Machine learning-based investigation of the cancer protein secretory pathway

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Abstract

Deregulation of the protein secretory pathway (PSP) is linked to many hallmarks of cancer, such as promoting tissue invasion and modulating cell-cell signaling. The collection of secreted proteins processed by the PSP, known as the secretome, is often studied due to its potential as a reservoir of tumor biomarkers. However, there has been less focus on the protein components of the secretory machinery itself. We therefore investigated the expression changes in secretory pathway components across many different cancer types. Specifically, we implemented a dual approach involving differential expression analysis and machine learning to identify PSP genes whose expression was associated with key tumor characteristics: mutation of p53, cancer status, and tumor stage. Eight different machine learning algorithms were included in the analysis to enable comparison between methods and to focus on signals that were robust to algorithm type. The machine learning approach was validated by identifying PSP genes known to be regulated by p53, and even outperformed the differential expression analysis approach. Among the different analysis methods and cancer types, the kinesin family members KIF20A and KIF23 were consistently among the top genes associated with malignant transformation or tumor stage. However, unlike most cancer types which exhibited elevated KIF20A expression that remained relatively constant across tumor stages, renal carcinomas displayed a more gradual increase that continued with increasing disease severity. Collectively, our study demonstrates the complementary nature of a combined differential expression and machine learning approach for analyzing gene expression data, and highlights key PSP components relevant to features of tumor pathophysiology that may constitute potential therapeutic targets.

Author summary

The secretory pathway is a series of intracellular compartments and enzymes that process and export proteins from the cell to its surrounding environment. Dysfunction of the
secretory pathway is associated with many diseases, including cancer, and therefore constitutes a potential target for novel therapeutic strategies. The large number of interacting components that comprise the secretory pathway pose a challenge when attempting to identify where the dysfunction originates or how to restore healthy function. To improve our understanding of how the secretory pathway is changed within tumors, we used gene expression data from normal tissue and tumor samples from thousands of individuals which included many different types of cancers. The data was analyzed using different machine learning algorithms which we trained to predict sample characteristics, such as disease severity. This training quantified the relative degree to which each gene was associated with the tumor characteristic, allowing us to predict which secretory pathway components were important for processes such as tumor progression—both within specific cancer types and across many different cancer types. The machine learning-based approach demonstrated excellent performance compared to traditional gene expression analysis methods and identified several secretory pathway components with strong evidence of involvement in tumor development.

1. Introduction

One of the most challenging features in diagnosing and treating cancer is its heterogeneity—the tissue of origin, gene mutation profile, patient, and local tumor environment are just a few of the many factors that can affect the pathophysiology and response to treatment of a particular cancer [1]. However, a core set of features exhibited by cancer cells establish a common thread despite other variations. Many of these shared features have been distilled into a set of “cancer hallmarks”, such as resisting cell death, activating invasion and metastasis, and avoiding immune destruction [2]. Furthermore, tumor cells acquire and sustain many of these hallmarks through interactions with each other and with neighboring “normal” cells, which together with the cancer cells form the tumor microenvironment [3]. An important system that links tumor cells to each other and to the microenvironment is the protein secretory pathway (PSP) [4]. Secreted and membrane proteins processed by the PSP contribute to critical tumor functions, such as facilitating communication among different cells residing in the microenvironment (and even with distant tissue sites in the body), and for construction and turnover of the tumor extracellular matrix. Collectively, these functions support a key role for the PSP in cancer physiology, making it an attractive target for potential therapeutic approaches.

Advancements in high-throughput molecular profiling technologies such as transcriptomics and proteomics have enabled extensive investigation and characterization of the human secretome [5] and its changes during the onset and progression of diseases such as cancer [6,7]. Although many components of the PSP that drive these important secretome changes have been studied individually, an investigation of how these constituents behave together as a system is lacking, particularly in the context of cancer. Recent efforts have begun to elucidate this system by exploring how PSP expression patterns compare to those of the secretome among different human tissues [8], and by developing genome-scale reconstructions of the PSP to mechanistically link these characteristics to the metabolic network [9]. We sought to further extend the investigation of the PSP through the application of machine learning (ML) approaches.

The efficacy of ML-based approaches in the investigation of omics datasets has been demonstrated in a number of recent studies [10–14]. For example, van IJzendoorn and colleagues
applied a random forest algorithm to three gene expression databases (TCGA, GTEx, and the French Sarcoma Group) to identify novel diagnostic markers for soft tissue sarcoma, which was validated with qRT-PCR in an independent experiment [11]. In another study, Wood and colleagues used L1-regularized logistic regression (Lasso) to develop a classifier for nonalcoholic fatty liver disease (NAFLD) based on phenotypic, genomic, and proteomic features [10]. Giordano and colleagues merged the results of three different approaches—Lasso regression, support vector machine (SVM) with recursive feature elimination, and a collection of randomized decision trees (Extra Trees)—to identify robust gene signatures for cigarette exposure, which harbor potential for biomarker development [15]. Furthermore, the MLSeq R package was developed to facilitate the use of over 90 different ML algorithms for the analysis of RNA-seq or microarray data, enabling the generation of classification models and identification of potential biomarkers [16].

