

1 **Methods for the targeted sequencing and analysis of integrons and their gene cassettes**  
2 **from complex microbial communities**

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18 antibiotic resistance

19 **Abstract**

20

21 Integrons are bacterial genetic elements that can integrate mobile gene cassettes. They are  
22 mostly known for spreading antibiotic resistance cassettes among human pathogens.

23 However, beyond clinical settings, gene cassettes encode an extraordinarily diverse range of  
24 functions important for bacterial adaptation. The recovery and sequencing of cassettes has

25 promising applications, including: surveillance of clinically important genes, particularly

26 antibiotic resistance determinants; investigating the functional diversity of integron-carrying

27 bacteria; and novel enzyme discovery. Although gene cassettes can be directly recovered

28 using PCR, there are no standardised methods for their amplification and, importantly, for

29 validating sequences as genuine integron gene cassettes. Here, we present reproducible

30 methods for the PCR amplification, sequence processing, and validation of gene cassette

31 amplicons from complex communities. We describe two different PCR assays that either

32 amplify cassettes together with integron integrases, or gene cassettes together within cassette

33 arrays. We compare the use of Nanopore and Illumina sequencing, and present bioinformatic

34 pipelines that filter sequences to ensure that they represent amplicons from genuine integrons.

35 Using a diverse set of environmental DNAs, we show that our approach can consistently

36 recover thousands of unique cassettes per sample and up to hundreds of different integron

37 integrases. Recovered cassettes confer a wide range of functions, including antibiotic

38 resistance, with as many as 300 resistance cassettes found in a single sample. In particular,

39 we show that class 1 integrons appear to be collecting and concentrating antibiotic resistance

40 genes out of the broader diversity of cassette functions. The methods described here can be

41 applied to any environmental or clinical microbiome sample.

42

43

44 **Introduction**

45

46 Integrons are bacterial genetic elements that can capture, mobilise and rearrange gene  
47 cassettes [1, 2]. They are mostly known for spreading a diverse repertoire of gene cassettes  
48 that collectively confer resistance to almost all classes of antibiotics [3]. Beyond clinical  
49 settings, however, integrons play a crucial role in bacterial evolution by rapidly generating  
50 genomic diversity [4, 5]. Functional integrons are characterised by their flagship gene, the  
51 integron integrase (*intI*), which encodes a site-specific tyrosine recombinase (IntI). IntI  
52 mediates the insertion of gene cassettes at the integron recombination site (*attI*), which acts as  
53 the insertion site of captured gene cassettes [6]. Gene cassettes, prior to their insertion, are  
54 circular molecules, which possess a cassette recombination site (*attC*). Their insertion  
55 involves IntI-mediated recombination between the *attI* site of the integron and the *attC* site of  
56 the cassette [7-11]. Multiple cassettes can be inserted to form a linear cassette array, which  
57 can vary in size from zero or one cassette to hundreds [12, 13]. IntI activity is induced by  
58 DNA damage, often triggered by environmental stress [14, 15]. Integrons can therefore  
59 provide genomic diversity at precisely the moment when it is needed the most, thus  
60 facilitating ‘adaptation on demand’ [16].

61 Recovery and sequence analysis of integron gene cassettes serve several purposes.

62 First, screening gene cassettes can provide a direct method for surveillance of resistance  
63 genes that are prevalent in an environment of interest. It has also been proposed that  
64 surveying gene cassettes can help detect novel functions that might be harmful to human  
65 health, such as increased pathogenicity or resistance to novel antibiotics [17]. In particular,  
66 class 1 integrons, due to their mobility, abundance and distribution [18, 19], are primed to  
67 play a crucial role in dissemination of these genes. Finally, exploring gene cassettes provides  
68 a window into the functional diversity of the bacterial pangenome. Gene cassettes have been

69 found to be extraordinarily abundant and diverse in every environment surveyed [20-26].  
70 Further, many cassettes with known functions act as single-gene/single-trait entities. As such,  
71 they need minimal integration into metabolic networks and can likely function in a relatively  
72 wide range of genomic contexts. These traits make them highly valuable commodities for  
73 synthetic biology and biotechnological applications, particularly for the discovery of diverse  
74 enzymatic activities [17].

75 Currently, gene cassettes can be recovered from genome sequencing of cultured  
76 isolates, whole metagenomic sequencing, or amplicon sequencing of *attC*-associated genes.  
77 Since most bacteria are yet to be cultured, cassettes identified from isolate genomes  
78 inevitably reflect only a small proportion of all gene cassettes, exacerbated by the fact that  
79 different strains of the same species can vary widely in cassette content [4]. Whole  
80 metagenomic sequencing, although potentially a less biased approach, can be challenging, as  
81 gene cassettes often exist at very low abundances and can contain repeat sequences. A  
82 targeted amplicon sequencing approach, however, can overcome these issues and could  
83 provide the most efficient method for recovering diverse gene cassettes from complex  
84 microbial communities [27].

85 Cassette-targeted amplicon sequencing has been used previously, with varying returns  
86 in gene cassette recovery [20-27]. As sequencing technologies have improved, the ability to  
87 capture a greater diversity of gene cassettes has also increased [20, 26]. However, such  
88 studies lack standardised methods for amplifying and, importantly, validating amplicon  
89 sequences as part of genuine cassettes arrays. Here, we present standardised and reproducible  
90 methods for amplifying, sequencing, and bioinformatic filtering of genuine gene cassettes  
91 from mixed microbial communities.

92 We applied two different PCR assays using DNA isolated from diverse environmental  
93 samples with the aim of recovering integron integrases and gene cassettes. PCR products

94 were sequenced with both long-read Nanopore (ONT) and short-read Illumina MiSeq  
95 sequencing technologies. Importantly, we present bioinformatic pipelines that filter  
96 sequences for complete *attC* sites or *intI* genes. We show that after filtering, we can  
97 consistently recover thousands of gene cassettes from a single sample. We find that recovered  
98 genes display a diverse suite of functional traits, including antibiotic resistance.

99

100 **Methods**

101

102 *Sample collection and DNA extraction*

103 Duplicate samples were collected from six different sites (3 x terrestrial and 3 x aquatic  
104 environments). Terrestrial sites consisted of urban parkland soil from Macquarie University  
105 (Sydney, New South Wales, Australia) [20], hot desert soil from Sturt National Park (North-  
106 western New South Wales) [28, 29], and Antarctic soil from Herring Island [20]. The aquatic  
107 sites consisted of river sediment (Lane Cove River, New South Wales), freshwater biofilms  
108 (Mars Creek, New South Wales) [30], and estuarine sediment (Paramatta River Estuary, New  
109 South Wales). From each of the 12 samples, DNA was extracted from 0.3 g of material using  
110 a standard bead-beating protocol [31]. Each resulting DNA sample was used as the template  
111 for two different PCR assays, described below, and all were subsequently sequenced using  
112 long-read Nanopore (ONT) and short-read MiSeq technologies (Fig. 1).

113

114 *PCR amplification and DNA sequencing*

115 For each sample, we conducted two different PCR assays (Fig. 1A). The first used the  
116 primers HS287 and HS286 [27], which target *attC* recombination sites in opposing directions  
117 to amplify intervening gene cassettes. The second primer set, intI-R / HS286, amplifies  
118 approximately 800bp of the integron integrase gene as well as adjacent gene cassettes. The

119 primer intI-R (5'- GCG AAC GAR TGB CGV AGV GTG TG -3') was designed to target  
120 diverse integron integrases and was based on an alignment of 174 complete *intI* sequences  
121 containing a functional catalytic site, as compiled by Cambray et al. [14]. Importantly, the  
122 last 6 bp of the 3' end of intI-R exactly matches 75% of aligned *intI* sequences. For  
123 amplification, we used Phusion Hot Start II DNA Polymerase (ThermoFisher Scientific,  
124 Waltham, MA, USA), which is a long-range DNA polymerase, chosen to facilitate the  
125 amplification of large segments of integron cassette arrays, known to reach more than 100  
126 kilobases in length [12]. The PCRs were carried out in 50 µL volumes containing a final  
127 concentration of 1 x GC Phusion Buffer, 0.2 mM dNTPs, 0.5 µM of each primer, 3% DMSO  
128 and 2 U of Phusion DNA polymerase. All PCRs were performed using GeneReleaser®  
129 (Bioventures, Murfreesboro, TN, USA) as previously described [32]. Triplicate PCRs were  
130 performed for each sample to increase the chances of capturing rare gene cassettes that might  
131 otherwise escape amplification due to the stochastic nature of PCR.

132 For the HS287 / HS286 primer set, the following thermal cycling program was used:  
133 98 °C for 3 min for 1 cycle; 98 °C for 10 s, 60 °C for 30 s, 72 °C for 3 min 30 s for 35 cycles;  
134 and a final extension step at 72 °C for 10 min. For the intI-R / HS286 primer set, the  
135 following thermal cycling program was used: 98 °C for 3 min for 1 cycle; 98 °C for 10 s, 65 °C  
136 for 30 s, 72 °C for 3 min 30 s for 35 cycles; and a final extension step at 72 °C for 10 min.  
137 PCR efficiency was assessed using 2% agarose gel electrophoresis. Triplicate PCRs were  
138 pooled and then purified with AMPure XP beads (Beckman Coulter, Danvers, MA, USA)  
139 according to the manufacturer's protocol.

