RESEARCH ARTICLE

Inferring Single Nucleotide Polymorphisms in MicroRNA Binding Sites of Lung Cancer-related Inflammatory Genes

Fei He¹, Ling-Ling Zheng², Wen-Ting Luo³, Rong Yang⁴, Xiao-Qin Xu⁵, Lin Cai^{1*}

Abstract

Single nucleotide polymorphisms located at microRNA (miRNA)-binding sites are likely to affect the expression of miRNA targets and may contribute to the susceptibility of humans to common diseases. Here 335 candidate lung cancer-related inflammatory genes were selected according to the existing literature and database. We identified putative miRNA-binding sites of 149 genes by specialised algorithms and screened SNPs in the 3'UTRs of these genes. By calculating binding free energy, we sorted 269 SNPs on the basis of the possibility of prediction. The proposed approach could help to easy the identification of functionally relevant SNPs and minimize the workflow and the costs.

Keywords: Algorithms- inflammatory- lung cancer- microrna- polymorphisms- 3'UTR

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Introduction

Since the first noted by the pathologist Rudolph Virchow who found inflammatory cells are present within tumors (Coussens et al., 2002), epidemiological evidences now indicate inflammation mediates oncogenesis and chronic inflammation contributes to about 25% of all human cancers (Hussain et al., 2007). Inflammation induces carcinogenesis maybe by the following mechanism (Ohnishi et al., 2013): reactive oxygen species (ROS), reactive nitrogen species (RNS), harmful endogenous genotoxic substances, generated from inflammatory and epithelial cells result in oxidative and nitrative DNA damage and are involved in the initiation and/or promotion of inflammation-mediated carcinogenesis.

Lung cancer is the most common malignant tumor and the primary reason of global neoplasm deaths (Jemal et al., 2011). Recently, there were also numerous of laboratory and clinical studies have extensively reported the relationship between inflammation and lung cancer, as well as a current review (Cho et al., 2011; Hattar K et al., 2013). Although the exact cause of lung inflammation leading to carcinogenesis is not known, there was the hypothesis (Houghton et al., 2008) demonstrated that chronic airway inflammation alters the bronchial epithelium and lung microenvironment and the expression of oncogenes and tumor suppressor genes might be induced to cause to neoplastic transformation.

Pulmonary inflammation plays a risk role in promoting development of lung cancer. Several kinds of conditions

bring about lung inflammation, including tobacco smoke, occupational hazards and pathogen infections. Cigarette smoke contains great amount of carcinogens and modulates inflammation and promotes chronic inflammation in the conducting airways by impairing innate host defense mechanisms (Lee et al., 2009; Lee et al., 2012). Another condition contributes to inflammation is pathogen. Infection triggers the inflammatory response which is a part of normal host defense, preceding tumor development. However, tumorigenic pathogens subvert host immunity and establish persistent infections associated with low-grade but chronic inflammation and then dysregulate inflammatory cytokines and transcription factors (Grivennikov et al., 2010).

Lung cancer cells use inflammatory cytokines and their receptors for tumor growth, invasion, migration and metastasis. MicroRNAs (miRNAs) are endogenous non-coding single-stranded RNAs of about 21-22 nucleotides in length, which can regulate gene translation and modulate gene expression at post-transcriptional level during the most biological and pathological processes (Lagos-Quintana et al., 2001). At present, many researchers (Saunders et al., 2007) considered that single nucleotide polymorphisms (SNPs) in the miRNA seed sequence have higher probability of affecting miRNAs function. Therefore, supposed that SNPs occur in the binding site between miRNAs and mRNAs (usually in 3' untranslated region, UTR) in genes of inflammation signaling pathways, which may weaken or reinforce the expression of miRNA target genes and then, modify

