Incorporation of a Genetic Factor into an Epidemiologic Model for Prediction of Individual Risk of Lung Cancer: The Liverpool Lung Project

Olaide Y. Raji1, Olorunsola F. Agbaje2, Stephen W. Duffy3, Adrian Cassidy1, and John K. Field1

Abstract

The Liverpool Lung Project (LLP) has previously developed a risk model for prediction of 5-year absolute risk of lung cancer based on five epidemiologic risk factors. SEZ6L, a Met430lle polymorphic variant found on 22q12.2 region, has been previously linked with an increased risk of lung cancer in a case-control population. In this article, we quantify the improvement in risk prediction with addition of SEZ6L to the LLP risk model. Data from 388 LLP subjects genotyped for SEZ6L single-nucleotide polymorphism (SNP) were combined with epidemiologic risk factors. Multivariable conditional logistic regression was used to predict 5-year absolute risk of lung cancer with and without this SNP. The improvement in the model associated with the SEZ6L SNP was assessed through pairwise comparison of the area under the receiver operating characteristic curve and the net reclassification improvements (NRI). The extended model showed better calibration compared with the baseline model. There was a statistically significant modest increase in the area under the receiver operating characteristic curve when SEZ6L was added into the baseline model. The NRI also revealed a statistically significant improvement of around 12% for the extended model; this improvement was better for subjects classified into the two intermediate-risk categories by the baseline model (NRI, 27%). Our results suggest that the addition of SEZ6L improved the performance of the LLP risk model, particularly for subjects whose initial absolute risks were unable to discriminate into “low-risk” or “high-risk” group. This work shows an approach to incorporate genetic biomarkers in risk models for predicting an individual’s lung cancer risk.

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Introduction

Interest in methods of assessing individual risk of developing diseases has continued to grow over the years, partly due to their usefulness in selection of high-risk individuals that would benefit in prevention or screening programs (1–3). Within the Liverpool Lung Project (LLP), we have previously developed and validated a predictive model for 5-year absolute risk of developing lung cancer for an individual with a specific combination of epidemiologic risk factors including smoking duration, previous diagnosis of pneumonia, prior diagnosis of malignant tumor, occupational exposure to asbestos, and family history of lung cancer (4). Recent advancement in genetic epidemiology leading to identification of genetic and molecular variants affecting the risk of disease means that genetic markers such as single-nucleotide polymorphisms (SNP) can be added to risk models for improved prediction of future risk of disease (5–7).

DNA pooling and high-throughput sequencing recently conducted as part of the Sequenom-Genefinder project has identified a candidate region in chromosome 22q12.2 that contained SNPs showing significant differences between lung cancer case-control subjects (8). This region overlies the seizure 6-like (SEZ6L) gene, with polymorphic Met430lle (SNP rs663048) identified in the region as a top candidate variant modulating the risk of lung cancer. The SEZ6L marker SNP was further validated in individual genotyping in two independent data sets: the LLP and the M.D. Anderson Cancer Study populations (8). The two studies independently found a significant effect of the variant with a combined 3-fold risk of lung cancer for the homozygotes mutant allele compared with the wild-type.

This study incorporates the SEZ6L SNP into the LLP risk model for predicting 5-year absolute risk of developing
lung cancer and assesses the predictive ability and improved accuracy of the extended model.

**Patients and Methods**

Data collected as part of the LLP case-control study were used in this study. The detailed LLP recruitment procedure and study protocol have been previously described elsewhere (4, 9). Briefly, incident cases of histologically or cytologically confirmed lung cancer, ages between 20 and 80 years, were included. Lung cancer included any of topographical subcategories of code C34 of the International Classification of Disease for Oncology 9th revision. Two population controls per case, matched on year of birth (±2 years) and gender, were selected from registers of general practitioners in Liverpool area. All participants were resident in the Liverpool area and provided written informed consent.

The current analysis is based on 200 histologically and cytologically confirmed lung cancer cases and 188 age-(±2 years) and sex frequency-matched controls. This group of subjects were among those used for developing the original LLP model and were genotyped as part of the validation study. The genotype was determined by extracting the genomic DNA from blood peripheral leukocytes using the Qiagen DNA Blood Mini Kit according to the manufacturer's instructions. Details of the genotyping process are provided elsewhere (8).