We applied differential expression (DE) analysis and 8 different ML methods on RNA-seq data from The Cancer Genome Atlas (TCGA) to identify genes encoding PSP machinery that are associated with clinical features including cancer status, tumor stage, and mutation profile. The classification performance of the ML algorithms was evaluated for each of the clinical features, and relevant PSP genes were identified by DE analysis and compared with those identified by ML. The analyses reveal PSP components that exhibit pan-cancer and cancer-specific roles, and demonstrate the complementarity of DE and ML methods in the analysis of omics data.

2. Results

2.1 Data retrieval and definition of PSP genes

We retrieved 11,053 RNA-seq samples and 9,375 mutation profiles from TCGA, spanning 10,198 individuals and 33 cancer types (S1 Table). Our analysis was focused on the subset of 575 genes encoding and/or regulating the human PSP machinery, as defined in the study by Feizi et al. [8]. Feizi and colleagues defined the PSP genes through literature surveys, KEGG pathway annotation information, and orthologs from a model of yeast protein secretion [17]. These 575 PSP genes encode for secretory processes such as folding, glycosylation, and trafficking, as well as protein-related stress responses (e.g., the unfolded protein response). We limited our study to this subset of genes to specifically investigate the behavior of the PSP in different cancer types, and infer which components (if any) appear to have a more pronounced role or association with sample characteristics, such as cancer status or tumor stage. This focused approach also served to reduce the feature space for the application of ML methods, which often face challenges when the number of features greatly exceeds the number of samples.

2.2 ML-based gene scoring

We implemented a gene scoring approach (Fig 1) whereby samples were grouped according to a known binary variable of interest (such as normal vs. tumor), and a ML classifier was trained to predict the group (class) of each sample based on the expression of its PSP genes. Classifiers were trained using 10-fold cross validation, and prediction performance was quantified by area under the receiver operating characteristic (ROC) curve (ROC AUC). The resulting feature importance scores of the trained classifier, which quantify roughly how useful each gene is in predicting sample class, were normalized by taking the absolute value and scaling to a range of 0 to 1. A consensus score for each gene was computed as the average score across the 8 different ML algorithms.

The ML algorithms used in the present study were random forests [18], extremely randomized trees (ExTrees) [19], adaptive boosting (AdaBoost) [20], extreme gradient boosted trees
(XGBoost) [21], linear discriminant analysis (LDA), Lasso regression [22], Ridge regression, and support vector machine [23]. We did not seek to include a comprehensive coverage of the available ML algorithms, as this would be infeasible and beyond the scope of the study. The algorithms were selected to include some of the most commonly employed methods for biological data [24,25], and to span different classes such as ensemble learning (random forests, ExTrees), boosting (AdaBoost, XGBoost), regularized logistic regression (lasso and ridge regression), and other common linear classifiers (LDA and SVM). Furthermore, algorithms were limited to those for which feature importance scores could be calculated.

All classification analyses were performed within individual cancer types before comparing or averaging with the results of other cancer types. This approach was implemented to avoid tissue-specific features that would likely dominate gene expression changes when including samples from different tissues of origin in the same analysis. In this way, we could effectively eliminate differences associated with tissue type, and instead highlight those driven by a class variable of interest, such as cancer status or tumor stage.

2.3 Mutation of tumor protein 53

We first sought to validate our ML gene scoring approach using a class variable for which the associated gene(s) are well-established. A mutation in the TP53 gene (encoding the p53 protein) is one of the most common mutations observed in human cancers, and the resulting loss or change in its activity as a tumor suppressor contributes to malignant progression [26]. Since p53 and its regulatory targets have been extensively characterized [27], we began our investigation with p53 mutation status as the class variable by which to group samples (non-mutated vs. mutated TP53). Genes known to be regulated by p53 are likely to exhibit a change in expression if p53 is mutated, and can therefore be used as positive controls to assess whether the scores from our ML approach are biologically meaningful. Of the 575 PSP genes considered in the study, 4 are known to be direct targets of p53 regulation: BCL2 Associated X (BAX), Heat Shock Protein Family A Member 4 Like (HSPA4L), Kinesin Family Member 23 (KIF23), and BCL2 Antagonist/Killer 1 (BAK1) [27–29]. It is therefore expected that an effective approach
should be able to identify some or all of these 4 genes as being associated with p53 mutation status.

Mutation data for the TP53 gene in TCGA subjects was obtained from whole-exome sequencing data and aligned with the RNA-seq data, enabling the classification of tumor RNA-seq samples in each cancer type as “mutated” or “non-mutated” in TP53. Cancer types with fewer than 10 samples in each class were discarded, leaving a total of 22 different cancer types. Each of the 8 ML algorithms were trained on the data to predict p53 mutation status based on PSP gene expression, and the resulting gene scores (S2 Table) and ROC AUC values (S3 Table) were calculated. In addition, DE analyses were performed between mutated and non-mutated samples for each cancer type, yielding a log2 fold-change and associated significance (p-value; adjusted for the false discovery rate (FDR)) for each PSP gene (S4 Table).

