Alternative Polyadenylation Patterns for Novel Gene Discovery and Classification in Cancer

Oguzhan Begik*, Merve Oyken*, Tuna Çinkilli Alıcan*, Tolga Can†‡ and Ayse Elif Erson-Bensan*‡

*Department of Biological Sciences, M.E.T.U., Ankara, 06800, Turkey; †Department of Computer Engineering, M.E.T.U., Ankara, 06800, Turkey; ‡Cancer Systems Biology Laboratory (CanSyL), M.E.T.U., Ankara, 06800, Turkey

Abstract

Certain aspects of diagnosis, prognosis, and treatment of cancer patients are still important challenges to be addressed. Therefore, we propose a pipeline to uncover patterns of alternative polyadenylation (APA), a hidden complexity in cancer transcriptomes, to further accelerate efforts to discover novel cancer genes and pathways. Here, we analyzed expression data for 1045 cancer patients and found a significant shift in usage of poly(A) signals in common tumor types (breast, colon, lung, prostate, gastric, and ovarian) compared to normal tissues. Using machine-learning techniques, we further defined specific subsets of APA events to efficiently classify cancer types. Furthermore, APA patterns were associated with altered protein levels in patients, revealed by antibody-based profiling data, suggesting functional significance. Overall, our study offers a computational approach for use of APA in novel gene discovery and classification in common tumor types, with important implications in basic research, biomarker discovery, and precision medicine approaches.

Introduction

Despite the flow of new information provided by genome and transcriptome sequencing studies, certain aspects of diagnosis, prognosis, and treatment of cancer patients are still important challenges to be addressed. Therefore, a better understanding of the complexity of cancer necessitates characterization of “less obvious but potentially important” changes that we generally fail to detect or consider to be noise in conventional experimental setups. From this perspective, gene expression studies face a key bottleneck; conventional methods are generally not tailored to detect nor quantify 3′ isoforms generated by alternative polyadenylation (APA) [1]. This may negatively impact our ability to discover cancer-related genes and comprehensively understand critical molecular mechanisms underlying disease progression.

APA isoforms are formed as a result of endonucleolytic cleavage of the nascent RNA at alternative poly(A) sites [2]. APA is tightly regulated and is responsive to proliferative, tissue-specific, or developmental cues [3]. APA-generated short or long 3′ untranslated region (UTR) isoforms harbor different cis-elements where microRNAs (miRNAs) and/or RNA-binding proteins bind [4]. Consequently, APA isoforms have different stability, localization, and translation efficiency, all of which significantly modulate protein levels and/or activity. Considering that majority of human genes have multiple poly(A) sites in their 3′-ends [5], APA constitutes an important but less understood layer of complexity in gene expression regulation. Recently, deregulation of APA has gained increasing interest in cancer research because APA emerges as a novel mechanism to activate oncogenes, generally by 3′UTR shortening and loss of repressive cis-elements. For example, 3′UTR shortening of CCND1 (Cyclin D1) mRNA prevents the miRNA-mediated repression and causes further increase in CCND1 levels, which correlate with decreased overall survival of patients [6]. Insulin-like growth factor 2 mRNA binding protein 1 (IGF2BP1) also goes through a shortening

Abbreviations: APA, alternative polyadenylation; poly(A), polyadenylation; miRNAs, microRNAs; 3′UTR, 3′ untranslated region; ER, estrogen receptor; SLR, short to long ratio; SAM, statistical analysis of microarrays; CfsSubsetEval, correlation-based feature selection subset evaluation; BFL, best first list; PCA, principle component analysis; IHC, immunohistochemistry; mRNA, messenger RNA

Address all correspondence to: Ayse Elif Erson-Bensan, Department of Biological Sciences, M.E.T.U., Ankara, 06800, Turkey.
E-mail: erson@metu.edu.tr
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of 3′UTR, and this shorter isoform is associated with profound oncogenic transformation [7]. In addition, our group reported hormone-responsive APA, where estrogen treatment resulted with upregulation and 3′UTR shortening of cell division cycle 6 (CDC6), a major regulator of DNA replication, in breast cancer cells. Thus, as a result of APA, the short CDC6 isoform was linked to higher CDC6 protein levels and increased S-phase entry [8].

While numerous cases of 3′UTR shortening have been linked to increased protein levels and oncogene activation [7], consequences of 3′ UTR shortening on protein levels and functions may be complex. It turns out that 3′UTR shortening may also lead to changes in secondary structure of the mRNA, exposing hidden cis-elements, this time leading to decreased protein levels [9]. In addition, 3′UTR isoforms can have different functions as scaffolds to tether RNA-binding proteins that alter the localization and even function of the translated protein [10]. Alternatively, in cases where proximal poly(A) signals are within introns or coding exons, truncated proteins can be generated with potentially different and/or opposing functions (reviewed in [11]). Hence, APA can contribute to the oncogenic phenotype through various mechanisms [7].