140 For long-read sequencing, the 24 PCR products (representing the 12 samples  
141 amplified with each primer set) were multiplexed in a single DNA library using the ONT  
142 Ligation Sequencing Kit (SQK-LSK109) and the ONT Native Barcoding Expansion Kits  
143 (EXP-NBD104 and EXP-NBD114) according to the manufacturer's protocol. The DNA

144 library was sequenced using a MinION MK 1B sequencing device on an R10.3 flow cell.  
145 Sequencing was allowed to run for 24 hours. Basecalling was carried out with Guppy v.4.3.4  
146 with default parameters using the high accuracy (HAC) basecalling model.

147 For short-read sequencing, the 24 PCR products underwent an Illumina DNA shotgun  
148 library preparation using the Nextera XT protocol and then sequenced with MiSeq 300 bp  
149 paired-end sequencing on a single lane. Illumina sequencing and library preparation were  
150 carried out at the Australian Genome Research Facility (Melbourne, Australia).

151

152 *Sequence processing and attC filtering: HS287 / HS286 PCRs*

153 To compare short- and long-read sequencing technologies, we sequenced HS287 / HS286  
154 PCRs on both Nanopore (ONT) and MiSeq platforms. The respective workflows and  
155 software used for sequence processing and filtering are summarised in Figure 1B.

156 For ONT sequences of the HS287 / HS286 PCRs, we first filtered reads based on  
157 average quality (q) scores. We removed any reads with an average q score below 10 using  
158 NanoFilt v2.8 [33] [parameters: -q 10]. Although each read spans the length of an entire  
159 amplicon, many amplicons represent overlapping subsections of larger potential templates.  
160 Thus, an assembly of these initial reads into larger cassette arrays was carried out using Canu  
161 v2.0 [34] [parameters: genomeSize=5m minReadLength=250 minOverlapLength=200  
162 corMinCoverage=0 corOutCoverage=20000 corMhapSensitivity=high  
163 maxInputCoverage=20000 batMemory=125 redMemory=32 oeaMemory=32 batThreads=24  
164 purgeOverlaps=aggressive]. Assembled contigs and unassembled reads were then extracted  
165 together using the tgStoreDump script within Canu [parameters: -consensus -fasta]. Any  
166 redundancies were removed using dedupe.sh, available from the BBTools package v35 [35]  
167 with default parameters. Consensus sequences were then corrected with 4 rounds of polishing  
168 using Racon v1.4.20 [36]. Each round of Racon polishing involved read mapping with

169 minimap2 v2.20-r1061 [37] [parameters: -x map-ont -t 24] and error correction with Racon  
170 [parameters: -m 8 -x 6 -g -8 -w 500 -t 24].

171 For MiSeq sequence data of the HS287 / HS286 PCRs, paired-end reads first  
172 underwent quality trimming and adapter clipping using Trimmomatic v0.38 [38] [parameters:  
173 -phred33 ILLUMINACLIP:adapters.fa:2:30:10 LEADING:3 TRAILING:3  
174 SLIDINGWINDOW:4:15 MINLEN:30] where ‘adapters.fa’ is a fasta-formatted file  
175 containing all commonly used Illumina adapter sequences. If only one end of paired reads  
176 had acceptable quality, it was used as a single read during assembly. Surviving paired-end  
177 reads and single reads were assembled together using SPAdes v3.14.1 [39-41] [parameters -k  
178 21,33,55,77,99,127 --only-assembler --careful].

179 The resulting ONT and MiSeq sequences were both filtered based on the presence of  
180 internal cassette recombination sites (*attCs*). Given the degenerate nature of the primers, off-  
181 target amplicons might constitute a significant portion of the reads. Filtering for sequences  
182 that have internal *attC* sites is thus an essential step when analysing cassette amplicon data. It  
183 should be noted that filtering for *attCs* in this way may discard some genuine amplicons  
184 which consist of single gene cassettes, since they do not possess a complete *attC* site (Fig.  
185 1A). Nevertheless, we consider that for obtaining meaningful ecological data, the removal of  
186 potential false positives is more important than the loss of some true positives. To filter for  
187 *attC* sites, we used an in-house script, attC-screening.sh (available:  
188 <https://github.com/timghaly/integron-filtering>), with default parameters. The script uses the  
189 HattCI [42] + Infernal [43] pipeline that has been previously described [44, 45]. In short,  
190 attC-screening.sh searches for the sequence and secondary structures conserved among *attCs*  
191 and retains any input sequence that has at least one *attC* site. The script can be used on data  
192 generated from any sequencing technology.

193

194 *intI-R / HS286 PCRs: sequence processing and IntI filtering*

195 All intI-R / HS286 PCRs were also sequenced on both ONT and MiSeq platforms (Fig. 1B).

196 For Nanopore sequencing, basecalled reads were first quality filtered using NanoFilt v2.8

197 [33] [parameters: -q 10]. Reads representing concatemers and chimeras were removed using

198 yacrd v0.6.2 [46] with default parameters for ONT data. Since all amplicons should be

199 anchored on one end to the *intI* gene, an assembly would not be suitable. Instead, we

200 clustered reads into amplicon ‘types’ using isONclust v0.0.6.1 [47]. Each cluster was then

201 individually corrected using isONcorrect v0.0.8 [48] with default parameters. Unlike error-

202 correction of genomic data, isONcorrect takes into account uneven coverage within the same

203 read as well as structural variation among similar reads from different clusters (e.g., reads

204 that represent true biological rearrangements of the same gene cassettes). From each

205 corrected cluster, a consensus sequence was then generated using spoa v4.0.7 [36]

206 [parameter: -r 0]. Any redundancies were removed using the BBTools v35 [35] script

207 dedupe.sh with default parameters.

208 MiSeq sequences were processed in the same manner as described above for the

209 HS287 / HS286 data. This involved quality trimming and adapter clipping using

210 Trimmomatic v0.38 [38], followed by an assembly of the reads using SPAdes v3.14.1 [39-

211 41].

212 The resulting ONT and MiSeq sequences were both filtered based on the presence of

213 IntI protein sequences. To detect sequences that encoded IntI, we used an in-house script,

214 intI-screening.sh (available: <https://github.com/timghaly/integron-filtering>), with default

215 parameters. The script uses a profile hidden Markov model (HMM) provided by Cury et al.

216 [13] to detect the additional domain that is unique to integron integrases, separating them

217 from other tyrosine recombinases [49]. The intI-screening.sh pipeline first uses Prodigal [50]

218 to predict all encoded protein sequences, and then screens them for the IntI-specific domain

219 using hmmsearch from the HMMER v3 software package [51]. Any sequences that do not  
220 contain a recognisable integron integrase are discarded. Similarly, intI-screening.sh can be  
221 used on data generated from any sequencing technology.

222

223 *Protein prediction and functional classification of gene cassettes*

224 Cassette open reading frames (ORFs) and their translated protein sequences were predicted  
225 using Prodigal v2.6.3 [50] in metagenomic mode [parameters: -p meta].

226 To assess the broad-scale functional diversity of gene cassettes, we used the Clusters  
227 of Orthologs Groups (COGs) database [52]. COG functions were assigned to cassette-  
228 encoded protein sequences using eggNOG-mapper v2.0.1b [53, 54] executed in DIAMOND  
229 [55] mode with default parameters. To detect cassette-encoded antimicrobial resistance genes  
230 (ARGs), we used ABRicate v0.8 [56] to search against the Comprehensive Antibiotic  
231 Resistance Database (CARD) [57] [Downloaded: 2021-Apr-21].

232

233 *Taxonomic classification of attC sites*

234 The gene cassettes of sedentary chromosomal integrons (SCIs) generally possess highly  
235 similar *attC* sites, and this conservation spans the SCIs of different bacteria within the same  
236 taxon [4, 58, 59]. We have recently modelled the conserved sequence and structure of *attC*  
237 sites from the chromosomal integrons of 11 bacterial taxa [44]. These included six  
238 Gammaproteobacterial orders (Alteromonadales, Methylococcales, Oceanospirillales,  
239 Pseudomonadales, Vibrionales, Xanthomonadales) and an additional five phyla  
240 (Acidobacteria, Cyanobacteria, Deltaproteobacteria, Planctomycetes, Spirochaetes). A  
241 covariance model (CM) was generated separately for each taxon, and this can be used to  
242 correctly identify the source taxon of *attC* sites with high specificity (98-100%) [44].

243 Here, we used an in-house script, attC-taxa.sh (available:  
244 <https://github.com/timghaly/attC-taxa>), with default parameters to detect any *attC* sites that  
245 have originated in the SCIs of one of the 11 taxa. The attC-taxa.sh pipeline incorporates all  
246 11 CMs and uses cmsearch [parameters: --notrunc --max] from the Infernal software package  
247 [43] to classify *attCs*. It is important to note that each taxon-specific model exhibits different  
248 sensitivities in detecting true positives and thus the relative proportion of different taxa  
249 cannot be compared within the same sample. However, the relative proportion of the same  
250 taxon can be compared between different samples.

