Modulation of Cystatin A Expression in Human Airway Epithelium Related to Genotype, Smoking, COPD, and Lung Cancer

Marcus W. Butler¹, Tomoya Fukui¹, Jacqueline Salit¹, Renat Shaykhiev¹, Jason G. Mezey¹⁄², Neil R. Hackett¹, and Ronald G. Crystal¹

Abstract

The cathepsin inhibitor Cystatin A (CSTA) has antiapoptotic properties linked with neoplastic changes in squamous cell epithelium, where it has been proposed as a diagnostic and prognostic marker of lung cancer. Notably, cystatin A is upregulated in dysplastic epithelium, prompting us to hypothesize that it might be modulated in chronic obstructive pulmonary disease (COPD), a small airway epithelial (SAE) disorder that is a risk factor for non–small cell lung cancer (NSCLC) in a subset of smokers. Here we report that genetic variation, smoking, and COPD can all elevate levels of CSTA expression in lung small airway epithelia, with still further upregulation in squamous cell carcinoma (SCC), an NSCLC subtype. We examined SAE gene expression in 178 individuals, including healthy nonsmokers (n = 60), healthy smokers (n = 82), and COPD smokers (n = 36), with corresponding large airway epithelium (LAE) data included in a subset of subjects (n = 52). Blood DNA was genotyped by SNP microarray. Twelve SNPs upstream of the CSTA gene were found to associate with its expression in SAE. Levels were higher in COPD smokers than in healthy smokers, who, in turn, had higher levels than nonsmokers. CSTA gene expression in LAE was also smoking-responsive. Using publicly available NSCLC expression data we also found that CSTA was upregulated in SCC versus LAE and downregulated in adenocarcinoma versus smoke-exposed SAE. All phenotypes were associated with different proportional expression of CSTA to cathepsins. Our findings establish that genetic variability, smoking, and COPD all influence CSTA expression, as does SCC, supporting the concept that CSTA may make pivotal contributions to NSCLC pathogenesis in both early and late stages of disease development. Cancer Res; 71(7): 1–10. ©2011 AACR.

Introduction

Cystatin A (CSTA; also referred to as stefin A, and acid cysteine proteinase inhibitor), a member of the class I or stefin subgroup of the 3 described cystatin families, is a 11 kDa single chain intracellular cysteine proteinase inhibitor capable of inhibiting papain and cathepsins B, H, and L, as well as the cysteine proteinase activity of the major house dust mite allergen Der p 1 (1–3). CSTA has antiapoptotic properties (4–6) and cystatin A expression has been linked with neoplastic changes in squamous cell epithelium with cystatin A levels paralleling the response to antitumor therapy (7–11). As with other members of the stefin family, cystatin A has been proposed as a marker and prognostic indicator of lung cancer (11, 12). In the lung, cystatin A expression is upregulated in dysplastic epithelium, and highly expressed in many lung cancers, especially squamous cell carcinoma (SCC; refs 7, 10, 12).

With this background, and based on the knowledge that the majority of lung cancers are derived from the small airway epithelium (SAE) of cigarette smokers (13–15), but that only a subset of smokers develop lung cancer (16, 17), we hypothesized that the expression of CSTA in the SAE is modulated by genetic variation and upregulated by smoking. Further, in the context that chronic obstructive pulmonary disease (COPD), a disease that starts in the small airways (13, 18–21), is a risk factor for lung cancer independent of smoking (22–26), we asked whether CSTA is upregulated in the SAE of smokers with COPD beyond that of smokers per se, and whether this upregulation is also dependent on genetic variation. In view of the fact that both smoking and COPD are stronger risk factors for SCC than for adenocarcinoma of the lung, and that most SCCs are derived from large airways (15, 27–30), we also examined CSTA expression in large airway epithelium (LAE) samples obtained in a subset of the study subjects, and compared our CSTA gene expression data to publicly available lung cancer microarray...
data derived using comparable methodologies (31), to see if CSTA is discordantly expressed in SCC and adenocarcinoma of the lung, relative to the SAE and LAE.

