BiQ Analyzer HiMod: an interactive software tool for high-throughput locus-specific analysis of 5-methylcytosine and its oxidized derivatives

Daniel Becker^{1,2}, Pavlo Lutsik^{1,*}, Peter Ebert², Christoph Bock^{2,3,4}, Thomas Lengauer² and Jörn Walter^{1,*}

¹Department of Genetics, Saarland University, Saarbrücken 66123, Germany, ²Department of Computational Biology and Applied Algorithms, Max-Planck Institute for Informatics, Saarbrücken 66123, Germany, ³CeMM Research Center for Molecular Medicine of the Austrian Academy of Sciences, Vienna 1090, Austria and ⁴Department of Laboratory Medicine, Medical University of Vienna, Vienna 1090, Austria

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ABSTRACT

Recent data suggest important biological roles for oxidative modifications of methylated cytosines, specifically hydroxymethylation, formylation and carboxylation. Several assays are now available for profiling these DNA modifications genome-wide as well as in targeted, locus-specific settings. Here we present BiQ Analyzer HiMod, a user-friendly software tool for sequence alignment, quality control and initial analysis of locus-specific DNA modification data. The software supports four different assay types, and it leads the user from raw sequence reads to DNA modification statistics and publicationquality plots. BiQ Analyzer HiMod combines wellestablished graphical user interface of its predecessor tool, BiQ Analyzer HT, with new and extended analysis modes. BiQ Analyzer HiMod also includes updates of the analysis workspace, an intuitive interface, a custom vector graphics engine and support of additional input and output data formats. The tool is freely available as a stand-alone installation package from http://biq-analyzer-himod.bioinf. mpi-inf.mpg.de/.

INTRODUCTION

DNA methylation is widely recognized as a key epigenetic mechanism playing a crucial role in development and disease (1,2). Sequencing of bisulfite-treated deoxyribonucleic acid (DNA) is the gold standard for base resolution mapping of DNA-methylation (3–5). Recently, oxidized derivatives of 5-methylcytosine (5mC) were identified that are assumed to have important biological function (6). Like

5mC, the oxidized derivatives occur predominantly at CpGdinucleotide cytosines (7). However, conventional bisulfite sequencing does not discriminate between different oxidized forms. Sodium bisulfite treatment converts unmodified cytosine, 5-carboxy (5caC) and 5-formylcytosyne (5fC) into deaminated forms, which are subsequently replaced by thymines during polymerase chainreaction (PCR) amplification, while methylated and hydroxymethylated (5hmC) cytosines remain unchanged (8). Several new methods have been suggested recently to overcome these limitations and transform the specific oxidative modifications into sequence-based signals. Oxidative bisulfite sequencing (oxBS-seq) involves an oxidation reaction to convert 5hmC to 5fC (9). A subsequent bisulfite reaction followed by the amplification step converts 5fC to thymine whereas the original 5mC is replaced by cytosines in PCR amplicons (Figure 1a). TET-assisted bisulfite sequencing (TAB-seq) utilizes the opposite scheme and applies an enzymatic oxidation by the TET protein to convert all 5mC to 5hmC and further to 5fC and 5caC, while the initially present 5hmC is protected by glycosylation prior to the bisulfite reaction (Supplementary Figure S1). The subsequent bisulfite treatment and PCR converts the initially present 5mC (as well as 5fC and 5caC) to thymine while the 5hmC is replaced with cytosines (10). Similar methods were developed for mapping 5fC and 5caC, such as formyl-chemically-assisted bisulfite sequencing (fCAB-seq) (11) (Figure 1b) and chemical-modificationassisted bisulfite sequencing (CAB-seq) (12) (Supplementary Figure S1). Following the modified bisulfite treatments of either oxBS-seq, TAB-seq, CAB-seq or fCAB-seq the DNA is amplified and sequenced. The abundance of any modification under inspection is estimated by comparing such sequencing results to data obtained by conventional bisulfite sequencing. Note that all modified bisulfite meth-

^{*}To whom correspondence should be addressed. Tel: +49 681 302 4367; Fax: +49 681 302 2703; Email: j.walter@mx.uni-saarland.de Correspondence may also be addressed to Pavlo Lutsik, Tel: +49 681 302 2881; Fax: +49 681 302 2703; Email: j.lutsik@mx.uni-saarland.de