Despite these potential impacts, APA-generated isoforms are generally underdetected simply because we do not look for them in conventional gene expression analyses. Here, we report a meta-analysis pipeline for APA isoform discovery to improve cancer-related gene discovery efforts. Identification of cancer specific APA isoforms is likely to have important implications in basic cancer research and biomarker discovery fields. We anticipate our proposed comprehensive approach to be applicable to other malignancies where expression datasets are available.

Materials and Methods

Datasets

For the discovery datasets, breast, colon, lung, ovarian, and prostate cancer patient data (GSE2109), as part of Expression Project for Oncology from National Center for Biotechnology Information Gene Expression Omnibus (GEO), were utilized. Data for gastric cancer and normal samples (GSE29272) were obtained from GEO. Breast cancer patient data included 318 patients: 69 patients (21.7%) were diagnosed as estrogen receptor negative (ER−), 146 (45.9%) were ER+, and 12 patients (3.8%) were diagnosed with triple-negative breast cancer. Colon cancer patient data included 249 cancer samples: 203 patients (81.5%) were diagnosed with adenocarcinoma, 30 (12%) with mucinous carcinoma, 15 (6%) with carcinoma arising in a villous adenoma, and 1 (0.4%) with signet ring cell carcinoma. Gastric cancer patient data included 134 patients: 62 patients (46%) were diagnosed with cardia adenocarcinoma, and 72 (54%) were diagnosed with noncardia adenocarcinoma. Lung cancer patient data had 105 samples: 32 patients (31%) were diagnosed with squamous cell carcinoma, 29 (28%) with lung adenocarcinoma, and 13 (13%) with bronchioalveolar carcinoma. Ovarian cancer patient data included 166 samples: 28 patients (16.9%) were diagnosed with papillary serous carcinoma, 27 (16.3%) with papillary serous adenocarcinoma, and 15 patients (9%) with endometrioid cancer. Prostate cancer patient data had 73 samples: 63 patients (86%) were diagnosed as acinar type adenocarcinoma and 10 (14%) as adenocarcinoma-NOS.

Detection and Quantification of APA Events

APADetect tool [12] was used to detect and quantify APA events in common cancers. CEL files of Human Genome U133A (HGU133A, GPL96) and U133 Plus 2.0 arrays (HGU133Plus2, GPL570) were analyzed to identify intensities of probes that were grouped based on poly(A) site locations extracted from PolyA_DB [13]. For each transcript, mean signal intensities of proximal and distal probe sets were calculated. The ratio of proximal probe set mean to the distal probe set was called the “short to long” ratio (SLR). SLR values of cancer samples were compared to those of corresponding normal tissue samples. Next, SLR values were further subjected to significance analysis of microarrays (SAM) [14], as implemented by the TM4 Multiple Array Viewer tool [15], for statistical significance after log normalization. A fold change filter further eliminated APA events below a determined threshold (SLR >1.5 for shortening events or SLR <0.66 for lengthening events). SLR values reported in at least 85% of the samples were included in the subsequent analysis and classification pipeline.

Feature Selection

Correlation-based feature selection subset evaluation (CfsSubsetEval) method was used to avoid overfitting and “curse of dimensionality” problems [16,17], as implemented in WEKA data mining software [18]. CfsSubsetEval assessed the performance of a subset of attributes (i.e., SLR values) based on predictive ability and redundancy. The subset space of all the attributes were searched using the BestFirst algorithm with default parameters in WEKA [19]. The attributes were evaluated using 10-fold cross validation. To increase specificity and sensitivity, we selected SLR values that were listed as best attributes in at least 5 of the 10 cross-validations. This group of APA events was identified as best first list (BFL) (Supplementary Tables 1, 2). Heatmap illustration of APA events in BFL was done with a hierarchical clustering implemented in Multiple Array Viewer tool [15]. For the hierarchical clustering based on Pearson correlation coefficient, average linkage-based gene tree with optimized gene leaf order was used as parameter. For a distance-based comparison of the samples, we constructed color-coded gene distance matrices for normal and cancer samples using Pearson correlation coefficient (Supplementary Figure 1).

