MethPat: a tool for the analysis and visualisation of complex methylation patterns obtained by massively parallel sequencing

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Abstract

Background: DNA methylation at a gene promoter region has the potential to regulate gene transcription. Patterns of methylation over multiple CpG sites in a region are often complex and cell type specific, with the region showing multiple allelic patterns in a sample. This complexity is commonly obscured when DNA methylation data is summarised as an average percentage value for each CpG site (or aggregated across CpG sites). True representation of methylation patterns can only be fully characterised by clonal analysis. Deep sequencing provides the ability to investigate clonal DNA methylation patterns in unprecedented detail and scale, enabling the proper characterisation of the heterogeneity of methylation patterns. However, the sheer amount and complexity of sequencing data requires new synoptic approaches to visualise the distribution of allelic patterns.

Results: We have developed a new analysis and visualisation software tool “Methpat”, that extracts and displays clonal DNA methylation patterns from massively parallel sequencing data aligned using Bismark. Methpat was used to analyse multiplex bisulfite amplicon sequencing data on a range of CpG island targets across a panel of human cell lines and primary tissues. Methpat was able to represent the clonal diversity of epialleles analysed at specific gene promoter regions. We also used Methpat to describe epiallelic DNA methylation within the mitochondrial genome.

Conclusions: Methpat can summarise and visualise epiallelic DNA methylation results from targeted amplicon, massively parallel sequencing of bisulfite converted DNA in a compact and interpretable format. Unlike currently available tools, Methpat can visualise the diversity of epiallelic DNA methylation patterns in a sample.

Keywords: DNA methylation, software, visualization, bisulfite, targeted amplicon, epigenetics, epiallele
Background
In mammals, the predominant and most widely studied DNA methylation mark occurs at CpG dinucleotide (CpG) palindromic sequences [1]. The vast majority of methods that investigate DNA methylation utilise bisulfite treatment of genomic DNA followed by PCR amplification to distinguish methylated from unmethylated CpG sites [2–5]. Bisulfite treatment discriminates methylated from unmethylated cytosines by selectively reacting with unmethylated cytosines to generate uracil. During the subsequent first step of PCR amplification, the uracils are read as thymine. Conversely, methylated cytosines do not react with the bisulfite reagent and remain as cytosines after PCR amplification [6]. DNA methylation readouts at single sites employing bisulfite conversion become analogous to genotyping assays by detecting either a cytosine or thymidine at the C position of a CpG site and are interpreted as methylated or unmethylated cytosines respectively.

An epiallele refers to a distinct pattern of methylation, typically over a short genomic region [7, 8]. In addition to the methylation state given for each CpG site, the pattern of DNA methylation of all CpG sites across the epiallelic or clonal template can also be characterised [7]. Indeed, in terms of biological function, CpG methylation should be often considered in an allelic fashion over multiple adjacent CpG sites [9, 10].

However, currently most studies summarise data into average percentage values at each CpG site thus losing the positional pattern information of DNA methylation across each clonal template [9]. Analysis platforms such as the Illumina Infinium BeadArray [11], bisulfite pyrosequencing [12] and SEQUENOM™ EpiTYPER™ [13] use bisulfite mediated chemistry to discriminate the methylation state of CpG sites but summarise measurements into percentage values across each CpG site or region of interest. Percentage methylation described in most DNA methylation studies hides important pattern and positional information of DNA methylation with potential functional and regulatory relevance [7]. It is only with clonal sequencing approaches [1, 14, 15], whole genome bisulfite sequencing [16] or reduced representation bisulfite sequencing [17], that the methylation state of individual CpG sites within a genomic DNA template can be readily measured in a digital sense, as methylated or not, allele by allele.

Imprinted regions of the genome such as IGF2/H19 and MEST typically display two epialleles, where one is completely methylated and the other is unmethylated. The loss of imprinting at such loci leads to syndromic complications [18, 19]. Average DNA methylation across these loci are typically presented as 50 % methylation but the pattern of DNA methylation at each epiallele is lost [7].

