Significant frequency of allelic imbalance in 3p region covering RARβ and MLH1 loci seems to be essential in molecular non-small cell lung cancer diagnosis

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Abstract The aim of the study was to investigate the influence of allelic imbalance (AI) in several loci of tumor suppressor genes in 3p region on the non-small cell lung cancer (NSCLC) development. We evaluated the frequency of loss of heterozygosity and/or microsatellite imbalance (LOH/MSI) and assessed their association with patients’ characteristics (age, gender, tobacco addiction) and NSCLC classification according to TNM/AJCC staging. To analyze the potential role of AI involved in NSCLC pathogenesis, we allelotyped a group of 74 NSCLC patients using 7 microsatellite markers. The highest frequency of LOH/MSI, however, not statistically significant, was observed in RARβ and MLH1 (p = 0.104 and p = 0.216, respectively) loci. The association between high LOH/MSI frequency in 3p region with male gender (p = 0.041) as well as with age (especially >60 years) for RARβ and MLH1 genes (p = 0.0001 and p = 0.020, respectively) was documented. Statistically significant increased frequency of MLH1 allelic loss in squamous cell carcinoma (SCC) versus non-squamous cell carcinoma (non-SCC) was observed (p = 0.01). Significant increase in LOH/MSI frequency in 3p region (mainly in FHIT and MLH1 loci) in correlation with cigarette addiction in a lifetime (≥40 years and ≥40 Pack Years) was also documented (p < 0.05). The highest LOH/MSI was revealed in RARβ locus in IA tumors (p = 0.0001), while the similarly high allelic loss of MLH1 correlated with III A/B tumors (p = 0.0002), according to AJCC staging. The obtained results demonstrate that AI is influenced by tobacco smoking and seems to be vital in the molecular diagnosis of NSCLC, especially of SCC subtype.

Keywords Non-small cell lung carcinoma (NSCLC) · Loss of heterozygosity (LOH) · Microsatellite instability (MSI) · Microsatellite markers · Molecular diagnosis

Introduction

Lung cancer is one of the leading cause of cancer mortality in most developed countries, especially in men between the ages of 50 and 70 [1, 2]. Based on the histological verification and tumor biology, lung cancer is classified into two major groups: (1) small cell lung carcinomas (SCLC), accounting for about 20 % and (2) non-small cell lung carcinoma (NSCLC), constituting approximately 75 % of all primary lung cancers. NSCLC is further divided into squamous cell carcinoma (SCC) and non-squamous cell
carcinoma (non-SCC) with distinctive subtypes: adenocarcinoma (AC) and large cell carcinoma (LCC) [3]. This group of lung tumors is biologically heterogeneous, and the molecular studies of NSCLC, including genomic microarray and allelic imbalance analysis, are promising to extend and improve standard pathological methods of lung tumor assessment [4–6].

It has been documented that tobacco smoking is the most important risk factor in lung cancer development. Epidemiological studies in European populations have indicated that the increased incidence of lung cancer is proportional to the amount of smoked cigarettes and may have significant effect on tobacco-related cancer (TRC) risk [7–9]. However, it is well known that lung cancer (mainly NSCLC) is a genetically complex disease, developing in a result of the accumulation of multiple genetic abnormalities. Numerous genetic/epigenetic factors, especially gene polymorphisms, copy number alteration or gene methylation profiles, have been extensively investigated in lung cancer [7, 10–14]. The aggressive cancer phenotypes and their impact on patient outcome are actually characterized by numerous of these molecular changes [15–18]. Microsatellite instability (MSI) and more frequently occurring loss of heterozygosity (LOH) have been identified as the initial event in lung carcinogenesis and recognized in multiple chromosomal regions: 1p, 2p, 2q, 3p, 4q, 5q, 6p, 6q, 7p, 7q, 8p, 9q, 10q, 11p, 13p, 13q, 17p, 18q, 19q, 21q and 22q [16, 19–21]. Among them, the microsatellite alterations in 3p are regarded as biomarkers in genetic classification of pathological stages of NSCLC, as well as markers having prognostic significance [17, 18, 21]. However, despite many molecular studies on loss of heterozygosity and/or microsatellite imbalance (LOH/MSI) in 3p, the important tumor suppressor genes (TSGs) accepted as candidate genes associated with lung tumorigenesis have not been unequivocally identified yet. It is important to focus on this issue as inactivation of TSGs is a vital genetic event during the initiation as well as progression of lung carcinogenesis. The aim of our study was to confirm whether LOH/MSI alterations in the selected gene loci in 3p (FHIT, RASSF1A, MLH1, RARβ, VHL) might have important diagnostic and/or prognostic value in NSCLC patients.

