Epigenome-Wide DNA Methylation Profiling in Colorectal Cancer and Normal Adjacent Colon Using Infinium Human Methylation 450K

Rashidah Baharudin 1, Muhiddin Ishak 1, Azliana Muhamad Yusof 2, Sazuita Saidin 1, Saiful Effendi Syafruddin 1, Wan Fahmi Wan Mohamad Nazarie 3, Learn-Han Lee 4,∗ and Nurul-Syakima Ab Mutalib 1,4,5,∗

1 UKM Medical Molecular Biology Institute (UMBI), Universiti Kebangsaan Malaysia, Kuala Lumpur 56000, Malaysia; iedabaharudin90@gmail.com (R.B.); muhiddin@ppukm.ukm.edu.my (M.I.); sazuita@ukm.edu.my (S.S.); effendisy@ppukm.ukm.edu.my (S.E.S.)
2 Cytogenetics & Molecular Diagnostics Laboratory, Pantai Premier Pathology Sdn Bhd, Pandan Indah, Kuala Lumpur 55100, Malaysia; azlianayusof@gmail.com
3 Faculty of Science and Natural Resources, Universiti Malaysia Sabah, Kota Kinabalu 88400, Malaysia; wanfahmi5785@gmail.com
4 Novel Bacteria and Drug Discovery Research Group, Microbiome and Bioresource Research Strength, Jeffrey Cheah School of Medicine and Health Sciences, Monash University of Malaysia, Subang Jaya 47500, Malaysia
5 Faculty of Health Sciences, Universiti Kebangsaan Malaysia, Kuala Lumpur 50300, Malaysia
∗ Correspondence: lee.learn.han@monash.edu (L.-H.L.); syakima@ppukm.ukm.edu.my (N.-S.A.M.); Tel.: +60-391-459-073 (N.-S.A.M.)

Abstract: The aims were to profile the DNA methylation in colorectal cancer (CRC) and to explore cancer-specific methylation biomarkers. Fifty-four pairs of CRCs and the adjacent normal tissues were subjected to Infinium Human Methylation 450K assay and analysed using ChAMP R package. A total of 26,093 differentially methylated probes were identified, which represent 6156 genes; 650 probes were hypermethylated, and 25,443 were hypomethylated. Hypermethylated sites were common in CpG islands, while hypomethylated sites were in open sea. Most of the hypermethylated genes were associated with pathways in cancer, while the hypomethylated genes were involved in the PI3K-AKT signalling pathway. Among the identified differentially methylated probes, we found evidence of four potential probes in CRCs versus adjacent normal; HOXA2 cg06786372, OPLAH cg17301223, cg15638338, and TRIM31 cg02583465 that could serve as a new biomarker in CRC since these probes were aberrantly methylated in CRC as well as involved in the progression of CRC. Furthermore, we revealed the potential of promoter methylation ADHFE1 cg18065361 in differentiating the CRC from normal colonic tissue from the integrated analysis. In conclusion, aberrant DNA methylation is significantly involved in CRC pathogenesis and is associated with gene silencing. This study reports several potential important methylated genes in CRC and, therefore, merit further validation as novel candidate biomarker genes in CRC.

Keywords: colorectal cancer; DNA methylation; adjacent normal colon; Infinium Human Methylation 450K; microarray

1. Introduction

Colorectal cancer (CRC) is a leading cause of morbidity and cancer death worldwide. In Malaysia, CRC is identified as the most common cancer in men and the second most common cancer in women [1]. This disease is highly heterogeneous, with varying responses to cancer therapy and prognosis. The heterogeneity of CRC evolved from multiple pathways, including Chromosomal Instability (CIN), Microsatellite Instability (MSI), and CpG Island Methylator Phenotype (CIMP) [2]. Unlike other pathways, CIMP is an epigenetic mechanism that is more dynamic [3] and often reversible in the presence of
inducing factors such as demethylating agents. The concept of CIMP was first introduced by Toyota et al. in 1999 and defined as widespread methylation in the CpG island of the genes [4]. The methylation process was described by the addition of the methyl group (CH$_3^-$) at the carbon 5 of the cytosine ring to form 5-methylcytosine, and the process is catalysed by the enzyme DNA methyltransferase (DNMT) in which S-adenosyl-methionine (SAM) acts as a methyl donor [5,6]. Aberrant methylation in CRC has correlated with the inactivation of tumour suppressor genes [7] and the activation of oncogenes [8] that function to control a variety of cellular processes, including apoptosis, proliferation, invasion, and migration [9]. Aberrant DNA methylation is a hallmark of cancer that occurs early in cancer development [10], increases with the progression of the disease, and is involved in the treatment response [11,12]. Therefore, DNA methylation may serve as a potential biomarker for cancer diagnosis, predicting patient prognosis, and monitoring response towards therapy. Several studies have been conducted to identify a methylation biomarker with high specificity and sensitivity to be used in the diagnosis of CRC. For instance, Freitas and his colleagues discovered promoter methylation of the three-genes panel (MGMT, RASSF1A, and SEPT9) in accurately diagnosed CRC with 96.6% sensitivity and 74% specificity of detection [13]. Furthermore, a meta-analysis from 38 studies has offered the potential diagnostic markers of hypermethylation SFRP1, SFRP2, NDRG2, and VIM genes in CRC patients [14].

As previously stated, the methylation status of specific genes may also predict the prognosis of the patients. An interesting study by Maija et al. 2013, discovered that the activation of oncogene KRAS along with promoter methylation of CDKN2A leads to more aggressive rectal cancers [15]. In addition, CHFR is another promising prognostic marker whereby promoter methylation of this gene indicates poor prognosis in stage II microsatellite stable CRC [16]. Numerous investigations have identified a therapeutic response mediated by DNA methylation [17–19]. In our previous research, we identified the aberrant methylation of five potential therapeutic targets, CCNE1, CCNDBP1, PON3, CHL1, and DDX43, involved in CRC chemoresistance [20]. Moreover, CRC patients with the hypermethylation of NKX6.1 [21], TFAP2E-DKK4 [22], and IGFBP3 genes [23] were unresponsive to 5-FU chemotherapy treatment. Patients with unmethylated RASSF1A [24] and SRBC [18] were highly responsive to oxaliplatin chemotherapy drugs than patients who exhibited aberrant methylation of these genes. However, the methylation of GPX3 was associated with oxaliplatin sensitivity [25]. The hypermethylation of BNIP3 reduces the sensitivity of CRC towards irinotecan chemotherapeutic drugs [26].

DNA methylation has been recognised as a potential biomarker in CRC; however, only a few methylation markers are currently used in cancer diagnostics. DNA methylation-based biomarkers are still relatively new, hence, careful assessments of the potential biomarkers are required to further validate them prior to being used in clinical diagnostics. Therefore, in this study, we aim to profile the DNA methylation in CRC and explore cancer-specific methylation biomarkers to provide evidence that can support their use in clinical practice.

2. Materials and Methods
2.1. Clinical Specimens

Fifty-four pairs (n = 108) of CRC and the respective adjacent normal tissues were collected from the UKM Medical Center, Malaysia. This study was conducted following the recommendations and approval of the Universiti Kebangsaan Malaysia Research Ethics Committee (Reference number: UKM 1.5.3.5/244/UMBI-004-2012). All subjects gave written informed consent for their participation following the Declaration of Helsinki. The tissues were dissected, snap-frozen and stored in liquid nitrogen prior to sectioning. All sectioned tissues were stained with Hematoxylin and Eosin (H&E). Only cancer tissues that contained more than 80% cancerous cells and normal adjacent tissues with less than 20% necrosis were subjected to the next step. Genomic DNA from frozen tissues was extracted using the QIAamp DNA mini kit according to the manufacturer’s instructions.
The quantification and purity of DNA for each sample were assessed using Qubit 2.0 fluorometer and Nanodrop 2000c Spectrometer (Thermo Fisher Scientific, Inc., Wilmington, DE, USA), respectively. Only samples with purity from 1.8 to 2.0 were selected for the microarray study.

2.2. DNA Methylation Profiling

Methylation profiling was performed on 108 samples (54 paired tumour–adjacent normal colon) using the Infinium Human Methylation 450K BeadChip, which covers 485,577 CpG dinucleotide sites distributed over the whole genome according to the manufacturer’s specification (Illumina, Inc., San Diego, CA, USA). Genomic DNA underwent bisulfite treatment to convert all unmethylated cytosine to uracil using EZ DNA methylation—Gold kit (Zymo Research, Inc., Irvine, CA, USA) following the manufacturer’s protocol prior to being subjected to profiling. Scanning of the BeadChips was performed on the iScan scanner (Illumina, Inc., San Diego, CA, USA).

2.3. Human Methylation 450K Data Analysis

The raw IDAT files were exported from the scanner, and quality control was performed using Genome Studio software version 2.0.4 (Illumina Inc.). The passed IDAT files from 108 samples were further analysed using the ChAMP R package [27] in a single analysis, and filters were applied to all datasets where CpG sites that had a detection p-value > 0.01 in each probe were excluded from further analysis. The data were normalised using the Peak-Based Correction (PBC) method [28] prior to the batch effect correction using ComBat [29]. The $\beta$-values were extracted, and statistical analysis was performed. The limma Bioconductor package was used to determine the differentially methylated CpG sites [30,31], and we applied Benjamini–Hochberg (BH) p-value < 0.05 to identify significant differentially methylated probes. Then, to determine the methylation status of the probes, we conducted further filtering based on the $\Delta\beta$ value of tumour versus normal, where $\geq 0.2$ was considered hypermethylated and $\Delta\beta \leq -0.2$ was hypomethylated. The heatmap was generated using the online Morpheus tool from the Broad Institute [32].

2.4. Functional Enrichment Analyses of Differentially Methylated Genes

The DMGs were then subjected to gene ontology enrichment analysis using the bioinformatics analysis tool, DAVID Bioinformatics Resources 6.8 [33,34], to identify the pathways involved in the hypermethylated and hypomethylated genes. Adjusted p-values < 0.05 were used as the cut-off criterion.

