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OncoLnc: Linking TCGA survival data to mRNAs, miRNAs, and IncRNAs

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OncoLnc is a tool for interactively exploring survival correlations, and for downloading clinical data coupled to expression data for mRNAs, miRNAs, or IncRNAs. OncoLnc contains survival data for 8,647 patients from 21 cancer studies performed by The Cancer Genome Atlas (TCGA), along with RNA-SEQ expression for mRNAs and miRNAs from TCGA, and IncRNA expression from MiTranscriptome beta. Storing this data gives users the ability to separate patients by gene expression, and then create publication-quality Kaplan-Meier plots or download the data for further analyses. OncoLnc also stores precomputed survival analyses, allowing users to quickly explore survival correlations for up to 21 cancers in a single click. This resource allows researchers studying a specific gene to quickly investigate if it may have a role in cancer, and the supporting data allows researchers studying a specific cancer to identify the mRNAs, miRNAs, and IncRNAs most correlated with survival, and researchers looking for a novel IncRNA involved with cancer lists of potential candidates. OncoLnc is available at http://www.oncolnc.org
OncoLnc: Linking TCGA survival data to mRNAs, miRNAs, and IncRNAs

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Abstract

OncoLnc is a tool for interactively exploring survival correlations, and for downloading clinical data coupled to expression data for mRNAs, miRNAs, or IncRNAs. OncoLnc contains survival data for 8,647 patients from 21 cancer studies performed by The Cancer Genome Atlas (TCGA), along with RNA-SEQ expression for mRNAs and miRNAs from TCGA, and IncRNA expression from MiTranscriptome beta. Storing this data gives users the ability to separate patients by gene expression, and then create publication-quality Kaplan-Meier plots or download the data for further analyses. OncoLnc also stores precomputed survival analyses, allowing users to quickly explore survival correlations for up to 21 cancers in a single click. This resource allows researchers studying a specific gene to quickly investigate if it may have a role in cancer, and the supporting data allows researchers studying a specific cancer to identify the mRNAs, miRNAs, and IncRNAs most correlated with survival, and researchers looking for a novel IncRNA involved with cancer lists of potential candidates. OncoLnc is available at http://www.oncolnc.org.

Main article text

Introduction

The Cancer Genome Atlas (TCGA) provides researchers with unprecedented amounts of molecular data along with clinical and histopathological information (http://cancergenome.nih.gov/). This data set has not only led to increases in our understanding of cancer (Ciriello et al. 2013; Hoadley et al. 2014), but its scale has also allowed for previously impossible projects such as a comprehensive cataloguing of the human transcriptome (Han et al. 2014; Iyer et al. 2015). However, the size and complexity of this unique data set makes it difficult for cancer researchers to access and fully utilize.

Multiple resources exist to help researchers download or explore TCGA data (Cerami et al. 2012; Gyorffy et al. 2013; Koch et al. 2015). Despite this, there is no simple tool that lets users explore the correlation of a gene's expression to survival in multiple cancers at once, provides users the ability to divide patients into any high expressing and low expressing groups for Kaplan-Meier analysis, allows for simple download of the survival data coupled to expression data, and uses modern gene definitions.

In addition, although the role of long noncoding RNAs (IncRNAs) in cancer is beginning to be appreciated (Yarmishyn & Kurochkin 2015), the Tier 3 TCGA mRNA files contain expression data for only the limited number of IncRNAs that were known at the initiation of the TCGA project. As a result, tools for exploring TCGA data will not contain many IncRNAs currently being studied. Although a platform has already been developed to fill this gap (Li et al. 2015), to help the scientific community study IncRNAs OncoLnc incorporates analyses and data for
MiTranscriptome beta lncRNAs, http://mitranscriptome.org/, in addition to Tier 3 TCGA mRNAs and miRNAs.

Materials and methods

Code, files, and software

All of the code necessary to reproduce Tables 1, 2, 3, S1, S2, and S3, and therefore the data in OncoLnc, along with a limited amount of raw data and intermediate files is located at https://github.com/OmnesRes/onco_lnc. The rest of the raw data can be downloaded from https://tcga-data.nci.nih.gov/tcga/ and http://mitranscriptome.org/. This code was run with Python 2.7.5, NumPy 1.7.1, and rpy2 2.5.6, and can require upwards of 6GB of RAM. OncoLnc runs on Django 1.8.2, Python 2.7, matplotlib 1.2.1, NumPy 1.7.1, rpy2 2.5.6, uses the SQLite3 database engine, and utilizes Bootstrap CSS and JavaScript, and Font Awesome icons.

