

Global Methylation Profiling of Lung Cancer Identifies Novel Methylated Genes¹

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Abstract

Epigenetic changes, including DNA methylation, are a common finding in cancer. In lung cancers methylation of cytosine residues may affect tumor initiation and progression in several ways, including the silencing of tumor suppressor genes through promoter methylation and by providing the targets for adduct formation of polycyclic aromatic hydrocarbons present in combustion products of cigarette smoke. Although the importance of aberrant DNA methylation is well established, the extent of DNA methylation in lung cancers has never been determined. Restriction landmark genomic scanning (RLGS) is a highly reproducible two-dimensional gel electrophoresis that allows the determination of the methylation status of up to 2000 promoter sequences in a single gel. We selected 1184 CpG islands for RLGS analysis and determined their methylation status in 16 primary non-small cell lung cancers. Some tumors did not show methylation whereas others showed up to 5.3% methylation in all CpG islands of the profile. Cloning of 21 methylated loci identified 11 genes and 6 ESTs. We demonstrate that methylation is part of the silencing process of *BMP3B* in primary tumors and lung cancer cell lines. *Neoplasia* (2001) 3, 314–323.

Keywords: non-small cell lung cancer, DNA methylation, RLGS, genome scanning, epigenetic.

Introduction

Lung cancer is the leading cause of cancer-related death in both males and females worldwide [1]. Clinically, lung cancer can be divided into two groups: small cell lung cancer (SCLC) and non-small cell lung cancer (NSCLC) [2,3]. The latter constitutes approximately 75% of all lung cancers [2] and includes squamous cell carcinoma (SCC), adenocarcinoma (AC) and large cell carcinoma (LCC) (approximately 30%, 40% and 15%, respectively in all lung cancer cases in North America) [3].

Molecular abnormalities in lung cancer affect both growth-promoting oncogenes and growth-inhibiting tumor

suppressor genes. So far mutations have been reported in the oncogene *K-RAS* [4–6] as well as in tumor suppressor genes *p53* [7], *CDKN2* [8–11] and *RB* [12]. In addition to genetic changes, the epigenetic change of DNA methylation, the addition of a methyl group to the cytosine ring in 5'-CpG-3' dinucleotides, may play a significant role during lung cancer development [13–17]. DNA methylation is established and maintained by a family of DNA methyltransferases [18] and affects chromatin organization as well as gene expression [19]. A well-studied example in lung cancer is the aberrant promoter methylation of the tumor suppressor gene, *CDKN2*, which correlates with gene silencing [8,10,20–22]. Because most of the reports describe methylation in single cancer genes, no measurement of the overall contribution of promoter methylation in lung cancer exists. As an initial step to address this question, Zochbauer-Muller et al. showed that numerous genes, including *retinoic acid receptor β -2* (*RAR β*), *tissue inhibitor of metalloproteinase 3* (*TIMP-3*), *CDKN2*, *O⁶-methylguanine-DNA-methyltransferase* (*MGMT*), *death-associated protein kinase* (*DAPK*), *E-cadherin* (*ECAD*), *p14^{ARF}* and *glutathione S-transferase P1* (*GSTP1*), were methylated at various degrees in a collection of 107 primary NSCLC [22].

Methylation changes in lung cancer appear to be early events and thus should be useful in improving early detection of potentially malignant cells [23–25]. For example, *CDKN2* promoter methylation is proposed as a biomarker for early detection of lung cancer and monitoring of prevention trials [23–25]. Using sensitive PCR-based methylation analysis, methylation in *CDKN2* and/or *MGMT*

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promoters was found in sputum of smokers up to 3 years before clinical diagnosis of squamous cell lung carcinoma [23–25].

There is a need for the identification of novel markers in lung cancer as well as the identification of cancer-related genes. Aberrantly methylated target sequences can guide a search for novel genes that may be useful biomarkers, as well as candidate cancer genes. In this study we use, for the first time, restriction landmark genomic scanning (RLGS) to determine frequencies of DNA methylation and to identify novel methylation targets in NSCLC samples. Transcription patterns for one methylated gene, *BMP3B*, were studied in greater detail in primary tumors as well as in lung cancer cell lines. We show that aberrant methylation of the CpG island of *BMP3B* downregulates gene transcription of this interesting gene product.

Materials and Methods

Primary Human NSCLC Samples and Cell Lines

The tumor samples were derived from patients here at The Ohio State University, James Cancer Hospital. Complete pathologic classification is available for all tumor samples studied. Tissues were collected through the Cooperative Human Tissue Network to maintain patients' confidentiality. Samples were subsequently stored in our lung tumor tissue bank. Sixteen frozen paired NSCLC tumors with normal adjacent tissue were selected for this study. We used four lung cancer cell lines, all obtained from ATCC. H23 was derived from an AC, H125 was derived from an adenosquamous carcinoma, H522 was derived from an AC, and H1115 was derived from an LCC that metastasized to the lymph node. All cell lines were cultured in RPMI-1640 medium (Gibco BRL, Rockville, MD) supplemented with 10% fetal bovine serum, 100 U/ml penicillin and 0.1 mg/ml streptomycin (Gibco BRL).

