Evolutionary Action score of TP53 (EAp53) identifies high risk mutations associated with decreased survival and increased distant metastases in head and neck cancer

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Abstract

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Disclaimers/Conflict of Interest: None
TP53 is the most frequently altered gene in head and neck squamous cell carcinoma (HNSCC) with mutations occurring in over two third of cases, but the prognostic significance of these mutations remains elusive. In the current study, we evaluated a novel computational approach termed Evolutionary Action (EAp53) to stratify patients with tumors harboring TP53 mutations as high or low risk, and validated this system in both in vivo and in vitro models. Patients with high risk TP53 mutations had the poorest survival outcomes and the shortest time to the development of distant metastases. Tumor cells expressing high risk TP53 mutations were more invasive and tumorigenic and they exhibited a higher incidence of lung metastases. We also documented an association between the presence of high risk mutations and decreased expression of TP53 target genes, highlighting key cellular pathways that are likely to be dysregulated by this subset of p53 mutations which confer particularly aggressive tumor behavior. Overall, our work validated EAp53 as a novel computational tool that may be useful in clinical prognosis of tumors harboring p53 mutations.

**Keywords**

Head and neck squamous cell carcinoma; TP53; prognostic biomarker

**Introduction**

Head and neck squamous cell carcinoma (HNSCC) is the 6th most common cancer worldwide and accounts for over 45,000 new cases annually in the United States (1, 2). Since TP53 is the most frequently mutated gene in HNSCC, genomic alterations in this gene are key events in the development and progression of this disease (3–6).

Multiple studies have demonstrated that TP53 mutations are prognostic for poor outcomes in HNSCC, yet molecular testing for TP53 alterations has not become routine in clinical practice (7–11). Although several classifications systems have been described, the main limitation of TP53 as a prognostic biomarker is the lack of a reliable system to accurately assess the functional and clinical impact of specific mutations (10). Whereas most alterations involving tumor suppressor genes render them nonfunctional through truncating mutations or deletions, p53 is unique in that there is a strong selection bias for missense mutations, particularly within the DNA-binding domain. P53 mutation can result in loss of wild type functions through either the loss of DNA-binding activity of p53 responsive elements or a dominant negative effect where the mutated allele binds and inhibits the remaining functional wild-type allele(12). Moreover, some mutant p53 displays oncogenic properties, termed “gain of function” (GOF), which are independent of wild-type p53 function(13). Accordingly, gain of function p53 mutants can enhance cell transformation, increase tumor formation in mice and confer cellular resistance to chemotherapy(14, 15).

While this GOF activity has been well characterized in cancer for five ‘hotspot’ or frequently altered p53 amino acids, 175, 245, 248, 273, and 282, our work indicates that non ‘hotspot’ mutations can also confer GOF activity(16). Therefore we hypothesized that there is a subset of mutations that are particularly deleterious to p53 function resulting in a GOF phenotype and are associated with adverse outcomes in patients with HNSCC.
In an effort to predict which TP53 mutations are highly deleterious, we extended the Evolutionary Trace (ET) approach, an extensively validated method to identify key functional or structural residues in proteins(17). This is achieved by assigning every sequence position a grade of functional sensitivity to sequence variations, defined by whether its evolutionary substitutions correlate with larger or smaller phylogenetic divergences. Residues with large ET grades typically cluster structurally into evolutionary ‘hot-spots’ that overlap and predict functional sites(18). In large scale validation studies, motifs made of top-ranked ET residues predict function in protein structures(19), accurately enough to anticipate enzyme substrates (20).

We have hypothesized that the ET method would assess the impact of TP53 missense mutations. The impact should be greater when the mutated residues are more evolutionarily sensitive to sequence variations, i.e. have a larger ET grade, and also when the amino acid change is least conservative, so the mutational impact is the largest. These two components were computed and combined into a single score, called Evolutionary Action EA(21). This action has been shown to correlate linearly with loss of protein function in test systems and with morbidity in Mendelian diseases, as well as apply across protein coding variations population-wide. To apply this Evolutionary Action to TP53 mutations in HNSCC, we further developed a scoring system (EAp53) to stratify TP53 missense mutations into high and low risk.