Random Forest

Random forest classifiers [20,21] were trained using SLR values in BFL. Both the selection of features and training of random forest classifiers were conducted using only the discovery (i.e., training) datasets. The classification accuracy was assessed in independent validation datasets. Use of random forest classifiers was also important for error balancing which can be critical for cancer studies as the number of control samples is usually smaller than the number of cancer samples. Confusion matrix for cancer type analysis was constructed as an output of random forest analysis.

Principle Component Analysis (PCA)

PCA [22] was performed to visualize the SLR-based separation between samples in a lower dimensional space. PCA, as implemented in WEKA, was used with default parameters. Dimensionality reduction was accomplished by choosing the top two principle components in the normal versus cancer separation and top three principle components in the cancer classification. PCA results were then visualized using GraphPad Prism 6 software and Gnuplot (http://gnuplot.sourceforge.net).

Ontology and Network Analysis

Significant APA events (SLRs <0.66 or >1.5) were analyzed by Gene Set Enrichment Analysis (GSEA) (http://www.broadinstitute.org/gsea/index.jsp) [23] and Molecular Signature Database [23]. Network database STRING (http://string-db.org) [24] was used to find potential networks in the APA-regulated transcript lists.
**Immunohistochemistry (IHC) Data**

Antibody-based protein profiling using IHC data from Human Protein Atlas database (http://www.proteinatlas.org) [25] was used to evaluate the protein levels of significant BFL genes. Staining intensities in normal tissues and cancer samples were presented as pathology-based annotation of protein expression levels (low, medium, high).

**Results**

Microarray data can be retrospectively analyzed to discover differential APA isoforms using algorithms based on probe intensities and poly(A) site location information. Earlier, we have developed and successfully implemented a meta-analysis tool for APA isoform discovery: “APADetect” [8,12] (described in “Materials and Methods”). To decipher deregulated APA profiles, we compiled a pool of cancer patient expression data representing six common cancer types (breast, colon, gastric, lung, ovary, and prostate cancers) [26] from the Expression Project for Oncology and National Center for Biotechnology Information GEO databases (Supplementary Table 3). Given that APA is tissue specific [27], independent datasets for corresponding normal tissues were included in the APA detection pipeline (Supplementary Table 4). Overall, we analyzed expression data for a total of 1045 cancer patients (318 for breast, 249 for colon, 134 for gastric, 105 for lung, 166 for ovarian, and 73 for prostate cancers) compared to 479 corresponding normal tissue samples (81 for breast, gastric, 105 for lung, 166 for ovarian, and 73 for prostate samples) for a total of 1045 cancer patients (318 for breast, 249 for colon, 134 for gastric, lung, ovarian, and prostate samples, respectively (Figure 1C, Supplementary Table 7).

To further test the discriminating power of these APA events, random forest classifiers trained in discovery datasets were then applied to an independent validation set (Supplementary Table 3). The true-positive rate for distinguishing cancer samples from their corresponding normal tissues was quite significant using these classifiers. The lowest false-positive rate was 10% (F value, 0.81) in gastric cancer, and the highest was 1.00 in prostate cancers (Supplementary Table 6). For the validation set, PCA showed that the first two principle components explained 45%, 58%, 68%, 61%, 56%, and 74% of the total variance in breast, colon, gastric, lung, ovarian, and prostate samples, respectively (Figure 3B, Supplementary Table 5). Gene distance matrices for normal and cancer samples using Pearson correlation coefficient are given in Supplementary Figure 1.

Using SLR Values for Normal Versus Cancer Discrimination

Following the discovery of global APA profiles, we investigated whether subgroups of identifier APA events can discriminate normal and cancer samples. Therefore, we used feature selection and classification methods available in WEKA data mining software [18]. CfsSubsetEval and random forest methods (see “Materials and Methods” for details) showed that normal and different cancer samples were distinguished using a total of 63 unique attributes (BFL) (Supplementary Tables 1 and 2). In particular, the number of attributes with classifier potential was 16 for breast, 9 for colon, 11 for gastric, 14 for lung, 11 for ovarian, and 13 for prostate. Several APA events (e.g., TOP2A, BGN, RPL13) were redundant in more than one cancer type. Heat maps illustrating color-coded levels of increased or decreased SLR values, representing 3’UTR shortening or lengthening, in cancer patients compared to normal tissues are shown in Figure 3A. Next, we applied PCA using SLR values of transcripts in BFL and produced visualizations in which key APA events between normal tissues and cancer types were represented by a point in the plane formed by two principal axes. In the discovery set, the first two principle components explained 41%, 56%, 68%, 72%, 71%, and 69% of the total variance in breast, colon, gastric, lung, ovarian, and prostate cancer samples, respectively (Figure 3B, Supplementary Table 5). Gene distance matrices for normal and cancer samples using Pearson correlation coefficient are given in Supplementary Figure 1.