The consensus ML gene scores were averaged across all cancer types to identify genes that were generally associated with the p53 mutation (Fig 2A and 2B). The top three genes were BAX, HSPA4L, and KIF23—three out of the four direct regulatory targets of p53—thus providing support for the biological relevance of the ML gene scores. Although the four p53 target genes were significantly differentially expressed (p53 mutated vs. non-mutated) in many of the cancer types, only KIF23 was among the top three when averaging DE gene scores across all cancers, and BAX in the top ten (S1 Fig). The ML gene scoring approach thus outperformed the DE method in identifying the PSP genes directly associated with the p53 mutation status.

Another gene exhibiting a high average ML score across cancer types was ASAP3. Although we did not find any literature supporting its direct regulation by p53, ASAP3 has been identified to promote cell migration and invasion by destabilizing cytoskeletal protein ACTG1. It is therefore possible that ASAP3 is indirectly affected by the activity of p53 as part of a broader tumorigenic program, or is simply correlated with the mutation.

Despite the excellent performance of the ML scoring approach in identifying relevant genes, this was not entirely reflected in the associated ROC AUC values. The average ROC AUC across all cancer and ML algorithm types was 0.74 ± 0.11 (mean ± standard deviation) (Fig 2C), where a value of 1 corresponds to a perfect predictor and 0.5 is no better than random. There were no clear differences between algorithms in terms of ROC AUC, though the regularized regression methods (Ridge and Lasso) exhibited slightly higher scores than most. A much larger difference was observed between cancer types, where the average ROC AUC for LGG, BRCA, and UCEC exceeded 0.85, but was at or below 0.60 for SKCM, OV, and LUSC (Fig 2D and 2E). Interestingly, BAX, HSPA4L, and KIF23 were often not among the top-scoring genes for the three cancer types with the highest ROC AUC values, which suggests that useful information can be extracted from the feature scores of trained ML classifiers despite a relatively poor corresponding ROC AUC.

2.4 Investigation of PSP genes associated with malignant transformation

After validating the ML gene scoring pipeline, we used the approach to evaluate the relative importance of each PSP component in distinguishing normal vs. tumor samples and identify genes that are likely to contribute to the tumor phenotype. Analogous to the p53 mutation analysis, samples for each cancer type were grouped according to cancer status (normal or tumor) and each of the 8 ML algorithms, as well as DE analysis, were used to score the 575 PSP genes. Cancer types without at least 10 samples in each group were excluded, yielding a total of 16 cancer types.

Unlike the classifiers trained on p53 mutation status, the ROC AUC for predicting normal vs. tumor samples based on PSP gene expression was high across all cancer types and ML methods, with an overall average of 0.98 ± 0.03 (S2 Fig). Only the LDA algorithm and the
ESCA cancer type tended to exhibit lower ROC AUC values relative to the others, but the lowest value for each was still greater than 0.80. This higher prediction performance for cancer status as compared to p53 mutation status was expected since there are more substantial expression changes between normal and tumor samples than there are between tumor cells differing in a single gene mutation.

2.4.1 Pan-cancer features

Inspection of the ML gene scores and DE analysis results revealed that kinesin-6 family proteins (\textit{KIF20A} and \textit{KIF23}), Crystallin Alpha B (\textit{CRYAB}), and several proteins belonging to the soluble N-ethylmaleimide-sensitive-factor attachment protein receptor (\textit{SNARE}) family (\textit{STX1A}, \textit{STX12}, \textit{STX11}, and \textit{VAMP2}) generally scored highly in both ML and DE approaches among the different cancer types (Figs 3A and 3B and S3), suggesting that these proteins may play an important role in tumor physiology. \textit{KIF20A} and \textit{KIF23} were among the top 3 genes
with the highest average ML consensus scores and exhibited a significant (FDR adjusted p-value < 0.01) expression increase in tumor compared to normal samples for all 16 cancer types except two renal carcinomas, KICH and KIRP (S4 Fig). Although they are associated with Golgi-to-ER retrograde transport and intracellular organelle transport, KIF20A and KIF23 play a critical role in mitosis and cytokinesis [30,31]. Inhibitors of these and other kinesin family proteins are undergoing clinical trials as anticancer therapeutics [32].

Although STX1A exhibited a similar expression increase across most cancer types as the kinesin-6 family proteins, the expression of STX11, STX12, and VAMP2 was significantly decreased in tumors across nearly all 16 cancer types. We further investigated the expression changes of the PSP genes belonging to the STX (Fig 3C) or VAMP (S5 Fig) gene families. There was a common restructuring pattern of STX expression among the different cancer types, involving a mixture of increases and decreases across the different STX genes, whereas VAMP genes tended to be more broadly decreased with the exception of VAMP1 and VAMP8. SNARE proteins, which include the STX and VAMP families, mediate the membrane fusion necessary for trafficking through the different steps of the secretory pathway [33]. SNAREs have been found to support many tumorigenic functions such as autophagy, cell invasion, and chemo-resistance, and thus constitute potential targets in anti-cancer therapies [34].