Materials and Methods

Study subjects

Healthy nonsmokers, healthy smokers, and smokers with COPD were assessed in the Weill Cornell National Institutes of Health Clinical and Translational Sciences Center and Department of Genetic Medicine Clinical Research Facility using protocols approved by the Weill Cornell Medical College Institutional Review Board. Prior to inclusion in the study, each individual provided written informed consent. No abnormality was found in the nonsmokers (n = 60) or the healthy smokers (n = 82) following a standardized screening evaluation composed of a history, physical examination, complete blood count, serum chemistries, coagulation profile, liver function tests, urine studies, chest X-ray, EKG, and full lung function studies. The COPD smokers (n = 36) were diagnosed in accordance with standard GOLD criteria (32). Blood was collected for genotyping on a random subset of individuals (nonsmokers, n = 44; healthy smokers, n = 48; COPD smokers, n = 20). Fewer subjects with genotyped samples were available than subjects with SAE gene expression data because of a significantly longer temporal interest of our laboratory in the study of airway epithelium gene expression than in genomic studies. LAE samples were not given as high a priority as SAE samples when individuals were bronchoscoped, reflecting the additional probe sets 200838_at (cathepsin H), 202295_s_at (cathepsin L), and 202087_s_at (cathepsin B), with the same platform that was used on the SAE and LAE samples. The proportional relationship among CSTA and cathepsin gene expression was examined in all samples using the additional probe set 204971_at on the Affymetrix HG-U133 Plus 2.0 chip, the same platform that was used on the SAE and LAE samples. All microarray data have been deposited at the Gene Expression Omnibus (GEO) site (ref. 35; accession number GSE22047). Genomic data from the Affymetrix Human SNP 5.0 were assessed using the BRLMM-P Analysis Tool (BAT) 2.0 software (Affymetrix) to determine genotype for cis-SNPs in the vicinity of CSTA on chromosome 3. Associations between CSTA gene expression levels and genotype for the 48 SNPs located within 100 kb of the gene were performed using PLINK, with permutation testing used to control for the effect of genetic ancestry. Haplotype associations with expression were performed using PLINK and Haploview, with linear regression modeling (see the Supplementary Materials and Methods section).

Collection of airway epithelium and assessment of gene expression

Fiberoptic bronchoscopy was used to obtain small (and for a random subset, large) airway epithelial cells as previously described (33, 34). Further details of the airway epithelial sampling, preparation of cDNA, hybridization of labeled cRNA to Affymetrix HG-U133 Plus 2.0 arrays and confirmation by TaqMan (Applied Biosystems) real-time RT-PCR (reverse transcriptase PCR) can be found in the Supplementary Materials and Methods section. No cell lines were used in the conduct of this research; all samples used were freshly collected and immediately processed for RNA or protein.

Correlation of genotypes and haplotypes with gene expression

Genomic DNA was extracted from stored blood samples using the Autogen FX robotic system in accordance with Autogen’s protocols (Autogen). Preprinted bar-coded labels were affixed to sample containers to minimize sample mix-ups, and critical steps in sample processing were only undertaken when 2 technicians were present. The Affymetrix Human SNP Array 5.0 platform was used to assess genotype using the manufacturer’s protocols. The focus was on all 48 SNPs found on the SNP array whose chromosomal location was within 100,000 base pairs either side of the CSTA gene. To avoid artifactual association of genotype with expression caused by SNPs with known minor allele frequency more than 5% situated within the target sequence of Affymetrix expression probe sets, the sequences of individual probes (obtained from NetAffx, Affymetrix) were cross-checked against the National Center for Biotechnology Information (NCBI) dbSNP build 129. Potential effects of copy number variation on associations of genotype with gene expression were assessed as outlined in the Supplementary Materials and Methods section.