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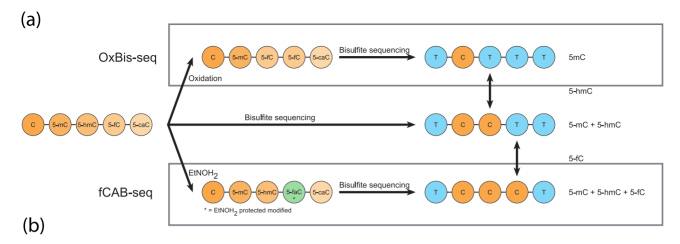


Figure 1. Principal scheme of oxBS-seq and fCAB-seq methods. (a) oxBS-seq. Oxidative bisulfite treatment followed by PCR leads to conversion of the oxidative modifications to thymines while only 5mC appears as cytosine in the sequencing reads. The cumulative level of 5hmC can be established for each CpG site by comparing to ordinary bisulfite sequencing. (b) In fCAB-seq 5fC is protected from the conversion and together with 5mC and 5hmC appears as cytosine after PCR. The bulk 5fC abundance is calculated by subtracting the cumulative levels of 5mC + 5hmC pair obtained from ordinary bisulfite sequencing.

ods are only partially modification-specific and mostly produce cumulative estimates of several different modifications.

While several tools have been developed for the analysis of conventional locus-specific bisulfite sequencing data of various scale (13–18), no software package currently supports an integrated analysis of data from multiplexed locus-specific high-throughput DNA modification profiling. For the genome-scale data the MLML tool has been designed to produce accurate DNA methylation and hydroxymethylation levels from preprocessed oxBS-seq, TAB-seq and other similar data (19).

To fill this gap we developed BiQ Analyzer HiMod (or shortly BiQ HiMod), a specialized interactive software package for the preprocessing, quality control and analysis of various DNA modifications in high-throughput targeted sequencing experiments. BiQ HiMod continues our line of computational tools for locus-specific DNA methylation analysis (14,15,18) and is an extension of the previously published BiQ Analyzer HT (14). It provides a userfriendly interface, comprehensive and customizable streamlined analysis pipeline, rich graphical and tabular output. The tool currently supports four experimental assays and can be easily extended to include the upcoming methods.

BiQ ANALYZER HiMod

Overview

BiQ Analyzer HiMod is a cross-platform interactive Java application, which can run on any system with properly installed and configured Java Runtime Environment. The package installation is performed via a simple 'click-through' installer. BiQ HiMod facilitates a fully interactive primary processing and analysis of sequencing data from locus-specific DNA modification profiling experiments, and currently supports oxBS-seq, TAB-seq, fCAB-seq and CAB-seq assays. The comprehensive and ergonomic graphical user interface provides means for multi-level overview of the complete multiplexed sequencing project, in which

a number of genomic loci of interest are sequenced in several biological samples. The underlying software architecture uses the fact that most of the basic data processing steps are invariant to the type of chemical treatment and thus the reads from ordinary and modified bisulfite procedure can be preprocessed independently. During the quality control cycle, the user refines the quality thresholds of the processing pipeline until a satisfactory quality level is reached for each read batch. After low-level processing, cumulative levels of each DNA modification are summarized based on the results of the individual preprocessing pipeline runs. Final results can be exported as tables, figures, alignments and genome browser tracks (see Supplementary text for a detailed list). Importantly, BiQ HiMod maintains the full functionality of the previous versions by supporting the analysis of conventional bisulfite sequencing data in an independent analysis mode.

Data preparation and project setup

BiQ HiMod is designed to import multiple data sets of individual sequencing reactions and loci. Today's data sets are often generated by NGS-based multiplexed locus-specific sequencing (16,23,24). In NGS approaches sequences are usually indexed by short sequence tags (using multiplexing library kits), pooled and submitted to a high-throughput sequencing platform, e.g. Illumina MiSeq or Ion Torrent. In such a setting BiQ HiMod requires the direct sequencing output to be demultiplexed using available third-party tools. We recommend the barcode splitter from Galaxy (20–22) as an adequate solution which suits researchers with minimal bioinformatics experience. Reads ready to be loaded must be in FASTA or FASTQ format with one file per each sample-locus-treatment combination.