2.5. Gene Expression Profile Analysis

The gene profiling analysis was performed using the level 3 IlluminaHiSeq RNAseqV2 mRNA dataset in patients diagnosed with CRC from The Cancer Genome Atlas (TCGA) data portal. A total of 157 samples were available for gene expression data, with 13 matched normal and tumour samples. The normalised RNAseq by Expectation-Maximization (RSEM) data was input into R programming software, and the limma Bioconductor package was subsequently used for the calculation of differentially expressed genes. The Benjamini–Hochberg (BH) [35] procedure was applied to identify significantly differentially expressed genes between CRC and normal colon tissues with the cut-off criterion of adjusted p-values < 0.05 and log2 fold change (FC) $\geq 1$.

2.6. Integrated Promoter Methylation and Gene Expression Profiling

In order to identify a set of genes whose expression is primarily and causally regulated by promoter DNA methylation in CRC, we performed an integrated analysis of genome-wide DNA methylation and a gene expression profile. The promoter region was defined as the genomic interval that begins 1500 bp upstream and 200 bp downstream of the transcription start site in the CG-rich region. Additionally, the gene expression data were classified into two groups according to their expression level in which differentially ex-
pressed genes were exhibited, log2 FC > 1 were considered upregulated, and log2 FC < −1 in CRC, compared to the normal colon, were considered downregulated. The promoter methylation mediated silencing genes were determined by overlapping the methylated promoter genes with the downregulated genes.

2.7. Protein-Protein Interaction

The interaction between the proteins encoded by top promoter methylated-silencing genes were determined using the Search Tool for the Retrieval of Interacting Genes (STRING) v11 database [36]. All parameters were set to defaults. In the search for the candidate biomarker, a few criteria were utilised, such as genes that posed a high level of methylation and low expression, genes that have strong interaction that can regulate other genes, as well as genes associated with predisposition to CRC.

2.8. Receiver Operating Characteristics (ROC) Curve of Genes

The diagnostic performance of the candidate biomarker was evaluated by the ROC curves. The area under the ROC curve (AUC) was constructed with a 95% confidence interval (95% CI) as an accuracy criterion for the examination of the candidate biomarker. The methylation value of candidate biomarkers in CRC cases was plotted against their corresponding control, and a perfect diagnostic marker had an AUC value of 1. All the analyses were generated using GraphPad Prism 8.0.2.

3. Results

3.1. Demography

Demography data of the 54 patients are presented in Table 1. The majority of the patients were female and above the age of 50 years old. Most of the patients were diagnosed with Duke’s B and positioned on the left side of the colon. Moreover, the majority of the tumour tissues were moderately differentiated.

| Characteristics            | Number of Patients |
|----------------------------|--------------------|
| **Gender**                 |                    |
| Female                     | 31                 |
| Male                       | 23                 |
| **Age (years)**            |                    |
| ≤50                        | 8                  |
| >50                        | 46                 |
| **Ethnicity**              |                    |
| Malay                      | 26                 |
| Chinese                    | 24                 |
| Indian                     | 4                  |
| **Duke’s Staging**         |                    |
| A                          | 5                  |
| B                          | 31                 |
| C                          | 18                 |
| **Tissue differentiation** |                    |
| Well                       | 21                 |
| Moderately                 | 27                 |
| Poorly                     | 3                  |
| Unknown                    | 3                  |
| **Location of the tumour** |                    |
| Right                      | 24                 |
| Left                       | 30                 |

Table 1. Demographic data of the 54 CRC patients.
3.2. Locations of Differentially Methylated Probes

We compared the differential methylation status of 54 CRC tissue samples with the 54 adjacent cancer-free colonic tissue samples. In order to explore epigenome-wide methylation profiles, probe filtering was performed to identify the differentially methylated probes with a detection adjusted \( p \)-value < 0.01 after FDR correction. This resulted in 157,846 probes for the downstream analysis. These probes were further classified as hypermethylated or hypomethylated based on the absolute average \( \beta \) value difference (\( \Delta \beta \)) at \( \geq 0.2 \) between CRC and normal adjacent tissues. This value represents 20% change in the methylation level. A total of 26,093 probes were identified (Figure 1A). Of these, 650 probes were hypermethylated and 25,443 probes were hypomethylated. Then, we stratified the probes into CpG island, shores, shelves, and open sea regions. From the 650 differentially hypermethylated probes, 331 probes (50.92%) were located in the island region, accounting for half of the total number of probes, 145 probes (22.31%) were located on the shore, 112 probes (17.23%) were in the open sea, and the remaining 62 probes (9.54%) were located in the shelf region (Figure 1B). In contrast, most of the hypomethylated probes were in the open sea area of the genome (\( n = 16,749; 65.83% \)), followed by the shore, shelf, and island region with 4022 (15.81%), 3657 (14.37%), and 1015 (3.99%) probes, respectively (Figure 1C).

![Figure 1. Differentially methylated probes in CRCs relative to its adjacent normal. (A) Distribution of significantly differentially methylated probes in a genomic region. (B) Hypermethylated probes and (C) Hypomethylated probes in genomic region. (D) Distribution of differentially methylated probes, (E) hypermethylated probes, (F) Hypomethylated probes with respect to genomic features, (G) distribution of methylation CpG sites in the human genome. The CpG island is surrounded by shores (within 2 kb sequence neighbouring the islands) with shelves flanked further from the shores. The open sea area is outside of the shelf region.](image)

Meanwhile, categorization based on genomic features revealed that most differentially methylated probes did not belong to any gene. The majority of these probes were in the intergenic region (IGR) (\( n = 10,150; 38.90% \)), closely followed by the gene body (\( n = 9211; 35.30% \)), TSS1500 (\( n = 2888; 11.07% \)), 5’UTR (\( n = 1524; 5.84% \)), 3’UTR (\( n = 1047; 4.01% \)), TSS200 (\( n = 762; 2.92% \)), and 1st exon (\( n = 511; 1.96% \)), as illustrated in Figure 1D. On closer inspection, the majority of the significantly hypermethylated loci were in the body (\( n = 250; 38.46% \)), followed by 5’UTR (\( n = 107; 16.46% \)), the intergenic region (\( n = 80; 12.31% \)), closely
followed by TSS200 ($n=78; 12\%$), TSS1500 ($n=72; 11.08\%$), 1st exon ($n=35; 5.38\%$), and lastly 3′UTR ($n=28; 4.31\%$) (Figure 1E). On the contrary, the significant hypomethylated loci were not associated with any genes ($n=10,150; 39.58\%$) or in the gene body ($n=9211; 35.22\%$), while the rest were mainly located in TSS1500, 5′UTR, 3′UTR, TSS200, and 1st exon (Figure 1F).

### 3.3. Methylation Level of Differentially Methylated Probes

Significant methylation differences of the 50 topmost significant differentially methylated probes were generated and illustrated through the heatmap in Figure 2. The five hypermethylated probes with the highest $\Delta \beta$ values were $SEPT9$ cg17300544 ($\Delta \beta = 0.353$), $HOXA2$ cg06786372 ($\Delta \beta = 0.342$), $HOXA3$ cg27539480 ($\Delta \beta = 0.330$), OPLAH cg17301223 ($\Delta \beta = 0.317$), and cg16179589 ($\Delta \beta = 0.315$). The list of the top 10 hypermethylated probes is displayed in Table 2, and information on the top 50 is provided in Supplementary Table S1. On the other hand, the five hypomethylated probes with the highest reduction of methylation were $ZBTB46$ cg20267897 ($\Delta \beta = -0.497$), cg15638338 ($\Delta \beta = -0.497$), $MATN4$ cg01268752 ($\Delta \beta = -0.496$), cg08550523 ($\Delta \beta = -0.495$), and $TRIM31$ cg02583465 ($\Delta \beta = -0.495$). The list of the top 10 hypomethylated probes is displayed in Table 3, and information on the top 50 hypomethylated probes is provided in Supplementary Table S2.

#### Table 2. Top 10 significant hypermethylated loci in CRC versus normal tissues.

| Gene     | Probes     | $\Delta \beta$ | Genomic Feature | Genomic Region |
|----------|------------|----------------|-----------------|----------------|
| $SEPT9$  | cg17300544 | 0.353          | TSS200          | Island         |
| $HOXA2$  | cg06786372 | 0.342          | Body            | Shore          |
| $HOXA3$  | cg27539480 | 0.330          | 3′UTR           | Shore          |
| OPLAH    | cg17301223 | 0.317          | Body            | Island         |
| na       | cg16179589 | 0.315          | IGR             | Shore          |
| IRF4     | cg17228900 | 0.314          | 5′UTR           | Island         |
| ADHFE1   | cg20912169 | 0.311          | 5′UTR           | Island         |
| $PRKAR1B$| cg18601167 | 0.310          | 5′UTR           | Shore          |
| $ZFHX3$  | cg02973693 | 0.308          | 5′UTR           | Shelf          |
| $HOXA2$  | cg00188704 | 0.307          | Body            | Shelf          |

na = not available. The probes in the IGR region were not annotated with a gene name.

#### Table 3. Top 10 significant hypomethylated loci in CRC versus normal tissues.

| Gene     | Probes     | $\Delta \beta$ | Genomic Feature | Genomic Region |
|----------|------------|----------------|-----------------|----------------|
| $ZBTB46$ | cg20267897 | -0.497         | 5′UTR           | Shore          |
| na       | cg15638338 | -0.497         | IGR             | Open sea       |
| $MATN4$  | cg01268752 | -0.496         | Body            | Shore          |
| na       | cg08550523 | -0.495         | IGR             | Open sea       |
| $TRIM31$ | cg02583465 | -0.495         | Body            | Open sea       |
| na       | cg17400812 | -0.493         | IGR             | Open sea       |
| na       | cg25506686 | -0.483         | IGR             | Open sea       |
| na       | cg12297066 | -0.477         | IGR             | Open sea       |
| $TM4SF19$| cg05445326 | -0.476         | TSS1500         | Open sea       |
| $OC90$   | cg03344782 | -0.470         | Body            | Open sea       |

na = not available. The probes in the IGR region were not annotated with a gene name.
Figure 2. Heatmap of 50 differentially methylated genes, consisting of 25 hypermethylated and 25 hypomethylated genes in CRCs in comparison to the adjacent normal tissues with a p-value < 0.05. Every row represents individual genes, and the column represents individual samples. The horizontal bars indicate the patient’s age at diagnosis and tumour information, such as the tumour’s location and stage. The colour in each small box constitutes the methylation level of the genes in which red boxes indicate genes with a high methylation level while blue boxes display genes with a low methylation level.