Data analysis and statistics for SAE expression and genomic data

Gene expression analyses on all samples were performed using the Microarray Suite 5.0 (MAS5) software (Affymetrix). Publicly available data from Kuner and colleagues (31) for 40 subjects with lung adenocarcinoma and 18 subjects with SCC of the lung were downloaded from the NCBI Web site (accession number GSE10245), using the gene expression data from the only CSTA probe set 204971_at on the Affymetrix HG-U133 Plus 2.0 chip, the same platform that was used on the SAE and LAE samples. The proportional relationship among CSTA and cathepsin gene expression was examined in all samples using the additional probe sets 200838_at (cathepsin B), 202295_s_at (cathepsin H), and 202087_s_at (cathepsin L). Gene expression data were normalized using GeneSpring version 7.2 software (Agilent Technologies) per chip and per gene across all samples. All microarray data have been deposited at the Gene Expression Omnibus (GEO) site (ref. 35; accession number GSE22047). Genomic data from the Affymetrix Human SNP 5.0 were assessed using the BRLMM-P Analysis Tool (BAT) 2.0 software (Affymetrix) to determine genotype for cis-SNPs in the vicinity of CSTA on chromosome 3. Associations between CSTA gene expression levels and genotype for the 48 SNPs located within 100 kb of the gene were performed using PLINK, with permutation testing used to control for the effect of genetic ancestry. Haplotype associations with expression were performed using PLINK and Haploview, with linear regression modeling (see the Supplementary Materials and Methods section).

Results

Association of CSTA small airway epithelial gene expression with genomic variation

The study population demographic findings and characteristics of brushed SAE samples and microarray expression probe performance are detailed in the Supplementary Results section. Based on the hypothesis that genetic variability might modulate the level of CSTA expression in the SAE, we evaluated the correlation of CSTA gene expression with genotypes of cis-SNPs within 100 kbp either side of the CSTA gene using
Nine of the 12 SNPs identified had a stronger association with the observed effect. For example, rs16832956, the SNP with the strongest genetic association, had low CSTA expression, with a >1.4-fold increase seen in CC genotypes (Fig. 1B, P < 1.6 × 10^{-4}). The SAE CSTA gene expression levels were confirmed by TaqMan RT-PCR (Fig. 1C, r² = 0.67, P < 10^{-4}).

**Table 1. Association of CSTA genotypes with CSTA gene expression in the small airway epithelium**

<table>
<thead>
<tr>
<th>SNP identity</th>
<th>Chr. 3 location</th>
<th>Minor allele frequency</th>
<th>All subjects (n = 112)</th>
<th>Healthy nonsmokers (n = 44)</th>
<th>Healthy smokers (n = 48)</th>
<th>COPD smokers (n = 20)</th>
</tr>
</thead>
<tbody>
<tr>
<td>rs1354162</td>
<td>123,436,767</td>
<td>0.040</td>
<td>0.02</td>
<td>NS</td>
<td>NS</td>
<td>4.6 × 10^{-3}</td>
</tr>
<tr>
<td>rs7652858</td>
<td>123,480,795</td>
<td>0.125</td>
<td>3.6 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs2134225</td>
<td>123,484,943</td>
<td>0.125</td>
<td>3.6 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs16832956</td>
<td>123,500,198</td>
<td>0.161</td>
<td>4.5 × 10^{-4}</td>
<td>NS</td>
<td>3.4 × 10^{-3}</td>
<td>0.010</td>
</tr>
<tr>
<td>rs1402200</td>
<td>123,505,107</td>
<td>0.411</td>
<td>7.3 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs5008830</td>
<td>123,513,152</td>
<td>0.089</td>
<td>1.0 × 10^{-3}</td>
<td>0.025</td>
<td>1.1 × 10^{-5}</td>
<td>0.038</td>
</tr>
<tr>
<td>rs2001548</td>
<td>123,515,479</td>
<td>0.101</td>
<td>1.7 × 10^{-3}</td>
<td>NS</td>
<td>3.4 × 10^{-5}</td>
<td>0.038</td>
</tr>
<tr>
<td>rs4678180</td>
<td>123,520,487</td>
<td>0.455</td>
<td>5.2 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs9864290</td>
<td>123,522,752</td>
<td>0.451</td>
<td>7.8 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs6803098</td>
<td>123,523,300</td>
<td>0.453</td>
<td>4.1 × 10^{-3}</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>rs9817571</td>
<td>123,538,670</td>
<td>0.099</td>
<td>0.03</td>
<td>NS</td>
<td>0.018</td>
<td>NS</td>
</tr>
<tr>
<td>rs9842752</td>
<td>123,539,955</td>
<td>0.076</td>
<td>0.03</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
</tbody>
</table>

Note: CSTA gene expression levels in small airway epithelium as assessed by Affymetrix HGU133 Plus 2.0 microarray expression probe set 204971_at were correlated with Affymetrix Human SNP Array 5.0 cis-SNPs situated within 100 kbp of the CSTA gene location on chromosome 3q21.