Cox models

The multivariate model run for each cancer and each data type is listed at the top of Tables S1, S2, and S3, and the code for running all of the Cox regressions is present in the GitHub repository. In general only primary solid tumors were included in analyses, and this is implemented by only using samples with "01" in the patient barcode. The exceptions are LAML, which is a blood derived cancer, and therefore has the designation "03", and SKCM, which contains primarily metastatic tumors, and therefore designations "01" and "06" were allowed for SKCM analyses. It is possible for a patient to have more than one sequencing file, and in these cases the counts were averaged. The TCGA data was downloaded on January 5th and 6th, 2016, and miRBase version 21 was used. More info can be found in the GitHub repository and in the text below.

Results

Overview of OncoLnc

OncoLnc stores over 400,000 analyses, which includes Cox regression results as well as mean and median expression of each gene. For the Cox regression results, in addition to p-values, OncoLnc stores the rank of the correlation. Different cancers contain very different p-value distributions (Anaya et al. 2016; Yang et al. 2014), and it is unclear what causes this difference. As a result, using one p-value cutoff across cancers is not possible, and the rank of the correlation is a simple way to measure the relative strength of the correlation. The rank is calculated per cancer, per data type. Tables 1-3 contain information about how many genes there are for each cancer and each data type.

The mRNA and miRNA identifiers used by TCGA are out of date, and the identifiers in OncoLnc have been manually curated using NCBI Gene: http://www.ncbi.nlm.nih.gov/gene, and
recent miRBase definitions: http://www.mirbase.org/. Over 2,000 mRNA symbols were
updated, and these are listed in Table S4. Genes which have had their Entrez Gene ID removed
from NCBI Gene, or could not be confidently mapped to a single identifier, are not included in
OncoLnc but are still included in Table S1.

Using OncoLnc is very straightforward. The preferred method of using OncoLnc is to submit a
gene at the home page, and this submission is not case sensitive. If a user submits a gene not in
the database they will be notified and provided with links to all the possible gene names and IDs.
Submission of a valid gene identifier will return correlation results for up to 21 cancers for
mRNAs and miRNAs, or 18 cancers for MiTranscriptome beta lncRNAs (Fig. 1). If a gene does
not meet the expression cutoff for the analysis, it will not be present in the database, and
therefore a user may receive less than the maximum possible number of results. For users using
OncoLnc on smaller devices, it is possible to perform a single cancer search. The link for this
search is on the home page, and the user must submit the TCGA cancer abbreviation along with
the gene of interest.

At the results page is a link to perform a Kaplan-Meier analysis for each cancer (Fig. 1). The
user will be asked how they would like to divide the patients. Patients can be split into any non-
overlapping upper and lower slices, for example upper 25 percent and lower 25 percent. Upon
submission users will be presented with a PNG Kaplan-Meier plot, a logrank p-value for the
analysis, and text boxes with the data that was plotted (Fig. 2). If a user simply wants all the data
for that cancer and that gene, the user can submit 100 for "Lower Percentile", and 0 for "Upper
Percentile".

Users then have the option to either go to a PDF of the Kaplan-Meier plot, or download a CSV
file of the data plotted. In both cases the file name will be the cancer, gene ID, lower percentile,
upper percentile, separated by underscores. Gene ID had to be used instead of gene name
because there are multiple HUGO gene symbol conflicts between TCGA Tier 3 mRNAs and
MiTranscriptome beta, as well as between TCGA mRNA HUGO gene symbols and updated
mRNA HUGO gene symbols. In the case that a user performs a search for a name with a
conflict, OncoLnc presents a warning message and instructs the user how to proceed.

mRNAs

Table 1 contains information about the patients for each Tier 3 mRNA study included in
OncoLnc, and how many gene analyses are present in OncoLnc for each study. Tier 3
RNASeqV2 was used for all 21 cancers, and expression was taken from the
"rsem.genes.normalized_results" files. As a result, the expression data in OncoLnc for Tier 3
mRNAs is in normalized RSEM values. Table 1 contains different numbers of genes for the
different cancers because an expression cutoff was used to determine if a gene would be included
in the analysis. For mRNAs this cutoff was a median expression greater than 1 RSEM, and less
than a fourth of the patients with an expression of 0.
The results of every Tier 3 mRNA Cox regression performed are included in Table S1. The Tier 3 expression files contain both a HUGO gene symbol and Entrez Gene ID for each gene, but these IDs and gene symbols are not current. To update the gene symbols I downloaded every human gene from NCBI Gene, and updated any symbol for which the Entrez Gene ID was still current. For genes that had deleted or changed Entrez Gene IDs I had to manually curate the IDs and symbols along with the updated names and symbols, and Table S4 lists genes which had either the symbol or ID changed. OncoLnc allows users to search mRNAs using either an updated HUGO gene symbol or Entrez Gene ID.

