

# Detecting DNA Base Modifications Using Single Molecule, Real-Time Sequencing

## Introduction

Base modifications are important to the understanding of biological processes such as gene expression, host-pathogen interactions, DNA damage, and DNA repair<sup>1</sup>. Single Molecule, Real-Time (SMRT®) sequencing has the potential to revolutionize the study of base modifications through direct detection of unamplified source material.

Traditionally, it has been a challenge to study the wide variety of base modifications that are seen in nature. Most high-throughput techniques focus only on cytosine methylation and involve both bisulfite sequencing to convert unmethylated cytosine nucleotides to uracil nucleotides, and comparison of sequence reads from bisulfite-treated and untreated samples<sup>2</sup>. SMRT sequencing, in contrast, does not require base conversion within the source material in order to detect base modifications. Instead, the kinetics of base addition is measured during the normal course of sequencing. These kinetic measurements present characteristic patterns in response to a wide variety of base modifications – researchers at PacBio have been able to observe unique kinetic characteristics for over 25 base modifications<sup>3</sup>.

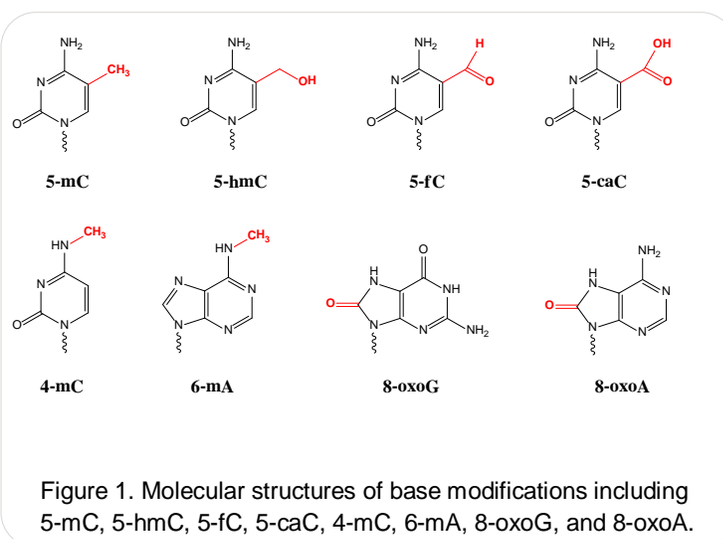
As a result of this breakthrough methodology to detect base modifications, it is now possible to sequence modifications other than 5-methylcytosine (5-mC). Bacterial modifications such as 6-methyladenine (6-mA), 4-methylcytosine (4-mC), or more recently identified eukaryotic modifications such as 5-hydroxymethylcytosine<sup>4</sup> (5-hmC), are also accessible to study using a single sequencing method on the PacBio® RS system.

As our understanding of kinetic information grows, the analysis of base modifications using SMRT technology will continue to become easier and faster.

## Types of Base Modification in Biology

DNA base modifications have a variety of functional roles which include:

- Epigenetic markers for influencing gene expression such as 5-mC, 5-hmC, 5-formylcytosine (5-fC), and 5-carboxylcytosine (5-caC).
- Bacterial identity markers for affecting host-pathogen interactions such as 6-mA, 4-mC, and 5-mC.
- Bacterial epigenetic markers for regulating DNA replication and repair and transcription regulation such as 6-mA.
- Products of DNA damage such as 8-oxoguanine (8-oxoG) and 8-oxoadenine (8-oxoA).



Today, the most commonly studied base modification is 5-mC, commonly referred to as “methylation” despite the fact that other bases are also naturally modified by methyl groups.

Methylation has gathered interest from researchers in a variety of disciplines including early developmental biology, cancer biology, and neurological disorders. Methylation and demethylation help regulate gene expression and have been linked to several human diseases through mechanisms such as deactivating tumor suppressors or activating oncogenes<sup>5</sup>.

Other modifications, such as 6-mA in bacteria, have been studied with lower resolution methods – such as chromatography or through methylation's protective effect against restriction endonucleases – because they are not easily accessible with standard sequencing techniques. This modification is associated with basic functions such as DNA replication and repair<sup>6</sup>. It is also common in protists and plants, and some studies suggest that it may also be present in mammalian DNA<sup>7</sup>.