*For analysis with the group of all subjects, P values represent Wald statistic following permutation analysis: after initial associations were identified, each association was further tested by performing 10^3 permutations within clusters of similar genetic ancestry, as defined by STRUCTURE, using PLINK software. For the subgroup analyses, t tests are shown.

Only those SNPs with P values for association of genotype with expression of <0.05 are shown. SNPs are listed in ascending order of location on chromosome 3. 5' to 3', the first 4 SNPs listed are within introns of the upstream CASR gene, the next 6 are intergenic, and the last 2 listed SNPs are located within introns of CSTA.

Chr, chromosome. Locations refer to NCBI Human Genome Reference Genome Assembly, Build 36.3

Minor allele frequencies observed in the study population.

All subjects with SAE gene expression data and genotype data. Healthy nonsmokers (n = 44), healthy smokers (n = 48), and COPD smokers (n = 20) combined. NS = not significant (P > 0.05).

Paired airway gene expression data and blood DNA SNP data from the subset of the study population for whom both gene expression and SNP data were available (healthy nonsmokers, n = 44; healthy smokers, n = 48; and COPD smokers, n = 20).

Ten SNPs located upstream of the CSTA gene, and 2 more within introns of the CSTA gene, displayed significant correlation of genotype with SAE expression levels of CSTA (Table 1, Fig. 1A). For the total combined group of subjects (healthy nonsmokers, healthy smokers and smokers with COPD), the most significant correlation of the levels of CSTA small airway gene expression was with the SNP rs16832956 (P < 5 × 10^{-4}).

Nine of the 12 SNPs identified had P < 10^{-2} in reference to the strength of their association with CSTA SAE gene expression. The 4 SNPs most upstream of CSTA were located within introns of the calcium-sensing receptor (CASR) gene, but none had any relationship of genotype with CASR gene expression levels (P > 0.8, not shown).

These associations persisted at this significance level after assessing 10^3 permutations within clusters of similar genetic ancestry suggesting that genotype rather than genetic ancestry was the cause of the observed effect. For example, G_ genotypes of rs16832956, the SNP with the strongest genetic association, had low CSTA expression, with a >1.4-fold ± 0.1-fold increase seen in CC genotypes (Fig. 1B, P < 1.6 × 10^{-4}). The SAE CSTA gene expression levels were confirmed by TaqMan RT-PCR (Fig. 1C, r² = 0.67, P < 10^{-4}).

**Influence of ancestral background on genetic associations with CSTA small airway epithelium gene expression**

Genetic ancestry could be a potential confounding variable limiting the generalizability of observations made in relation to the modulation of CSTA gene expression by genetic variability. However, the associations of rs16832956 genotypes with CSTA expression levels were similar among the 2 major ancestral groupings (P > 0.03, both ancestral groups), with no significant difference in allele frequency among those of African American ancestry versus subjects of European ancestry (Fig. 2A, G allele frequency 0.19 in African American ancestry group, G allele frequency 0.09 in European ancestry group, P > 0.08). Haplotypes, comprising SNPs surrounding the CSTA genomic location, were significantly associated with...
CSTA gene expression levels in SAE in a random subset of healthy
CSTA microarray-assessed SAE

either side of the gene. Shown are the correlations (Wald statistic) with
cis
the gene, same scale, are the corresponding chromosomal locations of 48
view of the

generations

rs16832956

123.40M
123.50M
123.60M
123.70M

rs16832956

rs16832956

rs16832956

rs16832956

rs16832956

CC
G

rs16832956 genotypes and
CSTA expression (microarray)

Normalized average relative expression (microarray 204951_at)

20
15
10
5
0

P < 10^(-4)

B

C

TaqMan confirmation of
CSTA gene expression

Normalized average gene expression (TaqMan)

P < 10^(-4)

r^2 = 0.67

Normalized average gene expression (microarray 204971_at)

