Methylation profiling of normal individuals reveals mosaic promoter methylation of cancer-associated genes.

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ABSTRACT:

Epigenetic silencing by promoter methylation of genes associated with cancer initiation and progression is a hallmark of tumour cells. As a consequence, testing for DNA methylation biomarkers in plasma or other body fluids shows great promise for detection of malignancies at early stages and/or for monitoring response to treatment. However, DNA from normal leukocytes may contribute to the DNA in plasma and will affect biomarker specificity if there is any methylation in the leukocytes. DNA from 48 samples of normal peripheral blood mononuclear cells was evaluated for the presence of methylation of a panel of DNA methylation biomarkers that have been implicated in cancer. SMART-MSP, a methylation specific PCR (MSP) methodology based on real time PCR amplification, high-resolution melting and strategic primer design, enabled quantitative detection of low levels of methylated DNA. Methylation was observed in all tested mononuclear cell DNA samples for the \textit{CDH1} and \textit{HIC1} promoters and in the majority of DNA samples for the \textit{TWIST1} and \textit{DAPK1} promoters. \textit{APC} and \textit{RARB} promoter methylation, at a lower average level, was also detected in a substantial proportion of the DNA samples. We found no \textit{BRCA1}, \textit{CDKN2A}, \textit{GSTP1} and \textit{RASSF1A} promoter methylation in this sample set. Several individuals had higher levels of methylation at several loci suggestive of a methylator phenotype. In conclusion, methylation of many potential DNA methylation biomarkers can be detected in normal peripheral blood mononuclear cells, and is likely to affect their specificity for detecting low level disease. However, we found no evidence of promoter methylation for other genes indicating that panels of analytically sensitive and specific methylation biomarkers in body fluids can be obtained.

INTRODUCTION

Recent research has led to the understanding that, despite the importance of genetic change, epigenetic mechanisms are perhaps the predominant drivers of cancer [1-3]. In particular, inactivation of genes by \textit{de novo} promoter methylation accompanied by global hypomethylation is the most common DNA lesion of cancer cells [2, 3].

The CpG dinucleotide is the principal unit of methylation in humans as its palindromic pairing with a CpG dinucleotide on the other strand enables the semiconservative replication of methylation by DNA methyltransferase 1. Regions of high CpG density
are referred to as CpG islands. CpG islands are often found spanning the promoter and first exon of genes, and these promoter region CpG islands are normally unmethylated. DNA methylation of promoter regions is strongly associated with transcriptional silencing of the downstream gene. This may in part be explained by the binding of proteins containing methyl binding domains which then initiate the formation of complexes that repress transcription [4].

De novo methylation of CpG islands is often an early event in cancer [5]. Thus aberrant promoter methylation of a number of genes shows great promise as biomarkers for early cancer detection [5-7]. Ideally methylation biomarkers should be detectable in readily accessible body fluids or tissues. However, the majority of DNA methylation biomarkers lack the sensitivity and specificity required for a diagnostic test [7,8].

Plasma and serum are the most studied body fluids with regard to DNA methylation biomarkers. Very sensitive and quantitative methods are needed for the detection of these biomarkers. However, tumour-derived DNA in body fluids may often be significantly contaminated with DNA from normal cells especially when the tumour is small. Though the source of the normal DNA is unclear, it is likely that DNA from normal leukocytes may affect biomarker specificity if methylated.

Thus, very sensitive and quantitative methods may be needed for the detection of these biomarkers in body fluids.

Biomarker sensitivity and specificity is also influenced by the choice of region for analysis and the choice of method for methylation analysis. Many different methods for the detection of DNA methylation are available [9, 10]. The most sensitive methods utilise methylation-specific PCR (MSP) [11] in which amplification is determined by the methylation of the region under the primers. Conventional MSP is widely used due to its simplicity. However, it is non-quantitative and is also prone to false positive results caused by incomplete bisulphite conversion or false priming events [5, 9, 12-14].

Thus, when it is necessary to detect rare methylated sequences, MSP may be unable to distinguish low-level methylation from low-level incomplete conversion or false priming. MethyLight, a variant of MSP using fluorescent TaqMan probes is more specific [15] as the use of the probe makes it quantitative and enables increased specificity.

We recently developed a highly sensitive probe-free method called Sensitive Melting Analysis after Real Time (SMART)-MSP [12], which can provide quantitative data, works well with DNA derived from formalin fixed
paraffin embedded (FFPE) tissues [16], and is capable of identifying many false positives resulting from false priming or incomplete conversion of unmethylated cytosines during the bisulphite conversion of the DNA template. This is possible due to the use of high-resolution melting analysis [17], which utilises intercalating dyes that can be used at saturating conditions without inhibiting the PCR [18].