### SNP marker and genotyping

The Sequenom-Genefinder pilot study aimed to identify polymorphic susceptibility variants for lung cancer and genotyped and tested 83,715 SNP, located primarily in gene-based regions. A subset of the LLP case-control study subjects were genotyped as part of the validation study. The genotype was determined by extracting the genomic DNA from blood peripheral leukocytes using the Qiagen DNA Blood Mini Kit according to the manufacturer's instructions. Details of the genotyping process are provided elsewhere (8).

### Statistical analysis

Pearson's χ² test was used to assess the relationship between the SNP marker and each of the epidemiologic risk factors included in the previously published LLP risk model. This test of association was done for all subjects together and for cases and controls separately.

The risk data were analyzed by conditional logistic regression. We refitted the original risk model for subjects with both genetic and epidemiologic data (n = 388) and compared the estimated coefficients to that of the original model (n = 1,275). We then fitted the model, including...
the SEZ6L SNP, to estimate the effect of the latter, adjusting for the SNP.

The previously published multivariable conditional logistic regression model was used to generate estimates of predicted 5-year absolute risk of lung cancer with and without genotype information. The baseline risk ($\alpha$—the constant term in the regression model) for the prediction of 5-year absolute risk using the model with genetic information was recalculated. The procedure for calculating the baseline $\alpha$ from age- and sex-specific lung cancer incidence rates for the Liverpool area has earlier been described (4); the only difference is that the linear combination of the $\beta$ coefficients in the probability model now includes information on the SNP genotype.

The area under the receiver-operating characteristic curve was calculated as a measure of discriminatory ability of the models and to formally compare the models with and without the SEZ6L SNP (10). We also calculated the net reclassification improvement (NRI) to quantify the improvement in the LLP risk model containing the SEZ6L. NRI cross-classifies subjects’ predicted risks from the original and extended models, assesses the proportions of subjects reclassified into new risk categories (case and control separately), and quantifies the correct movement in categories—upward for cases and downward for controls (11). Simple measures such as the true positive fraction (sensitivity) and false positive fraction (1 − specificity) were computed for the risk thresholds classifying the subjects into high-, intermediate-, and low-risk groups. Improvement in model calibration was also assessed by comparing the closeness of the predicted risks to the observed risks (goodness of fit) for each model using the Hosmer-Lemeshow’s $\chi^2$ statistic (12, 13) and the Akaike information criteria (14).

**Results**

**Subjects’ characteristics**

The distribution of the five epidemiologic and lifestyle characteristics of the 388 case-control subjects used in this study by SEZ6L genotype is presented in Table 1. The majority of subjects (55%) had SEZ6L genotype GG, 38% had heterozygote genotype TG, and approximately 7% had homozygote mutant genotype TT. The distribution by case-control status revealed a statistically significant increase in the proportion of cases with homozygote mutant genotype (10%) compared with controls (3%). There was no statistically significant relationship between any of the five epidemiologic factors and the SNP for either the combined or the separate (data not shown) analysis of the case and control subjects.

**Multivariable risk model**

Table 2 gives the odds ratios (OR) and 95% confidence intervals (95% CI) for the multivariate conditional logistic regression using different subsets of LLP subjects. The