Of the 26,093 differentially methylated CpGs identified from the probe-level test, 15,943 (61.1%) represented 6156 genes. There were 156 genes with overlapping methylation status; 5781 were uniquely hypomethylated, and 219 were uniquely hypermethylated (Figure 3A, Supplementary Table S3). The HOXA5 gene had the highest number of differentially methylated loci (n = 21), followed by HOXA3 (n = 13) and HOXA2 (n = 12), in which all loci were hypermethylated (Figure 3B). Twenty loci were in the HOXA5 islands, and one locus was at the shore. On the contrary, less than one-third of the hypermethylated loci in HOXA3 were at the island; almost half (46%) were at the shore. HOXA2 followed an almost similar trend, with the hypermethylated loci mainly at the shore and only one locus at the island. Another member of the HOX gene family, HOXA6, was also identified with three hypermethylated loci (data not shown). Other than these three genes, the majority (n = 284; 75.7%) of the genes had only one hypermethylated site.
Figure 3. Overview of the methylation status identified in 6156 genes. (A) A total of 156 genes with overlapping methylation status and 5781 genes exhibit hypomethylation; in contrast, 219 genes were hypermethylated. (B) Bar chart displaying genes that had the highest differentially hypermethylated loci. (C) The bar chart details the genes that had the highest differentially hypomethylated loci.

There were more hypomethylated genes as compared to the number of hypermethylated genes. 

SDK1, PTPRN2, and TNXB were the genes with the highest number of hypomethylated loci (n = 116, 83, and 83, respectively) (Figure 3C). As expected, the majority of the hypomethylated loci in SDK1 were at the open sea (81.8%), with PTPRN2 and TNXB following a similar pattern.
3.4. Pathway Enrichment Analysis of DMG

Next, the list of hypermethylated and hypomethylated genes were subjected to pathway enrichment analysis using DAVID Functional Annotation Bioinformatics Microarray Analysis. We discovered ten enriched pathways of hypermethylated genes that could be potentially associated with CRCs. The top five most enriched pathways were pathways in cancer, the PI3K/Akt signalling pathway, signalling pathways regulating the pluripotency of stem cells, Proteoglycans in cancer, and Melanoma (Table 4). However, none of the listed pathways was statistically significant. Most of the genes that were hypermethylated in the pathways in cancer were APC, CTBP2, NFKBIA, SMAD2, COL4A1, CDK6, DAPK2, FGF12, FGF14, FGF21, IGF1R, LAMB3, as well as PIK3R1.

Table 4. Five top pathways regulated by the hypermethylated genes in CRCs compared to adjacent normal tissues.

| Pathway                              | No. of Genes | p-Value | Fold Enrichment |
|--------------------------------------|--------------|---------|-----------------|
| Pathways in cancer                   | 14           | 0.6873  | 1.9923          |
| PI3K/Akt signalling pathway          | 13           | 0.6873  | 2.1073          |
| Signalling pathways regulating       |              |         |                 |
| pluripotency of stem cells           | 10           | 0.1544  | 3.9947          |
| Proteoglycans in cancer              | 9            | 0.6960  | 2.5167          |
| Melanoma                             | 6            | 0.6873  | 4.7262          |

Conversely, the top five significantly enriched pathways associated with hypomethylated genes were the PI3K/AKT signalling pathway, pathways in cancer, focal adhesion, cell adhesion molecules, and the RAS signalling pathways (Table 5). Hypomethylated genes were observed in the PI3K/AKT signalling pathways, namely CHRM2, TNXB, LAMA2, COL11A2, PIK3CD, PIK3CG, COMP, RPTOR, MYC, GNG7, PDGFD, AKT3, PDGFC, TNR, SYK, ANGPT1, ITGA4, IGF1, NGF, PTK2, RBL2, FGF14, CDK6, COL1A2, COL4A2, COL5A1, COL4A1, COL5A2, FGF18, COL6A3, COL6A6, SOS1, FGF1R, and CREB5.

Table 5. Five top pathways regulated by the hypomethylated genes in CRCs compared to adjacent normal tissues.

| Pathway                              | No. of Genes | p-Value | Fold Enrichment |
|--------------------------------------|--------------|---------|-----------------|
| PI3K/Akt signalling pathway          | 34           | 2.89 × 10^{-5} | 2.6902          |
| Pathways in cancer                   | 28           | 0.0254  | 1.9448          |
| Focal adhesion                       | 23           | 3.73 × 10^{-4} | 3.0477          |
| Cell adhesion molecules (CAMs)       | 21           | 2.89 × 10^{-5} | 4.0369          |
| Ras signalling pathway               | 19           | 0.0314  | 2.2949          |

3.5. Integrated Analysis of Promoter Methylation and Gene Silencing

The hypermethylation of the promoter region has been associated with the silencing of the genes; meanwhile, cancer-linked DNA hypomethylation has received little attention due to the association with repeated DNA elements. We analysed the relationship between promoter hypermethylation and gene expression by integrating the differentially methylated genes (DMGs) and differentially expressed genes (DEGs). We identified a total of 105 hypermethylated genes in the promoter regions (comprised of transcription start sites; TSS1500 and TSS200 within the CG rich region).

To address whether the promoter methylation plays a role in the regulation of the gene expression, we observed the expression level of promoter hypermethylated genes using RNAseq data from The Cancer Genome Atlas (TCGA) of 134 CRCs and 23 normal colon samples. Out of 105 promoters hypermethylated, only 31 genes overlapped with
DEGs from the TCGA datasets. The list of the promoter methylated genes, Δβ value, and corresponding expression level are displayed in Supplementary Table S4.

Then, from 31 overlapped genes, 28 genes exhibited hypermethylation associated with gene silencing in CRCs as compared to normal, whereas the three remaining genes displayed expression levels directly proportional to the methylation level.

The top ten hypermethylated-induced silencing of genes in the promoter region were ADHFE1 (Δβ = 0.299, log2 fold change = −3.531), HOXA5 (Δβ = 0.271, log2 fold change = −1.419), ZNF542 (Δβ = 0.261, log2 fold change = −1.675), ZNF334 (Δβ = 0.259, log2 fold change = −1.887), ZNF135 (Δβ = 0.258, log2 fold change = −1.720), USP44 (Δβ = 0.255, log2 fold change = −1.225), SFMBT2 (Δβ = 0.249, log2 fold change = −1.634), ADARB2 (Δβ = 0.248, log2 fold change = −1.158), ZNF582 (Δβ = 0.244, log2 fold change = −1.577), and ZNF132 (Δβ = 0.244, log2 fold change = −1.379). This is illustrated in Figure 4.

**Figure 4.** The methylation and expression level of the top ten hypermethylated genes are associated with gene silencing. The (A) ADHFE1, (B) HOXA5, (C) ZNF542, (D) ZNF334, (E) ZNF135, (F) USP44, (G) SFMBT2, (H) ADARB2, (I) ZNF582 and (J) ZNF132 genes were significantly hypermethylated and downregulated in CRC tumours versus normal tissues. The box plot displays the average methylation level of each gene across 56 pairs of CRC tissues and their adjacent normal tissues from our microarray analysis. The green box plot represents the methylation level of the respective genes in CRC tumour tissues, while the yellow box plot represents normal colon tissues. The expression level of the genes in CRC tissues versus normal is shown on the right side of the methylation box plot graph, using 134 CRC tissues and 23 normal colon tissues from the TCGA dataset. The green box plot graph indicates the level of expression of the corresponding genes in CRC tissues, whereas the yellow box plot graph indicates the level of expression of the genes in normal tissues. The statistical analysis between tumour versus normal was determined using a two-sided Student’s t-test (**** p-value < 0.0001).
3.6. Protein-Protein Interaction of Promoter Hypermethylated Genes

To explore the potential function of each protein encoded by the top ten promoter hypermethylated-silencing genes, we constructed a protein-protein interaction network using the STRING database Version 11.0 based on the homo sapiens association model. In this network, each node represents a protein, and each edge represents a physical interaction between two proteins.

The protein-protein interaction network of ADHFE1 resulted in 10 nodes and 18 edges with the enrichment $p$-value of 0.0181. The biological process of this protein is mainly involved in alcohol dehydrogenase activity, the retinol metabolism process, as well as ethanol oxidation.

Next, HOXA5 consisted of 10 nodes and 55 edges. The strong interaction of HOXA5 with other proteins provided the most significant enrichment $p$-value of $1 \times 10^{-16}$. This protein is involved in DNA binding transcription activator activity, embryonic skeletal system morphogenesis, and development. The following protein, SFMBT2, was connected to 10 proteins and 13 edges. The protein-protein interaction enrichment $p$-value of SFMBT2 was 0.224, and it is involved in chromatin binding, histone binding, as well as transcription corepressor activity.

The zinc-finger (ZNF) proteins ZNF135, ZNF582, ZNF132, and ZNF334 consisted of 11 nodes and 15 edges, 11 nodes and 14 edges, 11 nodes 12 edges with an enrichment $p$-value of 0.112, 0.15, 0.337, respectively. Among the zinc-finger families, ZNF334 displayed the least interaction with neighbouring proteins with 6 nodes and 5 edges, and the enrichment $p$-value of this protein was 0.572. ZNF families are involved in transcriptional regulation, ubiquitin-mediated protein degradation, signal transduction, and other cellular processes.

Another protein, namely USP44, is involved in regulating ubiquitin-protein ligase activity and cell division, specifically in the G2/M transition phase. USP44 protein was connected to 10 different proteins with 12 edges with an enrichment $p$-value of 0.353. On top of that, the protein interaction network of ADARB2 and ZNF542 could not be retrieved from the STRING database. Eight protein-protein interactions of promoter hypermethylated-silencing genes are displayed in Figure 5.