Besides demultiplexing, data preparation usually includes several additional standard steps, e.g. trimming of low quality parts of the sequence reads (often at the 3'-end) and—in case of paired-end sequencing—joining of overlapping mate reads. In the FAQ section of the BiQ HiMod

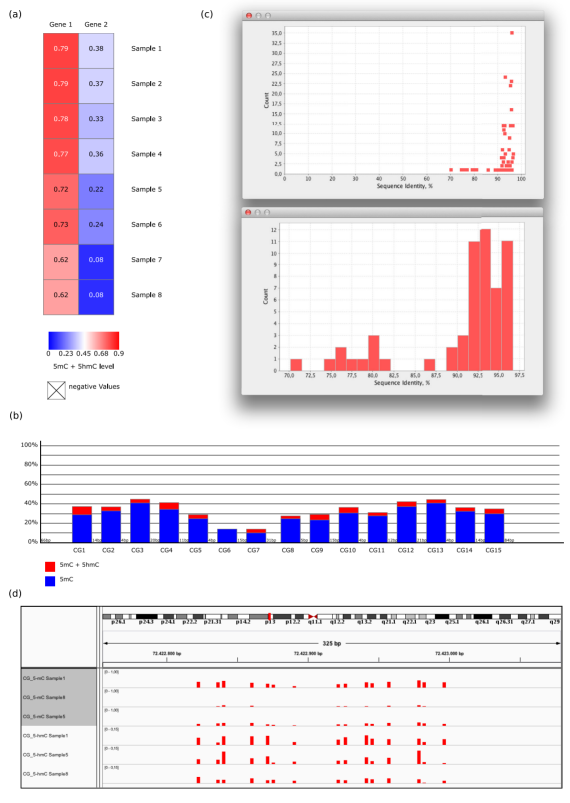


Figure 2. BiQ Analyzer HiMod visualization features. (a). Global heat map of average bulk (5mC + 5hmC) modification levels at sequenced loci across the samples. The default color code is given in the color legend (see Supplementary Figure S2 for more details). (b) Bar chart displaying stacked levels of two modifications at each CpG position in a single locus (see also Supplementary Figure S3). (c) Diagnostic histogram visualizing the distribution of a quality control metric — sequence identity of each read-reference alignment — simplifying the sequence identity cutoff selection. (d) Integration with the IGV genome browser showing genomic tracks with 5mC and 5hmC levels of three samples (the value range is much smaller for the 5hmC tracks).

Table 1. Information that can be extracted from each DNA modification profiling method supported by BiQ Analyzer HiMod

		Underlyin				
Method	Bisulfite treatment		Modified treatment		Measurements	
	C	T	\mathbf{C}	T		
oxBS-seq	5mC 5hmC	5fC 5caC C	5mC	5hmC 5fC 5caC C	5mC (SM) 5hmC (B) 5fC + 5caC + C (B)	
TAB-seq			5hmC	5mC 5fC 5caC C	5hmC (SM) 5mC (B) 5fC + 5caC + C (B)	
fCAB-seq			5mC 5hmC 5fC	5caC C	5fC (B) 5mC + 5hmC (B) 5caC + C (B)	
CAB-seq			5mC 5hmC 5caC	5fC C	5caC (B) 5mC + 5hmC (B) 5fC + C (B)	

SM—single-molecule resolution, B—bulk per-CpG measurement

Table 2. Benchmarking of BiQ Analyzer HiMod on artificially generated read batches of variable size

Number. of reads in each batch ^a		1000	2000	5000	10 000	20 000	50 000	
Running time, min.sec	Normal	0.15	0.32	1.52	5.20	16.48	155.15	
	w/o pattern maps ^b	0.15	0.26	1.06	2.13	4.25	25.14	

^aGiven are numbers of reads in each of two read batches (oxBS and conventional bisulfite) processed in every benchmarking step. Therefore, the actual number of processed reads was twice the given number in each step.

web site (http://biq-analyzer-himod.bioinf.mpi-inf.mpg.de/FAQ.php) we describe an example of such a data preparation workflow in a step-by-step way using third-party tools integrated into the Galaxy framework (20–22).

