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| Fatal: plus patch | vi |
|--|----|
| Fatal: TODO: Consider comparing A1 and A2 vs meta-PCNA and | |
| meta-ECM in TCGA – are $A1/A2$ better than the metas? Model | |
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| Fatal: TODO: Cohort recruitment and ethics | 33 |
| Fatal: TODO: Sample collection, preparation, and gene expression mi- | |
| croarrays | 34 |
| Fatal: give instantiation values for the algo somewhere | 47 |
| Fatal: Add the derivation in somewhere – perhaps an appendix. It's a | |
| pain in the arse so probs want to avoid the main text | 52 |
| Fatal: Add specific value of mindepth used | 53 |

Mah Dissertat'n

Mark Pinese

 $\label{eq:January 11, 2015} \text{ Build } 0.0.293$

ORIGINALITY STATEMENT

'I hereby declare that this submission is my own work and to the best of my knowledge it contains no materials previously published or written by another person, or substantial proportions of material which have been accepted for the award of any other degree or diploma at UNSW or any other educational institution, except where due acknowledgement is made in the thesis. Any contribution made to the research by others, with whom I have worked at UNSW or elsewhere, is explicitly acknowledged in the thesis. I also declare that the intellectual content of this thesis is the product of my own work, except to the extent that assistance from others in the project's design and conception or in style, presentation and linguistic expression is acknowledged.'

| Signed | |
|--------|--|
| Date | |

Acknowledgements

Da abstract.

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Software versions

Unless otherwise specified, the following versions of software were used in all work.

| bamtools | 2.2.2 |
|------------------------|----------------------------|
| bedtools | 2.18.2 |
| cd-hit | 4.6.1 MP Fatal: plus patch |
| FastQC | 0.10.1 |
| GATK | 3.1-1 |
| julia | 0.3.2 |
| MSigDB | 4.0 |
| muTect | 1.1.6-4-g69b7a37 |
| ncbi-blast | 2.2.29 |
| picard-tools | 1.109 |
| PROVEAN | 1.1.5 |
| Python | 2.7.8 / 3.4.1 |
| R | 3.1.1 |
| ahaz | 1.14 |
| depmixS4 | 1.3-2 |
| doParallelMC | 1.0.8 |
| Exact | 1.4 |
| GSVA | 1.14.1 |
| illumina Humanv 4.db | 1.24.0 |
| lumi | 2.18.0 |
| lumidat | 1.2.3 |
| nleqslv | 2.5 |
| NMF | 0.20.5 |
| nnls | 1.4 |
| ${ m org. Hs. eg. db}$ | 3.0.0 |
| | |

| ${\rm random} Forest$ | 4.6-10 |
|-----------------------|--------|
| Rsolnp | 1.14 |
| survival | 2.37-7 |
| samtools | 1.0 |
| SHRiMP | 2.2.3 |
| strelka | 1.0.14 |
| tabix | 1.0 |
| vcftools | 0.1.10 |
| VEP | 76 |

Conventions

Unless otherwise specified, the following conventions are used throughout this dissertation.

- \bullet Indices in algorithm pseudocode are 1-based.
- Logarithms (log) and exponentiations (exp) are to base e.
- Square brackets denote the Iverson bracket: [X] := 1 if X is true, else 0.
- x_+ indicates the value of the ramp function at real $x, x_+ := \max(0, x)$.

Chapter 1

Introduction

Chapter 2

A Preoperative Molecular Prognostic for Pancreas Cancer

Thesis: A preoperative prognostic tool for pancreas cancer can be developed to discriminate good between and poor prognosis patients more reliably than current methods.

Summary For those patients fortunate enough to be diagnosed with a resectable tumour, surgical removal of the primary cancer is the best first-line therapy for pancreas cancer. However, the significant morbidity associated with pancreas cancer resection makes it cruicially important to only operate on the patients who stand to benefit from the procedure. Identifying just those patients who will respond to resection remains a serious challenge in pancreas cancer treatment: current criteria to select patients for resection perform poorly, and consequently many patients will undergo a complex procedure, with serious effects on future quality of life, for little benefit. Tumour biomarkers have the potential to dramatically refine morphology-based staging criteria by supplying a direct readout of tumour biology, and recent technological developments have enabled the preoperative measurement of tissue biomarkers in pancreas cancer. The ability to measure pancreas cancer tissue biomarker levels preoperatively, combined with the enhanced information on disease state available from tissue biomarkers, finally enables the development of preoperative staging criteria that accurately identify pancreas cancer patients for resection. This chapter details the development and validation of a two-biomarker preoperative prognostic tool for resectable pancreas cancer, that can be used to define new resectability criteria to accurately select patients who stand to benefit from primary tumour resection.

2.1 Introduction

For patients with a resectable tumour and no known metastases, surgical removal of the primary tumour is the current recommended first-line therapy for pancreas cancer, and the only intervention offering the realistic possibility of a cure [13]. However, pancreas cancer resection is a major procedure, with the potential for serious complications, morbidity, and reduced quality of life following recovery [23]. Due to the substantial negative effects of surgery, the decision of whether or not to perform curative-intent resection should balance the risks of surgery against its expected benefits, tailored to each individual case.

Unfortunately, current practice guidelines recommend that curative-intent surgery be offered to all metastasis-free patients with a resectable tumour, with no consideration of personal benefit [13]. This blanket approach to selecting patients for curative resection has proven to be highly inadequate. Even following pathologically complete tumour removal and adjuvant chemotherapy, more than 70% of current pancreas ductal carcinoma patients will relapse with, and ultimately succumb to, distant metastases [4]. These occult metastases must have been present prior to removal of the primary tumour, yet were undetectable during initial investigations, and their presence means that any curative-intent resection was futile. As a result, the majority of 'curative' resections that are undertaken based on current selection criteria are performed on patients with occult metastases, have no hope of actually effecting a cure, and would not have been undertaken at all if the presence of the metastases had been known prior to surgery. Better methods for selecting patients for resection are urgently needed.

A number of pancreas cancer grading and schemes and prognostic tools have been described, but inconsistent performance, or a reliance on information that can only be known post-operatively, limits their use in pre-operative decisions. The level of serum carbohydrate antigen 19-9 (CA-19-9) is a well-characterised biomarker of pancreas cancer, with high levels correlating with

increased tumour burden, lower probability of resectability, increased post-resection recurrence, and worse prognosis [27, 3, 4, 32]. CA-19-9 levels are easily determined pre-operatively, but the use of this marker is complicated by a lack of consensus on threshold concentrations, the elevation of CA-19-9 levels by a number of conditions other than pancreas cancer, and the complete absence of this marker in approximately 10% of the general population [3]. Additionally, although CA-19-9 levels are statistically associated with post-resection recurrence by distant metastasis, a very low positive predictive value (PPV) renders the biomarker unhelpful when deciding whether or not to resect [27].

The current standard prognostic tool for pancreas cancer is the Memorial Sloan-Kettering Cancer Center (MSKCC) nomogram [9], which integrates a number of clinico-pathological variables (CPVs) to arrive at point estimates for survival post-resection. Unfortunately, its clinical utility is small: as it relies on information that is only available following resection, the MSKCC nomogram is only useful in a post-operative context, and cannot assist in pre-operative decisions to resect. This severely limiting reliance on postoperative variables was made necessary by the fact that most classical prognostic factors in pancreas cancer (such as lymph node infiltration, resection margin status, or histological grade [7]) can only be reliably measured following resection. Any prognostic tool for pancreas cancer that relies only on classical CPVs will likely share this same reliance on post-operative variables; an effective pre-operative prognostic conversely will need to be based on measurements other than the classical CPVs.

Tissue biomarkers can provide an almost direct window on the cellular state of tumour cells, and thus have the potential to predict cell behaviour far more reliably than macroscopic CPVs. Given that most pancreas cancer patients who undergo curative resection quickly recur due to occult metastases, biomarkers of metastasis have the potential to identify those patients who are likely to already have occult metastatic disease at the time of surgery, and thus better inform the decision to resect. Two such biomarkers of metastasis are the cancer cell levels of the epithelial to mesenchymal transition (EMT)-related S100A2 and S100A4 proteins, both of which are strongly predictive of outcome following resection, and appear to reflect the presence of a pro-metastatic invasive phenotype in the cancer [5, 47, 28]. Unfortunately, these tissue biomarkers have to date only been assessed in bulk tissue sam-

ples collected during surgery, and their utility, or even measurability, in a pre-operative setting, is untested.

Recent techological developments have made possible the pre-operative measurement of tissue biomarkers as a part of endoscopic ultrasound (EUS), a routine diagnostic modality for pancreas cancer. Immunohistochemical (IHC) staining has been successfully performed on fine needle aspirate (FNA) biopsies of pancreas neoplasms collected during EUS [36, 40, 43], and in principle EUS-FNA-IHC could form the basis of a pre-operative biomarker measurement methodology for routine use in pancreas cancer diagnosis. Although this proposed biomarker measurement approach is not currently in common use, it utilises only techniques that are commonly available in pancreas cancer treatment centres, and thus has the potential to be rapidly integrated into current diagnostic workflows, should biomarker measurements prove to be clinically useful.

The nexus of known biomarkers of metastatic behaviour, and new preoperatively applicable techniques to measure these biomarkers, presents an opportunity to address the pressing need for better criteria to select patients for pancreas cancer resection. As part of the APGI, as well as other work, the group has collected tissue measurements of S100A2 and S100A4 biomarkers, and detailed patient follow-up, for a large number of cases of pancreas cancer from a range of independent cohorts. These cases will be used to develop a prognostic predictive tool for outcome following resection, that uses tissue levels of S100A2 and S100A4 as major prognostic factors. This tool will rely on biomarker measurements made on tissue collected during resection, and thus will not be directly applicable pre-operatively. However, pilot study data will be used to show that levels of S100A2 and S100A4 measured by pre-operative EUS-FNA-IHC correlate well to tissue levels of the biomarkers measured on peri-operative specimens, and therefore that biomarker-based pre-operative prediction of patient outcome following surgery is possible in a clinical setting.

The majority of pancreas cancer resection procedures today are performed on patients who should never have been offered surgical resection at all. These patients have undetected metastases at the time of surgery, and will derive little benefit from a major operation that will have serious impacts on quality of life. Current tools for patient staging and estimation of prognosis are either ineffective at identifying patients at risk for occult metastases, or only applicable post-operatively, and thus cannot be used to inform the decision of whether or not to resect. It may be possible to identify pre-operatively those patients with occult metastases, by examining levels of known tissue biomarkers of pancreas cancer metastatic potential. This chapter will describe the development of a prognostic tool for pancreas cancer that makes use of these biomarkers to predict personalised outcome following surgery, and assist in making treatment decisions appropriate for each individual pancreas cancer patient.

2.2 Results

Data from a large retrospectively-acquired cohort, the New South Wales Pancreatic Cancer Network (NSWPCN) cohort, were used to derive a prognostic predictor for survival of pancreas cancer patients who underwent curative-intent resection. Discrimination and calibration of this predictor were then verified on TODO number independent surgical cohorts. Data from an EUS-FNA-IHC pilot study were then used to establish that pre-operatively assessed tissue biomarker levels reflected measurements from operative biopsies, and thus that the prognostic predictor developed here could be applied in a pre-operative decision setting.

Prognostic variables and biomarkers

As the aim was to develop a prognostic predictor that could be applied preoperatively, only the small subset of traditional CPVs that were deemed to be
measurable prior to resection were considered. These variables were: patient
sex and age at diagnosis, tumour location (dichotomised as head of pancreas
vs other location), and size of the tumour's longest pathological axis. All
but one of these variables is straightforward to measure pre-operatively, with
the exception being tumour size. In the training and validation sets, tumour
size was measured post-operatively from the resected tumour, whereas in preoperative application tumour size must be estimated by imaging techniques
such as computed tomography (CT) or EUS. The correlation between CT and
EUS estimates of tumour size, and actual size upon resection, is strong but not
perfect [2]. As no pre-operative measurements of tumour size were available
in the cohorts used for this work, pre-operative size was approximated by the
post-operative measurement. The implications of this approximation for the

prognostic tool developed here, as well as for future work, are considered in the discussion.

The dichotomised tissue levels of two pre-operatively assessable biomarkers, S100A2 and S100A4, were considered when building the prognostic tool. These biomarker levels were measured by immunohistochemistry (IHC) on tissue collected during resection, and thus

Scores for two pre-operatively assessable biomarkers, S100A2 and S100A4, were considered during prognostic development. Post-operative

TODO: Biomarkers. Specifically, why I didn't use CA-19-9.

Cohorts and Characteristics

General characteristics of the NSWPCN, TODO, and TODO cohorts are summarised in Table ??.

Prognostic model building and selection

Candidate prognostic models were constructed on the NSWPCN training data by iterative model fitting, evaluation, and refinement. To guard against over-fitting caused by this iterative process, the NSWPCN cohort was randomly split once into model building and testing sets. All model fitting and refinement described below was performed on the model building set, to yield three final candidate prognostic predictors. The performance of each of these three predictors was then assessed on the model test set, and the most parsimonious high-performing model was chosen as the final prognostic predictor, for subsequent external validation.

Model functional form and expanded terms The Cox proportional hazard (CPH) framework was used to assess functional form for the two continuous covariates: age at diagnosis, and maximum pathological axis size. local regression (LOESS) smooths of martingale residuals indicated a possible weak U-shaped relationship for age at diagnosis (TODO figref), and a knee-shaped form for size (TODO figref), with the knee at approximately 0 in centered units. In subsequent fits these forms were modelled by adding a quadratic term for centered age at diagnosis, and a size₊ ramp term for centered axis size. The original set of five linear prognostic terms, plus the two additional nonlinear terms, was denoted the expanded term set.

Proportional hazards assumption A Grambsch-Therneau test [17] on the CPH model fit using all expanded terms indicated that patient sex violated the proportional hazards (PHs) assumption (P = TODO, TODO fig resid plot) – in other words, the two sexes had significantly different baseline hazard forms. To account for this effect, all subsequent models were stratified by patient sex, so that the survival of male and female patients was modelled by two different baseline hazard functions. A repeated Grambsch-Thernau test on the stratified model indicated no further significant violations of PH.

Variable selection Genetic selection was used to identify the model with optimal Bayesian information criterion (BIC) from the set of all CPH models that use any combination of the expanded terms. Models with interactions between terms of up to degree two were considered, and a marginal term constraint was enforced, to ensure that interaction effects were only present in the model specification if the associated main effects were also. Stratification of baseline hazard by patient sex was used in all models. The identified optimal CPH model used three variables: tumour size (linear term only), S100A2 status, and S100A4 status, in addition to the sex stratum. This model was also identified by stepwise backward selection for optimal BIC, starting from the marginal CPH fit using all expanded terms. The final BIC-selected set of prognostic terms (tumour size linear term, S100A2 binary status, S100A4 binary status, and a patient sex stratum) was denoted the reduced term set.

Model CP1 A final prognostic CPH regression model was fit to the NSW-PCN model building data using only the reduced term set; this model was termed CP1. CP1 did not violate the PH assumption by the Grambsch-Therneau test (global P = TODO). Model residuals were inspected to identify possible outlier patients, and assess overall fit stability. Deviance residuals indicated no egregious outlier patients, and DFBETAS indicated TODO influential patients with $|DFBETAS_{i,j}| > \frac{2}{\sqrt{n}}$, where n = TODO. TODO: Stuff on these outliers.

Model GG1 Semiparametric Cox PH models such as CP1 provide a convenient framework for covariate testing and model diagnostics, but their unspecified baseline hazard term significantly complicates their use as prognostic predictors: patients can only be ranked by relative hazard, and absolute es-

timates of survival probabilities are unavailable. Although it is possible to approximate the baseline hazard in the Cox model, a more robust alternative is to use fully parametric models, in which the baseline hazard distribution is explicitly specified. The advantages of parametric models in terms of robustness and interpretability are offset by their more stringent assumptions: if the chosen baseline distribution is unsuited to the particular data to be fit, predictions from parametric models can be very poor. Given the potential benefits of parametric models for survival prediction, a parametric alternative to model CP1, termed model GG1, was investigated.

Model GG1, employing a generalised gamma (GG) survival distribution [12], was fit to the NSWPCN model building data by maximum likelihood. Guided by the model functional form and baseline hazard stratification indicated by the Cox model diagnostics, the GG distribution location parameter β was linearly dependent on all terms in the reduced set, but the shape parameters σ and λ were modelled as dependent on patient sex only. The goodness of fit of GG1 was investigated by examination of residuals, and graphical assessment of prediction accuracy. Deviance and DFBETAS residuals indicated no extreme outliers or unduly influential samples, and GG1 survival predictions matched empirical Kaplan-Meier (KM) estimates to within error across a range of covariate values and times Figure ??TODO fig). Do some QQ plots (eg http://www.sciencedirect.com/science/article/pii/S0531556508003264, or maybe http://onlinelibrary.wiley.com/doi/10.1002/sim.1592/pdf?)

Model RSF Regression models like CP1 and GG1 are familiar and readily interpretable, but are heavily dependent on the analyst identifying appropriate variables and functional forms. Ensemble tree models such as random forests [8] naturally and automatically model nonlinearity and arbitrary level interactions, and are tolerant of large numbers of irrelevant or collinear variables, albeit at the cost of very poor interpretability, and large data and computational requirements. Random forests have been adapted to model censored data [25], and can provide an alternative prognostic predictor that is very distinct from CP1 and GG1, and may be able to leverage data structure not used by these more classical models.

To investigate whether tree ensemble models could provide improved performance over classical approaches, a random survival forest model, termed RSF, was fit to the NSWPCN model building data. In contrast to CP1 and GG1, which used a reduced set of terms as predictors, RSF was supplied all preoperatively-assessable variables as covariates.

Model selection Predictive performance of the three prognostic models was compared on the holdout NSWPCN model test set, to select a single high-performing parsimonious model for external validation. For each model, prediction accuracy over time was assessed by the Brier score for censored data [16], and overall accuracy was measured by the integrated Brier score (IBS). All predictors were also compared against a no-information control predictor, formed as the KM estimate in the NSWPCN model building set of the marginal survival function. This no-information estimate, here denoted KM0, is the optimal survival predictor in the absence of any prognostic information or patient stratification; the prognostic models developed here must at least outperform KM0 to be of any utility at all. Brier scores and IBSs were calculated from multiple bootstrap draws of the holdout model test set, to supply rough assessments of score variability.

Traces of the Brier score over time indicated that all models produced survival estimates superior to KM0 from approximately 100 days to 600 days post-diagnosis (TODO figref). The performance of the models was similar, with RSF displaying slightly higher error rates at longer follow-up times, and models CP1 and GG1 demonstrating comparable performance. Bootstrapped differences between KM0 and model IBS further highlighted that although all models were superior to KM0, RSF had noticeably lower average performance than CP1 and GG1 (TODO fig), which were in turn nearly equivalent in terms of performance.

Of the three models assessed, CP1 and GG1 exhibited very similar best predictive performance on the holdout data. As there was no substantial difference in performance between CP1 and GG1, the simpler of the two models, GG1, was selected for external validation.

External validation

Web tool

TODO: In app, show +/- margin curves, to guide surgeons as to benefit from aggressive surgery.

Cohort characteristics

Development of a preoperative prognostic model

Validation of the prognostic model

Intro here on disc & calib. Also describe cohorts briefly.

Discrimination

Calibration

Web tool TODO – better name

2.3 Discussion

2.4 Methods

Cohort recruitment and ethics

Chapter 3

Signatures of Survival Processes in Pancreas Cancer

Thesis: Specific molecular processes control survival of patients with resectable pancreatic ductal adenocarcinoma, and these processes can be identified using gene expression data.

Summary Very little is known regarding the biological processes that control the survival of patients with pancreatic ductal adenocarcinoma (PDAC), the most common and aggressive form of pancreas cancer. As discussed in Chapter 2, the wide range of relative patient survival times that is observed in practice is not well explained by extrinsic factors such as age at diagnosis, and perhaps instead reflects differences in the biological processes operating within each tumour. Recent molecular profiling work [11] has identified possible molecular subtypes within the previously homogenous group of PDAC, but these subtypes have not achieved the maturity or clinical application of those in breast cancer, and their discovery and validation has been hampered by adhoc methodology, and the lack of large, well-curated cohorts of PDAC samples. The recently-compiled APGI cohort contains the largest group of clinically annotated PDAC samples, with accompanying gene expression (GEX) and high-quality follow-up data, in the world. It presents a unique opportunity to apply modern techniques for prognostic signature identification to the discovery of biological processes that drive the clinical course of pancreas cancer. These signatures may find application as prognostic tools in their own right, but more importantly can supply much-needed information on the fundamental biology of the one common cancer that has, to date, been almost entirely refractory to all the tools of modern molecular medicine.