The goals of this study were to evaluate the ability of EAp53 to identify a subset of TP53 mutations in HNSCC that are associated with the worst patient outcomes and to validate the impact of these mutations in laboratory based models. We found that EAp53 could identify mutations with GOF phenotypes, termed high risk, that are highly prognostic of poor overall survival, progression-free survival, and the development of distant metastasis in two patient cohorts. Furthermore, high risk mutations were found to be associated with increased cellular invasion, tumorigenicity, and propensity for distant metastases in both in vitro and in vivo models, thereby associating functionally significant TP53 mutations with outcomes of patients harboring high risk TP53 alterations. These oncogenic p53 mutations also had a distinct mRNA expression profile, which suggests high risk p53 mutants regulate unique cellular pathways at the transcriptional level. These findings highlight the need for further evaluation of the EAp53 scoring system as prognostic biomarker in both retrospective and prospective data sets. Finally this study emphasizes the need for continued investigation into the cellular pathways driving the oncogenic phenotype of these high risk p53 mutations which could lead to the identification of novel therapeutics targets and ultimately personalization of cancer treatment based on p53 mutational status.

**Methods**

**Patient Selection and Tissue Procurement for the EAp53 Method**

Training set: A cohort of patients was identified from the TCGA HNSC project that had HPV negative tumors (See supplemental methods for additional details) and underwent surgical resection alone (n=103) or surgery followed by post-operative radiation (n=65). Patient and tumor characteristics along with outcome data were extracted from TCGA HNSC supplemental data.
Validation set: A cohort of patients with HNSCC treated with surgery followed by post-operative radiation at The University of Texas MD Anderson Cancer Center (UTMDACC) from 1992 – 2002 was identified (n=96). Clinical records were reviewed retrospectively and TP53 gene status was determined according to a protocol approved by the institutional review board at UTMDACC. Patients who received chemotherapy were excluded. Clinical and pathologic factors were recorded, including patient age, sex, T and N stage, surgical margin status, and extracapsular lymph node extension (ECE). Survival outcomes including overall and disease free survival along with time to distant metastases were also determined.

DNA Isolation

Samples were isolated using three different methods depending on the platform used to perform the sequencing of TP53. The techniques used either snap-frozen tumor sample or formalin-fixed, paraffin-embedded tissue and the detailed description of the extraction method is included in the supplemental methods.

TP53 Sequencing

As mentioned above three different techniques were utilized to determine TP53 sequence. In each assay, the coding regions and surrounding splice sites from exons 2–11 of the TP53 gene were evaluated via direct sequencing from genomic DNA and a detailed description is included in the supplemental methods.

Calculation of the EAp53 scores

The evolutionary action (EA) scores for each TP53 mutation were calculated based on a simple model of the phenotype-genotype relationship, which hypothesizes that protein evolution is a continuous and differentiable process. Accordingly, the genotype (γ) and the fitness phenotype (φ) will be related by \( \phi = f(\gamma) \), and the phenotypic impact of any mutation at residue \( i \) (evolutionary action) will be the product of two terms: the sensitivity of p53 function to residue variations (\( \partial f / \partial \gamma_i \)) and the magnitude of the substitution (\( \Delta \gamma_i \)). The term \( \partial f / \partial \gamma_i \) was measured by importance ranks of the Evolutionary Trace method (22, 23), according to which, residues that vary amongst closer homologous sequences are ranked less important than those that only vary amongst distant homologous sequences. The magnitude of the substitution (\( \Delta \gamma_i \)) was measured by ranks of amino acid substitution odds(24), however, these odds were computed for different deciles of the evolutionary trace grade at the substituted position. We normalized the product to become percentile scores for p53 protein, for example, an EA score of 68 implied that the impact was higher than 68% of all possible amino acid substitutions in p53.

Statistical Classification by EAp53

Missense mutations were scored by EAp53 from 0–100 with higher scores representing more deleterious alterations. Wildtype p53 (wtP53) sequences were scored as zero since this is the normally functioning protein, see supplemental data for additional description of scoring system. Additionally to obtain scores for individual p53 mutations, an EAp53 server is available at http://mammoth.bcm.tmc.edu/EAp53.
Univariate Cox proportional hazards models were used to estimate hazard ratios and their corresponding *p*-values for all risk factors in the training set. The optimal threshold for EAp53 to stratify patients between favorable and poor outcomes was identified using the training data set, and the *p*-value for the estimated hazard ratio was adjusted (25). The threshold discretizes evolutionary action into low-risk and high-risk patients.

