

Cancer metabolic subtypes and their association with molecular and clinical features

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Abstract

Alterations of metabolic pathways in cancer have been investigated for many years, beginning way before the discovery of the role of oncogenes and tumor suppressors, and the last few years have witnessed a renewed interest in this topic. Large-scale molecular and clinical data on tens of thousands of samples allow us today to tackle the problem from a general point of view.

Here we show that transcriptomic profiles of tumors can be exploited to define metabolic cancer subtypes, which can be systematically investigated for association with other molecular and clinical data. We find thousands of significant associations between metabolic subtypes and molecular features such as somatic mutations, structural variants, epigenetic modifications, protein abundance and activation; and with clinical/phenotypic data including survival probability, tumor grade, and histological types.

Our work provides a methodological framework and a rich database of statistical associations, accessible from metaminer.unito.it, that will contribute to the understanding of the role of metabolic alterations in cancer and to the development of precision therapeutic strategies.

Introduction

The last few years have witnessed a renewed interest in the metabolism of cancer, with the reprogramming of energy metabolism by cancer cells having been recently added, together with immune evasion, to the fundamental oncogenic processes collected under the heading of “Hallmarks of Cancer” [1].

Today, next-generation sequencing and other high-throughput techniques provide us with the unprecedented opportunity to take the study of the role of metabolic alterations in cancer progression to a new level, complementing the in-depth study of specific instances of metabolic changes with unbiased assessments of cancer metabolism and its relationship with the other hallmarks. In particular the TCGA project provides us with a systematic repository of genomic, epigenomic, transcriptomic, and clinical data on tens of thousands of samples of many tumor types. For example, a recent study [2] demonstrated the close relationship between patterns of copy-number alterations and metabolic phenotypes. Another study [3] showed the clinical relevance of metabolic pathways, as reflected by gene expression profiles of metabolic genes, to patient survival.

In this work we undertake a systematic analysis of the associations between metabolic profiles of human tumors and their molecular and clinical features. Our aim is to underline the role of metabolism in precision

cancer medicine, by showing that metabolic profiling can classify tumors into classes that are significantly different in both their molecular-level features (somatic mutations, structural genomic variants, epigenetic modifications) and in their clinical aspects (survival probability, tumor grade, histological type). While intrinsically correlative in nature, this analysis can serve as a guide to mechanistic studies linking metabolism to other molecular features, and to the exploration of precision therapeutic approaches informed by tumor metabolism.

While, ideally, such a study should be based on direct assays of metabolite abundance in primary tumors, these are not yet available on a large scale. Therefore we use transcriptomic profiles of tens of thousands of samples of many tumor types as a proxy of metabolomic assays to classify them into metabolic subtypes. We then systematically correlate such subtypes with molecular and clinical data using appropriate statistical tests, thus generating a database of thousands of associations.

Results

Transcriptome-based classification of tumors into metabolic subtypes

For each tumor type with transcriptomic data available from the TCGA, and for each metabolic pathway defined as a set of genes by annotation databases such as KEGG [4,5], Reactome [6], and the Gene Ontology [7,8], we divided the patients into metabolic subtypes by clustering the samples using the expression profiles of the genes belonging to the pathway. Fig. 1 shows the KEGG pathways used in the analysis and the genes assigned to each pathway as a bipartite network.