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| Gene | rs | MAF | Common- variation | miRNA | $ \Delta\Delta G _{average}$ (kJ/mol) | Gene | rs | MAF | Common- variation | miRNA Iz | ∆∆Gl _{average} (kJ/mol) |
|-----------------|--------------------------|----------------|----------------------|---|--|------------------|------------|-------|----------------------|--|-------------------------------------|
| CD4 SARM1 | rs3213427 rs118138669 | 0.134 0.05 | T-C C-T | hsa-miR-1272 hsa-miR-31 | 24.2 9.44 | CDK6 | rs42377 | 0.116 | A-G | hsa-miR-487a hsa-miR-154 | 2.97 |
| | | | | hsa-miR-140 hsa-miR-3922 hsa-miR-4708 | | NFATC4 | rs10362 | 0.142 | G-T | hsa-miR-144 hsa-miR-1207-5p hsa-miR-486-3p | 2.97 |
| | | | | hsa-miR-4753 | | ITAL 1 | 10(05004 | 0.05 | TC | hsa-miR-625 | 2.0 |
| | | | | hsa-miR-4701 | | IFNA1 CVCL 12 | rs12685904 | 0.05 | T-G | hsa-miR-1911 | 2.9 |
| IRF1 | rs6873426 | 0.153 | G-T | hsa-miR-4696 hsa-miR-214 | 9.3 | CXCL12 | rs1801157 | 0.244 | G-A | hsa-miR-941 hsa-miR-149 | 2.9 |
| SMAD3 | rs8031627 | 0.453 | A-G | hsa-miR-596 | 6.5 | | | | | hsa-miR-631 | |
| CDKN1B | rs4251697 | 0.085 | G-A | hsa-miR-575 | 6.1 | IL1A | rs1304037 | 0.073 | A-G | hsa-miR-298 | 2.9 |
| KSR1 | rs1075952 | 0.293 | A-G | hsa-miR-3136 | 6 | 11111 | 151501057 | 0.075 | no | hsa-miR-548 | 2.9 |
| SLC44A2 | rs10948 | 0.326 | T-G | hsa-miR-489 | 6 | SMAD4 | rs28403611 | 0.067 | A-G | hsa-miR-4803 | 2.77 |
| GATA3 | rs1058240 | 0.061 | A-G | hsa-miR-95 | 5.7 | | | | | hsa-miR-552 | |
| AGT | rs7079 | 0.098 | C-A | hsa-miR-218-2 | 5.5 | | | | | hsa-miR-125a-3p | |
| PPM1A | rs6573305 | 0.05 | G-T | hsa-miR-4789-3p | 5 | CD4 | rs3829972 | 0.354 | G-A | hsa-miR-1206 | 2.75 |
| CCND1 | rs678653 | 0.091 | C-G | hsa-miR-602 | 5 | | | | | hsa-miR-663 | |
| GREM1 | rs7162202 | 0.244 | C-T | hsa-miR-3921 | 4.5 | GHR | rs2910875 | 0.321 | C-T | hsa-miR-103 | 2.7 |
| TXNIP | rs7212 | 0.144 | C-G | hsa-miR-296-3p | 4.4 | MAP2K7 | rs3745386 | 0.085 | A-G | hsa-miR-4252 | 2.68 |
| PRLR | rs392279 | 0.292 | A-G | hsa-miR-3164 hsa-miR-3118 | 4.35 | | | | | hsa-miR-4488 hsa-miR-4697-5p | 1 |
| | | | | hsa-miR-134 | | | | | | hsa-miR-4420 | |
| | | | | hsa-miR-4501 | | | | | | hsa-miR-637 | |
| SMAD3 | rs1052488 | 0.314 | T-C | hsa-miR-544 | 4.3 | | | | | hsa-miR-181a-2 | |
| EDARADD | rs61737025 | 0.092 | T-C | hsa-miR-362-3p | 4.23 | TLR4 | rs11536889 | 0.216 | G-C | hsa-miR-1208 | 2.65 |
| | | | | hsa-miR-425 | | | | | | hsa-miR-1236 | |
| | | | | hsa-miR-329 | | COL1A1 | rs1061947 | 0.081 | C-T | hsa-miR-1260 | 2.63 |
| TLR10 | rs11466661 | 0.317 | A-C | hsa-miR-3667 | 4.2 | | | | | hsa-miR-532-3p | |
| CFLAR | rs2881929 | 0.217 | G-T | hsa-miR-3678-3p | | | | | | hsa-miR-328 | |
| CUID | 2072016 | 0.000 | | hsa-miR-1245b-3 | • | MAPK14 | rs6457878 | 0.19 | C-T | hsa-miR-449a/b | 2.6 |
| GHR | rs2973016 | 0.092 | A-G | hsa-miR-564 | 3.9 | CDKN1A | rs1059234 | 0.453 | C-T | hsa-miR-3169 | 2.6 |
