RESEARCH ARTICLE

High Throughput Method for Analysis of Repeat Number for 28 Phase Variable Loci of Campylobacter jejuni Strain NCTC11168

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Abstract

Mutations in simple sequence repeat tracts are a major mechanism of phase variation in several bacterial species including Campylobacter jejuni. Changes in repeat number of tracts located within the reading frame can produce a high frequency of reversible switches in gene expression between ON and OFF states. The genome of C. jejuni strain NCTC11168 contains 29 loci with polyG/polyC tracts of seven or more repeats. This protocol outlines a method—the 28-locus-CJ11168 PV-analysis assay—for rapidly determining ON/OFF states of 28 of these phase-variable loci in a large number of individual colonies from C. jejuni strain NCTC11168. The method combines a series of multiplex PCR assays with a fragment analysis assay and automated extraction of fragment length, repeat number and expression state. This high throughput, multiplex assay has utility for detecting shifts in phase variation states within and between populations over time and for exploring the effects of phase variation on adaptation to differing selective pressures. Application of this method to analysis of the 28 polyG/polyC tracts in 90 C. jejuni colonies detected a 2.5-fold increase in slippage products as tracts lengthened from G8 to G11 but no difference between tracts of similar length indicating that flanking sequence does not influence slippage rates. Comparison of this observed slippage to previously measured mutation rates for G8 and G11 tracts in C. jejuni colonies detected a 2.5-fold increase in slippage products as tracts lengthened from G8 to G11 but no difference between tracts of similar length indicating that flanking sequence does not influence slippage rates. Comparison of this observed slippage to previously measured mutation rates for G8 and G11 tracts in C. jejuni indicates that PCR amplification of a DNA sample will over-estimate phase variation frequencies by 20-35-fold. An important output of the 28-locus-CJ11168 PV-analysis assay is combinatorial expression states that cannot be determined by other methods. This method can be adapted to analysis of phase variation in other C. jejuni strains and in a diverse range of bacterial species.

Introduction

Simple sequence repeat (SSR) tracts are highly mutable sequences due to the potential for slip strand mis-pairing during DNA replication [1]. Slippage results in insertion or deletion of one
or more repeats within the repetitive DNA tract. This combination of hypermutability and reversible mutations has driven evolution of SSR as a major mechanism of phase variation (PV) in several bacterial species [2,3]. PV describes a phenomenon of frequent, reversible alterations in expression of specific phenotypes [4]. These switches in expression are mediated by SSR located within the reading frame, core promoter or other regulatory sequences of a gene. The presence of multiple SSR-controlled phase-variable genes generates high levels of phenotypic variants within bacterial populations. Investigation of the fluctuations in population structure and adaptive benefits of these phenotypic variants requires detection of alterations in repeat number and correlation with expression state in multiple isolates from populations both prior to and post selection.

*Campylobacter jejuni* is a leading cause of human bacterial gastroenteritis, with contaminated meat products considered a main source of infection [5,6]. *C. jejuni* readily colonises the intestinal mucosa of a wide variety of wild and domestic birds and other animals. Infections in poultry are usually asymptomatic whilst human infection can result in significant inflammation and a profuse, bloody diarrhoea. As a commensal of poultry and a pathogen of humans, *C. jejuni* needs to rapidly adapt to fluctuations in host environments such as changing nutrient compositions and appearance of innate/adaptive immune effectors. Further selective pressures are caused by transmission through genetically and immunologically variable host populations and exposure to bacteriophages. *C. jejuni* is likely to use PV as a major mechanism for adaptation to these selective pressures.

Mononucleotide repeat tracts consisting of seven or more cytosine or guanine bases are an unexpected feature of the AT-rich genomes of *C. jejuni* [7]. These SSR are the main mechanism of PV in this species and an analysis of four genome sequences indicated the presence of 12 to 29 tracts per genome [8]. The majority of these loci are predicted to encode enzymes involved in modification of surface structures (i.e. lipooligosaccharide, flagella and capsule) but a few encode surface proteins or restriction enzymes [9–12]. There are 29 polyG/polyC tracts in *C. jejuni* NCTC11168. The majority (n = 23) of these tracts are located within the main part of the reading frames whilst the others are at the 3’ end of the reading frame (n = 1; cj0045), in pseudogenes (n = 2; cj0046 and cj0676), or in intergenic regions (n = 3; cj0565, cj0742 and cj1321) [9]. The tracts in the reading frame mediate ON or OFF switches in gene expression as shown for a sub-set of genes using reporter constructs or antibodies against the encoded protein or modified epitope [8, 13].

An important aim has been to understand the contributions of PV to specific phenotypes and to more generic behaviours such as colonisation of host animals. Some limited experimental characterisation has been performed on individual phase-variable genes, including locking the reading frame into an ON or OFF state [14–16]. A more generic approach has been to analyse the whole population by PCR using fluorescent primers and fragment analysis on an auto sequencer. The major and minor tract lengths are estimated from the relative sizes of peaks [17–19]. This approach is limited by the slippage that is known to occur in repetitive tracts during PCR [20]. An alternative approach is to perform next generation sequencing (NGS) and estimate numbers of variants from sequences spanning a repeat tract. However, NGS is not as high-throughput as PCR and the error rate of NGS increases as a function of repeat number, hence NGS has not yet been widely used to analyse PV [21]. Furthermore, while both of these approaches have utility for determining the PV states for individual genes neither can be used for determination of combinatorial expression states.

