Multiple Forms of Genetic Instability within a 2-Mb Chromosomal Segment of 3q26.3–q27 Are Associated with Development of Esophageal Adenocarcinoma

Lin Lin,1* Zhuwen Wang,1 Michael S. Prescott,1 Herman van Dekken,2 Dafydd G. Thomas,2 Thomas J. Giordano,2 Andrew C. Chang,1 Mark B. Orringer,1 Stephen B. Gruber,3 John V. Moran,1 Thomas W. Glover,4 and David G. Beer1*

1Department of Surgery, University of Michigan, Ann Arbor, Michigan
2Department of Pathology, University of Michigan, Ann Arbor, Michigan
3Department of Internal Medicine, University of Michigan, Ann Arbor, Michigan
4Department of Human Genetics, University of Michigan, Ann Arbor, Michigan
5Department of Pathology, Erasmus Medical Center, University Medical Center Rotterdam, The Netherlands

Gene amplification is one of the mechanisms to activate oncogenes in many cancers, including esophageal adenocarcinoma (EA). In the present study, we used two-dimensional restriction landmark genome scanning to clone a NotI/DpnII fragment that showed increased genomic dosage in 1 of 44 EAs analyzed. This fragment maps to 3q26.3–q27, and subsequent experiments identified two intrachromosomal amplicons within a 10-Mb DNA segment in 7 of 75 (9%) EAs. The distal amplified-core region maps centromeric to the PIK3CA locus, and a microsatellite (D3S1754) within this region exhibited significant instability (MSI), in stark contrast to the genomewide microsatellite stability found in EA. D3S1754-MSI arises in premalignant Barrett’s dysplastic cells and preceded amplification of the nascent MSI allele in the corresponding EA. Seven ESTs within the amplified-core were overexpressed in amplicon-containing EAs. One of these, EST AW513672, represents a chimeric transcript that initiated from an antisense promoter sequence in the 5′UTR of a full-length LINE-1 element (L1-5′ASP). Similar chimeric transcripts encoding portions of the MET oncogene and the BCAS3 gene also were overexpressed in EAs, suggesting that L1-5′ASP activation may occur at a broad level in primary EAs. Thus, the fine dissection of a 2-Mb amplified DNA segment in 3q26.3–q27 in EA revealed multiple genetic alterations that had occurred sequentially and/or concurrently during EA development. This article has supplementary material, available at http://www.interscience.wiley.com/jpages/1045-2257/suppmat.

INTRODUCTION

The incidence of esophageal adenocarcinoma (EA) has been increasing rapidly in many western countries (Devesa et al., 1998; Bollschweiler et al., 2001). Chronic gastroesophageal reflux (GER) has been proposed as a major risk factor for the development of Barrett’s metaplasia, and EA frequently is associated with adjacent Barrett’s metaplasia and/or dysplasia (Winters et al., 1987; Lagergren et al., 1999). The prognosis for patients with EA is poor, with a 5-year survival rate of only 10% (Farrow and Vaughan, 1996).

A number of genetic alterations have been observed in EAs. These include the frequent occurrence of somatic mutations and/or loss of heterozygosity (LOH) in the TP53 and CDKN2A tumor-suppressor genes in both EA and pre-malignant Barrett’s mucosa (Casson et al., 1991; Barrett et al., 1996). In addition, the use of comparative genomic hybridization (CGH) has enabled observation of multiple genomic losses and/or gains in EA (Riegerman et al., 2001). Through the use of two-dimensional restriction landmark genome scanning (2-D RLGS) and STS-amplification mapping, we previously reported finding intrachromosomal amplicons at multiple chromosomal locations in EAs and identifying candidate genes that may be involved in cancer development and/or progression (Hughes et al., 1998; Lin et al., 2000a; 2000b; Lin et al., 2002; Miller et al., 2003a). Gene amplification represents one essential mechanism for the activation of proto-oncogenes and is a tumor-specific event that does not occur in the normal human genome (Bishop, 1987). Therefore, identifying and characterizing these amplicons are important to delineate the molecular events that underlie EA tumorigenesis.

In the present article, we report the identification of a 2-Mb intrachromosomal amplicon at...
3q26.3–q27 in EAs. The amplified DNA fragment is associated with various forms of genomic instability, including regional microsatellite instability, overexpression of various cellular transcripts, and the induction of a chimeric transcript initiated from an antisense promoter located in the 5’UTR of a full-length LINE-1 element. The presence of these varied genetic alterations within a 2-Mb chromosomal segment in EA may suggest a common mechanism in cancer development but will certainly require additional studies of this and other cancer types.