Multivariate Cox proportional hazards model was created with all risk factors except the discretized evolutionary action factor. Rank-based procedure was used based on the Cox proportional hazards model to determine the best threshold for evolutionary action in a multivariate model (26). A Cox proportional hazard model using all risk factors as above along with the discretized evolutionary action factor is created to estimate hazard ratios for each risk factor and their *p*-values. The *p*-value for the discretized evolutionary action factor is adjusted to account for using the data to first determine the cut point. We remove covariates from the model one by one and repeat the above procedure until a model is developed containing risk factors that have hazard ratios which are significant and the *p*-values for the coefficient of the factors in the final model are < 0.05.

Next, the threshold established in the training dataset was applied to the validation dataset to classify TP53 mutations as either low or high risk. Using survival time, univariate Cox proportional hazards model were built for each risk factor using the validation data set to get estimates for hazards ratios and their *p*-values. No adjustments to the *p*-values were necessary. Cox proportional hazard model was also built containing all the risk factors including the EAp53 threshold determined in the test set and reducing the model by removing the covariate with the largest *p*-value in a stepwise manner until the final model only contained significant risk factors.

We used the discretized evolutionary action data to perform log-rank tests to determine differences in time to death between low-risk and high-risk TP53 patients. The above analyses for the validation data set were reproduced with disease-free survival as an outcome and time to metastases as an outcome.

**Site-directed Mutagenesis and TP53 Constructs**

Mutations with a variety of EA scores (Supplementary Table 3) were produced using a QuickChange-II Site-Directed Mutagenesis Kit (Agilent Technologies), starting with wildtype human p53 cDNA with polymorphic region 72 containing arginine previously cloned into a pBabe retroviral expression vector (pBABEpuro, Addgene). Primer sequences used for mutagenesis are shown in Supplementary Table 3. All mutagenesis steps were performed according to the manufacturer’s protocol. Mutations were confirmed by Sanger sequencing at the MD Anderson Cancer Center DNA core facility.

**Generation of the HNSCC Stable Cell Lines**

Two HNSCC cell lines UMSCC1 and PCI13 were selected for their lack of p53 expression due to a splice-site in UMSCC1 (hg19:chr17:7578370C>T) and a deletion in PCI13 (hg19:chr17:7579670_7579709del). UMSCC1 was provided by Dr. Thomas Carey (University of Michigan). PCI13 was acquired from Dr. Jennifer Grandis (University of...
Pittsburgh). The genomic identities of the cell lines were authenticated using short tandem repeat analysis (27). Cells stably expressing TP53 constructs were generated as described previously and a detailed description of the technique is included in the supplemental methods (28).

**Immunoblotting**

Western blotting was performed using standard techniques previously described (27) and primary antibodies to anti-p53 (Santa Cruz Biotechnology, sc-126), anti-p21 (Calbiochem, OP64) and beta-actin (sc81178, Santa Cruz Biotechnology) were used.

**Invasion Assay**

BD BioCoat Matrigel Invasion Assays were used following the manufacturer’s protocol and a detailed description is included in the supplemental methods. Each cell line was run in triplicate. 3T3 NIH cells were used as an invasion control as these cells characteristically migrate but do not invade. The low and high risk represents a composite of three mutations, F134C, A161S, and Y236C and four mutations, R175H, H179Y, C238F, G245D, respectively and the results represent two independent experiments.

**Cell Proliferation Assay**

Cell proliferation was determined using a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay as previously described(29). Briefly, the cells were seeded at different densities and grown in a medium containing 10% FBS in 96-well tissue culture plates. After a 24-hour attachment period, the plates were assigned to different time points to obtain linear optical density (O.D.). Cells were then incubated for 3 hours in medium containing 2% FBS and 0.25 mg/mL MTT, after which the cells were lysed in 200 µL dimethylsulfoxide (DMSO) to release the formazan. The conversion of MTT to formazan was quantified with an EL-808 96-well plate reader (BioTek Instruments, Winooski, VT) set at an absorbance of 570 nm. The OD values were then obtained and analysed to determine the % of cell viability.

**Orthotopic Nude Mouse Model of Oral Cavity Cancer**

All animal experimentation was approved by the Animal Care and Use Committee (ACUC) of the University of Texas MD Anderson Cancer Center. Our orthotopic nude mouse model of oral cavity cancer has been previously validated and a detailed description of the technique is included in the supplemental method (30). The results represent three independent experiments with the low and high risk mutations including two mutations F134C and A161S and four mutations, R175H, H179Y, C238F, and G245D respectively.