Two-Dimensional Separation by Restriction Landmark Genome Scanning (RLGS)

RLGS was performed as described previously [26,27]. In summary, high molecular weight DNA was digested with the methylation sensitive restriction enzyme *NotI* (Promega, Madison, WI), end-labeled by [α -³²P]dGTP and [α -³²P]dCTP (Amersham, Piscataway, NJ), and then digested using the restriction enzyme *EcoRV* (Promega). *NotI*–*EcoRV* DNA fragments were separated in a first dimension through a 0.8% agarose tube gel, followed by an in-gel digestion with a third restriction enzyme, *HinfI* (New England Biolabs, Beverly MA). Finally, the DNA was separated on a second dimension 5% polyacrylamide gel; the gel was dried and exposed to X-ray film for 5 to 10 days. RLGS profiles of primary tumors and normal adjacent lung tissue were superimposed to visually detect differences in the intensities and/or presence of the radiolabeled fragments.

RLGS Analysis

The fragments in the RLGS profiles have been named on our "Master RLGS profile" (see website E-mail: <http://pandora.med.ohio-state.edu/masterRLGS/>). The master RLGS profile, derived from normal peripheral blood lymphocyte DNA, is divided into 63 sections. Each RLGS fragment is given a unique identifier (e.g., 3C1) that relates to the position within the RLGS profile. Therefore, data sets from different patients can be compared to identify commonly changed fragments.

Cloning of RLGS Fragments

A human *NotI*–*EcoRV* plasmid library and library mixing gels were created previously to facilitate cloning of RLGS fragments [26,28]. These mixing gels allow the determination of an address for a library clone corresponding to the RLGS fragment by identifying enhancement in the plate, row, and column mixing gels [26]. Bacterial clones were cultured in LB medium with ampicillin to isolate plasmid DNA using Qiagen miniprep kit (Qiagen, Valencia, CA). The plasmid DNA was digested with *NotI* and *EcoRV* (Promega), end-labeled with [α -³²P]dGTP and [α -³²P]dCTP (Amersham). Labeled DNA, 5.2 and 10.4 pg per clone, were mixed with labeled peripheral blood lymphocytes (PBL) genomic DNA from a normal healthy donor, and subsequently separated in the two-dimensional RLGS mixing gel. Enhanced intensities of the RLGS fragment of interest in these mixing gels indicated that the *NotI*/*EcoRV* clone represents the RLGS fragment of interest.

Characterization of RLGS Fragments

The confirmed plasmid clones are sequenced with M13 forward and M13 reverse primers. DNA sequences from M13 forward primer were used to perform standard nucleotide–nucleotide BLAST searches, using nonredundant (nr) and high throughput genomic sequence (htgs) databases at release time March 1, 2001 (<http://www.ncbi.nlm.nih.gov/BLAST/>). When possible, 2-kb genomic sequences from both sides of the *NotI* site were used for subsequent BLAST searches for genes or ESTs. Chromosomal location of cloned DNA fragments were obtained either directly from the information given in Genebank (<http://www.ncbi.nlm.nih.gov/>), or by searching the OMIM database (<http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=OMIM>) using gene names, or by searching the BAC resource website (<http://www.ncbi.nlm.nih.gov/genome/cyto/hbrc.shtml>) that contains cytogenetic data of FISH-mapped and sequence-tagged BAC clones. The properties of CpG islands were determined using a web-based program <http://www.itba.mi.cnr.it/webgene/>, which is provided by the Institute of Advanced Biomedical Technologies (ITBA), Italy.

Southern Hybridization

Southern hybridization was performed as described previously [26]. Briefly, control samples, DNA from healthy donors, were digested by *EcoRV* alone and a *NotI*/*EcoRV*

combination. The control sample will show the expected fragment size in the *EcoRV* digestion and a smaller fragment in the double digestion. All tumor samples were digested by *NotI/EcoRV*. The probes were prepared by restriction enzyme digestion of the clone DNA, purification of the target fragments, and subsequent labeling by random priming using the Prime IT II kit (Stratagene, La Jolla, CA). Percentage of methylation was quantified by a phosphor-imager.

5-aza-2'-Deoxycytidine Treatment of Cell Lines

A total of 3×10^6 cells of each NSCLC cell line were seeded into T75 culture flasks and cultured with RPMI-1640 media overnight. 5-aza-2'-deoxycytidine media was freshly made each day from stock solution (10 mmol/l in 100% DMSO) (Sigma, St. Louis, MO) to a final concentration of 1 μ mol/l. Cells were cultured with 5-aza-2'-deoxycytidine medium for 24 hours and then washed with PBS twice and continued to culture with fresh medium without 5-aza-2'-deoxycytidine for 2 days (24-hour timepoint). Cells for the 72-hour timepoint were cultured in 5-aza-2'-deoxycytidine medium that was changed daily for 3 days, then washed with PBS twice and continued to culture in fresh medium without 5-aza-2'-deoxycytidine for one additional day.

Semiquantitative RT-PCR

Total RNA from cell lines with or without 5-aza-2'-deoxycytidine treatment, primary tumors and paired normal adjacent lung tissue was isolated using TRIzol (Gibco BRL) and purified by RNeasy Mini Kit (Qiagen). Three micrograms total RNA was reverse transcribed *in vitro* by random hexamer and oligo dT using SUPERSCRIP^T first-strand synthesis kit (Gibco BRL). cDNA was amplified by PCR. Primers for RT-PCR were designed from the published cDNA sequences. Forward and reverse primers are from different exons to avoid amplification from genomic DNA.