**MATERIALS AND METHODS**

**Tissue Collection**

Tumors and their associated normal tissue were obtained from patients undergoing esophagectomy or pulmonary resection at the University of Michigan Medical Center between 1992 and 2000. Patients provided written consent, and the project was approved by the University of Michigan Institutional Review Board. Patients in this study had no preoperative radiotherapy or chemotherapy. Tissue samples were frozen in liquid nitrogen and stored at −80°C until use.

**Cell Lines**

Two cell lines, Flo-1 and Bic-1, were derived from EA tissues in our laboratory. Het-1A is a human esophageal squamous cell line immortalized using SV40 and was kindly provided by Dr. Gary Stoner of Ohio State University.

**DNA Isolation and 2-D RLGS Gel Electrophoresis**

High-molecular-weight DNA was isolated as previously described (Blin and Stafford, 1976). All tumor portions used for DNA isolation were more than 70% tumor cells, as determined by frozen-tissue sectioning. Two-dimensional RLGS gel electrophoresis was performed as previously described (Kuick et al., 1995).

**Cloning of 2-D DNA Fragment**

The 2-D DNA fragment was purified and cloned as previously described (Lin et al., 2000b). Individual colonies were collected for DNA isolation (minipreps) using QIAprep® Spin Miniprep kit (Qiagen, Valencia, CA).

**Bioinformatic Analysis**

The sequences of the cloned fragments were analyzed by NCBI BLAST tools (www.ncbi.nlm.nih.gov). Precise chromosome location of the cloned 2-D fragment was determined by analyzing the resulting BAC sequences using NCBI’s bioinformatic tools, electronic PCR, and Map Viewer. Exon and gene prediction software, GraiLEXP (grail.lsd.ornl.gov/grailexp/) and GENSCAN (genes.mit.edu/GENSCAN.html), was used in conjunction with the GenBank EST database to determine expressed sequences within selected contigs.

**Comparative Genomic Hybridization (CGH) and Interphase Fluorescence in situ Hybridization (FISH) Assays**

CGH analysis was performed as described previously (Riegman et al., 2001). DNA loss was defined as chromosomal regions in which the mean green-to-red signal ratio was below 0.80, whereas gain was defined as regions in which the ratio was above 1.20. High-level amplification was seen as a distinct peak (ratio > 1.5). At least 8–10 metaphases were used per sample. Interphase FISH was assayed as described previously (Lin et al., 2002), except for the chromosome arm 14q probe (BAC clone R-356O9 DNA), which was cohybridized as a control with the target probe BAC AC076966, which includes the amplified 2-D sequence.

**STS Amplification Mapping Using Quantitative Genomic PCR (QG-PCR)**

STS/EST/gene markers in the 3q26–q27 regions were selected for QG-PCR, as previously described (Lin et al., 2000a). QG-PCR is a semi-quantitative PCR procedure that involves a multiplex PCR reaction in which a pair of primers from the control sequence, either GAPDH or the same chromosome centromeric/telomeric markers, were coamplified with the target genomic sequence in the same PCR reaction. Densitometry of PCR product signal ratios (Ts/c:Ns/c) for tumor (Ts/c, tumor STS fragment/tumor control) and normal (Ns/c, normal STS fragment/normal control) DNA was quantified using ImageQuant software (Amer sham Biosciences, Piscataway, NJ). When the paired normal was not informative, the other normal samples were compared with the tumor, and the ratio was calculated. Values ≥ 2.0 were considered indicative of DNA amplification, and values between 1.5 and 1.9 were considered to indicate copy number gain. All assays were repeated three times.