3.7. Receiver Operating Characteristics (ROC) Curve Analysis of Promoter Hypermethylated Genes

Next, to evaluate the diagnostic power of the genes as a biomarker, we performed the ROC analysis on the top ten promoter hypermethylated genes by measuring the specificity and sensitivity performance of the biomarkers ($p$-value < 0.0001) (Table 6). Among the top ten promoter hypermethylated genes, ADHFE1 had the highest discriminative power (AUC = 0.9088, 95% CI = 0.847 to 0.971), followed by SFMBT2 (AUC = 0.880, 95% CI = 0.818 to 0.942), and ZNF135 (AUC = 0.859, 95% CI = 0.879 to 0.933).
SFMBT2 was 0.224, and it is involved in chromatin binding, histone binding, as well as transcription corepressor activity.

The zinc-finger (ZNF) proteins ZNF135, ZNF582, ZNF132, and ZNF334 consisted of 11 nodes and 15 edges, 11 nodes and 14 edges, 11 nodes and 12 edges with an enrichment p-value of 0.112, 0.15, 0.337, respectively. Among the zinc-finger families, ZNF334 displayed the least interaction with neighbouring proteins with 6 nodes and 5 edges, and the enrichment p-value of this protein was 0.572. ZNF families are involved in transcriptional regulation, ubiquitin-mediated protein degradation, signal transduction, and other cellular processes.

Another protein, namely USP44, is involved in regulating ubiquitin-protein ligase activity and cell division, specifically in the G2/M transition phase. USP44 protein was connected to 10 different proteins with 12 edges with an enrichment p-value of 0.353. On top of that, the protein interaction network of ADARB2 and ZNF542 could not be retrieved from the STRING database. Eight protein-protein interactions of promoter hypermethylated-silencing genes are displayed in Figure 5.

**Figure 5.** Interaction network of eight selected proteins from promoter hypermethylated-gene silencing. STRING Database Version 11.0b was used to construct the functional protein association networks based on the homo sapiens association model. Medium confidence was set to 0.400, and the max number of interactors was no more than 10. (A) The ADHFE1 protein is significantly connected to 10 neighbouring proteins with 18 edges, (B) The HOXA5 is strongly connected to 10 neighbouring proteins with 55 edges, (C) The ZNF 334 protein has the least interaction with neighbouring proteins with 6 nodes and 5 edges, (D) The SFMBT2 protein is connected to 10 neighbouring protein with 13 edges, (E) The ZNF135 protein shows interaction with 11 neighbouring proteins and 15 edges, (F) The USP44 protein is connected to 10 neighbouring proteins with 12 edges, (G) The ZNF582 interacts with 11 neighbouring proteins with 14 edges, (H) The ZNF132 protein interacts with 11 neighbouring proteins with 12 edges.

| Gene     | Probes      | AUC      | 95% Confidence Interval | p-Value |
|----------|-------------|----------|-------------------------|---------|
| ADHFE1   | cg18065361  | 0.909    | 0.8470–0.9706           | <0.0001 |
| HOXA5    | cg19643053  | 0.776    | 0.6876–0.8639           | <0.0001 |
| ZNF542   | cg27477373  | 0.844    | 0.7656–0.9230           | <0.0001 |
| ZNF334   | cg10140114  | 0.826    | 0.7432–0.9090           | <0.0001 |
| ZNF135   | cg06454760  | 0.859    | 0.7855–0.9326           | <0.0001 |
| USP44    | cg13879483  | 0.829    | 0.7445–0.9140           | <0.0001 |
| SFMBT2   | cg02866454  | 0.880    | 0.8180–0.9420           | <0.0001 |
| ZNF582   | cg13916740  | 0.809    | 0.7198–0.8982           | <0.0001 |
| ZNF132   | cg03735888  | 0.842    | 0.7654–0.9184           | <0.0001 |
| ADARB2   | cg02899206  | 0.79     | 0.7055–0.8810           | <0.0001 |
In the previous analysis, we discovered HOXA5 exhibited a strong protein-protein interaction network; however, the ROC analysis for HOXA5 showed the lowest discriminative power with an AUC value of 0.776 to differentiate the CRC from normal mucosa tissues.

3.8. Methylation Level of ADHFE1 in Various Cancers

From the ROC analysis, we identified the potential of the ADHFE1 gene as a diagnostic marker in CRC. All probes identified in the promoter region of the ADHFE1 gene were hypermethylated, with cg18065361 exhibiting the highest methylation level in colorectal tumours versus normal tissues. The methylation status of the ADHFE1 gene in each probe in the promoter region is illustrated in Figure 6.

Then, we performed an in silico methylation analysis of this gene in various cancers using Wanderer software [37] to compare the diagnostic potential of this gene in CRC as well as other cancers. This software provides the methylation level of all the Human Methylation 450K probes in the ADHFE1 gene. Twenty-seven CpG loci in ADHFE1 were found to be significantly aberrant methylated in 308 CRCs versus 38 normal tissues, including our CpG locus of interest; cg18065361.

The methylation of ADHFE1 cg18065361 was also significantly hypermethylated in esophageal carcinoma and head and neck squamous cell carcinoma. However, among these two cancers, CRC showed a significant difference in the methylation level of ADHFE1 between tumours and normal tissues. Thus, based on our results and from the TCGA dataset, we concluded that the methylation of the ADHFE1 gene occurs more frequently in CRC than in other malignancies. Figure 7 presents the methylation status of the ADHFE1 gene in tumours versus normal tissues from various cancers using TCGA datasets.
Then, we performed an in silico methylation analysis of this gene in various cancers using Wanderer software [37] to compare the diagnostic potential of this gene in CRC as well as other cancers. This software provides the methylation level of all the Human Methylation 450K probes in the ADHFE1 gene. Twenty-seven CpG loci in ADHFE1 were found to be significantly aberrant methylated in 308 CRCs versus 38 normal tissues, including our CpG locus of interest; cg18065361. The methylation of ADHFE1 cg18065361 was also significantly hypermethylated in esophageal carcinoma and head and neck squamous cell carcinoma. However, among these two cancers, CRC showed a significant difference in the methylation level of ADHFE1 between tumours and normal tissues. Thus, based on our results and from the TCGA dataset, we concluded that the methylation of the ADHFE1 gene occurs more frequently in CRC than in other malignancies. Figure 7 presents the methylation status of the ADHFE1 gene in tumours versus normal tissues from various cancers using TCGA datasets.

Figure 7. In silico validation of ADHFE1 methylation in the TCGA dataset of selected cancers. The red box indicates the potential hypermethylated locus identified in our study. We displayed a concordant result with the TCGA dataset. Green probes indicate CpG island location, and the figure was generated using Wanderer software [36].

4. Discussion

We analysed the global methylation status of 54 paired CRC and the corresponding normal tissue samples. Our demographic data showed that most patients were diagnosed over 50 years of age. Many studies have reported an increase in the incidence rate of CRC among individuals aged more than 50 years old [1,38]. To the best of our knowledge, this study provides the largest epigenome-wide DNA methylation profiles in CRC–adjacent normal colon tissue pairs using the 450K BeadChip. The Cancer Genome Atlas [39] has data on the DNA methylation status in 308 CRCs; however, only 38 matched tumour–adjacent normal samples were included. In 2013, Naumov and colleagues performed genome-wide methylation profiling in 22 paired CRC and adjacent normal tissues in addition to 19 colon tissue samples from cancer-free donors [40]. Recently, Gu et al. analysed 12 pairs of CRC and adjacent normal tissues using the newest version of methylation chip; the MethylationEpic Beadchip [41].

Our study revealed 26,093 differentially methylated probes that were distributed over the CpG sites of the genome. The CpG sites were comprised of several regions such as CpG island, shores (2 kb upstream from the island), shelves (2 kb upstream from the shores), and open sea, which make up for the remaining genomic region [42]. Notably,
we discovered that most of the identified CpG sites were hypomethylated rather than hypermethylated, contrasting with previous findings in other cancers [43,44]. Our findings corroborated recent research by Gu and colleagues, which revealed that approximately 87% of differentially methylated CpG sites were hypomethylated, whereas just 13% were hypermethylated [41]. This may be explained by the genomic distribution of the Human-Methylation 450K microarray, which identified more probes in the open sea region (36.3%) than in CpG islands (30.9%), shores (23%), and shelves (9.7%) [45,46].

The CpG island is a region that is rich in CG sequences and often associated with the transcription start site. Our results showed that half of the hypermethylated probes were in the CpG island which is associated with promoter regions. This finding is supported by Sproul et al. where they showed that most of the cancers frequently exhibit hypermethylation at the CpG rich regions [47]. Conversely, hypomethylation often occurs in the open sea area of the genome. A similar pattern has been observed in another study where hypomethylation of CpG sites was enriched at the open sea area and intergenic region [48–50].

Various studies have reported septin 9 gene (SEPT9) methylation in CRC, highlighting the relevance of SEPT9 methylation in cancer [51–56]. SEPT9 is one of the widely studied hypermethylated genes in CRC, and our finding also supported its role in this cancer. We also uncovered many potential genes with interesting profiles. For instance, a group of Homeobox A (HOXA) cluster genes, the members of the HOX family, and an important gene in normal organ development was found to be significantly hypermethylated in CRC compared to the normal adjacent tissues. On top of that, four genes in this family, namely HOXA5, HOXA3, HOXA2, and HOXA6, were hypermethylated at multiple loci. Our discovery is supported by a recent study by Li and colleagues, who reported the hypermethylation of HOXA5, HOXA2, and HOXA6 [57]. Furthermore, HOXA5 methylation was shown to be associated with age, stage, and tumour status, while HOXA6 methylation was linked to age and KRAS mutation [57]. The HOXA family has been the subject of substantial research in cancer. Numerous clinical trials have been conducted on the HOXA genes, but none have focused exclusively on the HOXA2 gene. Recently, the methylation level of HOXA1 was used to accurately differentiate between cholangiocarcinoma and benign biliary stricture from brushed biliary samples in clinical trial NCT04568512 [58]. HOXA9 is among the biomarkers studied in the myeloid leukaemia clinical trial NCT03701295. The expression level of HOXA9 was measured in myeloid leukaemia after treatment with chemotherapeutic drugs, including Pinometostat and Azacitidine [59]. Despite the fact that several HOXA family genes have been translated into clinical trials, none have focused entirely on the HOXA2 gene. The relationship of HOXA2 with cancer progression is limited, and the role of HOXA2 in cancer prognosis and response to treatment is unknown. In 2019, Li and colleagues established a link between HOXA2 and age, cancer staging, lymphovascular invasion, and lymph node involvement in CRC [57]. However, this is the only study within the last decade that revealed the significance of HOXA2 in the clinicopathological characteristics of CRC. The identification of hypermethylation of the HOXA2 gene as a biomarker for CRC in our study adds to the evidence of HOXA2’s association with cancer. Hence, our research may aid in the clinical development of HOXA2. In addition, it will be interesting to assess the clinicopathological correlation with the methylation status of HOXA2 genes in our patients, which will be a subject for future research.