Next, because proximal poly(A) site usage is generally associated with higher protein levels due to loss of repressive cis-elements, we asked whether APA isoform variation has any potential effects on resulting protein levels in cancer patients. Therefore, protein levels of the test group of APA events were investigated in the Human Protein Atlas (http://www.proteinatlas.org/cancer) database [25] where protein expression data are derived from antibody-based protein profiling using IHC in normal tissue and cancer patient samples. For
Figure 1. (A) Study workflow. CEL files of microarray data for normal (479) and cancer samples (1045) were processed with the previously developed APADetect tool [12]. U133Plus2 and U133A platforms were simultaneously analyzed through APADetect, increasing potential sample numbers. Poly(A) genomic position information was extracted from PolyA_DB. Probe intensities grouped by poly(A) site positions were processed through probe, intensity, and distal filters, which exclude outliers. LogSLR matrix file is the output file where individual APA events were assigned an SLR. APA events, which were detected in at least 85% of samples, were then run through SAM. To identify the attributes that have the best ability to discriminate between normal and cancer samples, correlation-based feature selection was compiled with BestFirst algorithm and 10-fold cross validation. Significant attributes were used in random forest and principle component analyses for classifying cancers compared to normal tissues and specific cancer types among other cancers. (B) Volcano plots for all APA events detected in six different cancer types compared to corresponding normal tissues. X and Y axes represent SLR of APA events in cancer and normal samples. Red dots represent proximal poly(A) usage, blue dots represent distal poly(A) usage, and gray dots represent insignificant SLR values in cancer patients compared to normal tissue. Bar graphs show the number of APA events.
each cancer type, high SLRs in patients were found to be generally associated with varying degrees of high protein staining compared to normal tissues (Figure 5B). For example, increased proximal polyadenylation pattern of topoisomerase II alpha (TOP2A) was in agreement with high level of staining in breast and lung cancers. Another significant case of APA was detected for ribosomal protein L13 (RPL13) in breast, lung, and ovarian cancers where IHC data also suggested medium to high level of staining in all three types of cancer patients compared to normal tissue. Likewise, WD and tetratricopeptide repeats 1 (WDTC1) with 3′UTR shortening in lung cancers had high level of IHC staining in lung cancer patients.

We then investigated staining patterns of proteins whose transcripts did not appear to be regulated by APA (SLR = 1). A randomly selected group of (SLR = 1) transcripts had similar protein staining patterns in normal and in cancer samples, further strengthening our approach (Supplementary Figure 3).

Interestingly, high-level protein staining was also detected for some cases [e.g., SET nuclear proto-oncogene (SET)], for which 3′ UTR lengthening was observed in cancer cells, suggesting longer isoforms to enhance translation (Supplementary Figure 3A). This is consistent with a previous observation on 3′ UTR shortening to potentiate translational repression due to altered secondary structure of the mRNA, exposing miRNA binding sites only on

Figure 2. Network and ontology enrichment analysis for significant APA events. (A) STRING analysis illustrated multiple clusters, numbered according to GO Biological Process classes shown below. Unconnected proteins are excluded from the illustration. (B) GSEA analysis indicates enriched biological processes with highest log(P) values. FDR q values indicate the false-discovery rate for each biological process.
On the other hand, 3′UTR lengthening was detected in our analysis for proline rich nuclear receptor coactivator 1 (PNRC1), which is a nuclear receptor coactivator that also interacts with GRB2 (an adapter protein involved in growth factor/Ras-mediated signaling pathways) and suppresses GRB2-mediated Ras/MAP-kinase activation. Therefore, PNRC is a potential tumor suppressor candidate, and in fact, its expression is reported to be low in breast cancers [30]. Other cases of distal poly(A) site usage and corresponding protein levels in cancer samples are shown in Supplementary Figure 3A.

We identified another group of APA events where active polyadenylation sites reside in introns or coding exons [e.g., SLU7 homolog, splicing factor (SLU7)] (Supplementary Figure 4A). Selection of these sites suggests APA to be coupled to alternative splicing and lead to production of truncated proteins. In the case of SLU7, activation of the exonic poly(A) site Hs.435342.1.9 leads to a ~70 aa truncation at the C-terminus of the protein. Interestingly, SLU7 itself is a known splicing factor that binds to the C13orf25 primary transcript in which the polycistronic oncomiR miR-17-92 resides [31]. Hence, alterations of the protein length may have functional significance. Another case, a more redundant APA case, was for TOP2A which had increased selection of a proximal poly(A) site within Exon 26, Hs.156346.1.29 (Supplementary Figure 4B). Activation of this coding sequence poly(A) site leads to 384 amino acid truncation at the C-terminus. Therefore, exonic and intronic poly(A) activation cases are of great interest to investigate whether protein activities are altered, specifically when these proteins are being considered as drug targets. It is also important to be aware of these isoforms during the choice of antibodies for detection purposes.