Heterogeneous DNA methylation describes the phenomenon where different contiguous CpG sites have different levels of methylation. DNA methylation heterogeneity can arise in a variety of ways including but not limited to: (i) more than a single population of cells is analysed that differ in DNA methylation at the locus of interest, (ii) the locus of interest is imprinted i.e. two different epialleles are present in each cell or, (iii) the locus is inherently heterogeneous in its DNA methylation composition. It is only using clonal sequencing approaches with allelic outputs, high resolution melting (HRM) [7, 20], or a novel ligation mediated approach [10] that heterogeneous DNA methylation can be detected. It is also inferred by varying methylation at CpG sites e.g. from Pyrosequencing. Importantly, the number of methylated alleles can be substantially underestimated unless clonal approaches are used [20]. Clonal sequencing is currently the best method to investigate heterogeneous DNA methylation and the extent of epiallelic methylation patterns that exist within a single sample [15].

Until recently, it has been cost prohibitive to assess the complexity of methylation patterns, as large number of clones need to be individually sequenced to determine the extent of heterogeneous DNA methylation. As one clone represents a single epiallele, many tens to hundreds of clones need to be sequenced to gain a true representation of different epialleles in a sample. The introduction of massively parallel sequencing enables the sequencing of many thousands of DNA templates from multiple regions simultaneously providing a true representation of the diversity and extent of heterogeneous DNA methylation patterns derived from a given sample. However, as the number of clones sequenced increases, the ability to analyse and present this type of data then becomes a significant challenge, and at this time, there are very few software tools available to manage such data from massively parallel sequencing experiments [21, 22]. Some visualisation and analysis tools are available for Bisulfite Sanger Sequencing including BiQ Analyzer [23], MethVisual [24], QUMA [25], BISMA [26]. However, these tools do not scale up with massively parallel sequencing having been designed for Sanger sequencing. BiQ Analyser HiMod is a tool that enables visualisation of high throughput sequencing of 5-methylcytosine and other methyl-variant modifications [27] however, results are expressed in percentage methylation values masking allelic methylation patterns.

In this study, we have developed Methpat, a software tool which processes bisulfite sequencing data following
Bismark alignment [28] and summarises DNA methylation according to epiallelic methylation patterns. This software has been used to analyse multiplex bisulfite amplicon PCR coupled to massively parallel deep sequencing on a range of primary haematopoietic tissue samples and model cancer cell lines to observe the extent of heterogeneous DNA methylation. Methpat is also able to create publication-ready, compact visualisations of the summarised data showing heterogeneous DNA methylation patterns in a space efficient and comprehensible manner.

**Materials, methods and implementation**

Samples, library preparation, sequencing and sequence alignment. Details of sample preparation, library generation, sequencing and sequence alignment protocol employed are summarised in the Additional file 1. Human samples used in this study were approved for