SMRT sequencing is capable of detecting 6-mA as well as other common bacterial base modifications. As a result, the technology is expected to increase our understanding of a broad array of biological processes.

The potential benefits of detecting base modification, using SMRT sequencing, include:

- Single-base resolution detection of a wide

variety of base modifications (including those in Figure 1 and more).

- Single-molecule resolution over long-read distances.
- Unamplified double-stranded input DNA, which means that strand-specific modifications, such as hemimethylation, are detectable.
- Hypothesis-free base detection which allows discovery of unknown or unexpected modifications through the effects on sequencing kinetics (as described below).

### Studying Polymerase Kinetics with SMRT® Sequencing

SMRT Sequencing allows the observation of single DNA polymerases reading individual molecules of DNA in real time. Therefore, the kinetic characteristics of DNA polymerization are observable on a single-molecule basis. The kinetic characteristics, such as the time duration between two successive base incorporations, are altered by the presence of a modified base in the DNA template<sup>3</sup>. This is observable as an increased space between fluorescence pulses, which is called the interpulse duration (IPD), as shown in Figure 2.

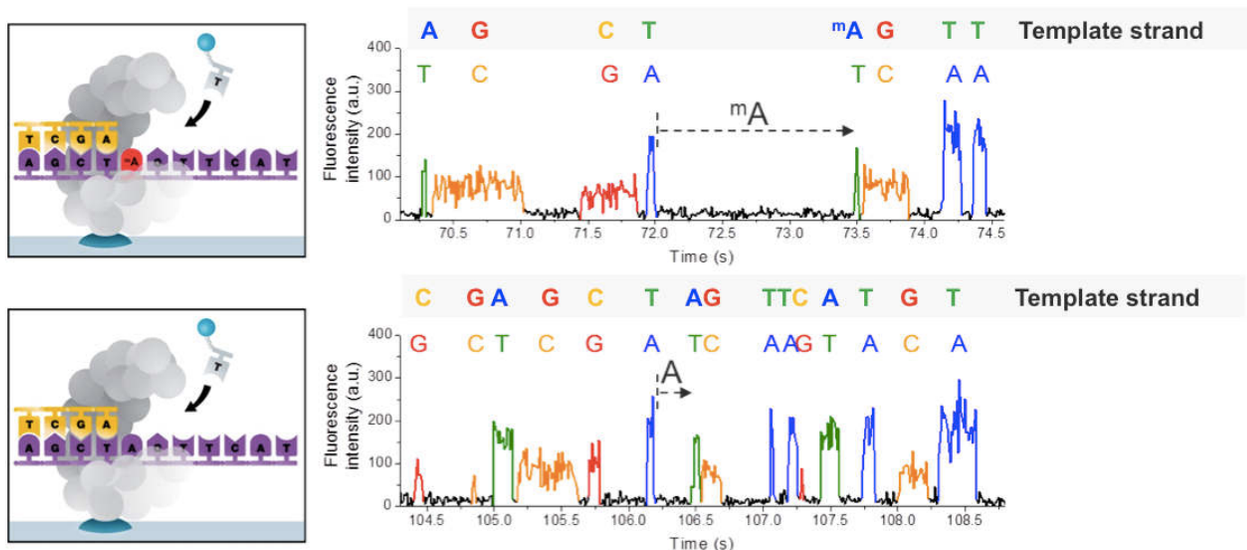


Figure 2. Principle of detecting modified DNA bases during SMRT sequencing. The presence of the modified base in the DNA template (top), shown here for 6-mA, results in a delayed incorporation of the corresponding T nucleotide, i.e. longer interpulse duration (IPD), compared to a control DNA template lacking the modification (bottom).<sup>3</sup>

These changes in the DNA polymerase speed, relative to an *amplified control* template lacking modified bases, can be measured for each template position to indicate the presence of modified bases in the DNA template. In order to quantify the change in IPD distributions between a control sample and a native sample, we define the *IPD ratio* as the ratio of the mean IPD at a site in the native sample to the mean IPD at the same site in the amplified control.