<table>
<thead>
<tr>
<th>Variables</th>
<th>LLP subjects with both EPI variables and SEZ6L SNP (n = 388)</th>
<th>Original LLP risk model using LLP case-control subjects (n = 1,275)</th>
<th>Model with SEZ6L only using all subjects genotyped (n = 463)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Model with EPI variables only</td>
<td>Model with EPI + SEZ6L variables</td>
<td></td>
</tr>
<tr>
<td>Smoking duration (y)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
<td>OR (95% CI)</td>
</tr>
<tr>
<td>1-19</td>
<td>1.84 (0.79-4.24)</td>
<td>1.69 (0.72-3.94)</td>
<td>2.16 (1.22-3.82)</td>
</tr>
<tr>
<td>20-39</td>
<td>5.32 (2.48-11.41)</td>
<td>5.08 (2.34-11.01)</td>
<td>4.28 (2.63-6.96)</td>
</tr>
<tr>
<td>Pneumonia (Yes)</td>
<td>1.25 (0.68-2.31)</td>
<td>1.24 (0.67-2.33)</td>
<td>1.82 (1.26-2.63)</td>
</tr>
<tr>
<td>Asbestos exposure (Yes)</td>
<td>2.35 (1.35-4.08)</td>
<td>2.32 (1.32-4.06)</td>
<td>1.89 (1.35-2.63)</td>
</tr>
<tr>
<td>Previous tumor (Yes)</td>
<td>1.62 (0.61-4.32)</td>
<td>1.58 (0.58-4.29)</td>
<td>1.97 (1.23-3.15)</td>
</tr>
<tr>
<td>Family history of cancer</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early onset (&lt;60 y)</td>
<td>2.16 (0.82-5.67)</td>
<td>2.34 (0.88-6.20)</td>
<td>2.02 (1.18-3.46)</td>
</tr>
<tr>
<td>Late onset (&gt;60 y)</td>
<td>0.87 (0.46-1.67)</td>
<td>0.93 (0.49-1.79)</td>
<td>1.19 (0.80-1.78)</td>
</tr>
<tr>
<td>SEZ6L genotype marker</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TG</td>
<td>1.17 (0.72-1.92)</td>
<td></td>
<td>1.19 (0.78-1.82)</td>
</tr>
<tr>
<td>TT</td>
<td>4.51 (1.56-13.02)</td>
<td></td>
<td>3.79 (1.45-9.86)</td>
</tr>
<tr>
<td>Goodness-of-fit statistic</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hosmer-Lemeshow (P)</td>
<td>6.24 (0.62)</td>
<td>5.54 (0.70)</td>
<td></td>
</tr>
<tr>
<td>AIC</td>
<td>418.12</td>
<td>413.38</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: EPI, epidemiological; AIC, Akaike information criteria.
estimated ORs and 95% CIs for models with and without SEZ6L seemed comparable, suggesting no serious confounding effect of SEZ6L on the relationship between each of the original epidemiologic risk factors and lung cancer risk. The parameter estimates for the model with reduced subjects were similar to that of the original LLP risk model, but with lesser precision. We therefore retained coefficients of the original model and incorporated the adjusted estimate for the SEZ6L SNP for the expanded model, but with the α’s recalculated as previously mentioned above.

**Predictive ability of the expanded risk model**

There was an improvement in the calibration when SEZ6L was incorporated into the risk model, as shown by the reduced Hosmer-Lemeshow χ² statistic of 5.54 (P = 0.70) compared with the 6.24 (P = 0.62) observed for the model without SEZ6L (Table 2). This finding was supported by the reduction in the Akaike information criteria value from 418.12 for the model without SEZ6L to 413.38 for the model with SEZ6L, suggesting improved fit for the extended model.

Figure 1 shows the receiver operating characteristic curve for the models with and without the SEZ6L SNP. A significant (P = 0.01) increase of about 4% in area under the receiver-operating characteristic curve was observed (from 0.72 to 0.75) for the model extended with SEZ6L SNP.

The sensitivity and specificity corresponding to the three different threshold classifying the subjects into low-risk (<0.91), intermediate-risk (0.91 to <5.12), and high-risk (>5.12) groups are shown in Table 3. The threshold values were defined from the predicted 5-year absolute risks for the original LLP control samples (n = 1,272), assuming the risk distribution in this group to be similar to that of the general Liverpool population. The upper threshold (5.12) corresponds to the value for the top 20% of predicted absolute risks in the population; individuals whose 5-year predicted absolute risk is or above this value are designated as “high-risk” group. The lower threshold value of 0.91 corresponds to the bottom 40% of absolute risks in the control population and demarcates the “low-risk” group. This definition of high-risk and low-risk groups was used in a cardiovascular diseases study (15).

**Table 3. Sensitivity and specificity for models with and without SEZ6L at specified risk thresholds**

<table>
<thead>
<tr>
<th>Risk threshold</th>
<th>LLP risk model using all subjects (n = 1,275)</th>
<th>Refitted LLP model using subject with EPI + SNP variables (n = 388)</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>se (Sp) (%)</td>
<td>No SEZ6L</td>
<td>With SEZ6L</td>
<td>No SEZ6L</td>
</tr>
<tr>
<td>0.91</td>
<td>86.1 (39.5)</td>
<td>83.0</td>
<td>84.0</td>
<td>43.1</td>
</tr>
<tr>
<td>2.50</td>
<td>67.8 (64.2)</td>
<td>65.0</td>
<td>67.0</td>
<td>67.6</td>
</tr>
<tr>
<td>5.12</td>
<td>49.9 (79.8)</td>
<td>50.5</td>
<td>50.0</td>
<td>83.5</td>
</tr>
</tbody>
</table>

Abbreviations: Se, sensitivity; Sp, specificity.
observed values for the two measures (sensitivity and specificity) were comparable for the two models in the two extreme groups (low risk and high risk); this means that knowledge of the SNP genotype had little or no effect for subjects already classified to be of low or high risk. In contrast, the model with SEZ6L seemed to discriminate better in the intermediate group and is particularly more specific among this group of subjects.