3.1 Introduction

Despite extensive research, PDAC remains a poorly-understood disease. Recent genomic profiling has revealed the genetic alterations that accompany the cancer [6], and a huge number of prognostic factors are known [22] (refer to Chapter 1 for further discussion on both points), but these findings have shed little light on the fundamental disease processes at work in individual tumours. This is a consequence of genetic and biomarker data being poorly-suited for understanding the biological state of a cell: although genetic alterations are central to the etiology of cancer, they give incomplete information on the pathways and systems actually active in a given tumour, and biomarkers supply non-causal readouts of cell state that are difficult to trace back to underlying biological processes.

Sitting between the regulatory function of transcription control, and the effector function of protein expression, GEX data integrate information from all aspects of cell condition, including genetic alterations, signalling pathway activity, and metabolic status. As such, it is unsurprising that GEX data are superior indicators of cell state, better than all other high-throughput measurement methods, such as protein expression or genetic alterations [38]. However, the involvement of GEX with so many biological inputs is also a weakness: typical differential expression studies will identify many hundreds of transcripts that vary between disease states, and the deconvolution of this complex set of hundreds of effects back to a small number of causative molecular processes remains challenging.

Historically, disease GEX profiling studies have largely refrained from attempting to infer the state of a few molecular processes from the many hundreds of differentially-expressed genes identified; notable early exceptions are for example [1, 29]. A number of factors are likely to have contributed to this reluctance: deconvolution methods require relatively large sets of high-quality measurements [33], early techniques were poorly-suited to the particular requirements of the GEX deconvolution problem, and the signature databases that assist the assignation of a biological annotation to the output from a deconvolution calculation (for example, the MSigDB [45]) are only now reaching

maturity, with some areas of biology still underrepresented.

A simple synthetic example illustrates the problem and process of GEX deconvolution, and the character of solutions produced by both classical and modern techniques. Consider a group of samples, each of which is in one of three distinct biological states: state A, state B, and an intermediate state. Which state a sample is in affects the expression of two genes, gene 1, and gene 2: state A is associated with higher gene 2 expression than gene 1 expression; state B with higher gene 1 expression than gene 2; and the intermediate state with low expression for both genes (Figure 3.1). From the figure it is apparent that samples lie along two lines in transcription space; these lines I term metagenes.

Accurately knowing the metagenes at work within a biological system considerably simplifies reasoning about transcription within the system. In the example of Figure 3.1, state A is associated with high metagene 1, state B with high metagene 2, and the transition state with low scores of both. Additionally, the loadings of genes on the metagenes themselves (the directions of the metagene arrows) provides information on transcriptional control within the system: metagenes define the axes along which cell state must move, and so provide a simpler and more accurate representation of cell state than the full set of gene expression measurements. Metagenes can also be considered to capture co-expressed modules of genes, with likely biological significance. The advantages of a metagene-centric perspective to interpreting GEX become increasingly apparent as more genes are considered, and when thousands of genes are measured per sample, deconvolving the highly complex patterns of expression of thousands of genes, to only tens of metagenes, represents a powerful reduction in complexity. However, in practical use deconvolution methods must operate in thousand dimensional spaces, rather than the two dimensions in this example, and the computational and methodological complexities involved, as well as the poor results yielded by traditional approaches, have limited the application of GEX deconvolution.

A number of techniques from the field of matrix factorization have been applied to the GEX deconvolution problem, first principal component analysis (PCA) [1], then independent component analysis (ICA) [30], and more recently the various forms of non-negative matrix factorization (NMF) (first used for GEX in [10]). A number of reports have highlighted the unsuitability of PCA for GEX deconvolution, and the relative superiority of ICA

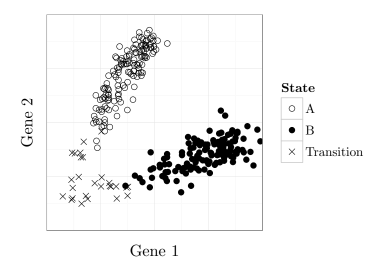


Figure 3.1: The gene deconvolution problem. Shown are the expression levels of two genes across three biological states, where each point represents the gene expression of a single sample in one of the three biological states. State A (hollow circles) is characterised by gene 2 > gene 1; state B (solid circles) by gene 1 > gene 2; and the intermediate state (crosses) by low levels of both genes. The challenge of gene deconvolution is to automatically infer, from unlabelled data (ie state is unknown), the dominant lines of gene expression (metagenes) along which most samples lie.

[29, 39, 46]; this is primarily due to the PCA requirement that metagenes be orthogonal [31], a situation that is not supported by our knowledge of biology, and results in bizarre artefacts such as PCA metagenes not actually being aligned with the expression pattern of any sample (Figure 3.2(a)). Although the results from ICA are more interpretable than those from PCA, they still do not consider that GEX is a non-negative process: it is impossible to have a concentration of mRNA that is less than zero, and therefore for best interpretability we wish metagenes to have non-negative 'expression' as well. ICA does not produce solutions satisfying this requirement, and more importantly its non-Gaussianity objective is not necessarily optimal for GEX deconvolution (Figure 3.2(b)), reducing its ultimate utility. NMF techniques have the potential to produce excellent GEX decompositions (Figure 3.2(c)), but are relatively new methods that have very high computational requirements, and often require careful tuning, making their effective application challenging.

In addition to the general technical challenges of GEX deconvolution, issues particular to pancreas cancer significantly complicate attempts to identify

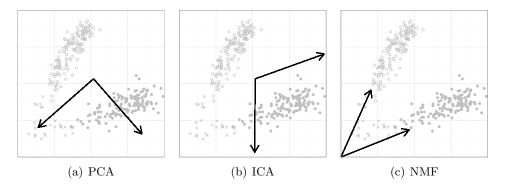


Figure 3.2: NMF produces a more accurate GEX decomposition than either PCA or ICA. Metagenes found by each method are shown as arrows. PCA (panel a) produces metagenes that don't match the expression pattern seen in any sample; these metagenes do not have a ready biological interpretation. ICA (panel b) accurately identifies one metagene, but the inappropriateness of the non-Gaussianity criterion for these data leads to an incorrect estimate of the other; although this solution is better than that of PCA, not all metagenes align well with biology. NMF (panel c) provides the best deconvolution; the metagenes identified closely match the expression patterns observed, and reflect the true structure of co-expression within the samples.

molecular processes at work within the tumours. Pancreas cancer is challenging to sample, and mRNA in the tissue degrades rapidly once extracted, complicating sample collection. Additionally, a feature of PDAC is the presence of a dense desmoplastic stromal reaction throughout the tumour, that is formed by genetically normal patient stroma cells [34]. The fraction of tumour cells that are actually cancerous varies by more than 10-fold between tumours [6], meaning that without careful correction, gene expression profiles are dominated by stromal cell fraction signals, and not true differential expression within a cell type. Microdissection has been used to separate cancer cells from surrounding stroma in order to simplify analysis [11], but current thought in the field is that the stroma in PDAC is an essential and enabling, if not in itself neoplastic, component of the tumour [34], and that the examination of cancer cell expression in isolation ignores the likely important interplay between the two major synergistic components of a tumour: transformed epithelial cells, and genetically normal stroma.

Due to these challenges to GEX deconvolution of PDAC, to date only one study (by Collisson *et al*, published in 2011) has reported a breakdown of PDAC GEX into a small number of biological modules [11]. This study exam-

ined microdissected cancer cells only, and found that the transformed epithelial cells of PDAC could be placed into three major categories, based on their patterns of gene expression. Tumours from these three categories followed distinct clinical courses, and cell lines exhibited category-specific sensitivity to therapeutic drugs. As the first report to identify potential clinically relevant molecular subtypes within PDAC, the Collisson study was a significant advance in the understanding of the molecular processes at play within what was previously considered a homogeneous disease. However, it also possesses shortcomings that limit its clinical utility.

Two main issues complicate the interpretation of the Collisson classes: microdissected cancer cells were used, and therefore stromal effects would be severely attenuated; and the deconvolution technique employed was tuned to achieve sample clustering, rather than GEX deconvolution. Consequently, although the Collisson classes could be a fundamental advance in the understanding of PDAC, they necessarily do not consider the full context of the disease, and potentially have artifically identified subgroups when in reality a smooth continuum of disease types may exist. Additionally, although the Collisson tumour subgroups were observed to follow different clinical courses, they were not explicitly generated to stratify patients by outcome, and so may not have captured the full biology underlying differential survival in PDAC.

A substantial gap remains in our molecular understanding of PDAC: little is known about the core molecular processes at work within both the cancer and stroma of different tumours, and almost nothing on those processes that control patient survival following diagnosis. Such a gap in knowledge is not merely of academic interest: a better understanding of the processes affecting patient survival can lead directly to improved methods for staging, may stratify patients for customised therapies, and even suggest targets for therapeutics capable of transforming a poor-prognosis cancer into a good-prognosis one. The primary obstacle for the identification of these survival-associated processes in PDAC is one of data: a large, high-quality dataset of GEX measurements and associated well-curated CPVs is needed. The APGI cohort addresses this data problem for the identification of fundamental survival processes in PDAC. As the largest cohort of PDAC samples, with accompanying GEX and curated CPVs, in the world, it can provide the data quality and cohort size required by modern GEX deconvolution techniques.

In this chapter I describe the application of NMF for the GEX decon-

volution of genes associated with outcome. The metagenes thus identified represent orthogonal coordinately-expressed sets of genes which I then map to biological annotations, identifying the fundamental processes that may be involved in controlling the clinical course of a patient's pancreas cancer. The results of this work are directly applicable as signatures of survival time following diagnosis of PDAC, identify discrete biological processes that appear to determine outcome with pancreas cancer, and highlight fertile future avenues for research into this poorly-understood disease.

3.2 Results

Survival-associated metagenes were identified by selecting the set of genes which had GEX associated with outcome in the APGI cohort, and then performing NMF factorization to deconvolve the full matrix of gene expression signals into a small set of metagenes. Metagenes were found to fall into patterns defining two axes of outcome-associated cell state. These prognostic axes were then tested for association with clinical course and other CPVs, as well as known general prognostic signatures, and their prognostic ability was validated in a range of cancers by testing in separate cohorts. The two prognostic axes were then correlated with biological process signatures to associate axis scores with the activity of biological processes.

Cohort characteristics and subsetting

228 unique patients from the APGI cohort had both GEX and follow-up data; for the discovery of metagenes specifically associated with PDAC survival these were subset to patients with histologically confirmed PDAC, who did not suffer perioperative mortality, and were treated within Australia. This subsetting produced a homogeneous 110-patient APGI discovery cohort, which was used for all metagene discovery work.

General characteristics of both the full APGI cohort, and the 110-patient PDAC APGI discovery cohort, are summarised in Table 3.1.

Two axes predict survival with resectable pancreatic cancer in multiple cancers

Probe selection In order to focus the GEX deconvolution method on finding outcome-associated metagenes, it was necessary to filter the full set of gene expression data to only contain those genes that were likely to be associated with patient survival.

Unsupervised filtering to remove lowly-expressed and redundant probes yielded APGI cohort gene expression measurements for 13,000 genes, of which 361 were identified to be associated with time from diagnosis to disease-specific death (DSD) by sure independence screening (SIS)-feature aberration at survival times (FAST), using a complementary pair subset selection (CPSS) wrapper to reduce false positive rate. 50 variable selection runs on permuted data gave a median number of selected genes of 87.5, resulting in an estimated false-discovery rate (FDR) for the selection procedure of approximately 25%. This relatively high FDR was a consequence of the lenient selection parameters used, in an attempt to ensure that even genes for which expression was only weakly prognostic, were included.

Prognostic genes factorized into six metagenes NMF was used to reduce the complex expression patterns of 361 survival-associated genes into a small number of metagenes. NMF aims to approximate a non-negative gene \times sample GEX matrix A by a product of low-rank non-negative matrices W and H, $A \approx WH$. The gene \times metagene matrix W, termed the basis matrix, stores the contribution of each gene's expression to each metagene, whereas the metagene \times sample matrix H, termed the coefficient matrix, contains the 'expression' of each metagene in each sample. The NMF procedure is highly sensitive to the choice of the rank of W and H (the number of metagenes) – an incorrect rank will lead to metagenes inappropriately being either combined, or split.

The expression of the 361 survival-associated genes across the 110 patients of the APGI PDAC cohort was decomposed into metagenes by the sparse non-negative matrix factorization, long variant (SNMF/L) NMF algorithm. The number of metagenes (factorization rank) was automatically estimated to be 6, being the lowest rank for which the improvement in estimation error achieved by adding the next rank, was less than that observed for permuted data (Figure 3.3).

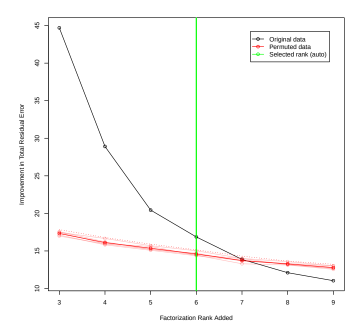


Figure 3.3: Automatic selection of factorization rank. SNMF/L was performed for varying ranks on either unpermuted data (black line) or data permuted within samples (red lines), and the improvement in total residual approximation error $||A - WH||_F$ calculated. The highest added rank for which the error improvement on unpermuted data exceeded that of permuted data plus two standard deviations (threshold shown by dotted red line) was the final selected rank (green line).

500 random restarts of rank 6 SNMF/L were then performed on the survival-associated gene matrix to yield the final factorization. The resultant clustering consensus matrix was stable (Figure 3.4), and the basis matrix W was reasonably sparse (Figure 3.5). Sparsity of the basis matrix is a desirable condition for this analysis, as it indicates that metagenes are largely distinct transcriptional modules, with little overlap in terms of shared transcripts with high loadings; SNMF/L was selected against alternative NMF algorithms as its design favours solutions with sparse W. A table of values of the basis matrix W is available as Appendix A on page 59.

Three metagenes together formed a prognostic model The transcription patterns of genes associated with survival in the APGI cohort could be

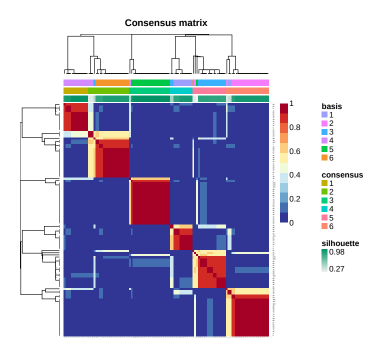


Figure 3.4: Clustering consensus matrix for the final rank-6 clustering. Colours indicate the stability of gene (in rows) and sample (in columns) clusters across random restarts of the factorization; at rank 6 this factorization was largely stable, with identical clusters assigned in all 500 random restarts to the majority of genes and samples.

decomposed into just six largely distinct metagenes. Due to the presence of false positives in the 361 screened input genes, some of the metagenes will have no strong association with outcome. To identify which of the six metagenes were ultimately predictive of patient survival, I performed LASSO regression on the 110-patient APGI discovery cohort data, using non-negative least squares (NNLS)-estimated coefficients of each of the six metagenes as marginal predictors of outcome. The LASSO regularization parameter λ was chosen by 10-fold cross-validation to be the highest value for which the mean test set partial likelihood deviance was within one standard error of the lowest mean value. This resulted in a final model in which three metagenes, MG1, MG2, and MG5, were selected as prognostic (Figure 3.6).

Prognostic metagenes define two axes of cell transcription Further investigation of the three prognostic metagenes revealed that they were associ-

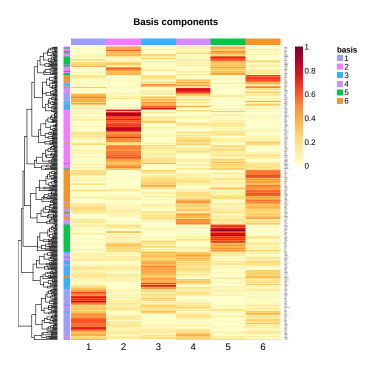


Figure 3.5: Basis matrix W of the final SNMF/L factorization. Rows represent genes, and columns metagenes, with cell colours proportional to the loading of a given gene on a given metagene. The loadings are sparse within rows, indicating that the metagenes are modular, each affecting the expression of largely distinct sets of target genes. A table of values of this basis matrix is available as Appendix A on page 59.

ated: APGI patient coefficients for pairs MG1 and MG5, and MG2 and MG6 (the latter not selected by the LASSO), were mutually exclusive (Figure 3.7, Kendall's τ test $P < 1 \times 10^{-6}$ for each pair). This suggested that both metagenes in each pair captured the signal of a single axis of cell behaviour, with one measuring activation of the axis, and the other deactivation. For subsequent work I therefore combined the signals of the metagenes within each axis, to give axis activity summaries: Axis A1 activity = MG1 coefficient – MG5 coefficient; Axis A2 activity = MG6 coefficient – MG2 coefficient. Activation values for axes A1 and A2 were uncorrelated, indicating that these axes were orthogonal processes operating in the APGI cohort tumours (Figure 3.8, Kendall's τ test P=0.21). Metagenes MG3 and MG4 also formed a mutually exclusive pair (not shown), but were not investigated further, as neither was determined to be prognostic by the metagene LASSO.

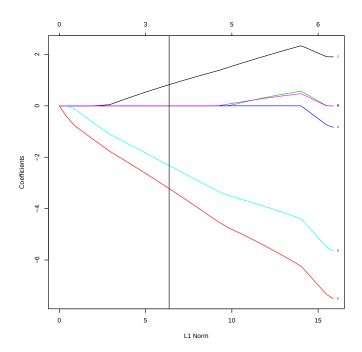


Figure 3.6: Coefficient vs penalty fit trajectories for the LASSO model predicting DSS from metagene expression. Each line represents the model coefficient for a metagene as the model is smoothly varied from a null model (L1 norm = 0), to a full unpenalised Cox fit (L1 norm \approx 16). The vertical line indicates the optimal value of L1 norm as selected by the 1SE criterion on 10-fold cross-validation; at this point in the trajectory only metagenes MG1, MG2, and MG5 contribute to prognosis estimates.

The PARSE score A repeat of the previous LASSO fit with 10-fold cross-validation (CV), this time using predictors of A1 activity, A2 activity, and the A1:A2 interaction, identified both A1 and A2, but not their interaction, as useful predictors of outcome. Coefficients from the LASSO fit were used to define a new risk score, the prognostic axis risk stratification estimate (PARSE), as PARSE score = $1.354 \times A1$ activity + $1.548 \times A2$ activity.

Exact calculation of the PARSE score requires the solution of a number of NNLS problems, which presents a potential barrier to use. An approximation to PARSE can be derived by relaxing the non-negative constraint; this approximation requires only a weighted mean of gene expression estimates, and is detailed in Appendix D on page 73.

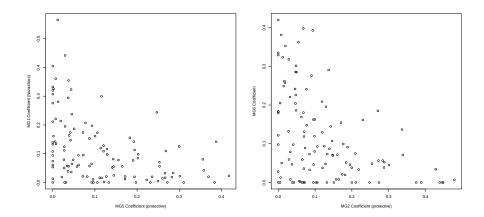


Figure 3.7: Prognostic metagenes form two axes of cell state. Metagene pairs MG1 and MG5, and MG2 and MG6, displayed mutually exclusive coefficient patterns in the APGI cohort, and could be combined to form just two axes of cell state.

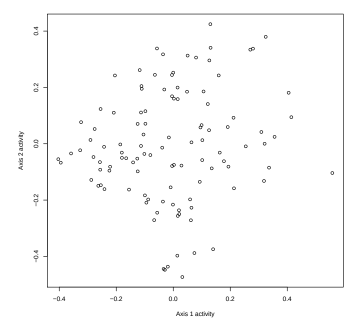


Figure 3.8: Prognostic axis signals are uncorrelated. Activity estimates of axes defined by highly correlated mutually exclusive metagene pairs (Axis A1 = MG1 - MG5, axis A2 = MG6 - MG2) were uncorrelated (Kendall τ test P=0.21), indicating that these axis signals encoded orthogonal outcomeassociated processes within tumours.