In this contribution, we have evaluated the methylation levels in normal peripheral blood mononuclear cells for a panel of genes (APC, BRCA1, CDH1, CDKN2A, DAPK1, GSTP1, HIC1, RARB, RASSF1A and TWIST1). These genes have been implicated as actual or potential tumour suppressor genes and are methylated in various cancers. These genes have potential as DNA methylation biomarkers for both early detection and post-therapeutic monitoring. By using SMART-MSP we were able to distinguish artefactual results from true positive results, and to obtain quantitative data for the studied panel of biomarkers. Thus, we were able to show that some are normally methylated at low levels in peripheral blood mononuclear cells, whereas methylation was undetectable in others.

RESULTS

Melting profiles of the SMART-MSP assays

The melting profiles of true positives for each assay were obtained by amplifying methylated standards. The melting temperature ($T_m$) was 78.3°C for APC, 76.5°C for BRCA1, 81.9°C for CDH1, 75.8°C for CDKN2A, 79.5°C for DAPK1, 80°C for GSTP1, 79.8°C for HIC1, 81.5°C for RARB, 78.2°C for RASSF1A and 79.8°C for TWIST1. Only bands of the expected size were observed after electrophoresis (data not shown).

The sensitivity and quantitative accuracy of the SMART-MSP assays

The sensitivity of the SMART-MSP assays was tested using a standard dilution series of methylated DNA into unmethylated DNA. The CDKN2A and the RASSF1A assays were sensitive to 0.1% methylated template with high reproducibility. The APC, BRCA1, CDH1, DAPK1, GSTP1, HIC1, RARB and TWIST1 assays were sensitive to 0.05% methylated template (data not shown).

All standards contained approximately equal amounts of template suitable for PCR after bisulphite

![Figure 2: Examples of false positive results.](image-url)
conversion, as evident from the similar C\(_r\) values obtained in the COL2A1 control assay. We used the Rotorgene software to obtain a standard curve for the dilution series for calculation of the correlation coefficient (r\(^2\)) and PCR efficiency (E) for each assay. The correlation coefficients (APC: r\(^2\) = 0.987, BRCA1: r\(^2\) = 0.985, CDH1: r\(^2\) = 0.992, CDKN2A: r\(^2\) = 0.998, DAPK1: r\(^2\) = 0.995, GSTP1: r\(^2\) = 0.985, HIC1: r\(^2\) = 0.995, RARB: r\(^2\) = 0.982, RASSF1A: r\(^2\) = 0.978, TWIST1: r\(^2\) = 0.995) indicated a strong linear relationship between C\(_r\) values and given concentrations for all assays. The PCR efficiency of most assays (APC: E = 1.89, CDH1: E = 1.88, CDKN2A E = 1.96, DAPK1: E = 1.86, GSTP1: E = 1.94, HIC1: E = 1.87, RARB: E = 1.92, TWIST1: E = 1.88) was approximately the same as the control assay (COL2A1: E = 1.90). The PCR efficiencies of the BRCA1 (E = 1.78) and RASSF1A (E = 1.98) assays were more different from the control assay, but as none of the samples were positive for these assays, quantification was not necessary.

**Figure 3:** Sequencing confirms that right-shifting was due to incomplete conversion. In the APC SMART-MSP assay seven non-CpG cytosines are found in between the primers (shown as upper case T’s when converted during the bisulphite modification). The last base of each primer and the region in between them are shown here. CpG sites are shown in bold. A. Sequencing results for the 100% methylated control. All non-CpG cytosines were converted during the bisulphite modification. B. Sequencing results for sample 48 from figure 2A. It is observed that five out of the seven non-CpG cytosines found in between the primers were not converted. This is causing the right-shift observed in figure 2A.

**Promoter methylation levels in the samples**

The COL2A1 control assay was used to normalise for DNA input after bisulphite conversion in the real-time PCR quantification [12]. The samples generally amplified simultaneously and relatively early with the COL2A1 assay. One sample (#45) that amplified late for the COL2A1 assay, (indicating a very low number of target bisulphite modified templates) was omitted from the analysis.

Quantitative data for all samples for each assay is shown in Table 2. None of the samples were positive for BRCA1, CDKN2A, GSTP1 or RASSF1A methylation in the regions analysed. The regions analysed for the other genes were detectably methylated in some or all of the individuals tested: HIC1: 100%, CDH1: 100%, TWIST1: 72%, DAPK1: 51%, RARB: 32%, APC: 23%.

