with long targets. Next, we generated ten thousand (read, target) sample pairs for each combination of read and target distributions. For each pair, we randomly "placed" a target start on the read with a uniform distribution across read positions, in line with how targets are distributed across DNA fragments in realistic single-molecule sequencing experiments. We then asked if the first part of the read (as seen by the adaptive sampling algorithm) would have detected the target. Since all simulated reads contain a target, failure to detect one in the first number of bases counts as a false negative (Figure S3). To estimate effect sizes on the false-negative rate (FNR) we fit a multivariate regression model using \texttt{brms} (see above) (Figure S3):

\begin{equation}
\text{FNR} \sim \text{Normal}(\mu_i, \sigma)
\end{equation}

\begin{equation}
\mu_i = \alpha + \beta_1 \log(\text{read\_length}) + \beta_2 \log(\text{target\_length})
\end{equation}

**List of abbreviations**

ARG .. antimicrobial resistance gene, ORF .. open reading frame, MAG .. metagenome-assembled genome, FNR .. false negative rate

**Ethics approval and consent to participate**

Not applicable; only microbial samples were used, which are not subject to ethical approval. Human DNA sequences were removed from the metagenomic stool dataset before analysis by filtering them against the recently published complete human reference genome "CHM13."

**Consent for publication**

Not applicable. The manuscript includes no specific details, images or videos relating to an individual person.

**Availability of data and materials**

All basecalled nanopore sequencing data has been deposited with the SRA, NCBI. Metagenomic reads are available under project ID PRJNA788147. Reads from isolate genomes, the Flongle flow cell and the experiment alternating between adaptive and standard state have been deposited under project ID PRJNA788148 under their respective sample ID (Raoultella ornithinolytica: SAMN23928631, Citrobacter freundii: SAMN23928632, Citrobacter amalonaticus: SAMN23928633).

Beyond standard analyses described in the methods, code and referenced assemblies (isolates, metagenome) for the following analyses are available from a dedicated code repository at [github.com/phiweger/adaptive](https://github.com/phiweger/adaptive) Putative horizontal
gene transfer, creation of the target database for adaptive sequencing, analysis of the experiment that switched adaptive sequencing on and off and how to specify a Bayesian regression model of the target count.

**Competing interests**

AV has received travel expenses to speak at Oxford Nanopore meetings. AV, CB and MH are co-founders of nanozoo GmbH and hold shares in the company.

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None received.

**Authors’ contributions**

AV designed the study. AV and ND collected and characterized all samples. MM and CB extracted DNA, prepared Nanopore libraries, and performed all sequencing experiments. AV, CB, and MH analyzed the data. All authors revised the manuscript critically and approved the article’s final version for publication. MWP and CB supervised the study.

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