251

#### 252 *ONT – MiSeq comparisons*

253 For comparisons of the cassette and integrase diversity recovered between ONT and MiSeq  
254 technologies, we first considered differences in sequencing depth. To do this, we randomly  
255 subsampled 50 Mb of raw reads from each sample using rasusa v0.3.0 [60] [parameters: --  
256 coverage 50 --genome-size 1Mb]. After subsampling, all sequence processing and filtering  
257 steps were repeated as described above.

258 All formal comparisons were made using two-sample T-tests (or Wilcoxon rank sum  
259 tests if the data were not normally distributed) using the rstatix v0.7.0 R package [61]. To  
260 determine if the data were normally distributed, Shapiro-Wilk tests were carried out using  
261 rstatix v0.7.0 [61], as well as visually inspected the distributions against their theoretical  
262 normal distributions using Q-Q plots generated with the R package ggpunr v0.4.0 [62].

263 To assess the overlap in recovered ORFs between ONT and MiSeq, we mapped the  
264 cassette ORFs from one technology to the reads of the other using minimap2 v2.20-r1061  
265 [37]. We considered the ORF to be present if it had a mean coverage depth of at least 1x that  
266 spanned at least 98% of the ORF. For ONT and MiSeq read mapping, we used the minimap2  
267 presets [-ax map-ont -t 8] and [-ax sr -t 8], respectively. Coverage statistics were extracted

268 from the mapping alignments using the ‘sort’ and ‘coverage’ programs within the SAMtools  
269 software package [63, 64].

270

271 **Results and Discussion**

272

273 Here, we present a stringent pipeline for PCR amplifying, sequencing and analysing integron  
274 integrases and gene cassettes from diverse microbial communities (Fig. 1). For this, we used  
275 two different PCR primer sets, HS287 / HS286 and intI-R / HS286 (Fig. 1A). The sample  
276 types consisted of a wide variety of soils (from an urban parkland, an Australian desert, and  
277 an Antarctic island), as well as river and estuarine sediments, and freshwater biofilms.

278 To assess the suitability of long- and short-read sequencing technologies, we  
279 sequenced amplicons from both PCR assays using Nanopore (ONT) and Illumina MiSeq  
280 platforms, respectively. Average ONT yield was 181 Mb (100 – 358 Mb per sample) for the  
281 HS287 / HS286 primer set and 216 Mb (62 – 502 Mb per sample) for the intI-R / HS286  
282 primer set. The average MiSeq yield was 418 Mb (228 – 720 Mb per sample) and 663 Mb  
283 (275 – 1,247 Mb per sample), respectively for these primer sets.

284 To ensure amplicons were part of genuine integrons, we filtered the HS287 / HS286  
285 data for *attC* sites, and the intI-R / HS286 data for IntI protein sequences (Fig. 1B). For the  
286 HS287 / HS286 data, an average of 23.8% and 19.0% of amplicon sequences were retained  
287 after filtering for ONT and MiSeq, respectively (Supplementary Fig. S1A). While, for the  
288 intI-R / HS286 data, an average of 1.2% and 1.5% of sequences remained after filtering for  
289 ONT and MiSeq, respectively (Supplementary Fig. S1B). The difference in proportions of  
290 sequences retained after filtering between ONT and MiSeq were not statistically significant  
291 for either primer set. The low proportion of surviving sequences for the intI-R / HS286 data is  
292 likely a result of the intI-R primer binding to other tyrosine recombinases. While many

293 sequences were filtered out, the data retained from this primer set, as described below,  
294 include a large, diverse set of both known and entirely novel integron integrases and gene  
295 cassettes.

296 The lengths of the recovered sequences for both primer sets were significantly larger  
297 for ONT sequencing compared to MiSeq (Supplementary Fig. S2). For the HS287 / HS286  
298 set, sequence lengths ranged from 500 bp to more than 25,304 bp for ONT, and 500 bp to  
299 19,244 bp for MiSeq. For the intI-R / HS286 data, sequence lengths ranged from 803bp to  
300 16,179bp for ONT and 800bp to 7,432bp for MiSeq.

301

302 *Recovered diversity of gene cassette ORFs*

303 We assessed the efficiency of both primer sets in recovering gene cassette open reading  
304 frames (ORFs). Among all 12 samples, the HS287 / HS286 primers amplified 33,854 and  
305 62,118 non-redundant cassette-encoded proteins when sequenced with ONT and MiSeq,  
306 respectively (Fig. 2A). After adjusting for sequencing depth, there was no significant  
307 difference in cassette recovery between the two sequencing technologies (Fig. 2B). On  
308 average, we observed that ~50% of cassette ORFs sequenced with one technology were also  
309 recovered by the other (Supplementary Fig. S3A).

310 The HS287 / HS286 primer set is preferred in order to recover the greatest diversity of  
311 gene cassettes. Indeed, the recovery rate of gene cassettes using the methods described here  
312 surpasses any previously described approach. Notably, Pereira et al. [45] conducted an  
313 impressive survey of gene cassettes from 10 terabases of metagenomic data obtained from 14  
314 public databases. Across all datasets, they identified an average of 0.03 unique cassette ORFs  
315 per 500 kilobases of assembled data. Here, we recover 218 and 265 ORFs per 500 kilobases  
316 of assembled data when sequenced with ONT and MiSeq, respectively. Although screening  
317 metagenomes may provide a relatively unbiased approach in analysing gene cassettes, it

318 clearly requires much deeper sequencing to recover sufficient cassette data for in-depth  
319 ecological or evolutionary analyses. Studies of integrons and their associated genetic cargo  
320 will therefore continue to benefit from the use of amplicon sequencing approaches, such as  
321 described in the present study.

322 For the intI-R / HS286 primer set, we recovered a total of 9,641 and 3,742 non-  
323 redundant cassette ORFs when sequenced with ONT and MiSeq, respectively (Fig. 2C). ONT  
324 sequencing recovered significantly more ( $P < 0.0001$ ) of the cassette ORF diversity of each  
325 sample than MiSeq (Fig. 2D). This was despite all ONT cassette ORFs being covered by the  
326 MiSeq reads (Supplementary Fig. S3B). This shows that although the MiSeq reads cover all  
327 the cassettes being amplified by the primers, recovery of the cassettes is sub-optimal, most  
328 likely due to difficulties in their assembly. In particular, different cassette arrays associated  
329 with the same or similar integron integrase are likely to be problematic for a short-read  
330 assembly approach. While the intI-R / HS286 primer pair does not recover as much diversity  
331 as the HS287 / HS286 set, it does provide additional key information on IntI diversity  
332 (discussed further below) and indicates which gene cassettes are associated with which *intI*  
333 genes.

334 The intI-R / HS286 primer set can also reveal which gene cassettes are located  
335 towards the start of a cassette array (Fig. 1A). This is of biological and ecological  
336 significance, since the first cassettes in arrays are the most recently inserted cassettes and are  
337 likely to be strongly expressed [65]. During environmental perturbations, integron integrase  
338 activity leads to the acquisition of novel and rearrangement of existing cassettes, inserting  
339 them at the start of the array where strong expression is guaranteed [17, 66, 67]. Selection  
340 fixes lineages with first-position cassettes that confer significant advantages. Thus, gene  
341 cassettes recovered from the intI-R / HS286 primer set might provide important ecological  
342 insights at the time of sampling.

343

344 *Recovered diversity of integron integrases*

345 Using the intI-R / HS286 primers, we recovered a total of 1,413 and 1,867 different integron  
346 integrase genes among the 12 samples when sequenced with ONT and MiSeq, respectively  
347 (Fig. 3A). There was no significant difference in integron-integrase recovery between the two  
348 sequencing technologies, with or without adjusting for sequencing depth (Figs. 3A-B). Both  
349 sequencing technologies could recover an impressive diversity of IntIs from the 12 samples.  
350 In comparison, a comprehensive screening of 2,484 bacterial genomes recovered only 215  
351 different IntIs [13]. This shows that despite the low rate of intI-R / HS286 sequences that are  
352 retained after filtering, a significant number of novel integron sequences are recovered.

353 To determine how many classes of integrons these IntIs represented, we sought to  
354 define the amino acid clustering threshold for an integron class. To do this, we used the most  
355 abundant and widely distributed IntI, the class 1 integron integrase (IntI1) [68]. Here, we  
356 iteratively set decreasing amino acid clustering thresholds for our library of IntIs using CD-  
357 HIT v4.6 [69, 70] [parameters: -n 5 -d 0 -g 1 -t 0]. We continued until all IntI1s in our dataset  
358 were grouped into a single cluster while ensuring all non-IntI1s were excluded  
359 (Supplementary Fig. S4A). This resulted in a 94% amino acid identity being selected as the  
360 most appropriate clustering threshold for IntI1s. Although this might not reflect the ideal  
361 threshold for all classes, it nevertheless provides a semi-quantitative approach to defining an  
362 integron class based on amino acid homology.