Some regions tended to be methylated at higher levels i.e. in a higher proportion of cells than others. The estimated methylation levels were highest for the HIC1 and CDH1 assays. TWIST1, APC, DAPK1 and RARB positive samples were methylated at lower levels. The mean methylation level of positive samples for each gene was: HIC1: 3.61%, CDH1: 1.26%, TWIST1: 0.43%, RARB: 0.18%, APC: 0.18%, DAPK1: 0.13%. The overall level of methylation observed in the positive samples was higher for the assays that were positive in a greater proportion of samples. Thus, it is possible that each gene’s methylation may represent a continuous variable with the more highly methylated genes having the whole range above the threshold of detection.

The range of methylation levels varied the most for the HIC1 assay (many samples were far from the mean). The methylation levels also varied considerably for the CDH1 and TWIST1 assays. Real-time amplification data and melting profiles for representative samples are shown for CDH1 and TWIST1 (Figure 1).

A tendency towards increasing methylation levels with increasing age was observed for most of the genes. However, more samples would be needed to adequately address this question. Two samples (10 and 11) showed overall elevated methylation levels for most of the assayed genes. These samples were not only methylated at high levels but also had a large number of methylated genes. In contrast, samples 26, 27, 35, 37, 39, 42, 46, 47 and 48 overall showed low methylation levels and methylation was restricted to a few genes (Table 2).

The reproducibility was not as good for the replicates showing the lowest levels of DNA methylation as evident from figure 1A, and occasionally only one replicate amplified. This can be readily explained by the limiting amounts of methylated template at the lowest methylation levels. The quantitative data in Table 2 is calculated from the mean values of the replicates. In situations where only one of the two replicates amplified, the methylation level estimated from that replicate was halved.
Identification of false positives caused by incomplete conversion

In the SMART-MSP assays used, only non-CpG cytosines are located between the primers. Thus, if these cytosines are not converted to uracil during the bisulphite modification, the amplicon will have a higher GC-content, melt later, and thus a right-shifting (due to a higher than expected melting temperature) of the melting profile will be observed. This, in combination with late amplification is a strong indicator of a false positive result [12].

Occasionally, amplified samples showed markedly right-shifted melting profiles, and were scored as false positives (Table 2). Examples from the APC, BRCA1, DAPK1 and RARB assays are shown in figure 2. Right-shifting was always associated with late amplification and poor reproducibility between replicates as would be expected for this type of false positive result [12]. This is because incompletely converted molecules are typically found at low levels. These samples would have been scored as positive results when using gel electrophoresis as in conventional MSP because gel electrophoresis sorts DNA molecules according to size, and does not evaluate the bases found in between the primers as the SMART-MSP assays do.

Samples were scored as a false positive result in situations where both replicates amplified and showed right-shifted melting and in situations where only one of the replicates amplified and showed a right-shifted melting profile while the other replicate did not amplify. Variant APC products were sequenced to confirm that right-shifting was due to incomplete conversion. An example is

| Gene | Primer sequences (CpG sites in bold and converted Cs as capital Ts or As) | Annealing temperature (°C) | Size (bp) | Non-CpG Cs between primers | Spanned region (UCSC Genome Browser, March 2006: NCBI36/hg18) |
|------|-------------------------------------------------------------------------------------------------|-----------------------------|----------|-----------------------------|-------------------------------------------------------------|
| APC  | F-"TcgtgagatcggagTtgaggc" R-"ccatcgAagAaAactccgacgc" | 68                          | 55       | 7                           | 112101367-112101421 Chr. 5                                  |
| BRCA1| F-t"tgtagtTgtcttTgTgtagTgc" R-"tccgcggtTtttcg" | 65                          | 49       | 1                           | 38530948-38530996 Chr. 17                                   |
| CDH1 | F-"ggtggggggTcgtgagTgtagTgc" R-"ccatcctaAaAaccgacgc" | 65/68                       | 58       | 3                           | 67328555-67328612 Chr. 16                                   |
| CDKN2A| F-"gTgTTgagTgTgtagTgtagTgc" R-"ccatcctaAaAaccgacgc" | 64                          | 72       | 6                           | 21964971-21965042 Chr. 9                                   |
| DAPK1| F-"aggaTagTcgggTgtagTgc" R-"tAacccgtAaAacccgtAacgc" | 67                          | 61       | 4                           | 89302618-89302678 Chr. 9                                   |
| GSTP1| F-"gctgagTgtagTgtagTgc" R-"ccatcctaAaAaccgacgc" | 67                          | 51       | 4                           | 67107679-67107729 Chr. 11                                  |
| HIC1 | F-"TTaggcccTgTgtagTgc" R-"ctAaAaAaAaccgacgc" | 66                          | 54       | 4                           | 1906662-1906714 Chr. 17                                    |
| RARB | F-"atgTcgggTgtagTgc" R-"gtcgggtTtttctaAaAaccgacgc" | 71                          | 64       | 3                           | 25444860-25444923 Chr. 3                                   |
| RASSF1A| F-"cggTTgggTcgggTtgagTgc" R-"AAacccgtAaAaccgacgc" | 68                          | 58       | 5                           | 50353240-50353297 Chr. 3                                   |
| TWIST1| F-"cgcgggTgtagTa" R-"aAaccgacgc" | 68                          | 59       | 4                           | 19124102-19124160 Chr. 7                                   |
| COL2A1| F-"gTaatgTgtagTgtagTgtagTgc" R-"ccatcctaAaAaccgacgc" | 65                          | 86       | 1                           | 46667210-46667295 Chr. 12                                  |