**miRNAs**

Table 2 contains information about the patients for each Tier 3 miRNA study included in OncoLnc, and how many gene analyses are present in OncoLnc for each study. Tier 3 miRNASeq was used for every cancer except GBM, which only had microarray data available. The results of every Cox regression performed are included in Table S2. Many of the miRBase IDs, or possibly read counts, present in Table S2 and OncoLnc will be different from the IDs and read counts in TCGA data files and available at other data portals for TCGA data. This is because I went through each expression file and updated the IDs and read counts. The "isoform.quantification" files contain both miRBase IDs as well accession numbers. In these files the 5p and 3p arms of miRNAs are referred to with the same ID, for example hsa-let-7b-5p and hsa-let-7b-3p would both be listed as hsa-let-7b. In order to update the names and read counts for the Tier 3 miRNAs I used the read counts assigned to each accession number to obtain reads per million miRNAs mapped for each accession number, and updated the ID with the current miRBase ID. When an accession number was not available I used the genomic coordinates provided to identify the accession number, and therefore ID. GBM names were updated using the "aliases" file from the miRBase FTP site, and if an alias could not be confidently identified the miRNA was not included in OncoLnc, but is still in Table S2.

As a result, all expression values in Table S2 and in OncoLnc are reads per million miRNA mapped for every cancer except GBM, which are microarray normalized values. The numbers of miRNAs in Table 2 differ because the miRNA may not have been in the expression files for that cancer, or may not have met the expression cutoff. An expression cutoff of a median of .5 reads per million miRNA mapped, and less than one fourth of the patients with 0 expression was used. OncoLnc allows users to search for miRNAs with either a miRBase version 21 mature accession number or ID.

**lncRNAs**

Table 3 contains information about the patients for each MiTranscriptome beta lncRNA analysis, along with how many lncRNAs are included in OncoLnc for each cancer. Normalized lncRNA
counts were downloaded from http://mitranscriptome.org/, and these were mapped to patient barcodes using the library information provided. MiTranscriptome beta contains over 8,000 of the most differentially expressed IncRNAs in the entire MiTranscriptome dataset, but the actual number of IncRNAs in OncoLnc for each cancer is far fewer due to the expression cutoff used: a median of .1 normalized counts, and less than a fourth of patients with 0 expression. Table S3 contains every IncRNA Cox regression performed, and these are all included in OncoLnc. OncoLnc allows users to search for MiTranscriptome beta IncRNAs using either a name or transcript ID.

**Discussion**

Depending on the researcher, OncoLnc should be used in different ways. If a researcher is studying a specific gene and looking for a cancer association, they should go to http://www.oncolnc.org and perform a search with their gene of interest. Instead of focusing on p-values, I would focus more on the rank of the correlations for the different cancers, and also on the sign of the Cox coefficients. A positive Cox coefficient indicates high expression of the gene increases the risk of death, while a negative Cox coefficient indicates the opposite. A gene with a high rank in multiple cancers (indicated by a low number, 1 being the best), and Cox coefficients with the same sign could be very interesting. It is also important to look at the level of expression of the gene. Different genes obviously require different levels of expression to exert their effects, but genes with expression near 0 should be dealt with caution. In addition, users can investigate the range of expression of the gene at the Kaplan-Meier plotting page. Genes that have large fold increases from the low expression to high expression group could be interesting candidates.

A researcher studying a specific cancer should download Tables S1, S2, and S3 to see which mRNAs, miRNAs, and IncRNAs are most correlated to survival for their cancer. Once they identify some genes of interest they can go to http://www.oncolnc.org to perform further analyses such as checking the range of expression of the gene, or if it is associated with survival in other cancers. Similarly, bioinformaticians looking to perform large scale analyses of prognostic genes can use these tables as a starting point, or if a user wants to change the Cox models they can use the GitHub code to alter the models.