The CRYAB gene exhibited the second highest ML consensus score on average across the 16 cancer types (Fig 3A), and was significantly differentially expressed (FDR-adjusted p < 0.01) in all but 3 cancer types. Unlike the kinesins whose expression was nearly always increased in tumor relative to normal tissue, CRYAB expression was significantly decreased in tumor for 10 cancer types and increased in only 3: LIHC, KIRC, and KIRP (S6 Fig). The mean CRYAB mRNA abundance of LIHC samples were the lowest of all cancer types (< 10 TPM) and thus the DE results are less reliable; however, both KIRC and KIRP exhibited among the highest expression of CRYAB in paired normal samples which further increased by 1.9- and 6.5-fold in their corresponding tumor samples, respectively. The main role of CRYAB is to form multimeric structures with other proteins to prevent aggregation, but it has also been shown to exhibit other activities such as protection from oxidative stress and apoptotic stimuli [35]. In the context of cancer, there does not appear to be a clear consensus as to whether CRYAB supports or suppresses tumorigenesis [35]. Many studies conclude a pro-tumorigenic effect of CRYAB and a positive correlation between its expression and tumor aggression [36], whereas others report a tumor-suppressive activity and/or decreased expression in more aggressive tumors [37,38]. Our results suggest that cancer type is one factor determining whether CRYAB exerts an inhibitory or supportive role in a tumor, and that renal carcinomas in particular may be susceptible to CRYAB-modulating therapies.

### 2.4.2 Cancer-specific features

Although the kinesins, SNAREs, and CRYAB were among the highest ML gene scores when averaging over all 16 cancer types, no genes were consistently high scoring in more than a few of the cancer types. An inspection of the top-scoring genes of each individual cancer type revealed that high-scoring genes were primarily cancer-specific (S7 Fig). For example, RAS oncogene family member 17 (RAB17) scored highly in prostate adenocarcinoma (PRAD) across nearly all ML algorithms with a consensus score of 0.82, whereas its score in all other cancer types ranged from 0.01 to 0.17. Members of the RAB family regulate vesicle trafficking and are known to both promote and suppress tumor growth, depending on the family member and cancer type [39]. Although increased expression of RAB25 has been shown to contribute to prostate cancer malignancy and recurrence [40], similar studies or observations involving RAB17 are lacking.
When investigating the highest-scoring genes for each individual cancer type, we observed a high frequency of genes associated with glycosylation, particularly for five cancer types: STAD, READ, COAD, KICH, and THCA (Fig 4). For each of these cancer types, 3 out of their top 5 scoring genes encoded some form of glycosylation activity, despite such activity accounting for less than 18% of the 575 PSP genes considered in this study.

Genes associated with O-linked peptide glycosylation (LARGE2, B3GNTL1, B4GALNT2, and the GALNT family) were associated with KICH, COAD, and to a lesser extent READ, whereas genes encoding N-linked glycosylation activity (DPM2, MAN1A1, TUSC3, and the ALG family) scored highly for the STAD and THCA cancer types. The expression and specific patterns of glycans dictate cellular functions such as adhesion, signal transduction, differentiation, and proliferation, and the alteration of such patterns is a hallmark of tumor physiology [41,42]. It is therefore logical that genes encoding these post-translational modifications (PTMs) scored highly in the ML classifiers distinguishing normal from tumor samples. Furthermore, the cancer-specificity of these high-scoring genes is likely a reflection of the specificity and complexity of the glycosylation machinery and its large repertoire of glycan patterns [43].

We further explored the overrepresentation of glycosylation activity among high-scoring genes in these cancer types by investigating expression changes among secretome genes. The secretome gene expression fold-changes (between tumor and paired normal samples) were retrieved from a previous secretome analysis of TCGA data [7] and analyzed together with post-translational modification information obtained from UniProt [44]. For each of the secretome genes, the total number of N- and O-linked glycosylation sites were summed to obtain the total number of glycosylation sites per gene. We calculated the Spearman correlation between the total glycosylation sites and the expression fold change of each secretome gene for the five cancer types that exhibited an enrichment of glycosylation activity among its high-scoring PSP genes (STAD, READ, COAD, KICH, and THCA) (Fig 4B). In addition, we performed an
enrichment analysis (hypergeometric test) to evaluate if the significantly differentially expressed (FDR-adjusted p < 0.05) secretome genes were enriched in those with several (5 or more) N- or O-glycosylation sites (Fig 4C). Although only COAD and READ were significantly correlated (albeit weakly; Spearman’s ρ ~ 0.1), all of the cancer types except READ exhibited an enrichment of differentially expressed glycosylated secretome genes, suggesting that there may be some tuning of the PSP to accommodate the changing PTM needs of its secretome clientele. However, this is a very simplified approach to quantify a far more complex relationship between the PSP and secretome, which combined with the modest results should be interpreted with caution.

2.5 Analysis of different tumor stages

We next focused on PSP gene expression changes between tumor stages to identify secretory pathway components that were associated with disease severity and tumor development. Most TCGA samples are annotated with tumor stage information which generally ranges from stage I to stage IV, enabling the investigation of transcriptomic changes as a function of disease severity. Primary tumor samples were therefore grouped into stages I, II, III, and IV within each cancer type. Although a regression formulation could be applied by assigning numerical values to each of the tumor stages (such as 1–4 for stages I–IV, respectively), this establishes an implicit assumption about the relative severity of the different stages, for which we did not have sufficient supporting evidence. Furthermore, a regression formulation prevents the use of the same binary classification scheme and set of ML algorithms throughout the study. See S1 Text and S8 Fig for further detail and an implementation of the regression-based approach.