| 07472 | 2744402 | 0.042 | тc | hsa-miR-4253 | 2.0 | | | | | hsa-miR-509-5p | |
| STAT3 | rs3744483 | 0.243 | T-C | hsa-miR-1255 | 3.9 | VDAS | | 0 222 | A.C. | hsa-miR-1288 | 2.55 |
| TAB2 | rs7896 | 0.1 | C-G | hsa-miR-3170 hsa-miR-4260 | 3.85 | KRAS | rs8720 | 0.233 | A-G | hsa-miR-374b | 2.55 |
| TLR6 | rs73236628 | 0.475 | C-T | hsa-miR-494 | 3.8 | PRLR | rs249522 | 0.233 | G-A | hsa-miR-885-5p hsa-miR-616 | 2.55 |
| PANX1 | rs1046805 | 0.256 | A-G | hsa-miR-10a | 3.7 | IKLK | 18249322 | 0.255 | 0-A | hsa-miR-4289 | 2.33 |
| FANAI | 1510 10005 | 0.250 | 110 | hsa-miR-4732-3p | | TNFSF11 | rs1054016 | 0.456 | G-T | hsa-miR-3613-3p | 2.5 |
| | | | | hsa-miR-212-3p | | MAPK11 | rs2072878 | 0.1 | A-G | hsa-miR-4300 | 2.47 |
| ETS2 | rs1051476 | 0.239 | C-G | hsa-miR-1293 hsa-miR-323-5p | 3.7 | | | | | hsa-miR-920 hsa-miR-939 | |
| GREM1 | rs17816260 | 0.208 | C-A | hsa-miR-190 | 3.6 | CCND2 | rs3217926 | 0.105 | T-C | hsa-miR-596 | 2.4 |
| GREAT | 1517010200 | 0.200 | en | hsa-miR-299 | 5.0 | SOCS5 | rs41379147 | 0.05 | C-T | hsa-miR-331-5p | 2.4 |
| MAP4K4 | rs12622613 | 0.092 | A-G | hsa-miR-4328 | 3.6 | CFLAR | rs7558475 | 0.016 | A-G | hsa-miR-3919 | 2.4 |
| VEGFA | rs3025053 | 0.098 | G-A | hsa-miR-324 | 3.45 | MAP3K2 | rs1129121 | 0.083 | C-T | hsa-miR-15 | 2.4 |
| CXCL12 | rs1029153 | 0.232 | T-C | hsa-miR-3130 hsa-miR-499 | 3.4 | | | | | hsa-miR-195 hsa-miR-16 | |
| | | | | hsa-miR-34 | | | | | | hsa-miR-497 | |
| ERK2(MAPK1) | rs6928 | 0.439 | G-C | hsa-miR-602 | 3.35 | MAP3K1 | rs702688 | 0.078 | T-C | hsa-miR-3682 | 2.4 |
| TI D 4 | 7072704 | 0.077 | C-G | hsa-miR-602 | 2.2 | NFATC4 | rs11848279 | 0.477 | A-G | hsa-miR-1285 | 2.4 |
| TLR4 | rs7873784 | 0.067 | G-C | hsa-miR-144 | 3.3 | | | | | hsa-miR-3680 | |
| MAPK13 STAT6 | rs2071863 | 0.389 | C-T | hsa-miR-34a | 3.3 | | | | | hsa-miR-3180-5p | |
| PPM1A | rs703817 rs2273623 | 0.337 0.134 | G-A A-G | hsa-miR-216a hsa-miR-4471 | 3.3 3.25 | RELB | rs28372683 | 0.1 | G-T | hsa-miR-612 hsa-miR-1224 | 2.4 |
| IIWIIA | 182273023 | 0.154 | A-0 | hsa-miR-1292 | 3.23 | KELD | 1820572005 | 0.1 | 0-1 | hsa-miR-596 | 2.4 |
| NLRC5 | rs27193 | 0.133 | A-G | hsa-miR-490-3p | 3.2 | | | | | hsa-miR-523 | |
| MAPK10 | rs1201 | 0.321 | A-G | hsa-miR-147 | 3.1 | | | | | hsa-miR-29c | |
| SMAD3 | rs8031440 | 0.463 | G-A | hsa-miR-619 | 3.03 | | | | | hsa-miR-138-1 | |
| | | | | hsa-miR-490-3p | - | | | | | hsa-miR-34a | |
| | | | | hsa-miR-1299 | | PAK1 | rs2844337 | 0.268 | T-G | hsa-miR-4761-5p | 2.4 |
| CCND2 | rs3217925 | 0.407 | C-T | hsa-miR-3202 | 3 | | | | | hsa-miR-320c | |
| MAPK10 | rs3527 | 0.05 | A-G | hsa-miR-466 | 3 | MAP2K1 | rs14303 | 0.108 | G-T | hsa-miR-3658 | 2.4 |
| ETS2 | rs1051475 | 0.225 | C-T | hsa-miR-4270 | 3 | JAK3 | rs79044512 | 0.075 | A-T | hsa-miR-502-5p | 2.37 |
| NR2C2 | rs28524664 | 0.092 | A-T | hsa-miR-129 | 3 | | | | | hsa-miR-659 | |
| | | | | | | | | | | hsa-miR-4796-3p | |