It should be noted that optimal annealing temperature may vary according to the PCR machine used and the other parameters of the PCR reaction.
shown for sample 48 in figure 3 (also shown in figure 2A). It is observed that five out of the seven non-CpG cytosines found in between the APC primers were not converted, whereas all seven sites were converted in the positive control.

**Verification of CDH1 results by MethyLight**

The CDH1 MethyLight assay was quantitatively accurate in the range from 100% down to 0.1% methylated template. The correlation coefficient of the MethyLight assay was: \( r^2 = 0.984 \), again indicating a strong linear relationship between \( C_T \) values and given concentrations. We tested a selection of the samples showing different levels of CDH1 methylation (sample 5, 6, 7, 10, 11, 13, 14, 19, 20, 36, 41, 43, 44, 46) using MethyLight to confirm that these positive results were not a result of false priming. In the MethyLight assay, CpG sites are also present in the probe sequence which increases the specificity for methylated templates. All of the samples, except sample 19 which was negative, were shown to be methylated at similar but slightly lower levels compared to the data obtained by the CDH1 SMART-MSP assay. MethyLight amplification data for six of the samples are shown in figure 4.

**CDH1 SMART-MSP using different annealing temperatures**

If the observed methylation with the CDH1 assay was due to a rare subpopulation of fully methylated cells, the samples would be estimated to be methylated at the same levels when using a higher annealing temperature. We tested a number of the samples showing different levels of CDH1 methylation (sample 2, 3, 5, 10, 11, 12, 15, 18, 20, 37) using 68°C as the annealing temperature instead of 65°C. Again, we found high reproducibility and non-shifted melting profiles (data not shown). However, we found that all samples were methylated at similar but slightly lower levels. These results and the MethyLight results indicate that there is some heterogeneity in the methylation. As the temperature is increased, tolerance of the primers binding to an imperfectly matched template is decreased.