Materials and methods

Biological material

The procedures used in the study were approved by the Ethical Committee of the Medical University of Lodz (RNN/140/10/KE). Informed written consent was received from each patient.

Lung tissue samples (100–150 mg) were obtained from patients (47 men and 27 women) who had undergone pulmonectomy or lobectomy at the Department of Thoracic Surgery, General and Oncologic Surgery, Medical University of Lodz, Poland, between July 2010 and June 2012. The studied biological material included 74 non-small cell lung carcinoma specimens and 74 matching macroscopically unchanged lung tissue samples received from the most distant site from the resected center of the primary lesion. Immediately after resection, tissue samples were collected in RNAlater® and frozen at −80 °C. The resected NSCLC specimens were postoperatively histopathologically evaluated and classified according to the AJCC staging as well as TNM classification (pTNM), according to the WHO Histological Typing of Lung Tumor and IASCLC Staging Project 7th ed. (2010) Cancer [22].

Characterization of the patients

The study involved 74 patients with diagnosed non-small cell lung carcinoma. The clinical characteristics of the studied patients, including their smoking habits, and histopathological verification of NSCLC samples are shown in Table 1. The smoking history was available for 71 patients: 5 patients were non-smokers, and 66 were smokers or former smokers. The amount of cigarettes smoked was presented as Pack Years (PYs) and was calculated according to the NCI Dictionary of Cancer Terms (1 Pack Year is equal to 20 cigarettes smoked per day for 1 year) [23].

DNA extraction

Isolation of genomic DNA from NSCLC samples and matching macroscopically unchanged lung tissues (reference DNA) was performed using QIAamp DNA Mini Kit (Qiagen, Germany), according to the manufacturer’s protocol. The quality and quantity of isolated DNA was spectrophotometrically assessed (Eppendorf BioPhotometr™ Plus, Eppendorf, Germany). DNA with a 260/280 nm ratio in range 1.8–2.0 was considered to be of high quality and used in further analysis.

Microsatellite analysis

The markers used for microsatellite analysis were selected from NCBI database (http://www.ncbi.nlm.nih.gov/genome/sts/sts) with supplementary mapping information, if necessary, provided in Cooperative Human Linkage Centre Database (http://www.chlc.org) or Genome Database (http://www.gdb.org). The chosen 7 microsatellite markers contained polymorphic microsatellite repeats, that is, (TG)n, (CA)n and (CAAA)n, and were linked to the chromosomal regions (3p14.2, 3p21.3, 3p22.2, 3p24.2 and 3p25.3) covering the loci...
of genes involved in significant processes of carcinogenesis, especially cell cycle regulation, proliferation and adhesion.

Pairs of DNA samples, that is, one sample obtained from the primary lesion and the other from unchanged lung tissue within the operational margin from the same patient, were amplified using primers for the studied microsatellite markers and AmpliTaq Gold®/C210 360 DNA Polymerase Kit (Applied Biosystems, USA). Reaction mixtures (total volume of 12.5 l) contained 30–40 ng DNA, 10× AmpliTaq Gold® 360 buffer (150 mM Tris–HCl, pH 8.3, 500 mM KCl), 360 GC Enhancer, 5 U/µl AmpliTaq Gold® 360 DNA Polymerase, 25 mM MgCl₂, 10 mM dNTPs, forward and reverse primers 0.5 µM each and nuclease-free water. All forward primers were labeled at 3’-end with fluorescent dye: 6-FAM, NED, PET or VIC. The temperatures of annealing were experimentally set for each pair of primers and were as follows: 45–47 °C (for D3S1317, D3S3611, D3S3615), 51–55 °C (for D3S1300, D3S1234) and 57–58 °C (for D3S1611, D3S1583). Negative and contamination controls were used for each marker. The chromosomal localization (region/gene) of the microsatellite markers and nucleotide sequences of primers used in the study are shown in Table 2.