**Discussion**

In cancer cells, deregulated APA can be an important source of isoform diversity which may be overlooked in conventional gene expression analyses. For example, sequencing is becoming the standard method for transcriptome analysis, but random priming and differential PCR amplification lead to read depletion near 3′ ends, negatively affecting APA isoform discovery [32] (also reviewed in [11]). Our proposed approach takes advantage of microarray platforms where probe sets are generally designed from 3′UTRs, making it possible to identify APA events via altered signal intensities near poly(A) sites. A drawback to microarray data use is that only
transcripts with probe sets divided by poly(A) sites are informative. Despite this limitation, an important advantage of our probe-based approach is rapid and simultaneous analysis of publicly available datasets. In addition, our pipeline presented here offers distinct and reproducible results to reveal APA-generated diversity in cancer cells [8,12].

Here, we show cancer-specific APA patterns. Our results indicate altered use of proximal or distal poly(A) sites in cancer transcriptome compared to normal tissue. While some of the APA-regulated genes we identified have already been implicated in tumorigenesis (e.g., TOP2A, EIF5B), a consequential link between APA and how these isoforms may contribute to tumorigenesis is not currently known. In addition, for certain oncogene activation or tumor suppressor inactivation cases, APA may provide insight into how protein levels/functions are altered in cancer cells. For example, shortened or lengthened 3′UTRs may lose or retain binding sites for trans-factors, which may impact the level and/or subcellular localization of the resulting protein. Alternatively, activation of intronic or coding sequence poly(A) sites may generate isoforms that differ at the 3′ends with different coding potentials.

We have to note the possibility that there may be other more proximal or distal poly(A) sites on transcripts; however, either they are not activated in that specific tissue or the probe distribution on microarrays does not allow detection of such poly(A) sites. Therefore, it is possible that APA isoforms may be more widespread in these and other cancer types. Indeed, an earlier work by Xia et al. presented APA isoform diversity in 358 TCGA pan-cancer tumor/normal pairs in

Figure 4. Classification of individual cancer types from a mixed pool of other cancers. (A) Our classifier model consisting of 21 APA events distinguished all 6 cancer types from each other. Confusion matrix demonstrates the performance of the classification model where percentage of correct calls, number of patients, and F values are indicated. (B) PCA showed cancer type discrimination represented by three principal axes (Supplementary Movie 1).

Figure 5. (A). BFL APA events and exemplary log SLR values for proximal and top distal poly(A) site usage cases plotted for each cancer type. Unpaired t test with Welch’s correction was used to compare group means. **** indicates statistical significance (P < .0001). (B) Protein levels detected by IHC in cancer and normal samples are shown. Data for antibody-based protein profiling was extracted from the Human Protein Atlas [25]. Sample size indicates the number of cancer and normal samples scored for staining intensities (low, medium, high) as pathology-based annotation of protein expression levels.
A) 11 APA Events

5' CDS EXON UTR 3'

47 APA Events

5 APA Events

B) Normal Samples
Cancer Samples (Proximal Poly(A) Selection)
Cancer Samples (Distal Poly(A) Selection)

Breast

Colon

Gastric

Lung

Prostate

Ovary

C) Normal High
Normal Medium
Normal Low
Normal Not Detected
Cancer High
Cancer Medium
Cancer Low
Cancer Not Detected

Sample size

Proximal Poly(A) Selection

Breast Colon Gastric Lung Ovary Prostate
a different set of tumor types (bladder urothelial carcinoma, head and neck squamous cell carcinoma, lung squamous cell carcinoma, lung adenocarcinoma, breast invasive carcinoma, kidney renal clear cell carcinoma, uterine corpus endometriosis carcinoma) using an APA-specific algorithm (DaPars) to analyze RNA-seq data [33].

In summary, our approach and results have two significant implications: 1) revealing APA isoform variation in cancer cells may help the discovery of as of yet unknown but potentially important novel cancer-related genes/pathways, and 2) tissue- and cancer type-specific APA may provide novel targets for diagnostic and prognostic purposes, which could help optimize patient outcomes using precision medicine approaches.

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Conflicts of Interest
The authors declare no conflict of interest.

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