Primer sequences were as follows: BMP3B forward: 5'-GGTGGACTTCGACAGACATCG-3'; BMP3B reverse: 5'-GATGGTGGCATGGTTGGATG-3', product size: 130 bp. GPI forward: 5'-GACCCCCAGTTCCAGAAGCTG-3'; GPI reverse: 5'-GCATCACGTCCTCCGTCACC-3', product size: 178 bp. In all reactions the forward primer for each pairs was end labeled by [γ -³²P]ATP with T4 kinase (Gibco BRL).

The semiquantitative radioactive RT-PCR was performed with optimized conditions for both the target gene and an internal control, *glucose-6-phosphate isomerase (GPI)*, in a single reaction tube. Amplification was stopped in the exponential range for both genes. The exponential range was determined by phosphorimager quantification of the PCR product band intensities from different amplification cycles. Each PCR reaction was carried out in 50 μ l final volume containing 5 μ l of $10 \times$ PCR buffer, 1.5 mM MgCl₂, 10 pmol of each primer, 200 μ M of each dNTP and 2.5 U Platinum Taq DNA polymerase (Gibco). The reactions were initiated with 95°C for 10 minutes to activate DNA polymerase and then followed by amplification. *BMP3B* was

amplified at 96°C for 20 seconds, 63°C for 15 seconds, and 72°C for 15 seconds for eight cycles before addition of GPI primers and additional 22 cycles.

Statistical Methods

Tests were performed for heterogeneity in methylation across patients and for preferential methylation of certain CpG island fragments, described in detail in Ref. [30]. Briefly, the heterogeneity test is based on a comparison of the mean methylation frequency to its variance in a chi-square statistic. Preferential methylation is assessed using a standard goodness-of-fit test [29] assuming that all spots are lost at equal true frequency. Empirical null distributions for both of these statistics were obtained by performing appropriate 10,000 random permutations of the fragment/patient data. Such an approach accounts for multiple testing (i.e., multiple fragments were examined) and does not rely on asymptotic distribution assumptions.

Results

Levels of Methylation in CpG Islands of NSCLC

RLGS profiles from 16 matched pairs consisting of lung tumors from NSCLC patients and matched normal lung tissue were prepared using the enzyme combination *NotI-EcoRV-HinfI* (Figure 1A). Each tumor profile was compared against the matching normal lung RLGS profile. Previously, we had shown that the loss of an RLGS fragment in the tumor compared with the matching normal is indicative of DNA methylation in the *NotI* site [30,31]. The total number of methylation events out of 1184 RLGS fragments analyzed, as well as clinical data for the patients are shown in Table 1. The range of methylation in these samples is from 0% to 5.3%. Although most (12 of 16) of the tumor samples showed methylation levels below 1%, 4 of 16 patients show levels of CpG island methylation above 2%. Of these, patients 5, 11, 14, and 17 show methylation frequencies of 4.9%, 5.3%, 2.4%, and 5.0%, respectively. No obvious correlation of overall methylation frequency and clinical data can be seen. This range of variation is greater than would be expected if all patients had the same underlying methylation rate. A chi-square test (see Statistical Methods section) shows significant heterogeneity in methylation levels across the patients ($P < .0001$). In addition, some CpG islands are preferentially methylated, as indicated by the number of fragments showing relatively high methylation frequency (Figure 2, $P < .0001$). Of the total 1184 analyzed RLGS fragments, 1036 were never methylated. A total of 76 fragments were methylated in only one tumor, 53 fragments were methylated in two tumors, 9 fragments (2E61, 2F43, 3D24, 3F82, 3F85, 4D8, 4E53, 4F15, and 4F58) were methylated in three tumors, 2 fragments (2D45 and 3F16) were methylated in four tumors, 1 fragment (3F28) was methylated in five tumors, 1 fragment (3G78) was methylated in seven tumors, 5 fragments (2C35, 3C1, 3E55, 4E1, and 1 RLGS fragment not present in the RLGS master