**Confirming that CDH1 is heterogeneously methylated by HRM**

To further test whether the observed CDH1 methylation is heterogeneous, we used the same primers as for the MethyLight assay, but with the dye SYTO-9 instead of the probe. This allows evaluation of the methylation status of the CpG sites between the primers including the two CpG sites covered by the MethyLight assay. If left-shifted melting profiles relative to the 100% methylated standard are observed this is an indication that the two CpG sites are heterogeneously methylated, due to a less GC-rich amplicon and/or heteroduplex formation.
| Sample (Age/Sex) | APC      | CDH1   | DAPK1   | HIC1    | RARB    | TWIST1   |
|-----------------|----------|--------|---------|---------|---------|----------|
| 1 (84/M)        | 0.09%    | 0.52%  | 0.16%   | 2.72%   | 0%      | 0.05%    |
| 2 (84/M)        | 0%       | 1.68%  | 0.14%   | 2.21%   | 0.06%   | 0.21%    |
| 3 (83/F)        | 0.06%    | 1.18%  | 0.17%   | 1.46%   | 0.13%   | 0.32%    |
| 4 (82/M)        | False positive | 0.64% | 0.06%   | 0.96%   | 0%      | 0.17%    |
| 5 (82/F)        | 0.24%    | 1.18%  | 0.14%   | 3.13%   | 0.06%   | 0.39%    |
| 6 (80/F)        | 0%       | 3.85%  | 0.32%   | 7.69%   | 0.30%   | 0.48%    |
| 7 (77/M)        | 0%       | 3.35%  | 0.11%   | 8.25%   | 0.28%   | 0.16%    |
| 8 (76/M)        | 0%       | 0.78%  | 0%      | 1.46%   | 0%      | 0.17%    |
| 9 (75/F)        | 0%       | 1.68%  | 0.18%   | 12.50%  | 0%      | 0.09%    |
| 10 (74/M)       | 0.21%    | 4.42%  | 0.39%   | 12.50%  | 0.48%   | 2.06%    |
| 11 (71/M)       | 0.52%    | 5.83%  | 0.17%   | 20.31%  | 0.45%   | 7.18%    |
| 12 (71/F)       | 0%       | 1.79%  | 0.07%   | 1.92%   | 0%      | 0.07%    |
| 13 (71/F)       | 0%       | 2.21%  | 0%      | 4.74%   | 0%      | 0%       |
| 14 (69/M)       | 0%       | 1.27%  | 0%      | 1.92%   | 0%      | 0.05%    |
| 15 (68/M)       | 0%       | 1.27%  | 0%      | 14.36%  | 0.21%   | 0.42%    |
| 16 (68/M)       | 0.09%    | 0.73%  | 0%      | 3.13%   | 0.10%   | 0.15%    |
| 17 (66/F)       | False positive | 1.11% | False positive | 2.06% | 0%      | 0.16%    |
| 18 (66/M)       | 0%       | 1.18%  | 0%      | 5.83%   | 0.07%   | 0.24%    |
| 19 (61/F)       | 0%       | 1.56%  | 0.09%   | 1.79%   | False positive | 0.05% |
| 20 (61/F)       | 0.08%    | 0.96%  | 0%      | 2.92%   | 0.05%   | 0.36%    |
| 21 (60/M)       | 0.10%    | 0.59%  | 0.05%   | 2.37%   | 0.13%   | 0.05%    |
| 22 (58/M)       | False positive | 0.84% | 0.06%   | 1.46%   | 0%      | 0%       |
| 23 (56/M)       | 0%       | 0.45%  | 0%      | 0.73%   | 0%      | 0.26%    |
| 24 (55/M)       | 0%       | 0.84%  | 0.06%   | 3.35%   | 0%      | 0.21%    |
| 25 (54/F)       | 0%       | 1.03%  | 0.05%   | 1.27%   | 0.06%   | 0.18%    |
| 26 (52/F)       | 0%       | 0.36%  | 0%      | 1.67%   | False positive | 0%    |
| 27 (50/F)       | False positive | 0.48% | 0%      | 1.36%   | 0%      | 0.06%    |
| 28 (50/M)       | 0%       | 1.27%  | 0.21%   | 12.50%  | 0%      | 0%       |
| 29 (49/F)       | 0%       | 0.73%  | 0%      | 1.46%   | 0%      | 0.12%    |
| 30 (48/M)       | 0%       | 0.28%  | 0%      | 4.12%   | 0%      | 0%       |
| 31 (45/F)       | 0.26%    | 2.06%  | 0.06%   | 1.67%   | 0%      | 0.10%    |
| 32 (44/M)       | 0%       | 1.92%  | 0.13%   | 3.35%   | 0.16%   | 0.10%    |
| 33 (42/F)       | 0.06%    | 1.03%  | 0.18%   | 1.10%   | 0%      | 0.15%    |
| 34 (39/F)       | 0%       | 0.36%  | 0%      | 1.92%   | 0%      | 0.16%    |
| 35 (37/M)       | 0%       | 0.48%  | 0%      | 1.36%   | 0%      | 0%       |
| 36 (33/M)       | 0%       | 1.56%  | 0%      | 2.21%   | 0%      | 0.11%    |
| 37 (33/F)       | 0%       | 0.52%  | 0%      | 1.27%   | 0%      | 0%       |
| 38 (33/M)       | False positive | 0.89% | 0%      | 1.56%   | 0%      | 0.11%    |
| 39 (29/F)       | 0%       | 0.28%  | 0%      | 1.46%   | 0%      | 0%       |
| 40 (27/M)       | 0%       | 0.52%  | 0%      | 2.21%   | 0%      | 0%       |
| 41 (26/F)       | 0.09%    | 1.18%  | 0.06%   | 1.67%   | 0%      | 0.06%    |
| 42 (26/M)       | 0%       | 0.45%  | 0%      | 0.78%   | 0%      | 0.05%    |
| 43 (26/F)       | 0%       | 0.96%  | 0.08%   | 1.18%   | False positive | False positive |
| 44 (25/M)       | 0%       | 1.68%  | 0.13%   | 2.06%   | 0.13%   | 0.10%    |
| 45 (23/M)       | No data  | No data | No data | No data | No data | No data  |
| 46 (23/F)       | 0%       | 0.36%  | 0%      | 0.78%   | 0%      | 0%       |
| 47 (23/M)       | 0%       | 0.63%  | 0.06%   | 1.79%   | 0%      | False positive |
| 48 (?/? )       | False positive | 0.24% | 0%      | 1.18%   | False positive | 0%    |
DISCUSSION