Validation of the PARSE score External validation confirmed that the PARSE score was prognostic in other cohorts, including in cancers other than PDAC. PARSE score was significantly prognostic in PDAC cohorts GSE28735 [49] (LRT P=0.0149) and The Cancer Genome Atlas (TCGA) paad (LRT P=0.0156), but not in GSE21501 [44] (LRT P=0.115). When assessed against all TCGA cancers for which at least 50 patients had both an event and complete RNASeq data, the PARSE score was also significantly prognostic for head and neck squamous cell carcinoma, kidney renal clear cell carcinoma, lower grade glioma, and lung adenocarcinoma, at a 5% familywise error rate (FWER) (Table 3.2, column a). This significant result reflected the ability of PARSE score to stratify patients into risk groups in a range of solid tumours, as illustrated in Figure 3.9.

Meta-PCNA is a 130-gene signature of cell proliferation that has been found to be generally prognostic in a number of cancer cohorts [48]. To exclude the possibility that PARSE score simply recapitulated the known meta-PCNA signature, I examined whether PARSE contributed additional prognostic information to meta-PCNA in the large TCGA cohorts. In TCGA kidney renal clear cell carcinoma, lower grade glioma, and lung adenocarcinoma, there was significant evidence that the PARSE score provided prognostic information beyond that given by meta-PCNA, at a 5% FWER (Table 3.2, column b).

Table 3.1: Characteristics of the full APGI patient cohort, and the homogenous PDAC-only subset used for signature discovery. Ordinal variables are shown as median, with quartiles in parentheses. Categorical variables for which percentages do not add up to 100% indicate the presence of minor unlisted categories. Abbreviations: AAC - ampullary adenocarcinoma; IPMN - intraductal papillary mucinous neoplasm; PNET - pancreatic neuroendocrine tumour; PR - Puerto Rico

| Characteristic | | Full APGI | Discovery |
|------------------------|------------|--------------------|--------------------|
| Number of patients | | 228 | 110 |
| Gender | Male | 54.8% | 54.6% |
| Ethnicity | Caucasian | 92.3% | 95.4% |
| | Asian | 6.4% | 4.6% |
| | African | 0.9% | 0% |
| Treatment country | Australia | 86.0% | 100% |
| | USA / PR | 12.7% | 0% |
| Age at diagnosis | (years) | 68 (60 - 75) | 67 (61 - 73) |
| Procedure | Whipple | 63.2% | 71.8% |
| Excision margin status | R0 | 76.8% | 62.7% |
| | R1 | 20.6% | 22.7% |
| | R2 | 2.6% | 14.6% |
| Histological type | PDAC | 61.8% | 100% |
| | AAC | 11.0% | 0% |
| | IPMN | 5.7% | 0% |
| | PNET | 5.7% | 0% |
| Histological grade | 1 | 12.0% | 7.3% |
| | 2 | 55.8% | 64.6% |
| | 3 | 30.1% | 27.3% |
| | 4 | 2.1% | 0.8% |
| Location | Head | 64.0% | 84.6% |
| | Ampulla | 11.4% | 0% |
| | Tail | 11.0% | 8.2% |
| | Body | 5.7% | 6.4% |
| Length | (mm) | 33.0 (24.5 - 45.0) | 35.0 (28.0 - 45.0) |
| Invasion | Perineural | 70.3% | 88.1% |
| | Vascular | 62.4% | 67.9% |
| Node involvement | | 69.3% | 77.1% |
| Disease-specific death | | 52.6% | 63.6% |
| Length of follow-up | (days) | 614 (366 - 888) | 632 (402 - 912) |

Table 3.2: The PARSE score is prognostic in a range of TCGA cancers. P-values are from likelihood ratio tests either comparing a Cox model with PARSE score as a linear predictor, to a null model (a); or a Cox model with PARSE and meta-PCNA scores as linear predictors, against one with meta-PCNA alone (b). Shaded cells are significant at a 5% FWER following Holm's correction. TCGA study codes: glm: glioblastoma multiforme; hnsc: head and neck squamous cell carcinoma; kirc: clear cell kidney carcinoma; lgg: lower grade glioma; luad: lung adenocarcinoma; lusc: lung squamous cell carcinoma; ov: ovarian serous cystadenocarcinoma.

| TCGA study | Number of events | Number of patients | Risk score P-value (a) | Improvement P-value (b) |
|------------|------------------|--------------------|---------------------------|----------------------------|
| gbm | 54 | 143 | 0.2287 | 0.1587 |
| hnsc | 124 | 367 | 8.08E-3 | 0.0108 |
| kirc | 153 | 497 | 2.03E-12 | 2.89E-3 |
| lgg | 53 | 272 | 1.49E-5 | 7.85E-3 |
| luad | 106 | 431 | 8.34E-6 | 1.04E-4 |
| lusc | 117 | 395 | 0.9624 | 0.4110 |
| ov | 115 | 251 | 0.0238 | 0.0178 |

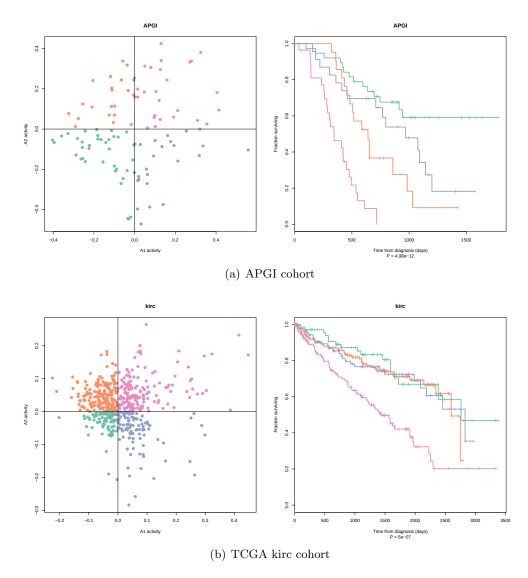


Figure 3.9: PARSE score axes define patient subgroups with differing outcome in a range of solid tumours. Activities for axes A1 and A2 of the PARSE score were calculated on the labelled cohorts, and patients split into four subgroups based on the sign of A1 and A2 activities (left panels). The four subgroups thus defined displayed significantly differing clinical courses (right panels). (continued...)

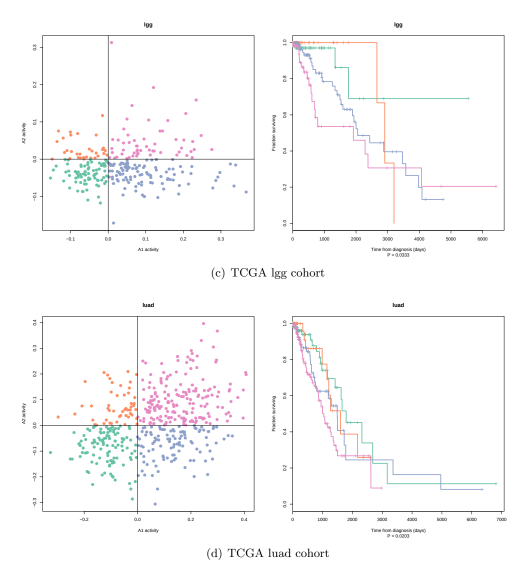


Figure 3.9: (Concluded). PARSE score axes define patient subgroups with differing outcome in a range of solid tumours. Activities for axes A1 and A2 of the PARSE score were calculated on the labelled cohorts, and patients split into four subgroups based on the sign of A1 and A2 activities (left panels). The four subgroups thus defined displayed significantly differing clinical courses (right panels).

PARSE identifies proliferation and EMT as fundamental processes controlling survival in PDAC

To link the two prognostic axes that form the PARSE score with potential underlying biology, axis activities on the APGI discovery cohort were compared

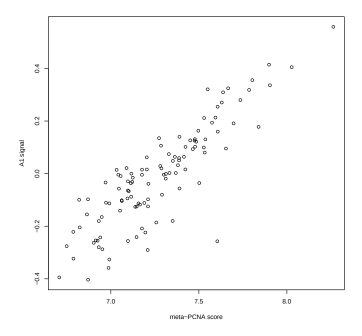


Figure 3.10: Axis A1 signal is closely associated with the meta-PCNA signature. A1 signal and meta-PCNA [48] scores were as evaluated on the APGI training set; Kendall's $\tau = 0.663$, n = 110, linear model $R^2 = 0.740$.

to clinical variates, known survival signatures, and scores for signatures from the molecular signatures database (MSigDB) [45].

MSigDB correlations, as well as comparisons to a general proliferative signature, revealed that the PARSE axis A1 (MG1 - MG5) primarily reflected the proliferative state of cells. A1 signal was very strongly correlated with meta-PCNA [48] score (Kendall's $\tau=0.663,\,n=110$, Figure 3.10), a relationship supported by its close association to cell cycle-related MSigDB signatures (Appendix B on page 70). A1 signal was also significantly positively correlated with qPure [42] estimates of cancer cell fraction in the tumour (Kendall's $\tau=0.284,\,n=110$, Table 3.3), although the strength of this association was marginal (linear model $R^2=0.155$).

Among the clinical variables tested, PARSE axis A2 (MG6 - MG2) correlated with stromal content and tumour grade: conditions of high A2 signal were associated with higher stromal content, higher grade, and shorter survival. A2 signal was positively correlated with tumour microscopic pathological grade (Holm-corrected P = 0.0067, 50 tests performed), although this

Table 3.3: Association P-values between metagenes and CPVs. P-values were either from Kendall τ tests, in the case of continuous or large ordinate clinical variates, or from ANOVA, in the case of categorical variates. Only three associations were significant at a 5% FWER level by Holm's correction; these are highlighted.

| Variable | Axis 1 | Axis 2 | |
|----------------------------|-----------------------|-----------------------|--|
| Age at diagnosis | 0.925 | 0.666 | |
| Ethnicity | 0.771 | 0.113 | |
| Gender | 0.158 | 0.010 | |
| Histological subtype | 0.697 | 0.157 | |
| Invasion | | | |
| Perineural | 0.095 | 0.225 | |
| Vascular | 0.650 | 0.071 | |
| Pack years smoked | 0.356 | 0.275 | |
| Pathological grade | $2.39{	imes}10^{-3}$ | 1.30×10^{-4} | |
| Cancer cell fraction | 2.13×10^{-4} | 4.11×10^{-4} | |
| Recurrence site | | | |
| Bone | 0.789 | 0.413 | |
| Brain | 0.430 | 0.062 | |
| Liver | 0.160 | 0.105 | |
| Lung | 0.390 | 0.713 | |
| Lymph nodes | 0.933 | 0.870 | |
| Mesentery | 0.933 | 0.121 | |
| Omentum | 0.139 | 0.082 | |
| Other | 0.193 | 0.161 | |
| Pancreatic bed | 0.887 | 0.530 | |
| Pancreas remnant | 0.534 | 0.184 | |
| Peritoneum | 0.916 | 0.015 | |
| Staging: M | 0.441 | 0.425 | |
| Staging: N | 0.252 | 0.263 | |
| Staging: T | 0.264 | 0.427 | |
| Staging: Overall stage | 0.061 | 0.236 | |
| Tumour location | 0.177 | 0.139 | |
| Tumour longest axis length | 0.844 | 0.171 | |

dependence was weak: on average, A2 signal was 0.1103 higher in grade 3 or 4 tumours over grade 1 or 2, with $R^2 = 0.119$. A2 signal was also negatively associated with tumour cancer cell fraction, the opposite of the positive relationship observed for axis A1, despite signal in both axes being positively associated with poor prognosis. This reveals a potential context dependency in the influence of stromal content on survival, where high stromal content

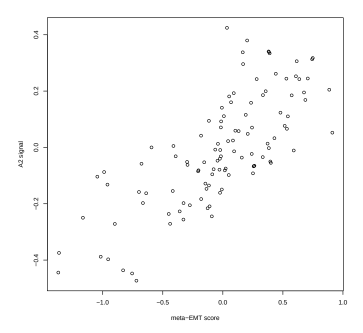


Figure 3.11: Axis A2 signal is closely associated with a signature of the EMT. A2 signal and meta-EMT [18] scores were as evaluated on the APGI training set; Kendall's $\tau = 0.568$, n = 110, linear model $R^2 = 0.557$.

of a tumour may indicate either good or poor prognosis, depending on which underlying axis is responsible.

A number of MSigDB signatures were associated with A2 signals, among them integrins, extracellular matrix (ECM) processes, and a signature for LEF1-mediated EMT (Appendix C on page 72). Prompted by the strong positive correlation between A2 and the LEF1 overexpression signature, I investigated the association between A2 signal and score for a general signature of EMT, meta-EMT [18]. meta-EMT and A2 signals were strongly positively correlated (Kendall's $\tau=0.568,\ n=110$, linear model $R^2=0.557,\ 3.11$), even when cancer cell fraction was taken into account (LRT $P=9.4\times10^{-14}$), strongly indicating that A2 signal predominantly encodes EMT activity. A potential link between A2 and inflammation may also be present: A2 signal was strongly positively correlated with the gene set variation analysis (GSVA) score for MSigDB GNF2_PTX3 (Kendall's $\tau=0.593$, Appendix C on page 72), a proxy for expression of the acute phase response protein pentraxin 3.

3.3 Discussion

1

At the molecular level, the phenomenon of cancer has long been recognised as a composite of many processes [20], however the relative importance of each process to a particular type of cancer has been largely uncertain. In pancreas cancer, a huge number of individual biomarkers are known [22], and some attempts have been made to stratify cancers into empirical molecular subtypes [11], but no studies have provided a comprehensive analysis of which basic hallmarks of cancer are actually important in determining patient outcome. This work fills that gap in knowledge, and is the first to exhaustively identify proliferation and the EMT as the major molecular processes that control survival of patients with pancreas cancer.

- PARSE = meta-PCNA + meta-EMT (+ immune + stroma?).
- Context-dependency of stroma signal
- The folly of clustering
- GSE21501 why didn't it validate?
- Broader implications of pan-cancer survival signature
- Limitations not necessarily complete. But can I be sure that the major players have been captured?
- Relevance to future work.

3.4 Methods

Cohort recruitment and ethics

2

 $^{^1\}mathrm{MP}$ Fatal: TODO: Consider comparing A1 and A2 vs meta-PCNA and meta-ECM in TCGA – are A1/A2 better than the metas? Model complexity is the same so therefore can just compare partials – woo

²MP Fatal: TODO: Cohort recruitment and ethics

Sample collection, preparation, and gene expression microarrays

3

Data preprocessing

Microarray quality control and normalization Illumina data (IDAT) files were read into Bioconductor lumi structures using the lumidat package. Seven arrays were excluded on the basis of poor signal, due to fewer than 30% of probes on these arrays having detection P-values of less than 0.01. The remaining 234 microarrays represented a range of tumour types, and were normalized as one batch using the lumi package. Normalization proceeded serially as: RMA-like background subtraction (lumiB method "bgAdjust.affy"), variance stabilizing transform (VST) (lumiT method "vst"), and quantile normalization (lumiN method "quantile").

Unsupervised probe selection Probes were excluded if they met any of the following criteria: fewer than 10% of samples with expression P-values of less than 0.01, a probe quality (from the illuminaHumanv4PROBEQUALITY field in Bioconductor package illuminaHumanv4.db) not equal to 'perfect' or 'good', missing gene annotation, or a standard deviation of normalized expression values across all samples of less than 0.03. The choice of this latter threshold is expected to yield approximately a 5% false probe rejection rate, based on an analysis of the variation between technical replicate samples. In cases where multiple post-filter microarray probes mapped to the same gene, only the probe with the highest standard deviation, as evaluated across all samples that passed quality checks, was retained. The effect of these combined filtering steps was to reduce the number of features under consideration from 47,273 probes to 13,000, one per gene.

Sample selection From the full set of 234 tumour samples that passed quality checks, eight were from four samples that had each been arrayed twice, and two were from patients with multiple conflicting CPV data. The two with conflicting CPV data were excluded from further study, and the eight replicated

³MP Fatal: TODO: Sample collection, preparation, and gene expression microarrays

samples were averaged, after multidimensional scaling (MDS) indicated that each replicate pair had very similar expression.

The 228 APGI patients for which GEX and clinical data were available were subset further to yield a homogeneous PDAC cohort, suitable for the discovery of the survival-associated processes specific to PDAC. 141 of 228 patients had pathologically confirmed PDAC; of these, five were judged to have suffered a perioperative death, and were not considered further. 110 of the 136 remaining patients were treated in hospitals in Australia, 23 in the USA, two in Italy, and one in Puerto Rico. To eliminate the potential for country-specific gene expression patterns to interact with possible differential survival between countries, only the Australian subset of the cohort was retained, resulting in 110 patients in the final APGI discovery cohort.

Summary The above preprocessing steps yielded matched CPV and resected tumour GEX data for 13,000 genes across 110 patients.

Outcome-associated gene selection

Genes that were associated with DSS were identified by SIS-FAST [15], with a CPSS wrapper to reduce the false positive rate [41]. FAST statistics for time from diagnosis to DSD were calculated using R package ahaz on standardized log-scale expression values; genes which had an absolute statistic value exceeding 7 were selected by the inner SIS-FAST procedure. The outer CPSS wrapper selected genes which were returned by at least 80% of 100 complementary paired SIS-FAST runs. Gene selection FDR was estimated by permutation: 50 repeats of the full gene selection procedure were performed on data in which patients had been randomly shuffled, and the FDR was estimated as the median number of genes selected in permuted runs, divided by the number of genes selected by the unpermuted procedure.

Rank estimation and metagene factorization

The gene \times patient expression matrix of outcome-associated genes was decomposed into metagenes by the SNMF/L procedure of [26], as implemented in R package NMF. SNMF/L is a variant of NMF, a class of procedures that decomposes a non-negative matrix A into a product of non-negative matrices W and H, $A \approx WH$. W and H typically have rank much less than A, the

effect of NMF then being to effectively reduce a large gene \times sample matrix A into smaller matrices, the gene \times metagene basis matrix W, and metagene \times sample coefficient matrix H. SNMF/L was chosen from the many NMF variants available for its design that favours solutions with sparse W: SNMF/L factorizations tend to associate each gene with a small number of metagenes, a situation that matches our biological expectation that, for most genes, expression of that gene is only associated with a small number of biological processes.

As NMF is a linear factorization, the VST-transformed expression matrix A was approximately linearized by elementwise exponentiation, $a_{i,j} \leftarrow 2^{a_{i,j}}$. To reduce the influence of large variations in baseline expression on the factorization, each row (gene) of A was then independently linearly scaled to lie between zero and one, $a_{i,j} \leftarrow (a_{i,j} - \min(a_{i,*})) \div (\max(a_{i,*}) - \min(a_{i,*}))$, where $a_{i,*}$ denotes row i of A.

Factorization rank was estimated following [14]: for test ranks ranging from 2 to 9, 5 SNMF/L decompositions were performed, each on a version of the transformed expression matrix in which rows (genes) had been independently permuted within each column (sample). Approximation error for each decomposition was calculated as $||A - WH||_F$, and the reduction in approximation error with increasing rank was compared between factorizations of the original data, and those of the 5 permuted data matrices. The highest rank for which the improvement in error achieved by adding that rank to the factorization on the original data, exceeded the improvement seen by adding that rank on the permuted data, taking into account permutation noise, was selected as the final factorization rank. Specifically, let the improvement in approximation error that results in choosing a rank i decomposition over a rank i-1 decomposition, on the unpermuted data, be $\Delta_i = ||A - W_{i-1}H_{i-1}||_F - ||A - W_iH_i||_F$. Equivalently, define Δ_i^{*j} to be the improvement observed when rank i is added to the factorization of A^{*j} , the j^{th} permutation of the data matrix: $\Delta_i^{*j} = \|A^{*j} - W_{i-1}^{*j} H_{i-1}^{*j}\|_F - \|A^{*j} - W_i^{*j} H_i^{*j}\|_F$. Denote the mean and standard deviation of Δ_i^* across all 5 permutations of the data matrix, for each i, as $\overline{\Delta_i^*}$ and $SD(\Delta_i^*)$, respectively. Then, the final selected rank k was selected as $k = \max(\{i : \Delta_i > \overline{\Delta_i^*} + 2SD(\Delta_i^*)\}).$

Following rank estimation, a final factorization of the data was performed using only the identified rank, and a larger number of random algorithm restarts, as described below. Subsequent work used this final factorization.

The SNMF/L algorithm requires parameters α and η to control regularization; for all factorizations $\alpha = 0.01$, and $\eta = \max(A)$. The default convergence criteria of the NMF package were used.