**Microsatellite Instability Screening, MMR Gene Mutation, and Tissue Microarray Analyses**

Thirty microsatellite (MS) markers were chosen (Table 1) including the five recommended by the...
### Table 1. Microsatellite Markers Investigated

<table>
<thead>
<tr>
<th>Marker</th>
<th>Chromosome (map location in Mb)</th>
<th>Repeat type</th>
<th>Repeat nucleotide</th>
<th>Repeat length (bp)</th>
<th>MSI (%)</th>
</tr>
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<tr>
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<td>3 (178.8)</td>
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<td>(TAGA)n</td>
<td>98</td>
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<td>(CA)n</td>
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<td>0/48</td>
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<td>Bat-26</td>
<td>2 (47.6)</td>
<td>Monorepeat</td>
<td>(A)n</td>
<td>26</td>
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<td>Direpeat</td>
<td>(CA)n</td>
<td>60</td>
<td>0/44</td>
</tr>
<tr>
<td>Bat-21</td>
<td>3 (37.0)</td>
<td>Mono- + direpeats</td>
<td>(A)n + (TA)n</td>
<td>20 + 26</td>
<td>1/46 (L61)</td>
</tr>
<tr>
<td>D3S3603</td>
<td>3 (181.2)</td>
<td>Direpeat</td>
<td>(CA)n</td>
<td>56</td>
<td>0/12</td>
</tr>
<tr>
<td>AFM072yb7</td>
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<td>Direpeat</td>
<td>(CA)n</td>
<td>42</td>
<td>0/12</td>
</tr>
<tr>
<td>D3S3609</td>
<td>3 (183.5)</td>
<td>Tetrarepeat + trirepeats</td>
<td>(TAGA)n + (TTA)n</td>
<td>66 + 42</td>
<td>0/12</td>
</tr>
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<td>C1_4_1</td>
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<td>(CAA)n</td>
<td>32</td>
<td>0/42</td>
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<td>74</td>
<td>1/44 (163)</td>
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<td>52</td>
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<td>50</td>
<td>2/44 (D66, T67)</td>
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<td>(TAAA)n + (CA)n</td>
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<td>(CA)n + (TATATG)n</td>
<td>58 + 54</td>
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<td>(CAG)n</td>
<td>59</td>
<td>0/75</td>
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<td>(CA)n</td>
<td>36</td>
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<td>(CA)n</td>
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<td>0/75</td>
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<td>44</td>
<td>0/75</td>
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<td>Tetrarepeat</td>
<td>(TGAA)n</td>
<td>26</td>
<td>0 (44)</td>
</tr>
<tr>
<td>D17S250</td>
<td>17 (37.5)</td>
<td>Di- + direpeats</td>
<td>(TA)n + (CA)n</td>
<td>53 + 33</td>
<td>2/75 (L86, T67)</td>
</tr>
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<td>Direpeat</td>
<td>(CA)n</td>
<td>40</td>
<td>0 (44)</td>
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<td>Tetrarepeat</td>
<td>(GAAA)n</td>
<td>184</td>
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<tr>
<td>D2S119</td>
<td>2 (44.1)</td>
<td>Tetrarepeat + Hexarepeats</td>
<td>(CA)n + (CATA)n</td>
<td>50 + 70</td>
<td>0 (44)</td>
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<td>DYS19</td>
<td>Y (9.1)</td>
<td>Tetrarepeat</td>
<td>(TAGA)n</td>
<td>66</td>
<td>0/44</td>
</tr>
</tbody>
</table>

*Recommended MSI markers for diagnosis and research by the NCI MSI Detection Workshop (Boland et al., 1998).

**These tumors have two positive MSI markers including D3S1754 among the 75 tumors examined.

*Marker D3S1754 is at 178.8-Mb of 3q (as of February 2004) and is within the core-amplified domain. EAs positive for MSI with this marker are A07, D06, D66, L63, I63, M55, M59, W61, L86, M60, M55, S08, B95, V65, L86, V61.

**Bat-21 spans the sequence between 3’ end of intron 11 and 5’ site of exon 12 of human MLH1 gene. PCR fragment of Bat-26 intersects intron 4 and exon 5 of the MSH2 gene.

**Markers D17S250 (37.5-Mb) and D17S1787 (40.1-Mb) are 0.8 and 1.8-Mb telomeric, respectively, to the ERBB2 (38.3 Mb) oncogene. ERBB2 represents the most frequently amplified amplicon (21.8%) in esophageal adenocarcinoma (Miller et al., 2003b).