**Tail Vein Model**

The experimental metastatic tail vein injection model was performed using standard techniques previously described and a detailed description is included in the supplemental methods (31). The results represent four independent experiments with PCI13 and UMSCC1 cell lines harboring low risk mutations: F134C or A161S, high risk mutations: R175H, C238F or G245D, pBabe, or wtp53.
mRNA expression arrays

Total RNA was isolated from cell lines by using Tri-reagent and hybridized to Affymetrix GeneChip Human Exon 1.0ST Arrays (Affymetrix) according to manufacturer’s instructions and a detailed description is included in the supplemental methods. The one hundred and two p53 target genes were identified and validated through rigorous literature review (L Donehower unpublished data). The difference in expression between pBabe and each of other groups was calculated and heatmaps were generated depicting these expression patterns target gene and 49 selected genes.

Quantitative Reverse Transcription PCR (RT-qPCR) Analyses

Validation of the mRNA expression array for two TP53 target genes (p21 and Notch1) was performed by RT-qPCR. Total RNA was isolated from HNSCC cell lines using Trizol method. Reverse transcription was performed using the high capacity cDNA Reverse Transcription kit (Applied Biosystem) according to the manufacturer’s protocol and a detailed description is included in the supplemental methods.

Results

Patient characteristics and follow-up in the training and validation set

Analysis of the TCGA HNSCC cohort identified 168 patients with non-HPV-associated tumors harboring either missense mutations or wildtype p53 for our training cohort. The majority of these patients presented with advanced T stage (70%) while only 30% of patients presented with advanced neck disease (Supplemental Table 1). The median survival time was 2.96 years and median follow up time was 2.49 years.

Ninety-six patients treated at MD Anderson Cancer Center for HNSCC through surgical resection and post-operative radiation comprised an independent validation set for the EAp53 scoring system. The majority of patients had advanced T (76%) or N stage (54%), and nearly half had aggressive pathologic features (Supplemental Table 1). The median survival time and median follow-up time was 5.49 and 7.66 years respectively.

TP53 mutation in the patient cohorts

One hundred and forty-three TP53 missense mutations were identified in 127 patients in the training set while fifty missense mutations were identified in 47 patients in the validation cohort (Supplementary Table 2 and 3). The majority of these missense mutations were within the DNA-binding domain. Furthermore 34% and 38% of the mutations were located in the well-characterized TP53 hotspot sites, 175, 245, 248, 273, 282 in the training and validation sets, respectively.

Determination of EAp53 Threshold, Risk Models and Survival Analysis

Age, disease site, T-stage, N-stage, and TP53 missense mutations were assessed as risk factors for poor overall survival in the training set. The initial comparison analyzed three groups independently, low EAp53 score, high EAp53 score, and wildtype p53. Univariate analysis in the training set revealed that the low EAp53 score mutations, i.e. low risk, and wild type were not statistically different whereas the high EAp53 score mutations, termed
high risk mutations, appeared to be distinct from the other two groups (Table 1 and Figure 1). Given the similar outcomes, patients with tumors having low risk mutations were combined with wtp53. After univariate analyses an EA threshold of greater than 74.39 identified a group of 82 patients with high risk mutations associated with a significantly decreased overall survival compared to 86 patients in the low risk group, \( p=0.008 \) (Table 1). A rank-based procedure based on the Cox proportional hazards ratios in a multivariate model was then applied to the training set, and identified an EA threshold of 77.78, \( p=0.009 \) (Table 2). Because this model relies on a threshold specific to the TCGA data, the range from 74.39 – 77.78 should contain the valid threshold. Therefore we applied a threshold of 75 to the validation set to determine low and high risk individuals. With this threshold established, this prognostic model was validated in an independent data set and patients with HNSCC harboring high risk mutations \((n=33)\) had significantly decreased overall and disease free survival relative to the low risk group \((n=63)\), \( p=0.030 \) and \( p=0.011 \) (Table 1 and 2). Furthermore this survival difference appeared to be associated with the development of distant metastases, \( p=0.059 \) (Table 2). Additionally, when patients that achieved locoregional control were analyzed, a log rank test revealed tumors with high risk mutations were still associated with increased rate of distant metastases, \( p=.00006 \). Additionally in a multivariate analysis using backward variable selection, the final model identified N stage \(\geq N2b\), \( p=.0064 \), and high risk mutations, \( p=.05 \) as predictors of distant metastases.