The quality of PCR products were analyzed in 2 % agarose gel electrophoresis, after bromide ethidium staining. In order to perform the capillary electrophoresis, 0.5 µl of PCR product was mixed with 0.25 µl GS500-LIZ Size Standard and Hi-Di™ Formamide (both reagents: Applied Biosystems, USA) up to the final volume of 10 µl. The obtained mixture was denatured for 5 min at 95 °C and subsequently cooled on ice for 3 min. The separation in capillary electrophoresis was conducted in 3130xl Genetic Analyzer (Applied Biosystems, Hitachi, USA), and the allele presence was assessed using GeneMapper Software v 4.0, according to the manufacturer’s protocol. The informativeness of the studied samples (heterozygosity) was confirmed when two distinct alleles were detected in the reference sample (DNA from unchanged lung tissue from the same patient). Evaluation of LOH/MSI was performed by calculating the ratio of the fluorescence intensity of the alleles from unchanged lung tissue sample (N, normal, that is, control sample) to the fluorescence intensity of the alleles from NSCLC sample (T, tumor). For each informative tumor–normal DNA pair (paired T and N samples), an allelic imbalance ratio (AIR) was calculated, based on the maximum allele peak heights (fluorescence intensity), as follows: normal allele 1:normal allele 2/tumor allele 1:tumor allele 2 (N1:N2/T1:T2), according to the previously published protocol [24]. LOH in tumor samples was considered indicative when AIR value was less than 0.67 or greater than 1.35 (according to the criteria of GeneMapper Software v 4.0). MSI in tumor DNA was considered indicative if one or more additional alleles were present in tumor DNA sample, as compared with the control DNA sample.

LOH/MSI frequency was calculated as a percentage of LOH/MSI alteration in relation to all informative loci (heterozygous DNA) and according to all analyzed tumors’ and patients’ variables.

Table 1

| Clinical and pathological features | n (%) |
|-----------------------------------|-------|
| Mean age (total), 65 ± 8.433 (range 47–87) | 74 |
| Men, 65 ± 8.129 (range 47–87) | 47 (63.5) |
| Women, 63 ± 8.823 (range 42–79) | 27 (36.5) |
| Age groups | |
| <60 | 20 (27) |
| 60–70 | 32 (43) |
| >70 | 22 (30) |
| Smokers, n total = 71 | 66 (93) |
| Smoking period | |
| <40 years | 37 (52) |
| ≥40 years | 29 (41) |
| Amount of cigarettes smoked | |
| 10–15 cigarettes per day | 6 (8) |
| 20 cigarettes per day (1 pack) | 43 (61) |
| 30–40 cigarettes per day (1.5–2 packs) | 17 (24) |
| Pack years (PYs) | |
| <40 PYs | 30 (42) |
| ≥40 PYs | 36 (51) |
| Histopathological type of NSCLC | |
| Squamous cell carcinoma (SCC) | 40 (54) |
| Non-squamous cell carcinoma (non-SCC) | 34 (46) |
| Adenocarcinoma (AC) | 27 (37) |
| Large cell carcinoma (LCC) | 7 (9) |
| pTNM | |
| T1 | 20 (27) |
| T2 | 34 (46) |
| T3–4 | 20 (27) |
| AJCC | |
| AJCC IA | 14 (19) |
| AJCC IB | 14 (19) |
| AJCC II A | 14 (19) |
| AJCC II B | 9 (12) |
| AJCC IIIA/IIIB | 23 (31) |