The presence of multiple phase-variable genes in *C. jejuni* genomes raises the potential for combinatorial effects particularly where genes are involved in modification of the same macromolecule [9]. In order to investigate this phenomenon, the tract lengths of each phase-variable gene within individual cells need to be determined. Single colonies, derived from a population...
by plating out serial dilutions, can be utilised as surrogates of single cells under the assumption that switches in the repeat tract are infrequent during growth of the colony. As switching rates are ~1×10⁻³ mutations per division [8], the major tract length of each gene in each colony will be the same as the initial, derivative single cell. The major tract length can then be determined for colony lysates by PCR and fragment analysis. A multiplex PCR and fragment analysis assay was previously developed for six of the polyG/polyC tracts of C. jejuni strain NCTC11168 and used to determine expression states for six genes in samples from in vitro and in vivo passage experiments [8]. A manual analysis protocol was utilized for assessing tract length and expression state from the fragment sizes output of the fragment analysis assay. Expression states were determined for 30 colonies derived from each population as well as DNA extracted from the total population. Estimates of the major tract length and expression state were similar for each sample indicating that the method was robust for determining expression states and was utilized for analysis of combinatorial expression states [8].

A scale up of the six-gene PV assay to incorporate all 28 phase-variable loci of C. jejuni required navigation of a series of complicated issues. Some of the phase-variable loci contain identical sequences in and around the repeat tracts necessitating development of locus-specific primers and careful design of multiplex PCR reactions. A 28 multiplex fragment analysis assay was required in order to minimise cost and maximise throughput. Similarly, high throughput and accuracy necessitated an automated extraction process for conversion of fragment analysis data into tract lengths and expression states. This paper describes a rapid and accurate 28-locus-CJ11168 PV-analysis assay whose data outputs include user-friendly formats readily accessible to further analysis in downstream applications.

**Materials and Methods**

**Part I—Multiplex PCR amplification of 28 phase-variable loci**

**Bacterial DNA extraction.** C. jejuni colonies were grown on Campylobacter blood-free selective agar plates (CCDA, Oxoid) for 48 to 72 hours at 37°C in 5% oxygen, 10% carbon dioxide and 85% nitrogen. For DNA extraction, single colonies were re-suspended in 100 μl molecular-grade water in 0.5 ml PCR tubes or 96-well plates and heated to 95°C for 5 min. Tubes were then briefly vortexed before centrifugation for 2 min. The aqueous part (containing bacterial DNA) was transferred into a 96-well plate and stored at -20°C.

**Primer design.** Primers were designed to generate PCR products spanning each repeat tract (Table 1) and having sizes between 80 and 470 base pairs in length (Table 2). One primer in each primer pair was labelled with a fluorescent dye (FAM, VIC or NED; Table 1). Non-labelled and FAM-labelled primers were supplied by Eurofins MWG or SIGMA, VIC- and NED-labelled primers were supplied by Applied Biosystems.

Primers were designed so that the products of each primer pair could be distinguished by either fluorescent label or size of PCR product (Table 2). Any PCR products that gave overlapping peaks during fragment length analysis were clearly distinguishable by different dye colours. Three pairs of genes (cj1295 and cj1296; cj1305 and cj1306; cj1421 and cj1422) and a group of three genes (cj1310, cj1318 and cj1335) have regions of identical or high sequence identity. Gene-specific primers were designed for cj1295, cj1296, cj1305, cj1306 and cj1310 genes. These primers differed at 1 or more positions including the 3′ terminal nucleotide. For cj1421-cj1422 and cj1318-cj1335, a common unlabelled primer was combined with a fluorescently-labelled gene-specific primer whose fluorescent tag differed for each gene of the two paired loci, so that similarly-sized PCR products could be easily distinguished.

Repeat numbers for control samples and for validation were determined by dideoxy sequencing of PCR products. These products were generated by amplification with non-
Table 1. Phase-variable genes and repeat tract specific PCR primers.