Following the data preparation one needs to set up an analysis project and import the data. For a relatively small project this can be done directly using the GUI controls. For larger data sets one should prepare a spreadsheet-like text file, (see our example on the program web site). Samples are defined by text labels, while the genomic loci exhibit one-to-one mapping to their genomic reference sequences, supplied as single- or multi-sequence FASTA files. We recommend extracting sequence information via the Galaxy 'Extract Sequence' tool starting from a BED file with genomic coordinates of each amplicon, such that the generated FASTA records contain genomic location information. This information will be used further on, in order to bring the analysis into the genomic context and generate genome browser tracks.

Finally, sequencing reads can be imported directly from the genome-wide sequencing experiment (e.g. wholegenome or reduced-representation bisulfite sequencing), if the data are available as aligned sequencing reads (BAM files). BiQ HiMod facilitates loading the reads of a corre-

sponding region based on coordinate information and analyzing them in the context of the corresponding genomic reference. This feature of BiQ HiMod can be useful for quickly scanning locus-specific read distributions in regions of interest based on large genome-wide WGBS or RRBS datasets provided they are accessible as BAM files.

Primary processing pipeline

The software backend was subjected to several modifications compared to the predecessor version of the tool. The FASTQ files parsing is now supported using the data import library of the BioJava project (23). Loaded sequence reads are independently aligned to the corresponding reference sequence using the bisulfite-modified semi-global implementation of Needleman–Wunsch algorithm, which was further tweaked to improve robustness of the modification site alignment (see Supplementary Text and (14) for more details). The set of filtering criteria was extended to include sequencing quality scores, and default thresholds are set to ensure that only high quality bases are considered. The full list of quality metrics and DNA modification statistics are given in the Supplementary Text. All processing and analysis options remain fully customizable and can be refined after the initial analysis.

^bThe high computation time for large read sets is based on the size of the produced single read-resolution pattern maps. The user can decide which graphics to generate and by choosing to disable pattern maps the computation time can be significantly reduced.

Quantifying the modification levels

A quantitative assessment of the relative abundances of different modifications depends on the comparison of two independent sequencing results. This comparative mode is a core feature implemented in BiQ Analyzer HiMod. However, such comparisons have two main limitations: (i) except for one modification type one can obtain estimates of modification differences at single CpG positions only as an average across all sequences and not per sequence (ii) in most comparisons combinations of several modification types are not separable from each other. For illustration, we consider the results of the oxBS-seq method. Here, the reads from the standard bisulfite treatment contain unconverted cytosines which were methylated or hydroxymethylated in the original DNA molecule. The read cytosines from the oxidative bisulfite treatment can only be mapped to the 5mC of the original DNA. All in all, oxBS-seq yields a single-molecule resolution map of 5mC, bulk per-CpG average levels of 5hmC and a combination of 5fC, 5caC and unmodified cytosines. Table 1 summarizes the information which can be extracted from each of the data types. For every type of analysis BiO HiMod takes into account all potential modifications and summarizes the information accordingly.

The estimation of bulk modification levels in BiQ Analyzer HiMod is performed by a simple subtraction of converted cytosine frequencies. In case of low abundance of the target modification, in particular 5fC and 5caC, the true biological changes of modification levels may overlap with the experimental and the technical error range, sometimes leading to negative DNA modification levels. To overcome such difficulties, on the experimental side we recommend gathering data from several independent experiments using very high sequencing coverage. Moreover, the statistical significance of the observed variation should be tested. Summarized data can be exported in bedGraph format and loaded into downstream statistical software, e.g. MLML or Rn-Beads (http://rnbeads.mpi-inf.mpg.de), for correction and proper statistical evaluation.

Visualization and data export

The frontend of BiQ HiMod provides capabilities for the top-down level-by-level exploration of a sequencing experiment. On the top level the global overview of the analysis project is given by the samples versus loci heat maps visualizing average modification abundances for each samplelocus pair (Figure 2a). More detailed profiles of each studied locus are provided by the locus-wide bar plots visualizing the modification levels at each CpG position with realistic relative genomic distances (Figure 2b). On the bottom level, sequence pileup, read-level cytosine pattern map and data tables are available for the most in-depth exploration and quality control similar to the analysis mode of BiQ Analyzer HT. The newly introduced diagnostic plots simplify the selection of the quality control thresholds by visualizing read batch-wide distributions of major analysis metrics, e.g. score and sequence identity of the read-reference alignment, bisulfite conversion rate, number of missing or mutated modification sites etc. (Figure 2c).