NCI guidelines of MSI in cancer detection (Boland et al., 1998). Another five markers were of the (TAGA)n type, analogous to marker D3S1754. Forward primers were 32P-labeled, and PCR was applied to the DNA from normal–tumor pairs. Additional novel MS fragments within the 3q and ERBB2 amplicons were designed using the Repeat Masker Server (repeatmasker.genome.washington.edu) and analyzed. Primer sequences are given in the Supplementary Table (supplementary material for this article can be found at http://www.interscience.wiley.com/jpages/1045-2258/suppmat). The GenBank accession numbers of microsatellite fragments 3q12136-3 and 17q21-1 are DQ157857 and DQ157856, respectively. Mutation analysis of MSH3 and MSH6 was performed according to the modifications from Yin et al. (1997).

A tissue microarray (TMA) block was constructed according to Kononen et al. (1998). The TMA contained multiple cores of 64 EA resections from 59 patients, 8 lymph node metastases, 8 dysplastic Barrett’s mucosas, 11 Barrett’s mucosas, and 10 normal controls from various tissues. Sections from 59 patients, 8 lymph node metastases, 8 dysplastic Barrett’s mucosas, 11 Barrett’s mucosas, and 10 normal controls from various tissues. Sections were incubated with antibodies against MLH1 (1:100 dilution; BD Biosciences, San Diego, CA; cat. no. 554073) or MSH2 (1:100 dilution; Oncogene Research Products, Boston, MA; cat. no. NA27). Microwave citric acid epitope retrieval was performed for 20 min for both antibodies. Each slide was lightly counterstained with hematoxylin.

*Recommended MSI markers for diagnosis and research by the NCI MSI Detection Workshop (Boland et al., 1998).
Southern Blot and Array CGH Analyses

EST AW513672 (nt5635-2) and PIK3CA were used as probes and hybridized to Southern membranes containing six pairs of normal–EA DNA using standard hybridization and washing conditions. Array CGH was performed as previously described (Pinkel et al., 1998).

RNA Isolation and Quantitative RT-PCR

Total RNA was isolated using Trizol reagent (Invitrogen, Carlsbad, CA). All the RNA samples were treated with DNase I (Promega, Madison, WI) prior to performing reverse transcription. Two micrograms of total RNA was reverse-transcribed (Invitrogen, Carlsbad, CA) and primed by both (dT)18 and random hexamers in a 20-ul reaction volume. One microliter of the cDNA products underwent RT-PCR using GAPDH as a coamplified internal control. The RT-PCR products were resolved on 8% denaturing PAGE gels and analyzed using ImageQuant (Amersham Biosciences).

Reverse Northern Blot Analysis

Eight BAC clones covering the chromosome 3q26.3–q27 region from 178.4- to 180.6-Mb were selected. BAC clone RP11-245C23 (BAC3K) was kindly provided by Dr. Steve Scherer, Baylor College of Medicine. All the remaining BACs were purchased from BACPAC Resources (Children’s Hospital, Oakland Research Institute, Oakland, CA). BAC DNA was prepared as previously described (Lin et al., 2002) and digested using the EcoRI restriction enzyme. Tumor or normal RNA from patient M28 was reverse-transcribed with oligo(T) and [α-32P]dCTP for direct incorporation for the synthetic cDNA, then hybridized to the arrayed BAC membrane.

Affymetrix Microarray Assays

Forty-six samples including nondysplastic and dysplastic Barrett’s mucosas and EAs were subjected to gene expression profiling using Affymetrix U133A chips (Affymetrix, Santa Clara, CA) as previously described (Giordano et al., 2001).

DNA Sequencing

All IMAGE cDNA clones were purchased from Open Biosystems (Huntsville, AL), and plasmid DNA was prepared using a QIAprep® Spin Miniprep kit (Qiagen). The GenBank accession numbers of the full-length insert sequencing are AY679731 for IMAGE 2737847, AY679732 for IMAGE 2344756, and AY769439 for IMAGE 4345107. DNA was sequenced by the University of Michigan Sequencing Core.

RESULTS

A Cloned Restriction Fragment Amplified in EA Is Mapped to 3q26.3–q27

We used two-dimensional (2-D) RLGS to investigate genomic amplification in 44 primary EAs. More than 2,000 individual NotI/HinFI restriction fragments were visualized and used to compare the 2-D images of normal and tumor DNA (Fig. 1). In one patient (EA I63), we identified a NotI/HinFI