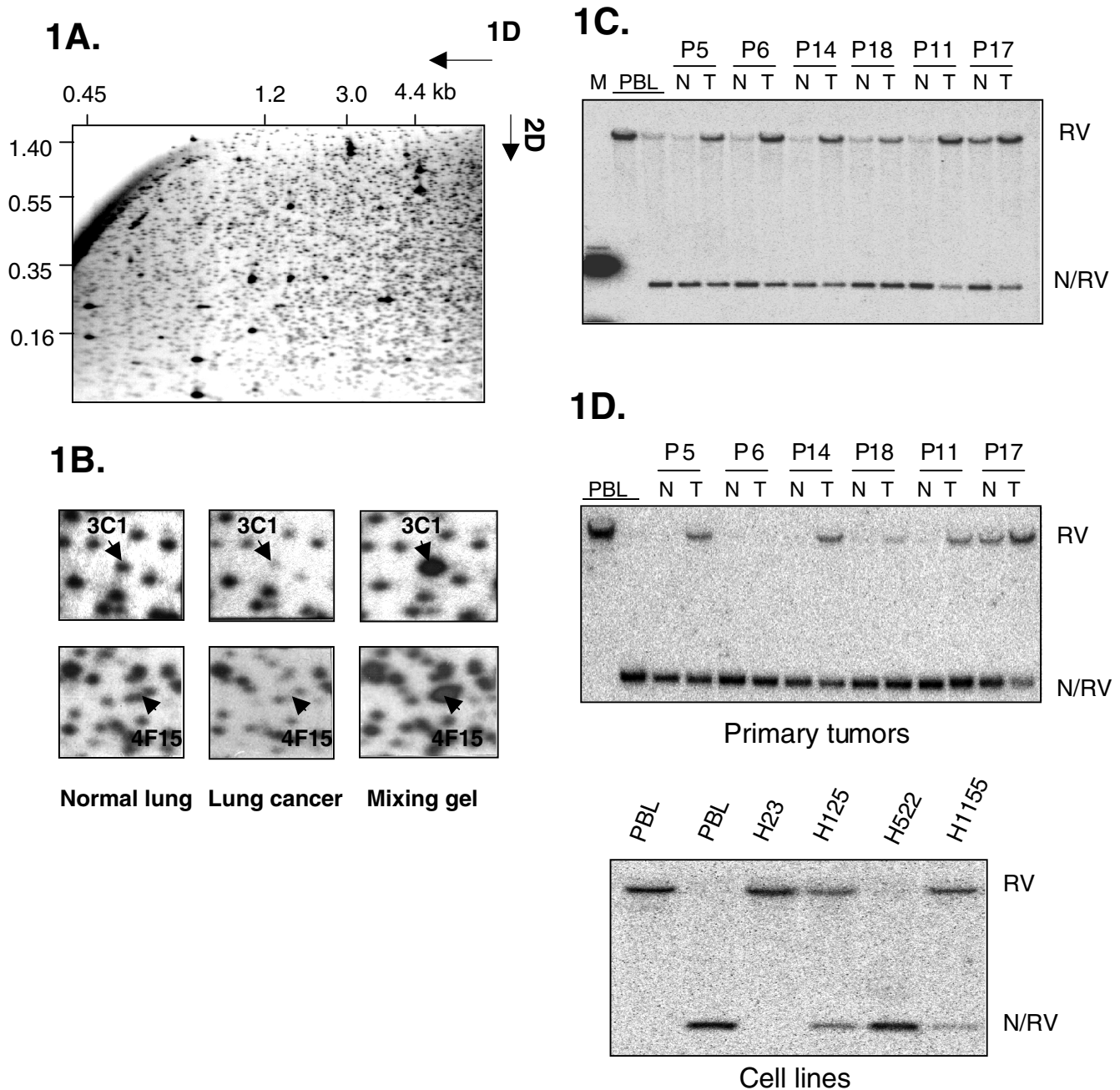


Figure 1. RLGS and confirmation of methylation by Southern hybridization. (A) Representative RLGS profile of normal lung DNA using the enzyme combination *NotI*–*EcoRV*–*HinI*. Fragment sizes for the first dimension (1D) and second dimension (2D) are given in kilobases. (B) Sections of RLGS profiles from normal lung and lung tumor from patient 17 highlighting RLGS fragment 3C1 (GNAL) and 4F15 (BMP3B) (arrows). While the lung cancer profile shows decreased intensity of 3C1 and 4F15, the mixing gel shows enhancement, indicating that the correct *NotI*–*EcoRV* clone was isolated from the library. Only differences in RLGS fragments 3C1 and 4F15 were indicated, other changes between normal and tumor profiles were not marked by arrows. (C) and (D) Southern blot analysis of primary lung cancers and matched normal adjacent tissue. Hybridization was used to confirm methylation of RLGS fragment 3C1, GNAL (C) and 4F15, BMP3B (D) in RLGS profiles of primary lung cancer. The peripheral blood lymphocyte (PBL) DNA in the first lane of each blot was digested with *EcoRV* only. All other tumor (T) and normal adjacent lung (N) DNA samples were double digested by *EcoRV/NotI*. The majority of the DNA is digested by *NotI* and results in the smaller *NotI*–*EcoRV* (N/RV) fragment. Tumor samples show a much higher degree of methylation. M: indicates the marker lane. H23, H125, H522, and H1155 are lung cancer cell lines.

profile) were methylated in eight tumors, and 1 fragment (1F22) was methylated in nine tumor samples (Figure 2).

No correlation between global methylation frequency and any of the clinical parameters including tumor stage, differentiation, histopathological classification, age or gender could be detected. However, it is interesting to note that a

subset of methylation events correlate with the histopathological features of the tumors. For example, of the 50 CpG island sequences that are methylated at least twice in tumors with clear histopathology (excluding patients No. 3, 10, and 15), 1 is found only in the two LCCs, no CpG islands was found methylated only in the four ACs, and 2 are found

Table 1. DNA Methylation in 16 Patients with Lung Cancer.