363 Using a 94% clustering threshold, we recovered a total of 984 and 1,646 integron  
364 classes among our dataset when sequenced with ONT and MiSeq, respectively  
365 (Supplementary Fig. S4B). There was no significant difference in integron class recovery  
366 between the two sequencing technologies, with or without adjusting for sequencing depth  
367 (Figs. 3C-D). In addition, we examined the most prevalent integron classes; defined here as

368 those being IntIs that were present in at least one-third of all samples (Table 1). This  
369 identified ten prevalent IntI classes, found to be 60-70% similar to endogenous IntIs from  
370 diverse bacterial phyla (Table 1). Not surprisingly, class 1 integrons were the only class to be  
371 found in every sample, including those from Antarctica and outback Australia.

372

373 *Functional diversity of gene cassettes*

374 Here we show that gene cassette ORFs largely encode proteins of unknown functions (Fig.  
375 4A). This is in agreement with previous functional analyses of gene cassettes [5, 20, 24, 25].  
376 On average, only ~20% of cassette-encoded proteins amplified with HS287 / HS286 could be  
377 assigned a COG functional category, approximately half of which could be assigned a  
378 category of known function (Fig. 4A). The dominant COG categories were ‘Transcription’,  
379 ‘Replication, recombination and repair’, and ‘Amino acid transport and metabolism’ (Fig.  
380 4B). We show that our methods are capable of recovering gene cassettes that confer a wide  
381 range of traits spanning many functional classes.

382

383 *Cassette-encoded antibiotic resistance*

384 For a more specific functional characterisation, we focused on antimicrobial resistance, since  
385 these phenotypes are often conferred by integron gene cassettes in clinical settings [3, 71,  
386 72]. Interestingly, we found that for either primer set, ONT sequencing could recover many  
387 more ARGs than MiSeq, the latter recovering no ARGs for most samples (Fig. 5A). In  
388 contrast, ONT sequencing recovered as many as 300 ARG cassettes within a single sample.  
389 We suspect that this discrepancy is an artefact caused by the high similarity between different  
390 ARG types, and multiple arrangements of the same ARGs in class 1 cassette arrays that  
391 makes their assembly difficult from short-read data. In total, we recovered 106 different  
392 ARGs from both primer sets when sequenced with ONT (Fig. 5B). Almost all ARG cassettes

393 encoded proteins known to confer resistance to  $\beta$ -lactam and aminoglycoside antibiotics,  
394 these being the most commonly observed integron-mediated resistance types [3].

395 Upon examining all cassette ORFs associated with class 1 integron integrases  
396 recovered using intI-R / HS286 primers (Table 1), we found that 162 of 462 (34.6%) were  
397 known ARGs. In comparison, only 586 of the 10,385 (5.6%) total cassette ORFs amplified  
398 with this primer set were known ARGs. These findings show that class 1 integrons are  
399 collecting and concentrating ARG cassettes out of the broader diversity of cassette functions.  
400 This enrichment strongly supports the idea that class 1 integrons are key vectors for  
401 acquisition and dissemination of antibiotic resistance [3, 73, 74].

402

403 *Taxonomic classification of attC sites*

404 We could identify the likely taxonomic sources of 5,998 *attCs* (18.8%) and 10,257 *attCs*  
405 (20%) sequenced with ONT and MiSeq, respectively. For taxonomic classification, we used  
406 models that capture the sequence and structural homology of chromosomal *attCs* from 11  
407 different taxa. These included six Gammaproteobacterial orders (Alteromonadales,  
408 Methylococcales, Oceanospirillales, Pseudomonadales, Vibrionales, Xanthomonadales) and  
409 an additional five phyla (Acidobacteria, Cyanobacteria, Deltaproteobacteria, Planctomycetes,  
410 Spirochaetes). It should be noted that although the specificity (ability to reject false positives)  
411 of each model is very high (98-100%), they exhibit a wide range of sensitivities (proportion  
412 of true positive detected) [44]. Therefore, relative abundance of each taxon cannot be  
413 compared within the same sample, however, the same taxon can be compared between  
414 different samples. It also indicates that the relative abundances of each taxon are likely to be  
415 lower-bound estimates.

416 Here, we show that the relative abundance of each taxon varied across the different  
417 sampled environments (Fig. 6). For example, both the Cyanobacteria- and Methylococcales-

418 type *attCs* were most abundant in urban parkland soil (Fig. 6C-D), while *Vibrionales*-type  
419 *attCs* were more abundant in estuarine sediments and freshwater biofilm samples (Fig. 6F  
420 and Supplementary Figure S5 for a comparison of all 11 taxa). Such data can provide useful  
421 information on the taxonomic contribution to gene cassette pools among different samples.

422

## 423 **Conclusions**

424

425 Here, we present experimental and bioinformatic methods for the PCR amplification, DNA  
426 sequencing and analysis of integrons from microbial communities. We describe approaches  
427 using two different PCR assays and compare the outputs from ONT and MiSeq sequencing.  
428 We find that, relative to sequencing depth, ONT generally outperforms or performs the same  
429 as MiSeq regarding the recovery of gene cassettes and integron integrases. Most notably,  
430 ONT outperforms MiSeq in the recovery of complete ARG gene cassette sequences. We also  
431 find that the primer set HS287 / HS286 is efficient at amplifying a wide range of gene  
432 cassettes, encompassing extensive *attC* and functional diversity. However, the intI-R / HS286  
433 primer set can provide additional useful information in linking gene cassettes with an  
434 integron class. For example, we show that class 1 integrons are collecting and concentrating  
435 ARGs relative to the broader cassette pool.

436 Our described methods can recover key information on the diverse pool of gene  
437 cassettes that are helping drive adaptation and niche specialisation in bacteria [4, 16, 67].  
438 Such an approach allows us to investigate the potential traits that are available to integron-  
439 carrying bacteria, and to understand the role that gene cassettes play in mediating  
440 evolutionary responses under environmental or clinical selection pressures. In addition, the  
441 large proportion of cassettes with unknown functions provides an important resource for the  
442 discovery of novel enzymatic activities [17].

443

444 **Data availability**

445 All sequence data generated in this study are available from the NCBI SRA database under  
446 the BioSample accessions SAMN21354384 to SAMN21354431. All BioSamples are linked  
447 to the NCBI BioProject PRJNA761546.

448

449 **Code availability**

450 The code used for filtering sequences to ensure that they represent amplicons from genuine  
451 integrons are available at <https://github.com/timghaly/integron-filtering>. The code used to  
452 predict the taxonomic sources of gene cassette recombination sites (*attCs*) is available at  
453 <https://github.com/timghaly/attC-taxa>.