Figure 1. Identification of intra-chromosome amplification in EA DNA using RLGS. The numbers represent the restriction sizes in 1-D (4,900 bp) and 2-D (260 bp for NotI/HinFI digestion and 340 bp for NotI/DpnII) gels. An enlarged area of the respective target fragment is shown in the panels underneath. (A) Restriction NotI/HinFI fragments from normal DNA of patient I63 were resolved on 2-D RLGS gels. (B) Two-dimensional gel image of EA I63. A comparison between normal and tumor gel images reveals a fragment that increased in density in the EA (arrow in B) relative to the corresponding normal (arrow in A). (C) NotI/DpnII digestion was performed for cloning purposes. Fragment C (arrow) was cloned directly from the 2-D gel. The very intense fragments visible on these gels represent ribosomal DNA, present in multiple copies in the genome.
DNA fragment that was amplified in the primary EA (Fig. 1B), but not its matched normal DNA (Fig. 1A). A corresponding 343-bp fragment from NotI/DpnII restriction digestion (Fig. 1C) was cloned, and the sequence of the cloned fragment was found to be identical to a sequence within BAC clone AC076966, which maps to chromosome band 3q27.1 (www.ncbi.nlm.nih.gov). We then performed CGH analysis on 3 of the 44 EAs including the EA (I63) that demonstrated the amplified NotI/DpnII fragment in RLGS assay (Fig. 2A). CGH verified that the EA I63 DNA had an increased genomic dosage at 3q26–q27 (Fig. 2A). Moreover, interphase FISH performed on EA I63 demonstrated an increased copy number when probe AC076966 DNA was hybridized and compared with non-amplified EA and the metaphase control (Fig. 2B).

**Two Amplicons Identified in 3q26.3–q27 Region**

To characterize and fine-map the 3q26.3–q27 amplification, we designed a series of STS/EST/gene markers extending more than 5-Mb in both directions from the location of the 2-D fragment. We applied these markers to 75 pairs of EAs and their matched normal DNA, which included most of the 44 EA samples we had analyzed in the 2-D RLGS assays. STS-amplification mapping revealed that 7 of 75 (9.3%) EAs, 3 of 33 (9.0%) lung adenocarcinomas, and 1 of 24 (4.2%) esophageal squamous carcinomas demonstrated amplification at 3q26.3–q27 (Fig. 3A). Two amplification units were identified in the 10-Mb segment (Figs. 3A and 4). The proximal 3q amplicon encompassed the TERC and SKIL loci, whereas the distal amplified-core region mapped between D3S3096 and WI-13792 (Figs. 3A and 4). EA M28 exhibited the highest copy number (>10-fold) at marker D3S1754 (Fig. 3B), but was not amplified at the PIK3CA locus (Fig. 3A). DNA copy number remained unchanged at control markers on the chromosome 3 centromere and 3p telomere, respectively, indicating that the amplicon in all affected tumors was intrachromosomal (Fig. 3C). These centromeric or telomeric sequences of chromosome 3 were used as...
internal controls for the multiplex PCR reactions, similar to the use of GAPDH as the internal control (Fig. 3A).

Array CGH of four pairs of EA and matched Barrett’s DNA from the pool of 75 DNA samples used in this study revealed two peaks exhibiting an increase in DNA dosage in EA B05 at 3q26–q27 (Fig. 3D), consistent with the mapping results presented in Figure 3A. The centromeric peak was linked to the TERC and SKIL loci (BAC clone RP11-141C22); and the copy number of the PIK3CA locus was unchanged in this EA (Fig. 3D). The telomeric peak lay outside the 10-Mb mapping region in EA B05 (Fig. 3D). Southern blot analysis of EA M28 and B05 demonstrated increased DNA dosage at 179.7-Mb using probe AW513672 but not with the probe containing PIK3CA (data not shown). Together, the above data identified two amplicons within the 10-Mb region in a subset of EAs, and the telomeric amplified-core domain may likely exclude PIK3CA as a candidate gene for amplification in EA (Fig. 4).

Frequent Microsatellite Instability Found at D3S1754 and Regional MSI Precedes Nascent Allele Amplification during EA Tumorigenesis

Analysis of the distal 3q26.3–q27 amplicon in the EAs also revealed that amplification often affected only one allele, with the second allele either unchanged or deleted in the EA genome (Fig. 5A). These results might account for the inability to detect an increase in DNA copy number when a nonpolymorphic marker was used.