Human 5-oxoprolinase, OPLAH, was shown to be hypermethylated in our study, which had a significant impact on the gene’s downregulation, suggesting a possible contribution to CRC through the dysregulation of gene expression. Numerous investigations have also revealed that OPLAH is frequently hypermethylated in CRC versus normal tissues [40,60]. Despite a paucity of information on OPLAH methylation in cancer, several patents have been filed for its applicability in cancer detection. Recently, OPLAH was identified as one of the biomarkers that have been patented for the diagnosis of lung cancer (patent number: US 11028447 B2) [61] and CRC (patent number: US 11078539 B2) [62]. Taken together, our findings suggest that hypermethylated OPLAH has a role in
the identification of CRC in the Malaysian population. In the future, the methylation status of *OPLAH* can be determined in blood, urine, and saliva, thereby establishing *OPLAH* as a non-invasive biomarker and accelerating the translation of molecular evidence to clinical practice.

Next, we discovered that *TRIM31* displayed a global loss of DNA methylation in CRC tissues. DNA hypomethylation is a common epigenetic alteration observed in human oncogenes. *TRIM31* is an oncogene that has been shown to be overexpressed in different types of cancer, including pancreatic [63], acute myeloid leukaemia [64], hepatocellular carcinoma [65], breast [66], and CRC [67]. Whilst *TRIM31* is commonly overexpressed in cancers, the mechanism by which *TRIM31* is overexpressed remains unknown. Our study is the first to demonstrate that *TRIM31* is hypomethylated in CRC. DNA hypomethylation may enhance gene expression by allowing transcription factors to bind to the promoter region of the gene. Therefore, we postulated that the overexpression of *TRIM31* in CRC is mediated by a global loss of DNA methylation. Nonetheless, additional research is necessary to confirm the association between DNA hypomethylation and *TRIM31* overexpression. In addition, increased *TRIM31* expression was associated with an aggressive phenotype and poor prognosis in pancreatic cancer. Moreover, this gene lowered the chemosensitivity of pancreatic cancer to gemcitabine, a commonly used chemotherapy treatment for pancreatic cancer [63]. In CRC, the high expression of *TRIM31* promotes cancer invasion and metastasis [67]. According to the findings presented, the overexpression of *TRIM31* had a role in the development, invasion, and metastasis of cancer, as well as resistance to chemotherapy.

Barrow and colleagues [68] performed an epigenome-wide analysis of DNA methylation in CRC patients with different smoking statuses, and among the significant hypomethylated genes were the *SDK1*, *PTPRN2*, and *TNXB* genes. In our study, we identified 116 hypomethylated *SDK1* loci and 83 hypomethylated loci in each of the *PTPRN2* and *TNXB* genes. These three genes also contained the highest number of hypomethylated loci. Sidekick cell adhesion molecule 1 (*SDK1*) hypomethylation was also reported in sporadic colorectal cancer [69] and is concordant with our findings. Receptor-type tyrosine-protein phosphatase N2 (*PTPRN2*) hypomethylation, on the other hand, has been rarely reported. The *TNXB* (tenascin X) gene was first implicated in Ehlers–Danlos syndrome [70], but its role in several human cancers have been established, including nasopharyngeal [71] and mesothelioma [72]. Recent evidence further supports its role in malignancy, whereby *TNXB* is indicated as one of the triple-evidenced genes, which exhibit aberrant methylation, differentially expressed and associated with somatic mutation, hence, displaying the superior predictive ability in cancer diagnosis and prognosis [73].

The relationship between DNA methylation and gene expression is highly complex, and traditionally, DNA methylation-silenced gene expression was primarily affected in the CpG island promoter region [74–76] since the regulation of expression is controlled by transcription factors that bind to the promoter [77]. To gain further insight into the role of promoter methylation in silencing the gene expression, we performed an in silico analysis of gene expression profiles of normal and CRC samples from the TCGA datasets. We focused on the methylation profile of the promoter methylated genes and the association with the gene expression. Our analysis showed supporting evidence that genes associated with hypermethylated promoters display reduced gene expressions in CRC patients [78,79]. The addition of a methyl group at the promoter of the genes inhibits the binding of the transcription factor to the promoter region, hence, initiating the activation of genes [80].

A protein-protein interaction network was constructed to identify the regulator protein among the selected promoter hypermethylated. *HOXA5* and *ADHFE1* had strong protein-protein interactions with neighbouring proteins to form a regulatory network. The strong regulatory protein may influence the expression of neighbouring proteins, which contribute to the pathogenesis of CRC. Furthermore, the strong connection among the proteins is likely to form a protein complex and often direct the biological processes [81,82]. Our findings discovered the interaction of *HOXA5* with *HOXB6* and *HOXB7*. The downregulation of
HOXA5 increased the expression of HOXB6 and HOXB7, which were associated with poor clinical outcomes in cancer patients [83–85]. On the other hand, the hypermethylation of ADHFE1 further reduced the expression of neighbouring proteins, ADH6, ADH7, as well as ADH1A, in which the genes were associated with the patient’s prognosis and cancer pathogenesis [86–88].

Taken together, this summarises the potential of HOXA5 and ADHFE1 as promising biomarkers in CRC. However, when we performed ROC analysis for the top ten hypermethylated promoters mediating gene silencing, ADHFE1 showed the highest discriminative values that significantly differentiated the CRCs from the normal colonic tissues. Compared with HOXA5, the better discrimination of normal and cancer tissues by ADHFE1 signifies the great potential for this gene as a methylation marker to indicate pathological changes. ADHFE1, known as Alcohol Dehydrogenase Ion Containing 1, is a member of the iron-activated alcohol dehydrogenase family [89]. Consistent with previous studies, we observed that the CpG island methylation status of the ADHFE1 promoter was higher in CRC tissues in contrast to their adjacent normal mucosa, and the loss expression of ADHFE1 in CRC was associated with promoter methylation [90,91].

Our additional methylation analysis using the Wanderer database further strengthens the diagnostic potential of hypermethylated ADHFE1 in detecting CRC as we observed that the methylation of ADHFE1 displayed significant differences in CRCs versus normal tissues as compared to other cancers, for instance, glioblastoma multiforme (GBM), breast-invasive carcinoma, esophageal carcinoma, lung adenocarcinoma, head and neck squamous cell carcinoma, prostate adenocarcinoma, and thyroid carcinoma. The potential of ADHFE1 as early detection of CRC was also discovered by Fan and his colleagues, whereby they observed hypermethylated ADHFE1 in colorectal adenoma [92]. According to Moon et al., the hypermethylation of the ADHFE1 gene promotes cell proliferation in CRC [91]. This finding was supported by Hu and colleagues, who demonstrated that the hypermethylation of the ADHFE1 gene enhances CRC proliferation via altering cell cycle progression [89]. ADHFE1 acted as a tumour suppressor gene in esophageal squamous cell carcinoma and was reported to be hypermethylated in a Chinese Han population [93]. More recently, the downregulation of the ADHFE1 gene has been linked to decreased cancer survival [94]. Additionally, this study discovered that ADHFE1 might contribute to cancer progression through its interactions with signalling pathways such as energy metabolism, DNA replication, and the cell cycle. With the evidence provided, we believed the promoter methylation-mediated downregulated gene of ADHFE1 could be one of the potential DNA methylation biomarkers in detecting CRC. However, the diagnostic roles of ADHFE1 will be subjected to further validation in a larger patient cohort in our country. It will be interesting if the methylation of ADHFE1 can be detected in the blood for an early, accurate, and non-invasive blood-based biomarker.

5. Conclusions

This relatively extensive methylation study has revealed several potentially important genes in CRC that may be potential biomarker candidates. The hypomethylated and hypermethylated genes reported in this study are relevant to carcinogenesis and are in concordance with other studies. We also offer the first evidence for the potential of HOXA2 cg06786372 OPLAH cg17301223, cg15638338, and TRIM31 cg02583465 as diagnostic biomarkers for CRC. On top of that, we also discovered the potential of promoter methylation ADHFE1 as a biomarker for CRC diagnosis. This is also the first insight into the epigenome-wide methylation profile of the cancer-adjacent normal colon in Malaysian CRC patients to complement the majority of data available from other populations. The new knowledge from this study can be utilised to advance our understanding of CRC methyomics; however, some of the findings need further investigations to confirm the involvement of the candidate genes in CRC.
Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/diagnostics12010198/s1. Table S1: Top 50 hypermethylated genes in CRC compared to the normal adjacent colon, Table S2: Top 50 hypomethylated genes in CRC compared to the normal adjacent colon, Table S3: List of differentially methylated genes and Table S4: The list of promoter hypermethylation genes with $\Delta \beta$ value and their corresponding expression level.

Author Contributions: Conceptualisation, N.-S.A.M.; methodology, R.B., M.I., A.M.Y. and S.S.; software, R.B.; formal analysis, N.-S.A.M. and R.B.; writing—original draft preparation, R.B.; writing—review and editing, N.-S.A.M., L.-H.L., W.F.W.M.N. and S.E.S.; supervision, N.-S.A.M. and S.E.S.; project administration, N.-S.A.M.; funding acquisition, N.-S.A.M. All authors have read and agreed to the published version of the manuscript.