| Sample          | Mapping Efficiency | Unique Hits | Methylated CpG | Methylated CHG | Methylated CHH | Total C's analysed |
|-----------------|--------------------|-------------|----------------|----------------|----------------|--------------------|
| 293             | 52.2 %             | 7539        | 64.9 %         | 0.2 %          | 0.3 %          | 316211             |
| 40424           | 55.3 %             | 9414        | 37.5 %         | 0.2 %          | 0.2 %          | 351086             |
| 910046          | 42.0 %             | 7060        | 32.6 %         | 0.2 %          | 0.3 %          | 299795             |
| 12a-cd19        | 14.9 %             | 48648       | 47.9 %         | 0.4 %          | 0.5 %          | 1933767            |
| 12a-cd34        | 30.3 %             | 85049       | 36.5 %         | 0.1 %          | 0.2 %          | 3703147            |
| 12a-cd45        | 32.4 %             | 109173      | 32.6 %         | 0.1 %          | 0.2 %          | 4714744            |
| 12acd33         | 36.2 %             | 161885      | 32.8 %         | 0.2 %          | 0.2 %          | 6997070            |
| 6-mdar53        | 54.6 %             | 201660      | 84.4 %         | 0.8 %          | 1.3 %          | 9179816            |
| 6-c-d19         | 7.9 %              | 22258       | 77.8 %         | 0.2 %          | 0.3 %          | 777739             |
| 6-c-d33         | 27.9 %             | 20071       | 35.2 %         | 0.2 %          | 0.2 %          | 851116             |
| 6-c-d34         | 19.5 %             | 36928       | 49.7 %         | 0.2 %          | 0.2 %          | 1628107            |
| 6c-d45          | 33.0 %             | 31087       | 39.5 %         | 0.1 %          | 0.2 %          | 1314281            |
| 9a-cd19         | 21.2 %             | 39352       | 48.7 %         | 0.2 %          | 0.3 %          | 1638757            |
| 9a-cd33         | 31.9 %             | 125884      | 35.8 %         | 0.2 %          | 0.2 %          | 5459419            |
| 9a-cd34         | 26.2 %             | 77870       | 43.4 %         | 0.2 %          | 0.2 %          | 3321993            |
| 9a-cd45         | 46.6 %             | 28085       | 29.8 %         | 0.2 %          | 0.2 %          | 1211803            |
| 9awholeblood    | 31.5 %             | 97532       | 30.8 %         | 0.2 %          | 0.2 %          | 4081834            |
| bclf            | 49.3 %             | 9107        | 32.7 %         | 0.2 %          | 0.4 %          | 398977             |
| caco            | 19.6 %             | 129536      | 78.1 %         | 0.2 %          | 0.2 %          | 4512574            |
| dg75            | 51.7 %             | 10827       | 57.2 %         | 0.3 %          | 0.3 %          | 489096             |
| ekvax           | 23.0 %             | 115915      | 63.1 %         | 0.2 %          | 0.2 %          | 4494359            |
| hela            | 43.1 %             | 41650       | 55.9 %         | 0.2 %          | 0.2 %          | 1731811            |
| hepg2           | 39.2 %             | 24667       | 63.4 %         | 0.3 %          | 0.3 %          | 971693             |
| ht1080          | 40.7 %             | 4586        | 67.0 %         | 0.2 %          | 0.4 %          | 176188             |
| htab2-col       | 30.9 %             | 45576       | 79.9 %         | 0.2 %          | 0.2 %          | 1863098            |
| jvl             | 31.3 %             | 18814       | 42.7 %         | 0.2 %          | 0.2 %          | 771188             |
| k562            | 49.7 %             | 144791      | 55.5 %         | 0.3 %          | 0.3 %          | 6230391            |
| ls174t          | 41.2 %             | 3691        | 57.2 %         | 0.2 %          | 0.3 %          | 151722             |
| mcf7            | 30.0 %             | 87404       | 71.6 %         | 0.8 %          | 0.8 %          | 3786412            |
| mda-mb231-bag   | 29.0 %             | 94811       | 77.3 %         | 1.0 %          | 1.1 %          | 4171147            |
| nalm6           | 43.6 %             | 37669       | 85.8 %         | 0.2 %          | 0.2 %          | 1569041            |
| nccit           | 44.0 %             | 31656       | 45.7 %         | 0.4 %          | 0.3 %          | 1406165            |
| ovca8           | 32.3 %             | 46864       | 63.4 %         | 0.3 %          | 0.3 %          | 1917527            |
| sknas           | 21.6 %             | 275040      | 27.7 %         | 0.1 %          | 0.2 %          | 11313285           |
| u231            | 14.0 %             | 123302      | 74.8 %         | 0.4 %          | 0.2 %          | 4389352            |
Fig. 1 (See legend on next page.)
Methpat—a tool to summarise epiallelic DNA methylation patterns

We have developed the software tool, Methpat, to summarise and visualise the resultant epiallelic DNA methylation patterns from multiplex bisulfite amplicon experiments. Source code is available on GitHub (http://bjpop.github.io/methpat/). Methpat takes the output from bismark_methylation_extractor and summarises the methylation state of each CpG site within each amplicon template sequenced. DNA methylation patterns are then counted and their abundance is summarised into a tab delimited text file amenable for further downstream statistical analyses. Methpat also outputs a standalone HTML file that provides a visualisation of the DNA methylation pattern of each amplicon of interest and a visual summary of their abundance in each sample. A range of visualisation settings are customisable so that the end-user can change the settings to facilitate interpretation of the data and generate publication-ready figures. These options include presenting pattern counts as a percentage of the total, as absolute count or log-scaled counts (Additional file 2: Figure S1). Patterns can be arranged in order either by count abundance or by DNA methylation state. Colours within the visualisation can also be modified (Additional file 3: Figure S2), and the image saved as a PNG file for presentation or publication.