miRNA targeting activities and modulate the level of inflammation in response to various inflammatory stimuli. Many studies evaluated the possible associations between lung cancer and polymorphisms (Chen et al., 2013; Cheng et al., 2013; Zu et al., 2013; Kim et al., 2014).

In this study, we selected lung cancer-related genes that belong to inflammation pathways responding to microorganism and cigarette smoking and hoped to catalogue SNPs, which might affect the expression levels of the target genes. It will provide data for the followup studies on susceptibility or prognosis, functional verification and build evidence for diagnosis and treatment of lung cancer.

Materials and Methods

Selection of candidate genes

We focused on the genes which related to lung cancer participated the inflammatory reaction to microorganisms, such as chlamydia pneumonia, mycobacterium tuberculosis or human immunodeficiency virus (HIV) and tobacco smoking. Candidate genes are retrieved according to the

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| | | | A for the Gene | | 0 | | | |
|------------------------------|------------------|--------------------------|---------------------------------|----------------------|--------------------------|------------------------------|----------------------|--------------------------|
| miRNA | Gene | rs | miRNA | Gene | rs | miRNA | Gene | rs |
| hsa-miR-103 | GHR | rs2910875 | hsa-miR-297 | FCER1A | rs79965525 | hsa-miR-509-5p | CDKN1A | rs1059234 |
| hsa-miR-106b | EGFR | rs884225 rs117378779 | hsa-miR-297 | CXCL9 | rs1050176 | hsa-miR-509-5p | PANX1 | rs12800562 |
| lisa-mik-1000 | PSTPIP1 PRLR | rs371913 | hsa-miR-299 hsa-miR-299 | GREM1 IL12A | rs17816260 rs568408 | hsa-miR-510 hsa-miR-510 | RIPK1 MAPK9 | rs17548629 rs34095777 |
| hsa-miR-1197 | RIPK1 | rs17548629 | hsa-miR-30 | CCNA2 | rs1507987 | hsa-miR-512-5p | IL4R | rs2074570 |
| | NLRX1 | rs45439091 | hsa-miR-30 | F2R | rs1801719 | hsa-miR-512-5p | MAPK9 | rs34095777 |
| | TICAM1 | rs1046673 | hsa-miR-300 | SOCS6 | rs7231397 | hsa-miR-518a-5p | SMAD3 | rs3743342 |
| hsa-miR-1202 | EDARADD | rs6428955 | hsa-miR-300 | ERK2(MAPK1) | rs2276008 | hsa-miR-518a-5p | PANX1 | rs12800562 |
| hsa-miR-1208 | MAPK8IP3 TLR4 | rs2575329 rs11536889 | hsa-miR-302 hsa-miR-302 | INSR STAT3 | rs1366600 rs1053023 | hsa-miR-519d hsa-miR-519d | XIAP PRLR | rs17330644 rs371913 |
| hsa-miR-1208 | PAK1 | rs2729762 | hsa-miR-302 | KRAS | rs7973450 | hsa-miR-520 | INSR | rs1366600 |
| hsa-miR-1224 | RELB | rs28372683 | hsa-miR-302 | GREM1 | rs33963919 | | TIRAP | rs8177375 |
| hsa-miR-1224 | TOLLIP | rs3168046 | hsa-miR-3128 | SLC44A2 | rs2288902 | | GREM1 | rs33963919 |
| hsa-miR-1233 | CFLAR | rs13035714 | hsa-miR-3128 | GREM1 | rs3743104 | | MMP2 | rs7201 |
| hsa-miR-1233 | MAPK10 | rs958 | hsa-miR-3147 | PAK1 | rs2729762 | hsa-miR-527 | SMAD3 | rs3743342 |
| hsa-miR-1236 hsa-miR-1236 | TLR4 TIRAP | rs11536889 rs8177375 | hsa-miR-3147 hsa-miR-3180-5p | TXNIP NFATC4 | rs4755 rs11848279 | hsa-miR-527 hsa-miR-544 | PANX1 SMAD3 | rs12800562 rs1052488 |
| hsa-miR-1250 | IRF1 | rs6894655 | hsa-miR-3180-5p | NLRC5 | rs3751705 | 115a-1111K-J++ | KRAS | rs12245 |
| hsa-miR-125 | MAPK10 | rs958 | hsa-miR-32 | PANX1 | rs12800562 | | CDK2 | rs2069415 |
| hsa-miR-1255 | STAT3 | rs3744483 | hsa-miR-32 | FCER1A | rs79965525 | hsa-miR-545 | CCNA2 | rs1507987 |
| hsa-miR-1255 | GBP1 | rs2624 | hsa-miR-329 | EDARADD | rs61737025 | hsa-miR-545 | EDARADD | rs6428955 |
| hsa-miR-1260 | COL1A1 | rs1061947 | | EDARADD | rs61736989 | hsa-miR-548 | IL1A | rs1304037 |
| hsa-miR-1260 | TOLLIP | rs3168046 | | BTK | rs700 | | NLRP3 | rs10802502 |
| hsa-miR-1275 hsa-miR-1275 | PAX5 CSF1R | rs3739437 rs2066934 | hsa-miR-338-5p | MALT1 XIAP | rs2319974 rs17334746 | hsa-miR-552 | MMP2 SMAD4 | rs7201 rs28403611 |
| hsa-miR-1275 | NR2C2 | rs28524664 | hsa-miR-338-5p | NLRP3 | rs10754558 | hsa-miR-552 | SMAD4 SMAD3 | rs117707762 |
| hsa-miR-129 | BTK | rs1057403 | hsa-miR-34a | MAPK13 | rs2071863 | hsa-miR-561 | CASP8 | rs13113 |
| hsa-miR-129-5p | COL1A1 | rs75713851 | hsa-miR-34a | RELB | rs28372683 | | KRAS | rs9266 |
| hsa-miR-129-5p | TNFSF10 | rs11720451 | hsa-miR-3609 | SMAD3 | rs3743342 | | SMAD3 | rs117707762 |
| hsa-miR-132 | MAP3K7 | rs3734657 | | XIAP | rs17330644 | hsa-miR-570 | ECSIT | rs1062958 |
| hsa-miR-132 | IL2RA PANX1 | rs12722604 rs12800562 | hsa-miR-3611 | PRLR TLR6 | rs371913 rs5743831 | | INSR IL33 | rs1052371 rs1048274 |
| hsa-miR-137 hsa-miR-137 | INSR | rs3745550 | hsa-miR-3611 | PRLR | rs379899 | | BCL2 | rs3744935 |
| hsa-miR-144 | TLR4 | rs7873784 | hsa-miR-3616-5p | XIAP | rs9856 | hsa-miR-573 | XIAP | rs9856 |
| | CDK6 | rs42377 | 1 | PRLR | rs379899 | | PRLR | rs379899 |
| | CDK6 | rs42035 | | PRLR | rs371913 | | PRLR | rs371913 |
| hsa-miR-147 | MAPK10 | rs1201 | hsa-miR-362-3p | EDARADD | rs61737025 | hsa-miR-575 | CDKN1B | rs4251697 |
| hsa-miR-147 | CXCL12 | rs1065297 | hsa-miR-362-3p | MALT1 DANK1 | rs2319974 | hsa-miR-575 | VEGFA | rs3025040 |