Funding: This study is supported by the University Research Grant Scheme from the Universiti Kebangsaan Malaysia (GUP-2018-070), and the Higher Institution Centre of Excellence (HiCOE) grant (HiCOE 10-64-01-005) from the Ministry of Higher Education.

Institutional Review Board Statement: This study was carried out in accordance with the recommendations of the Universiti Kebangsaan Malaysia Research Ethics Committee (Reference number: UKM 1.5.3.5/244/UMBI-004-2012). The protocol was approved by the Universiti Kebangsaan Malaysia Research Ethics Committee.

Informed Consent Statement: This study was written informed consent from all subjects. All subjects gave written informed consent in accordance with the Declaration of Helsinki.

Data Availability Statement: The raw data of microarray methylation can be obtained in GEO under accession GSE193535.

Acknowledgments: The authors thank Monash University Malaysia (Selangor, Malaysia) for supporting the article processing charges and to the Department of Surgery and Department of Pathology, Hospital Canselor Tuanku Muhriz, for assisting in the specimen collection and histology verification.

Conflicts of Interest: The authors declare no conflict of interest.

References
1. Azizah, A.M.; Hashimah, B.; Nirmal, K.; Siti Zubaidah, A.R.; Puteri, N.A.; Nabihah, A.; Azlina, A.A. Malaysia National Cancer Registry Report (MNCR) 2012–2016. Natl. Cancer Registry NCI 2019, 3, 34–37.
2. Evrard, C.; Tachon, G.; Randrian, V.; Karayan-Tapon, L.; Tougeron, D. Microsatellite Instability: Diagnosis, Heterogeneity, Discordance, and Clinical Impact in Colorectal Cancer. Cancers 2019, 11, 1567. [CrossRef] [PubMed]
3. Hughes, L.A.E.; Melotte, V.; de Schrijver, J.; de Maat, M.; Smit, V.T.H.B.M.; Bovee, J.V.M.G.; French, P.J.; van den Brandt, P.A.; Schouten, L.J.; de Meyer, T.; et al. The CpG Island Methylator Phenotype: What’s in a Name? Cancer Res. 2013, 73, 5858–5868. [CrossRef] [PubMed]
4. Toyota, M.; Ahuja, N.; Herman, J.G.; Baylin, S.B.; Issa, J.-P.J. CpG Island Methylator Phenotype in Colorectal Cancer. Proc. Natl. Acad. Sci. USA 1999, 96, 8681–8686. [CrossRef] [PubMed]
5. Creighton, S.D.; Stefanelli, G.; Reda, A.; Zovkic, I.B. Epigenetic Mechanisms of Learning and Memory: Implications for Aging. Int. J. Mol. Sci. 2020, 21, 6918. [CrossRef] [PubMed]
6. Ab Mutalib, N.-S.; Baharuddin, R.; Jamal, R. Chapter 18—Epigenome-Wide Analysis of DNA Methylation in Colorectal Cancer. In Computational Epigenetics and Diseases; Wei, L.K., Ed.; Translational Epigenetics; Academic Press: Cambridge, MA, USA, 2019; Volume 9, pp. 289–310. ISBN 978-0-12-814513-5.
7. Struhl, K. Is DNA Methylation of Tumour Suppressor Genes Epigenetic? eLife 2014, 3, e02475. [CrossRef] [PubMed]
8. Vadakedath, S.; Kandi, V. DNA Methylation and Its Effect on Various Cancers: An Overview. J. Mol. Biomark. Diagn. 2016, s2, 2. [CrossRef]
9. Wang, L.-H.; Wu, C.-F.; Rajasekaran, N.; Shin, Y.K. Loss of Tumor Suppressor Gene Function in Human Cancer: An Overview. CPB 2018, 51, 2647–2693. [CrossRef]
10. Takeshima, H.; Yamada, H.; Ushijima, T. Chapter 5—Cancer Epigenetics: Aberrant DNA Methylation in Cancer Diagnosis and Treatment. In Oncogenomics; Dammacco, F., Silvestris, F., Eds.; Academic Press: Cambridge, MA, USA, 2019; pp. 65–76. ISBN 978-0-12-811785-9.
11. Amabile, G.; Di Ruscio, A.; Müller, F.; Welner, R.S.; Yang, H.; Ebralidze, A.K.; Zhang, H.; Levantini, E.; Qi, L.; Martinelli, G.; et al. Dissecting the Role of Aberrant DNA Methylation in Human Leukaemia. Nat. Commun. 2015, 6, 7091. [CrossRef]
12. Jelinek, J.; Gharibyan, V.; Estecio, M.R.H.; Kondo, K.; He, R.; Chung, W.; Lu, Y.; Zhang, N.; Liang, S.; Kantarjian, H.M.; et al. Aberrant DNA Methylation Is Associated with Disease Progression, Resistance to Imatinib and Shortened Survival in Chronic Myelogenous Leukemia. PLoS ONE 2011, 6, e22110. [CrossRef]
13. Freitas, M.; Ferreira, F.; Carvalho, S.; Silva, F.; Lopes, P.; Antunes, L.; Salta, S.; Diniz, F.; Santos, L.L.; Videira, J.F.; et al. A Novel DNA Methylation Panel Accurately Detects Colorectal Cancer Independently of Molecular Pathway. *J. Transl. Med.* 2018, 16, 45. [CrossRef]

14. Mojtabanezhad Shariatpanahi, A.; Yassi, M.; Nouraie, M.; Sahebkar, A.; Varshoea Tabrizi, F.; Kerachian, M.A. The Importance of Stool DNA Methylation in Colorectal Cancer Diagnosis: A Meta-Analysis. *PloS ONE* 2018, 13, e0200735. [CrossRef]

15. Kohonen-Corish, M.R.J.; Tseung, J.; Chan, C.; Currey, N.; Dent, O.F.; Clarke, S.; Bokey, L.; Chapuis, P.H. KRAS Mutations and CDKN2A Promoter Methylation Show an Interactive Adverse Effect on Survival and Predict Recurrence of Rectal Cancer. *Int. J. Cancer* 2014, 134, 2820–2828. [CrossRef] [PubMed]

16. Cleven, A.H.G.; Derks, S.; Drahit, M.X.G.; Smits, K.M.; Melotte, V.; Van Neste, L.; Tournier, B.; Jooste, V.; Chapusot, C.; Weijenberg, M.P.; et al. CHFR Promoter Methylation Indicates Poor Prognosis in Stage II Microsatellite Stable Colorectal Cancer. *Clin. Cancer Res.* 2014, 20, 3261–3271. [CrossRef]

17. Bosch, L.J.W.; Trooskens, G.; Snaebjornsson, P.; Coupé, V.M.H.; Mongera, S.; Haan, J.C.; Richman, S.D.; Koopman, M.; Tol, J.; de Meyer, T.; et al. Decoy Receptor 1 (DCRI) Promoter Hypermethylation and Response to Irinotecan in Metastatic Colorectal Cancer. *Oncotarget* 2017, 8, 63140–63154. [CrossRef]

18. Moutinho, C.; Martínez-Cardús, A.; Santos, C.; Navarro-Pérez, V.; Martínez-Balibrea, E.; Musulen, E.; Carmona, F.J.; Sartore-Bianchi, A.; Cassingena, A.; Siena, S.; et al. Epigenetic Inactivation of the BRCA1 Interactor SRBC and Resistance to Oxaiplatin in Colorectal Cancer. *J. Natl. Cancer Inst.* 2013, 106, djt322. [CrossRef]

19. Zhang, X.; Shimodaira, H.; Soeda, H.; Takahashi, H.; Ouchi, K.; Inoue, M.; Takahashi, M.; Takahashi, S.; Ishioka, C. CpG Island Methylation Phenotype Is Associated with the Efficiency of Sequential Oxaiplatin- and Irinotecan-Based Chemotherapy and EGFR-Related Gene Mutation in Japanese Patients with Metastatic Colorectal Cancer. *Int. J. Clin. Oncol.* 2016, 21, 1091–1101. [CrossRef] [PubMed]

20. Baharudin, R.; Ab Mutalib, N.-S.; Othman, S.N.; Sagap, I.; Rose, I.M.; Mohd Mokhtar, N. Identification of Predictive DNA Methylation Biomarkers for Chemotherapy Response in Colorectal Cancer. *Front. Pharmacol.* 2017, 8, 47. [CrossRef]

21. Sy, C.; Cc, K.; Cc, W.; Cw, H.; Jm, H.; Ch, H.; Yc, C.; Yl, S.; Yw, L. NKX6.1 Hypermethylation Predicts the Outcome of Stage II Colorectal Cancer Patients Undergoing Chemotherapy. *Genes Chromosom. Cancer* 2018, 57, 268–277. [CrossRef]

22. Ebert, M.P.A.; Tänzer, M.; Balluff, B.; Burgermeister, E.; Kretzschmar, A.K.; Hughes, D.J.; Tetzner, R.; Lotfon-Day, C.; Rosenberg, R.; Reinacher-Schick, A.C.; et al. TFA2P2-DKK4 and Chemoresistance in Colorectal Cancer. *N. Engl. J. Med.* 2012, 366, 44–53. [CrossRef] [PubMed]

23. Perez-Carbonell, L.; Balaguer, F.; Toiyama, Y.; Egoavil, C.; Rojas, E.; Guarinos, C.; Andreu, M.; Llor, X.; Castells, A.; Jover, R.; et al. IGFBP3 Methylation Is a Novel Diagnostic and Predictive Biomarker in Colorectal Cancer. *PLoS ONE* 2014, 9, e104285. [CrossRef]

24. Sun, X.; Yuan, W.; Hao, F.; Zhuang, W. Promoter Methylation of RASSF1A Indicates Prognosis for Patients with Stage II and III Colorectal Cancer Treated with Oxaiplatin-Based Chemotherapy. *Med. Sci. Monit.* 2017, 23, 5389–5395. [CrossRef]

25. Pelosof, L.; Yerram, S.; Armstrong, T.; Chu, N.; Danilova, L.; Yanagisawa, B.; Hidalgo, M.; Azad, N.; Herman, J.G. GPX3 Promoter Methylation Predicts Platinum Sensitivity in Colorectal Cancer. *Epigenetics* 2017, 12, 540–550. [CrossRef]