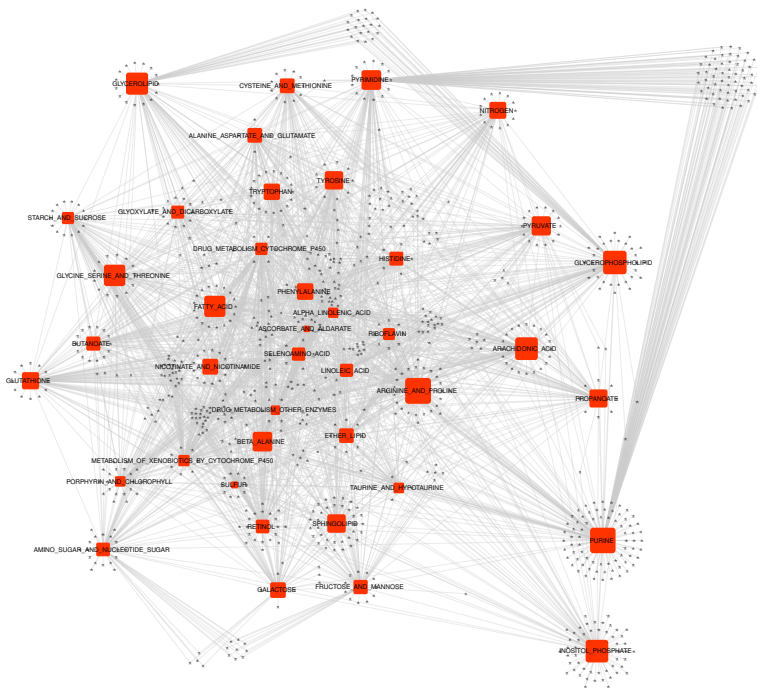


Figure 1: The KEGG pathways (red squares) used to cluster tumor samples and the respective genes (grey triangles) represented as a bipartite network. The size of the squares is proportional to the number of associated genes.

Compared to differential expression analyses used e.g. in [3], clustering by metabolic pathways correctly takes into account the fact that the genes associated to a pathway are often involved in either anabolic or catabolic processes. Therefore the activation of the pathway in a subset of patients is typically reflected by the transcriptional activation of the former and repression of the latter. For example, in Fig. 2, we show the clustering of low-grade glioma samples based on the “arachidonic acid metabolism” KEGG pathway: Each

cluster is characterized by both up- and downregulated genes, a structure that would not be captured by simple differential expression of the pathway genes as a whole.