We found no evidence for methylation of the CDKN2A, GSTP1, RASSF1A and BRCA1 promoter regions assayed in this set of peripheral blood mononuclear cell DNA samples. It is instructive that these biomarkers for which we found no methylation have been successfully used in a number of other studies. As examples, methylation of the CDKN2A promoter has been detected in sputum of smokers up to 3 years before they were diagnosed with cancer [19, 20], GSTP1 has been used in prostate cancer detection [21], and RASSF1A can be detected in serum from gastric colorectal adenocarcinoma patients [22]. Women with detectable BRCA1 methylation in the peripheral blood have an increased risk of early onset breast cancer [23].

However, we have shown that significant levels of mosaic promoter CpG island methylation are present in peripheral blood mononuclear cells derived from normal individuals for a number of other genes whose methylation is implicated in cancer. These results have several important implications.

Firstly, they indicate that the assays that show considerable background methylation will be insufficiently specific for use as DNA methylation biomarkers for peripheral blood–based early detection or monitoring. Similar considerations apply for monitoring of body fluids or tissues that have significant a haematopoietic component but these will require further analysis of background methylation in other contributing tissues. Thus, our results support the use of assays based on BRCA1, CDKN2A, GSTP1 and RASSF1A methylation as specific biomarkers. However, assays based on APC, CDH1, DAPK1, HIC1, RARB and TWIST1 methylation may prove to be less specific biomarkers, especially if non-quantitative methods are used.

One reservation about this study using peripheral blood mononuclear cells is that the neutrophils which comprise the major leukocyte component of blood were not analysed. Future studies will need to investigate blood fractions more intensively as well as investigating variations in other tissues.

Some of the genes with substantial background methylation in this study have been used for DNA methylation biomarkers. For example, TWIST1 and RARB methylation was used to detect breast cancer cells in ductal lavages [24]. Whereas, the assays used were different from ours and may have been more specific under the reaction conditions used, it is clear that each assay for a methylation biomarker needs to be thoroughly tested against population controls and appropriate tissues as a critical step in its validation.

Among the panel of genes that we have found to be methylated, CDH1 [25] and DAPK1 [26] have previously been reported to be methylated in the blood in a high proportion of normal individuals. CDH1 was reported to be heterogeneously methylated as also found in the present study. The use of less sensitive methods and/or the use of different blood fractions and a lack of sufficient normal controls may explain why the other genes for which we observed significant methylation have not previously been shown to be methylated in normal blood. Furthermore, positives obtained by conventional MSP may have been considered as false positive results. In this study, false positive results that are likely to be caused by incomplete conversion were readily identified. By contrast, identification of false positives is not possible using conventional MSP.

The use of a whole genome amplified (WGA) unmethylated control allowed us to determine the appropriate PCR conditions for minimising false priming events. However, even after optimisation the WGA control occasionally amplified. This was associated with a lack of reproducibility in the replicates, very late amplification, and right-shifted melting profiles. As no CpG sites were placed in between most pairs of SMART-MSP primers, this allowed us to conclude that the observed right-shifting is caused by incomplete conversion of some of the non-CpG cytosines found between the primers. This was verified by sequencing for the APC SMART-MSP assay.

Similar false positive results likely to be due to the amplification of very low-level incompletely converted templates were observed in the samples. When these false positive reactions were analysed by electrophoresis, a fragment of the same size as the 100% methylated positive control was observed. These samples would have scored as positive by conventional MSP, however, the number of non-CpG cytosines between the primers influences the likelihood of detecting these false positives. In our BRCA1 samples, only one non-CpG cytosine is located between the primers, due to restrictions in assay design caused by the BRCA1 pseudogene[23]. Amplification occurred in 3 samples and the melting profiles were right-shifted by 0.6-0.8ºC. Given the size of the amplicon a shift in this range is expected for a single C to T change [27,28]. Thus, these samples were scored as false positive results. However, for the other assays with more non-CpG cytosines between the primers, the vast majority of samples we scored as false positive results showed much larger right-shifts (generally between 1ºC to 4ºC) indicating that several of the cytosines were not converted.