| Gene      | Gene size (bp) | Location of Repeat Tract | In-frame Repeat Number | Primer name       | Sequence (5’ → 3’)                      |
|-----------|----------------|--------------------------|------------------------|-------------------|----------------------------------------|
| cj0031    | 3,732          | +2,572                   | 9G                     | cj0031-fwd-FAM cj0032-rev | GCCCTTTGATCTCATCATCAGGG GCCAAACCTCCCCTATTCCT |
| cj0045c   | 723            | +709                     | 11G<sup>2</sup>        | cj0045-fwd-FAM cj0045-rev | TTTTACATGAACCCCCAGAG CTTAAAGTGCCAGGCTGG |
| cj0046    | 1,407          | +610                     | (11G)                  | cj0046-fwd-NED cj0046-rev | TCAAAATTCTCAGAACCCCCAGAG TAGAACTATTAGCCGCT |
| cj0171    | 744            | +243                     | 8G                     | cj0171-fwd-NED cj0171-rev | TGGTTTGAGGAATGGGTTGGG GCTCCTTCATTAGATTC |
| cj0275    | 1,251          | +694                     | 8G                     | cj0275-fwd-NED cj0275-rev | ATACTCCTGAGTATAGGGG AAACTCAAATGGCCTACCC |
| cj0565    | 729            | -58                      | (10G)                  | cj0565-fwd cj0565-rev-VIC | AATTTCCAATCCCCTATGAGTA TGGTTGAGGAATGGGTTG |
| cj0617    | 609            | +559                     | 10G                    | cj0617-fwd-VIC cj0617-rev | TGGTAATGACTCTAGGTCAGGA AACAACTAATCTCAAGG |
| cj0628 (capA) | 3,432        | +501                     | 11G/10G<sup>3</sup>    | CapA-fwd-FAM CapA-rev | TATTTCTCTAATGAGCCAC GAGGAACATTCACCCCTTC |
| cj0676    | 1,656          | +854                     | 10G                    | cj0676-fwd-NED cj0676-rev | ATGCTTCTCCCTATGAGGTCG TGGTTTAAACCCAAATTC |
| cj0685c   | 1,350          | +877                     | 9C                     | cj0685-fwd-FAM cj0685-rev | GATACGAGAATAAATCTCTCAAATC GAGAAATCCCCATACCAAG |
| cj1139c   | 909            | +330                     | 8G                     | cj1139-fwd-VIC cj1139-rev | GACACTTCACTTTATATCCG AAAATGTGATGCT |
| cj1144c   | 843            | +294                     | 10G                    | cj1144-fwd-NED cj1144-rev | GATGTGTGTTGACCTTG TGGTGACCTCCACCTTCAG |
| cj1295    | 1,305          | +143                     | 9G                     | cj1295-fwd-NED cj1295-rev | TTCCCTACTCCCTAGGATATC ATAGGCTTCTTAAATCC |
| cj1296    | 792            | +309                     | 10G                    | cj1296-fwd-NED cj1296-rev | ATAAAGTGTGACTCTAAAGGC CAGCAAAAGAAAATAGGG |
| cj1305c   | 1,215          | +579                     | 9G                     | cj1305-fwd-VIC cj1305-rev | CAACCTTTATATCCCTACTTATGGAG AAAAGCGACCCCAGATTATC |
| cj1306c   | 1,224          | +579                     | 9G                     | cj1306-fwd-NED cj1306-rev | TTAATTTCTCTTGCGTGGGAAA AAAATGTGATGCT |
| cj1310c   | 1,212          | +579                     | 9G                     | cj1310-fwd-NED cj1310-rev | GACAAATATTCTCTTATAAG TGGTGAAACTCTTCTATTCAG |
| cj1318    | 1,947          | +167                     | 11G                    | cj1318-fwd-FAM cj1318-35-rev | TGCTGCTCTCTCTACTTGAGGC GTGGTACACTCTTACCCCA |
| cj1321    | 540            | -37                      | 11G                    | cj1321-fwd-VIC cj1321-rev | AAAAGGAGATGATGCTTCC CCCCCCTCTATGATAGGC |
| cj1326    | 672            | +252                     | 9G                     | cj1326-fwd-VIC cj1326-rev | CTTGTTGAGAATTGATAGTACA TTGAGAGATGATGCTTCC |
| cj1335    | 1,944          | +168                     | 11G                    | cj1335-fwd-NED cj1335-35-rev | CACAAATTTGTTCTCTACCCAAAG GTTGGAACCTTCTTATTCAG |
| cj1342c   | 1,239          | +560                     | 9G                     | cj1342-fwd-NED cj1342-rev | TGGCAATCTTGCTTCAACCC GCAAAATGCGCACTTTATCC |
| cj1420c   | 771            | +393                     | 9G                     | cj1420-fwd-VIC cj1420-rev | GCTATGCTTCTTTATGAGGAC CTCAATGTGGCGGAGATTC |
| cj1421c   | 1,836          | +87                      | 9G                     | cj1421-22-fwd cj1421-rev-VIC | TTGGTTATTAATGTTGGGGAAA TCAAAACCATATTATATTCATTTTCT |
| cj1422c   | 1,875          | +87                      | 9G                     | cj1422-22-fwd cj1422-rev-NED | TTGGTTATTAATGTTGGGGAAA AATTGATTTGCTTCCAGGAA |
| cj1426c   | 849            | +294                     | 10G                    | cj1426-fwd-VIC cj1426-rev | TATACGGCAGAATCCAAAGG GATAATCCTTGGCCGCCAC |
| cj1429c   | 924            | +291                     | 10G                    | cj1429-fwd-VIC cj1429-rev | ATGGGAGATGATGCTTCC CCCCCCTCTATGATAGGC |

(Continued)
labelled versions of the fragment analysis primers or, for the loci with short PCR products, additional primers located further from the repeat tract. DNA sequencing reactions were performed with Phusion high-fidelity DNA polymerase (Thermo Fisher).

**Design of multiplex PCR reactions.** A series of six PCR reactions were designed to amplify between three and six loci (Table 2). Primers for genes with high sequence identity or using a common primer were assigned to different reactions in order to prevent false-priming or competition between amplifications. Reaction mix C was designed as a six locus reaction but was split into two reactions (C1 and C2) due to inefficient amplification of some genes. Primers

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**Table 2. Multiplex PCR primer mixes.**

| Primer mix | Locus | Dye colour | PCR product size |
|------------|-------|------------|------------------|
| A          | capA  | Blue       | 457              |
|            | cJ0031| Blue       | 221              |
|            | cJ0045| Blue       | 280              |
|            | cJ0685| Blue       | 128              |
|            | cJ1326| Blue       | 165              |
|            | cJ1342| Blue       | 392              |
| B          | cJ1318| Blue       | 332              |
|            | cJ1420| Blue       | 109              |
|            | cJ1426| Blue       | 209              |
|            | cJ1429| Blue       | 89               |
|            | cJ1437| Green      | 275              |
| C1         | cJ0275| Yellow     | 215              |
|            | cJ1296| Yellow     | 104              |
|            | cJ1306| Yellow     | 304              |
| C2         | cJ0171| Yellow     | 282              |
|            | cJ1144| Yellow     | 173              |
|            | cJ1335| Yellow     | 337              |
| D          | cJ0565| Green      | 357              |
|            | cJ0617| Green      | 167              |
|            | cJ1139| Green      | 200              |
|            | cJ1305| Green      | 220              |
|            | cJ1321| Green      | 149              |
|            | cJ1421| Green      | 287              |
| E          | cJ0046| Yellow     | 353              |
|            | cJ0676| Yellow     | 206              |
|            | cJ1295| Yellow     | 159              |
|            | cJ1310| Blue       | 365              |
|            | cJ1422| Yellow     | 399              |

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1. Values in brackets indicate the arbitrary number of repeats coded as ON because the repeat tract is intergenic or within a pseudogene.