0.0
1.0
2.0
3.0
4.0
5.0
6.0

Figure 1. Genetic modulation of CSTA SAE gene expression. A, magnified
view of the CSTA gene region, with direction of transcription (arrow). Below
the gene, same scale, are the corresponding chromosomal locations of 48
c/c-SNPs on the Affymetrix Human SNP Array 5.0 found within 100 kb
either side of the gene. Shown are the correlations (Wald statistic) with
microarray-assessed SAE CSTA gene expression in 112 healthy
nonsmokers, healthy smokers, and smokers with COPD. B, microarray
normalized average expression values of CSTA for genotypes of SNP
rs16832956. Shown are data for all 112 genotyped subjects with CSTA
SAE gene expression on the ordinate. G_ = data for the combined
genotypes GG and CG. C, TaqMan RT-PCR confirmation of microarray
CSTA gene expression levels in SAE in a random subset of healthy
nonsmokers (n = 23), healthy smokers (n = 28), and COPD smokers
(n = 13).

CSTA gene expression for specific ancestral subgroups exam-
ined separately (Fig. 2B and C), with the strongest association
seen in the group of individuals of European ancestry for
haplotype GAGGGACCCGCT (Fig. 2C, P < 0.008; see the Sup-
plementary Results section and Supplementary Table II).

Effect of smoking status and COPD on CSTA gene
expression in the SAE

Gene expression levels of CSTA in SAE were significantly
higher in the group of healthy smokers (n = 82) compared
with healthy nonsmokers (n = 60, P < 0.04, pairwise Student’s t
test), but were even more upregulated in smokers with COPD
(n = 36) compared with the healthy smokers (Fig. 3A, P < 10^{-3},
pairwise Student’s t test; P < 10^{-4} by analysis of variance for all
3 groups). The association of CSTA expression with phenotype
was confirmed by RT-PCR, and was explained neither by
genetic ancestry nor by age, though there was evidence of a
possible dose–response relationship (Fig. 3B–D; see the Sup-
plementary Results section). The effect of genotype on SAE
CSTA gene expression seen in the total genotyped study
population, was also observed when healthy nonsmokers,
healthy smokers, and COPD smokers were examined sepa-
rately (see the Supplementary Results section, Supplementary
Fig. 1A and B).

Western analysis of cystatin A protein expression

Whole-cell lysates of SAE from healthy nonsmokers, healthy
smokers, and smokers with COPD were quantitatively
assessed for cystatin A expression using Western analysis.
Western analysis revealed increased CSTA protein expression
in healthy smokers compared with nonsmokers, with even
higher expression in smokers with COPD compared with the
healthy smokers (Fig. 4A), which was confirmed quantitatively
by densitometry (Fig. 4B, P < 0.04 by pairwise comparisons by
Student’s t test; P < 10^{-2} by analysis of variance for all 3
phenotypic groups).

CSTA gene expression in LAE, SCC, and
adenocarcinoma

Most lung cancers, especially adenocarcinoma, are derived
from the SAE, and yet it is well established that smoking and
COPD are more strongly linked to SCC than to adenocarci-
nomas (14, 28, 30). SCCs tend to arise in the more central
airways (15). A comparison of CSTA expression was made
among SAE, LAE, and non–small cell lung cancer (NSCLC)
specimens (see the Supplementary Results section for details
of the study population demographics and characteristics of
brushed LAE samples). CSTA expression levels in LAE of
healthy nonsmokers were similar to the upregulated levels
seen in the SAE of COPD smokers. LAE CSTA expression was
smoke-responsive (Fig. 5A, P < 10^{-3}) and but further upre-
gulated in SCC (Fig. 5A, P < 10^{-2}, all comparisons). In contrast,
adenoarcinoma CSTA gene expression levels were sig-
ificantly downregulated compared with all but the SAE of
healthy nonsmokers (Fig. 5A; see the Supplementary Results
section).

In view of the marked upregulation of CSTA observed in
SCC subjects, a comparison was made of the ratio of CSTA
gene expression to expression levels of the 3 known cathepsin targets (B, H, and L) of CSTA within individuals. These expression ratios differed significantly among most phenotypic and tissue groupings, except for adenocarcinoma versus nonsmoker SAE (see the Supplementary Results section, Fig. 5B–D).