Statistical analysis

Chi-square test ($\chi^2$) was used to assess the association between total LOH/MSI frequency (%) and chromosomal regions or markers. Nonparametrical statistical tests (Kruskal–Wallis or U Mann–Whitney test) were used to
assess the association between total LOH/MSI frequency in 3p region and clinical variables of patients (age at diagnosis, gender, smoking habits), characteristics of the tumor (NSCLC histological subtype, tumor size according to TNM classification, tumor preoperative and postoperative staging according to AJCC classification). Statistical significance was determined at the level of \( p < 0.05 \). The results are presented as mean or median ± SEM and ±SD values. For calculations, Statistica for Windows v. 10 was applied.

**Results**

Allelic imbalance and microsatellite instability

Pairs of DNA specimens obtained from 74 NSCLC patients (cancerous and macroscopically unchanged tissue samples from each patient) were available for LOH/MSI analysis using a panel of 7 microsatellite markers. All studied DNA samples from NSCLC specimens were informative for at least two studied loci. The informativeness of the markers was assessed to be in the range of 42.66–90.66 % (mean 69.90 % ± 17.19). LOH/MSI changes were observed for all (7/7, 100 %) microsatellite markers.

The obtained results indicate that in all studied samples, the frequency of LOH/MSI was in the range of 24.32–41.93 % (mean 33.85 % ± 6.17), depending on the marker. Representative examples of LOH/MSI in DNA derived from NSCLC samples are shown in Fig. 1.

**Total LOH/MSI frequency in particular loci**

Total LOH/MSI frequency (%) for each individual marker in NSCLC samples was assessed. The highest total frequency values were observed for D3S1583 marker (41.93 %; 13/31 informative loci) in 3p24.2 chromosomal region, that is, in \( RASSF1A \) locus, and for D3S1611 marker (40.38 %; 21/52 informative loci) in 3p22.2, that is, in \( MLH1 \) locus. The lowest LOH/MSI incidence (24.32 %; 9/37 informative loci) was identified for D3S3615 marker, localized in 3p21.3 \( (RASSF1A \) locus). Genetic instabilities of LOH/MSI type were also observed in \( FHIT \) locus for D3S1234 and D3S1300 markers, as well as in \( VHL \) locus for D3S1317 and D3S3611 markers, with similar frequencies: 36.36 % (20/55 informative loci), 32.76 % (19/58 informative loci), 31.37 % (19/60 informative loci) and 29.85 % (20/67 informative loci), respectively (see Fig. 2a).

Statistical analysis of LOH/MSI in particular gene loci revealed that the observed increase in LOH/MSI frequencies for D3S1583 \( (RASSF1A \) and D3S1611 \( (MLH1 \) were not statistically significant \( (p = 0.104, \chi^2 = 2.65 \text{ and } p = 0.216, \chi^2 = 1.53, \text{ respectively}) \). The lowest frequency of total LOH/MSI observed for D3S3615 \( (RASSF1A \) locus) was also not statistically significant \( (p = 0.052; \chi^2 = 3.76). \)

Correlation of total LOH/MSI frequency in 3p region with clinicopathological parameters

Total LOH/MSI frequency in 3p region was analyzed for all 7 markers in relation to histopathological characteristics of tumors (according to TNM and AJCC classifications and NSCLC subtypes) as well as clinical features of patients: gender, age at time of diagnosis.

Statistically significant differences in LOH/MSI frequency between all studied histopathological NSCLC subtypes (AC, LCC, SCC) were documented \( (p = 0.0006, \text{ ANOVA Kruskal–Wallis test}) \). Taking into account a small number of LCC tumors \( (n = 7) \), we compared SSC versus non-SCC. The increased frequency of LOH/MSI in SCC group for all studied 3p loci was found (see Fig. 2b).