Further analysis of the 3q26–q27 amplicon revealed relatively frequent MSI at marker D3S1754 (16.0%) in EAs, which was in stark contrast to the genomewide microsatellite stability (MSS) and/or low frequency of MSI (MSI-L, 2.6%, \( P < 0.005, \) standard \( \chi^2 \) test) commonly observed in EA (Fig. 5B and Table 1). D3S1754 resides in the distal amplified core and showed the highest DNA copy number in EA M28 (Fig. 3B). Interestingly, amplification was detected in the newly replicated MSI allele in EA M28 (Fig. 5B), prompting us to investigate the possible relationships between...
We did not find mutations in \textit{MSH3} and \textit{MSH6}. Both MLH1 and MSH2 were abundantly expressed at the protein level as shown by immunohistochemistry on tissue microarrays (TMA) in D3S1754-MSI(+) EAs (data not shown). We concluded that amplification may not be responsible for D3S1754-MSI because: (a) D3S1754-MSI occurring in premalignant dysplastic Barrett’s mucosa and preceding the genomic amplification that arose only in the EA though allelic imbalance could be observed as early as in the premalignant Barrett’s DNA (Fig. 5C); (b) only 2 of 7 amplified EAs were D3S1754-MSI(+) and 2 of 12 D3S1754-MSI(+) EAs developed 3q amplification (Figs. 4 and 5B); (c) other microsatellites within both the 3q26.3–q27 and \textit{ERBB2} amplified-core regions were stable in the D3S1754-MSI(+) EAs (data not shown); and (d) D3S1754-MSI was positive in four MS-unstable (MMR\textsuperscript{+}) colon cancers, whereas it was negative in all five MS-stable colon cancers examined (Fig. 5D). Thus, microsatellite D3S1754 is highly unstable in both EA (MMR\textsuperscript{+}) and colon cancers regardless of MMR proficiency.

Furthermore, the identical genotype of the nascent D3S1754-MSI alleles between Barrett’s dysplasia and the corresponding EAs provided molecular evidence that EA is derived directly from Barrett’s dysplasia by clonal expansion (Fig. 5C).

\textbf{Seven ESTs Mapped within Distal 3q Amplified Core Are Overexpressed in EA}

We next implemented a reverse Northern blot assay to identify differentially expressed transcripts within the amplified-core region. Two of eight arrayed BAC DNAs that map within the amplified-core showed increased expression in EA M28 RNA (Fig. 6A). Fifteen ESTs, including eight within the two BACs (BAC1 and BAC3, Fig. 6A), were screened by quantitative RT-PCR (Fig. 6B and C). Overexpression of seven ESTs was detected in EAs containing the distal amplicon (Fig. 6B and C). We also used U133A microarrays to analyze expression-profiling data at 3q26.3–q27 in 46 samples including samples of Barrett’s metaplasia, dysplasia, and EA. Although increased \textit{PIK3CA} expression was found in 3 of 15 EAs, none of these three contained the 3q amplicon, suggesting increased expression of \textit{PIK3CA} resulted from transcriptional alterations (data not shown). These results supported the previous notion that \textit{PIK3CA} was not part of distal 3q amplicon. One of the overexpressed ESTs, \textit{n5635-3}, is a part of \textit{KCNMB2}. EST
nt5635-7 is a portion of a 1.25-kb transcript (GenBank accession number AY686686; Fig. 7A). Sequence analysis of seven IMAGE cDNA clones analogous to L1-5’ASP/EAAS2 from various cell or tissue sources validated the observed RT-PCR products from the EA samples (Fig. 7B and C; and see GenBank IDs in Materials and Methods section). We also determined that L1-5’ASP/EAAS2 was spliced from 5–8 exonic structures all residing within the distal amplified core (Fig. 7A and C). As shown, L1-5’ASP/EAAS2 is differentially expressed not only in EA cells but also in primary EAs (Fig. 7B). In addition, we also identified nine CpG islands in this L1-5’ASP (Fig. 7D).

Given the above data, we next investigated the possible consequence of L1 5’ASP activity in other EA amplicons. We selected from the GenBank database two L1-5’ASP-associated cDNAs in known amplicons, L1-5’ASP/MET (BF208095), at 7q31.3, and L1-5’ASP/BCAS3 (AU123136), at 17q23, encoding portions of the MET oncogene and the BCAS3 gene, respectively (Fig. 8). BCAS3 is a gene of unknown function that commonly is amplified in breast cancer (Barlund et al., 2002). RT-PCR analyses revealed that L1-5’ASP/MET and L1-5’ASP/BCAS3 were differentially expressed in both primary EAs and EA cell lines (Fig. 8A). Interestingly, the expression of L1-5’ASP/MET was found only in EAs containing MET amplification and in two EA cell lines (Fig. 8A). L1-5’ASP in L1-5’ASP/MET and L1-5’ASP/BCAS3 were both found within the introns of the two genes, initiating two truncated gene forms, respectively (Fig. 8B).