26. Shimizu, S.; Iida, S.; Ishiguro, M.; Uetake, H.; Ishikawa, T.; Takagi, Y.; Kobayashi, H.; Higuchi, T.; Enomoto, M.; Mogushi, K.; et al. Methylation of BNP3 Gene in Colorectal Cancer Progression. *Oncol. Lett.* 2010, 1, 685–687. [CrossRef]

27. Tian, Y.; Morris, T.J.; Webster, A.P.; Yang, Z.; Beck, S.; Feber, A.; Teschendorff, A.E. ChAMP: Updated Methylation Analysis Pipeline for Illumina BeadChips. *Bioinformatics* 2017, 33, 3982–3984. [CrossRef]

28. Fortin, J.-P.; Labbe, A.; Lemire, M.; Zanke, B.W.; Hudson, T.J.; Fertig, E.J.; Greenwood, C.M.; Hansen, K.D. Functional Normalization of 450k Methylation Array Data Improves Replication in Large Cancer Studies. *Genome Biol.* 2014, 15, 503. [CrossRef] [PubMed]

29. Johnson, W.E.; Li, C.; Rabinovic, A. Adjusting Batch Effects in Microarray Expression Data Using Empirical Bayes Methods. *Biostatistics* 2007, 8, 118–127. [CrossRef]

30. Ritchie, M.E.; Phipson, B.; Wu, D.; Hu, Y.; Law, C.W.; Shi, W.; Smyth, G.K. Limma Powers Differential Expression Analyses for RNA-Sequentialing and Microarray Studies. *Nucleic Acids Res.* 2015, 43, e47. [CrossRef] [PubMed]

31. Wettenhall, J.M.; Smyth, G.K. LimmaGUI: A Graphical User Interface for Linear Modeling of Microarray Data. *Bioinformatics* 2004, 20, 3705–3706. [CrossRef]

32. Heatmap of 50 Differentially Methylated Genes. Available online: https://software.broadinstitute.org/morpheus/ (accessed on 26 December 2021).

33. Huang, D.W.; Sherman, B.T.; Lempicki, R.A. Systematic and Integrative Analysis of Large Gene Lists Using DAVID Bioinformatics Resources. *Nat. Protoc.* 2009, 4, 44–57. [CrossRef] [PubMed]

34. Huang, D.W.; Sherman, B.T.; Lempicki, R.A. Bioinformatics Enrichment Tools: Paths toward the Comprehensive Functional Analysis of Large Gene Lists. *Nucleic Acids Res.* 2009, 37, 1–13. [CrossRef]

35. Benjamini, Y.; Hochberg, Y. Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. *J. R. Stat. Soc. Ser. B* 1995, 57, 289–300. [CrossRef]

36. Szklarczyk, D.; Morris, J.H.; Cook, H.; Kuhn, M.; Wyder, S.; Simonovic, M.; Santos, A.; Doncheva, N.T.; Roth, A.; Bork, P.; et al. The STRING Database in 2017: Quality-Controlled Protein-Protein Association Networks, Made Broadly Accessible. *Nucleic Acids Res.* 2017, 45, D362–D368. [CrossRef] [PubMed]
37. Diez-Villanueva, A.; Mallona, I.; Peinado, M.A. Wanderer, an Interactive Viewer to Explore DNA Methylation and Gene Expression Data in Human Cancer. *Epigenet. Chromatin* **2015**, *8*, 22. [CrossRef] [PubMed]

38. Siegel, R.L.; Miller, K.D.; Fedewa, S.A.; Ahnen, D.J.; Meester, R.G.S.; Barzi, A.; Jemal, A. Colorectal Cancer Statistics, 2017. *CA A Cancer J. Clin.* **2017**, *67*, 177–193. [CrossRef] [PubMed]

39. Cancer Genome Atlas Network Comprehensive Molecular Characterization of Human Colon and Rectal Cancer. *Nature* **2012**, *487*, 330–337. [CrossRef]

40. Naumov, V.A.; Generozov, E.V.; Zaharjevskaya, N.B.; Matushkina, D.S.; Larin, A.K.; Chernyshov, S.V. Genome-Scale Analysis of DNA Methylation in Colorectal Cancer Using Infinium HumanMethylation450 BeadChips. *Epigenetics* **2013**, *8*, 921–934. [CrossRef]

41. Song, M.-A.; Tiirikainen, M.; Kwee, S.; Okimoto, G.; Yu, H.; Wong, L.L. Elucidating the Landscape of Aberrant DNA Methylation in Hepatocellular Carcinoma. *PLoS ONE* **2013**, *8*, e55761. [CrossRef]

42. Gu, S.; Lin, S.; Ye, D.; Qian, S.; Jiang, D.; Zhang, X.; Li, Q.; Yang, J.; Ying, X.; Li, Z.; et al. Genome-Wide Methylation Profiling Reveals Novel Epigenetic Signatures in Squamous Cell Lung Cancer. *BMC Genom.* **2017**, *18*, 901. [CrossRef]

43. McInnes, T.; Zou, D.; Rao, D.S.; Munro, F.M.; Phillips, V.L.; McCall, J.; Black, M.A.; Reeve, A.E.; Guilford, P.J. Genome-Wide Methylation Analysis Identifies a Core Set of Hypermethylated Genes in CIMP-H Colorectal Cancer. *BMC Cancer* **2017**, *17*, 228. [CrossRef] [PubMed]

44. McInnes, T.; Zou, D.; Rao, D.S.; Munro, F.M.; Phillips, V.L.; McCall, J.; Black, M.A.; Reeve, A.E.; Guilford, P.J. Genome-Wide Methylation Profiling Reveals Cancer-Associated Changes within Early Colonic Neoplasia. *Oncogene* **2017**, *36*, 5035–5044. [CrossRef] [PubMed]

45. Fiedler, D.; Hirsch, D.; El Hajj, N.; Yang, H.H.; Hu, Y.; Sticht, C.; Nanda, I.; Belle, S.; Rueschoff, J.; Lee, M.P.; et al. Genome-Wide DNA Methylation Analysis of Colorectal Adenomas with and without Recurrence Reveals an Association between Cytosine-Phosphate-Guanine Methylation and Histological Subtypes. *Genes Chromosom. Cancer* **2019**, *58*, 783–797. [CrossRef] [PubMed]

46. Jin, P.; Kang, Q.; Wang, X.; Yang, L.; Yu, Y.; Li, N.; He, Y.; Han, X.; Hang, J.; Zhang, J.; et al. Performance of a Second-Generation Methylated SEPT9 Test in Detecting Colorectal Neoplasm. *J. Gastroenterol. Hepatol.* **2015**, *30*, 830–833. [CrossRef]

47. Töth, K.; Gałamb, O.; Spisák, S.; Wichmann, B.; Sipos, F.; Valcz, G.; Leiszter, K.; Molnár, B.; Tulassay, Z. The Influence of Methylated Septin 9 Gene on RNA and Protein Level in Colorectal Cancer. *Pathol. Oncol. Res.* **2011**, *17*, 503–509. [CrossRef] [PubMed]

48. Wasserkort, R.; Kalmárné, A.; Valcz, G.; Spisák, S.; Krispin, M.; Tóth, K.; Tulassay, Z.; Szedziewski, A.Z.; Molnár, B. aberrant Septin 9 DNA Methylation in Colorectal Cancer is Restricted to a Single CpG Island. *BMC Cancer* **2013**, *13*, 398. [CrossRef]

49. Tosetti, K.; Wasserkort, R.; Sipos, F.; Kalmár, A.; Wichmann, B.; Leiszter, K.; Valcz, G.; Juhász, M.; Miheller, P.; Patai, Á.V.; et al. Detection of Methylated Septin 9 in Tissue and Plasma of Colorectal Patients with Neoplasia and the Relationship to the Amount of Circulating Cell-Free DNA. *PLoS ONE* **2014**, *9*, e115415. [CrossRef] [PubMed]

50. Prachayakul, V; Rugivarodom, M.; Nopparoorsri, P.; Cheirsilpa, K.; Chang, A.; Kamolhan, T.; Boonyaaruntrate, T.; thuwajit, C.; Thuwajit, P. Diagnostic Power of DNA Methylation Markers Suggestive of Cholangiocarcinoma in ERCP-Based Brush Cytology. *Gastrointest. Endosc.* **2022**, *85*, 123–130.e1. [CrossRef]

51. National Cancer Institute (NCI). A Phase Ib/II Study of the Histone Methyltransferase Inhibitor Pinometostat in Combination with Azacitidine in Patients with 11q23-Rearranged Acute Myeloid Leukemia. 2021. Available online: [clinicaltrials.gov](http://clinicaltrials.gov) (accessed on 26 December 2021).