Total LOH/MSI occurrence in 3p region was significantly more often in women as compared to men \( (p = 0.041; \text{ U Mann–Whitney test}) \). Regarding patient’s age at time of diagnosis, they were divided into the following age groups: (1) up to 60 years, (2) 60–70 years, (3) over 70 years and significant differences were found between them \( (p = 0.006, \text{ ANOVA Kruskal–Wallis test}) \). Statistically significant increase in LOH/MSI frequency was revealed in patients under the age of 60 years as compared with patients aged 60–70 years \( (p = 0.002, \text{ U Mann–Whitney test}) \) (see Fig. 3a).

Statistical analysis confirmed the significantly higher LOH/MSI frequency in SCC group as compared with non-SCC \( (p = 0.007, \text{ U Mann–Whitney test}) \) (see Fig. 3b).

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**Table 2** The chromosomal localization (region/gene) of the microsatellite markers, marker ID and nucleotide sequences of primers used in the study

| Chromosomal region (gene) | Marker ID | Nucleotide sequence of primers |
|---------------------------|-----------|--------------------------------|
| 3p14.2 \( (FHIT) \)       | D3S1234   | \( F \) CCTTGGAGGACAAGGCAAGAC—F |
|                           |           | \( R \) GaCCATAGGCACGGCTA—R     |
|                           | D3S1300   | \( F \) AGCTCACATCTGTCAGCT—F    |
|                           |           | \( R \) GCAATTCCTCCAGATG—R       |
| 3p21.3 \( (RASSF1A) \)    | D3S3615   | \( F \) TGGAAGGTAAGCAAGAC—N     |
|                           |           | \( R \) TCTCAGGGAAGAC—N          |
| 3p22.2 \( (MLH1) \)       | D3S1611   | \( F \) CCCCAGGCTGCAC—V         |
|                           |           | \( R \) AGCTGAGACTACAGGC—V       |
| 3p24.2 \( (RAR\beta) \)   | D3S1583   | \( F \) AGCTTTGATTATAGGTCTACA—N |
|                           |           | \( R \) TGGTCTTAATAGGCAC—R       |
| 3p25.3 \( (VHL) \)        | D3S1317   | \( F \) TACAAGTTCAGTTGGAACC—F   |
|                           |           | \( R \) CCTCCAGGCCATACAGC—R      |
|                           | D3S3611   | \( F \) GCATCTCTGCTGGAGCAT—V    |
|                           |           | \( R \) TCAAGAAGCTGTTGGG—R       |

\( F \), 6-FAM; \( P \), PET; \( N \), NED; \( V \), VIC

\( ^a \) \( F \) forward (sense), \( R \) reverse (antisense)

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Fig. 1 LOH/MSI analysis in NSCLC specimens (3130xl Genetic Analyzer, GeneMapper Software v. 4.0; Applied Biosystems, Hitachi). 1a T—homozygous DNA from tumor sample (sample no. 35, D3S1583 marker), 1b N—homozygous DNA from macroscopically unchanged lung tissue from the same patient, 2a T—heterozygous DNA from tumor sample (sample no. 12, D3S1611 marker), 2b N—heterozygous DNA from macroscopically unchanged lung tissue from the same patient, 3a T—LOH in DNA from tumor sample (sample no. 5, D3S1234 marker), 3b N—heterozygous DNA from macroscopically unchanged lung tissue from the same patient, 4a T—MSI in DNA from tumor sample (sample no. 72, D3S1300 marker), 4b N—heterozygous DNA from macroscopically unchanged lung tissue from the same patient.

Fig. 2 LOH/MSI frequencies (%) in NSCLC for all 7 studied microsatellite markers in a NSCLC samples; b in individual NSCLC histopathological subtypes (SCC vs. non-SCC)

(a)  (b)
No association was found between the total LOH/MSI frequency in 3p region and tumor clinical staging: pTNM classification (pT1, pT2, pT3–pT4), as well as AJCC classification (IA–B, IIA–B, IIIA–IIIB) \( (p > 0.05; \text{Kruskal–Wallis test}). \)

Correlation of LOH/MSI frequency in particular loci with clinicopathological parameters

Loss of heterozygosity and/or microsatellite imbalance (LOH/MSI) frequency (%) was analyzed separately for each locus in relation to clinical features of patients: gender, patient’s age at time of diagnosis as well as histopathological characteristics of tumors (according to TNM and AJCC classifications and NSCLC subtypes).