454

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458

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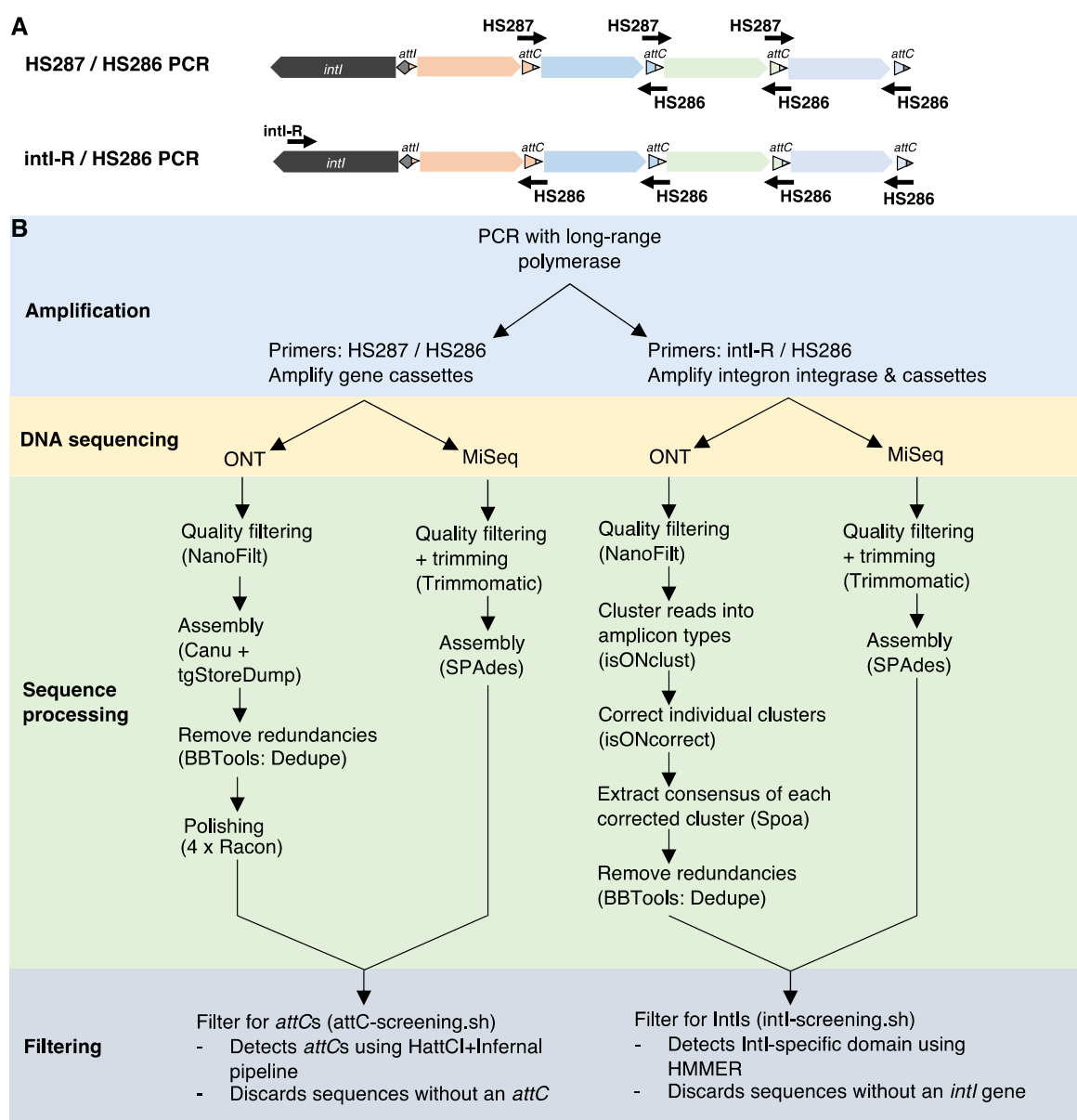
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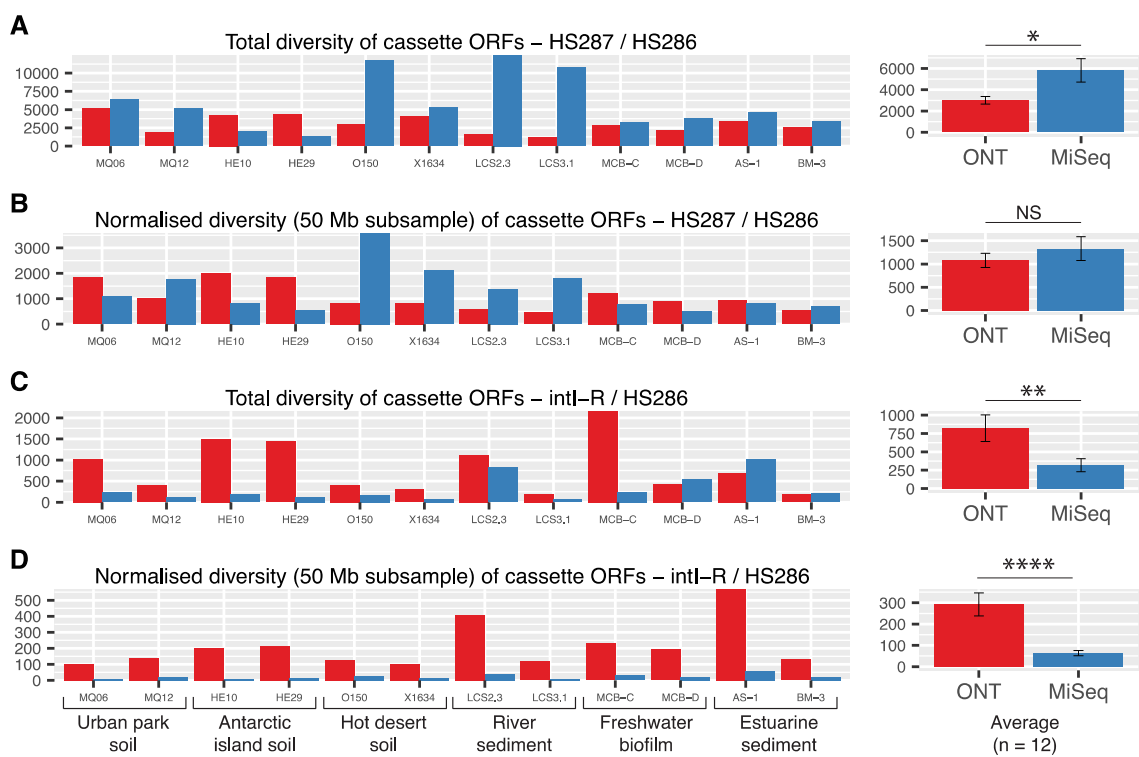
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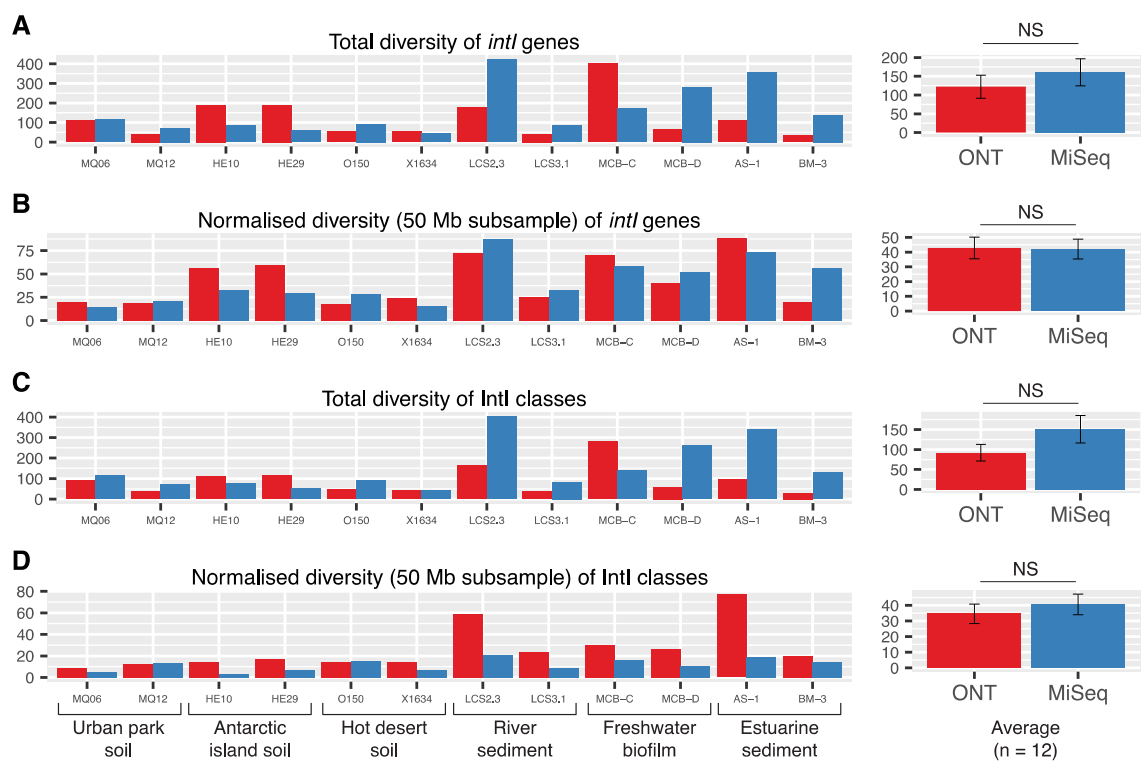
667 **Figures and Table**