SNMF/L may not necessarily find a global optimum factorization; to address this, multiple random initializations of matrix W were made from Uniform(0, max(A)), the SNMF/L procedure was run to convergence, and the result with lowest approximation error was retained. 50 random restarts were used during rank estimation runs, and 500 for the final factorization; examination of approximation error distributions for these repeated runs indicated that these values were conservative, and factorizations were robust to the choice of random start.

Estimating metagene coefficients on new cohort data

To apply the signatures developed in this work to GEX data other than those from the APGI training set, the following procedure was used. GEX measurements from the new cohort were subset to the 361 outcome-associated genes identified by CPSS-SIS-FAST (these genes are listed in Appendix A on page 59), and transformed to a linear scale if necessary. Linear measurements were then scaled within genes to between zero and one, as was performed for metagene factorization. Genes for which no expression data were available (the genes being either filtered out in preprocessing or not measured at all) were assigned scaled expression values of zero. These manipulations yielded a gene \times sample matrix A' with rows matching the gene \times metagene basis matrix W from SNMF/L. The metagene \times sample coefficient matrix H' for the new cohort was then estimated by NNLS implemented in R package nnls, solving for each column of $a'_{*,i}$ of A' the optimization problem $h'_{*,i} = \operatorname{argmin}_x \|Wx - a'_{*,i}\|_2$, where $h'_{*,i}$ denotes column i of H'. Values of the W matrix used are available as Appendix A on page 59.

For consistency, the above procedure was used to estimate metagene coefficients H for the discovery APGI cohort, as well as all validation cohorts.

⁴Note that this parameter α is denoted β in the R NMF package; I use the symbol α here for consistency with [26]

Calculation of the PARSE score on new cohort data

Given metagene coefficients estimated as above, axis activity scores were calculated as Axis A1 activity = MG1 coefficient-MG5 coefficient; Axis A2 activity = MG6 coefficient-MG2 coefficient. PARSE scores were then made by combining axis activity estimates, as PARSE score = $1.354 \times A1$ activity + $1.548 \times A2$ activity.

Although not used in this work, a simplified procedure for the approximate calculation of PARSE scores was also developed; see Appendix D on page 73 for details.

External validation of outcome-associated metagenes

Gene expression data for accessions GSE21501 and GSE28735 were down-loaded as processed series matrix data from the National Center for Biotechnology Information (NCBI) Gene Expression Omnibus (GEO). Survival times, censoring indicators, clinical covariates (for GSE21501), and probe expression estimates were extracted from the series matrix files. Probes were annotated with gene symbols using the associated GPL annotation files, and probes with no gene annotation were discarded. If multiple probes mapped to the same gene symbol, only the probe with the highest standard deviation across all samples in a data set was retained. Finally, only probes with a standard deviation within the top 20th percentile within a data set were kept for metagene scoring.

Gene expression and outcome data for all TCGA cancers were downloaded from the public TCGA open-access repository at https://tcga-data.nci.nih.gov/tcgafiles/ftp_auth/distro_ftpusers/anonymous/tumor/, on 18 November 2014. RNASeq Version 2 Level 3 expression estimates (on an approximately linear scale) from Illumina HiSeq machines only were used, without further processing. Expression estimates were scaled within genes to between 0 and 1 separately within each TCGA cancer type. For reasons of statistical power, only TCGA cancers for which at least 50 patients had both complete RNASeq expression data, and an event, were considered in validation. Cohort paad was included despite it not meeting this criterion, to allow validation against another PDAC cohort.

For each validation data set, metagene coefficients, axis activities, and PARSE scores, were calculated as described above. Prognostic performance of the PARSE score was tested within each validation data set using likelihood ratio tests comparing a Cox model using PARSE score as the sole linear covariate, with an intercept-only Cox model.

GSVA scoring

The expression of gene sets from the MSigDB [45] were estimated on the APGI cohort using a modification of the GSVA method [21]. GSVA with default settings was used to estimate expression scores for all MSigDB gene sets in the full $13,000 \times 228$ VST-scaled APGI GEX data matrix. MSigDB contains both undirected gene sets such as metabolic pathways, in which members of the set are not expected a-priori to move in concert, and directional signatures, with paired *_UP and *_DN components that would be expected to change in coordinated and opposite patterns. Conventional analyses based on MSigDB ignore this distinction, but for this work I combined paired directional signatures to yield an overall signed estimate of signature activity. For undirected signatures, GSVA activity estimates were simply calculated using parameter abs.ranking=TRUE. In the case of paired signatures, GSVA scores were estimated separately for the *_UP and *_DN sets using parameter abs.ranking=FALSE, and the signed combined activity *_SIGNED was calculated as the *_DN score subtracted from the *_UP score. This procedure resulted in summarised activity estimates for 8,138 gene sets, many of which were highly correlated.

Gene sets with highly correlated activity scores were collapsed into compound summary sets as follows. Pairwise Pearson correlation distances between all scores were calculated as $d_{i,j} = \frac{1}{2}(1 - \cos(s_i, s_j))$, and were used to cluster gene sets using R hclust and complete linkage. R cutree identified clusters of highly similar gene sets, using a distance threshold of 0.02; gene set activities within each cluster were merged by taking median values across all samples, to form a new merged gene set activity estimate. Following merging, 7,633 single and compound gene set activity estimates remained across 228 samples.

meta-PCNA and meta-ECM score calculation

Scores for the meta-PCNA signature were calculated from GEX data as described in [48]. To estimate meta-ECM scores, log-scale GEX data were me-

dian centered, and then median values across samples were calculated for all genes in the two lists of [18] Table S3, to yield EMT-overexpressed, and EMT-underexpressed, gene list median expression estimates per sample. The meta-ECM score was then calculated as the EMT-overexpressed median value, less the EMT-underexpressed median value.

Prognostic axis functional characterization

Clinical variate comparisons Prognostic axis activities calculated on the APGI data were tested for association with a restricted set of the available APGI CPVs, as outlined in Table 3.4. Numeric variables were tested for association with each axis by Kendall's τ test; factor and boolean variables using ANOVA with the CPV as the explanatory variable. 50 tests in total were performed (25 variables, 2 axes), and P-values were corrected together using the Holm-Bonferroni procedure [24]. Corrected P-values of less than 0.05 were considered significant.

MSigDB signature score comparisons Kendall correlation coefficients were calculated between axis activity estimates and GSVA scores for MSigDB gene sets, on the APGI expression dataset. A subset of the full MSigDB was used, as outlined in Table 3.5. Absolute correlations of greater than 0.5 were deemed substantive and reported for further characterisation.

Attribution of work

Data for the APGI discovery cohort were generated as part of the APGI project, under the umbrella of the International Cancer Genome Consortium (ICGC). The generation of these data was a huge team effort, of which I only played a small part. However, both conception of the project, and all steps subsequent to raw data generation, from low level processing of IDAT files through to analysis planning, signature development, testing, and interpretation, were performed solely by me.

Table 3.4: CPVs tested for association with prognostic axis signals.

| Clinical variate | Trno | |
|----------------------------|---------|--|
| Ciliical variate | Type | |
| Age at diagnosis | Ordinal | |
| Ethnicity | Factor | |
| Gender | Boolean | |
| Histological subtype | Factor | |
| Invasion: | | |
| Perineural | Boolean | |
| Vascular | Boolean | |
| Pack years smoked | Ordinal | |
| Pathological grade | Boolean | |
| Recurrence found in: | | |
| Bone | Boolean | |
| Brain | Boolean | |
| Liver | Boolean | |
| Lung | Boolean | |
| Lymph nodes | Boolean | |
| Mesentery | Boolean | |
| Omentum | Boolean | |
| Other | Boolean | |
| Pancreas remnant | Boolean | |
| Pancreatic bed | Boolean | |
| Peritoneum | Boolean | |
| Staging: M | Boolean | |
| Staging: N | Boolean | |
| Staging: T | Factor | |
| Staging: Overall stage | Factor | |
| Tumour location | Boolean | |
| Tumour longest axis length | Ordinal | |

Table 3.5: The subset of MSigDB signatures tested for association with axis activities. Within each MSigDB class, only those matching the indicated inclusion pattern were tested. * represents a wildcard; — matches nothing.

| MSigDB class | Signature name inclusion pattern |
|--------------|----------------------------------|
| c1 | _ |
| c2 | KEGG_*, PID_*, REACTOME_* |
| c3 | * |
| c4 | GNF2_*, MORF_* |
| c5 | * |
| c6 | * |
| c7 | * |

Chapter 4

Comparative genomics

Outline ideas:

- Introduction / overview:
 - The use of models in PC (very brief)
 - Specific models used in PC, with strong focus on the most common (KPC), and derivates. Cover ease-of-use briefly.
 - Current knowledge re: how appropriate the models are. Consider histology, genetic features, disease progress (incl. metastatic potential), response to therapy. Highlight gap in genetic information, and relevance to response to therapy.
 - Brief overview of known genetic features of human disease. Raise possibility of subtypes.
 - Wrap-up with overview of project:
 - 1. Collect matched tumour-normal DNA from a range of GEMMs.
 - 2. Sequence and determine conserved model-specific and general patterns of somatic mutation.
 - 3. Compare observed patterns to human disease.
 - * Are genetic features of human disease recapitulated generally in the models?
 - * Does a single model match the genetic features of human disease much better than the others?
 - * Do specific models serve as simulations of certain subtypes of human disease?

- Overall thesis for this work:

Matching patterns of genetic alterations in mouse models of pancreatic cancer to those seen in human disease can inform researchers as to which models are generally best, and which best match specific patient types.

Sub-theses:

- * The patterns of mutations seen in common mouse models of pancreatic cancer match those consistently seen in human disease.
- * Different mouse models possess different mutation spectra, and models may be close fits to specific genetic subtypes of patients.
- Results
 - 1. Somatic SNV and indels
 - 2. CNV and LOH
- Conclusion

4.1 Methods

Models

Sample Origin and Processing

Sequencing

QC

Mapping

For initial mapping, all lanes were processed independently. SHRiMP was used to map colourspace reads to the mm10 genome using 'all-contigs' and 'single-best-mapping' options. Unpaired reads in the source fastq files were mapped as single reads; paired reads were mapped with pair mode 'opp-in', and a per-fastq insert size distribution estimated from a normal distribution fit to insert sizes of the first 10,000 reads. Likely duplicate reads were marked using Picard MarkDuplicates on each individual lane binary sequence alignment / map file (BAM), using an optical duplicate pixel distance parameter of 10.

Lane BAMs were progressively merged: first, duplicate lane BAMs for a given mouse and sample type (tumour or normal) were combined, then tumour and normal BAMs for a given mouse, and finally combined tumour-normal BAMs for all mice. Prior to each level of merging, the Genome analysis toolkit (GATK) was used to separately perform local alignment and base quality score recalibration (LA-BQSR) on each input BAM. Finally, the full experiment BAM file was recalibrated with LA-BQSR, and then split by mouse and sample type for analysis, yielding 62 paired tumour and normal final BAMs.

Somatic SNV and Indel Detection

muTect and Strelka were used separately to detect somatic single nucleotide variants (SNVs) and insertion / deletion events (indels) in individual mouse tumour and normal BAMs. muTect was supplied default parameters; Strelka used the parameter settings given in listing 4.1; these are the default parameters as recommended for use with the BWA mapper, with the exception that in this work isSkipDepthFilters was set to 1.

Listing 4.1: Strelka configuration file used for SNV / indel detection

```
[user]
isSkipDepthFilters = 1
maxInputDepth = 10000
depthFilterMultiple = 3.0
snvMaxFilteredBasecallFrac = 0.4
snvMaxSpanningDeletionFrac = 0.75
indelMaxRefRepeat = 8
indelMaxWindowFilteredBasecallFrac = 0.3
indelMaxIntHpolLength = 14
ssnvPrior = 0.000001
sindelPrior = 0.000001
ssnvNoise = 0.0000005
sindelNoise = 0.000001
ssnvNoiseStrandBiasFrac = 0.5
minTier1Mapq = 20
minTier2Mapq = 5
ssnvQuality_LowerBound = 15
sindelQuality_LowerBound = 30
isWriteRealignedBam = 0
binSize = 25000000
```

CNV and LOH Detection

Overview:

- Very brief background of CNV and LOH in tumours, and the possibility of detection from NGS data. Maybe pull in the hallmarks paper, or perhaps specific PC / GEMM examples.
- Brief overview of existing techniques and why unsuited?
 - CNV:
 - * Exome pulldown complication
 - * Ill-posed nature of problem
 - * Human-specific methods
 - * Outbred population-specific methods
 - LOH:
 - * That Bayesian thing. Unfortunately affected by CNV, which is unknown.

Loss of heterozygosity at individual loci

This work took a simple approach to identify loci with significant evidence of loss of heterozygosity (LOH) in a tumour sample: locate high-confidence heterozygous loci in matched normal DNA, and then test only these heterozygous loci for a significant change in allelic fraction between matched tumour and normal samples. In regions of the genome with ploidy 2n and below, such allelic imbalance is indicative of LOH, even in the presence of unknown levels of diploid genome contamination.

Identifying heterozygous loci in normal DNA High-confidence heterozygous loci in normal DNA were identified by comparing posterior genotype likelihoods using a Bayesian model comparison (BMC) approach. BMC is a procedure for deciding which of two competing models is better favoured by the observed data; here the two models are, for a given locus: 'the locus is homozygous' (model HOM), and 'the locus is heterozygous' (model HET). The likelihoods of these two models (assessed on the reads observed at a locus) can be used to calculate a Bayes factor, which encodes which of the two models is better supported by the data at that locus, and how strongly. More

formally, we partition the ten possible diploid genotypes at a locus into two classes, Hom and Het:

$$Hom = \{AA, CC, GG, TT\} \tag{4.1}$$

$$Het = \{AC, AG, AT, CG, CT, GT\}$$

$$(4.2)$$

The two models, HOM and HET, may be written

$$HOM: G \in Hom$$
 (4.3)

$$HET: G \in Het$$
 (4.4)

where G is the true genotype at the locus. The Bayes factor K comparing HOM and HET is then

$$K = \frac{\mathcal{L}(HET)}{\mathcal{L}(HOM)} \tag{4.5}$$

$$= \frac{Pr(D|G \in Het)}{Pr(D|G \in Hom)} \tag{4.6}$$

$$= \frac{Pr(D|G \in Het)}{Pr(D|G \in Hom)}$$

$$= \frac{\sum_{g \in Het} Pr(D|G = g)Pr(G = g|G \in Het)}{\sum_{g \in Hom} Pr(D|G = g)Pr(G = g|G \in Hom)}$$

$$(4.6)$$

with D being the reads at the locus. We make the simplifying assumption that all genotypes in each of *Hom* and *Het* are equally likely, so that all $Pr(G = g|G \in X) = \frac{1}{\|X\|}$ for $X \in \{Hom, Het\}$. Then

$$K = \frac{\frac{1}{\|Het\|} \sum_{g \in Het} Pr(D|G = g)}{\frac{1}{\|Hom\|} \sum_{g \in Hom} Pr(D|G = g)}$$
(4.8)

$$= \frac{\frac{1}{\|Het\|} \sum_{g \in Het} \mathcal{L}(G = g|D)}{\frac{1}{\|Hom\|} \sum_{g \in Hom} \mathcal{L}(G = g|D)}$$

$$(4.9)$$

encodes the weight of evidence for the observed read data D favouring a locus being heterozygous over homozygous, and a value exceeding a given threshold is taken as significant evidence that the locus under consideration is heterozygous.

An implementation of the above heterozygous locus detection method is given in algorithm 1. The input posterior genotype likelihoods $\mathcal{L}(G=g|D)$ are supplied by samtools mpileup -q 20 -Q 20 -v -u operating on per-mouse normal sample BAMs, and the minimum value of K for a locus to be called as heterozygous is $\exp(minscore)$. Two additional filters are also employed in the algorithm: a locus is not reported as heterozygous if either the total read depth at the locus is less than *mindepth*, or if the difference in samtools-supplied log likelihood between the top two genotypes is less than *mindelta* nats. The latter filter is used to exclude any problem loci with an apparent triallelic state. ¹

Identifying tumour LOH at known normal heterozygous loci Given a set of loci that are known to be heterozygous with high confidence in the normal DNA of a given mouse, it is straightforward to test for LOH in the tumour DNA of the same mouse, provided the tumour ploidy at the locus is 2n or less. Considering only a single heterozygous locus, reads from a normal DNA sample will predominantly be for the two bases constituting the heterozygous genotype, possibly with a small number of reads from other bases due to sequencing or mapping errors. The number of reads for the two genotype bases may be quite different, as the exome capture processing step may favour one allele over the other, and lead to allelic bias in the observed read fractions. However, under the null hypothesis of no LOH and no mutation at the locus in the tumour DNA, if the tumour ploidy at the locus is 2n or less, then the relative proportions of reads for the two genotype bases should be the same in both the tumour and the normal samples. This null hypothesis can be tested using a contingency test comparing two binomial proportions; for this work I used the two sided Z-pooled test as implemented in R package Exact.

In the general case with potential normal cell contamination of the tumour sample, it is not possible to use allelic imbalance as an indicator of LOH if the local copy number exceeds two. For example, in the triploid case, a LOH haplotype AAA, and a non-LOH haplotype AAB, both exhibit allelic imbalance. For this reason, allelic imbalance calls from the above test must be interpreted in the context of local copy number variation (CNV) estimates from the next procedure, and LOH calls only made if allelic imbalance is detected in regions of copy number 2n or less.

Copy number variation at individual loci

Problem description Considering a single locus, either a single nucleotide or a contiguous stretch of DNA, the expected number of reads from a sequencing experiment that map to that locus is proportional to the copy number of

¹MP Fatal: give instantiation values for the algo somewhere

Data: Total sequence depth at the locus D, minimum depth for call mindepth, list of alternate alleles A, list of Phred-scaled genotype likelihoods L, minimum likelihood difference in nats between top two genotypes mindelta, minimum Bayes factor in nats for heterozygous to be called over homozygous minscore.

Result: A boolean: true if the locus is called heterozygous, false if it is not.

```
begin
    if D \leq mindepth then
     return false;
    end
    // Convert Phred-scaled likelihoods to nats
    for i \leftarrow 1 to ||L|| do
        L_i \longleftarrow -\frac{1}{10}\log(10)L_i;
    end
    // Ensure the likelihood difference between the two most
        likely genotypes is at least mindelta.
    L^* \longleftarrow L sorted in decreasing order;
    if L_1^* - L_2^* \leq mindelta then
       return false;
    end
    // Calculate combined likelihoods for heterozygous and
        homozygous genotypes
    switch ||A|| do
        case 2
            L_{het} \longleftarrow L_2;
            L_{hom} \longleftarrow \log \left(\frac{1}{2} \sum_{i \in \{1,3\}} \exp(L_i)\right);
        \mathbf{end}
        case 3
            L_{het} \longleftarrow \log \left( \frac{1}{6} \sum_{i \in \{2,4,5,7,8,9\}} \exp(L_i) \right);
L_{hom} \longleftarrow \log \left( \frac{1}{4} \sum_{i \in \{1,3,6,10\}} \exp(L_i) \right);
        end
        case default
         return false;
        end
    endsw
    // Compute the Bayes factor for heterozygous vs
        homozygous, and compare to the threshold
    if L_{het} - L_{hom} \leq minscore then
        return false;
    end
    return true;
end
```

Algorithm 1: Determine a locus is heterozygous

the locus in the DNA input for sequencing. Based on this relationship it is – in principle – possible to estimate copy number from sequencing data, however a number of complicating factors are present, related to sequence 'mappability', exon capture affinity, sample contamination, and problem indeterminancy.

There are many regions in mammalian genomes for which it is challenging to map reads. These regions may be either poorly characterised themselves in the reference genome, or may be sufficiently like other parts of the genome for an unambiguous mapping to be impossible with the short and error-prone reads produced by next-generation sequencing (NGS) technologies. Most processing pipelines discard such ambiguous reads, with the net effect that difficult-to-map regions of the genome have much lower read depth than would be expected based on the quantity of DNA for those regions present as input to the sequencing procedure. Copy-number analysis techniques need to take this 'mappability' bias into account, or regions of reference DNA that are challenging to map may falsely be reported to undergo copy number loss.

A similar effect to 'mappability' bias is additionally present in datasets generated by exome sequencing. The process of exome enrichment necessarily favours certain regions of the reference genome (hopefully, the exome), over others. This enrichment is always imperfect: some non-target DNA will persist through the procedure, and not all target regions will be retained to the same degree. The ultimate effect of the exome enrichment procedure is to introduce an additional per-locus bias, 'exon capture affinity' that requires correction before copy number calls can be made. Unlike for 'mappability bias', ignoring exon capture affinity bias can lead to either false copy number loss or false copy number gain calls.