Table 4 gives the joint distribution of the classified predicted risks from each model. Overall, approximately one quarter of cases (49 of 200) and about 20% of controls (37 of 188) had their predicted risks reclassified into other risk groups when SEZ6L was incorporated into the risk predictive model. This reclassification showed improvement (upward shift) in approximately 14% of cases and became worse (downward shift) for 11%, resulting in a net gain of about 3%. The net gain was higher for controls (9%) with overall improvement in risks (downward shift) for 14% and worst performance (upward shift) for only 5% of controls. These figures thus correspond to a statistically significant NRI of approximately 12% ($P = 0.03$). Concentrating only on those subjects whose initial risks are in the two intermediate groups (cases, 66; controls, 75), 20 of 30 reclassified cases and 15 of 21 reclassified controls had improved risks, resulting in a better NRI of 27%.

**Discussion**

We have evaluated the contribution of the SEZ6L gene, a genetic marker variant identified at 22q12.2 region, to prediction of individual absolute risk of developing lung cancer within a 5-year period using the recently developed LLP risk model. Our results suggest a modest increase in the overall predictive ability of the model with the SEZ6L SNP genotype. A greater effect was confined to subjects whose initial risks from the baseline model were in the medium-risk category. For these subjects, a NRI of around 27% was recorded, demonstrating the usefulness of the SNP in risk prediction.

The results presented in this article are in line with recent suggestions on taking predictive risk models of lung cancer to the next level where mediating genetic biological markers are included to allow a more accurate prediction of interindividual lung cancer risks. This expansion of risk factors in predictive risk models beyond the traditional epidemiologic data has been elegantly pursued in cardiovascular diseases, wherein biological assay data and some genetic variants had been added into risk score models for prediction of absolute risk of cardiovascular disease (15, 16). Similar efforts in cancer studies include addition of mammographic density data to the Gail model for breast cancer (17), inclusion of assay data for DNA repair capacity and mutagen sensitivity into the Spitz lung cancer risk models for former and current smokers (18), and recent combination of a panel of low-risk SNPs with demographic data to form a simple algorithm for lung cancer risk prediction (7).

It is known that a single gene would normally convey a small risk and confer a small cumulative addition to prediction (19). However, the combined effect of a number of genetic risk factors may be substantial, and indeed, the addition of this single SNP added more to the area under the curve than, for example, history of a previous non-lung malignancy, one of the LLP risk model parameters. The internal improvement in accuracy is encouraging, but the next step is to apply the combined LLP including the SEZ6L SNP on independent data sets for external validation.

We noted that the SEZ6L SNP variant (rs663048) was not among the top-ranked genes reported in a genome-wide association study identified as lung cancer susceptibility genes (20), nor was it statistically significant in a recent meta-analysis of lung genome-wide association study SNPs (21). However, biological evidence supports the importance of the SEZ6L gene in lung cancer oncogenesis. Frequent allelic losses in non–small-cell lung carcinomas have been reported on chromosomes 22q, a region where the SEZ6L gene resides, indicating the presence of tumor
suppressor gene(s) at the location (22). Also, the SEZ6L gene is a Met430Ile amino acid substitution that has been predicted to be functional by both SIFT and PolyPhen, suggesting a protein disturbing functional role (8). SEZ6L has also been found to be highly hypermethylated in primary colorectal tumor (23) and gastric carcinoma (24), linking the mutation to the progression of other neoplasms. Also, it should be noted that we observed the effect in three independent studies, two case-control studies with individual genotyping, and one DNA pooling study.

Risk prediction models may potentially play a significant role in future control of lung cancer, as prevention and screening could be targeted at those at increased risk of developing the disease thereby improving diagnosis, treatment, and survival (2). In addition, the usefulness of risk prediction models in the design of public screening program or clinical trial for diagnostic or treatment procedure has recently been shown (25). These necessitate accurate prediction of risk. Recent development in molecular and genetic epidemiology has thus encouraged the inclusion of validated susceptibility genes into existing risk models to improve their predictive ability. Apart from accurate prediction of absolute risks, the expanded risk models may be reserved for individuals with intermediate risk, where the decision for classification into high or low risk group is ambiguous.

Our results show in principle how a genetic risk marker can be incorporated into an epidemiologic risk prediction model. In the example here, inclusion of the SEZ6L SNP shows a significant improvement in risk prediction. This work provides “proof of concept” in taking lung cancer risk prediction models to the next level.

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