Patient No.	Methylation events out of 1184 CpG islands	% Methylation	Age	Gender	Tumor stage	Differentiation	Tumor type
1	5	0.4	45	F	T2N1, IIB	Well	AC
2	3	0.3	72	M	T1N0, IA	Moderate	SCC
3	7	0.6	56	F	T2N1, IIB	Poor	LCC with features of AC
5	58	4.9	75	M	T2N0, IB	Moderate	SCC
6	7	0.6	62	F	T1N0, IA	Poor	SCC
7	2	0.2	68	M	T3N2, IIIA	Undifferentiated	LCC
8	0	0	66	M	T2N0, IB	Well	AC
9	5	0.4	78	F	T1N0, IA	Poor	SCC
10	1	0.1	69	M	T2N1, IIB	Poor	N/A
11	63	5.3	67	F	T2N0, IB	Poor	LCC
13	0	0	76	M	T1N1, IIA	Poor	SCC
14	28	2.4	61	M	N/A	Poor	SCC
15	4	0.3	70	F	T3N0, IIB	N/A	LCC with SCC and AC
16	2	0.2	79	F	T1N0, IA	N/A	AC (with clear cell features)
17	59	5.0	81	M	N/A	Well	AC
18	8	0.7	63	F	N/A	Poor	SCC

SCC, squamous cell carcinoma; AC, adenocarcinoma; LCC, large cell carcinoma. N/A, no data available.

methyated exclusively in the seven SCCs. Other methylation events (a total of 16 CpG islands) are shared between all three subtypes. In addition seven methylation events are found in AC and LCC, nine are shared between AC and SCC and a total of 15 CpG islands are methyated in both SCC and LCC (Table 2). Validation of these methylation events in larger sample sets is required to unambiguously identify tumor-type-specific methylation events.

Cloning of RLGS Fragments and Properties of Cloned Fragments

To further characterize some of the methyated fragments in NSCLC, the *NotI*/*EcoRV* plasmid clone library mixing gels were used to locate the corresponding clones [26]. Clones with expected insert sizes were analyzed in RLGS mixing gels to confirm that the correct fragment was cloned. Figure 1B shows two examples for RLGS fragments 3C1 and 4F15.

A total of 21 fragments were cloned and sequenced. Twelve of these clones (2D14, 2D20, 2C35, 2E24, 2E61, 3B36, 3C1, 3E55, 3F16, 3F50, 3F82, and 4E53) have been identified as methylation targets in other types of malignancies [30–32]. BLAST searches identified homologies to 11 genes and six EST sequences (Table 3). Two sequences (3F16 and 3F82) show high homology to *DNA-binding protein A (DBPA)* and mouse *early B-cell factor 3*, respectively and may represent either pseudogenes or novel gene family members. The remaining two showed homology to genomic sequences. We found that 20 of 21 *NotI* sites are located within CpG islands. Six of the 11 CpG islands with homology to genes are located in the 5' end of the genes. The CpG islands identified in *insulin promoter factor 1 (IPF1)*, *orthodenticle (drosophila) homolog 1 (OTX1)*, *HOX11*, *T-box brain 1*, *monocarboxylate transporter 3 (MCT3)* are located in the middle or 3' end of the genes. Chromosomal location of all the fragments were derived from database searches of the human draft sequence of the human genome, Genbank or OMIM database. The detailed information for these 21 fragments, including fragment addresses on master profile, total methylation frequency in primary tumors, CpG island properties and location in the

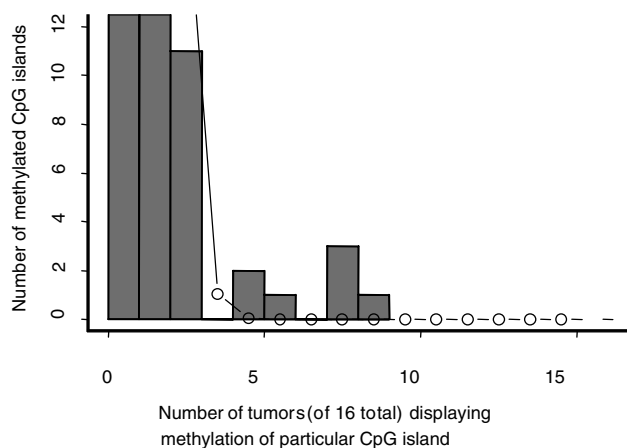


Figure 2. Global methylation patterns in lung cancer are nonrandom. Frequency distribution of methylation events in CpG islands within 16 NSCLCs.

Table 2. Distribution of CpG Islands Methyated at Least Twice in Various Histopathological Subgroups of Non-Small Cell Lung Cancer.

Subgroup	Number of methyated CpG islands
LCC	1
AC	0
SCC	2
LCC+AC	7
AC+SCC	9
LCC+SCC	15
LCC+AC+SCC	16

LCC, large cell carcinoma; AC, adenocarcinoma; SCC, squamous cell carcinoma.