2. G11 would allow the translation to extend by another 15 amino acids, overlapping the start of cJ0044c; G9 and G10 (consensus) have little effect.

3. capA fragment varies between 11168 strains due to variation in the polyT tract immediately upstream of polyG tract (6T in the lab strain and the hypermotile 11168H strain, 5T in the chicken-adapted 11168ca strain).

[^1]: Dye colour as annotated in output files from PeakScanner™ although yellow is displayed as black in this program.

[^2]: PCR product size, as determined from the *C. jejuni* NCTC11168 genome sequence (7).
were stored as combined mixes of six to twelve primers at a concentration of 2 μM for each oligonucleotide.

**PCR amplification and GeneScan assays.** PCR reactions were performed using GoTaq polymerase (Promega) or KAPA Taq (KAPA Biosystems) with standard conditions (Table 3) in 96-well PCR plates sealed with an adhesive film (Thermo Fisher, AB-0558). A sub-set of samples from each plate were checked on 2% TAE agarose gels to confirm amplification. PCR products from each reaction were pooled together and an A-tailing reaction was performed (Table 3) to ensure uniform addition of an untemplated adenine to all PCR products. Samples were stored at -20°C, as required, between steps.

**Part II—Multiplex GeneScan and fragment length analysis**

A 1.5 μl aliquot of pooled A-tailed PCR products for a specific colony was added to 9.25 μl of deionized formamide and 0.25 μl of a GeneScan ladder (i.e. GS500LIZ or GS600LIZ). Test samples were arrayed in 96-well PCR plates with PCR and fragment analysis control samples (see below). Samples were subject to capillary electrophoresis on an ABI3100 analyzer. Output files (*.fsa) generated by the autosequencer were loaded into Peak Scanner™ software. Determination of fragment size was performed using default settings, with GS500LIZ or GS600LIZ specified as size standard and 'PP—Primers Present' in the 'Analysis Method' column. An example of PeakScanner™ graphical view is shown in Fig 1A.

**Part III—Automated determination of repeat number and expression state**

**Data extraction from PeakScanner using PSAnalyse.** A complete output data set, including detailed information about all the peaks from each sample in the 96 well plate, was extracted from PeakScanner™ as a tab-delimited text file using the 'Export Combined Table' command. We have developed a Perl script (PSAnalyse) that will accept this output file of PeakScanner™ and analyse each sample to determine the fragment length, peak height, repeat number and expression state of the major peak for each of the 28 phase-variable loci of *C. jejuni* strain NCTC11168. Additionally the script will output the ratio of the major peak to flanking peaks. The repeat numbers and ON/OFF states for each PV locus are generated by comparison.

### Table 3. PCR amplification and A-tail reactions.

| Step | Reaction mix for GoTaq (Promega) | Reaction mix for KAPA Taq (KAPABiosystems) | Reaction in PCR cycler |
|------|----------------------------------|---------------------------------------------|------------------------|
| **Multiplex PCR** | 5x Buffer—2 μl | 10x Buffer A—1 μl | 94°C—5 min |
| | dNTPs (10 mM)—0.5 μl | dNTPs (10 mM)—0.5 μl | 25 cycles: |
| | MgCl₂ (25 mM)—1.2 μl | MgCl₂ (25 mM)—1.8 μl | 94°C—30 s |
| | Primer mix (2 μM)—1 μl | Primer mix (2 μM)—1 μl | 50°C—30 s |
| | GoTaq polymerase—0.1 μl | KAPA Taq polymerase—0.1 μl | 72°C—60 s |
| | DNA template—1 μl | DNA template—1 μl | 72°C—5 min |
| | Water—to 10 μl | Water—to 10 μl | 10°C—hold |
| **Pooling of PCR products and A-tail reaction** | **A-tail mix:** | **A-tail mix:** | Transfer to a new 96-well plate: |
| | 5x Buffer—0.8 μl | 10x Buffer A—0.4 μl | - 4 μl A-tail mix |
| | MgCl₂ (25 mM)—1 μl | MgCl₂ (25 mM)—0.4 μl | - 2 μl each multiplex PCR |
| | GoTaq polymerase—0.05 μl | KAPA Taq polymerase—0.05 μl | reaction |
| | Water—to 4 μl | Water—to 4 μl | (total volume 16 μl) |

Protocol for amplification of 28 PV sites by multiplex PCR in 96-well plates, including the A-tailing step before mixing of samples with Size Standard for GeneScan analysis.

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to control samples with known repeat numbers and expression states. Control samples were located on the same plate as the test samples. For ease of use, we also developed a user-friendly C# frontend allowing use of the script via text boxes/drop-down menus as an alternative to entry from the command line.

**Fig 1. Analysis of multiplex fragment analysis data in PeakScanner.** (A) PeakScanner application layout. Labels 1–6 refer to: 1, Sample Name, well location in the 96-well plate; 2, Size Standard, name of size standard as selected from the drop-down menu; 3, Analysis Method, analysis method name, e.g. PP—Primers Present, assumes primers have not been removed from the sample; 4, Size, length of PCR fragment for a specific peak in base pairs; 5, Height, height of each peak indicating fluorescent dye signal strength; 6, Dye Legend, displayed dye colours and relative scale. (B) Enlarged section of Panel A showing how similarly-sized PCR products can be distinguished using different dye colours and the occurrence of flanking peaks around a major peak.