Read-level and summarized modification data can be exported as tab-delimited files for downstream statistical analysis and visualization in favorite statistical software packages, e.g. R/Bioconductor, SPSS and Excel, or custom bioinformatic pipelines. Furthermore, BiQ HiMod feeds into large-scale experimental projects by exporting its results as genome browser tracks. Methylation values for each analyzed CpG-site can be saved in bedGraph format importable into genome viewers and browsers, such as IGV (24) or UCSC Genome Browser (25) (http://genome.ucsc. edu/index.html) (see example in Figure 2d). This allows for a convenient display of amplicon validation data in the context of genome-wide epigenetic data tracks, such as DNA methylation, histone modification, chromatin accessibility maps etc. Finally, the summarized data in bedGraph format can be loaded into our in-house genome-wide DNA methylation data analysis package RnBeads (http://rnbeads.mpiinf.mpg.de/) for statistically rigorous evaluation and comparison to corresponding genome-scale profiles in case such are available.

Software architecture, GUI improvements and the new graphics engine

BiQ HiMod builds upon the modular and robust design of its predecessor tool, BiQ Analyzer HT, with a multistep data processing pipeline, as its backend, combined with an interactive graphical user interface at the frontend. The backend takes raw sequencing reads and reference sequence information as input, performs a series of preprocessing steps—data loading, alignment, quality filtering, cytosine conversion calling—and generates the resulting tables and graphics for each sample-locus combination. The backend output is written directly to the hard drive and can be found in the project directory. This results in efficient use of the operating memory and guards against information loss. The interactive frontend, implemented in Java standard AWT/Swing framework, enables the setup of a largescale multiplexed next-generation sequencing (NGS) analysis project for one of the supported profiling methods, coordination of the pipeline runs, and summarization of the results at the sample, locus and project levels as well as export of the results.

BiQ HiMod expands the basic form of the GUI design of the predecessor tool, BiQ Analyzer HT, by introducing several major improvements. First, the project summary view has been turned into an ultimate control panel, affording an exhaustive overview without the need of frequent panel switching. It includes a project summary table, presenting the most crucial information about each read batch, the project-level heat maps visualizing average modification levels and locus-wide bar charts dynamically generated for each selected read batch. Second, the detailed views of individual read batches were updated and now include the summary pages presenting the levels of each modification or group of modifications as tables and plots.

BiQ HiMod features a completely new custom graphical engine generating high-quality vector images in scalable vector graphics (SVG) format. Most of the plotting parameters, including the flexible color scheme, are fully useradjustable. The exported SVG files can be further edited via one of the popular image packages without any loss of quality or directly incorporated into manuscripts and reports.

VALIDATION ON ARTIFICIAL AND REAL BIOLOGI-CAL DATA AND PERFORMANCE ASSESSMENT

In order to test BiQ HiMod and assess its performance we generated artificial read batches for each supported type of experiment by simulation (see Supplementary Text 1 for details). We estimated the accuracy and robustness of BiQ HiMod results under different conditions.

We then applied BiQ HiMod to reprocess the earlier published oxBS-seq data set obtained by sequencing of repetitive elements in two types of mouse ES cells (26). In brief, serum cultured murine E14 ES cells and 2i-medium cultured ES cells from three different time points—1, 3 and 7 days of cultivation—were sampled, two replicates of each. After oxidative and conventional bisulfite treatment, hairpin amplicons of regions within two repeat types—intracisternal A particle, also known as IAP, and long interspersed elements, also known as LINE-1 or L1—were constructed as described in an earlier study (27). The amplicons were sequenced on the MiSeq platform from Illumina using paired end sequencing (2 \times 150). The BiO HiMod results document an accumulation of 5hmC at LINE1 but not at IAP elements during the first 72 h of cultivation in 2i-medium (Supplementary Figure S7).

We also benchmarked the runtime of BiQ Analyzer Hi-Mod by analyzing batches of 1000 to 50 000 artificially generated reads. The tests were carried out on a machine with a 2.4 GHz Intel Core i5 Haswell dual core processor and 2 GB of RAM. The results of the benchmark, given in Table 2, show that moderate-sized analyzes can be performed in a reasonable amount of time on an average commodity laptop. For data from large-scale experiments amounting to several millions or more sequence reads, we recommend running BiQ HiMod on more powerful workstation computers with several multicore CPUs and a few dozen GB of RAM, where the program can benefit of its parallel computation capabilities.