| hsa-miR-15 | MAP3K2 SARM1 | rs1129121 rs739439 | hsa-miR-363 hsa-miR-363 | PANX1 FCER1A | rs12800562 rs79965525 | hsa-miR-578 hsa-miR-578 | BTK CCND2 | rs1057403 rs3217929 |
| | CXCL12 | rs1804429 | hsa-miR-3646 | CCND2 | rs3217923 | hsa-miR-579 | NFKBIA | rs8904 |
| hsa-miR-151-5p | COL1A1 | rs75713851 | hsa-miR-3646 | SMAD5 | rs3206633 | hsa-miR-579 | TNFSF10 | rs1131535 |
| hsa-miR-151-5p | CCNE1 | rs3218073 | hsa-miR-3667 | TLR10 | rs11466661 | hsa-miR-591 | OSM | rs2070889 |
| hsa-miR-155 | CSF1R | rs3828609 | hsa-miR-3667 | SARM1 | rs2239910 | hsa-miR-591 | VEGFA | rs3025039 |
| hsa-miR-155 | SPI1 | rs1057233 | hsa-miR-367 | PANX1 | rs12800562 | hsa-miR-592 | SLC44A2 | rs2288902 |
| hsa-miR-16 hsa-miR-16 | MAP3K2 SARM1 | rs1129121 rs739439 | hsa-miR-367 hsa-miR-369 | FCER1A IRF1 | rs79965525 rs6894655 | hsa-miR-592 hsa-miR-596 | ERK2(MAPK1) SMAD3 | rs13515 rs8031627 |
| hsa-miR-17 | INSR | rs1366600 | hsa-miR-369 | CXCL9 | rs10337 | 118a-1111K-390 | CCND2 | rs3217926 |
| hsa-miR-17 | PRLR | rs371913 | hsa-miR-374 | CCL2 | rs13900 | | RELB | rs28372683 |
| hsa-miR-181 | CD4 | rs16932921 | | IRF1 | rs6894655 | hsa-miR-602 | CCND1 | rs678653 |
| hsa-miR-181 | KRAS | rs9266 | | ETS2 | rs530 | hsa-miR-602 | ERK2(MAPK1) | rs6928 |
| hsa-miR-182 | RAC1 | rs9374 | 1 | CXCL9 | rs10337 | hsa-miR-603 | E2F1 | rs3213180 |
| hsa-miR-182 | GREM1 | rs3743104 | hsa-miR-381 | SOCS6 | rs7231397 | | MAPK9 | rs34095777 |
| hsa-miR-190 hsa-miR-190 | GREM1 GREM1 | rs17816260 rs33963919 | hsa-miR-381 hsa-miR-383 | ERK2(MAPK1) IL12A | rs2276008 rs568408 | hsa-miR-612 | MALT1 NFATC4 | rs2319974 rs11848279 |
| hsa-miR-193 | CXCL2 | rs9131 | hsa-miR-383 | IL1211 IL1R1 | rs3732131 | hsa-miR-612 | TXNIP | rs4755 |
| hsa-miR-193 | ERK1(MAPK3) | rs7542 | hsa-miR-411 | CTSB | rs9009 | hsa-miR-625 | NFATC4 | rs10362 |
| hsa-miR-194 | MAP3K7 | rs3734657 | hsa-miR-411 | HSP90AA1 | rs1059623 | hsa-miR-625 | PAX5 | rs3739437 |
| | CCND1 | rs7177 | hsa-miR-424 | SARM1 | rs739439 | hsa-miR-637 | MAP2K7 | rs3745386 |
| 1 | CDK6 | rs42035 | hsa-miR-424 | CD80 | rs17281703 | | PSTPIP1 | rs117378779 |
| hsa-miR-195 hsa-miR-195 | MAP3K2 SARM1 | rs1129121 rs739439 | hsa-miR-4253 hsa-miR-4253 | GHR TNFSF4 | rs2973016 rs16845543 | hsa-miR-658 | NLRP12 CD8A | rs10410581 rs1051386 |
| hsa-miR-195 | CD80 | rs17281703 | hsa-miR-4255 | NLRC5 | rs3751705 | hsa-miR-658 | MEFV | rs2741918 |
| hsa-miR-205 | CCNA2 | rs1507987 | hsa-miR-4257 | GBP1 | rs2296883 | hsa-miR-663 | CD4 | rs3829972 |
| hsa-miR-205 | SARM1 | rs2239910 | hsa-miR-4260 | TAB2 | rs7896 | hsa-miR-663 | MAPK8IP3 | rs118077547 |
| hsa-miR-205 | F2R | rs1801719 | hsa-miR-4260 | STAT6 | rs324015 | hsa-miR-665 | RIPK1 | rs17548629 |
| hsa-miR-20b | CD27 | rs1059501 | hsa-miR-4266 | NFKBIA | rs2273650 | | STAT5A | rs3198502 |
| hsa-miR-20b hsa-miR-212 | PRLR MAP3K7 | rs371913 rs3734657 | hsa-miR-4266 hsa-miR-4270 | CTSB ETS2 | rs9009 rs1051475 | hsa-miR-877 | TLR8 TIRAP | rs5744088 rs8177375 |
| hsa-miR-212 | CD80 | rs1599795 | hsa-miR-4270 | OSM | rs1051475 rs2070890 | 115a-1111 K -0 / / | KRAS | rs8177375 rs712 |
| hsa-miR-212 | IRF1 | rs6873426 | hsa-miR-4276 | F2R | rs1801719 | | MYD88 | rs7744 |
| | MAP2K2 | rs6629 | hsa-miR-4276 | TAB2 | rs2744434 | hsa-miR-885-5p | KRAS | rs8720 |
| | SARM1 | rs739439 | hsa-miR-4327 | INSR | rs3745551 | hsa-miR-885-5p | CCND3 | rs9529 |
| hsa-miR-216a | STAT6 | rs703817 | hsa-miR-4327 | CD4 | rs7901 | hsa-miR-888 | SOCS5 | rs4953419 |
| hsa-miR-216a | NLRP4 SMAD5 | rs302457 | hsa-miR-4658 | ERK2(MAPK1) MEEV | rs1063311 rs450021 | | MAPK10 ATE1 | rs17011312 rs820125 |
| hsa-miR-216b | SMAD5 MEFV | rs3206633 rs2741918 | hsa-miR-4658 hsa-miR-4795-5p | MEFV PRLR | rs450021 rs379899 | hsa-miR-92a | ATF1 PANX1 | rs829125 rs12800562 |
| | MAX | rs4902357 | hsa-miR-4795-5p | PRLR | rs371913 | | FCER1A | rs79965525 |
| hsa-miR-217 | RIPK2 | rs16900627 | hsa-miR-4796-3p | JAK3 | rs79044512 | hsa-miR-92b | PANX1 | rs12800562 |
| | PANX1 | rs12800562 | r | XIAP | rs17330644 | | PSTPIP1 | rs117378779 |
| | KSR1 | rs2241906 | | PRLR | rs371913 | hsa-miR-93 | INSR | rs1366600 |
| hsa-miR-223 | CCND1 | rs7177 | hsa-miR-486-3p | NFATC4 | rs10362 | | XIAP | rs17330644 |
| | FCER1A | rs79965525 | hsa-miR-486-3p | EGFR | rs884225 | hee m ¹ D 020 | PRLR MADE 11 | rs371913 |
| hea miP 25 | BTK PANY 1 | rs12800562 | hsa-miR-492 | IKBKE CD80 | rs10836 rs57271503 | hsa-miR-939 | MAPK11 OSM | rs2072878 |
| hsa-miR-25 | PANX1 FCER1A | rs12800562 rs79965525 | hsa-miR-492 hsa-miR-494 | CD80 TLR6 | rs57271503 rs73236628 | | OSM CCND1 | rs2070890 rs7177 |
| hsa-miR-490-3p | NLRC5 | rs27193 | hsa-miR-494 | CREBBP | rs9392 | | MAPK9 | rs1127580 |
| hsa-miR-490-3p | SMAD3 | rs8031440 | hsa-miR-497 | MAP3K2 | rs1129121 | hsa-miR-96 | IL12A | rs568408 |
| hsa-miR-296-3p | CD27 | rs1059501 | hsa-miR-497 | CD80 | rs17281703 | | RAC1 | rs9374 |
| hsa-miR-296-3p | TXNIP | rs7212 | | | | | | |