We instead chose to analyze the tumor stages within each cancer type in a pairwise manner by training ML classifiers on all possible pairs of tumor stages (for example, I vs. II, I vs. III, II vs. IV, etc.) to predict a sample’s stage based on its corresponding PSP gene expression profile. Cancer types without at least 10 samples in at least 2 tumor stages were discarded, yielding a total of 20 cancer types.

The ROC AUC values for the ML classifiers of tumor stage were substantially lower than those trained to separate normal vs. tumor samples, where many performed no better than random (ROC AUC ~ 0.5) (Figs 5A and S9). This was expected given that physiological differences between tumor stages are relatively subtle when compared to those between normal and cancerous tissue. Although our initial analysis with the p53 mutation ML classifiers suggested that feature importance scores can still provide some meaningful information despite relatively low ROC AUC values, we expect that such scores will largely degrade into random noise when approaching very poor values near (or below) 0.5. We therefore continued the analysis using only the top 10 performing cancer types based on their average ROC AUC values across the different ML algorithms and pairs of tumor stages (Fig 5A).

Among the PSP genes exhibiting the highest average consensus ML scores across the different cancer types and tumor stages, hyaluronan synthase 3 (HAS3) and KIF20A were some of the most prominent (Figs 5B and S10). Hyaluronan, an extracellular matrix polysaccharide, is enriched in the matrix surrounding virtually all epithelial tumors [45], and has been shown to promote tumor malignancy and metastasis by increasing cell invasiveness and anchorage-independent growth [46]. Consistent with the malignant function of hyaluronan, HAS3 exhibited a high consensus ML score for the TGCT, THCA, and STAD cancer types, suggesting an association between HAS3 expression and tumor stage. This was reflected in the gene expression profiles of TGCT tumor samples, which showed increasing expression of HAS3 and HAS2 with increasing tumor stage (S11 Fig). The expression of HAS3 also increased among later tumor stages of STAD, though after an initial decrease from stage I. Conversely, THCA
did not exhibit a substantial difference in HAS3 expression as a function of tumor stage, but a positive relationship was observed between THCA tumor stage and HAS1 expression (S11 Fig).

The top-scoring gene on average across the 10 cancer types was KIF20A, which exhibited particularly high scores for distinguishing tumor stages of KIRP, and to a lesser extent those of KIRC and ACC (S10 Fig). Interestingly, the renal carcinomas (KIRP, KICH, and KIRC) were among the few cancer types for which the kinesins (KIF23 and KIF20A) were either not significantly DE or exhibited a very low consensus ML score (less than 0.1 on average) when comparing normal to tumor tissues. This distinction becomes clear when comparing the expression of KIF20A between normal tissue and different tumor stages for each cancer type (Fig 5C). Most cancer types exhibit a sharp increase in KIF20A expression between normal and tumor samples that remain relatively constant across the different tumor stages, whereas renal carcinomas display a more gradual change in KIF20A expression that continues to increase with increasing tumor stage. Another noteworthy characteristic is the relatively low baseline expression of KIF20A in the normal kidney tissue samples compared to other cancer types, where even stage IV kidney carcinoma tumor samples exhibit expression levels that are comparable to the normal expression of other tissue types. This highlights the impact that tissue-specific differences can have on gene expression, which can mask potentially important changes associated with disease characteristics. Although KIF20A has been implicated in the development of many other cancer types [32,47,48], its involvement in renal carcinoma has not been addressed. The expression dynamics observed here suggest that KIF20A may support more invasive and metastatic functions associated with later stages of renal carcinoma, and thus constitutes a potential therapeutic target for this cancer type.
3. Discussion

The secretory pathway and its products are essential to the viability of eukaryotic organisms, but the deregulation of secretory machinery can support detrimental processes such as those driving tumorigenesis [7,49]. The identification of PSP components exhibiting oncogenic or tumor suppressive activities can aid in the development of novel anti-cancer therapies that aim to restore healthy PSP function through the modulation of these components. We therefore conducted a focused investigation of the PSP transcriptional changes associated with malignant transformation and tumor progression across many different cancer types. This allowed us to identify patterns in PSP expression that were common to carcinogenesis independent of cancer type, as well as explore secretory elements that exhibited cancer type specific behavior.