Survival analysis using the Kaplan Meier method revealed high risk \( TP53 \) mutations were associated with poor outcomes for overall survival, disease-free survival, and distant metastasis, \( p=0.04, 0.04, \) and \( 0.09 \) respectively (Figure 2a-c).

**High Risk \( TP53 \) mutations are more invasive in vitro and tumorigenic in vivo**

To determine the impact of the high risk \( TP53 \) mutations in pre-clinical models, several low and high risk \( TP53 \) mutant transcripts were stably expressed in two HNSCC cell lines that do not endogenously express p53 due to bi-allelic alterations including somatic mutations (Supplementary Figure 1). Cell lines expressing high risk \( TP53 \) mutations, including R175H, H179Y, C238F, and G245D were significantly more invasive than cells expressing wtp53, p53 null (pBabe), or low risk mutations, A161S, F134C, and Y236C, \( p=0.001, p=0.05, \) and \( p=0.03 \) respectively (Figure 3 Panel A). Assessment of cell motility by scratch assay revealed cells bearing high risk \( TP53 \) mutants, R175H, G245D, and C238F were significantly more motile than cells expressing wtp53, pBabe, or low risk \( TP53 \) mutation, A161S, \( p=0.04, p=0.01, \) and \( p=0.002 \) respectively (Figure 3 Panel B). In contrast to the increased cellular invasion and motility observed in the high risk \( TP53 \) mutations, the rate of proliferation was independent of \( TP53 \) status (Supplemental Figure 2). From these results, we concluded that the expression of high risk mutations is associated with greater migration and invasion, suggesting a gain of function phenotype.

To validate our in vitro data, the isogenic p53 mutant cell lines were introduced into an orthotopic nude mouse tongue cancer model. Although neither the rate nor time of tumor formation were different across various p53 statuses, cell lines that expressed high risk \( TP53 \) mutations formed significantly larger tumors compared to cells that expressed wtp53, pBabe, or low risk p53 mutations (Supplemental Table 4, Figure 3 Panel c and d).
**High Risk TP53 mutations have higher propensity for lung metastases**

To directly evaluate the contribution of high risk p53 mutations in the development of distant metastases, we used the experimental metastatic tail vein injection model. Animals injected with tumor cells carrying high risk mutations, R175H, C238F or G245D, had significantly more grossly visible pulmonary metastases when compared to those injected with cells with low risk p53 mutations, F134C or A161S, p=0.01, pBabe, p=0.008, and wtp53, p=0.004 (Figure 4 Panel a & b). Additionally animals injected with high risk mutant bearing cells had higher numbers of microscopic metastases (Figure 4 Panel c & d).

**High risk p53 mutations have a distinct expression profile of known p53 target genes**

In an effort to identify genes or pathways responsible for the increased tumorigenicity and metastases associated with high risk TP53 mutations, we performed mRNA expression arrays of cells harboring either wtp53, pBabe, low risk mutation, A161S, or high risk mutation, C238F. After standardizing gene expression to the control p53 null cell line, pBabe, an analysis of 102 p53 target genes revealed wtp53 cells had an elevated level of expression in 61% (62/102) of the targets genes whereas the high risk mutant expressing cells suppressed expression in the majority of the same genes. In contrast, the expression pattern of cells expressing a low risk mutation was more similar to that of wtp53 cells as revealed by the hierarchal clustering (Figure 5 Panel a). Upon further analysis of the entire expression array, 49 genes were differentially expressed with wtp53 and the high risk mutation having elevated and suppressed levels of expression, respectively, relative to p53 null, pBabe cells (Figure 5 Panel b). Within these 49 genes, three p53 target genes p21, Notch1 and BTG2 were shown to have this differential expression profile. Quantitative RT-PCR validated the elevated level of expression of these three genes in cells expressing wtp53 or low risk p53 mutations, A161S and Y236C, and the significantly decreased expression in cells bearing high risk mutations, C238F and G245D, relative to cells without p53 expression (Figure 5 Panel c).

**Discussion**

An appropriate method for classifying TP53 mutations and assessing the functional impact of these alterations has been sought for decades, yet a suitable classification scheme remains elusive. Therefore, we have utilized a novel computational method that considers evolutionary variations to predict the functional impact of TP53 mutations. We hypothesized this model could identify missense mutations with the greatest functional impact which would translate to decreased patient survival outcomes. Our analysis of two patient datasets along with in vitro and in vivo models supports TP53 mutational status as a prognostic biomarker in HNSCC.