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information from the website (http://www.sabiosciences. com/Cytokines_Inflammation.php), BioCarta and KEGG pathways which are two common databases that provide displays of gene interactions for human cellular processes (http://cgap.nci.nih.gov/Pathways) and literatures (Shen et al., 2011; Yu et al., 2011; McMillan et al., 2011 ;Lee et al., 2012) commonly acknowledged as important inflammatory genes associated with smoking.

Materials

For this work, we used the followings web sites for prediction the miRNAs binding sites: 1) miRBase (http:// www.mirbase.org/); 2) miRanda (http://www.microrna. org/); 3) PicTar (http://pictar.mdc-berlin.de/); 4) Diana-MicroT v3.0 (http://diana.cslab.ece.ntua.gr/microT/); 5) TargetScan Human 6.0 (http://www.targetscan.org/); 6) Patrocles (http://www.patrocles.org/); 7) PolymiRTS Database 3.0 (http://compbio.uthsc.edu/miRSNP/); 8) microRNA-related SNP (http://www.bioguo.org/ miRNASNP/).

For selection of 3'UTR: 9) UCSC genome browser (http://www.genome.ucsc.edu).

For searching for SNP in the target sites: 10) SNP database (dbSNP 136; http://www.ncbi.nlm.nih.gov/SNP/).

For calculation of the Gibbs binding free energy: 11) RNAhybrid 2.2 (http://bibiserv.techfak.uni-bielefeld.de/ rnahybrid/submission.html).

Procedure

For each gene, we proceeded as follows:

1) The 3'UTR was identified according to the UCSC genome browser; 2) Putative miRNA-binding sites within the 3'UTR of each gene were identified by five specialized algorithms approaches as mentioned at materials 1.2.3.4.5; 3) The polymorphisms falling within the miRNA-binding sites identified as in stage 2 were searched in SNP database; 4) The polymorphisms were directly predicted by inserting the gene name into the web sites as recited above at materials 6.7.8; 5) The SNPs selection was performed based on frequencies reported for Chinese and the criterion was the SNPs having the minor allele frequency (MAF) lower than 0.05 were excluded. Because in future, we will apply case-control association study on this ethnic group to do the further research; 6) The algorithm of material 11 was run to assess the binding free energy (expressed as ΔG (kJ/mol), Gibbs free energy) for both for the common and the variant alleles identified as in stages 3 and 4; 7) The SNPs for their abilities to affect the binding of the miRNAs with their targets were evaluated by calculating as variation of ΔG (i.e., $\Delta \Delta G$), which was expressed as the difference in the energies between the two alleles was computed and used as the parameter for the assessment of the impact that each polymorphism shows for a given miRNA target site.

Results

335 candidate genes that were inflammation-related genes were retrieved from sabiosciences website, BioCarta and KEGG pathways and literatures, which contain the **3604** *Asian Pacific Journal of Cancer Prevention, Vol 15, 2014*

reactive genes under the stimulus of tobacco smoke and microorganisms.

Currently, it is difficult to judge which algorithm produces the most reliable or sensitive target predictions. A union of the result from five algorithm databases was performed for obtaining more reliable genes. Among the 335 candidate genes, 149 genes had miRNA target sequences in their 3'UTR regions, which were predicted by five algorithms. 32 SNPs didn't show any binding free energy at the miRNA target binding site. The remaining 142 genes showed 269 SNPs in the predicted miRNA-binding sites based on criterion (It only list the SNPs, whose $|\Delta\Delta G|_{average} \ge 2.37$ kJ/mol in Table 1 for lack of space).

For some genes, several miRNAs were predicted as the target sites, and others may be predicted to be targeted by only one miRNA. In order to account for these differences, as parameter for predicting the biological impact of each polymorphism, the average of the absolute values of $\Delta\Delta$ Gs should be used for each SNP (expressed as $|\Delta\Delta$ Glaverage). In order to give a priority list of SNPs having an impact on miRNA binding, we ranked the values of $|\Delta\Delta$ Glaverage and classified the SNPs in three groups corresponding to quartile. The first grade ($|\Delta\Delta$ Glaverage ≥ 2.37 kJ/mol) is composed of SNPs having a predicted high impact on the biology of the miRNA binding sites. The second grade ($0.60 < |\Delta\Delta$ Glaverage ≤ 2.37 kJ/mol) is composed of SNPs with a predicted mild biological activity, whereas within the last ($|\Delta\Delta$ Glaverage < 0.60kJ/mol) belong SNPs maybe with weakest activity.

For the all 269 SNPs, there were 202 miRNAs predicted for binding more than one SNP (Table 2). CDK6 rs42377 and rs42035 were found to have the shared target gene of hsa-miRNA-144; GREM1 rs17816260 and rs33963919 were found to have the shared target gene of hsa-miRNA-190; EDARADD rs61737025 and rs61736989 were found to have the shared target gene of hsa-miRNA-329; PRLR rs379899 and rs371913 were found to have the shared target gene of hsa-miRNA-329; pRLR rs379899 and rs371913 were found to have the shared target gene of hsa-miRNA-329; PRLR rs379899 and rs371913 were found to have the shared target gene of hsa-miRNA-3616-5p, hsa-miRNA-4795-5p and hsa-miRNA-573.