The comparison of LOH/MSI frequencies between the studied microsatellites loci in men confirmed statistically significant low LOH/MSI frequency in 3p21.3 locus (D3S3615 marker; RASSF1A) \( (p = 0.0017, \chi^2 = 9.88) \) as compared with other markers.

Regarding the above mentioned age groups, statistically significant low LOH/MSI for D3S3615 marker (RASSF1A) in older patients, that is, at the age of 60–70 years \( (p = 0.0003, \chi^2 = 13.23) \) and >70 years \( (p = 0.0001, \chi^2 = 29.98) \) were revealed. In the same age groups, the highest LOH/MSI incidence was observed for D3S1234 marker (FHIT) and D3S1583 (RARβ) markers. Statistically significant high LOH/MSI frequency for D3S1611 marker (MLH1) in patients aged 60–70 years \( (p = 0.0198, \chi^2 = 5.43) \) and for D3S1583 marker (RARβ) in patients aged >70 years \( (p = 0.0001, \chi^2 = 24.04) \) were found.

According to TNM staging, the studied tumor samples were divided into three groups (pT1, pT2, pT3–T4). The highest frequency of LOH/MSI in the pT1 group was observed for D3S1234 marker (FHIT), and it was statistically significant \( (p = 0.0268, \chi^2 = 4.91) \).

Comparing the LOH/MSI frequencies in all specimens according to the AJCC classification (IA–B, IIA–B, IIIA–IIIB), statistically significant high LOH/MSI frequencies for D3S1583 marker (RARβ) in AJCC IA–B group \( (p = 0.0001, \chi^2 = 14.10) \) and for D3S1611 marker (MLH1) in AJCC IIA–B group \( (p = 0.0002, \chi^2 = 14.10) \) were found.

Statistically significant low LOH/MSI frequency was revealed for both D3S3615 (RASSF1A; \( p = 0.0018, \chi^2 = 9.73 \)) and D3S3611 (VHL; \( p = 0.0001, \chi^2 = 18.35 \)) markers in pT1 (TNM classification) and in IA–B group (AJCC classification; RASSF1A; \( p = 0.0015, \chi^2 = 10.06; \) VHL \( p = 0.034, \chi^2 = 4.50 \)).

**Fig. 3** Box-and-whisker plots, representing mean LOH/MSI frequencies in the studied groups, according to: a age \( (p = 0.006, \text{Kruskal–Wallis test}); \) b NSCLC histopathological subtypes \( (p = 0.007, \text{U Mann–Whitney test}); \) c the period of smoking \( (p = 0.007, \text{U Mann–Whitney test}); \) d PYs (number of cigarettes smoked in a lifetime) \( (p = 0.004, \text{U Mann–Whitney test}) \).
Regarding NSCLC histopathological subtypes (SCC and non-SCC), statistically significant high LOH/MSI frequency was observed for D3S1611 marker (MLH1) in SCC group \( (p = 0.011, \chi^2 = 6.47) \) and for D3S1583 marker (RAR\( \beta \)) in non-SCC group \( (p = 0.008, \chi^2 = 6.84) \).

Total LOH/MSI frequency in 3p region and tobacco smoking

Regarding smoking history, patients were divided into two groups, taking into account the duration of smoking \(<40 \) and \( \geq 40 \) years), and the two other groups, taking into account the number of cigarettes smoked in a lifetime, assessed as PYs \(<40 \) and \( \geq 40 \) PYs). The increased total LOH/MSI frequency significantly correlated with the longest smoking history and with the increased PYs. Statistically significant increase in LOH/MSI frequency in 3p region was observed in case of patients who had been smoking for more than 40 years and more than 40 PYs \( (p = 0.007 \) and \( p = 0.004 \), respectively; U Mann–Whitney test) (see Fig. 3c, d).