Accessing and interpreting the information embedded within omics data is non-trivial due to its high volume and dimensionality, and has traditionally been limited to a few methods, such as DE analysis and principal component analysis (PCA) [50]. We therefore sought to deepen the investigation by applying different machine learning (ML) approaches to provide a more detailed understanding of PSP behavior in tumors. However, ML methods generally struggle when the number of features (genes) greatly exceeds the number of samples, which is often the case for RNA-seq or other omics datasets and is referred to as the curse of dimensionality [51]. The ML methods were therefore well-suited for this focused study because the number of features was greatly reduced by including only the 575 PSP genes in the analyses. We note that the approach is not specific to these PSP genes and could be applied to any gene subset of interest, such as those encoding metabolism, the cell cycle, or the immune response. There is also often a risk of over-fitting or a high frequency of false positives when using a data-driven approach such as ML. We therefore implemented 8 different types of ML algorithms in our analyses and used a normalized average consensus score that combined the results of each algorithm. This ensured that genes identified as relevant to a given biological class variable were robust to the choice of ML algorithm or effects of overfitting.

The use of several different ML algorithms also enabled comparison of their predictive performance for each of the investigations, as quantified by ROC AUC. Although some ML algorithms (such as regularized regression and extreme gradient boosted trees) tended to outperform others (such as LDA and adaptive boosting) among the different class variables and cancer types, the difference was marginal and far from significant. This further supported using a consensus score that combined the output of the 8 different methods with equal weighting because no method consistently outperformed the others.

We used a well-studied feature in cancer biology—the mutation of p53—to evaluate the performance of the ML approach in terms of identifying biologically relevant features, and to compare with DE analysis. The highest consensus ML gene scores were exhibited by known regulatory targets of p53 in the 575 PSP genes (BAX, HSPA4L, and KIF23), providing confidence in the biological relevance of the ML results. Although the DE analysis identified these genes as important, the genes were not ranked as highly as other PSP genes. A reason for why the ML methods outperformed DE analysis in this case could be the capacity of the ML algorithms to capture interactions between genes and their expression patterns in different samples, whereas DE analysis estimates a fold-change and confidence for each gene individually.

A recurring gene of importance in our analyses was KIF20A, which was remarkably among the top-scoring genes for both ML and DE approaches and for all class variables (p53 mutation status, normal vs. tumor, and different tumor stages). This is consistent with the abundance of studies that have identified KIF20A to be highly expressed, linked to tumor aggressiveness, correlate with poor survival, diagnostic, and/or prognostic in many different cancer types [47], which support a critical and diverse role of the protein in general tumor development and
progression. There are however a lack of studies identifying any role or association of \textit{KIF20A} with renal carcinoma other than a co-expression network analysis of clear cell renal cell carcinoma (ccRCC) by Yuan and colleagues, in which \textit{KIF20A} was identified as one of six hub genes associated with ccRCC progression \cite{52}. We observed a substantial difference in the pattern of \textit{KIF20A} expression among normal and stage-stratified tumor samples in all renal carcinomas (KICH, KIRP, KIRC) compared to other cancer types; \textit{KIF20A} expression in renal carcinomas increased more gradually with increasing tumor stage rather than a sharp increase between normal and tumor that remained relatively constant across stages. We cannot speculate from this data alone as to the cause for the different dynamics, but it may indicate that anti-cancer treatments targeting \textit{KIF20A} could exhibit variable efficacy with tumor stage for renal carcinomas.

Our investigation demonstrates the efficacy of using a consensus ML-based gene scoring approach to predict biologically relevant features from a focused set of genes, and highlights the utility of using such an approach to complement and support the results of DE analysis. Furthermore, we present several PSP-associated proteins and protein families that exhibit a robust association with malignant transformation and tumor progression, and thus hold potential as targets in the development of anti-cancer therapeutics.

4. Methods

Analysis and figure scripts

The scripts used to perform the analyses and generate the figures presented here, as well as all analysis outputs, are available on GitHub: https://github.com/SysBioChalmers/CancerProteinSecretionML. Data files too large to host on GitHub were deposited on Zenodo: https://doi.org/10.5281/zenodo.3978373.

RNA-Seq and mutation data retrieval

Transcriptomic (RNA-seq) and mutation annotation data was retrieved from TCGA using the TCGAbiolinks R package \cite{53}. Raw gene counts and normalized (FPKM) gene counts were retrieved for 33 available cancer types. Mutation annotation information was obtained using the MuTect2 variant calling pipeline \cite{54}, and processed such that each gene in each sample was classified as mutated if it was modified in any way (insertion, deletion, missense, silent, etc.), otherwise it was classified as non-mutated. To avoid leakage of information between groups when training the ML classifiers, duplicate tumor samples originating from the same patient were removed such that there was only one tumor sample per patient.

Differential expression analysis

Differential expression analysis was performed on raw gene counts using the edgeR R package \cite{55}. Samples were grouped according to a binary class variable of interest (e.g., \textit{p53} mutation status, cancer status, or tumor stage), and all genes were included except for those that had fewer than 10 counts in 70% of the samples of the smallest group. Genes excluded from an analysis due to low counts were automatically assigned a log2 fold-change of zero and a \textit{p}-value of one. The design matrix included only information regarding group membership of each sample. We did not account for patient identity when performing the normal vs. primary tumor analysis because it would require the exclusion of many tumor samples which were used in the ML analyses. The expression fold-changes and associated significance (FDR-adjusted \textit{p}-values) were calculated prior to filtering out non-PSP genes. Analyses were
performed on each cancer type individually, where cancer types with fewer than 10 samples in a group were excluded.