Discussion

In this study, we finally identified 269 SNPs within the miRNA-binding sites of 142 genes. Firstly, we chose 335 genes from inflammatory response pathway, including toll-like receptors (TLRs) signaling pathway, JAK/ STAT signaling pathway, NF- $\alpha\beta$ pathway and NOD-like receptors (NLRs) pathway and so on. Such genes play a key role on active immunity and passive immunity and whether in the extrinsic or intrinsic, inflammatory cells and molecules can impact the genomes of cancer cells through a variety of mechanisms. Then by detecting SNPs in miRNA target sites, we got 149 genes. Finally, we obtained 269 SNPs in 142 genes. And we also found that there were 164 SNPs shared the predicted binding miRNAs.

Among these SNPs, some were located in the seen regions of miRNA target sequences, whereas some were in other regions. Since 1993 the first discovery of miRNA, a great number of studies had classified genetic polymorphisms, which will affect miRNA regulation by various molecular mechanisms into three categories. First are polymorphisms within precursor miRNAs (premiRNAs); second are polymorphisms in miRNA-targetmRNA sites and third are variations in miRNA machinery genes (Mishra et al., 2009). It indicated that because SNPs in miRNA-target-mRNA sites are more likely to be under positive selection pressure, they tend to be deleterious, due to differences among various populations and contribute to diseases (Saunders et al., 2007). In current study, we focused SNPs in this region and it will be better for evaluating potential causative SNPs.

Due to the expensive experimental expenses and not suitable approaches, so many researchers conducted bioinformatics prediction and statistical analyses to investigate the diseases associated SNPs in miRNA target sites. Some studies introduced bioinformatics methods to identify a set of SNPs within miRNAs binding sites of genes (Ding et al., 2011; Song et al., 2014) and others verified the specific SNP by molecular experiments (Bhat et al., 2013). It is well-known that the functions of SNPs in miRNA-binding sites were miRNAs regulation and miRNAs or target genes expression. At present, there are many available algorithms and databases to predict miRNA target genes. The commonly rule in the method is that nucleotides 6-8 in the 5' end of the miRNA (called as 'seed' sequences) provide the maximal binding free energy of the miRNA-target duplex and the G:T pairing is admitted in the miRNA-target (John et al., 2004; Tomari et al., 2005). The most studies predict SNPs within miRNA binding sites and assess the potential functions of SNPs in 3'UTR via well-developed algorithms, based on the differences in the alignment scores and variations in binding free energy (Betel et al., 2010; Liu et al., 2012). In addition, some studies investigated the effects of SNPs according to the secondary structures of the miRNA binding sites by using RNAfold (Hariharan et al., 2009). And others employed a linear model to assess the effects of SNPs on the gene expression phenotypes (Richardson et al., 2011; Zhang, 2012).

Using different programmes and databases, we obtained different quantity of lung cancer-related inflammatory genes that have miRNA-binding sites. It is difficult to judge which SNPs or miRNAs are likely to play more roles in lung cancer development without experiments being done, we combined results from eight databases to increase the accuracy of the analysis. In recent years, more and more databases have been used to explore the SNPs in miRNA-binding sites, which including miRanda, PicTar, MirSNP, TargetScan Human, miRNASNP, Patrocles, DIANA-microT and PolymiRTS Database. The more databases predicted the SNP, the more likely it would be the true target SNP (Song et al., 2014).

Currently, there were numbers of studies focus on the association between SNPs in miRNA binding-sites of specific gene and diseases some about susceptibility and others about prognosis. Now genome wide association studies (GWA study, or GWAS) reported scores of diseases related SNPs were in non-coding region. The significance of the association may be brought up by still unknown mechanisms or by linkage disequilibrium (LD)

with functional polymorphisms. Thus, the regulation of miRNAs on target genes may work. Some researchers (Richardson et al., 2011) basing on the list of SNPs from GWAS, or in strong LD with a GWAS SNP performed a genome-wide scan of SNPs that abrogate or create miRNA recognition element (MRE) seed sites (MRESS) and identified high priority candidate SNPs for functional studies and for disease risk prediction. Other researchers (Wu et al., 2011) discovered the relationship between specific genes expression or miRNA level and diseases, so they thought that SNPs within miRNA target sites may influence their encoded target-mRNAs and their downstream effectors and they would predict the 3'UTR of these genes contains potential MREs and verify that these mutations maybe as a key in regulating gene expression. At present, all studies the on gene variation in miRNA-binding regions were performing several kinds of arithmetic or getting help from various websites. But there was no uniform standard and procedures in selecting SNPs, the results of predicting not only need evaluate in the association study also confirm by the functional research. So we will do the further investigation to certificate these SNPs.

In our results, we found some SNPs were reported have association with cancer or other disease which may impact function of genes by combing different miRNAs because of changing sequences. This paper provided the basis for a reasoned algorithm-driven selection of SNPs. It is important to address that all the polymorphisms predicted supports future investigations to validate these results in well-characterised populations by functional assays or case-control association studies. The proposed approach could help to ease the identification of functionally relevant SNPs and minimize the workflow and the costs.