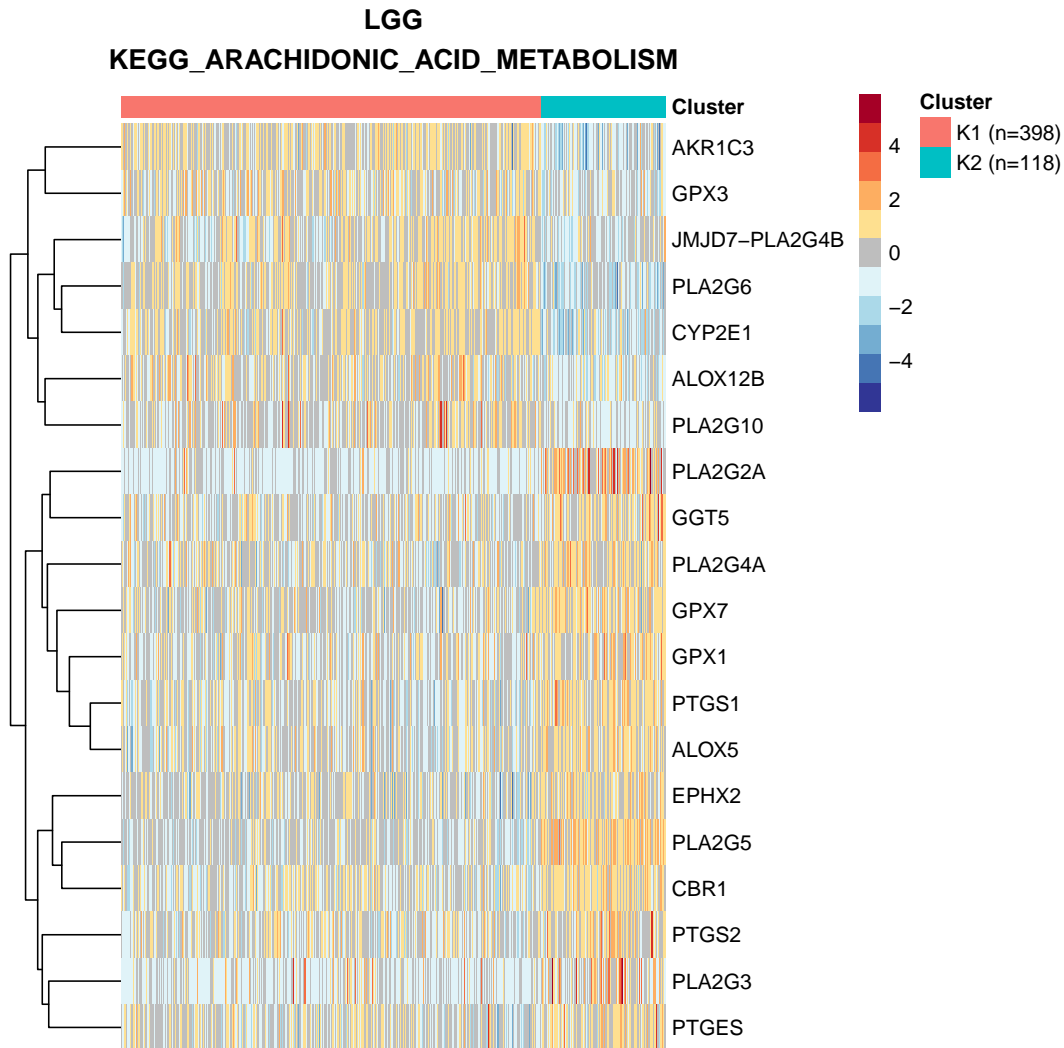


Figure 2: Clustering of low-grade glioma patients using the genes involved in arachidonic metabolism according to KEGG. Only the 20 genes most differentially expressed between the two clusters are shown.

To perform the clustering we used partitioning around medoids (PAM) [9], a more robust method compared to k-means [10] with respect to the presence of outliers. Cluster silhouette analysis allowed an unsupervised choice of the number of clusters. A total of 345 metabolic gene sets were used, as described in the Methods, on 38 tumor classes, thus performing a total of 13110 clusterings. In the majority of cases (9654, 73.6%) the samples were subdivided into $k = 2$ clusters, while in other cases we obtained up to 10 clusters. These clusters will be referred to as “metabolic subtypes.” The tumor types analyzed with the number of samples for which expression data were available are shown in Table 1.

Table 1: Number of samples used for each tumor class.

ID	Tumor_type	Samples
ACC	Adrenocortical Carcinoma	79
BLCA	Bladder Urothelial Carcinoma	408

ID	Tumor_type	Samples
BRCA	Breast Invasive Carcinoma	1093
CESC	Cervical Squamous Cell Carcinoma and Endocervical Adenocarcinoma	304
CHOL	Cholangiocarcinoma	36
COAD	Colon Adenocarcinoma	285
COADREAD	(= COAD + READ)	379
DLBC	Lymphoid Neoplasm Diffuse Large B-cell Lymphoma	48
ESCA	Esophageal Carcinoma	184
GBM	Glioblastoma Multiforme	153
GBMLGG	(= GBM + LGG)	669
HNSC	Head and Neck Squamous Cell Carcinoma	520
KICH	Kidney Chromophobe	66
KIPAN	(= KICH + KIRC + KIRP)	889
KIRC	Kidney Renal Clear Cell Carcinoma	533
KIRP	Kidney Renal Papillary Cell Carcinoma	290
LAML	Acute Myeloid Leukemia	173
LGG	Brain Lower Grade Glioma	516
LIHC	Liver Hepatocellular Carcinoma	371
LUAD	Lung Adenocarcinoma	515
LUNG	(= LUAD + LUSC)	1016
LUSC	Lung Squamous Cell Carcinoma	501
MESO	Mesothelioma	87
OV	Ovarian Serous Cystadenocarcinoma	303
PAAD	Pancreatic Adenocarcinoma	178
PCPG	Pheochromocytoma and Paraganglioma	179
PRAD	Prostate Adenocarcinoma	497
READ	Rectum Adenocarcinoma	94
SARC	Sarcoma	259
SKCM	Skin Cutaneous Melanoma	103
STAD	Stomach Adenocarcinoma	415
STES	(= STAD + ESCA)	599
TGCT	Testicular Germ Cell Tumors	150
THCA	Thyroid Carcinoma	501
THYM	Thymoma	120
UCEC	Uterine Corpus Endometrial Carcinoma	176
UCS	Uterine Carcinosarcoma	57
UVM	Uveal Melanoma	80

It is worth asking whether different metabolic gene sets cluster the patients of the same tumor in significantly different ways. To answer this question we computed the normalized mutual information between the cluster assignments obtained in each tumor with each metabolic gene set. The results are shown in Fig. 3, and suggest that, in most cases, different metabolic gene sets cluster tumor patients in clearly distinct ways.