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After input of the tab-delimited text file into PSAnalyse, the user sets cut-offs for peak 'height' (normal setting for efficient PCR reactions is 1,000), 'ratio to flanking peaks' (default setting is 1.5), 'tract error' (distance of observed peak size from expected peak size; allows for small well-to-well variations in migration of PCR products; default setting is 0.33 bp) and 'scan width' (distance from expected peak size for identification of a peak; default setting is 3.5 bp). The user then identifies the location of the control samples and assigns the matching peak set file, which contains the expected fragment size and fluorescent tag for each locus, and enables the program to assign each peak to a specific locus. The 'peakset' files also defines the repeat number (or numbers) associated with an ON expression state. The user also sets the calibration file (calib file), which contains the previously determined repeat numbers for the specific control sample present in the control well. The user can also input a CSV file with the names of each sample for each well of the analysed plate otherwise sample names are based on the tab-delimited text file exported from PeakScanner™. When run, PSAnalyse generates a series of output files (Table 4) containing combined and separated data for each sample. Note that expression states are coded as 0 for OFF and 1 for ON while datapoints that fail quality control checks are indicated with a '?' and missing data are shown with an 'N'.

**Accession number.** PSAnalyse and the control and test sequencing data are available on Dryad with accession number:- doi:10.5061/dryad.k9b6f.

**Results and Discussion**

Analysis of the multiple phase-variable genes of *C. jejuni* strains requires detection of indels in hypermutable polyG tracts and of the combinatorial effects of expression states. We have therefore developed a high throughput method—the 28-locus-CJ-11168 PV analysis assay—that involves multiplex PCR and fragment analysis of 28 PV loci from single colonies of *C. jejuni* strain NCTC11168. The method involves semi-automated analysis of PCR fragment sizes, repeat numbers and ON/OFF states for each locus. The method can be split into three stages:- (1) amplification of 28 PV sites by multiplex PCR using fluorescently labelled primers; (2) fragment length analysis by capillary electrophoresis and PeakScanner™; and (3) automated calling of repeat numbers and assignment of ON/OFF states for each gene using a custom script, PSAnalyse.

**Development and testing of the multiplex PCR and GeneScan assay**

The crucial features of the primer design for the multiplex PCR assay were:- high specificity of at least one primer within a pair for the target locus; and ensuring that no two PCR products
had both the same or similar size (<15 base pairs difference) and the same fluorescent label. This design ensured that every PCR product was distinguishable by size, dye colour or both. The 28-locus-CJ-11168 PV analysis assay utilised this approach to encompass 28 of the polyG/polyC tracts present in this strain (Table 1). One of the intergenic repeats, located at the 3’ end of two convergent genes (cj0742 and cj0743c), was excluded in order to reduce the potential for generating overlapping PCR products and due to the limited potential for this repeat to influence gene expression. The primers for cj1305 were observed to have additional binding sites in cj1306 and a downstream region resulting in >4 kbp product. Amplification was, however, inefficient and did not influence production of the expected 219 bp fragment for the SSR within cj1305.

To validate the assay, a series of lysates were prepared from single colonies of C. jejuni strain NCTC11168. These lysates were analysed using the multiplex PCR/fragment analysis assay and repeat numbers were assigned for each gene using PSAnalyse. A random set of these lysates were then re-amplified using locus-specific primers without fluorescent labels (2–13 per locus) and then these PCR products were subject to dideoxy sequencing (S1 Table). The sequence data was analysed by BLAST to confirm that the correct locus had been amplified and then repeat number was determined from observation of the trace files. For the genes with regions of high identity (cj1295 and cj1296; cj1305 and cj1306; cj1421 and cj1422; and cj1310, cj1318 and cj1335), there was only a limited number of sequence differences and these were examined in the trace files to confirm locus specificity. Due to the short length of some of these PCR products, tract length was validated by PCR amplification using primers located in non-identical flanking sequences and sequence analysis of the whole PCR fragment. In all cases, products were obtained from the expected locus and the observed repeat number matched the predictions from the 28 locus PV-analysis assay (S1 Table).

Development of multiplex PCR and GeneScan assays

Primers for multiple loci were combined in order to reduce the number of required PCRs (Table 2). Initially five primer mixes were designed to amplify five or six loci each. Primers for genes with high sequence identity or using a common primer were assigned to different reactions in order to prevent false-priming or competition between amplifications. Small differences in PCR efficiency were accommodated by setting the minimum peak height for detection of fragments by PSAnalyse below the height for the least efficient PCR reaction. Four of the primer mixes generated the expected products at similar efficiencies. Reaction mix C exhibited low PCR efficiencies for some loci. PCR efficiency was improved by splitting mix C into two reactions (C1 and C2) of three genes.

PCR products from the six PCR reactions were combined into a single sample and arrayed on a 96-well PCR plate. Two types of controls were included on this plate. A set of PCR products amplified from a genomic DNA preparation with known repeat numbers (i.e. determined by dideoxy sequencing) and a previously prepared set of fluorescently-labelled products. The former controls for variation in the efficiencies of PCR reactions and the latter for plate-to-plate variations in the migration of products on the autosequencer. Samples were subject to electrophoresis on an autosequencer.

Development and implementation of a custom script for automated PV state analysis

Each well of the fragment analysis plate can be analysed and manually interrogated in PeakS-canner™ to extract PCR fragment size. Comparison of these values to the control data enables the repeat number and expression state to be determined. As this process is laborious and
subject to error, an automated programme, PSAnalyse, was developed to extract data from
PeakScanner™ output files. These data sets contain multiple, spurious peaks as a result of non-
specific primer binding, primer-dimers, and background ‘noise’ during detection. In order to
extract only the relevant information, the program was designed to search for fragments within
specific regions of the scans performed by the autosequencer. Thus the basic principle of PSA-
alyse is recognition of the highest peak of the correct colour (i.e. fluorescence) within a user-
defined range (e.g. 3 nucleotides plus or minus) of the expected size (pre-set values contained
in Peakset files). The features of this major peak and of the flanking peaks (size, height and area
under the peak) are extracted and compared (see Fig 1B for an example of major and flanking
peaks for a locus). Peaks that exceed user-defined cut-offs for height and ratio to flanking
peaks are accepted as the observed peak for that locus otherwise peaks are flagged as problem-
atic (allowing for the user to re-examine the PeakScanner™ trace file). The repeat number and
gene expression state for each locus are assigned based on size relative to control samples.