Currently, TP53 mutational status is not incorporated into the clinical evaluation of HNSCC patients, even though previous reports have identified TP53 mutations associated with poor patient outcomes(8, 10, 11). These previous systems either heuristically stratified mutations based on their potential impact on the p53 molecule and could not be validated in these two cohorts (Supplemental Table 5 and 6 and Supplemental Figure 3 and 4) or stratified TP53 mutations by presence of a copy number loss using a multi-tiered genomic approach(11).
contrast, EAp53 is based on a formal model of evolution which has been shown to perform well against other approaches (17, 21), and has been refined here to specifically evaluate the clinical impact of TP53 mutations. Our results show that patients with HNSCC harboring high risk TP53 mutations had significantly worst overall survival, which was validated in a second patient cohort.

In addition to the development and validation of a novel method for stratifying TP53 mutations, we evaluated several low and high risk p53 mutations in two isogenic HNSCC cell lines in both in vitro and in vivo models. These results corroborated that EAp53 can identify a subset of high risk mutations that are more invasive and tumorigenic than the loss of p53, implying an oncogenic or GOF phenotype. Additionally as seen in the patient cohort, the high-risk mutations had a greater propensity for distant metastases in an experimental metastatic model. Furthermore the low risk mutations were most similar to wtp53 in both in vivo and in vitro models suggesting that these alterations may maintain some residual tumor suppressive functions. While previous classification systems in HNSCC have identified mutations associated with poor outcomes, these studies have not correlated the potential functional impact of these mutations at the cellular level (7, 8, 10). Additionally previous functional studies in yeast that determined the transactivational activity of TP53 mutations, classified most alterations as nonfunctional including many hotspot mutations considered to have GOF properties (32). Given the limitations of these previous studies, we were able to identify patients with tumors harboring mutations associated with decreased survival and confirm these same alterations have oncogenic properties in pre-clinical assays.

Analysis of mRNA expression arrays revealed cells with wtp53 had elevated levels of expression in the majority of TP53 target genes while cells bearing high risk mutations had suppressed levels of expression in these same genes which may partially explain the gain of function properties observed with these mutations. The low risk mutations modulated the expression of TP53 target genes more similarly to wtp53 again, implying some residual wildtype function. Suppressed target gene expression in mutations that have GOF phenotypes corroborates previous studies that have postulated alteration of the p53 transcriptome is a potential mechanism of GOF activities (33, 34).

In conclusion, the EAp53 system appears to identify high risk mutations associated with decreased survival and increased development of distant metastases in HNSCC patients which is corroborated in both in vitro and in vivo studies of invasion, tumorigenicity, and development distant metastases. In order to adopt EAp53 into clinical practice, it will be necessary to confirm the prognostic utility in prospective clinical trials with HNSCC patients managed with the therapeutic standard of care. Furthermore continued investigation into the cellular pathways driving the oncogenic phenotype of these high risk p53 mutations is necessary which could lead to the identification of novel therapeutics targets and ultimately personalization of cancer treatment based on p53 mutational status.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.
Acknowledgments

Funding

This work was supported by the U.T. M.D. Anderson Cancer Center PANTHEON program, the U.T. M.D. Anderson Cancer Center Sisters Institution Network Fund, the National Institute of Health Specialized Program of Research Excellence Grant (P50CA097007), the National Institute of Health (R01 DE14613), Cancer Prevention and Research Institute of Texas (CPRIT) (RP120258), National Research Science Award Institutional Research Training Grant (T32CA60374), the National Institute of Health Program Project Grant (CA06294), and the Cancer Center Support Grant (CA016672). O.L. gratefully acknowledges support from NIH R01 GM079656 and R01 GM066099, and NSF DBI 1356569 and DBI 0851393. PK was supported by the Pharmacoinformatics Training Program of the Keck Center of the Gulf Coast Consortia (NIH Grant No. 5 R90 DK071505).