**DISCUSSION**

Genomic amplification is often detected in cancer (Knuutila et al., 1998) and is an important mechanism for activating proto-oncogenes, resulting in high-level expression of the selected gene products (Bishop, 1987). Two-dimensional RLGS allows the comparison of greater than 2000 restriction fragments between normal and tumor DNA isolated from the same patient (Kuick et al., 1995), and cloning of an affected 2-D RLGS spot leads directly to the identification and localization of amplicons in the cancer genome (Hughes et al., 1998; Lin et al., 2000b). Genomic amplification at 3q25–q27 has been observed in several human cancers, suggesting the presence of genes that may have roles in the development or progression of multiple cancer types. In most studies, CGH defined the minimal amplified region as a large DNA segment of several megabases, making it difficult to identify a specific candidate gene responsible for tumor development (Ma et al., 2000; Guan...
et al., 2001; Imoto et al., 2001; Singh et al., 2001; Redon et al., 2002; Heselmeyer-Haddad et al., 2003). In the present study, we were able to fine-map the core amplified domain to a less than a 1-Mb region at chromosome band 3q26.3–q27. Seven ESTs within this amplified-core domain were found to be overexpressed or only expressed in EAs. Genomic amplification was found to be involved in only one allele when a group of polymorphic microsatellites was examined in EAs. The allelic amplification observed may support the common disease/common variant (CD/CV) hypothesis (Lander, 1996) and, most importantly, suggests the necessity of using both microsatellite polymorphisms and single-nucleotide polymorphisms (SNPs) to identify the risk allele within a specific EA amplicon.

We observed specific microsatellite instability (MSI) at marker D3S1754, which is in stark contrast to the genomewide microsatellite stability (MSS) and/or low frequency of MSI commonly observed in EA (in the present study and from Muzeau et al., 1997; Kulke et al., 2001). Interestingly, this unstable D3S1754 demonstrated the highest DNA copy number in the 3q26.3–q27 amplicon in tumor M28 (Fig. 3B). MSI may result from a pronounced deficiency of mismatch repair (MMR; Strand et al., 1993), yet genomewide MSS in EA indicates intact MMR. Consistent with this idea, we did not find mutations in MSH3 and MSH6 and demonstrated that both MLH1 and MSH2 were abundantly expressed at the protein level in D3S1754-MSI(+) EAs. The increased DNA polymerase slippage in the amplified region that results in frequent MSI at marker D3S1754, therefore, might be attained by an MMR-independent mechanism. Moreover, microsatellites selected from the ERBB2 amplicon were demonstrated to be MSI negative in EAs, suggesting that genomic amplification may not be associated with the origination of the marker-specific MSI found.
with D3S1754 in this study. ERBB2 is the most frequently amplified gene in EA (Miller et al., 2003b). Previous studies have shown that mismatch repair is microsatellite repeat-unit size dependent. Although the mutation rates are similar in MS tetra-, di-, and monorepeat fragments, Sia et al. (1997) showed that repair of tetraneucleotide repeats may be less efficient than repair of monorepeat microsatellites in yeast models. Nonetheless, sporadic microsatellite mutations, elevated microsatellite instability at selected tetraneucleotide repeats (EMAST), in particular, are distinct forms of MSI because of a lack of MMR dependence and have been reported in many cancers (Ahrendt et al., 2000; Danaee et al., 2002; Catto et al., 2003). Studies have shown that endogenous production of oxygen free radicals and carcinogen-induced DNA damage may promote instability of microsatellite sequences (Jackson et al., 1998; Slegos et al., 2002). D3S1754, at 3q26.3–q27, is a (TAGA)$_{98}$ tetraneucleotide microsatellite. Taken together, D3S1754-MSI may reflect genetic and/or environmental insults from gastroesophageal reflux and/or inflammation in EA, and the resultant sporadic MSI may be a result of less efficient mismatch repair of this particular tetraneucleotide repeat. Interestingly, colorectal cancers with MMR-dependent MSI have been found to be near-diploid and do not show the increased rates of chromosome losses and gains that are characterized as chromo-