Small right-shifts below 0.6ºC were observed for some samples (Figure 1B and D). This was a consequence...
of the bisulphite conversion procedure being done using columns for the standards and using plates for the samples. The precise kit used for bisulphite modification affects the melting results probably as a result of different salt concentrations (unpublished results). This correctable fault points out the importance of using the exact same kit for template production for all high resolution melting experiments.

We observed the highest false positive rates in the APC and RARB assays. This can be explained by the primers for these assays containing fewer non-CpG cytosines and thus selecting against incomplete conversion less efficiently. For this reason, we recommend designing primers containing many non-CpG cytosines and preferably right at the 3’ end for at least one of the two primers as is the case for our CDH1 primers.

If primers select strongly against incomplete conversion by the inclusion of multiple Ts derived from non-CpG Cs (as our CDH1 primers do) and the detected methylation levels are relatively high, it may be of interest to have CpG positions in between the primers, since this allows assessment of whether the methylation between the primers is heterogeneous or homogeneous by HRM analysis. If a complex melting pattern extending to the left is observed this is an indication that the CpG sites in between the primers are not consistently methylated. In our CDH1 SMART-MSP assay, we have shown that this can be assessed by running the assay at different annealing temperatures. If methylation levels are estimated to be lower when using higher annealing temperatures, this indicates that some of the CpG sites under the primers are not co-methylated. We interpret our CDH1 results as indicating that the CpG site close to the 5’ end of the forward primer does not need to be co-methylated when using 65°C annealing temperature. However, at 68°C, methylation levels were estimated to be slightly lower, indicating that this site now needs to be methylated for amplification to occur. Thus, when heterogeneously methylated DNA is analysed by quantitative MSP based methods, the estimated methylation levels will be dependent on the annealing temperature and the number and positions of the CpG sites in the primers. For very accurate quantification, methods capable of reading the methylation status of individual epialleles, such as digital MS-HRM and sequencing [29], digital MethyLight [30], clonal bisulfite genomic sequencing [31], and next generation sequencing of bisulfite modified DNA [32] will be necessary [33].

Interestingly, the observed individual variation of methylation levels for the studied healthy individuals, indicates that it may be of interest to further study the concept that individual variation in methylation propensity can be found in normal peripheral blood mononuclear cells. The results indicate that some individuals may have a significant burden of epimutations as they have increased methylation over a large number of methylatable promoter regions. In particular, samples 10 and 11 showed higher levels of methylation than the rest of the samples and had the most genes methylated as well. By contrast, samples 26, 27, 35, 37, 39, 42, 46, 47 and 48 showed low methylation levels and the methylation was restricted to only a few genes. The reasons for this remain unknown. Importantly, our results eliminate the trivial interpretation that the apparent hypermethylator phenotype is an artifact due to incomplete bisulfite conversion. Defects in methyl metabolism remain a possibility [34].

The varying levels of background methylation may reflect cancer predisposition. Each methylated allele represents an epimutation and if such epimutations affect genes whose inactivation predisposes to cancer development (as is the case for this gene panel), and these epimutations also occur in the target tissues for those cancers, the risk of developing cancer will be increased [34]. Prospective studies will be needed to determine if elevated methylation levels in fact predispose to cancer, and a larger sample cohort will be needed to address these questions appropriately. Interestingly, several studies have recently shown that DNA from peripheral blood may be differentially methylated between cancer patients and normal controls [35-37].

In conclusion, because DNA from normal leukocytes contributes to the templates prepared from body fluids that can be non-invasively used for PCR based analysis [6, 38], biomarkers methylated in normal blood may be less specific, especially when using non-quantitative methods. Thus, methylation levels should be assessed quantitatively for discrimination of low-level methylation in healthy cells from methylated DNA derived from cancer cells.

MATERIALS AND METHODS

Samples

The investigations were performed after approval by the Peter MacCallum Cancer Centre Ethics of Human Research committee (Projects 02/70 and 02/26). Peripheral blood samples from blood donors were obtained after informed consent from the Australian Red Cross Blood Service.