Results

Methpat analysis of DNA methylation demonstrates a wide diversity of DNA methylation patterns

DNA methylation of FOXP3 in primary haematopoietic cells

The promoter region of FOXP3 was analysed for DNA methylation to validate the amplicon next generation sequencing, bioinformatics analysis and Methpat visualisation pipeline. Amplicons obtained from whole blood and subpopulations of cells from bone marrow were analysed from a single individual, from which, a diverse range of
Fig. 2 (See legend on next page.)
DNA methylation states and their abundance was observed. Analysis of whole blood showed that although the majority of epialleles were either completely methylated or completely unmethylated at CpG sites (Fig. 1), there were a diverse array of methylation patterns present (62 in total). This could reflect the cellular composition of whole blood, such that a number of cell types exist with a variable DNA methylation state at FOXP3. In contrast, DNA extracted from CD34, CD19 and CD33 positive subpopulations were found to be largely methylated at FOXP3. The CD45 positive compartment was unmethylated (Fig. 1). This was in line with previous investigations on similar sample types [31].

**Methpat can visualise imprinted loci**

The extent of DNA methylation at a known imprinted locus, **MEST**, was investigated. This locus also served as a PCR amplification bias control as the DNA methylation status of the MEST region can be precisely defined. The extent of DNA methylation at a known imprinted locus, **MEST**, was investigated. This locus also served as a PCR amplification bias control as the DNA methylation state of the MEST region is expected to be ~50% in normal whole blood, HeLa, HePG2, JWl, MCF7 and NCCIT. In contrast, Caco was hypermethylated at this locus. Interestingly, in wildtype whole blood and the cell lines HEpg2, JWl, and NCCIT, the completely methylated epiallele could be observed but was at very low abundance compared to the unmethylated epiallele (Fig. 4). We confirmed that these alleles did not arise from incomplete bisulfite conversion artefacts as all non-CpG cytosines were converted to thymidine.

**Methpat visualisation of mitochondrial genome DNA methylation**

Bisulfite amplicon primers to the mitochondrial DNA D-loop regulatory sequence were included in the analysis to determine the DNA methylation state of the mitochondrial genome. The predominant epiallele was found to be unmethylated across most samples analysed; however, there was a significant range in the abundance of epialleles with variable DNA methylation state across all samples (Fig. 5, Additional file 7: Figure S5), suggesting that DNA methylation of the mitochondrial genome was present [37] but appeared to be independent of the disease status of the sample. This is in keeping with recent observations of mitochondrial genomic DNA methylation in human cells [38, 39]. We again confirmed that these alleles did not arise from incomplete bisulfite conversion artefacts as all non-CpG cytosines were converted to thymidine.

**Discussion**

Most studies investigating DNA methylation using conventional sequencing approaches represent DNA methylation into percentage values at each CpG site and in turn, do not show important positional information encoded within the epiallelic DNA methylation patterns. A comparison of features between methylation visualisation tools is summarised in Table 2. We have developed a new software tool called Methpat that processes output files from Bismark to visualise DNA methylation sequencing data by epialleles. Methpat facilitates visualisation of high throughput sequencing data after Bismark analysis and does not attempt to determine the success of a particular experiment. This is left to the
Fig. 3 Methpat visualisation of DNA methylation at the RASSF1A gene promoter region. Methylation of RASSF1A is present in cancer cell lines (Caco, HEPG2 and NALM6) with the exception of HeLa. Examples of RASSF1A methylation in whole blood and a normal lymphoblast cell line (JWL) are also shown.
**Fig. 4** Methpat visualisation of DNA methylation at the *CDKN2A* gene promoter region

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investigator to interpret the metrics from Bismark prior to Methpat visualisation. We demonstrate the utility of Methpat by examining the DNA methylation pattern abundance and epiallelic DNA methylation states that are lost when DNA methylation is summarised as percentage DNA methylation.

Methpat operates on Bismark output files and further summarizes this data into an interactive visualization that can be quickly interpreted within a web-browser. It can be executed locally to generate an HTML file which can be hosted remotely through the Internet or visualized locally on the most common web browsers (Chrome, Safari, Firefox, Internet Explorer). This feature which is unique to Methpat, is a major advantage. At this stage, Methpat does not have capability as a “genome-browser” to look at DNA methylation patterns at a genome-scale because it was designed for targeted deep sequencing of amplicons, however, we have made the source code available for further development by the research community to further improve Methpat (http://bjpop.github.io/methpat/).