CONCLUSIONS AND OUTLOOK

BiO Analyzer HiMod is the first interactive software package for preprocessing, quality control and initial analysis of various DNA modifications from four different experimental methods. Developed on the basis of BiQ Analyzer and BiQ Analyzer HT (14,15), it features a thoroughly reworked GUI with a large number of analysis plots as well as a new customizable vector graphics engine, additional data import and export formats. The tool enables a comprehensive analysis workflow starting from raw sequence reads up to publication-quality plots and genome browser tracks. BiQ HiMod exploits modular and flexible software design allowing for painless further extensions to support upcoming experimental assays, such as MAB-seq (28). The standardized, comprehensive and user-friendly software paradigm implemented by BiQ HiMod will help the tool find numerous applications in basic biology and biomedical research.

AVAILABILITY

The BiQ Analyzer HiMod installation package is freely available from http://biq-analyzer-himod.bioinf.mpi-inf. mpg.de/.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online including [1,2].

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REFERENCES

- 1. Smith, Z.D. and Meissner, A. (2013) DNA methylation: roles in mammalian development. *Nat. Rev. Genet.*, **14**, 204–220.
- 2. Bergman, Y. and Cedar, H. (2013) DNA methylation dynamics in health and disease (vol 20, pg 274, 2013). *Nat. Struct. Mol. Biol.*, **20**, 1236–1236
- Harris, R.A., Wang, T., Coarfa, C., Nagarajan, R.P., Hong, C., Downey, S.L., Johnson, B.E., Fouse, S.D., Delaney, A., Zhao, Y. et al. (2010) Comparison of sequencing-based methods to profile DNA methylation and identification of monoallelic epigenetic modifications. *Nat. Biotechnol.*, 28, U1097–U1194.
- Bock, C., Tomazou, E.M., Brinkman, A.B., Mueller, F., Simmer, F., Gu, H., Jaeger, N., Gnirke, A., Stunnenberg, H.G. and Meissner, A. (2010) Quantitative comparison of genome-wide DNA methylation mapping technologies. *Nat. Biotechnol.*, 28, U1106–U1196.
- 5. Bock, C. (2012) Analysing and interpreting DNA methylation data. *Nat. Rev. Genet.*, **13**, 705–719.
- Branco, M.R., Ficz, G. and Reik, W. (2012) Uncovering the role of 5-hydroxymethylcytosine in the epigenome. *Nat. Rev. Genet.*, 13, 7–13.
- Kohli, R.M. and Zhang, Y. (2013) TET enzymes, TDG and the dynamics of DNA demethylation. *Nature*, 502, 472–479.
- 8. Huang, Y., Pastor, W.A., Shen, Y., Tahiliani, M., Liu, D.R. and Rao, A. (2010) The behaviour of 5-hydroxymethylcytosine in bisulfite sequencing. *PLoS One*, **5**, e8888.
- 9. Booth, M.J., Branco, M.R., Ficz, G., Oxley, D., Krueger, F., Reik, W. and Balasubramanian, S. (2012) Quantitative sequencing of 5-methylcytosine and 5-hydroxymethylcytosine at single-base resolution. *Science*, 336, 934–937.
- Schuler, P. and Miller, A.K. (2012) Sequencing the sixth base (5-hydroxymethylcytosine): selective DNA oxidation enables base-pair resolution. *Angew. Chem., Int. Ed. Engl.*, 51, 10704–10707.
- Song, C.X., Szulwach, K.E., Dai, Q., Fu, Y., Mao, S.Q., Lin, L., Street, C., Li, Y., Poidevin, M., Wu, H. et al. (2013) Genome-wide profiling of 5-formylcytosine reveals its roles in epigenetic priming. Cell., 153, 678–691.
- 12. Lu,X., Song,C.X., Szulwach,K., Wang,Z., Weidenbacher,P., Jin,P. and He,C. (2013) Chemical modification-assisted bisulfite sequencing (CAB-Seq) for 5-carboxylcytosine detection in DNA. *J. Am. Chem. Soc.*, **135**, 9315–9317.
- Kumaki, Y., Oda, M. and Okano, M. (2008) QUMA: quantification tool for methylation analysis. *Nucleic Acids Res.*, 36, W170–W175.
- Lutsik, P., Feuerbach, L., Arand, J., Lengauer, T., Walter, J. and Bock, C. (2011) BiQ Analyzer HT: locus-specific analysis of DNA