A DE gene score was formulated to enable comparison with the ML gene score described below. The FDR-adjusted p-values of PSP genes for a given comparison and cancer type were log-transformed, negated, and normalized to a range of 0 to 1.

**ML model training and gene scoring**

All ML methods were implemented in python using the scikit-learn package [56] or the XGBoost package [21]. Gene expression values were converted to transcripts per million (TPM), and natural log transformed after adding a pseudocount of 1 TPM to avoid logarithm of zero. Samples were grouped according to a binary class variable of interest (e.g., p53 mutation status, cancer status, or tumor stage), and cancer types were analyzed individually, where cancer types with fewer than 10 samples in a group were excluded. Non-PSP genes and genes with a median expression below 0.1 TPM among both sample groups were also excluded.

For each cancer type and class variable, 8 classification models were trained using each of the 8 ML algorithms (random forests, ExTrees, AdaBoost, XGBoost, LDA, lasso regression, ridge regression, and SVM). Default parameters were used for each algorithm when available. For the tree-based methods, the number of estimators was set to the recommended value of the square root of the number of features, rounded down to the nearest integer. For the ridge and lasso regression methods, the “saga” solver was used with a maximum of 10,000 iterations. Training was performed using stratified 10-fold cross validation, such that each fold contained approximately the same proportion of samples from each group. Feature importance scores were extracted from each trained model, and normalized by taking the absolute value and scaling to a range of 0 to 1. A consensus score was calculated for each gene by averaging the normalized importance scores obtained from each of the 8 algorithms.

We note that our primary interest was to determine the relative importance of features (genes), as quantified by the gene scores, rather than developing predictive models. We therefore used all available samples when training each classifier and did not exclude any samples for a separate test set, meaning that the reported ROC AUC values are likely higher than what one would expect if the trained model predictions were evaluated using an independent test set of samples. The ROC AUC values were determined using stratified 10-fold cross validation, where the reported values are the mean of the 10 folds.

**ML model scoring metric**

Some of the cancer types and biological features investigated in this study involve imbalanced classes, where there are more samples belonging to one class than the other. Although metrics such as area under the precision-recall curve (PR AUC), average precision-recall, or the F1 statistic are often well-suited for heavily imbalanced datasets, these methods are sensitive to the choice of “positive” class and will vary upon class swapping. If the positive class is the majority class, then even a very poor model can exhibit a high PR AUC, for example.

In our study, it is not always trivial or meaningful to define the positive class, such as when comparing tumor stages. Even in cases where it may appear more straightforward to select the positive class (e.g., normal vs. tumor), some cancer types contain majority normal samples, whereas others contain majority tumor samples. If we for example define tumor samples as the positive class, the PR AUC would be inflated for cancer types with more tumor samples, and vice versa. We therefore chose to use the ROC AUC as the scoring metric because it is sufficiently resistant to class imbalances and is unaffected by class swapping, both of which are relevant features of the present dataset.
Classification of tumor stages

Unlike the p53 mutation or cancer status classification analyses, there were more than two possible classes when investigating tumor stages (I, II, III, and IV). The tumor stages were therefore analyzed in pairs to facilitate the use of the same binary classification pipeline used for the other analyses. Specifically, within each cancer type, all possible pairwise combinations of tumor stages were compared. Therefore, if a cancer type had a sufficient number of samples (10 or more) in all 4 tumor stages, a total of 6 classification analyses were performed: stage I vs. II, I vs. III, I vs. IV, II vs. III, II vs. IV, and III vs. IV. The formulation and application of an alternative regression-based approach for the tumor stages is presented in S1 Text.

Identification of p53 regulatory targets

To identify the set of PSP genes known to be directly regulated by p53, we conducted an extensive literature search in combination with an investigation of the STRING database [57]. We first searched for primary literature and review papers that identified or summarized the known p53 gene regulatory targets [27,28], and filtered these target genes to include only those overlapping with the PSP gene list. Furthermore, we extracted all indexed experimental interactions between the 575 PSP genes and p53 from the STRING database, using 0.4 as a minimum required interaction score to yield a list of 12 proteins with evidence of p53 interaction (S5 Table). We then performed an additional literature search on each of the 12 proteins to determine which were confirmed to be direct regulatory targets of p53. A total of 4 PSP genes were identified from the literature search as confirmed targets: BAX, HSPA4L, KIF23, and BAK1 [27–29]. We note that BCL2 and PARK2 were among the 12 genes identified from the STRING database for which some evidence of p53 regulation could be found, but there were contradictory studies showing that BCL2 is not regulated by p53 [58], and a lack of support for p53-mediated regulation of PARK2 in humans [27].

Secretome glycosylation correlation and enrichment analysis

Secretome gene expression changes between normal and tumor samples were analyzed together with their post-translational modification annotations. The list of secretome genes and their differential expression results (tumor vs. paired normal, fold-changes and FDR-adjusted p-values) were retrieved from S2 Table of the Robinson et al. cancer secretome study [7]. Post-translational modification information was retrieved from UniProt [44] for each of the secretome genes, specifically the number of N-glycosylation and O-glycosylation sites per gene. For 5 different cancer types (STAD, COAD, READ, KICH, and THCA), we calculated the Spearman correlation between the total number of N- and O-glycosylation sites and the normal vs. tumor log2 expression fold-change among all secretome genes. A hypergeometric test was also performed to assess whether differentially expressed secretome genes (FDR-adjusted p < 0.05) in these cancer types were significantly enriched with highly glycosylated members, defined here as those with 5 or more (95th percentile) total N- or O-glycosylation sites.