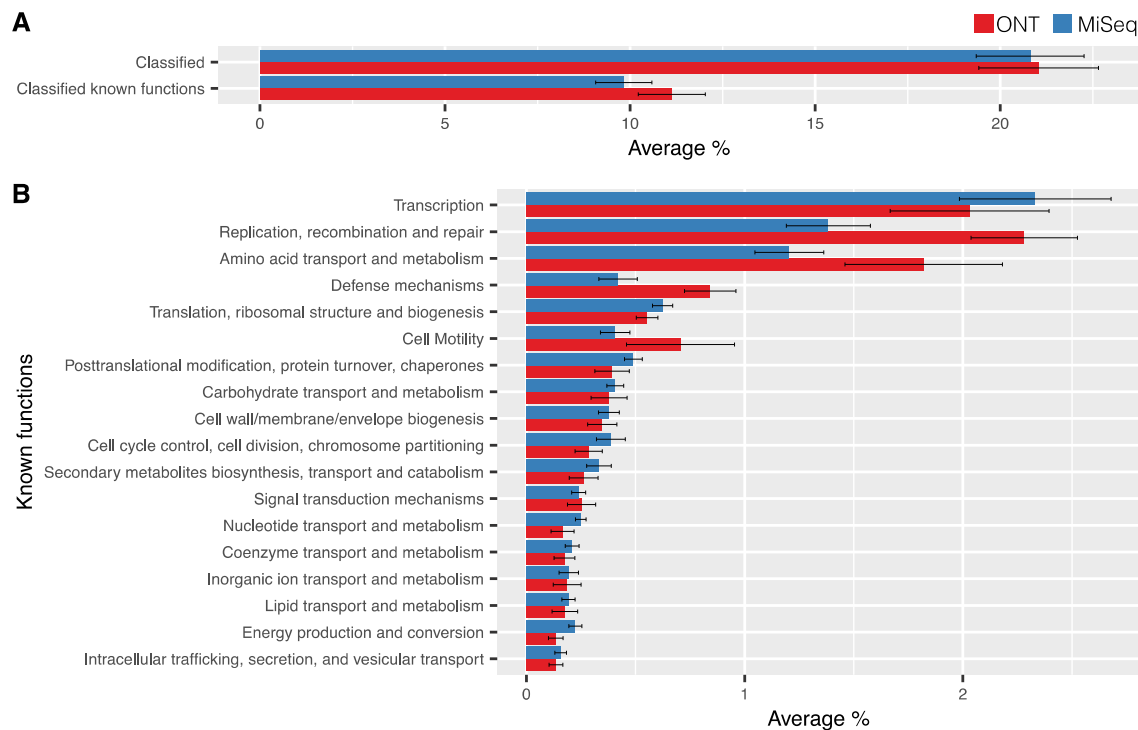
668 **Figure 1. Experimental and bioinformatic workflow for gene cassette amplicon**  
669 **sequencing.** (A) Components of integrons amplified by the two PCR assays. The primer set  
670 HS287 / HS286 targets cassettes that lie between two *attC* sites. Potentially any gene  
671 cassette(s) can be amplified by this primer set. The primer set intI-R / HS286 targets diverse  
672 integron integrases (*intI*) and cassette recombination sites (*attC*). The resulting amplicons  
673 include ~800 bp of *intI* and at least the first cassette(s) of an array. (B) The bioinformatic  
674 steps and software (in parentheses) used to process and filter amplicon data. Methods are  
675 shown for both primer sets sequenced with either Nanopore (ONT) or Illumina MiSeq.



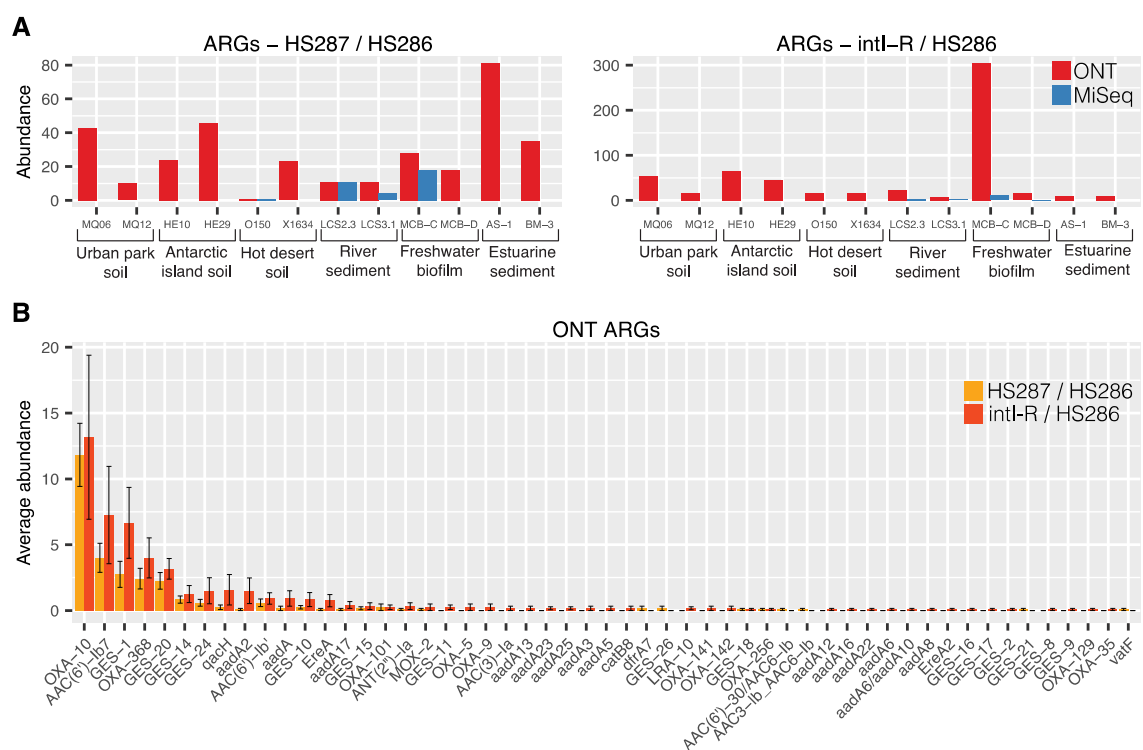
676 **Figure 2. Diversity of recovered gene cassette ORFs.** Redundancy was removed using a  
677 100% amino acid identity of translated protein sequences. **(A)** Total non-redundant cassette  
678 ORFs amplified using the primers HS287 / HS286. **(B)** Cassette ORF diversity was  
679 normalised for HS287 / HS286 sequencing depth (based on a 50 Mb subsample of sequence  
680 reads). Total **(C)** and normalised **(D)** cassette ORF diversity are shown for the intI-R / HS286  
681 primer set. Average ( $\pm 1$  S.E) diversity for each analysis are shown on the right-hand side of  
682 each panel. The degree of statistical significance is shown by asterisks as determined by two-  
683 sample T-tests or Wilcoxon rank sum tests (depending on the normality of the data). NS:  
684 P>0.05, \*: P <0.05, \*\*: P<0.01, \*\*\*: P<0.001, \*\*\*\*: P<0.0001.



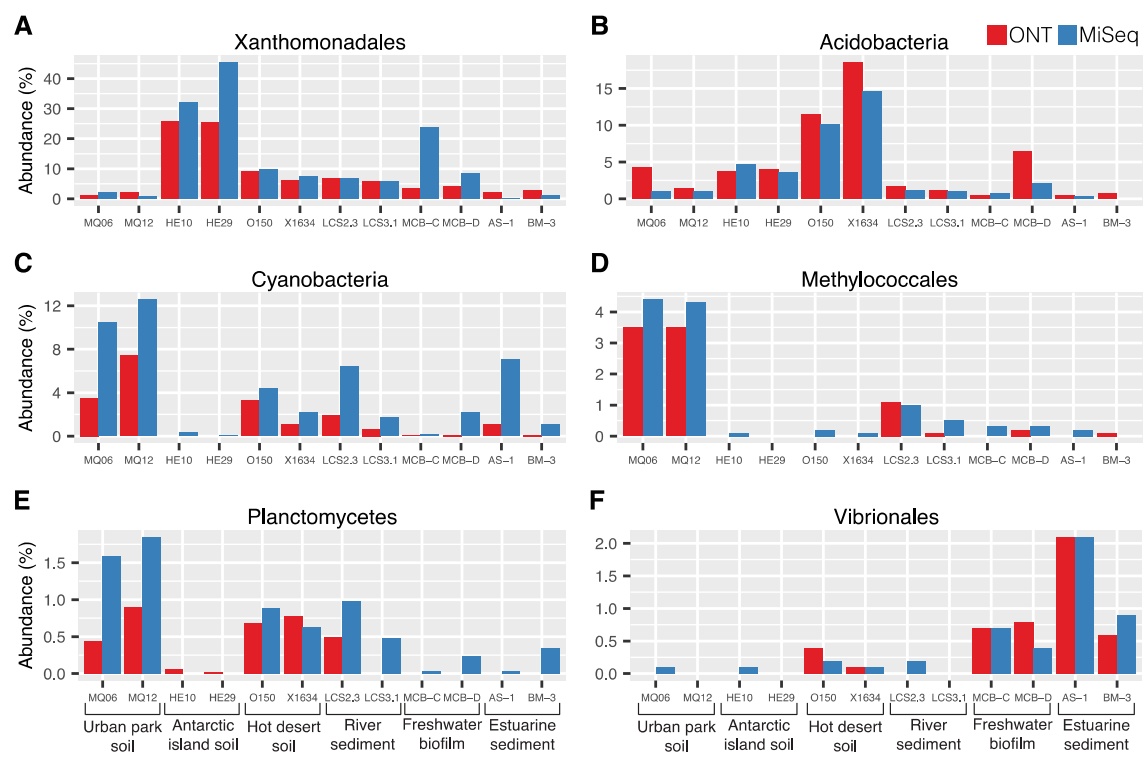
685 **Figure 3. Diversity of integron integrases recovered by the intI-R / HS286 primer set.**  
686 (A) Total non-redundant (100% amino acid identity) integron integrases (IntIs) recovered.  
687 (B) IntI diversity was normalised for sequencing depth (based on a 50 Mb subsample of  
688 sequence reads). Total (C) and normalised (D) diversity of IntI classes (using a 94% amino  
689 acid clustering threshold) are shown. Average ( $\pm 1$  S.E) diversity for each analysis are shown  
690 on the right-hand side of each panel. Differences between Nanopore (ONT) and Illumina  
691 MiSeq technologies were not significant (NS) as determined by Wilcoxon rank sum test.