References


Figure 1. EA\textit{p}53 identifies two functionally distinct groups of p53 mutations

Patients with tumors harboring high risk EA\textit{p}53 (n=83) mutations have a decreased overall survival relative to low risk EA\textit{p}53 (n=44) mutations and \textit{wtp}53 (n=41). In contrast, patients with low risk EA\textit{p}53 mutations appear to have similar survival outcomes to \textit{wtp}53.
Figure 2. EAp53 can identify patients with decreased survival outcomes in HNSCC
Log rank tests of Kaplan Meir survival plots for a cohort of patients treated with surgery followed by radiation therapy validated that EAp53 can identify patients with high risk mutations that are associated with a decreased overall and disease free survival, \( p=0.041 \) and \( 0.036 \) respectively (Panel A and B). Additionally high risk EAp53 mutations appear to be associated with an increased rate of distant metastases \( p=0.092 \) (Panel C).
Figure 3. High Risk TP53 mutations are more invasive in vitro and tumorigenic in vivo
Matrigel invasion assays were performed on PCI13 isogenic HNSCC cell line expressing either high or low risk mutations TP53 mutations, wtp53, pBabe empty vector control. The low and high risk series are a composite of three mutations, F134C, A161S, and Y236C and four mutations, R175H, H179Y, C238F, G245D, respectively and the results represent two independent experiments (Panel A). Scratch assays were performed on PCI13 isogenic HNSCC cell line expressing wtp53, pBabe empty vector control, low risk mutation A161S(shown), or high risk mutations, G245D, R175H, C238F(shown). Percent scratch closure represents two independent experiments and the mean closure is represented for each p53 status. PCI13 and UMSCC1 isogenic cell lines expressing either wtp53, pBabe, low risk mutations, A161S or F134C or high mutations, R175H, H179Y, C238F, or G245D were introduced into the orthotopic model of tongue cancer (Panel C and D).
Figure 4. High risk TP53 mutations are associated with increased development of lung metastases
UMSCC1 and PCI13 isogenic cell lines harboring either wtp53, pBabe, low or high risk mutations were injected into tail vein of nude mice. The low and high risk series are a composite of two mutations, A161S or Y236C and three mutations, R175H, C238F, or G245D, respectively and the results represent two independent experiments. Data labels in columns correspond to number of animals in each group. Macroscopic nodules can be appreciated in the representative images from the lungs of mice top from the pBabe, low and high risk groups (Panel B). The frequency of microscopic lung metastases was assessed by hematoxylin and eosin in animals from the wtp53, pBabe, low risk and high risk groups (Panel C & D).
Figure 5. Suppression of downstream TP53 target genes by high risk TP53 mutations

Heat map depicting expression for 102 TP53 target genes relative to pBabe (Panel A). Analysis of the entire expression array relative to the pBabe identified 49 genes that had significantly different levels of expression in both the wildtype and the high risk mutation. This analysis identified three known TP53 targets genes, p21, Notch1, and BTG2 as significantly overexpressed and under expressed in the wildtype and high risk mutation respectively (Panel B). Quantitative RT-PCR of p21 and Notch1 in wtp53, pBabe, two low risk mutations, A161s and Y236C, and two high risk mutations, C238F and G245D (Panel C). † and ‡ indicate significant overexpression and suppressed expression relative to pBabe respectively.
Table 1

Univariate analysis of overall outcomes in the training and validation sets

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<td>2.25</td>
<td>** 0.008</td>
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<td>121</td>
<td>1.14</td>
<td>(0.69, 1.90)</td>
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<td>N Stage</td>
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<td>52</td>
<td>0.97</td>
<td>(0.59, 1.59)</td>
<td>0.9005</td>
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<td>8</td>
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<td>(0.94, 5.87)</td>
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</table>

*Low risk is defined at EAp53 score of less than 74.39 in the training set and less than 75 in validation set and includes wildtype p53*
Table 2

Cox proportional hazard regression afterward backward selection and adjusting for EA score for overall survival in training set. The threshold of 75 was then set and the same analysis was performed on the validation set for overall survival, disease free survival, and time to distant metastasis in the validation set.

<table>
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<tr>
<th>Dataset</th>
<th>Characteristic</th>
<th>HR</th>
<th>95% CI</th>
<th>p-value</th>
<th>Adjusted for</th>
<th>Model Likelihood Ratio p-value</th>
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<td>EA score ≥77.78</td>
<td>2.30</td>
<td><strong>0.0087</strong></td>
<td>0.0002</td>
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<td><strong>Validation</strong></td>
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<tr>
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<td>EA score ≥75</td>
<td>1.79</td>
<td>(1.06, 3.04)</td>
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<td>(1.16, 3.25)</td>
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<td>N stage ≥2b</td>
<td>0.001</td>
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