Figure 3. Image from SMRT® View software using the *in silico* control showing detection of 6-mA at GATC motifs using IPD ratios. For each template position (x-axis), the ratio (y-axis, purple bar up for forward strand and orange bar down for reverse strand) of average interpulse durations (IPDs) for each strand of the native DNA to the control is plotted. Excursions from the baseline indicate the presence of a modified base (6-mA, in this example, marked with indicators at the top along with a seven-base context in 5' -> 3' orientation), slowing down the polymerase at the position of the modification<sup>9</sup>.

An alternative method calculates IPD ratios based on a computational model rather than using an unmodified control sample. This model is referred to as the “*in silico* control”. The model must be trained to recognize the kinetics of a specific sequencing chemistry. The *in silico* control has obvious benefits in reducing the time required to perform an initial experiment and avoids using an amplified control.

The local sequence context of unmodified DNA influences polymerase dynamics. As such, the *in silico* control uses a predicted value of the mean IPD at each reference position that is based on the local sequence context.

Kinetic effects are not necessarily limited to just the nucleotide incorporation opposite the modified base position in the DNA template. This is because the DNA polymerase is in intimate contact with the DNA over an extended region of approximately twelve

bases, and the modified base can impart effects on the polymerase dynamics at several positions over this region. This results in kinetic “signatures” which can aid in identifying the type of base modification.

For example, for the three common bacterial identity markers<sup>8</sup> (see Figure 4):

- 5-mC has typical characteristic kinetic signals two and six bases downstream of the methylated position
- 4-mC just at the position of the modification
- 6-mA at the modified position and often five bases downstream

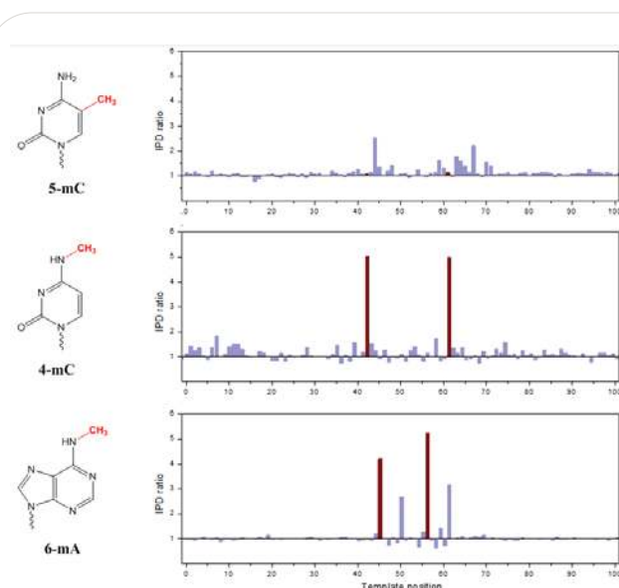


Figure 4. Kinetic signatures for the three common bacterial identity markers 5-mC, 4-mC, and 6-mA. The methylated template positions are highlighted in red. Note that the kinetic signatures vary both in magnitude as well as in the length of the region over which the polymerase dynamics are affected.

Different signal magnitudes translate to different amounts of sequencing-fold coverage required to obtain similar confidence levels for modification detection.

The current *in silico* control algorithms have proven reliable for studying 6-mA in bacteria, for example in methylome motif analysis. However, when studying subtle kinetic effects such as those for native 5-mC, using an amplified control is more accurate than

using the *in silico* control. Future commercial products are expected to improve use of the sequence context to better identify and interpret modification signals.

Based on the measurement and analysis principle outlined above, a typical experiment for detecting DNA base modifications can be designed as follows:

- Obtain native DNA of interest and prepare a SMRTbell™ library for SMRT sequencing (see the Pacific Biosciences Sample Preparation and Sequencing Guide).
- If using an amplified control design, use a small aliquot of the original DNA sample and perform a whole-genome amplification (WGA) reaction to obtain the control DNA sample lacking any base modifications. Then prepare a SMRTbell library for this sample.
- Perform adequate SMRT sequencing (for both samples when using an amplified control) to obtain the necessary coverage needed for the modifications under study, and sufficient overall coverage to characterize the genome (or portion of the genome) of interest.
- Use the bioinformatics tools to perform kinetic analysis for base modification<sup>9</sup>.