The importance of the ability to perform survival correlations with IncRNAs must be emphasized. There are multiple techniques for identifying protein coding genes that are involved in cancer because mutations that occur in protein coding genes can result in missense mutations, and methods have been developed for identifying which of these mutations are drivers as opposed to simply passengers (Carter et al. 2009; Kaminker et al. 2007; Youn & Simon 2011). In contrast, because it is unclear how mutations will affect IncRNA function, methods to identify IncRNAs involved in cancer must rely on IncRNA expression. As a result, OncoLnc is one of the few resources available for finding IncRNAs involved in cancer, and if a IncRNA researcher is searching for a novel IncRNA to study, Table S3 would be a good place to start.
When using OncoLnc it is important to remember that the correlations observed, regardless of p-value, are still only correlations. Perhaps the largest limitation of OncoLnc is that the Cox models do not account for intra-cancer subtypes. For example, GBM and BRCA both have well-established subtypes (Brennan et al. 2013; Perou et al. 2000). If the expression of a gene correlates with cancer subtypes, and those subtypes correlate with survival, subtype would be a confounding variable. As subtype definitions for the different cancers improve a future version of OncoLnc may be able to incorporate the subtypes in the Cox models.

An analysis is only as good as the data available, and the Tier 3 TCGA RNA-SEQ analyses were performed with outdated software and transcript information. There have been some attempts to reanalyze both the TCGA mRNA RNA-SEQ data and miRNA-SEQ data (Kuo et al. 2015; Rahman et al. 2015). In the event that TCGA or the scientific community releases a gold standard analysis of TCGA data, a future version of OncoLnc could incorporate this data.

Current data portals for TCGA data only allow users to view the results for one cancer at a time, may or may not offer Cox regression results, do not allow for complete control over separating patients during Kaplan-Meier analysis, and do not allow for download of the data used in the analysis. To my knowledge OncoLnc is the only online resource for TCGA data that includes these features, is the only resource that uses modern gene definitions for TCGA mRNA and miRNA data, and is the only resource for survival analysis of MiTranscriptome beta lncRNAs. In addition, current methods for survival analysis rely on a p-value cutoff of .05 for significance, which may lead to either the study of genes not actually correlated with survival or missing genes that are correlated with survival depending on the cancer. By storing the results of the correlation for every gene, OncoLnc can provide a context for the significance of a correlation. As a result, used correctly OncoLnc can not only increase the sensitivity of finding genes involved in cancer, but also the specificity. This combination of ease of use, results for complex analyses, and tools for exploring and downloading data make OncoLnc an invaluable resource for cancer researchers.

Additional Information and Declarations

Competing Interests

I offer services at omnesres.com which may involve web development or analysis of TCGA data.

Author Contributions

I conceived the project, downloaded and analyzed all data, developed OncoLnc, wrote the manuscript, and maintain the GitHub repository and any associated material at omnesres.com.

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Table 1 (on next page)

Characteristics of the Tier 3 mRNA datasets in OncoLnc

Age at diagnosis is in years, and is an average. The events indicate the number of deaths in the dataset. Median survival is in days and could not be calculated for COAD, KIRP, OV, READ, and UCEC.
| Cancer | Patients | Male/Female | Age at Diagnosis | Events | Median Survival | Genes in OncoLnc |
|--------|----------|-------------|------------------|--------|----------------|-----------------|
| BLCA   | 403      | 296/107     | 68.03            | 177    | 1008           | 16339           |
| BRCA   | 1006     | 11/995      | 58.34            | 135    | 3941           | 16602           |
| CESC   | 264      | 0/264       | 48.23            | 59     | 4086           | 16330           |
| COAD   | 440      | 235/205     | 66.58            | 85     | NA             | 16378           |
| ESCA   | 144      | 126/18      | 60.51            | 59     | 801            | 16790           |
| GBM    | 152      | 99/53       | 59.84            | 27     | 1426           | 16783           |
| HNSC   | 497      | 364/133     | 61.24            | 207    | 1732           | 16614           |
| KIRC   | 523      | 341/182     | 60.56            | 167    | 2764           | 16638           |
| KIRP   | 285      | 210/75      | 61.45            | 44     | NA             | 16399           |
| LAML   | 151      | 81/70       | 54.40            | 92     | 577            | 15227           |
| LGG    | 510      | 282/228     | 43.02            | 124    | 2835           | 16781           |
| LIHC   | 360      | 244/116     | 59.41            | 126    | 1694           | 15824           |
| LUAD   | 492      | 225/267     | 65.32            | 176    | 1492           | 16748           |
| LUSC   | 489      | 362/127     | 67.23            | 169    | 2224           | 16942           |
| OV     | 294      | 0/294       | 59.19            | 42     | NA             | 16893           |
| PAAD   | 175      | 96/79       | 64.37            | 92     | 607            | 17177           |
| READ   | 159      | 88/71       | 64.58            | 22     | NA             | 16472           |
| SARC   | 259      | 118/141     | 60.71            | 98     | 1991           | 16197           |
| SKCM   | 459      | 284/175     | 58.14            | 215    | 2454           | 16030           |
| STAD   | 379      | 247/132     | 65.49            | 146    | 1043           | 16885           |
| UCEC   | 541      | 0/541       | 63.95            | 90     | NA             | 16670           |
Table 2 (on next page)