Discussion

The cytosolic cysteine protease inhibitor cystatin A, coded for by the CSTA gene, has attracted interest with a number of reports relating expression of CSTA protein levels to neoplastic states in a variety of tissues including those of epithelial origin (7–12, 36). Immunohistochemical data have suggested that CSTA is expressed at higher levels in dysplastic human bronchial epithelium, in SCC in particular, and less often in adenocarcinoma of the lung, compared with normal bronchial epithelium (11, 12). In light of the fact that COPD, a disease that arises in the small airways, is a risk factor for lung cancer independent of the risk attributable to smoking, but only occurs in a minority of smokers (16–20, 22–27, 29), we asked: does genetic variation modulate CSTA gene expression levels in the human SAE; is the expression of CSTA haplotypes, African American ancestry

rs16832956

allele frequency

Figure 2. Effect of ancestry on genetic modulation of SAE CSTA expression. A, CSTA expression based on genotypes of rs16832956 parsed by the 2 major ancestral groups, African American and European. Ordinate—normalized average relative CSTA gene expression. Allele frequencies as shown. B, analysis of haplotypes derived from local CSTA cis-genotypes in subjects of African American genetic ancestry. Analyses performed on the subset of 56 subjects of African American ancestry with available genotype data. The genomic sequence and frequencies of the 6 haplotypes blocks are shown in the upper portion of the figure, corresponding to the named SNPs indicated below. In the haplotype map, the darker the shading, the stronger the linkage disequilibrium (LD) between adjacent SNPs, and vice versa. The haplotypes in blocks 3 and 4 (highlighted in bold type, with the highlighted constituent SNPs shown beneath) were significantly associated with CSTA gene expression (see Supplementary Table II). The relative location and direction of the CSTA gene is indicated. The relative location of the SNP most strongly associated with CSTA gene expression, rs16832956, is indicated (vertical arrow). C, CSTA haplotype association data for 35 subjects of European genetic ancestry. The indicated haplotype in block 2, composed of the 12 SNPs shown in bold, was the only haplotype significantly associated with CSTA gene expression (see Supplementary Table II).
**CSTA** gene expression levels in small airway epithelium (SAE) of healthy nonsmokers, healthy smokers, and smokers with COPD. A, average relative mRNA expression values of **CSTA** (healthy nonsmokers, \(n = 60\); healthy smokers, \(n = 82\); and COPD smokers, \(n = 36\)) as detected by microarray. B, **CSTA** microarray expression was confirmed at the transcript level using TaqMan RT-PCR. C, average relative expression level of **CSTA** as a function of age and smoking history. D, average relative expression level of **CSTA** in SAE as a function of ancestry and phenotype.

Figure 3. **CSTA** gene expression levels in small airway epithelium (SAE) of healthy nonsmokers, healthy smokers, and smokers with COPD. A, average relative mRNA expression values of **CSTA** (healthy nonsmokers, \(n = 60\); healthy smokers, \(n = 82\); and COPD smokers, \(n = 36\)) as detected by microarray. B, TaqMan RT-PCR confirmation of the association of SAE **CSTA** gene expression levels with phenotype in a subset of healthy nonsmokers (\(n = 23\)), healthy smokers (\(n = 28\)), and smokers with COPD (\(n = 13\)). C, average relative expression level of **CSTA** for the same subjects as in A, with genetic ancestry on the x-axis. Two-factor ANOVA: \(P < 10^{-5}\) for phenotype; \(P > 0.05\) for the factor "ancestry". Interaction among phenotype and ancestry \(P > 0.5\). D, average relative expression level of **CSTA** in SAE as a function of age and smoking history. Left ordinate: **CSTA** gene expression, bar plots; right ordinate: smoking history of the 2 groups of smokers (in pack-years), dot and whisker plots and asterisk \(P\) values. Each of the phenotypic groups is divided by mean age (of that group). Error bars, standard error. \(P\) values are from pairwise student t tests, except where otherwise indicated.

**CSTA** in SAE influenced by both smoking and by COPD; and since smoking and COPD each have a propensity toward SCC rather than adenocarcinoma (28–30), is **CSTA** expression differentially influenced by SCC versus adenocarcinoma of the lung relative to SAE and LAE expression levels in the noncancerous samples?