Total LOH/MSI frequency in particular loci and tobacco smoking

Total LOH/MSI frequency in each studied microsatellite locus considering patients’ smoking history, that is, the smoking period \(<40 \) and \( \geq 40 \) years) and the PYs \(<40 \) and \( \geq 40 \) PYs), was investigated.

Significantly high LOH/MSI frequency in 3p region, mainly for D3S1300 (FHIT) and D3S1611 (MLH1) markers in correlation with cigarette addiction in a lifetime \( \geq 40 \) years \( (p = 0.015, \chi^2 = 5.88 \) and \( p = 0.009, \chi^2 = 6.82, \) respectively) and \( \geq 40 \) PYs \( (p = 0.015, \chi^2 = 5.91 \) and \( p = 0.039, \chi^2 = 4.22, \) respectively), was found. Interestingly, statistically significant low LOH/MSI frequency for one marker, that is, D3S3615 (RASSF1A) was revealed in correlation with cigarette addiction in a lifetime \( \geq 40 \) years \( (p = 0.020, \chi^2 = 5.35, \) as well as \( \geq 40 \) PYs \( (p = 0.002, \chi^2 = 9.57) \). The results are shown in Table 3.

Discussion

In our study, total LOH/MSI frequency in NSCLC samples was found in the range of 24–42%, depending on the marker. The discrepancies between our results and those of others, who report 50–80% frequency of LOH in NSCLC [25–27], may result from different markers used for LOH/MSI analysis, method of detection and evaluation of LOH/MSI, as well as from the population-based differences. However, we confirmed the implication of smoking as a very important causative factor in lung carcinogenesis supporting the hypothesis that cigarette smoking—both current and former—might induce molecular alterations in genes localized in 3p [27–30]. Additionally, we documented significantly higher total LOH/MSI frequency in SCC versus non-SCC subtype of lung cancer that is most closely associated to smoking.

Frequent allelic losses in 3p in lung carcinogenesis suggest the presence of multiple TSGs in this chromosomal region; however, only few genes have strong evidence supporting their candidacy as important in lung cancer [31, 32]. RASSF1A gene, located in 3p21.3, frequently shows loss of expression in lung cancer cells. Two mechanisms of RASSF1A inactivation have been confirmed in lung tumors, namely LOH and promoter hypermethylation [26, 33]. Our study revealed rather low level of allelic loss (about 24%) in RASSF1A locus, as compared to other studied genes. It could support the hypothesis of other investigators who suggest the lesser importance of LOH in RASSF1A inactivation in NSCLC tumorigenesis and the prevalence of epigenetic modifications [33, 34]. Our results may suggest the role of another, as yet unidentified, 3p21.3 TSG gene/s important in NSCLC. Indeed, as so far besides RASSF1A, at least 7 other candidate TSGs (CACA2D2, PL6, 101F6, NPRL2, BLU, TUSC2 and HYAL2) have been identified in the 600-kb 3p21.3 homozygous deletion region [27, 32]. It still remains to be elucidated which gene/genes localized in this particular chromosomal region play a vital role and which mechanisms (promoter hypermethylation and/or LOH) resulting in TGS silencing are pivotal in lung carcinogenesis.

Another candidate gene in our study, FHIT, is located in the FRA3B fragile site at 3p14.2. Loss of FHIT expression is observed in lung cancer and pre-neoplastic lesions [30, 35, 36]. As found by Toledo et al. [37], LOH at FHIT gene in NSCLC is associated with high proliferation and low apoptotic level. In our study, LOH frequency in FHIT gene (33–36%) was lower than that reported by others (44–58%) [38, 39]; however, we observed significant increase in LOH/MSI frequency in FHIT locus in correlation with cigarette addiction in a lifetime. It is in accordance with the observation of other authors and may support the hypothesis that cigarette smoking could induce molecular alterations of FHIT [29, 35]. Regarding other studied correlations, we did not recognize any associations between FHIT loss of heterozygosity and patient’s clinical features, outcome or metastatic behavior of tumor but we confirmed the association of frequent FHIT LOH with small size (pT1) of tumors, confirming the role of FHIT in the initiation of lung tumorigenesis. This is in agreement with the results of others [35, 40, 41].