The ‘scan width’ setting enables detection of phase-variants that have shifted by a number
of nucleotides from the starting number for a given experiment. The polyG/polyc tracts of C.
jejuni strain NCTC11168 were observed to increase or decrease by one nucleotide during
examination of PV rates without selection and by up to 4 nucleotides during selection experi-
ments [8]. As discussed below, the scanning range can be widened but must prevent overlaps
between loci labelled with the same dye colour.

A common source of flagged errors for individual tracts was for the expected size of the
PCR fragment to exceed the limit by up to 0.2 bp (note the PeakScanner reports fragment sizes
to four decimal places). These errors arise due to differences in migration of fragments and
could be corrected by inspecting surrounding samples to determine if a sample was aberrant or
part of a trend for a series of samples. Another frequent observation was for differences
between odd and even wells in the first plate of a series when using the GS500LIZ size standard
(resulting in data flagged as an error due to observed peaks being 0.5 bp larger than control
peaks). This difference was attributed to warming of the plate and to use of a 48 capillary
machine so that odd wells of the first plate were injected whilst the plate was still cold. Inclu-
sion of control samples in both odd and even wells controlled for this difference. This problem
was eliminated in later analyses by switching to the GS600LIZ size standard, which is not sub-
ject to variations in fragment size. Another problematic observation was of the presence of
occasional spurious peaks in wells containing no samples. The majority of these peaks could be
eliminated by setting the minimum peak height to 1,000. This setting resulted in loss of data
from weak PCRs and recovery of this data had to be assessed against the potential for inaccu-
crate sizing of a small sub-set of peaks.

The expression states for each locus were based on an analysis of the whole genome
sequence for C. jejuni strain NCTC11168 (Table 1) [7]. For the genes with tracts located in the
reading frame, the ON expression state was determined as the repeat number associated with
an intact reading frame. In general, a single repeat number was provided in the peakset file as a
determinant of the ON state. In some cases, variants were observed that had switched by three
nucleotides to the next available ON state but were being reported as OFF by our early version
of PSAnalyse. Additional columns were added to the peakset files for the other expression
states and PSAnalyse was modified to search for these expression states or to treat as ON all
values differing from the given ON state by a multiple of three nucleotides/repeats.

An exception to our generalised approach to expression states was observed for the capA
gene. Ashgar et al. [15] had reported (and was confirmed by dideoxy sequencing) that variants
of NCTC11168 differed by one nucleotide within the A-tract, that is adjacent to the G-tract
within the capA reading frame, resulting in an alteration in the reading frame. Peakset files for
different variants were utilised with the ON repeat number for capA being either 10 or 11
nucleotides. Another exception is that the ON number of repeats was arbitrarily set for *cj0045* (the tract in this gene is located at the 3’ end of the gene), pseudogenes and intergenic repeats. Coding of some of the variation for these loci with the simplified binary code facilitated further analyses.

**Comparison of single colonies versus total DNA extracts for estimation of ON state proportions in a mixed population**

Previous investigations reporting ON/OFF states of individual PV genes by the fragment analysis method have analysed the relative ‘area under peak’ for each amplified PV site. In this approach, the DNA template was the total DNA extracted from the whole population of a given sample. This method is, however, subject to error due to slippage during PCR amplification of long homopolymeric tracts [20].

We compared how data generated from analysing individual colonies differed from analysis of total DNA extracts of a population. A set of 16 samples were selected for this analysis (these samples were isolated using selective media from *C. jejuni-*infected birds at 52 days post-inoculation; Lango-Scholey *et al.*, in preparation). DNA was extracted from large sweeps of confluent growth or >100 colonies (‘total’ DNA) and in addition from 30 individual colonies derived from dilutions of the same population. Tract lengths for individual colonies were assigned based on the major peak using PSAnalyse and then a % ON value was determined by dividing colonies with an ON repeat number by total number of colonies. For ‘total’ samples we calculated relative areas under peaks for tracts of different sizes and representing different ON and OFF states and then calculated the % ON value by dividing the area for ON peak(s) by the total area under all peaks (note that most samples contained one major peak and two minor peaks; see S1 File). The difference between the two analysis methods was assessed by subtracting the two values.

A set of 448 values for % ON (28 genes in 16 samples) was obtained by both the whole population (i.e. a sweep) and single colony analysis methods. Fig 2 shows the linear regression line (blue line) between % ON values for the two sampling methods. Divergence from the expected correlation (black line) exhibits two opposing trends, under- or over-estimation of %ON by the total colony method relative to the other method at high and low %ON states, respectively. The intersection between the lines for observed and expected values occurs at ~33%, which corresponds to the known relationship between ON and OFF states of 1:2 for intragenic SSR-mediated PV. Examination of values for individual genes (S1 Fig) did not show any consistent trends indicating that the differences are independent of gene function.