DNA extraction and bisulphite modification

Mononuclear cells from 3-5 mL of peripheral blood were prepared using Lymphoprep (Nyegaard, Oslo, Norway) and DNA was extracted using a salting out method [39]. Universal Methylated DNA (Chemicon, Millipore, Billerica, MA) was used as a fully methylated control. Whole genome amplified DNA from peripheral blood cells of normal individuals was used as unmethylated DNA as described previously.
SMART-MSP primer design

The amplicons were designed to allow the HRM analysis to assess the conversion status of non-CpG cytosines between the primers [12]. The primers overlaid a minimum of 2 CpG sites with one of the cytosines of a CpG site placed at or adjacent to the 3’ end. Non-CpG cytosines were included in the primer sequences to select against incompletely converted sequences, and at least one of these was placed as close to the 3’ end as possible. The primer sequences, genomic regions spanned, amplicon sizes and the annealing temperatures are found in Table 1. The primers for DAPK1 and CDKN2A (p16\(^{NK4a}\)) were previously published [12].

PCR and HRM conditions for the SMART-MSP assays

PCR cycling and HRM analysis were performed on the Rotor-Gene 6000™ (Corbett Research, Sydney, Australia). SYTO® 9 was used as the intercalating dye (Life Technologies, Carlsbad, CA). The reaction mixtures consisted of 25 ng of bisulphite modified template (pre bisulphite conversion amount), 1x PCR buffer, 2.5 mmol/L MgCl\(_2\), final (3 mmol/L in the CDH1 assay), 200 nmol/L of each primer, 200 µmol/L of each dNTP, 5 µmol/L of SYTO 9, 0.5U of HotStarTaq (Qiagen) in a final volume of 20 µL.

The PCR comprised one cycle of 95°C for 15 min, followed by 45 cycles of 95°C for 20 s, annealing at the appropriate temperature (Table 1) for 30 s, 72°C for 30 s, and one cycle of 95°C for 1 min. HRM was performed from 60°C to 90°C, with a temperature increase at the rate of 0.2°C per second for all assays. The annealing temperature was experimentally determined for each assay to ensure only methylated templates were amplified. For each assay, a standard dilution series was run to assess the quantitative accuracy and sensitivity. Fully methylated and fully unmethylated control (WGA product), unmodified control, and no template control were also included in every run. All samples were analysed in duplicate.

The CDH1 MethyLight assay

The same forward primer was used for the MethyLight assay as for the SMART-MSP assay. The reverse primer sequence was: 5’-cgctaattacataaatattctatcg-3’. The probe sequence was FAM-5’-tcgctgtgctgagggtctctg-3’-BHQ (IDT). The two CpG sites between the primers are covered by the probe. The reaction mixtures consisted of 25 ng of bisulphite modified DNA, 1x PCR buffer, 250 mmol/L of probe, 3 mmol/L MgCl\(_2\), 200 mmol/L of each primer, 200 µmol/L of each dNTP, and 0.5U of HotStarTaq (Qiagen) (5U/µL) in a total volume of 20 µL. The PCR comprised one cycle of 95°C for 15 min, 45 cycles of 95°C for 20 s and 64°C for 40 s. PCR was performed on the Rotor-Gene 6000. All samples were analysed in duplicate. In some reactions the probe was omitted and the primers were used for SMART-MSP to allow assessment of the methylation status of the two CpG sites in the region of the probe by HRM analysis.

Real-Time PCR quantification

The COL2A1 control assay amplifying a CpG free region was used to normalise for DNA input after bisulphite conversion in the real-time PCR quantification [12]. The relative 2\(^{-\Delta\Delta CT}\) quantification approach [40] was used. The C\(_T\) value of the control COL2A1 assay (Table 1) is subtracted from the C\(_T\) value of the target gene for the calibrator sample (the 100% methylated standard). For each sample, this value is then subtracted from the value resulting from the C\(_T\) value of the target gene minus the C\(_T\) value for the COL2A1 control assay. For this approach to be valid, the amplification efficiencies of the target and the control must be approximately equal [40]. The take-off values (defined as the cycle at which the second derivative is at 20% of the maximum level) given by comparative quantification (using the Rotor-Gene 6000 Series Software, version 1.7.61) were used as C\(_T\) values in the calculations.

Sequencing

Sequencing was used to verify a higher melting temperature was due to incomplete conversion of some of the non-CpG cytosines in between the APC SMART-MSP primers. For this purpose, a second amplification was performed with m13 tagged APC SMART-MSP primers. These products were sequenced using BigDye terminator chemistry v3.1 on an ABI 3730 (Applied Biosystems).

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