Using fiberoptic bronchoscopy, pure populations of SAE were obtained from 60 carefully phenotyped healthy nonsmokers, 82 healthy smokers, and 36 smokers with COPD, and for the majority, both SAE **CSTA** gene expression and corresponding blood **CSTA** genotypes were assessed by microarray. The data demonstrate that **CSTA** gene expression levels in the human SAE are modulated by 12 SNPs within the vicinity of the gene, a genetic association that is not confounded by genetic ancestry. There is an association of local cis-haplotypes with **CSTA** gene expression for 4 haplotypes in the case of subjects of African American ancestry and 1 haplotype in those of European ancestry. In addition, healthy smokers have higher **CSTA** gene expression levels than healthy nonsmokers, with even higher levels observed in smokers with COPD compared with the healthy smokers independent of pack-years, albeit with a suggestive evidence of a dose–response relationship for smokers. The microarray expression was confirmed at the transcript level by quantitative TaqMan RT-PCR and at the protein level by Western analysis. The genetic modulation of SAE **CSTA** gene expression was a consistent finding when the 3 phenotypic groups were examined separately. Finally, tumor tissues from individuals with SCC of the lung have the highest relative levels of **CSTA** gene expression, but tumor tissue from adenocarcinoma subjects have substantially lower levels of **CSTA** than in any of the other tissues and phenotypes examined (except for being similar to healthy nonsmokers). Interestingly, the phenotypic states examined (healthy, COPD, lung cancer) are each associated with significantly different proportional gene expression of **CSTA** to cathepsins B, H, and L. These observations are in keeping with the concept that the **CSTA** gene plays a role in the evolution of the bronchial epithelium of peripheral and central airways from normal to disease, and suggests a
complex interplay among genetics, smoking, COPD, and lung cancer histologic subtype in relation to CSTA gene expression. CSTA appears to have a more plausible connection to the specific evolution of healthy smoker and COPD smoker airway epithelium into SCC rather than into adenocarcinoma, in keeping with the known stronger relationship of the separate risk factors of smoking and COPD with SCC than with adenocarcinoma of the lung (27–30).

Small airway epithelium, smoke-induced lung disease, and genetics

There is now a large body of evidence pointing to the SAE (defined as bronchi of ≥6 generations, <2 mm in diameter) as the earliest site of pathologic involvement in COPD, the primary site of airflow limitation in this disorder, and the site of development of most NSCLC (13–15, 18–21). Consistent with these observations in the major smoke-induced lung diseases, morphologic changes are found in small airways of asymptomatic smokers with normal lung function (37, 38). COPD is a relevant phenotype to study in the progression from healthy airway epithelium to lung cancer, an airway epithelial-derived disease, because COPD is itself an independent risk factor for the development of lung cancer, with a contribution to lung cancer risk that is separate from the risk attributable to cigarette smoking (22–26). Since only 15% to 20% of smokers will develop COPD, together with evidence of familial clustering in COPD and NSCLC and twin concordance for the risk for COPD (15–17, 22, 39–41), it is likely that genetic variation influences disease risk. The fact that SCC is more common than adenocarcinoma in smokers and COPD subjects also raises the question of the role of the LAE in the pathogenesis of these tumor subtypes. Although the current study was not designed to focus on LAE gene expression/genotype relationships, it is interesting to note that the smoke-responsive nature of CSTA expression in SAE was also observed in LAE. However, the levels of CSTA in healthy smoker LAE are still considerably less than in SCC subjects and taken together with the smoke-responsiveness, are therefore less likely to reflect mere constitutive expression of CSTA in a LAE-derived tumor. Although the observations of genetic modulation of CSTA gene expression and the upregulation of such expression in SAE from COPD smokers compared with healthy smokers is intriguing, the present study was not designed to address the question of whether or not CSTA expression is genetically associated with COPD. Testing this hypothesis would require the genotyping of a large number of subjects to have sufficient statistical power to reach a definitive conclusion.

Although the data show some evidence for a dose–response relationship between intensity of smoke exposure and CSTA gene expression, disease also influences the expression levels, as evidenced by the greater relative increase in CSTA levels in COPD and SCC versus healthy smokers compared with the smaller increase in healthy smokers versus nonsmokers, even though the latter comparison would have the greater difference in smoking exposure. Also, reciprocal effects of disease state and pack-years on CSTA gene expression in SAE were observed in the healthy smoker and COPD smoker age-defined subgroups.