52. Ishak, M.; Baharudin, R.; Mohamed Rose, I.; Sagap, I.; Mazlan, L.; Mohd Azman, Z.A.; Abu, N.; Jamal, R.; Lee, L.-H.; Ab Mutalib, N.S. Genome-Wide Open Chromatin Methylation Profiles in Colorectal Cancer. *Biomolecules* **2020**, *10*, 719. [CrossRef] [PubMed]
61. Allawi, H.; Lidgard, G.P.; Giakoumopoulou, M.; Ahlquist, D.A.; Taylor, W.R.; Mahoney, D. Detection of Neoplasia by Analysis of Methylated DNA 2021. U.S. Patent No. 11,028,447, 23 November 2017.
62. Ahlquist, D.A.; Taylor, W.R.; Mahoney, D.; G.P.; Allawi, H.T. Detecting Colorectal Neoplasm 2021. U.S. Patent No. 11078539 B2, 3 August 2021.
63. Yu, C.; Chen, S.; Guo, Y.; Sun, C. Oncogenic TRIM31 Confers Gemicatibarine Resistance in Pancreatic Cancer via Activating the NF-KB Signaling Pathway. *Tumorsciences* 2018, 8, 3224–3236. [CrossRef]
64. Xiao, Y.; Deng, T.; Ming, X.; Ju, J. TRIM31 Promotes Acute Myeloid Leukemia Progression and Sensitivity to Daunorubicin through the Wnt/β-Catenin Signaling. *Biosci. Rep.* 2020, 40, BSR20194334. [CrossRef]
65. Guo, P.; Ma, X.; Zhao, W.; Huai, W.; Li, H.; Qiu, Y.; Zhang, Y.; Han, L. TRIM31 Is Upregulated in Hepatocellular Carcinoma and Promotes Disease Progression by Inducing Ubiquitination of TSC1–TSC2 Complex. *Oncoene* 2018, 37, 478–488. [CrossRef]
66. Guo, Y.; Li, Q.; Zhao, G.; Zhang, J.; Yuan, H.; Feng, T.; Ou, D.; Gu, R.; Li, S.; Li, K.; et al. Loss of TRIM31 Promotes Breast Cancer Progression through Regulating K48- and K63-Linked Ubiquitination of P53. *Cell Death Dis.* 2021, 12, 1–13. [CrossRef]
67. Wang, H.; Yao, L.; Gong, Y.; Zhang, B. TRIM31 Regulates Chronic Inflammation via NF-KB Signal Pathway to Promote Invasion and Metastasis in Colorectal Cancer. *Am. J. Transl. Res.* 2018, 10, 1247–1259.
68. Barrow, T.M.; Klett, H.; Toth, R.; Böhm, J.; Gigic, B.; Habermann, N.; Scherer, D.; Schrotz-King, P.; Skender, S.; Abbenhardt-Martin, C.; et al. Smoking Is Associated with Hypermethylation of the APC 1A Promoter in Colorectal Cancer: The ColoCare Study. *J. Pathol.* 2017, 243, 366–375. [CrossRef]
69. Farkas, S.A.; Vymetalkova, V.; Vodickova, L.; Vodicka, P.; Nilsson, T.K. DNA Methylation Changes in Genes Frequently Mutated in Sporadic Colorectal Cancer and in the DNA Repair and Wnt/β-Catenin Signaling Pathway Genes. *Epigenomics* 2014, 6, 179–191. [CrossRef]
70. Petersen, J.W.; Douglas, J.Y. Tenascin-X, Collagen, and Ehlers-Danlos Syndrome: Tenascin-X Gene Defects Can Protect against Adverse Cardiovascular Events. *Med. Hypotheses* 2013, 81, 443–447. [CrossRef] [PubMed]
71. Hui, L.; Zhang, J.; Ding, X.; Guo, X.; Jang, X. Identification of Potentially Critical Differentially Methylated Genes in Nasopharyngeal Carcinoma: A Comprehensive Analysis of Methylation Profiling and Gene Expression Profiling. *Oncoll. Lett.* 2017, 14, 7171–7178. [CrossRef]
72. Yuan, Y.; Nymoen, D.A.; Stavnes, H.T.; Rosnes, A.K.; Wu, C.; Nesland, J.M.; Davidson, B. Tenascin-X Is a Novel Diagnostic Marker of Malignant Mesothelioma. *Am. J. Surg. Pathol.* 2009, 33, 1673–1682. [CrossRef]
73. Fan, S.; Tang, J.; Li, N.; Zhao, Y.; Ai, R.; Zhang, K.; Wang, M.; Du, W.; Wang, W. Integrative Analysis with Expanded DNA Methylation Data Reveals Common Key Regulators and Pathways in Cancers. *NPJ Genom. Med.* 2019, 4, 1–11. [CrossRef] [PubMed]
74. Lim, W.-J.; Kim, K.H.; Kim, J.-Y.; Jeong, S.; Kim, N. Identification of DNA-Methylated CpG Islands Associated With Gene Silencing in the Adult Body Tissues of the Ogye Chicken Using RNA-Seq and Reduced Representation Bisulfite Sequencing. *Front. Genet.* 2019, 10, 346. [CrossRef]
75. Lim, D.H.K.; Maher, E.R. DNA Methylation: A Form of Epigenetic Control of Gene Expression. *Obstet. Gynaecol.* 2010, 12, 37–42. [CrossRef] [PubMed]
76. Nowacka-Zawisza, M.; Wiśniew, E. DNA Methylation and Histone Modifications as Epigenetic Regulation in Prostate Cancer (Review). *Oncol. Rep.* 2017, 38, 2587–2596. [CrossRef] [PubMed]
77. Heberlé, É.; Bardet, A.F. Sensitivity of Transcription Factors to DNA Methylation. *Essays Biochem.* 2019, 63, 727–741. [CrossRef] [PubMed]
78. Moarii, M.; Boeva, V.; Vert, J.-P.; Reyal, F. Changes in Correlation between Promoter Methylation and Gene Expression in Cancer. *BMC Genom.* 2015, 16, 1–14. [CrossRef]
79. Xu, W.; Xu, M.; Wang, L.; Zhou, W.; Xiang, R.; Shi, Y.; Zhang, Y.; Piao, Y. Integrative Analysis of DNA Methylation and Gene Expression Identified Cervical Cancer-Specific Diagnostic Biomarkers. *Signal Transduct. Target. Ther.* 2019, 4, 1–11. [CrossRef] [PubMed]
80. Bommarito, P.A.; Fry, R.C. Chapter 2–1—The Role of DNA Methylation in Gene Regulation. In *Toxicoepigenetics*; McCullough, S.D., Dolinoy, D.C., Eds.; Academic Press: Cambridge, MA, USA, 2019; pp. 127–151. ISBN 978-0-12-812433-8.
81. Spirin, V.; Mirny, L.A. Protein Complexes and Functional Modules in Molecular Networks. *Proc. Natl. Acad. Sci. USA* 2003, 100, 12123–12128. [CrossRef] [PubMed]
82. Kovác, I.A.; Luck, K.; Spirohn, K.; Wang, Y.; Pollis, C.; Schlabach, S.; Bian, W.; Kim, D.-K.; Kishore, N.; Hao, T.; et al. Network-Based Prediction of Protein Interactions. *Nat. Commun.* 2019, 10, 1240. [CrossRef] [PubMed]
83. Liao, W.-T.; Jiang, D.; Yuan, J.; Cui, Y.-M.; Shi, X.-W.; Chen, C.-M.; Bian, X.-W.; Deng, Y.-J.; Ding, Y.-Q. HOXB7 as a Prognostic Factor and Mediator of Colorectal Cancer Progression. *Clin. Cancer Res.* 2011, 17, 3569–3578. [CrossRef] [PubMed]
84. Tu, W.; Zhu, X.; Han, Y.; Wen, Y.; Qiu, G.; Zhou, C. Overexpression of HOXB7 Is Associated with a Poor Prognosis in Patients with Gastric Cancer. *Oncol. Lett.* 2015, 10, 2967–2973. [CrossRef] [PubMed]
85. Yang, S.; Lee, J.Y.; Hur, H.; Oh, J.H.; Kim, M.H. Up-Regulation of HOXB Cluster Genes Are Epigenetically Regulated in Tamoxifen-resistant MCF7 Breast Cancer Cells. *BMP Rep.* 2018, 51, 450–455. [CrossRef] [PubMed]
86. Liu, X.; Li, T.; Kong, D.; You, H.; Kong, F.; Tang, R. Prognostic Implications of Alcohol Dehydrogenases in Hepatocellular Carcinoma. *BMC Cancer* 2020, 20, 1204. [CrossRef]
87. Wu, M.; Li, Q.; Wang, H. Identification of Novel Biomarkers Associated With the Prognosis and Potential Pathogenesis of Breast Cancer via Integrated Bioinformatics Analysis. *Technol. Cancer Res. Treat.* 2020, 19, 1533033821992081. [CrossRef]
88. Sexton, R.E.; Hallak, M.N.A.; Uddin, M.d.H.; Diab, M.; Azmi, A.S. Gastric Cancer Heterogeneity and Clinical Outcomes. Technol. Cancer Res. Treat. 2020, 19, 19. [CrossRef]

89. Hu, Y.-H.; Ma, S.; Zhang, X.-N.; Zhang, Z.-Y.; Zhu, H.-F.; Ji, Y.-H.; Li, J.; Qian, X.-L.; Wang, Y.-X. Hypermethylation Of ADHFE1 Promotes The Proliferation Of Colorectal Cancer Cell Via Modulating Cell Cycle Progression. Onco Targets Ther. 2019, 12, 8105–8115. [CrossRef] [PubMed]

90. Tae, C.H.; Ryu, K.J.; Kim, S.-H.; Kim, H.C.; Chun, H.-K.; Min, B.-H.; Chang, D.K.; Rhee, P.-L.; Kim, J.J.; Rhee, J.C.; et al. Alcohol Dehydrogenase, Iron Containing, 1 promoter Hypermethylation Associated with Colorectal Cancer Differentiation. BMC Cancer 2013, 13, 142. [CrossRef]

91. Moon, J.W.; Lee, S.K.; Lee, Y.W.; Lee, J.O.; Kim, N.; Lee, H.J.; Seo, J.S.; Kim, J.; Kim, H.S.; Park, S.-H. Alcohol Induces Cell Proliferation via Hypermethylation of ADHFE1 in Colorectal Cancer Cells. BMC Cancer 2014, 14, 377. [CrossRef]

92. Fan, J.; Li, J.; Guo, S.; Tao, C.; Zhang, H.; Wang, W.; Zhang, Y.; Zhang, D.; Ding, S.; Zeng, C. Genome-Wide DNA Methylation Profiles of Low- and High-Grade Adenoma Reveals Potential Biomarkers for Early Detection of Colorectal Carcinoma. Clin. Epigenetics 2020, 12, 56. [CrossRef] [PubMed]

93. Wang, C.; Pu, W.; Zhao, D.; Zhou, Y.; Lu, T.; Chen, S.; He, Z.; Feng, X.; Wang, Y.; Li, C.; et al. Identification of Hyper-Methylated Tumor Suppressor Genes-Based Diagnostic Panel for Esophageal Squamous Cell Carcinoma (ESCC) in a Chinese Han Population. Front. Genet. 2018, 9, 356. [CrossRef]

94. Chen, Q.; Wu, Q.; Peng, Y. ADHFE1 Is a Correlative Factor of Patient Survival in Cancer. Open Life Sci. 2021, 16, 571–582. [CrossRef] [PubMed]