**Table 3.** Cloned RLGS Fragments Altered in RLGS Profiles of 16 Primary Tumors.

RLGS master address	No. of tumors methylated	Methylation found in following subgroups	CpG island	Location of CpG island in gene	Gene or EST	Accession number	Chromosomal location
2C35	8	AC/LCC/SCC	Yes		EST BG142595	AL139281, NT_024073.1	10p12
3C1	8	AC/LCC/SCC	Yes	5' end	<i>GNAL</i>	U55180	18p11.21-pter
3E55	8	AC/LCC/SCC	Yes	3' end	<i>IPF1</i>	NM_000209	13q12.1
3G78	7	AC/LCC/SCC	Yes	5' end	<i>TAL1</i>	AL135960	1p32
2D20	4	AC/LCC/SCC	Yes	middle	<i>OTX1</i>	AB037501	2p13
3F16	4	AC/LCC/SCC	Yes		Homologous to <i>EBF</i>	AL354950, NT_024100.1	10q26
3F50	4	LCC/SCC	Yes	3' end	<i>HOX11</i>	AJ009794	10q24
2E61	3	SCC	Yes		EST AJ230817	AL354000	17p11.2
3B55	3	LCC/SCC	Yes	3' end	<i>T-box brain 1</i>	XM_002531	2q23-27
3F82	3	LCC/SCC	Yes		Homologous to <i>DBPA</i>	M24069	
4D8	3	AC/LCC/SCC	No		EST BF928282	AC012118, NT_010641.1	17q25.1
4E15	3	LCC/SCC	Yes		EST BE247619	AC003959	5q
4E53	3	AC/LCC/SCC	Yes			AL353195, NT_009829.1	13q12.2
4F15	3	LCC/SCC	Yes	5' end	<i>BMP3B</i>	D49493, NT_008757.1	10q11.21-11.23
5E25	3	AC/LCC/SCC	Yes		<i>Formin 2 like</i>	AL359918, NT_004771.1	1q44
2D14	2	AC/LCC	Yes	5' end	<i>CD8 β1 Chain</i>	S87068	2p12
3B36	2	AC/SCC	Yes	5' end	<i>CYP1b1</i>	XM_002576	2p21-22
3D44	2	AC/SCC	Yes			AL355304, NT_019429.1	6q23.1-6q24.3
5C32	2	AC/SCC	Yes	5' end	<i>CD34</i>	M81938	1q32
2E24	1	AC	Yes	middle	<i>MCT3</i>	AF132611	22q12.3-13.2
3E34	1	SCC	Yes		EST AI631157	AC000068	22q11.2

AC, adenocarcinoma; LCC, large cell carcinoma; SCC, squamous cell carcinoma.

genes, BLAST search results, and chromosomal location of the genes, are listed in Table 3.

RLGS Fragment Loss Is Due to DNA Methylation

Southern hybridization was performed as a more sensitive method to estimate the degree of methylation of the *NotI* site from RLGS fragments lost in tumors. Southern hybridization has the ability to detect methylation as low as 5% to 10%, whereas RLGS allows the detection of 30% methylation [31]. Southern blot hybridization was also performed to evaluate the possibility that the RLGS fragment loss was due to homozygous deletion. Corresponding *NotI/EcoRV* plasmid clones for four RLGS fragments were used as probes on Southern blots. These four clones included three fragments with homology to known genes 3C1 (*G-α-olfactory, GNAL*), 3E55 (*insulin promoter factor 1, IPF1*), and 4F15 (*Bone morphogenetic protein 3B, BMP3B*). In addition, fragment 2C35 with homology to an EST sequence was used. Representative Southern blots for RLGS fragment 3C1 (*GNAL*) and 4F15 (*BMP3B*) are shown in Figure 1C and D, respectively. Tumor samples and paired normal adjacent lung tissue DNAs were digested with *NotI/EcoRV*. The detection of a fragment in the *NotI/EcoRV* digests equal in size to the one detected with *EcoRV* only was scored as a methylation event in the tumor DNA. The hybridization shows that most tumor samples have a much higher degree of methylation of the *NotI* site than the normal counterparts. Some of the normal tissue DNAs show a small fraction of methylated sequences. These normal lung tissues were matched controls derived from the lung cancer patients and it is possible that they may contain contaminating preneoplastic cells. No homozygous deletions were detected

because the hybridization signals are present in all tumor samples. Methylation of the *NotI* site in the 5' end of *GNAL* was observed in 10 of 16 patients. Methylation of 4F15 (*BMP3B*), was detected in five of six tumors and thus confirmed the methylation events found by RLGS. Tumor-specific or increased methylation was observed in 11 of 16 primary NSCLC for fragment 2C35 and 11 of 16 for 3E55 (*IPF1*) (Figure 1C and D and data not shown).

Aberrant Transcription of *BMP3B* in Primary NSCLC and NSCLC Cell Lines

Radioactive, semiquantitative RT-PCR reactions were performed to determine the expression levels of *BMP3B* in six primary NSCLC samples and their paired normal lung tissue. *BMP3B* was found hypermethylated in a CpG island that is located in the 5' end of the gene (Figure 3A) and was selected for further analysis. RT-PCR was performed under optimized conditions for both the target gene and the internal control gene *glucose-6-phosphate isomerase (GPI)*. *BMP3B* was expressed in normal lung, whereas expression of *BMP3B* in all studied tumor samples was reduced (Figure 3B). Interestingly, tumors from patients Nos. 2 and 6, which did not show methylation of the *NotI* site exhibited very low levels of *BMP3B* expression, suggesting heterogeneous methylation of the CpG island. However, it is also possible, and maybe even more likely, that *BMP3B* is silenced by other mechanisms including mutations, deletions and/or LOH.