As an example, the total population method detected small peaks flanking the main peak of *cj0275* and predicted that ~9% of cells in these populations were in the OFF state (data not shown). However, the only repeat number detected in 390 single colonies was G8 (ON) and indeed no variants have been detected so far during analysis of >2,000 colonies isolated from infected birds (Lango-Scholey *et al.* in preparation). Similarly, for the 11 birds in which *cj1429* was mainly in an ON state (i.e. G10), the total population analysis predicted that 77% of colonies were in an ON state whereas 93% (279 out of 299) of the single colonies were in this state. These results suggest that the detection of peaks correlating with OFF repeat numbers in the total population analysis were generated by PCR slippage rather than representing actual phase variants. Thus, PCR slippage appears to lead to inaccurate assessments of the proportion of cells in an ON or OFF state when PCR and fragment analysis of total DNA preparations are utilised for assessment of PV states in a population. The greatest differences are observed in populations exceeding 99% ON or 99% OFF (with values ranging up to 25% lower).
Analysis of PCR slippage during investigation of repeat numbers in single colonies

In order to further investigate how repeat number influences slippage during PCR, we determined the areas for the main and flanking peaks in analyses of single colonies from a single 96-well plate (see S2 File). Single colonies are expected to have low numbers of phase variants as the previously measured PV rates indicate that phase variants would, on average, be generated at a frequency of \(<0.05\) per colony for tracts of G8 to G12 [8]. The area under the flanking peaks was found to increase from 14% with G8 tracts to 39% with G11 and to increase as a function of repeat number (Fig 3A). This correlation was consistent both between different PV loci (Fig 3B) and between different tract lengths (S2 Table) of the same gene (Two-way ANOVA, \(p<0.001\)). This indicates that repeat number but not flanking sequence is the main determinant of switching rate. As the switching frequencies were determined to be 0.004 for a G8 tract in \(cj1139\) and 0.02 for a G11 tract in \(capA\) [8], these results indicate that analysis of a mixed population will tend to overestimate the frequencies of phase variants by 20–35 fold for all repeat numbers.

General applicability of the PV analysis assay

In order to assess whether these primer sets would have utility for analysis of SSR in other \(C. jejuni\) strains, we performed a BLAST analysis of primer conservation in six additional genome sequences (S3 Table). The current primer sets would work or need minor adjustments on 4–12 phase-variable loci per genome whereas other genes would require new primer sets. Primer design is however relatively straightforward and could be rapidly extended to other strains.
The PSAnalyse program would work on any other similar data set and would not require any adaptation. The only requirement would be determination of repeat number for each locus (from individual sequences of each locus for a control DNA sample of the specific strain) and expression states (obtained by analysis of either a whole genome sequence or of individual gene sequences). This data set would enable derivation of a calibration file. The Peakset data could then be generated by running fragment analysis assays with the control DNA sample to determine the association between fragment size and repeat number.

In summary, the PV analysis assay and the computer program, PSAnalyse, described here have general applicability for determining repeat number and associated expression state in any organism wherein this phenomenon is observed.

Fig 3. The effect of repeat number on PCR slippage as detected by fragment analysis of individual colonies with an ON number of repeats. Fragment length analysis was performed for 11 repeat tracts on a set of single colonies (n = 90). The relative area under each peak (i.e. the flanking peaks and the major peak) was calculated by dividing the area under the peak by the total area under all peaks for that specific locus and sample. These values were multiplied by 100 to obtain percentage values for the relative area under each peak. (A) Dotplot of individual values for the major peak of each locus separated by tract length. The number of loci analysed for each tract length were as follows: G8, 257; G9, 214; G10, 431; G11, 77; G12, 9. (B) Average values for each locus. Peaks for each locus were separated by the numbers of repeats in the polyG tract of the major peak with -1 and +1 indicating flanking peaks having one less or one more G residue. Horizontal line, average for relative area under the peak; n, number of colonies.

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Supporting Information

S1 Fig. Comparison of single colony and total population analyses of the % ON state for individual genes of a *Campylobacter jejuni* strain NCTC11168 population. Bacterial populations were obtained by plating serial dilutions of caecal samples from chickens infected for 52 days with *C. jejuni* strain NCTC11168H (Lango-Scholey et al., unpublished data). The total population is a sweep obtained from a low dilution plate while single colonies were obtained from high dilution plates. The displayed values represent the difference between the % ON values for the total population (as determined from the fragment analysis by dividing the area of the peaks for an ON number of repeats by the total area under all peaks) and multiple colony analysis (as determined from an analysis of between 11 and 30 colonies with the number of colonies with an ON repeat number being divided by the total number of colonies analysed) as obtained for each gene from 16 individual birds. The top panel shows the genes in which the majority of the population was in the ON state while the bottom panel shows the majority OFF state genes. Average, mean %ON state for a gene. Tracts, repeat number of major peak observed in total population analysis (note that not all birds had the same tract length for each gene hence multiple values are obtained). Positive and negative values indicate that the % ON is higher or lower, respectively, in the 'total DNA' as compared to the single colony analysis. Pink, negative 16–33% difference; Red, >33% negative difference; Green, 16–33% positive difference; Blue, >33% positive difference.

(PPTX)

S1 File. Percentage ON state data for total population analyses.
(XLSX)

S2 File. Slippage data obtained from single colony analyses.
(XLSX)

S1 Table. Confirmation of predicted repeat numbers from fragment analysis by dideoxy sequencing of PCR products.
(DOCX)

S2 Table. Slippage in repeat tracts during single colony analyses.
(DOCX)

S3 Table. Conservation of primer bindings sites in selected *C. jejuni* strains.
(DOCX)

Author Contributions

Conceived and designed the experiments: CDB MAJ. Performed the experiments: LLS AW JA. Analyzed the data: LLS JA CDB. Contributed reagents/materials/analysis tools: JA. Wrote the paper: LLS CDB.