Contamination of tumour DNA is a universal problem in solid tumour sequencing. This contamination may be with non-cancerous diploid DNA, or alternate cancer genotypes present in the same sample, or both. In the case of CNV estimation based on read depth, the presence of contaminating diploid DNA causes a shrinkage of the observed CNV profile towards that of diploid cells, and reduces the signal-to-noise ratio (SNR) of the copy number estimates. CNV callers aware of this effect must take this effect into account in their calls, and may also be required to estimate the fraction of contaminating normal DNA. In tumour samples containing multiple tumour genotypes, with varying locus copy numbers, CNV estimates are for the mean copy number of the genotypes, weighted by their prevalence in the sample. In such cases,

deconvolution of the signal into its component genotypes based on a single sample of the tumour is impossible without the benefit of additional external information.

Ultimately, without knowledge of the number of cells input into the sequencing procedure, CNV estimation from NGS data is a fundamentally indeterminate problem. This is easily seen by considering the case of a hypothetical fully haploid tumour: the read counts of all loci will be completely consistent with those of a normal diploid sample. Without observing that the quantity of DNA present per input tumour cell is half that of a diploid cell, the haploid tumour and diploid normal samples would be completely indistinguishable. Information on the number of cells used for extraction is very seldom available, and so in almost all cases additional assumptions are required to assign absolute copy number to NGS read depth data.

Taking all the above complications into account, I developed an organism-agnostic CNV detection procedure for exome or whole genome sequencing (WGS) data that uses NGS read depths as input.

CNV model and test development The mathematical setup of the procedure is as follows. We reserve upper case variable symbols for random variables, and use lower case equivalents for observed values of these random variables. Consider m disjoint loci on the reference genome; these loci may be individual base pairs or contiguous regions. For a single matched tumour-normal sample pair, let the number of reads that were mapped to locus $i \in \{1 \dots m\}$ be n_i for the normal sample, and t_i for the tumour sample. Denote the total read depths at all examined loci as d_N and d_T , $d_N = \sum_{i=1}^m n_i$, $d_T = \sum_{i=1}^m t_i$. To consider normal DNA contamination effects, we suppose that the tumour sample is actually a mixture of normal cell diploid DNA, and cancer cell DNA, where the fraction of cancer cells in the sample is the unknown quantity $f \in (0,1]$. Loci are subject to differential exome enrichment, locus size, and mapping biases, which are combined into the single per-locus quantity b_i , such that $\langle N_i \rangle \propto b_i$, $\langle T_i \rangle \propto b_i$, and $\sum_{i=1}^m b_i = 1$.

We model the process of reads in NGS as a Bernoulli scheme, and use the weak dependence between read depths at different sites to derive a per-locus Poisson approximation. In this model the sequencer has a fixed s total physical sites available for sequencing; in the SOLiD 4 system these sites correspond to positions on the sequencing slide. Some of these sites yield observed sequence

that is then mapped and used to estimate read depth, however many of them do not produce sequence reads, either because they are never populated with DNA, or because they fail low-level quality checks. We suppose that these failed sites occur independent of the DNA sequence, and at a rate of r_F among all available sites. Then, a given physical sequencing site can either fail to yield sequence, with probability r_F , or it can produce observed sequence for one of m loci, each at probability $(1 - r_F)b_i$, for $i \in \{1 \dots m\}$. This per-site categorical distribution, when sampled for each of s independent sites, results in a multinomial distribution on read depths,

$$(N_F, N_1, \dots, N_m) \sim Multi(s, (r_F, (1 - r_F)b_1, \dots, (1 - r_F)b_m))$$
 (4.10)

where N_F is the number of failed sites (not observed), and N_i is the number of reads observed for locus i. The multinomial distribution induces a negative dependency on the number of reads observed at different loci, as the total read count s is fixed. However, for m large, or site failure rate r_F large[35], these negative dependencies are small, and

$$N_i \stackrel{.}{\sim} Pois\left(s(1-r_F)b_i\right)$$
 (4.11)

The quantity $s(1 - r_F) = \langle D_N \rangle$ is unknown, and we approximate it with the observed value d_N . Therefore, the final approximate model for read depth in the normal sample is

$$N_i \stackrel{.}{\sim} Pois\left(d_N b_i\right)$$
 (4.12)

For the tumour sample, the expression for the Poisson rate parameter is more complex than in the normal case, as locus copy number is no longer assumed constant. Ignoring for the moment the possibility of diploid DNA contamination in the tumour sample (i.e. let f = 1), and following the derivation used in the normal case, we find that the number of reads at locus i in pure tumour sample is distributed as

$$T_i \stackrel{f=1}{\sim} Pois\left(d_T b_i c_i k_{pure}\right) \tag{4.13}$$

where c_i is the copy number of locus i in the tumour DNA, relative to diploid cells. $k_{pure} = 1/\sum_j b_j c_j$ is a normalization factor that ensures $\langle \sum T_i \rangle = d_T$. Now considering possible diploid DNA contamination, if tumour cells are

present at a fraction f, with the remainder diploid cells, the tumour locus read count is distributed as

$$T_i \stackrel{.}{\sim} Pois(kd_Tb_i(1+f(c_i-1)))$$
 (4.14)

Here k is no longer a simple normalization factor like k_{pure} , but is a value that involves sample purity and cancer cell DNA content. ²

The variable k is more than a convenient normalization constant: it encodes the signal expected of diploid loci in the tumour cells, and therefore controls the absolute copy numbers called by the procedure. To see this, observe that the pure tumour ploidy signal is $c_i k$, and therefore that tumour ploidy relative to 2n, c_i , is completely confounded with k. As noted earlier, without knowing the number of input cells in the tumour sample, it is impossible to determine absolute ploidy from NGS depth data, and so there is no way to conclusively determine the correct value for k. In this work I used the heuristic that the most common ploidy in a tumour cell should be diploid, and therefore selected values for k to ensure that the most common CNV call would be diploid (ie No CNV). This heuristic will almost certainly be wrong in cases, but is necessary given the fundamentally indeterminate nature of the CNV problem. Interpretation of the results of this CNV calling procedure must take into account the possibility that k is mis-specified, and that all CNV calls should be shifted appropriately.

Given the above approximate Poisson distributions for normal and tumour read depths as a function of locus ploidy, I developed a per-locus CNV test based on a ratio test for two Poisson-distributed random variables. Let R_i be the ratio of the read appearance rates at locus i in tumour and normal samples,

$$R_{i} = \frac{kD_{T}b_{i}\left(1 + f\left(c_{i} - 1\right)\right)}{D_{N}b_{i}} \tag{4.15}$$

$$= \frac{D_T}{D_N} \left(k \left(1 + f \left(c_i - 1 \right) \right) \right) \tag{4.16}$$

Then, the null hypothesis of no CNV at locus i, $H_0: c_i = 1$, is equivalent to a hypothesis on R_i ,

$$H_0: R_i = \frac{D_T}{D_N}k \tag{4.17}$$

²MP Fatal: Add the derivation in somewhere – perhaps an appendix. It's a pain in the arse so probs want to avoid the main text.

We test this hypothesis on R_i using the W_5 statistic of [19],

$$W_5(X_0, X_1) = \frac{2\left(\sqrt{X_0 + 3/8} - \sqrt{r_{H0}(X_1 + 3/8)}\right)}{\sqrt{1 + r_{H0}}}$$
(4.18)

where $r_{H0} = \frac{d_T}{d_N} \hat{k}$. This statistic is asymptotically normally distributed, so the one-sided copy number gain P-value $(H_1 : c_i > 1)$ is

$$p_{gain} = 1 - \Phi\left(w_5(t_i, n_i)\right) \tag{4.19}$$

where w_5 is the observed value of the statistic W_5 , and Φ is the cumulative distribution function of the standard normal distribution. W_5 is symmetric, so the one-sided P-value for copy number loss is

$$p_{loss} = \Phi\left(w_5(t_i, n_i)\right) \tag{4.20}$$

and the combined two-sided P-value for CNV at locus i is

$$p_{CNV} = \begin{cases} 2p_{loss} & \text{if } t_i/n_i < r_{H0} \\ 2p_{gain} & \text{if } t_i/n_i \ge r_{H0} \end{cases}$$

$$(4.21)$$

 ${f CNV}$ detection procedure Pseudocode for the implementation of perlocus CNV detection is given in algorithm 2. 3

Combining calls from adjacent loci

CNV and LOH are broad genomic events that typically affect many adjacent loci together, yet the methods presented in the preceding sections consider each locus in isolation. By examining loci separately, we disregard important information: that the CNV and LOH status of nearby loci is strongly correlated. Intuitively, by leveraging these local correlations and combining results from neighbouring loci, we can achieve more accurate CNV and LOH detection than if each locus were considered alone.

A number of approaches could be used to smooth LOH and CNV calls and share information between neighbouring loci; in this work I chose the hidden Markov model (HMM) formalism and extended the Pounds-Morris FDR estimator[37] to the locality-sensitive case. The Pounds-Morris procedure fits the observed distribution of test P-values to a mixture of Uniform and Beta distributions. The Uniform distribution models the expected distribution of

³MP Fatal: Add specific value of mindepth used

Data: An m-vector of normal locus read depths n, an m-vector of tumour locus read depths t, minimum normal sample depth mindepth.

Result: An m-vector of floats: for each locus, the one-sided P-value for CNV loss at that locus, p_{loss} .

```
begin
    d_N \longleftarrow \sum_{i=1}^m n_i; 
d_T \longleftarrow \sum_{i=1}^m t_i;
     // Estimate k so that the modal ploidy signal will be
          called as diploid
     s \leftarrow \{(t_i/d_T) \div (n_i/d_N) : i \in \{1 \dots m\} \land n_i \ge mindepth\};
     \hat{S} \longleftarrow KDE(s);
     \hat{k} \longleftarrow mode(\hat{S});
     // Calculate P-values. W_5 and \Phi are as defined in the
    r_{H0} \longleftarrow \frac{d_T}{d_N} \hat{k};
     p \leftarrow m-vector of NAs;
     for i \leftarrow 1 to m do
          if n_i \geq mindepth then
             p_i \longleftarrow \Phi\left(W_5(t_i, n_i)\right);
          end
     end
    return p;
```

Algorithm 2: Calculate CNV loss P-values

end

P-values under the null hypothesis, whereas the Beta distribution approximately fits the highly left-skewed distribution of P-values expected of tests for which the null hypothesis is false. After the observed distribution of P-values has been fit to the Beta-Uniform mixture model, the FDR associated with a given P-value can be estimated from the densities of the Beta and Uniform component distributions at that P-value.

The original Pounds-Morris procedure considers all tests as equivalent, and thus integrates no locality information, but for the LOH case combining the procedure with the locality-sensitive HMM is straightforward (figure 4.1). The HMM moves between two discrete states: *No LOH*, and *LOH*. The *No LOH* state emits a Uniform distribution of P-values, as expected under the null hypothesis of no LOH, whereas the *LOH* state emits a left-skewed Beta distribution of P-values, approximating the P-value distribution observed for

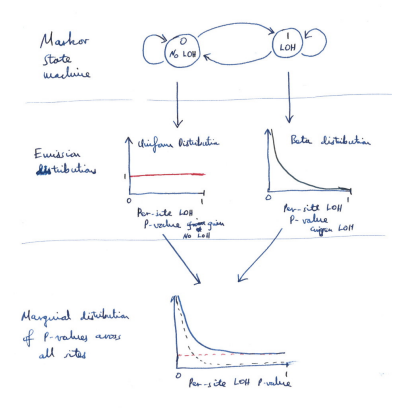


Figure 4.1: Locality-sensitive FDR estimation of LOH calls using a Markov chain Beta-Uniform mixture model.

loci at which the null hypothesis is false. Observed P-values at a chain of adjacent loci are fit to the HMM by standard algorithms implemented in R package depmixS4, and the posterior probability of a locus being in state No LOH directly gives the locality-adjusted FDR for that locus. In cases where too few extreme P-values are present to reliably estimate the parameters of the Beta distribution, the fit becomes unstable and FDR estimates potentially unreliable. To handle this situation gracefully, the method fits both the full No LOH / LOH model, and a restricted No LOH only model, and selects the model with the superior BIC.

Extension of the procedure to the CNV case requires three states: Diploid, Loss, and Gain (figure 4.2). We take advantage of the W_5 statistic's symmetry and fit the HMM to the one-sided p_{loss} CNV P-values; CNV loss is then indicated by P-values near zero, and CNV gain by P-values near one. The

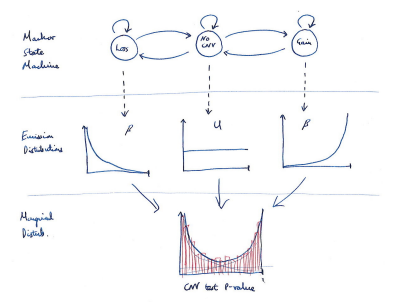


Figure 4.2: Locality-sensitive FDR estimation of CNV calls using a Markov chain double-Beta-Uniform mixture model.

Loss and Gain states are modelled by Beta distributions, left-skewed in the Loss case, and right-skewed in the Gain case. The posterior probability of a locus being in state Diploid then gives the overall FDR for a CNV call at that locus. BIC model selection is performed as for the LOH case, except in this case four models are compared: Diploid, Diploid / Loss, Diploid / Gain, and Loss / Diploid / Gain.

Although the given procedure is simple in formulation, some additional complexities were required for a practical implementation, all related to the high degree of flexibility of the Beta distribution. The Uniform distribution is a special case of the Beta distribution, and therefore in cases where the distribution of P-values is near Uniform (ie. all sites appear to satisfy the null hypothesis), the fitting problem is ill-posed. This issue was resolved by enforcing Beta parameters $\alpha \leq 0.95$ for LOH and CNV loss detection, and $\beta \leq 0.95$ for CNV gain detection. For FDR correction of CNV P-values, structural zeros were placed on the probabilities of direct transitions between Loss and Gain states (figure 4.2); although such transitions are biologically plausible, they were found to contribute to unstable fits in noisy data.

Chapter 5

Conclusion

Appendices

Appendix A

Basis matrix W for the six survival-associated metagenes

| | MG1 | MG2 | MG3 | MG4 | MG5 | MG6 |
|---------|--------|--------|--------|--------|--------|--------|
| A4GALT | 0.0295 | 0.0000 | 1.2977 | 0.0788 | 0.3625 | 0.5232 |
| A4GNT | 0.0000 | 0.7419 | 0.0483 | 0.0539 | 0.3720 | 0.0666 |
| ABHD16A | 0.6623 | 0.7249 | 0.0000 | 0.0000 | 0.5217 | 0.2210 |
| ABHD5 | 0.1481 | 0.7473 | 0.0000 | 0.7478 | 0.3988 | 1.1727 |
| ABLIM1 | 0.0145 | 0.9135 | 0.3159 | 0.0000 | 0.6066 | 0.3419 |
| ACE | 0.0333 | 0.8332 | 0.0536 | 0.0000 | 0.0000 | 0.1814 |
| ACKR3 | 0.0029 | 0.0000 | 0.3821 | 0.3591 | 0.2080 | 0.5772 |
| ACYP2 | 0.2481 | 0.8949 | 0.0000 | 0.2334 | 0.8454 | 0.4110 |
| ADH1A | 0.0730 | 0.4440 | 0.0052 | 0.1009 | 0.6614 | 0.0000 |
| ADM | 0.0000 | 0.0000 | 0.5168 | 0.5137 | 0.0000 | 0.3570 |
| AGRP | 0.0000 | 0.0000 | 0.0000 | 0.6786 | 0.0000 | 0.1744 |
| AKIP1 | 0.6365 | 0.2394 | 0.6036 | 0.7118 | 0.7849 | 0.7168 |
| AKR1A1 | 0.2470 | 1.0849 | 0.2633 | 0.2921 | 0.6588 | 0.4524 |
| ALDH5A1 | 0.0988 | 0.9930 | 0.5463 | 0.0566 | 0.8968 | 0.2222 |
| ALOX5AP | 0.0525 | 0.0084 | 0.0147 | 1.2654 | 0.3441 | 0.7138 |
| AMOT | 0.0653 | 0.8246 | 0.1374 | 0.5176 | 0.4311 | 0.5705 |
| ANGPTL2 | 0.0000 | 0.0000 | 0.3694 | 0.8726 | 0.1807 | 0.9222 |
| ANGPTL4 | 0.1789 | 0.0000 | 0.4156 | 0.0461 | 0.0260 | 0.3906 |
| ANKLE2 | 0.7503 | 0.1422 | 0.6238 | 0.5082 | 0.1879 | 0.3839 |
| ANKRD22 | 0.4067 | 1.3536 | 0.1731 | 0.2672 | 0.0381 | 0.2229 |
| ANKRD37 | 0.0562 | 0.1817 | 0.2150 | 0.7249 | 0.0129 | 0.5715 |
| | | | | | | |

| ANLN | 1.1696 | 0.2368 | 0.0796 | 0.0772 | 0.0000 | 0.7203 |
|----------|--------|--------|--------|--------|--------|--------|
| APCDD1 | 0.0000 | 0.1375 | 0.1494 | 0.1308 | 0.5957 | 0.8366 |
| APCS | 0.0000 | 0.0306 | 0.1569 | 0.1001 | 0.1638 | 0.3521 |
| ARFGAP3 | 0.0252 | 0.2988 | 0.5370 | 0.8377 | 0.4872 | 0.5353 |
| ARHGAP24 | 0.0628 | 1.0614 | 0.0157 | 0.7487 | 1.1007 | 0.6209 |
| ARHGEF19 | 0.0837 | 0.0833 | 1.2033 | 0.5242 | 0.4520 | 0.5071 |
| ARL4C | 0.0000 | 0.0171 | 0.3025 | 0.4910 | 0.2953 | 1.2264 |
| ARSD | 0.1550 | 1.2389 | 0.1919 | 0.0000 | 0.2154 | 0.1439 |
| ASPM | 1.1736 | 0.3897 | 0.2026 | 0.1743 | 0.0380 | 0.0396 |
| ATAD2 | 0.9358 | 0.0696 | 0.1136 | 0.0265 | 0.1092 | 0.3070 |
| ATF7IP2 | 0.0000 | 0.2019 | 0.1165 | 0.0000 | 0.0319 | 0.0000 |
| ATL3 | 0.6429 | 0.0252 | 0.1566 | 0.4867 | 0.2467 | 0.2863 |
| AURKB | 1.0027 | 0.1107 | 0.1351 | 0.0000 | 0.0096 | 0.0000 |
| AXIN2 | 0.0000 | 0.5221 | 0.4413 | 0.1313 | 0.8077 | 0.2911 |
| B3GALTL | 0.3601 | 0.3276 | 0.5636 | 0.3806 | 0.4898 | 0.7750 |
| BAMBI | 0.1091 | 0.0034 | 0.8430 | 0.3931 | 0.2428 | 0.1686 |
| BBS2 | 0.2474 | 1.1417 | 0.0000 | 0.2202 | 1.0006 | 1.1598 |
| BCKDK | 0.2186 | 0.2923 | 0.8654 | 1.0655 | 0.4050 | 0.1090 |
| BCL11B | 0.1982 | 0.9231 | 0.2260 | 0.2401 | 0.4151 | 0.0000 |
| BIRC5 | 1.3802 | 0.1694 | 0.3679 | 0.5452 | 0.0000 | 0.2427 |
| BOC | 0.0000 | 0.0000 | 0.3211 | 0.0000 | 1.6086 | 0.0000 |
| BTN3A1 | 0.6641 | 0.7077 | 0.0729 | 0.2544 | 0.9928 | 0.2964 |
| C1orf56 | 0.0000 | 0.8742 | 0.0000 | 0.3677 | 0.1145 | 0.3590 |
| C1QTNF6 | 0.0000 | 0.0000 | 0.5885 | 0.6205 | 0.2234 | 0.9726 |
| C2orf70 | 0.1081 | 1.0889 | 0.0206 | 0.0000 | 0.0000 | 0.0000 |
| C5orf46 | 0.0000 | 0.0000 | 0.0000 | 1.0562 | 0.1278 | 1.0438 |
| C9orf152 | 0.2087 | 1.3686 | 0.0000 | 0.3548 | 0.0206 | 0.0000 |
| CA8 | 0.0000 | 0.6859 | 0.0502 | 0.0094 | 0.0536 | 0.0000 |
| CACHD1 | 0.0000 | 0.6891 | 0.0153 | 0.0000 | 1.0768 | 0.4880 |
| CADPS2 | 0.2591 | 1.2923 | 0.0000 | 0.5506 | 1.0209 | 0.5729 |
| CAMK1G | 0.0940 | 0.2377 | 0.0000 | 0.0316 | 0.8847 | 0.0000 |
| CAPN6 | 0.0000 | 0.7541 | 0.0000 | 0.2282 | 0.6418 | 0.0000 |
| CARHSP1 | 0.7535 | 0.5316 | 0.8652 | 0.8993 | 0.2633 | 0.0000 |
| CATSPER1 | 0.1179 | 0.0000 | 0.9199 | 0.0000 | 0.0000 | 0.1046 |
| CAV1 | 0.4195 | 0.0000 | 0.1925 | 0.0801 | 0.2714 | 0.8420 |
| CCDC88A | 0.0000 | 0.1729 | 0.4668 | 0.0109 | 0.8006 | 1.0201 |
| | | | | | | |