Characteristics of the Tier 3 miRNA datasets in OncoLnc

Age at diagnosis is in years, and is an average. The events indicate the number of deaths in the dataset. Median survival is in days and could not be calculated for COAD, KIRP, READ, and UCEC.
| Cancer | Patients | Male/Female | Age at Diagnosis | Events | Median Survival | Genes in OncoLnc |
|--------|----------|-------------|------------------|--------|----------------|------------------|
| BLCA   | 404      | 297/107     | 68.02            | 177    | 1036           | 512              |
| BRCA   | 988      | 11/977      | 58.35            | 131    | 3941           | 479              |
| CESC   | 267      | 0/267       | 48.27            | 59     | 4086           | 501              |
| COAD   | 426      | 226/200     | 66.48            | 84     | NA             | 476              |
| ESCA   | 144      | 125/19      | 60.61            | 59     | 801            | 494              |
| GBM    | 561      | 343/218     | 57.94            | 67     | 2648           | 507              |
| HNSC   | 501      | 363/138     | 61.30            | 208    | 1732           | 514              |
| KIRC   | 506      | 331/175     | 60.48            | 165    | 2764           | 448              |
| KIRP   | 286      | 210/76      | 61.52            | 44     | NA             | 430              |
| LAML   | 164      | 88/76       | 54.05            | 100    | 518            | 374              |
| LGG    | 506      | 278/228     | 43.07            | 123    | 2660           | 486              |
| LIHC   | 362      | 248/114     | 59.41            | 125    | 1791           | 485              |
| LUAD   | 490      | 226/264     | 65.35            | 175    | 1492           | 493              |
| LUSC   | 467      | 346/121     | 67.43            | 160    | 2224           | 519              |
| OV     | 470      | 0/470       | 59.85            | 92     | 3128           | 467              |
| PAAD   | 175      | 96/79       | 64.37            | 92     | 607            | 494              |
| READ   | 154      | 84/70       | 64.23            | 22     | NA             | 495              |
| SARC   | 259      | 119/140     | 60.85            | 98     | 1991           | 455              |
| SKCM   | 438      | 271/167     | 58.01            | 207    | 2470           | 535              |
| STAD   | 400      | 260/140     | 65.54            | 155    | 1043           | 495              |
| UCEC   | 534      | 0/534       | 63.91            | 87     | NA             | 518              |
Table 3 (on next page)

Characteristics of the MiTranscriptome beta analyses in OncoLnc

Age at diagnosis is in years, and is an average. The events indicate the number of deaths in the dataset. Median survival is in days and could not be calculated for COAD, KIRP, and UCEC.
| Cancer | Patients | Male/Female | Age at Diagnosis | Events | Median Survival | Genes in OncoLnc |
|--------|----------|-------------|------------------|--------|----------------|-----------------|
| BLCA   | 120      | 86/34       | 67.37            | 61     | 706            | 4322            |
| BRCA   | 766      | 8/758       | 58.03            | 111    | 3941           | 4708            |
| CESC   | 106      | 0/106       | 48.22            | 26     | 3046           | 4493            |
| COAD   | 117      | 52/65       | 69.64            | 24     | NA             | 3302            |
| GBM    | 144      | 94/50       | 59.56            | 24     | 1426           | 4524            |
| HNSC   | 288      | 211/77      | 61.40            | 133    | 1762           | 4314            |
| KIRC   | 457      | 299/158     | 60.75            | 156    | 2764           | 5191            |
| KIRP   | 73       | 51/22       | 59.78            | 17     | NA             | 4627            |
| LAML   | 20       | 15/5        | 54.75            | 10     | 580            | 3940            |
| LGG    | 217      | 123/94      | 42.82            | 65     | 2660           | 4875            |
| LIHC   | 65       | 40/25       | 60.97            | 41     | 1005           | 3610            |
| LUAD   | 320      | 148/172     | 65.72            | 118    | 1357           | 4636            |
| LUSC   | 330      | 244/86      | 67.16            | 112    | 2284           | 4979            |
| OV     | 369      | 0/369       | 59.63            | 69     | 3128           | 4901            |
| READ   | 42       | 22/20       | 66.67            | 8      | 1581           | 3310            |
| SKCM   | 255      | 159/96      | 56.82            | 148    | 2192           | 3893            |
| STAD   | 148      | 93/55       | 65.72            | 56     | 940            | 4619            |
| UCEC   | 274      | 0/274       | 63.12            | 39     | NA             | 3706            |
Example of OncoLnc search results