We demonstrated the importance of calculating epiallelic abundance on the imprinted locus MEST, where we showed two predominant populations of epiallelic DNA methylation patterns, one completely methylated and the other completely unmethylated. Such patterns cannot be interpreted with percentage values at each CpG site as heterogeneous DNA methylation or, a sample containing a heterogeneous population of cells with variable DNA methylation states could give rise to the same percentage value [7]. Using Methpat to visualise the diversity of epialleles enables the inference at least of the existence of heterogeneous DNA methylation, or, the detection of heterogeneous populations of cells as demonstrated by investigating FOXP3 in whole blood and subpopulations of the haematopoietic compartment.

Of interest, in some model cancer cell lines, we observed a wide and diverse range of methylated epialleles. Having ruled out to the best of our ability any bisulfite conversion or PCR amplification artefacts, our results suggest that even within apparently homogeneous
Table 2 Alternative DNA methylation Analysis and Visualisation Tools

| Software     | Program Language and Implementation | Analysis Process | Visual Output                                                                 | Input file                                      | Output file                                                                 | Epiallelic Counts | Experiment Quality Check |
|--------------|------------------------------------|------------------|-------------------------------------------------------------------------------|------------------------------------------------|------------------------------------------------------------------------------|--------------------|--------------------------|
| Methpat      | Python, pip install, URL available to install files locally | Summarises Bismark output | Interactive HTML and summary text file of epiallele counts, Scalable PNG file  | Bismark methylation extractor output, user-defined BED format file            | HTML and tab delimited text file                                            | Yes                | No, leverages Bismark    |
| Bismark      | command line, Python, requires bwa | Performs alignment to bisulfite reference genome | None, generates BAM files for visualisation with SeqMonk or IGV                 | fastq file                                             | BAM and tab delimited text files                                            | No                 | Yes calculates C to T conversion |
| BSPAT        | Java/JS web interface              | Visualisation and summarisation of Bismark output | PNG file and UCSC Genome Browser file                                           | Bismark output, fastq files                           | Text file summary, PNG and UCSC Genome Browser BED file                      | Yes                | No                       |
| MPFE         | R library, Bioconductor            | Calculates probabilities that epialleles are true | R image outputs                                                               | Table of read counts from bisulfite sequencing data | Derived statistics and plots                                                | Yes                | Yes                      |
| Methylation plotter | R library, shiny interactive web application | Visualises beta DNA methylation values | Interactive webpage with setting options to adjust a static image of DNA methylation values for each sample. PNG and PDF output. | Text file containing matrix of sample vs beta value at each CpG of interest | PDF and PNG image file                                                      | No                 | No                       |
| RnBeads      | R library, Bioconductor            | Processes summary data from other software for visualisation | Interactive HTML and UCSC Genome browser track hub files. PNG files             | BED file                                               | HTML summary                                                               | No                 | Yes                      |
| coMET        | R library, Webserver for analysis  | For EWAS studies. Analyses derived matrix files | Image files of plots with genomic locations.                                   | Text matrix files                                       | Image files                                                               | No                 | No                       |

EWAS epigenome-wide association studies using Illumina Infinium HM450 BeadArrays
cell lines, the methylation state at a subset of gene promoters analysed is heterogeneous. This could be due to the nature of cell culture where the phenomenon of increasing DNA methylation is observed with increasing passage [40, 41], plasticity, or the setting of epigenetic memory of a sub-population of cells in the culture [42]. The detection of completely methylated epialleles of the CDKN2A gene promoter in whole blood and in other samples interrogated supports the validity of our approach, and indicates that Methpat provides a new tool to enable the detection of low level DNA methylation [43, 44]. The functional and biological implications of our current findings remain unclear, however, further investigation with appropriate specimens using Methpat is warranted.