Tumor stage processing

Tumor stages in TCGA are often provided with sub-stage detail, such as stage IIa, stage IIb, etc. We merged such annotations to achieve only four different stages: I, II, III, and IV. The merging was performed to avoid increasingly large numbers of pairwise stage comparisons, as well as groups with very few samples.
Supporting information

S1 Text. Regression analysis of tumor stages.

S1 Fig. Consensus ML and DE gene scores for p53 mutation status. Histogram of (A) mean ML gene scores and (B) mean DE gene scores across all available cancer types, where the PSP genes known to be directly regulated by p53 are labeled. Boxplots of (C) consensus ML gene scores and (D) DE gene scores for the top 10 scoring genes on average. Clustered heatmaps showing the (E) consensus ML gene scores and (F) DE gene scores for individual cancers for the top 10 scoring genes on average.

S2 Fig. ROC AUC values for the prediction of cancer status by the trained ML models. (A) Density histogram of all ROC AUC values across different cancer types and ML algorithms. Boxplots showing the ROC AUC values grouped by (B) cancer type or (C) ML algorithm.

S3 Fig. Consensus ML and DE gene scores for cancer status. Histogram of (A) mean ML gene scores and (B) mean DE gene scores across all available cancer types. Boxplots of (C) consensus ML gene scores and (D) DE gene scores for the top 10 scoring genes on average. Clustered heatmaps showing the (E) consensus ML gene scores and (F) DE gene scores for individual cancers for the top 10 scoring genes on average.

S4 Fig. Expression fold-change and significance of PSP genes belonging to the KIF family from the DE analysis of normal vs. tumor. Color indicates fold-change magnitude and direction, whereas circle size indicates significance (FDR-adjusted p-value).

S5 Fig. Expression fold-change and significance of PSP genes belonging to the VAMP family from the DE analysis of normal vs. tumor. Color indicates fold-change magnitude and direction, whereas circle size indicates significance (FDR-adjusted p-value).

S6 Fig. Expression of CRYAB in normal and tumor tissue samples across different cancer types. Cancer types are grouped according to whether CRYAB significantly (FDR-adjusted p-value < 0.01) changed in expression between normal and tumor, and whether that change was a decrease or increase.

S7 Fig. Heatmap of the consensus ML gene scores for cancer status. The heatmap includes all available cancer types and the top 5 scoring genes of each type. For visual aid, rows and columns were clustered such that high-scoring genes for each cancer tend to lie along or near the diagonal. The colorbar to the left of the heatmap indicates the function associated with each gene.

S8 Fig. Regression-based analysis of tumor stage. (A) Histogram of mean ML gene scores across all cancer types. (B) Consensus ML gene scores for the top 10 scoring genes on average. (C) Density histogram of negative mean squared error (Neg. MSE) values across different cancer types and ML algorithms. Boxplots showing the negative MSE values grouped by (D) cancer type or (E) ML algorithm. (F) Heatmap of consensus ML gene scores, showing the top 5
scoring genes of each cancer type. The colorbar to the left of the heatmap indicates the function associated with each gene.

S9 Fig. ROC AUC values for the prediction of tumor stage by the trained ML models. (A) Density histogram of all ROC AUC values across different cancer types and ML algorithms. Boxplots showing the ROC AUC values grouped by (B) cancer type, (C) ML algorithm, or (D) all possible tumor stage pairs. (E) Heatmap showing the consensus ML gene scores for the top 10 scoring genes on average, including all possible tumor stage pairs.

S10 Fig. Consensus ML and DE gene scores for tumor stage. Histogram of (A) mean ML gene scores and (B) mean DE gene scores across the 10 cancer types with the highest average ROC AUC values. Boxplots of (C) consensus ML gene scores and (D) DE gene scores for the top 10 scoring genes on average. Clustered heatmaps showing the (E) consensus ML gene scores and (F) DE gene scores for individual cancers for the top 10 scoring genes on average.

S11 Fig. Expression of PSP genes belonging to the HAS family among different tumor stages. Boxplots present the expression of HAS1, HAS1, and HAS3 in different tumor stages of STAD, TGCT, and THCA cancer types. Note that the stage IV expression levels are lacking for TGCT because no stage IV samples were available for this cancer type.

S1 Table. TCGA cancer abbreviations and sample metadata.

S2 Table. Consensus ML gene scores for p53 mutation, cancer status, and tumor stage.

S3 Table. Model score values of each ML algorithm for predicting p53 mutation, cancer status, and tumor stage.

S4 Table. Differential expression log2 fold-changes, FDR-adjusted p-values, and gene scores for p53 mutation, cancer status, and tumor stage.

S5 Table. Evaluation of potential p53 regulatory targets identified by the STRING database.

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