To investigate the effect of CpG island methylation on the transcription of the associated genes in more detail, we used three NSCLC cell lines that are methylated in the *NotI* site of *BMP3B* promoter region. H23, H125, and H1155 show more than 50% methylation in the *BMP3B* promoter. The cells

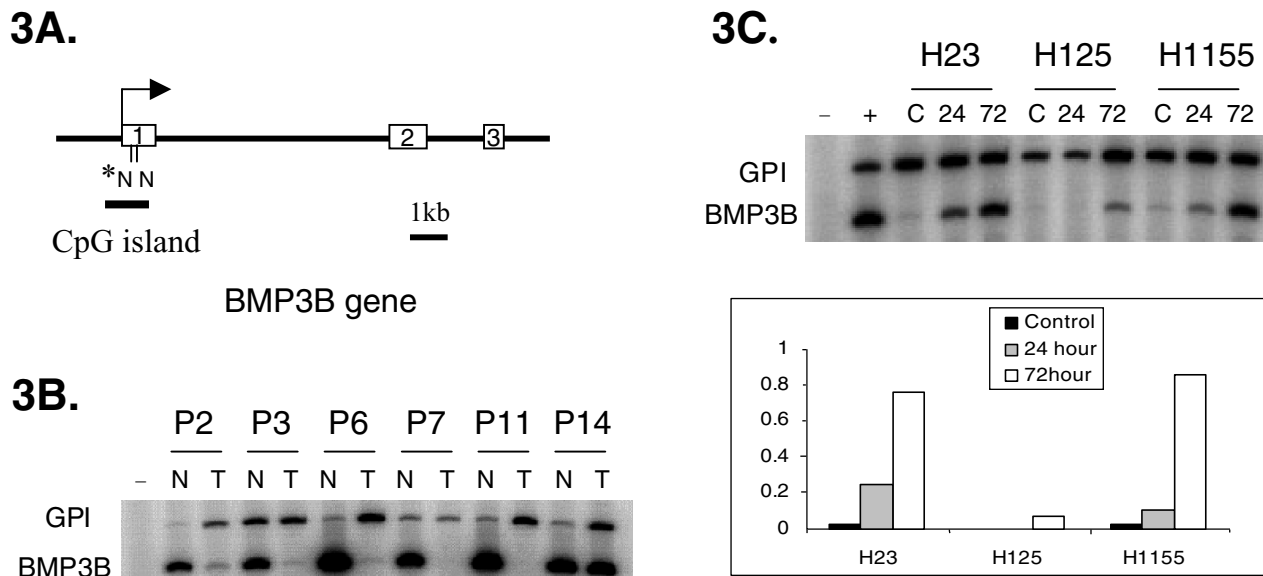


Figure 3. Abnormal transcription of *BMP3B* NSCLC samples. (A) Schematic representation of the genomic structure of *BMP3B*. The location of the *NotI* site relative to exon 1 and the extent of the CpG island are displayed. The star indicates the *NotI* site identified by RLGS. (B) Semiquantitative radioactive RT-PCR was used to determine the relative expression levels of *BMP3B* in six patients with lung cancer. Relative expression levels of RNAs obtained from tumor (T) and normal adjacent lung (N) were determined by comparing the intensities to the internal control, *GPI*. Negative control (-) contains water. *GPI*: Glucose-6-phosphate isomerase. (C) Radioactive semiquantitative RT-PCR was used to determine the relative expression of *BMP3B* levels in RNAs from three cell lines (H23, H125, and H1155) after treatment with 5'-aza-2'-deoxycytidine. The upper panel shows the RT-PCR results and the lower panel shows the relative expression levels of *BMP3B* compared to the internal control *GPI*. Cell lines were treated with 1 μ mol/15' -aza-2'-deoxycytidine for either 24 hours (24) or 72 hours (72). Untreated control cell lines (C) were harvested in exponential growth phase. Y-axis gives the relative ratio of band intensity for the target gene/*GPI* quantified by phosphorimager. Negative control (-) contained water, positive control (+) was normal lung tissue (NL).

were treated with two different timepoints (24 and 72 hours) of 5-aza-2'-deoxycytidine (Figure 3C). Similar to the primary tumors, all three cell lines (H23, H125, H1155) did not show any detectable level of *BMP3B* expression at baseline. However, expression was induced after 5-aza-2'-deoxycytidine treatment in all three cell lines (Figure 3C).

Discussion

The methylation scanning properties of RLGS have previously been used for the identification of imprinted genes in the mouse genome [33,34], as well as for the identification of methylated sequences in various human malignancies [30,31,35] but not lung cancer. Genome-wide scans for methylated sequences in lung cancer have been performed by two different strategies including the use of a methylated DNA binding column [36] and arbitrarily primed PCR [37]. However, these techniques either showed a bias for methylated repetitive sequences or were limited in the number of analyzed sequences. In contrast, RLGS, a highly reproducible two-dimensional gel electrophoresis, is a genome-wide scan of DNA methylation changes in CpG islands. Established cloning protocols utilizing an arrayed plasmid clone library facilitate the rapid identification of genomic sequences corresponding to the methylated targets [26,28]. RLGS is based on the digestion of genomic DNA with the methylation sensitive restriction enzyme *NotI*, which can only digest unmethylated genomic DNA and does not rely on prior knowledge of the gene sequence [26,31,35].