We investigated mitochondrial DNA methylation and believe our analysis is one of the first accounts of characterising epiallelic DNA methylation within the D-loop regulatory region of the mitochondrial genome. Our study confirms observations of DNA methylation within the mitochondria [37–39]. Given there can be many thousands of copies of the mitochondrial genome per cell, it is not possible at this stage to determine the providence of the methylation states we have identified. The issue of heteroplasmy for mutations in the mitochondrial genome [45] apply for DNA methylation and techniques to address heteroplasmy could be applied to investigate DNA methylation within the mitochondrial genome further [46]. By visualising DNA methylation patterns within the mitochondrial genome, Methpat can facilitate insight towards new biomarkers of disease [47].

While our current strategy and experimental results are unable to resolve PCR amplification artefacts (over-representation of particular sequence reads because of amplification), incorporation of unique molecular identifiers [48] could resolve this in future studies.

Conclusions
In summary, we demonstrate the feasibility of multiplex bisulfite amplicon deep sequencing to identify the extent of DNA methylation epialleles in a range of human samples. We have developed a software tool, called Methpat, which enables the summarisation and visualisation of DNA methylation sequencing data in the context of epiallelic information.

Availability of data and materials
The raw amplicon sequencing data, Bismark alignments and Methpat output files associated with this manuscript have been published with the DOI 10.1186/s13742-015-0098-x.

Methpat software can be obtained from this URL: (http://bjpop.github.io/methpat/)

Additional files

- **Additional file 1:** Sample preparation, library preparation and sequencing methods. (DOCX 132 kb)
- **Additional file 2:** Figure S1. Example of a screenshot of Methpat visualisation. A. Epiallele representation of the patterns of DNA methylation for respective amplicon in respective sample. B. Count histogram, the abundance of each epiallele represented in A. C. Genomic co-ordinate and position of CpG of interest. D. Proportion of DNA methylation at each CpG position. E. Save button, export visualisation as PNG file. F. Amplicon of interest. G. Legend depicting DNA methylation status. (PNG 202 kb)
- **Additional file 3:** Figure S2. Example of a screenshot of the settings page for each Methpat visualisation. A number of parameters can be changed and the visualisation replotted for ease of interpretation. (PNG 92 kb)
- **Additional file 4:** Figure S3. IGV screenshot of two amplicon regions used in this study that target DNA sequences with no CpG sites within the RANBP17 locus. Therefore it is expected that all cytosines within this region of interest are completely converted by bisulfite treatment. This is shown here for MCF7 and MDA-MB-231-BAG. (PNG 135 kb)
- **Additional file 5:** Table S2. Bisulfite PCR primers used in this study. (XLS 15 kb)
- **Additional file 6:** Figure S4. Diverse and wide ranging epiallelic DNA methylation patterns of RASSF1A in MCF7 and NCCIT model cancer cell lines. (PNG 434 kb)
- **Additional file 7:** Figure S5. Epiallelic DNA methylation patterns of the D-loop regulatory region of the mitochondrial genome. (PNG 163 kb)
- **Additional file 8:** Table S1. Human Samples used in this study. (XLS 8 kb)
- **Additional file 9:** Table S3. Amplicon details required for Methpat input (hg19 coordinates). (XLS 9 kb)
- **Additional file 10:** Description of Methpat options. (DOCX 118 kb)

Competing interests
XZ is a salaried employee of BioInfoRx Inc. MP is a salaried employee of BioResearch Software Consultants. NW is currently a salaried employee of Pacific Edge Biotechnology Limited however, performed this work prior to joining Pacific Edge. Next generation sequencing reagents used in this study were kindly supplied by Illumina Australia Pty Ltd as part of their MiSeq Pilot Sequencing Grant Program.

Authors’ contributions
NCW designed the study, performed the experiments, analysed the data and wrote the paper, BIP developed the software and wrote the paper, ILC conceptualised the study, designed the study and wrote the paper, SQW designed the study, performed initial pilot experiments and wrote the paper, THM designed the study, performed initial pilot experiments and wrote the paper, XZ analysed the data and created the pilot visualisation software and wrote the paper, MP analysed the data and created the pilot visualisation software and wrote the paper, SE performed the experiments, analysed the data and wrote the paper, SRD performed the experiments, analysed the data and wrote the paper, AD conceptualised the study, designed the study, analysed the data and wrote the paper. All authors read and approved the final manuscript.

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