## Examples of Current Applications

Direct DNA sequencing of base modifications requires that the following general considerations be fulfilled during the experimental design:

1. Unamplified DNA is necessary. Amplification of DNA through PCR, WGA, or other techniques, will result in the loss of base modifications. Therefore, enough native DNA must be available to make a sequencing library. With version 2 sequencing reagents ("C2 chemistry"), the sample requirement for preparing a ~500 bp SMRT sequencing library is 250 ng of input DNA.
2. As DNA input requirements are reduced over time, a smaller or targeted region of a genome is expected to be sufficient for studies without amplification requirements.

3. The different magnitudes of kinetic signal, for the different base modifications, translate to different amounts of sequencing fold coverage required to obtain high confidence levels of detection.

We recommend the minimum sequencing fold coverage *per strand* as follows<sup>9</sup>:

4-methylcytosine	25x
5-methylcytosine	250x
5-hydroxymethylcytosine	250x
glucosylated 5-hydroxymethylcytosine <sup>1</sup>	25x
hydroxymethylcytosine enriched with the Hydroxymethyl Collector™ Kit <sup>2</sup>	5x
6-methyladenine	25x
8-oxoguanine	25x

4. The genome, or portion of a genome, to be interrogated also has to be compatible with the current throughput performance of the PacBio RS. With C2 Chemistry, the base throughput of a two hour run on the PacBio RS is approximately 100 Mb. Studying methyladenine in *E. coli* requires 4 to 6 SMRT Cells per sample, which can be run in one afternoon. Over time, we expect SMRT technology throughput to increase, making larger genome sizes more practical to sequence for base modifications.

Applications particularly suitable for SMRT sequencing based on the criteria above include:

- Sequencing of *E. coli* to find the sites of 5-mC, 4-mC, and 6-mA modification when studying virulence, gene expression, or pathogen-host interactions. See the *Pacific Biosciences Technical Note - Detecting DNA Base Modifications* for more detailed information on strategies for experimental design and motif analysis in bacteria.
- Isolating and purifying mitochondrial, chloroplast or other small genomes, containing known or novel DNA base modifications.
- Enriching portions of a larger genome (e.g., isolation of DNA regions modified with 5-hmC,

<sup>1</sup> Generated by T4 Phage β-glucosyltransferase (Josse, J. and Kornberg, A. (1962) J. Biol.Chem., 237, 1968-1976)

<sup>2</sup> Available from Active Motif  
(<http://www.activemotif.com/catalog/775/hydroxymethyl-collector-trade>)

and enriched by specific biotinylation chemistries of 5-hmC and subsequent streptavidin pulldown). Normalizing IPDs with the *in silico* control is particularly useful for this type of experiment, where generating a completely overlapping set of amplified control sequence data can be a challenge.

## Conclusion

SMRT sequencing is the only commercially available technology capable of measuring the kinetics of base incorporation during a sequencing run. This kinetics information can be used to identify sites in the target DNA that have been chemically modified in a variety of ways (including methylation, formylation, carboxylation, and more). These base modifications are associated with several biological processes including gene expression and DNA oxidative damage.

Most of these modifications have not been extensively studied due to difficulties in adapting experimental techniques to a higher throughput sequencing method – we expect that these modifications will be possible to study with SMRT sequencing. Examples of novel studies that may be performed on the PacBio *RS* today include full-genome bacterial modification studies and hydroxymethylcytosine studies in mammalian genome regions enriched for that modification.

We anticipate that over time the range of target genomes that will be addressable by SMRT sequencing will improve and ultimately address regions as small as single genes as well as larger whole genomes. We further expect that the advances afforded by SMRT sequencing will enhance genetic studies into gene regulation, DNA damage and repair, bacterial virulence, and other important biological pathways. Novel DNA modifications could potentially be discovered, expanding the utility of sequencing to even broader areas of study.

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9. SMRT Analysis version 1.3.1 or higher, and/or "DNA Modification Detection with SMRT Sequencing using R" found at <https://github.com/PacificBiosciences/R-kinetics>.

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