Our analysis included also one of the genes implicated in the DNA mismatch repair system, that is, human MutL
homolog (hMLH1), located in 3p22.2. Microsatellite instability in this chromosomal region is confirmed in lung cancer where it is recognized in 38–68 % of NSCLC patients [42, 43]. The results of our study confirm high frequency of hMLH1 LOH (40.38 %). Based on our own findings and those of others, it may be concluded that AI in this locus (probably combined with epigenetic alteration) seems to be one of major events involved in lung carcinogenesis [43–45]. The significant correlation between hMLH1 LOH and advanced stage of tumors (III A/B) found in our study might reflect the role of hMLH1 in the late phase of carcinogenesis due to the accumulation of unpaired DNA lesions. Additionally, our results indicate significantly increased frequency with heavy smokers, assessed by daily tobacco uptake and total smoking exposure [42].

The gene encoding retinoic acid receptor beta (RARβ) has been found to be downregulated in lung tumors, suggesting its role lung carcinogenesis [46]. Frequent allelic losses in RARβ locus have been confirmed in lung tumors, NSCLC cell lines and in lung cancer precursor lesions [47, 48]. In our analysis, the frequency of LOH in RARβ locus was shown to be the highest among the studied loci, reaching nearly 42 %. This is in agreement with the results obtained by others [49]. Allelic loss in RARβ locus was also observed in smokers [47, 50]. In our study, we did not confirm this observation; however, the significant association between LOH in RARβ locus and stage I NSCLC found in our study might confirm the role of this gene in suppressing the lung tumorigenicity at its early stage.

The other candidate gene located in 3p region and included in our study is the von Hippel–Lindau (VHL) TSG. We used two markers flanking VHL locus (3p25.3). The observed frequency of AI found in the studied NSCLC samples, that is, about 30 %, appeared to be lower than that reported by others (46–63 %) [38, 39]. In some studies, higher LOH frequency was found in squamous cell carcinoma as compared with adenocarcinoma tumors [38, 51], although not confirmed in another study [39] neither in our analysis. Additionally, VHL LOH was more frequent in tumors from smokers as compared to those from non-smokers [38], which has not been confirmed in our study.

Reassuming, it should be stressed that respiratory epithelium carcinogenesis is a multifactorial process which includes inherited and acquired genetic changes (e.g., LOH/MSI), as well as epigenetic alterations. Additionally, cigarette smoking recognized as a one of the pivotal factors in lung cancer development, assessed in correlation with 3p region allelic losses provides controversial results. Therefore, the assessment of LOH/MSI in particular TSG loci separately and its correlation with smoking addiction might confirm the diagnostic and/or prognostic value of some genes (as in case of RARβ and hMLH1 in our study) and the influence of cigarette smoking on gene alteration (FHIT and hMLH1 in our study)—which seems to be promising.

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Table 3 LOH/MSI frequencies (%) in NSCLC patients regarding their smoking habits

| Marker | D3S1234 | D3S1300 | D3S3615 | D3S1611 | D3S1583 | D3S1317 | D3S3611 | U Mann–Whitney test p |
|--------|---------|---------|---------|---------|---------|---------|---------|-----------------------|
| Smoking time period (years) | | | | | | | | |
| <40 | 35 | 16 | 18 | 28 | 31 | 24 | 27 | 0.007 |
| ≥40 | 41 | 55 | 30 | 56 | 42 | 40 | 31 | |
| Pack years (PYs) | | | | | | | | |
| <40 PYs | 33 | 8 | 19 | 21 | 21 | 26 | 22 | 0.004 |
| ≥40 PYs | 41 | 54 | 25 | 52 | 45 | 37 | 34 | |
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