The Cox coefficient and p-value are from the gene term in precomputed multivariate Cox regressions. The FDR correction is performed per cancer analysis per data type, and in this example the correction would have involved around 16,000 genes for each cancer. The rank is also performed per cancer per data type. In this example DONSON is the 3rd most highly correlated gene in KIRC.

| Cancer | Cox Coefficient | P-Value | FDR Corrected | Rank | Median Expression | Mean Expression | Plot Kaplan? |
|--------|-----------------|---------|----------------|------|-------------------|----------------|--------------|
| BLCA   | -0.005          | 9.50e-01| 9.79e-01       | 15844| 410.38            | 478.94         | Yes Please!  |
| BRCA   | 0.192           | 2.80e-02| 3.08e-01       | 1494 | 268.08            | 341.28         | Yes Please!  |
| CESC   | -0.152          | 2.50e-01| 5.95e-01       | 6770 | 665.25            | 681.12         | Yes Please!  |
| COAD   | -0.123          | 2.40e-01| 5.44e-01       | 7106 | 279.33            | 296.04         | Yes Please!  |
| ESCA   | -0.123          | 3.60e-01| 9.77e-01       | 6053 | 334.31            | 379.24         | Yes Please!  |
| GBM    | -0.224          | 3.20e-01| 9.62e-01       | 5406 | 352.85            | 389.29         | Yes Please!  |
| HNSC   | 0.009           | 9.00e-01| 9.64e-01       | 15462| 275.38            | 330.77         | Yes Please!  |
| KIRC   | 0.574           | 8.00e-12| 4.45e-08       | 3   | 148.77            | 169.63         | Yes Please!  |
| KIRP   | 0.588           | 9.90e-05| 1.95e-03       | 813  | 187.15            | 209.66         | Yes Please!  |
| LAML   | 0.234           | 5.50e-02| 4.05e-01       | 2051 | 381.27            | 401.45         | Yes Please!  |
| LGG    | 0.333           | 4.10e-04| 1.60e-03       | 4393 | 203.47            | 230.77         | Yes Please!  |
| LIHC   | 0.188           | 5.50e-02| 2.32e-01       | 3738 | 300.61            | 354.61         | Yes Please!  |
| LUAD   | 0.180           | 2.30e-02| 1.29e-01       | 2946 | 253.81            | 295.14         | Yes Please!  |
| LUSC   | -0.09           | 2.40e-01| 8.35e-01       | 4762 | 329.73            | 380.92         | Yes Please!  |
| OV     | -0.028          | 8.50e-01| 9.92e-01       | 14385| 253.27            | 303.44         | Yes Please!  |
| PAAD   | 0.05            | 6.40e-01| 7.93e-01       | 13828| 187.31            | 196.42         | Yes Please!  |
| READ   | -0.168          | 4.30e-01| 9.24e-01       | 7545 | 269.13            | 278.58         | Yes Please!  |
| SARC   | 0.255           | 2.20e-02| 1.54e-01       | 2273 | 391.13            | 470.87         | Yes Please!  |
| SKCM   | 0.146           | 3.80e-02| 1.51e-01       | 3999 | 348.03            | 375.7          | Yes Please!  |
| STAD   | -0.059          | 4.90e-01| 8.02e-01       | 10245| 304.86            | 327.4          | Yes Please!  |
| UCEC   | 0.009           | 9.30e-01| 9.95e-01       | 15404| 278.83            | 315.72         | Yes Please!  |
Example of OncoLnc Kaplan-Meier results

A) Submitting non-overlapping percentiles will return a logrank p-value for the analysis and a PNG image with the option to generate a PDF of the plot. B) Below the Kaplan-Meier image will be the data that was plotted along with an option to download a csv file.