| CCL19 | 0.0000 | 0.0000 | 0.0000 | 0.0000 | 0.9529 | 0.0000 |
|----------|--------|--------|--------|--------|--------|--------|
| CCNB1 | 1.4334 | 0.4638 | 0.1274 | 0.2506 | 0.0155 | 0.3645 |
| CCR7 | 0.0569 | 0.0000 | 0.0000 | 0.0000 | 1.0524 | 0.0000 |
| CD70 | 0.0870 | 0.0000 | 0.2096 | 0.3612 | 0.0000 | 0.4343 |
| CDA | 0.2927 | 0.0000 | 0.3408 | 0.0000 | 0.0000 | 0.6991 |
| CDC45 | 0.9608 | 0.0779 | 0.1086 | 0.3364 | 0.0336 | 0.0000 |
| CDK12 | 0.1906 | 0.2755 | 0.0000 | 0.0788 | 0.8330 | 0.0000 |
| CDK2 | 1.0635 | 0.2517 | 0.0111 | 0.5230 | 0.3310 | 0.3338 |
| CEBPB | 0.0729 | 0.0654 | 1.2909 | 0.5287 | 0.5065 | 0.8131 |
| CEP55 | 1.4198 | 0.3340 | 0.0000 | 0.1690 | 0.0000 | 0.4555 |
| CFDP1 | 0.3512 | 0.5466 | 0.7440 | 0.6706 | 0.0000 | 0.2594 |
| CHAF1B | 0.9890 | 0.2957 | 0.1997 | 0.0187 | 0.5165 | 0.0960 |
| CHEK1 | 1.5161 | 0.1621 | 0.0000 | 0.0034 | 0.1080 | 0.2731 |
| CHN2 | 0.0000 | 0.4963 | 0.0000 | 0.3389 | 0.4366 | 0.0000 |
| CIDEC | 0.0279 | 0.0000 | 0.4258 | 0.2777 | 0.0038 | 0.0000 |
| CIDECP | 0.1140 | 0.0232 | 0.5161 | 0.2795 | 0.1093 | 0.0000 |
| CKAP2L | 1.7829 | 0.2230 | 0.2724 | 0.0319 | 0.0000 | 0.0884 |
| CLEC3B | 0.0589 | 0.0691 | 0.1151 | 0.0110 | 0.8063 | 0.0000 |
| CNIH3 | 0.0000 | 0.0591 | 0.0000 | 0.3178 | 0.0000 | 0.6014 |
| CNNM1 | 0.0000 | 0.8666 | 0.4109 | 0.0000 | 0.0897 | 0.0000 |
| COL12A1 | 0.0000 | 0.1328 | 0.0340 | 0.5329 | 0.1874 | 1.6461 |
| COL5A3 | 0.0000 | 0.0000 | 0.1816 | 0.0351 | 0.0660 | 1.0286 |
| COL7A1 | 0.0000 | 0.0000 | 0.5858 | 0.0000 | 0.0000 | 0.5878 |
| COLGALT1 | 0.3987 | 0.1554 | 0.6227 | 0.4286 | 0.1646 | 0.8792 |
| COLGALT2 | 0.0000 | 0.6011 | 0.0000 | 0.0199 | 0.0000 | 0.0000 |
| COX4I2 | 0.0000 | 0.1744 | 0.0740 | 0.0000 | 0.9855 | 0.3346 |
| CSNK1D | 0.2122 | 0.3756 | 1.5627 | 0.4799 | 0.1570 | 0.2284 |
| CST6 | 0.0651 | 0.0000 | 0.2022 | 0.0000 | 0.0690 | 0.6328 |
| CTSL | 0.3897 | 0.0000 | 0.1976 | 1.1757 | 0.4702 | 0.2240 |
| CTSV | 0.3015 | 0.0439 | 0.2623 | 0.0203 | 0.0194 | 0.1819 |
| CYP2S1 | 0.3223 | 1.0232 | 0.1543 | 0.0000 | 0.0927 | 0.0000 |
| DCAF8 | 0.0000 | 1.1369 | 0.4818 | 0.1094 | 0.5277 | 0.1875 |
| DCBLD2 | 0.4024 | 0.0000 | 0.1236 | 0.0000 | 0.1426 | 0.8437 |
| DCUN1D5 | 1.3599 | 0.0751 | 0.0000 | 0.8575 | 0.9561 | 0.7193 |
| DENND1A | 0.8191 | 0.0000 | 0.2458 | 0.1898 | 0.0000 | 0.1782 |
| DERA | 1.1839 | 0.1952 | 0.4571 | 0.6042 | 0.2890 | 0.3195 |

| DHRS9 | 0.0000 | 0.0000 | 0.9957 | 0.3426 | 0.0000 | 0.1699 |
|-----------|--------|--------|--------|--------|--------|--------|
| DKK1 | 0.4779 | 0.0000 | 0.2976 | 0.1847 | 0.0000 | 0.0242 |
| DNAJC9 | 0.7779 | 0.1108 | 0.3734 | 0.1159 | 0.1329 | 0.1528 |
| DPY19L1 | 0.3414 | 0.3625 | 0.2993 | 0.5360 | 0.0781 | 0.5087 |
| DSG2 | 0.4320 | 0.5696 | 0.1794 | 0.5147 | 0.0387 | 0.7066 |
| DSG3 | 0.1766 | 0.0000 | 0.2140 | 0.0000 | 0.0000 | 0.5384 |
| DYNC2H1 | 0.0000 | 1.6131 | 0.1497 | 0.0000 | 0.7591 | 0.6693 |
| E2F7 | 1.0366 | 0.0000 | 0.0315 | 0.0222 | 0.0000 | 0.5360 |
| EDIL3 | 0.0000 | 0.0000 | 0.0000 | 0.8576 | 0.0121 | 0.8163 |
| EIF2AK3 | 0.1806 | 1.2690 | 0.0000 | 0.3842 | 0.6143 | 0.3321 |
| ELMOD3 | 0.0000 | 1.1608 | 0.6902 | 0.3859 | 0.5348 | 0.0874 |
| EMP3 | 0.2499 | 0.0000 | 0.4619 | 0.1582 | 0.2170 | 0.5646 |
| ENO2 | 0.3608 | 0.3375 | 0.7898 | 0.0339 | 0.0000 | 0.9442 |
| EPHX2 | 0.0000 | 0.5912 | 0.1080 | 0.1660 | 0.6761 | 0.0000 |
| ERRFI1 | 0.1599 | 0.0301 | 0.5475 | 0.3478 | 0.2866 | 0.7895 |
| EXOSC8 | 0.9336 | 0.6010 | 0.2789 | 1.0216 | 0.3682 | 0.1481 |
| EYA3 | 0.0000 | 0.0869 | 0.5323 | 0.0000 | 0.0000 | 0.9120 |
| FAH | 0.6763 | 0.4158 | 0.3555 | 0.2131 | 0.3240 | 0.3914 |
| FAM120AOS | 0.1803 | 1.0488 | 0.0000 | 0.2845 | 0.7143 | 0.5698 |
| FAM134B | 0.0000 | 0.8232 | 0.0000 | 0.2342 | 0.2083 | 0.0000 |
| FAM189A2 | 0.0000 | 1.0020 | 0.0000 | 0.0213 | 0.1143 | 0.0000 |
| FAM83A | 0.2461 | 0.0000 | 0.1165 | 0.0000 | 0.0000 | 0.2211 |
| FAM91A1 | 0.9811 | 0.1968 | 0.1603 | 0.7865 | 0.0000 | 0.2703 |
| FBXO22 | 0.5017 | 0.3643 | 0.0000 | 0.5761 | 0.0000 | 0.3137 |
| FBXW8 | 0.2492 | 0.2604 | 0.6553 | 0.9331 | 0.1844 | 0.3307 |
| FEM1B | 0.3031 | 0.3008 | 0.0000 | 0.0017 | 0.0838 | 1.4170 |
| FER | 0.4975 | 0.1005 | 0.1802 | 0.4440 | 0.1792 | 0.8664 |
| FGB | 0.0000 | 0.0000 | 0.0170 | 0.3212 | 0.0000 | 0.0818 |
| FGD6 | 0.5544 | 0.0000 | 0.1308 | 0.1418 | 0.0000 | 0.4991 |
| FGG | 0.0548 | 0.0379 | 0.0000 | 0.1372 | 0.0068 | 0.2157 |
| FHDC1 | 0.1771 | 1.2361 | 0.2174 | 0.0189 | 0.0000 | 0.0512 |
| FLRT3 | 0.7913 | 0.1342 | 0.5121 | 0.2846 | 0.2220 | 0.3125 |
| FRZB | 0.0889 | 0.2374 | 0.0000 | 0.5404 | 1.4969 | 0.0017 |
| FSCN1 | 0.3709 | 0.0737 | 1.0622 | 0.1342 | 0.1423 | 0.7358 |
| FST | 0.0000 | 0.0000 | 0.1578 | 0.0000 | 0.0414 | 0.4947 |
| FYN | 0.0127 | 0.5194 | 0.1203 | 0.1287 | 1.6862 | 0.8654 |

| GAB2 | 0.0435 | 0.7351 | 0.3850 | 0.6361 | 1.3628 | 0.2664 |
|-----------|--------|--------|--------|--------|--------|--------|
| GABPB1 | 0.7363 | 0.1963 | 0.0000 | 0.7422 | 0.2159 | 0.6724 |
| GAPDH | 0.4758 | 0.3945 | 0.8305 | 0.2369 | 0.0000 | 0.7231 |
| GATA6 | 0.0534 | 0.8827 | 0.0860 | 0.1396 | 0.1932 | 0.0000 |
| GATC | 1.0220 | 0.1104 | 0.0000 | 0.4818 | 0.0723 | 0.4716 |
| GIMAP2 | 0.1486 | 0.7215 | 0.0000 | 0.6567 | 0.7701 | 0.0000 |
| GINS2 | 1.0803 | 0.1777 | 0.3933 | 0.0729 | 0.0000 | 0.0000 |
| GNPAT | 0.1710 | 0.9518 | 0.1369 | 0.4352 | 0.1758 | 0.1925 |
| GOLM1 | 0.0000 | 0.7145 | 0.1203 | 0.0488 | 0.0000 | 0.0000 |
| GPC3 | 0.0980 | 0.2322 | 0.0000 | 0.0000 | 1.2713 | 0.0000 |
| GPR176 | 0.4324 | 0.3072 | 0.0000 | 0.7415 | 0.3745 | 0.5882 |
| HIPK2 | 0.2587 | 1.2502 | 0.0694 | 0.2371 | 0.5213 | 0.0000 |
| HJURP | 1.3269 | 0.2436 | 0.2326 | 0.0210 | 0.0000 | 0.0000 |
| HRASLS2 | 0.3273 | 0.0000 | 0.3045 | 0.2167 | 0.0000 | 0.0000 |
| HSP90B1 | 0.5274 | 0.4642 | 0.7758 | 0.8972 | 0.2977 | 0.3795 |
| HSPB6 | 0.0000 | 0.1493 | 0.1298 | 0.0000 | 1.3081 | 0.3131 |
| ICAM2 | 0.5013 | 0.1959 | 0.4755 | 0.3105 | 0.4043 | 0.1342 |
| IDH2 | 0.7131 | 0.4322 | 0.3970 | 0.2145 | 0.3314 | 0.2342 |
| IFT140 | 0.0000 | 1.0890 | 0.5193 | 0.0000 | 0.2592 | 0.0662 |
| IGFBP1 | 0.2708 | 0.0000 | 0.2323 | 0.0327 | 0.0000 | 0.0058 |
| IGLL3P | 0.1660 | 0.1496 | 0.0000 | 0.0000 | 0.7633 | 0.0000 |
| IKBIP | 0.2893 | 0.0000 | 0.3028 | 1.1219 | 0.1455 | 0.4694 |
| IL1R2 | 0.0377 | 0.2543 | 0.4285 | 0.2301 | 0.0000 | 0.0605 |
| IL20RB | 0.2578 | 0.0000 | 0.3094 | 0.0000 | 0.0000 | 0.6805 |
| IL33 | 0.2369 | 0.0436 | 0.0000 | 0.1304 | 0.6759 | 0.0000 |
| ITGA5 | 0.0000 | 0.0000 | 0.4758 | 0.2666 | 0.1206 | 0.6815 |
| ITPKB | 0.0000 | 0.8315 | 0.6059 | 0.0000 | 1.1923 | 0.6724 |
| KANK4 | 0.0000 | 0.0000 | 0.1981 | 0.4683 | 0.0000 | 1.2292 |
| KCNQ3 | 0.0000 | 0.1296 | 0.1721 | 0.7768 | 0.0916 | 0.5160 |
| KCTD10 | 0.3776 | 0.1324 | 0.2867 | 0.4387 | 0.5081 | 0.7943 |
| KCTD5 | 0.3848 | 0.5133 | 1.1253 | 0.6056 | 0.0000 | 0.0000 |
| KIAA0513 | 0.0828 | 1.0351 | 0.1715 | 0.3220 | 0.5910 | 0.0000 |
| KIAA1549L | 0.3755 | 0.0812 | 0.2646 | 0.6647 | 0.1501 | 0.6423 |
| KIF14 | 1.1244 | 0.3648 | 0.1952 | 0.4293 | 0.0000 | 0.1264 |
| KIF20A | 1.3726 | 0.2864 | 0.2082 | 0.2320 | 0.0000 | 0.2888 |
| KIF2C | 0.7952 | 0.1329 | 0.1096 | 0.0074 | 0.0000 | 0.0000 |

| KLHL5 | 0.4215 | 0.1645 | 0.0000 | 0.3538 | 0.6955 | 1.1410 |
|--------------|--------|--------|--------|--------|--------|--------|
| KNTC1 | 1.0718 | 0.1383 | 0.4419 | 0.0827 | 0.1499 | 0.2787 |
| KRT17 | 0.2860 | 0.0000 | 0.3863 | 0.1586 | 0.1201 | 0.5074 |
| KRT6A | 0.1386 | 0.0000 | 0.1202 | 0.0000 | 0.0000 | 0.4668 |
| KRT6C | 0.1187 | 0.0000 | 0.0000 | 0.0000 | 0.0000 | 0.1640 |
| KRT7 | 0.4597 | 0.0020 | 0.5620 | 0.0000 | 0.1354 | 0.4370 |
| KYNU | 0.6104 | 0.0894 | 0.0693 | 0.5431 | 0.0000 | 0.2790 |
| LAMA5 | 0.3670 | 0.0772 | 1.0234 | 0.0000 | 0.3418 | 0.1832 |
| LCNL1 | 0.1072 | 0.2829 | 0.0115 | 0.2669 | 0.5289 | 0.0000 |
| LDHA | 0.6526 | 0.4664 | 0.0000 | 0.3186 | 0.0504 | 1.1696 |
| LETM2 | 0.4402 | 0.0000 | 0.3924 | 0.0000 | 0.0000 | 0.2831 |
| LGALS9B | 0.1106 | 1.0239 | 0.0000 | 0.0000 | 0.3463 | 0.4913 |
| LINC01184 | 0.6331 | 0.8045 | 0.0000 | 0.3418 | 0.8076 | 0.0000 |
| LMO3 | 0.0000 | 0.1062 | 0.0000 | 0.0090 | 1.1796 | 0.0136 |
| LMTK2 | 0.7364 | 0.3642 | 0.3100 | 0.5254 | 0.0204 | 0.2425 |
| LOC100506562 | 0.5772 | 0.2935 | 0.6002 | 0.6045 | 0.1075 | 0.1108 |
| LOX | 0.2078 | 0.0000 | 0.0806 | 0.3896 | 0.0866 | 0.9212 |
| LYNX1 | 0.0337 | 0.0000 | 0.2575 | 0.1651 | 0.0000 | 0.0951 |
| MAP3K8 | 0.1984 | 0.0000 | 0.0681 | 0.3075 | 0.5588 | 0.4348 |
| MARCKSL1 | 0.1504 | 1.3374 | 0.2978 | 0.0000 | 0.0000 | 0.2627 |
| MARS2 | 0.7481 | 1.0181 | 0.0000 | 0.4007 | 0.4981 | 0.0000 |
| MC1R | 0.1042 | 0.1313 | 1.0794 | 0.8656 | 0.4740 | 0.1335 |
| MCEMP1 | 0.0000 | 0.0000 | 0.0000 | 0.6056 | 0.0000 | 0.2992 |
| MCM10 | 1.1446 | 0.1414 | 0.0000 | 0.0141 | 0.0000 | 0.0808 |
| MCM4 | 1.2790 | 0.1411 | 0.3090 | 0.0254 | 0.0103 | 0.1276 |
| MCOLN2 | 0.1988 | 0.2778 | 0.0000 | 0.0000 | 0.9442 | 0.0000 |
| MELK | 1.0177 | 0.2864 | 0.0000 | 0.2322 | 0.0133 | 0.2208 |
| MEOX1 | 0.0000 | 0.0536 | 0.1642 | 0.0438 | 0.9639 | 0.0000 |
| MIF | 0.4348 | 0.3316 | 0.9576 | 0.4402 | 0.0008 | 0.6845 |
| MIR99AHG | 0.0371 | 0.2791 | 0.3859 | 0.4466 | 1.7947 | 0.2232 |
| MME | 0.0009 | 0.0000 | 0.0640 | 0.4532 | 0.0419 | 0.5791 |
| MRAP2 | 0.0430 | 0.7825 | 0.0000 | 0.2177 | 0.2314 | 0.0000 |
| MRPL24 | 0.1643 | 1.1324 | 0.2156 | 0.1207 | 0.2213 | 0.1778 |
| MTRNR2L1 | 0.2795 | 0.5589 | 0.4897 | 0.0719 | 0.5523 | 0.0000 |
| NACC2 | 0.5312 | 0.0000 | 0.7176 | 0.2474 | 0.0000 | 0.1055 |
| NAMPT | 0.3355 | 0.0000 | 0.0493 | 0.7543 | 0.3154 | 0.3500 |