- methylation by high-throughput bisulfite sequencing. Nucleic Acids Res., 39, W551-W556.
- 15. Bock, C., Reither, S., Mikeska, T., Paulsen, M., Walter, J. and Lengauer, T. (2005) BiO Analyzer: visualization and quality control for DNA methylation data from bisulfite sequencing. Bioinformatics, 21, 4067-4068.
- 16. Rohde, C., Zhang, Y., Reinhardt, R. and Jeltsch, A. (2010) BISMA-fast and accurate bisulfite sequencing data analysis of individual clones from unique and repetitive sequences. BMC Bioinformatics, 11, e230.
- 17. Hetzl, J., Foerster, A.M., Raidl, G. and Mittelsten Scheid, O. (2007) CyMATE: a new tool for methylation analysis of plant genomic DNA after bisulphite sequencing. Plant J., 51, 526–536.
- 18. Schuffler, P., Mikeska, T., Waha, A., Lengauer, T. and Bock, C. (2009) MethMarker: user-friendly design and optimization of gene-specific DNA methylation assays. Genome Biol, 10, eR105.
- 19. Qu,J., Zhou,M., Song,Q., Hong,E.E. and Smith,A.D. (2013) MLML: consistent simultaneous estimates of DNA methylation and hydroxymethylation. Bioinformatics, 29, 2645–2646.
- 20. Blankenberg, D., Von Kuster, G., Coraor, N., Ananda, G., Lazarus, R., Mangan, M., Nekrutenko, A. and Taylor, J. (2010) Galaxy: a web-based genome analysis tool for experimentalists. Curr. Protoc. Mol. Biol., Chapter 19, Unit 19 10 11-21.
- 21. Giardine, B., Riemer, C., Hardison, R.C., Burhans, R., Elnitski, L., Shah, P., Zhang, Y., Blankenberg, D., Albert, I., Taylor, J. et al. (2005) Galaxy: a platform for interactive large-scale genome analysis. Genome Res., 15, 1451-1455.
- 22. Goecks, J., Nekrutenko, A., Taylor, J. and Galaxy, T. (2010) Galaxy: a comprehensive approach for supporting accessible, reproducible, and

- transparent computational research in the life sciences. Genome Biol., 11 eR86
- 23. Holland, R.C., Down, T.A., Pocock, M., Prlic, A., Huen, D., James, K., Foisy, S., Drager, A., Yates, A., Heuer, M. et al. (2008) BioJava: an open-source framework for bioinformatics. Bioinformatics, 24, 2096-2097.
- 24. Robinson, J.T., Thorvaldsdottir, H., Winckler, W., Guttman, M., Lander, E.S., Getz, G. and Mesirov, J.P. (2011) Integrative genomics viewer. Nat. Biotechnol., 29, 24-26.
- 25. Kent, W.J., Sugnet, C.W., Furey, T.S., Roskin, K.M., Pringle, T.H., Zahler, A.M. and Haussler, D. (2002) The human genome browser at UCSC. Genome Res., 12, 996-1006.
- 26. Ficz, G., Hore, T.A., Santos, F., Lee, H.J., Dean, W., Arand, J., Krueger, F., Oxley, D., Paul, Y.-L., Walter, J. et al. (2013) FGF Signaling Inhibition in ESCs Drives Rapid Genome-wide Demethylation to the Epigenetic Ground State of Pluripotency. Cell Stem Cell. 13, 351-359.
- 27. Arand, J., Spieler, D., Karius, T., Branco, M.R., Meilinger, D., Meissner, A., Jenuwein, T., Xu, G., Leonhardt, H., Wolf, V. et al. (2012) In vivo control of CpG and non-CpG DNA methylation by DNA methyltransferases. PLoS Genet., 8, e1002750.
- 28. Hu, X., Zhang, L., Mao, S.Q., Li, Z., Chen, J., Zhang, R.R., Wu, H.P., Gao, J., Guo, F., Liu, W. et al. (2014) Tet and TDG mediate DNA demethylation essential for mesenchymal-to-epithelial transition in somatic cell reprogramming. Cell Stem Cell, 14, 512-522.