Figure 7. Chimeric L1 5’ASP/cellular transcripts were identified and found to be differentially expressed in EAs. (A) Diagram of seven amplified ESTs showing increased expression in the EAs in the 0.5-Mb amplified-core region. Numbers above each open or solid bar represent the EST series nt5635 examined in this study (Fig. 6B and C). The arrow above the number shows the transcription orientation. The Sequence of nt5635-2 (EAAS2) is identical to the 3’ end of a multiexon transcript, initiated by L1-5’ASP (solid gray bars). A 6.2-kb full-length L1 retrotransposon is mapped adjacent to EAAS2 in an opposite transcription direction. (B) RT-PCR products resolved on a 1.0% agarose gel. Differentially expressed L1-5’ASP/EAAS2 as well as transcription variants are shown. (C) Schematic of transcription variants of L1-5’ASP/EAAS2. r with arrows indicate forward and reverse primers for RT-PCR. Numbers representing each splicing structure are indicated. Uppercase indicates exon sequences, and lowercase indicates introns. Gray bars represent L1-5’ASP sequences. (D) Sequence alignment between the L1.2 promoter sequence (M80343) and the 104-bp L1-5’ASP sequence of the 5’ASP/EAAS2 transcript. A short line above highlights each of nine CpG islands.
some instability (GIN) (Jallepalli and Lengauer, 2001). Our data from CGH and STS-amplification mapping demonstrated that intrachromosomal amplification occurs at 3q26.3–q27 (Figs. 2A, 3A, 3C, 3D, and 4), and aneuploidy of chromosome 3 was excluded (Fig. 3C). Intrachromosomal amplification may arise from the breakage–fusion–bridge (BFB) cycle (Coquelle et al., 1997). In contrast, CIN appears to involve gross chromosomal changes that lead to aneuploidy in cancers and has been demonstrated to be affected by multiple genes related to cell mitosis (Jallepalli and Lengauer, 2001).

Among the seven ESTs amplified and overexpressed at the 3q26.3–q27 amplicon in EAs, EST AW513672 is a 3′ sequence of cDNA clone AF279780, which originally was cloned from a cDNA library made from an NTera2D1 teratocarcinoma cell line that actively expresses full-length L1 RNA (Speek, 2001). L1s are the most abundant mobile elements in the human genome, and their retrotransposition has been associated with genomic instability in transformed cell lines (Ostertag and Kazazian, 2001; Gilbert et al., 2002; Symer et al., 2002). Our data indicate that L1-5′ASP-originated cellular transcripts are not peculiar to cultured cells, for example, in EA cell lines as shown in the present study, but also are present in a subset of primary EAs. We hypothesize that epigenetic changes (i.e., possibly hypomethylation and/or histone acetylation) that occur during EA progression may lead to derepression of the L1-5′ ASP. Hypomethylation of the L1 5′UTR and derepression of L1 expression have been reported in a variety of human malignant cells as well as in primary cancers (Alves et al., 1996; Chalitchagorn et al., 2004). Consistent with this notion, we identified nine CpG islands in this L1-5′ASP. We also demonstrated that two other L1-5′ASP/cellular transcripts within known amplified regions, the L1-5′ASP/MET oncogene and the L1-5′ASP/BCAS3, were found to be differentially expressed in EAs. These findings suggest that derepression of L1-5′ ASP may occur at a broad level in the EA genome and could possibly play a role in EA development. However, the relationship between genomic amplification and L1-5′ASP derepression merits further investigation.

In the present study, we examined specific genetic alterations between EA and the corresponding normal squamous epithelia. Barrett's specimens also were compared with paired normal and tumor samples for genetic alterations in EA development. We and others have provided molecular evidence that Barrett's dysplasia, although premalignant, may lead to EA by direct clonal expansion (this study and Barrett et al., 1999). In addition, our results have demonstrated that sequential alterations of sporadic MSI and genomic amplification occur during EA development. A chimeric L1-5′ASP/cellular transcript also was differentially expressed within this amplified-core region in a subset of EAs. All three distinct alterations oc-
curred concurrently within a 2-Mb region of chromosome band 3q26.3–q27 in EA.

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REFERENCES