692 **Figure 4. COG functional analysis of cassette-encoded proteins recovered with the**  
693 **HS287 / HS286 primer set. (A)** Average ( $\pm 1$  S.E) percentage of proteins per sample (n=12)  
694 that can be classified into functional categories. On average ~20% of cassette-encoded  
695 proteins can be classified by a COG category, half of which fall into categories of known  
696 function. **(B)** The average ( $\pm 1$  S.E) proportion of proteins within a sample assigned to each  
697 of the known functional categories.



**Figure 5. Abundance and diversity of antibiotic resistance gene (ARG) cassettes. (A)**  
 Abundance of ARGs recovered from either primer set. (B) The average ( $\pm$  1 S.E) abundance of each ARG type recovered from Nanopore (ONT) sequencing per sample (n=12).



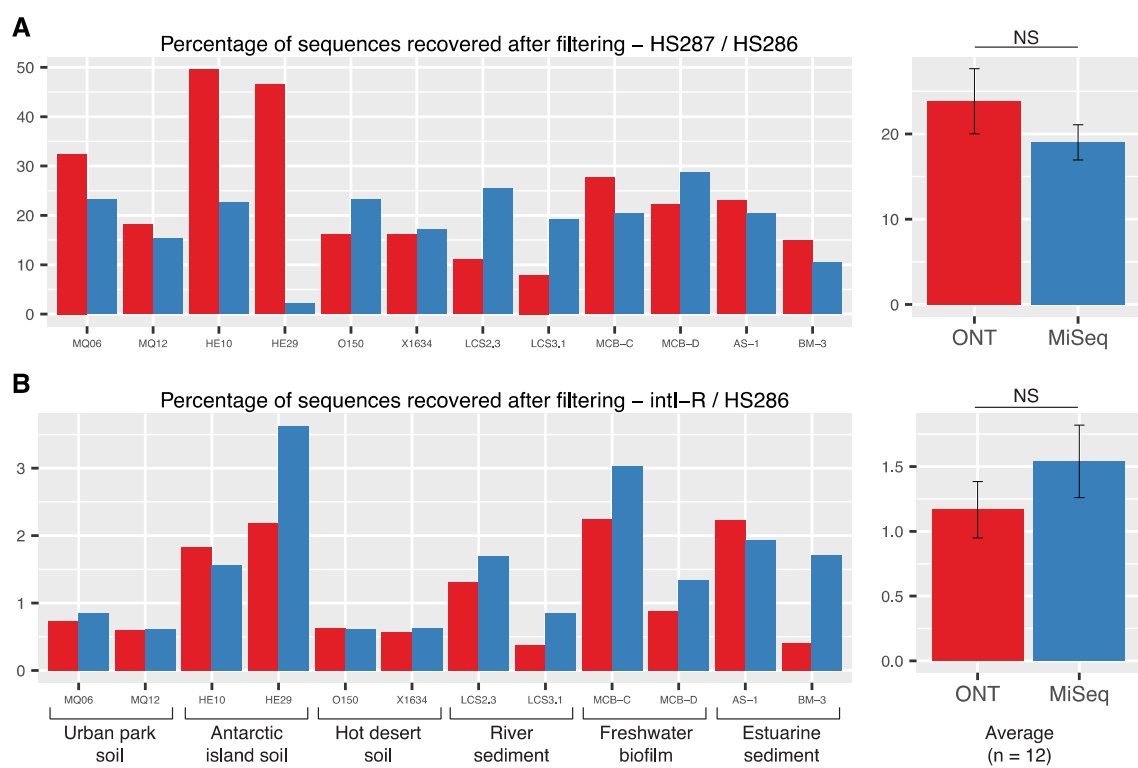
701 **Figure 6. Taxonomic classification of gene cassette recombination sites (*attCs*).**  
702 Taxonomic predictions are based on a selection of six (A-F) of the eleven available  
703 taxonomic models of chromosomal *attCs*. Each figure panel shows the proportion of *attCs*  
704 across each sample that exhibit sequence and structure conserved among that taxon. For a  
705 comparison of all eleven taxa, see Supplementary Figure S5.

706

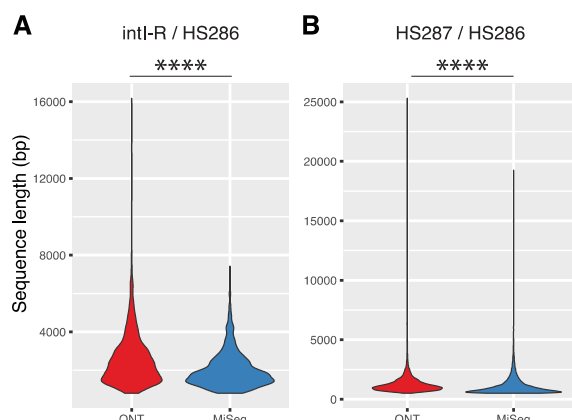
**Table 1. Most prevalent integron integrase (IntI) classes**

Number of IntIs in cluster/class	Percentage of samples present in	BLASTP taxa	BLASTP amino acid %
94	100	Class 1 integron - Multispecies	99.3
71	91.7	Xanthomonadales (Rhodanobacteraceae & Xanthomonadaceae)	~70
19	91.7	Multiple phyla (Deltaproteobacteria, Nitrospinae, Chloroflexi)	~60
26	66.7	Xanthomonadaceae ( <i>Lysobacter</i> , <i>Vulcaniibacterium</i> , <i>Thermomonas</i> , <i>Luteimonas</i> , <i>Pseudoxanthomonas</i> )	~70
10	58.3	Betaproteobacteria	~80
13	41.7	'IntI1-like' - Multispecies	~91
13	41.7	Rhodanobacteraceae	~75
12	41.7	Xanthomonadales (Rhodanobacteraceae & Xanthomonadaceae)	~74
8	41.7	Xanthomonadales (Rhodanobacteraceae & Xanthomonadaceae)	~72
7	41.7	Planctomycetes	~67