Analysis of a published gene expression data set that used microarray technology to study lung cancer tissues did not show a significant effect of smoking status on CSTA gene expression (42). Others have noted that CSTA levels are higher in early-detected lung cancers compared with lung cancers presenting later in the disease course, despite higher pack-year values in the latter (43). These disparate observations highlight the need for further study to clarify if there is an effect of smoking-intensity on CSTA gene expression in lung airway epithelium and cancers derived from the airway epithelium. The observations suggest that CSTA expression is not uniformly smoke-responsive and is likely to be specific to cell populations. Controlling for the influence of age while examining pack-year consumption is challenging and would likely require large data sets, because more often than not, individuals with higher pack-year histories will also be older. From the data in the present study, it seems more likely that it is the intensity of smoking rather than age that exerts the greater influence on CSTA expression levels, as evidenced by the concordance among the magnitudes of pack-year difference and the difference in SAE CSTA levels in comparing the 2 smoking phenotypic groups (healthy smokers and COPD smokers) parsed by median age.
Cystatin A in lung disease

In the normal human airway epithelium, cystatin A is detected in the basal cells by immunohistochemistry, with more extensive staining in preneoplastic bronchial epithelium (12). Strong staining for cystatin A has also been observed in lung cancer tissue, especially SCC but also in adenocarcinoma and large cell carcinoma (11, 12). For SCC, cystatin A expression was less marked in less well-differentiated tumors, in which case lower cystatin A expression was also associated with tumor recurrence (12). Quantitative upregulation of CSTA protein has been noted in human lung cancer tissues versus control lung using ELISA, and univariate survival analyses showed that individuals with higher levels of cystatin A had a better survival probability, suggesting that CSTA is upregulated in lung tumors to counteract potentially harmful tumor-associated activity (11, 12). The present study confirms the previously published immunohistochemical data showing an upregulation of cystatin A in SCC versus normal airway epithelium, but also reveals a downregulation of CSTA in adenocarcinoma relative to levels in the healthy state, challenging data from another cohort that suggested an upregulation of cystatin A in some (but not in all) bronchogenic adenocarcinomas compared with normal airway epithelium (12).

Whereas all of the functions of CSTA are not clear, other than its known cysteine protease inhibitory effects, there is some evidence that it inhibits apoptosis in the presence of stimuli such as UVB radiation and viruses (4–6). Although deficiency states have been described arising from mutations in the genes that code for cystatins B and C, resulting in phenotypes of progressive myoclonus epilepsy and hereditary cystatin C amyloid angiopathy respectively, to date no functional mutations have been reported for CSTA in humans (44, 45).

The putative role of CSTA as a tumor suppressor is supported by observations in other tissues, including breast, prostate, and esophageal tumors, with evidence that cystatin A can inhibit tumor cell growth, angiogenesis, invasion, and metastasis (9, 36, 46). The observations in the present study that CSTA is downregulated in adenocarcinoma of the lung with evidence of less-well–opposed cysteine protease activity versus other smoke-exposed phenotypes are compatible with a tumor suppressor role in adenocarcinoma, but with a different function in the progression of some smokers to COPD and SCC, perhaps reflecting an “excess” of cystatin relative to its cognate proteases, the cathepsins. In contrast with these observations, a homozygous mutant mouse model involving chromosomal deletion of csta, the murine homolog of human CSTA, together with 3 other genes, showed no
phenotypic abnormality, the animals were not overtly susceptible to spontaneous or irradiation-induced tumor formation, and had evidence of compensatory gene expression of genes phylogenetically related to Csta (47).

Based on reports that cystatin A is upregulated in dysplastic airway epithelium and in lung cancer, the findings of the present study, which shows that the regulation of CSTA expression in SAE is influenced by genetic variability, smoking, and COPD, and that CSTA is differentially expressed by lung cancer histologic subtypes, suggests that each of these factors should be controlled for when considering the use of CSTA as a marker related to the pathogenesis of lung cancer. The progressive rise in CSTA expression observed in disease states (i.e., COPD and SCC) does not appear to be offset by a corresponding rise in cognate cathepsin levels in the same tissue from any given individual, an observation that may have implications for disease pathogenesis if intracellular cysteine protease-anti-protease homeostasis is relevant, particularly in genetically predisposed smokers.

Disclosure of Potential Conflicts of Interest

There are no potential conflicts of interest to disclose.

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