References

1. Levinson G, Gutman GA. Slipped-strand mispairing: a major mechanism for DNA sequence evolution. Molecular Biology and Evolution. 1987; 4: 203–21. PMID: 3328815
2. Moxon R, Bayliss C, Hood D. Bacterial contingency loci: the role of simple sequence DNA repeats in bacterial adaptation. Annual Review of Genetics. 2006; 40: 307–33. PMID: 17094739
3. Bayliss CD. Determinants of phase variation rate and the fitness implications of differing rates for bacterial pathogens and commensals. FEMS Microbiology Reviews. 2009; 33: 504–20. doi: 10.1111/j.1574-6976.2009.00162.x PMID: 19222587
4. Moxon ER, Rainey PB, Nowak MA, Lenski RE. Adaptive evolution of highly mutable loci in pathogenic bacteria. Current Biology: CB. 1994; 4: 24–33. PMID: 7922307
5. Hermans D, Van Deun K, Martel A, Van Immerseel F, Messens W, Heyndrickx M, et al. Colonization factors of Campylobacter jejuni in the chicken gut. Veterinary Research. 2011; 42: 82. doi: 10.1186/1297-9716-42-82 PMID: 21714866
6. Young KT, Davis LM, Dirtha VJ. Campylobacter jejuni: molecular biology and pathogenesis. Nature reviews Microbiology. 2007; 5: 665–79. PMID: 17703225
7. Parkhill J, Wren BW, Mungall K, Kettle JM, Churcher C, Basham D, et al. The genome sequence of the food-borne pathogen Campylobacter jejuni reveals hypervariable sequences. Nature. 2000; 403: 665–8. PMID: 10688204
8. Bayliss CD, Bidmos FA, Anjum A, Manchev VT, Richards RL, Grossier JP, et al. Phase variable genes of Campylobacter jejuni exhibit high mutation rates and specific mutational patterns but mutability is not the major determinant of population structure during host colonization. Nucleic Acids Research. 2012; 40: 5876–89. doi: 10.1093/nar/gks246 PMID: 22434884
9. Aidley J, Bayliss CD. Repetitive DNA: a major source of genetic diversity in Campylobacter populations? In: Sheppard SK, Méric G, editors. Campylobacter Ecology and Evolution. Portland: Caister Academic Press; 2014.
10. Kim JS, Artymovich KA, Hall DF, Smith EJ, Fulton R, Bell J, et al. Passage of Campylobacter jejuni through the chicken reservoir or mice promotes phase variation in contingency genes Cj0045 and Cj0170 that strongly associates with colonization and disease in a mouse model. Microbiology. 2012; 158: 1304–16. doi: 10.1099/mic.0.057158-0 PMID: 22343355
11. Karlyshev AV, Linton D, Gregson NA, Wren BW. A novel paralogous gene family involved in phase-variable flagella-mediated motility in Campylobacter jejuni. Microbiology. 2002; 148: 473–80. PMID: 11832511
12. Guerry P, Szymanski CM, Prendergast MM, Hickey TE, Ewing CP, Pattarini DL, et al. Phase variation of Campylobacter jejuni 81–176 lipooligosaccharide affects ganglioside mimicry and invasiveness in vitro. Infect Immun. 2002; 70: 787–93. PMID: 11796612
13. Linton D, Gilbert M, Hitchen PG, Dell A, Morris HR, Wakarchuk WW, et al. Phase variation of a beta-1,3 galactosyltransferase involved in generation of the ganglioside GM1-like lipo-oligosaccharide of Campylobacter jejuni. Molecular Microbiology. 2000; 37: 501–14. PMID: 10931344
14. Hitchen P, Brzostek J, Panico M, Butler JA, Morris HR, Dell A, et al. Modification of the Campylobacter jejuni flagellin glycan by the product of the Cj1295 homopolymeric-tract-containing gene. Microbiology. 2010; 156: 1953–62. doi: 10.1099/mic.0.038091-0 PMID: 20338909
15. Ashgar SS, Oldfield NJ, Wooldridge KG, Jones MA, Irving GJ, Turner DP, et al. CapA, an autotransporter protein of Campylobacter jejuni mediates association with human epithelial cells and colonization of the chicken gut. Journal of Bacteriology. 2007; 189: 1856–65. PMID: 17172331
16. van Alphen LB, Wuhrer M, Bleumink-Pluym NM, Hensbergen PJ, Deelder AM, van Putten JP. A functional Campylobacter jejuni maf4 gene results in novel glycoforms on flagellin and altered autoagglutination behaviour. Microbiology. 2008; 154: 3385–97. doi: 10.1099/mic.0.2008/019919-0 PMID: 18957592
17. Fox KL, Dowideit SJ, Erwin AL, Srikhanta YN, Smith AL, Jennings MP. Haemophilus influenzae phase-variants have evolved from type III DNA restriction systems into epigenetic regulators of gene expression. Nucleic Acids Research. 2007; 35: 5242–52. PMID: 17675301
18. Jerome JP, Bell JA, Plovanich-Jones AE, Barrick JE, Mansfield LS. Standing genetic variation in contingency loci drives the rapid adaptation of Campylobacter jejuni to a novel host. PloS One. 2011; 6: e16399. doi: 10.1371/journal.pone.0016399 PMID: 21283682
19. Wilson DL, Rathinam VA, Qi W, Wick LM, Landgraf J, Bell JA, et al. Genetic diversity in Campylobacter jejuni is associated with differential colonization of broiler chickens and C57BL/6J IL10-deficient mice. Microbiology. 2010; 156: 2046–57. doi: 10.1099/mic.0.035717-0 PMID: 20360176
20. Shinde D, Lai Y, Sun F, Arnheim N. Taq DNA polymerase slippage mutation rates measured by PCR and quasi-likelihood analysis: (CA/GT)n and (A/T)n microsatellites. Nucleic Acids Research. 2003; 31: 974–80. PMID: 12560493
21. Bidmos FA, Bayliss CD. Genomic and global approaches to unravelling how hypermutable sequences influence bacterial pathogenesis. Pathogens. 2014; 3: 164–84. doi: 10.3390/pathogens3010164 PMID: 25437613