| NCAPD2 | 1.3843 | 0.4110 | 0.1605 | 0.1233 | 0.2041 | 0.3231 |
|------------------|--------|--------|--------|--------|--------|--------|
| NCAPG | 1.6056 | 0.4449 | 0.0000 | 0.0000 | 0.0000 | 0.5243 |
| NELFE | 0.9382 | 0.2255 | 0.5894 | 0.8561 | 0.3602 | 0.0798 |
| NEURL2 | 0.6888 | 0.1217 | 0.0000 | 0.2556 | 0.7216 | 0.4336 |
| NFIA | 0.1194 | 0.8389 | 0.0000 | 0.3854 | 1.5045 | 0.2708 |
| NFIX | 0.0000 | 0.8819 | 0.1383 | 0.0000 | 1.3919 | 0.7968 |
| \overline{NMB} | 0.2126 | 0.1909 | 0.6634 | 0.7944 | 0.0000 | 0.3640 |
| NPM1 | 0.0000 | 1.0465 | 0.0000 | 0.0029 | 0.0826 | 0.0446 |
| NR0B2 | 0.0000 | 0.8362 | 0.0000 | 0.0000 | 0.1422 | 0.0000 |
| NRP2 | 0.1462 | 0.0000 | 0.4996 | 0.0000 | 0.0000 | 0.0534 |
| NUP155 | 1.1296 | 0.4140 | 0.0620 | 0.3285 | 0.2288 | 0.4554 |
| OAZ1 | 0.8583 | 0.5931 | 0.6573 | 1.1219 | 0.5151 | 0.5871 |
| ORC1 | 0.9777 | 0.3231 | 0.1638 | 0.9547 | 0.1157 | 0.0101 |
| P2RY2 | 0.1789 | 0.0331 | 0.7738 | 0.2163 | 0.0000 | 0.5005 |
| P2RY8 | 0.2334 | 0.0728 | 0.0000 | 0.2788 | 1.6555 | 0.0000 |
| P4HA1 | 0.0430 | 0.1009 | 0.4121 | 0.8384 | 0.0000 | 0.5460 |
| P4HA2 | 0.3225 | 0.1659 | 0.1245 | 0.5449 | 0.1088 | 0.7371 |
| PAX8 | 0.7680 | 0.0000 | 0.5631 | 0.0000 | 0.0000 | 0.0000 |
| PAX8-AS1 | 0.5656 | 0.0447 | 0.3435 | 0.0750 | 0.0071 | 0.0000 |
| PBXIP1 | 0.0000 | 0.5144 | 0.4130 | 0.0000 | 0.4392 | 0.1667 |
| PCDH20 | 0.0000 | 0.4318 | 0.0000 | 0.1465 | 0.0000 | 0.0000 |
| PCF11 | 0.2613 | 0.9351 | 0.2527 | 0.0950 | 1.1086 | 0.4077 |
| PCOLCE2 | 0.0000 | 0.0076 | 0.1188 | 0.5379 | 0.0000 | 0.0542 |
| PDLIM7 | 0.1954 | 0.0000 | 0.4086 | 0.3731 | 0.1144 | 0.6779 |
| PEX11B | 0.1066 | 1.3518 | 0.0000 | 0.5264 | 0.2883 | 0.2455 |
| PFKFB4 | 0.5485 | 0.2199 | 0.6769 | 0.4272 | 0.1428 | 0.2854 |
| PGAM5 | 0.9213 | 0.0000 | 0.3859 | 0.4866 | 0.0000 | 0.0000 |
| PGBD3 | 0.6174 | 0.3626 | 0.4335 | 0.2008 | 0.5630 | 0.7384 |
| PHACTR3 | 0.1489 | 0.0000 | 0.3225 | 0.1416 | 0.0026 | 0.0728 |
| PHLDA1 | 0.0838 | 0.1387 | 0.7170 | 0.1250 | 0.6249 | 1.5017 |
| PHOSPHO2 | 0.3445 | 1.0681 | 0.0000 | 0.4652 | 0.4054 | 0.0514 |
| PIGL | 1.0637 | 0.1481 | 0.5587 | 0.3049 | 0.2423 | 0.0000 |
| PLAC9 | 0.0707 | 0.0000 | 0.0000 | 0.1090 | 1.2901 | 0.0766 |
| PLAU | 0.2139 | 0.0000 | 0.2764 | 0.0000 | 0.0249 | 0.8793 |
| PLEKHS1 | 0.0000 | 0.6411 | 0.3407 | 0.0862 | 0.2791 | 0.0176 |
| PLIN2 | 0.3057 | 0.0000 | 0.0818 | 1.0167 | 0.4683 | 0.2095 |

| PLIN3 | 0.3365 | 0.2607 | 0.9673 | 0.9320 | 0.1395 | 0.4103 |
|----------|--------|--------|--------|--------|--------|--------|
| PLOD1 | 0.0595 | 0.0000 | 1.2074 | 0.7504 | 0.3668 | 0.8026 |
| PLOD2 | 0.1489 | 0.0922 | 0.2366 | 0.2919 | 0.1729 | 0.8899 |
| POC1A | 1.3753 | 0.3309 | 0.3179 | 0.4709 | 0.0000 | 0.0000 |
| POLA2 | 0.8413 | 0.2234 | 0.3296 | 0.1331 | 0.2137 | 0.0000 |
| POP5 | 0.5635 | 0.5070 | 1.5160 | 0.2263 | 0.1092 | 0.1799 |
| POU2AF1 | 0.0611 | 0.4732 | 0.0000 | 0.0007 | 0.9240 | 0.0000 |
| PP7080 | 0.1047 | 0.9680 | 0.0000 | 0.0371 | 0.0000 | 0.0000 |
| PPAPDC1A | 0.0000 | 0.0000 | 0.0000 | 0.7582 | 0.0000 | 1.2230 |
| PPM1H | 0.0000 | 0.8512 | 0.4600 | 0.2700 | 0.2363 | 0.0000 |
| PPP1R12B | 0.1652 | 0.3193 | 0.7825 | 0.6308 | 0.0253 | 0.4910 |
| PPP1R14B | 0.3673 | 0.2586 | 0.7846 | 0.0000 | 0.3651 | 0.5928 |
| PPP1R3C | 0.0000 | 0.0160 | 0.1325 | 0.3710 | 0.0256 | 0.2554 |
| PPY | 0.0000 | 0.4957 | 0.0000 | 0.0805 | 1.0771 | 0.0000 |
| PRC1 | 0.9560 | 0.3521 | 0.0407 | 0.0375 | 0.0000 | 0.3200 |
| PRDM16 | 0.0000 | 1.1224 | 0.0000 | 0.0000 | 0.5289 | 0.0867 |
| PREP | 0.0587 | 0.9830 | 0.3047 | 0.1977 | 0.0203 | 0.0000 |
| PRKCDBP | 0.2571 | 0.0000 | 1.0161 | 0.5090 | 0.2613 | 0.5936 |
| PRMT7 | 0.1393 | 1.5003 | 0.4373 | 0.0000 | 0.1793 | 0.2230 |
| PROSER2 | 0.9335 | 0.1760 | 0.4026 | 0.3736 | 0.2680 | 0.3965 |
| PRR11 | 0.8207 | 0.0503 | 0.2272 | 0.0000 | 0.0000 | 0.0934 |
| PTGES | 0.5703 | 0.0160 | 0.5702 | 0.0681 | 0.0000 | 0.5634 |
| PTPN21 | 0.2722 | 0.1714 | 0.3219 | 0.4864 | 0.2674 | 0.8423 |
| PXDN | 0.0000 | 0.0000 | 0.3795 | 0.5917 | 0.3108 | 1.1884 |
| PYGL | 0.0808 | 0.0000 | 0.3079 | 0.3384 | 0.1413 | 0.7445 |
| RAB31 | 0.1110 | 0.0000 | 0.2586 | 0.8745 | 0.7552 | 1.1882 |
| RACGAP1 | 1.3720 | 0.3729 | 0.1382 | 0.1936 | 0.0734 | 0.3348 |
| RALGAPB | 0.9974 | 0.5032 | 0.2879 | 0.7587 | 0.2585 | 0.7977 |
| RAP1GAP | 0.0000 | 1.0067 | 0.4657 | 0.2773 | 0.7542 | 0.0000 |
| RASL11B | 0.0000 | 0.1852 | 0.0682 | 0.2236 | 1.2121 | 0.3095 |
| RAVER2 | 0.1985 | 0.9070 | 0.0534 | 0.0890 | 0.2667 | 0.0577 |
| RBMS2 | 0.6118 | 0.1541 | 0.0000 | 0.4022 | 0.3184 | 0.8946 |
| RERE | 0.0485 | 0.7372 | 0.6212 | 0.0026 | 0.9874 | 0.4207 |
| RERGL | 0.2378 | 0.0000 | 0.0000 | 0.1054 | 1.1842 | 0.0000 |
| RFC5 | 1.0809 | 0.2444 | 0.0000 | 0.5248 | 0.1556 | 0.3147 |
| RFK | 0.0000 | 0.6594 | 0.1169 | 0.0000 | 0.4342 | 0.2100 |
| | | | | | | |

| RFX2 | 0.0000 | 0.2219 | 0.2372 | 0.0000 | 0.4551 | 0.2959 |
|----------|--------|--------|--------|--------|--------|--------|
| RGS3 | 0.2370 | 0.1243 | 0.0000 | 0.8096 | 0.2269 | 0.3212 |
| RGS5 | 0.0000 | 0.4317 | 0.0455 | 0.0788 | 0.5794 | 0.0934 |
| RHOF | 0.7466 | 0.1749 | 0.4760 | 0.1428 | 0.0000 | 0.5878 |
| RMND5A | 0.2696 | 0.1188 | 0.2601 | 0.7065 | 0.0000 | 0.0750 |
| RNF103 | 0.0344 | 1.2504 | 0.1672 | 0.5545 | 0.2894 | 0.0635 |
| RPA2 | 0.4727 | 0.6964 | 0.7005 | 0.4129 | 1.4239 | 0.2443 |
| RPIA | 0.4609 | 1.3515 | 0.2200 | 0.1918 | 0.4584 | 0.0000 |
| SAMD5 | 0.1340 | 0.5397 | 0.0000 | 0.0000 | 0.0860 | 0.0000 |
| SCGB2A1 | 0.0000 | 0.8288 | 0.0000 | 0.1826 | 0.1547 | 0.0000 |
| SCYL2 | 0.7048 | 0.3901 | 0.0000 | 0.9782 | 0.4060 | 0.9614 |
| SDIM1 | 0.0000 | 0.0455 | 0.2422 | 0.0000 | 0.5017 | 0.0000 |
| SEC23IP | 0.3380 | 1.2955 | 0.0000 | 0.5310 | 0.3578 | 0.4605 |
| SELENBP1 | 0.0000 | 1.2032 | 0.3621 | 0.2011 | 0.2603 | 0.0000 |
| SEPW1 | 0.0349 | 0.9518 | 1.2360 | 0.0000 | 0.6293 | 0.5568 |
| SERPINB3 | 0.0000 | 0.0000 | 0.1755 | 0.1787 | 0.0000 | 0.0506 |
| SERPINH1 | 0.0000 | 0.0115 | 0.3898 | 0.2169 | 0.4300 | 1.0203 |
| SERTAD2 | 0.2931 | 0.1441 | 0.8991 | 0.9858 | 0.4859 | 0.4437 |
| SGSM1 | 0.0000 | 0.9290 | 0.0817 | 0.0211 | 0.8410 | 0.0000 |
| SH3GL1 | 0.1173 | 0.1075 | 1.0090 | 1.2494 | 0.2155 | 0.0000 |
| SLAMF9 | 0.0435 | 0.0000 | 0.0000 | 0.6663 | 0.0000 | 0.0657 |
| SLC12A2 | 0.0380 | 0.9089 | 0.3449 | 0.0968 | 0.4855 | 0.1821 |
| SLC15A1 | 0.0000 | 0.0000 | 0.4779 | 0.0000 | 0.0569 | 0.0565 |
| SLC16A3 | 0.1282 | 0.3828 | 1.1047 | 0.4222 | 0.0000 | 0.9957 |
| SLC2A1 | 0.1786 | 0.1209 | 0.9980 | 0.4099 | 0.0000 | 0.7045 |
| SLC2A3 | 0.0000 | 0.0000 | 0.3369 | 0.7592 | 0.3268 | 0.7204 |
| SLC30A3 | 0.4502 | 0.5017 | 0.0822 | 0.2136 | 0.6568 | 0.0654 |
| SLC40A1 | 0.0000 | 0.8927 | 0.0000 | 0.5789 | 0.2440 | 0.1550 |
| SMOX | 0.3692 | 0.2900 | 1.4313 | 0.9987 | 0.1840 | 0.0000 |
| SNORA11D | 0.0849 | 0.2729 | 0.4795 | 0.4375 | 0.0039 | 0.2687 |
| SNRPB | 0.9900 | 0.0786 | 0.4143 | 0.9037 | 0.0238 | 0.0000 |
| SOBP | 0.0000 | 0.1979 | 0.8103 | 0.1044 | 1.3581 | 0.0039 |
| SOD2 | 0.5780 | 0.1207 | 0.0000 | 0.4656 | 0.4023 | 0.1652 |
| SPHK1 | 0.2590 | 0.0000 | 0.2748 | 0.0907 | 0.6221 | 1.4095 |
| SPIN4 | 0.8495 | 0.3236 | 0.7960 | 0.3855 | 0.2224 | 0.3985 |
| SPOCD1 | 0.0000 | 0.0000 | 0.1782 | 0.2094 | 0.0000 | 0.7594 |
| | | | | | | |

| SPOCK1 | 0.1196 | 0.0000 | 0.0293 | 0.5189 | 0.3390 | 1.2727 |
|------------|--------|--------|--------|--------|--------|--------|
| SPP1 | 0.0294 | 0.0805 | 0.0000 | 1.0413 | 0.3073 | 0.7357 |
| ST3GAL2 | 0.3414 | 0.0000 | 0.8015 | 1.0746 | 0.4432 | 0.0000 |
| ST6GAL1 | 0.1717 | 0.8423 | 0.0000 | 0.2289 | 0.6651 | 0.0916 |
| ST6GALNAC1 | 0.0396 | 0.9957 | 0.0803 | 0.1154 | 0.0000 | 0.1050 |
| STAT5B | 0.0000 | 0.9053 | 0.3202 | 0.0618 | 1.3050 | 0.2213 |
| STK39 | 0.1526 | 0.9966 | 0.2351 | 0.1373 | 0.0838 | 0.1226 |
| SUGCT | 0.0000 | 0.0321 | 0.0000 | 0.6297 | 0.1256 | 0.9331 |
| SULF2 | 0.1725 | 0.1513 | 0.4552 | 0.1878 | 0.3858 | 0.7665 |
| SYNE2 | 0.0000 | 0.8824 | 0.2432 | 0.0000 | 0.2767 | 0.2763 |
| TAF5L | 0.2232 | 1.0626 | 0.1753 | 0.2440 | 0.2327 | 0.2249 |
| TARBP2 | 0.6779 | 0.3829 | 1.2178 | 0.6116 | 0.1843 | 0.0000 |
| TCEA3 | 0.0000 | 0.8898 | 0.2645 | 0.0922 | 0.6204 | 0.0000 |
| TCTA | 0.0000 | 0.7508 | 0.8167 | 0.0875 | 0.9836 | 0.0178 |
| TGFBI | 0.1874 | 0.0000 | 0.1522 | 0.1879 | 0.0548 | 0.9986 |
| THSD7B | 0.0859 | 0.2031 | 0.0000 | 0.2900 | 0.9574 | 0.1114 |
| TLE4 | 0.0509 | 0.8787 | 0.0746 | 0.3315 | 0.8984 | 0.4660 |
| TM9SF3 | 0.0000 | 1.0785 | 0.2190 | 0.0000 | 0.1641 | 0.2114 |
| TMED1 | 0.2561 | 0.3378 | 1.1457 | 0.8311 | 0.4929 | 0.2755 |
| TMEM26 | 0.0407 | 0.0237 | 0.1028 | 0.4886 | 0.2223 | 1.4490 |
| TMTC4 | 0.0000 | 1.2865 | 0.3348 | 0.2090 | 0.1995 | 0.2756 |
| TNFRSF10D | 0.1474 | 0.1117 | 0.6603 | 0.4579 | 0.0000 | 0.1751 |
| TNFRSF17 | 0.0258 | 0.0455 | 0.0000 | 0.0803 | 0.5772 | 0.0000 |
| TNFRSF6B | 0.6268 | 0.0000 | 0.0684 | 0.1841 | 0.0000 | 0.3940 |
| TOM1 | 0.0000 | 0.1032 | 1.4892 | 0.8140 | 0.6813 | 0.5236 |
| TOM1L2 | 0.1892 | 0.0000 | 0.6276 | 0.3305 | 0.0489 | 0.2346 |
| TOR2A | 0.0000 | 0.9859 | 0.4755 | 0.2012 | 0.5273 | 0.0000 |
| TPD52L2 | 0.6311 | 0.1617 | 1.3107 | 0.6501 | 0.4351 | 0.2322 |
| TPX2 | 1.3192 | 0.1540 | 0.0351 | 0.1488 | 0.0392 | 0.1087 |
| TRAPPC2 | 0.5080 | 1.0792 | 0.0000 | 0.4917 | 0.6155 | 0.1418 |
| TREM1 | 0.0472 | 0.0000 | 0.0870 | 0.7055 | 0.0000 | 0.3006 |
| TRERF1 | 0.4920 | 0.2861 | 0.3810 | 0.1345 | 0.0517 | 0.1346 |
| TRIM2 | 0.1310 | 1.1544 | 0.3127 | 0.3092 | 0.3595 | 0.0000 |
| TSTD1 | 0.1685 | 1.2229 | 0.4834 | 0.0685 | 0.4502 | 0.0191 |
| TUBA1C | 1.3100 | 0.5454 | 0.5360 | 0.5305 | 0.2711 | 0.5032 |
| TWIST1 | 0.0000 | 0.0000 | 0.1970 | 0.9070 | 0.1202 | 1.2015 |
| | | | | | | |

| UFC1 | 0.0000 | 1.1861 | 0.2466 | 0.4651 | 0.2997 | 0.0000 |
|---------|--------|--------|--------|--------|--------|--------|
| UHRF2 | 0.1520 | 0.2931 | 0.3251 | 0.4968 | 0.6565 | 1.1025 |
| UPP1 | 0.5505 | 0.0000 | 0.7864 | 0.4294 | 0.1567 | 0.1100 |
| USP30 | 0.5449 | 0.1353 | 0.3862 | 0.0000 | 0.0771 | 0.0000 |
| VPS35 | 0.3941 | 1.3902 | 0.0000 | 0.5311 | 0.0000 | 0.2457 |
| VSTM2L | 0.3176 | 0.0000 | 0.9398 | 0.0000 | 0.0509 | 0.0656 |
| WNT2B | 0.0885 | 0.1107 | 0.0000 | 0.0139 | 0.4530 | 0.0000 |
| XXYLT1 | 0.2408 | 0.0000 | 1.0488 | 1.0782 | 0.4595 | 0.8654 |
| ZBED2 | 0.1569 | 0.0000 | 0.1800 | 0.0000 | 0.0000 | 0.6435 |
| ZFPM1 | 0.0000 | 1.2172 | 0.2917 | 0.0000 | 0.4340 | 0.1504 |
| ZNF185 | 0.2542 | 0.1747 | 1.0210 | 0.4834 | 0.0000 | 0.7221 |
| ZNF565 | 0.0701 | 0.2851 | 0.0717 | 0.0569 | 0.2393 | 0.0768 |
| ZNF658 | 0.0000 | 0.8769 | 0.0000 | 0.0000 | 0.9099 | 0.2753 |
| ZPLD1 | 0.0000 | 0.0000 | 0.1873 | 0.0325 | 0.0294 | 0.1074 |
| ZSCAN16 | 0.3012 | 1.4502 | 0.0000 | 0.0175 | 0.5146 | 0.5090 |
| ZSCAN32 | 0.3467 | 1.1558 | 0.4982 | 0.3027 | 0.7286 | 0.2378 |

Appendix B

MSigDB signatures correlated with axis A1

Table B.1: MSigDB signatures substantially correlated with activity of the prognostic axis A1.