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References

- Betel D, Koppal A, Agius P, et al (2010). Comprehensive modeling of microRNA targets predicts functional nonconserved and non-canonical sites. *Genome Biol*, **11**, 90.
- Bhat IA, Pandith AA, Bhat BA, et al (2013). Lack of association of a common polymorphism in the 3'-UTR of interleukin 8 with non small cell lung cancer in Kashmir. *Asian Pac J Cancer Prev*, **14**, 4403-8.
- Chen Z, Xu L, Ye X, et al (2013). Polymorphisms of microRNA sequences or binding sites and lung cancer: a meta-analysis and systematic review. *PLoS One*, **8**, 61008.
- Cheng M, Yang L, Yang R, et al (2013). A microRNA-135a/b binding polymorphism in CD133 confers decreased risk and favorable prognosis of lung cancer in Chinese by reducing CD133 expression. *Carcinogenesis*, **34**, 2292-9.

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- Cho WC, Kwan CK, Yau S, et al (2011). The role of inflammation in the pathogenesis of lung cancer. *Expert Opin Ther Targets*, **15**, 1127-37.
- Coussens LM, Werb Z (2002). Inflammation and cancer. *Nature*, **420**, 860-7.
- Ding J, Gao Y, He Y, et al (2011). Screening SNPs residing in the microRNA-binding sites of hepatocellular carcinoma related genes. *Int J Data Min Bioinform*, **5**, 1-21.
- Grivennikov SI, Greten FR, Karin M (2010). Immunity, inflammation, and cancer. *Cell*, **140**, 883-99.
- Hattar K, Savai R, Subtil FS, et al (2013). Endotoxin induces proliferation of NSCLC *in vitro* and *in vivo*: role of COX-2 and EGFR activation. *Cancer Immunol Immunothe*, **62**, 309-20.
- Hariharan M, Scaria V, Brahmachari SK. dbSMR: a novel resource of genome-wide SNPs affecting microRNA mediated regulation. *BMC Bioinformatics*, **10**, 108.
- Houghton AM, Mouded M, Shapiro SD (2008). Common origins of lung cancer and COPD. *Nat Med*, 14, 1023-4.
- Hussain SP, Harris CC (2007). Inflammation and cancer: an ancient link with novel potentials. *Int J Cancer*, **121**, 2373-80.
- Jemal A, Bray F, Center MM, et al (2011). Global cancer statistics. *CA Cancer J Clin*, **61**, 69-90.
- John B, Enright AJ, Aravin A, et al (2004). Human MicroRNA targets. *PLoS Biol*, 2, 363.
- Kim M, Chen X, Chin LJ, et al (2014). Extensive sequence variation in the 3' untranslated region of the KRAS gene in lung and ovarian cancer cases. *Cell Cycle*, **13**.
- Lagos-Quintana M, Rauhut R, Lendeckel W, et al (2001). Identification of novel genes coding for small expressed RNAs. *Science*, **294**, 853-8.
- Lee G, Walser TC, Dubinett SM (2009). Chronic inflammation, chronic obstructive pulmonary disease, and lung cancer. *Curr Opin Pulm Med*, **15**, 303-7.
- Lee J, Taneja V, Vassallo R (2012). Cigarette smoking and inflammation: cellular and molecular mechanisms. *J Dent Res*, **91**, 142-9.
- Liu C, Zhang F, Li T, et al (2012). MirSNP, a database of polymorphisms altering miRNA target sites, identifies miRNA-related SNPs in GWAS SNPs and eQTLs. BMC Genomics, 13, 661.
- McMillan DH, Baglole CJ, Thatcher TH, et al (2011). Lungtargeted overexpression of the NF-kappaB member RelB inhibits cigarette smoke-induced inflammation. *Am J Pathol*, **179**, 125-33.
- Mishra PJ1, Bertino JR (2009). MicroRNA polymorphisms: the future of pharmacogenomics, molecular epidemiology and individualized medicine. *Pharmacogenomics*, **10**, 399-416.
- Ohnishi S, Ma N, Thanan R, et al (2013). DNA damage in inflammation-related carcinogenesis and cancer stem cells. *Oxid Med Cell Longev*, 387014.
- Richardson K, Lai CQ, Parnell LD, et al (2011). A genome-wide survey for SNPs altering microRNA seed sites identifies functional candidates in GWAS. *BMC Genomics*, **12**, 504.
- Saunders MA, Liang H, Li WH (2007). Human polymorphism at microRNAs and microRNA target sites. *Proc Natl Acad Sci USA*, 104, 3300-5.
- Shen N, Gong T, Wang JD, et al (2011). Cigarette smoke-induced pulmonary inflammatory responses are mediated by EGR-1/ GGPPS/MAPK signaling. *Am J Pathol*, **178**, 110-8.
- Song CQ, Zhang JH, Shi JC, et al (2014). Bioinformatic prediction of SNPs within miRNA binding sites of inflammatory genes associated with gastric cancer. *Asian Pac J Cancer Pre*, **15**, 937-43.
- Tomari Y, Zamore PD (2005). Perspective: machines for RNAi. *Genes Dev*, **19**, 517-29.

- Wu Y, Xiao Y, Ding X, et al (2011). A miR-200b/200c/429binding site polymorphism in the 3' untranslated region of the AP-2alpha gene is associated with cisplatin resistance. *PLoS ONE*, 6, 29043.
- Yu H, Zhao H, Wang LE, et al (2011). An analysis of single nucleotide polymorphisms of 125 DNA repair genes in the Texas genome-wide association study of lung cancer with a replication for the XRCC4 SNPs. DNA Repair, 10, 398-407.
- Zhang W, Edwards A, Zhu D, et al (2012). miRNA-mediated relationships between Cis-SNP genotypes and transcript intensities in lymphocyte cell lines. *PLoS One*, 7, 31429.
- Zu Y, Ban J, Xia Z, et al (2013). Genetic variation in a miR-335 binding site in BIRC5 alters susceptibility to lung cancer in Chinese Han populations. *Biochem Biophys Res Commun*, **430**, 529-34.