Associations between metabolic subtypes and molecular and clinical features

We then proceeded to systematically explore the statistical associations between metabolic subtypes and several molecular and clinical features of the corresponding tumors, classified into 9 variable classes. Statistical tests appropriate to the nature of the variables describing such features were chosen as described in the Methods. Bonferroni correction was applied separately to each variable type but to all tumor types together. Thus, for example, the Bonferroni correction for the association with miRNA expression takes into account

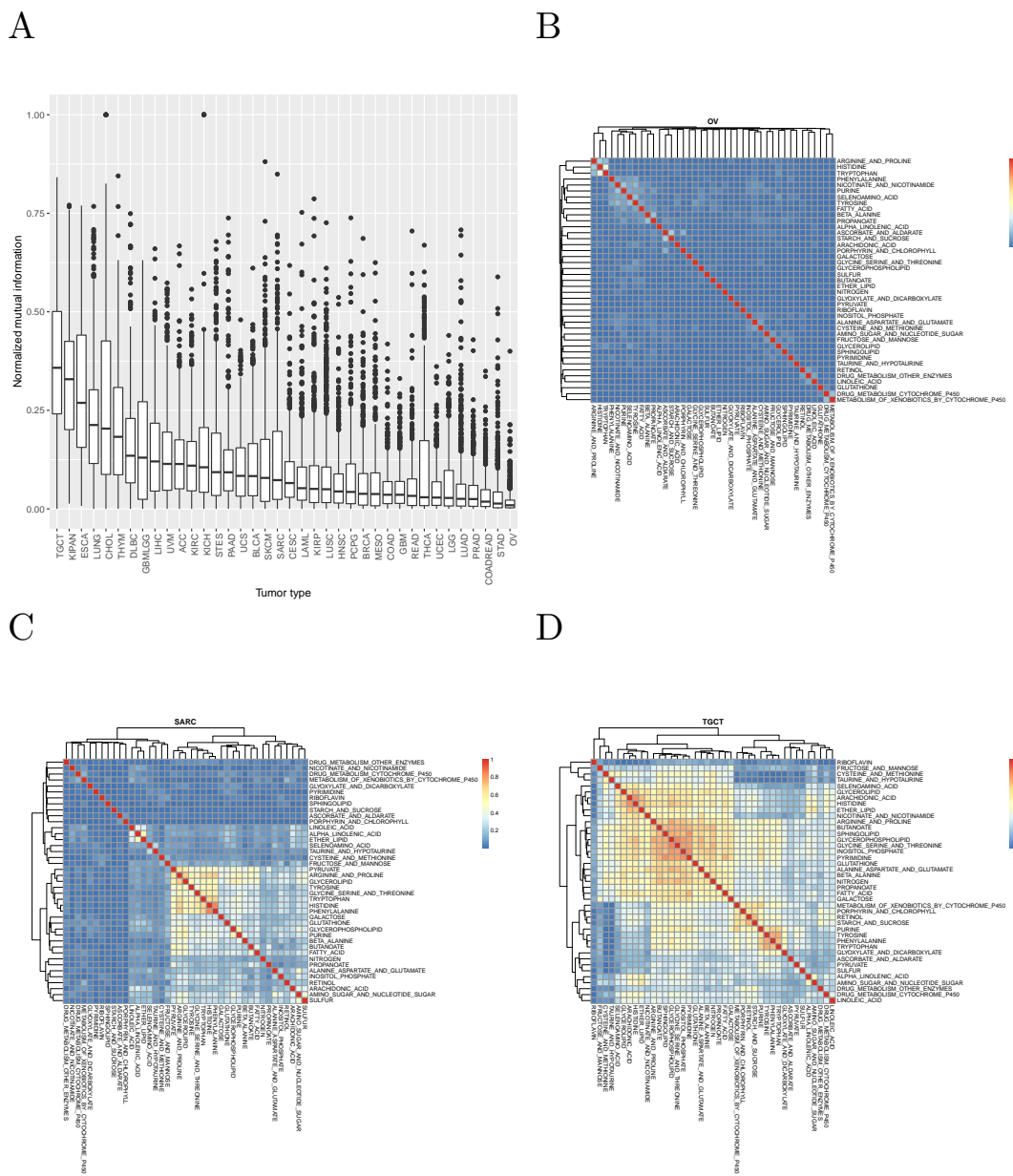


Figure 3: A: Normalized mutual information between clusterings of samples using KEGG metabolic gene sets. B, C, D: examples of tumor types with low, intermediate, and high mutual information.