We have used RLGS to determine the contribution of CpG island hypermethylation in NSCLC and to identify novel methylation targets. Although the importance of DNA methylation in lung tumor development had been demonstrated in several reports [23,38-40], the overall extent was previously unknown. Under the assumption that all RLGS fragments represent promoter sequences we are, for the first time, able to demonstrate that up to 5.3% of all promoter regions, or 1537 of the estimated total 29,000 CpG islands [41,42] in the tumor genome could be methylated. The variability in the range of methylation shows that NSCLC represents a heterogeneous group not only with respect to the genetic defects identified but also on the epigenetic level.

Promoter methylation in cancer-related genes is well known in lung cancer and is correlated with gene silencing in genes involved in cell cycle (*CDKN2*) [8-11], apoptosis (*DAP*) [43], metastasis *H-cadherin* [44] and (*TIMP-3*) [45], differentiation (*RAR β*) [46], DNA repair (*MGMT*) [47], and the recently identified candidate tumor suppressor gene (*RASSF1A*) with homology to the RAS family [38]. In this study, we have identified 21 additional genomic loci including 11 genes and six ESTs with aberrant CpG island methylation in NSCLC. None of the cloned genes has been reported to be methylated in lung cancer previously. However, it is interesting to note that CpG island 3F16 is approximately 200 kb away from *MGMT* (located in sequence contig NT_024100.1). *MGMT* is a DNA repair gene that was previously found to be methylated in 21% of NSCLCs [22]. This finding would suggest a more regional

effect of methylation, similar to aberrant methylation found in chromosome 17p11.2 in the major breakpoint cluster region for medulloblastomas [30]. Another interesting target sequence is 4D8 located in chromosome 17q25.1. Chromosome 17q had previously been implicated with frequent (42%) loss of heterozygosity in NSCLC [48], suggesting that both genetic and epigenetic mechanisms could be involved in the silencing of a putative tumor suppressor gene in this region. Other fragments are derived from chromosome segments in 1p (3G78), 5q (4E15), 10q (3F50), 13q (3E55) 18p (3C1), and 22q (2E24), all within regions for which either LOH or homozygous deletions have been reported in lung cancer [49,50]. Whether any of the newly identified genes meet the expected criteria for tumor suppressor genes remains to be determined and will be the focus of future work.

We focused our studies on RLGS fragment 4F15, derived from a CpG island in the promoter region of the *BMP3B* gene and determined the relation of CpG island methylation to transcription. *BMP3B* is located in chromosome 10q11.21–11.23, a region that shows 20% to 30% LOH in NSCLCs and 51% LOH in SCLCs [51] (see online http://www.helsinki.fi/~lgl_www/LOSS/Respiratory.html) and thus located in a candidate tumor suppressor region. *BMP3B* is a member of the transforming growth factor β (TGF- β) superfamily, originally identified due to their osteoinductive capacity. Members of this family are usually involved in the regulation of cell growth/differentiation during development and were shown to be dysregulated in various human malignancies. Interestingly other members of the BMP family have been shown to induce apoptosis during organ development [52–54]. In addition, two BMP family members, *BMP4* and *BMP2*, have been shown to induce apoptosis in multiple myeloma cell lines [55] or hematopoietic cells [56], respectively. *BMP2* was also shown to suppress the transformed phenotype in the human lung carcinoma cell line A549 [57].

BMP3B is highly expressed in human adult lung, brain, skeletal muscle, pancreas, and testis, an expression pattern that distinguishes it from the closest family member *BMP3* [58]. *BMP3B* knockout mice did not show any detectable abnormalities, suggesting a redundant function with that of other members of the TGF- β family [59]. However, adult rat lung tissue does not express *BMP3B* suggesting the possibility of different functions of *BMP3B* in rodents and humans [60]. We have shown that methylation in *BMP3B* is correlated with transcriptional repression. The repression is reversible by treatment with the demethylating agent 5-aza-2'-deoxycytidine. Thus, our data suggest a causal relationship between methylation of the *BMP3B* promoter and transcriptional repression. We found *BMP3B* downregulated in all NSCLC patient samples and cell lines, even in those without methylation in the *NotI* site, suggesting that methylation patterns in the CpG island are heterogeneous. This assumption was confirmed by COBRA analysis testing the methylation status in four *Bst*UI (CGCG) restriction sites in the promoter (Dai et al. unpublished). Alternatively, other mechanisms (e.g., LOH or mutations) of gene silencing

could be present. Additional work to study the complete genetic and epigenetic mutation spectrum of *BMP3B* in lung tumorigenesis is underway.

The identification of multiple targets for methylation opens the exciting possibility to use these methylation events as biomarkers for the early detection of lung cancer in sputum as demonstrated by others [23]. In addition, our data also indicate the possibility that certain methylation events may be specific for lung cancer or subtypes within this group and thus could serve as potential markers for the molecular classification of lung cancers and different disease stages. Validation of methylation events as possible markers for early diagnosis, as predictive markers for survival or markers that classify subtypes will require larger sets of patient samples.

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