MSigDB set

- $c5.M.PHASE/c5.MITOSIS/c5.M.PHASE_OF_MITOTIC_CELL_CYCLE$
- c5.REGULATION_OF_MITOSIS
- c5.CELL_CYCLE_PROCESS/c5.MITOTIC_CELL_CYCLE/c5.CELL_CYCLE_PHASE
- c5.SPINDLE
- c4.MORF_BUB1B
- c6.CSR_LATE_UP.V1_SIGNED
- c5.SPINDLE_POLE
- c2.PID_PLK1_PATHWAY
- c5.ORGANELLE_PART/c5.INTRACELLULAR_ORGANELLE_PART
- c2.REACTOME_CELL_CYCLE/c2.REACTOME_CELL_CYCLE_MITOTIC
- c2.REACTOME_CYCLIN_A_B1_ASSOCIATED_EVENTS_DURING_G2_M_TRANSITION
- c2.REACTOME_MITOTIC_PROMETAPHASE
- c2.KEGG_CELL_CYCLE
- c5.CHROMOSOME_SEGREGATION
- c4.MORF_FEN1
- $c2.REACTOME_G1_S_SPECIFIC_TRANSCRIPTION$
- ${\tt c2.REACTOME_ACTIVATION_OF_THE_PRE_REPLICATIVE_COMPLEX/c2.REACTOME_ACTIVE_CATIVE_COMPLEX/c2.REACTO$
- c2.REACTOME_E2F_ENABLED_INHIBITION_OF_PRE_REPLICATION_COMPLEX_FORMATIO
- ${\tt c2.REACTOME_E2F_MEDIATED_REGULATION_OF_DNA_REPLICATION}$
- c5.CELL_CYCLE_GO_0007049
- c2.REACTOME_KINESINS
- c3.V\$ELK1_02
- $c5.SPINDLE_MICROTUBULE$
- ${\tt c5.MITOTIC_CELL_CYCLE_CHECKPOINT}$
- c2.REACTOME_CELL_CYCLE_CHECKPOINTS/c2.REACTOME_G1_S_TRANSITION/c2.REACT
- c4.MORF_ESPL1
- c4.MORF_BUB1
- c4.MORF_BUB3/c4.MORF_RAD23A
- c5.CONDENSED_CHROMOSOME
- c4.MORF_RFC4/c4.MORF_RRM1
- c2.BIOCARTA_G2_PATHWAY
- c3.SCGGAAGY_V\$ELK1_02
- c2.PID_AURORA_A_PATHWAY
- c5.MITOTIC_SISTER_CHROMATID_SEGREGATION/c5.SISTER_CHROMATID_SEGREGATION
- c4.MORF_UNG
- c2.PID_FOXM1PATHWAY
- c4.MORF_GSPT1
- c2.REACTOME_METABOLISM_OF_NUCLEOTIDES

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- c2.PID_ATR_PATHWAY
- c2.BIOCARTA_MCM_PATHWAY
- c4.MORF_CCNF
- $c5.CELL_CYCLE_CHECKPOINT_GO_0000075$
- $c5. MITOTIC_SPINDLE_ORGANIZATION_AND_BIOGENESIS/c5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/c5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENESIS/C5. SPINDLE_ORGANIZATION_AND_BIOGENE$
- c4.MORF_EI24
- c5.DOUBLE_STRAND_BREAK_REPAIR
- c4.GNF2_PA2G4/c4.GNF2_RAN
- c2.REACTOME_G2_M_DNA_DAMAGE_CHECKPOINT
- c2 KEGG PVRIMIDINE METAROLISM

Appendix C

MSigDB signatures correlated with axis A2

Table C.1: MSigDB signatures substantially correlated with activity of the prognostic axis A2.

GeneSet

- c2.PID_INTEGRIN1_PATHWAY
- c2.PID_INTEGRIN3_PATHWAY
- c2.PID_UPA_UPAR_PATHWAY
- c4.GNF2_PTX3
- c2.KEGG_ECM_RECEPTOR_INTERACTION
- c2.PID_INTEGRIN5_PATHWAY
- c4.GNF2_MMP1
- $c2.REACTOME_EXTRACELLULAR_MATRIX_ORGANIZATION/c2.REACTOME_COLLAGEN_FOR COLLAGEN_FOR COLLAGEN_F$
- c5.AXON_GUIDANCE
- c2.KEGG_FOCAL_ADHESION
- c2.PID_SYNDECAN_1_PATHWAY
- ${\tt c2.REACTOME_CELL_EXTRACELLULAR_MATRIX_INTERACTIONS}$
- c2.PID_INTEGRIN_CS_PATHWAY
- c5.TISSUE_DEVELOPMENT
- c5.COLLAGEN
- c6.CORDENONSI_YAP_CONSERVED_SIGNATURE
- c6.LEF1_UP.V1_SIGNED
- c2.REACTOME_INTEGRIN_CELL_SURFACE_INTERACTIONS
- c5.AXONOGENESIS/c5.CELLULAR_MORPHOGENESIS_DURING_DIFFERENTIATION
- c6.STK33_NOMO_SIGNED
- ${\rm c7.GSE17721_CTRL_VS_CPG_12H_BMDM_SIGNED}$
- $c7.GSE1460_INTRATHYMIC_T_PROGENITOR_VS_THYMIC_STROMAL_CELL_SIGNED$

Appendix D

Approximate calculation of PARSE scores

Exact calculation of PARSE score requires the solution of a number of NNLS problems, which complicates application. The NNLS solutions can be approximated with conventional least squares solutions, ultimately transforming the calculation of an approximate PARSE score into a simple weighted sum of gene expression measurements.

Recall that NMF finds factorizations of the form A=WH, with all elements of A, W, and H, being non-negative. In the reverse problem of PARSE calculation, A and \widehat{W} are supplied, and H is to be estimated. I propose an approximation that removes the requirement that H be non-negative, $H \approx \widehat{W}^+A$, where \widehat{W}^+ is the Moore-Penrose pseudoinverse of \widehat{W} . By combining this approximation with the linear combination of metagene coefficients that forms the PARSE score, we can approximate PARSE as a simple weighted sum of gene expression measurements:

$$P = LH \tag{D.1}$$

$$\approx L\widehat{W}^+ A$$
 (D.2)

$$= kA \tag{D.3}$$

where P is the vector of PARSE score values, L is the metagene loadings for the PARSE score, $L = (1.354 - 1.548 \ 0 \ 0 - 1.354 \ 1.548)$, and k is a row vector of gene loadings for calculation of an approximate PARSE score. Approximation of P by kA appears excellent; when tested on APGI gene expression measurements,

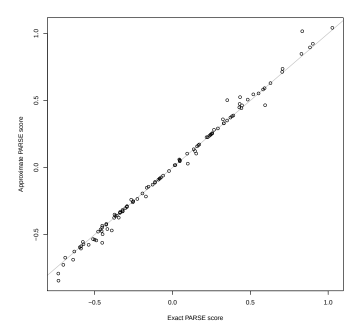


Figure D.1: The linear PARSE score approximation $P \approx kA$ closely matches the exact version calculated using NNLS, when evaluated on APGI GEX data.

the approximation closely matched the more laborious exact NNLS solution (Figure D.1).

To use the approximation in practice, perform the following steps:

- 1. Prepare a gene \times sample matrix of linear expression estimates A, in which values for each row (gene) have been scaled to encompass the range 0 to 1.
- 2. Subset A to only the genes present in the k table (below), and arrange rows of A so that they exactly match the order of rows of k. If genes present in k are missing from A, insert all-zero rows for these genes into A.
- 3. Calculate approximate PARSE scores P as P = kA. This is equivalent to, for each column (sample) of A, multiplying each entry of the column of A with the corresponding entry of k, and summing the results.

The loading vector for the calculation of approximate PARSE score, k^T , follows.

| | Value |
|----------|----------|
| A4GALT | 0.00418 |
| A4GNT | -0.01632 |
| ABHD16A | 0.00143 |
| ABHD5 | 0.01227 |
| ABLIM1 | -0.01392 |
| ACE | -0.00556 |
| ACKR3 | 0.00802 |
| ACYP2 | -0.01298 |
| ADH1A | -0.01845 |
| ADM | 0.00122 |
| AGRP | -0.00509 |
| AKIP1 | 0.00545 |
| AKR1A1 | -0.01321 |
| ALDH5A1 | -0.02452 |
| ALOX5AP | -0.00179 |
| AMOT | -0.00825 |
| ANGPTL2 | 0.01178 |
| ANGPTL4 | 0.01365 |
| ANKLE2 | 0.01205 |
| ANKRD22 | -0.00941 |
| ANKRD37 | 0.00474 |
| ANLN | 0.04364 |
| APCDD1 | 0.01244 |
| APCS | 0.00602 |
| ARFGAP3 | -0.01070 |
| ARHGAP24 | -0.02524 |
| ARHGEF19 | -0.00476 |
| ARL4C | 0.02609 |
| ARSD | -0.01466 |
| ASPM | 0.01593 |
| ATAD2 | 0.02602 |
| ATF7IP2 | -0.00405 |
| ATL3 | 0.00972 |
| AURKB | 0.01869 |

- AXIN2 -0.01658
- B3GALTL 0.01113
 - BAMBI -0.00680
 - BBS2 0.00587
 - BCKDK -0.02452
 - BCL11B -0.02161
 - BIRC5 0.02419
 - BOC -0.03047
 - BTN3A1 -0.00868
 - Clorf56 -0.00865
- C1QTNF6 0.01572
 - _____
 - C2orf70 -0.01360
 - C5orf46 0.01559
 - C9 or f152 -0.02152
 - CA8 -0.01129
- $CACHD1 \quad \text{-}0.01313$
- CADPS2 -0.02136
- CAMK1G -0.01790
- CAPN6 -0.02615
- CARHSP1 -0.01515
- CATSPER1 0.00163
 - CAV1 0.02989
- CCDC88A 0.01480
 - CCL19 -0.01715
 - $\begin{array}{cc} CCNB1 & 0.03071 \end{array}$
 - CCR7 -0.01775
 - $CD70 \quad 0.00954$
 - CDA = 0.02792
 - CDC45 0.01256
 - CDK12 -0.01624
 - CDK2 0.01546
 - CEBPB 0.00404
 - CEP55 0.03755
 - CFDP1 -0.00617
 - CHAF1B 0.00920
 - CHEK1 0.03669

- CHN2 -0.02051
 CIDEC -0.00596
 CIDECP -0.00684
 - $CKAP2L \quad 0.03545$
 - CLEC3B -0.01500
 - CNIH3 0.01413
 - CNNM1 -0.01611
- COL12A1 0.04098
- COL5A3 0.03177
- COL7A1 0.01688
- COLGALT1 0.02272
- COLGALT2 -0.00903
 - COX4I2 -0.00943
 - CSNK1D -0.01128
 - CST6 0.02032
 - CTSL -0.01263
 - CTSV 0.00987
 - CYP2S1 -0.01044
 - DCAF8 -0.02374
 - DCBLD2 0.03351
 - DCUN1D5 0.02056
- DENND1A 0.01898
 - DERA 0.01568
 - DHRS9 -0.00454
 - DKK1 0.00649
 - DNAJC9 0.01385
 - DPY19L1 0.00749
 - DSG2 0.01463
 - DSG3 0.02070
- DYNC2H1 -0.01537
 - E2F7 0.03923
 - EDIL3 0.01326
- EIF2AK3 -0.02073
- 21121110 0.02010
- ELMOD3 -0.03300
 - ENO2 0.02998

0.01550

EMP3

- EPHX2 -0.02392
- ERRFI1 0.01597
- EXOSC8 -0.00850
 - EYA3 0.02671
 - $FAH \qquad 0.01035$
- FAM120AOS -0.00980
 - FAM134B -0.01945
 - FAM189A2 -0.01692
 - FAM83A 0.01202
 - FAM91A1 0.01341
 - FBXO22 0.00649
 - FBXW8 -0.00891
 - FEM1B 0.04785
 - FER 0.02675
 - FGB -0.00252
 - $FGD6 \quad 0.02545$
 - FGG 0.00548
 - FHDC1 -0.01380
 - FLRT3 0.01416
 - FRZB -0.03715
 - FSCN1 0.02159
 - FST 0.01504
 - FYN -0.01133
 - GAB2 -0.03742
 - $GABPB1 \quad 0.01929$
 - GAPDH 0.02073
 - GATA6 -0.01780
 - GATC 0.02661
 - GIMAP2 -0.03176
 - GINS2 0.01713
 - GNPAT -0.01458
 - GOLM1 -0.01171
 - 0.011.1
 - GPC3 -0.02419
 - GPR176 0.00563
 - HIPK2 -0.02620
 - HJURP 0.02296

HRASLS20.00196HSP90B1-0.00641 HSPB6-0.01586ICAM2-0.00232IDH2 0.00528IFT140-0.02068IGFBP10.00427IGLL3P-0.01241**IKBIP** -0.00033 IL1R2-0.00660 ${\rm IL}20{\rm RB}$ 0.02671IL33-0.00991ITGA50.01407**ITPKB** -0.01390KANK40.03261KCNQ3 0.00040KCTD10 0.01501KCTD5-0.01440KIAA0513-0.02989 ${\rm KIAA1549L}$ 0.01354KIF14 0.01477KIF20A 0.02967KIF2C 0.01417KLHL5 0.02641KNTC1 0.02375KRT17 0.01644KRT6A 0.01795KRT6C 0.00798KRT70.01916**KYNU** 0.01181LAMA50.00174LCNL1 -0.01571LDHA 0.04004LETM20.01687LGALS9B -0.00232LINC01184 -0.01837

LMO3 -0.02246

LMTK2 0.00804

LOC100506562 -0.00290

LOX 0.02695

LYNX1 0.00001

MAP3K8 0.00338

MARCKSL1 -0.00884

MARS2 -0.01442

MC1R - 0.02281

MCEMP1 0.00025

 $MCM10 \quad 0.02451$

MCM4 0.02708

MCOLN2 -0.01684

MELK 0.02067

MEOX1 -0.01961

MIF = 0.01560

MIR99AHG -0.03712

MME = 0.01102

MRAP2 -0.01810

MRPL24 -0.01395

MTRNR2L1 -0.01563

NACC2 0.00733

 $\begin{array}{ll} NAMPT & 0.00071 \end{array}$

NCAPD2 0.02756

NCAPG 0.04487

NELFE -0.00390

NEURL2 0.01012

NFIA -0.03387

NFIX -0.01186

NMB -0.00205

NPM1 -0.01520

NR0B2 -0.01468

NRP2 0.00250

NUP155 0.02330

OAZ1 -0.00134

ORC1 -0.00199

P2RY2 0.01288P2RY8 -0.03043 P4HA10.00225P4HA20.01770PAX8 0.01350PAX8-AS10.00830PBXIP1 -0.01174PCDH20 -0.00861PCF11 -0.01710PCOLCE2 -0.00752PDLIM7 0.01678PEX11B -0.02280PFKFB4 0.00525PGAM5 0.00973PGBD3 0.01700PHACTR3 0.00172PHLDA1 0.03330PHOSPHO2 -0.02129**PIGL** 0.00833PLAC9 -0.02093PLAU 0.03213PLEKHS1 -0.01672PLIN2 -0.01174PLIN3 -0.00506 PLOD1 0.00369PLOD2 0.02261POC1A 0.01507POLA20.00692POP5-0.00224POU2AF1 -0.02222PP7080-0.01242PPAPDC1A 0.02867PPM1H-0.02311PPP1R12B 0.00096PPP1R14B 0.01352

0.00125

PPP1R3C

 PPY -0.02787PRC1 0.02492 ${\rm PRDM16}$ -0.02289PREP -0.01799 PRKCDBP0.00755PRMT7 -0.01665PROSER2 0.01761PRR11 0.01859**PTGES** 0.02681PTPN21 0.01723**PXDN** 0.02281**PYGL** 0.01714RAB31 0.01316RACGAP1 0.02957RALGAPB 0.02214RAP1GAP -0.03483RASL11B -0.01808RAVER2-0.01352RBMS2 0.02834RERE -0.01635RERGL -0.01801RFC50.01848RFK-0.01090 RFX2 -0.00264RGS3 -0.00319RGS5 -0.01505RHOF 0.02828RMND5A -0.00614RNF103-0.03019RPA2-0.02756RPIA -0.02226SAMD5 -0.00655SCGB2A1 -0.01773SCYL20.01826SDIM1 -0.01083

-0.01125

SEC23IP

SELENBP1 -0.02707

SEPW1 -0.01161

SERPINB3 -0.00201

SERPINH1 0.02086

SERTAD2 -0.00995

SGSM1 -0.02933

SH3GL1 -0.02784

SLAMF9 -0.00761

SLC12A2 -0.01821

SLC15A1 -0.00139

SLC16A3 0.01842

SLC2A1 0.01424

SLC2A3 0.00438

SLC30A3 -0.01126

SLC40A1 -0.02146

SMOX -0.02258

SNORA11D -0.00256

SNRPB 0.00276

SOBP -0.03269

SOD2 0.00120

SPHK1 0.03861

SPIN4 0.01254

SPOCD1 0.02117

SPOCK1 0.03046

SPP1 0.00175

ST3GAL2 -0.02187

ST6GAL1 -0.02118

ST6GALNAC1 -0.01232

STAT5B -0.03172

STK39 -0.01196

SUGCT 0.01833

SULF2 0.01494

SYNE2 -0.00968

TAF5L -0.01213

TARBP2 -0.01019

TCEA3 -0.02679

- TCTA -0.03326
- TGFBI 0.03259
- THSD7B -0.01931
 - TLE4 -0.01794
- TM9SF3 -0.01255
- TMED1 -0.01796
- TMEM26 0.03659
- TMTC4 -0.01797
- TNFRSF10D -0.00315
- TNFRSF17 -0.01180
- $TNFRSF6B \quad 0.02308$
 - TOM1 -0.01640
 - TOM1L2 0.00266
 - TOR2A -0.02926
 - TPD52L2 -0.00579
 - TPX2 0.02590
- TRAPPC2 -0.01920
- 110111 02 -0.01320
 - TREM1 -0.00073
- $TRERF1 \quad 0.00581$
 - TRIM2 -0.02689
 - TSTD1 -0.02503
- TUBA1C 0.02053
- $TWIST1 \quad 0.02246$
 - UFC1 -0.03123
 - UHRF2 0.01445
 - UPP1 0.00182
 - USP30 0.00629
 - VPS35 -0.01219
- $VSTM2L \quad 0.00352$
- WNT2B -0.00812
- XXYLT1 0.00341
 - ZBED2 0.02396
 - ZFPM1 -0.02180
- ZNF185 0.01435
- ZNF565 -0.00565
- ZNF658 -0.01988

ZPLD1 0.00165 ZSCAN16 -0.00720 ZSCAN32 -0.02184

Glossary

APGI Australian Pancreatic Cancer Genome Initiative. iv, 5, 12, 17–22, 24, 26, 28–30, 32, 35, 37, 39, 40, 73, 74

BAM binary sequence alignment / map file. 43, 44, 46

BIC Bayesian information criterion. 8, 55, 56

BMC Bayesian model comparison. 45

CA-19-9 carbohydrate antigen 19-9. 3, 4

CNV copy number variation. v, 47, 49, 50, 52–56

CPH Cox proportional hazard. 7, 8

CPSS complementary pair subset selection. 19, 35, 37

CPV clinico-pathological variable. iv, 4, 6, 17, 18, 31, 34, 35, 40, 41

CT computed tomography. 6

CV cross-validation. 23

DSD disease-specific death. 19, 35

DSS disease-specific survival. iii, 23, 35

ECM extracellular matrix. 32

EMT epithelial to mesenchymal transition. 4, 29, 32, 33

EUS endoscopic ultrasound. 5, 6

FAST feature aberration at survival times. 19, 35, 37

FDR false-discovery rate. 19, 35, 54–56

FNA fine needle aspirate. 5, 6

FWER familywise error rate. 25, 27

GATK Genome analysis toolkit. 44

GEO Gene Expression Omnibus. 38

GEX gene expression. iii, 12–19, 35, 37, 39, 74

GG generalised gamma. 9

GSVA gene set variation analysis. 32, 39, 40

HMM hidden Markov model. 53–55

IBS integrated Brier score. 10

ICA independent component analysis. 14–16

ICGC International Cancer Genome Consortium. 40

IDAT Illumina data. 34, 40

IHC immunohistochemical. 5

IHC immunohistochemistry. 5–7

indel insertion / deletion event. 44

KM Kaplan-Meier. 9, 10

LA-BQSR local alignment and base quality score recalibration. 44

LASSO least absolute shrinkage and selection operator. iii, 21–23

LOESS local regression. 7

LOH loss of heterozygosity. 45, 47, 53, 54, 56

MDS multidimensional scaling. 35

MSigDB molecular signatures database. i, ii, iv, 13, 30, 32, 39–41, 70–72

MSKCC Memorial Sloan-Kettering Cancer Center. 4

NCBI National Center for Biotechnology Information. 38

NGS next-generation sequencing. 49, 50, 52

NMF non-negative matrix factorization. iii, 14–20, 35, 36, 73

NNLS non-negative least squares. 21, 23, 37, 73, 74

NSWPCN New South Wales Pancreatic Cancer Network. 6–10

PARSE prognostic axis risk stratification estimate. ii–iv, 23, 25, 27–30, 38, 39, 73, 74

PCA principal component analysis. 14–16

PDAC pancreatic ductal adenocarcinoma. 12, 13, 16–19, 25, 26, 29, 35, 38

PH proportional hazard. 8

PPV positive predictive value. 4

SIS sure independence screening. 19, 35, 37

SNMF/L sparse non-negative matrix factorization, long variant. iii, 19, 20, 22, 35–37

SNR signal-to-noise ratio. 49

SNV single nucleotide variant. 44

TCGA The Cancer Genome Atlas. iv, 25, 27–29, 38

VST variance stabilizing transform. 34, 36, 39

WGS whole genome sequencing. 50

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