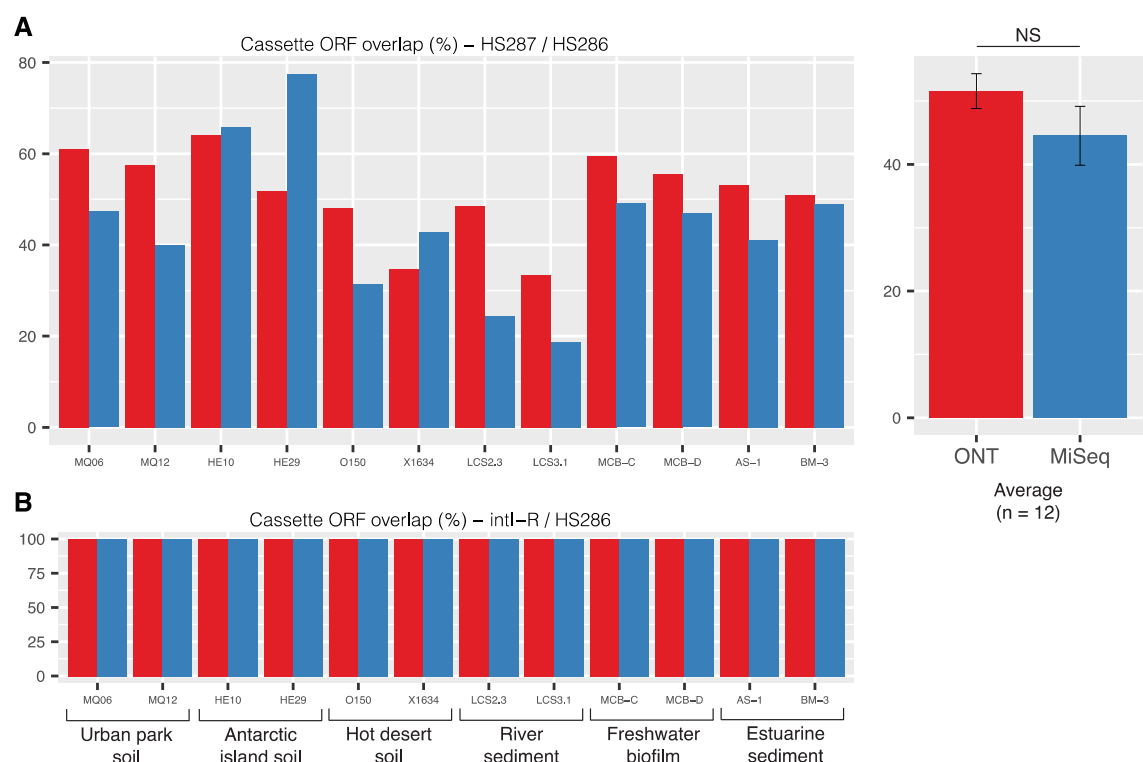
707 **Supplementary Figures**



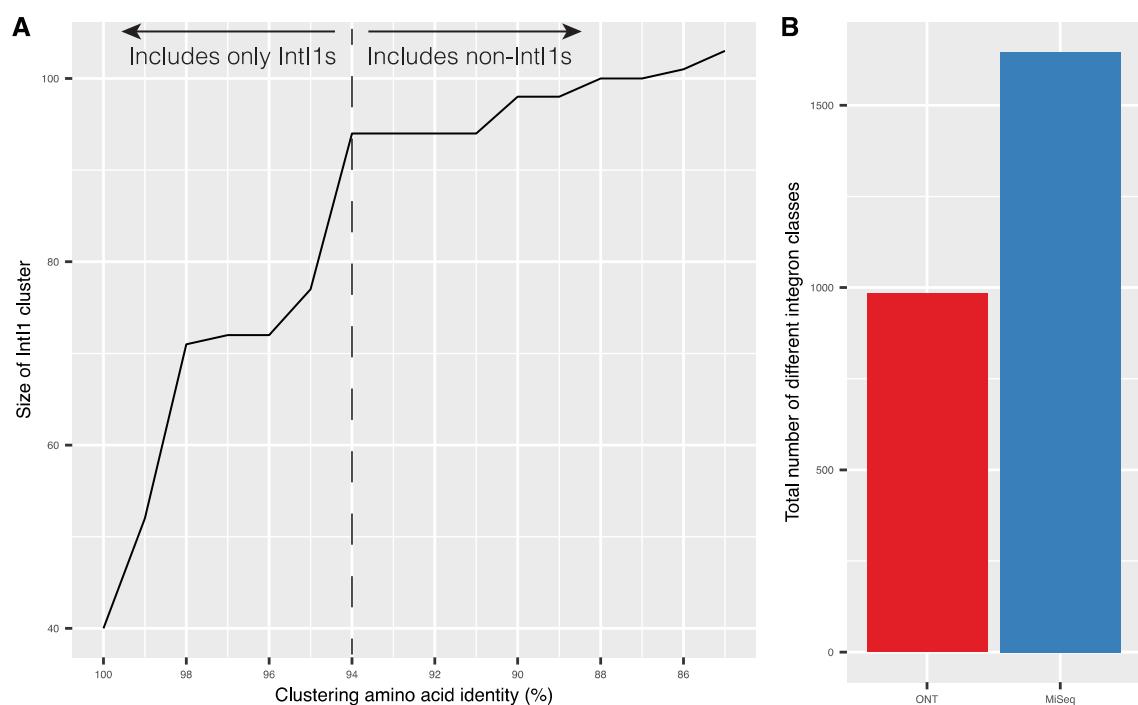
708 **Figure S1. Percentages of sequences recovered after bioinformatic filtering.** Filtering  
709 involved screening (A) HS287 / HS286 sequences for cassette recombination sites (*attC*s)  
710 and (B) intI-R / HS286 sequences for integron integrase (IntI) encoding genes to ensure that  
711 they represented amplicons of genuine integrons. Average ( $\pm 1$  S.E) diversity for each  
712 analysis are shown on the right-hand side of each panel. Differences between Nanopore  
713 (ONT) and Illumina MiSeq technologies were not significant (NS) as determined by two-  
714 sample T-tests.  
715



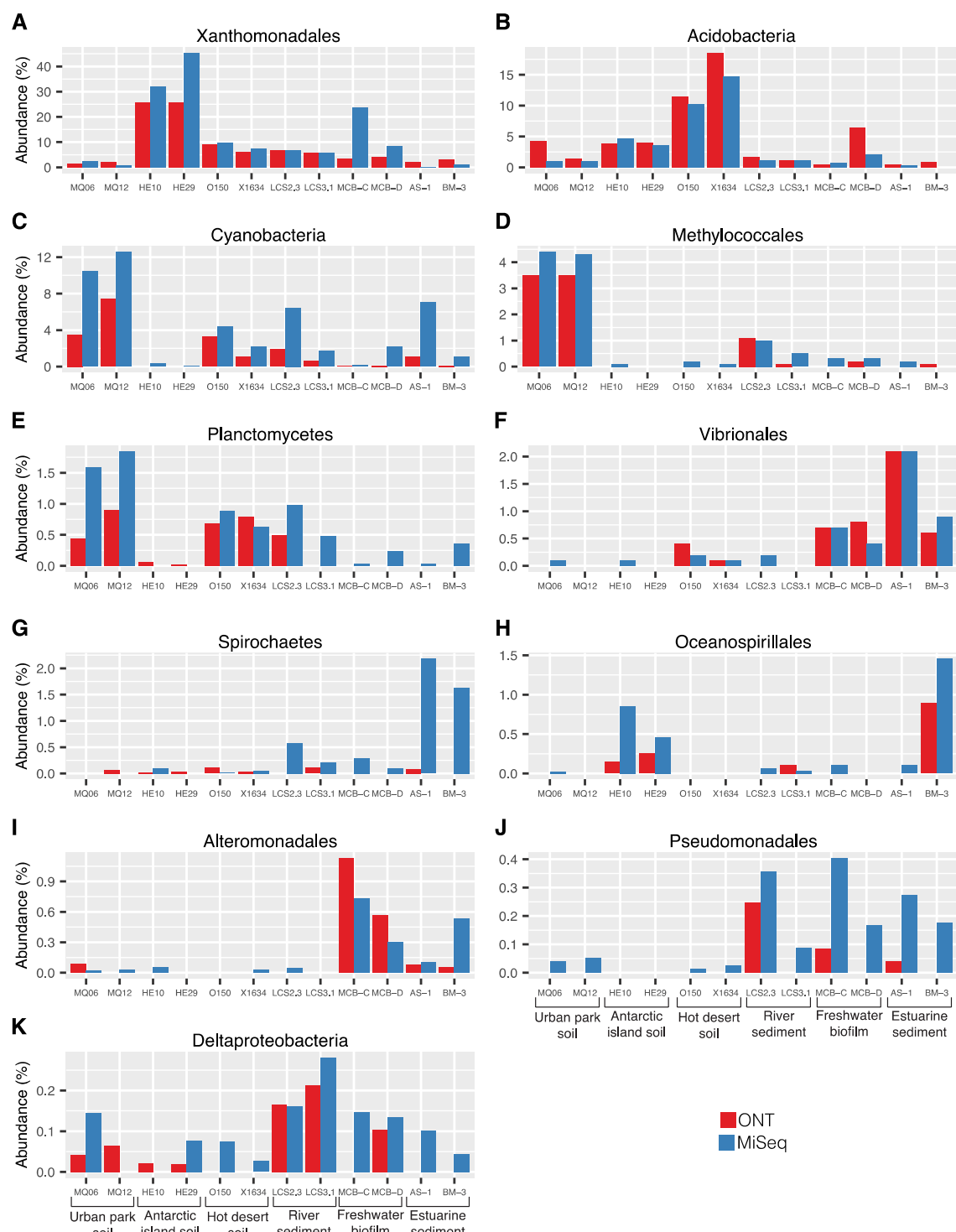
716 **Figure S2. Sequence lengths recovered by each sequencing technology.** The violin plots  
717 show the range of sequence lengths (bp) for (A) intI-R / HS286 and (B) HS287 / HS286  
718 primer sets. The width of each curve represents the relative density of datum points across the  
719 ranges. For both primer sets, the average length of recovered amplicons (n=12) is  
720 significantly larger (Wilcoxon rank sum test, P<0.0001) when sequenced with Nanopore  
721 (ONT) compared MiSeq sequencing.



722 **Figure S3. Overlap in recovered ORFs between Nanopore (ONT) and MiSeq**  
723 **sequencing technologies.** The percentage of ORFs recovered from one sequencing  
724 technology that were covered by reads from the other technology are shown for (A) HS287 /  
725 HS286 and (B) intI-R / HS286 primer sets. ORFs considered to present in the opposite  
726 sequencing technology had to have a mean coverage depth of at least 1x that spanned at least  
727 98% of the ORF. The average ( $\pm$  1 S.E) percentage overlap for HS287 / HS286 data is shown  
728 on the right-hand side of panel (A). There was no significant (NS) difference between ONT  
729 and MiSeq (Two-sample T-test, P=0.209).



730 **Figure S4. Diversity of integron classes.** (A) The amino acid clustering threshold for  
731 integron classes was determined using class 1 integron integrases (IntI1) present in our  
732 dataset. Decreasing amino acid clustering thresholds were iteratively set until all IntI1s were  
733 grouped in the same cluster and all non-IntI1s were excluded. A protein sequence was  
734 considered to be IntI1 if it aligned with any previously characterised class 1 integron in  
735 GenBank using BLASTP (>98% amino acid identity and >70% subject cover). An amino  
736 acid clustering threshold of 94% was found to include all IntI1s (n=94) and exclude all non-  
737 IntI1s. (B) The total number of integron classes (based on a 94% amino acid clustering  
738 threshold) recovered for all samples (n=12).



739 **Figure S5. Taxonomic classification of gene cassette recombination sites (attCs).**  
740 Taxonomic predictions are based on all eleven (A-K) available taxonomic models of  
741 chromosomal attCs. Each figure panel shows the proportion of attCs across each sample that  
742 exhibit sequence and structure conserved among that taxon.  
743