all tests of association between all miRNAs and all metabolic gene sets across all tumor types. The molecular and phenotypic variables considered are shown in the following table:

Table 2: Molecular and phenotypic variables analyzed for association with metabolic subtypes

class	description	type
CNA	Copy number alterations	molecular
Methylation	Global DNA methylation	molecular
miRNA	microRNA expression	molecular
Mutations	Specific point mutations	molecular
Gene Mutations	Gene-level mutations	molecular
RPPA	Protein expression	molecular
Clinical	Clinical and histologic parameters	phenotypic
OS	Overall survival	phenotypic
RFS	Recurrence-free survival	phenotypic

In the following, to limit redundancy, we focus on the results obtained with the KEGG annotation database (a total of 99621 significant associations). Figure 4A shows the distribution of these associations among tumor types. Note that the number of significant associations for a given tumor type will depend in general on the number of available samples: indeed the Spearman correlation coefficient between number of associations and number of samples is 0.755. In Figure 4B we show the number of associations by KEGG metabolic pathway, suggesting a particular relevance of the metabolism of aminoacids and fatty acids in classifying tumors into classes characterized by different molecular and/or clinical characteristics.

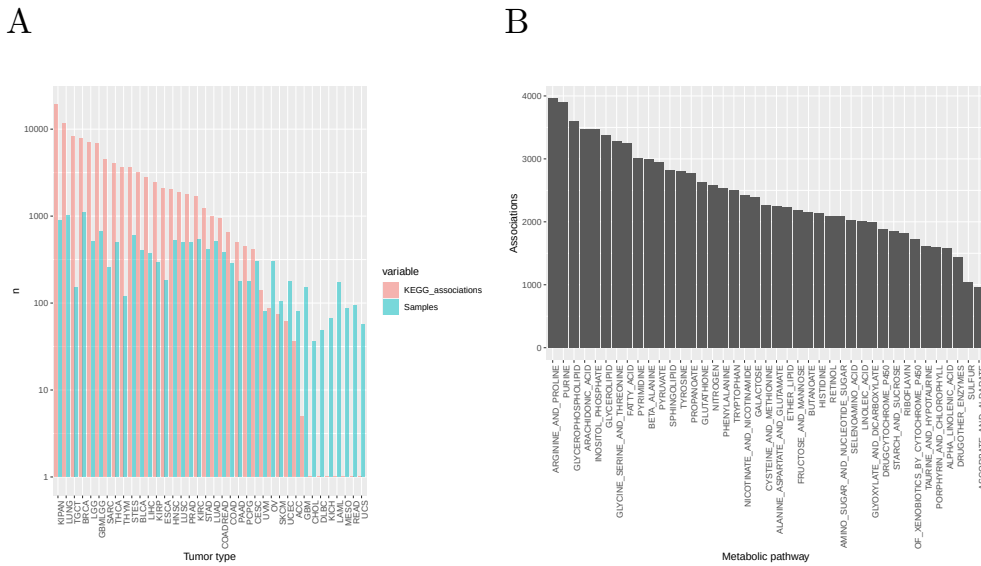


Figure 4: (A) Number of samples and of significant associations with molecular and clinical features by tumor type. (B) Number of significant associations by KEGG metabolic pathway.

Recurrent associations

Of particular interest are recurrent associations, that is associations that are statistically significant in more than one tumor type (not considering the tumor types built from the union of smaller types, such as LUNG,

KIPAN, etc.).

We found a total of 14937 such recurrent associations. Table 3 shows the most recurrent associations for each class of clinical and molecular variables. Notably, the only variable classes with no recurrent associations are point mutations: All the 45 associations with specific mutations and the 102 associations with gene-level mutations are specific of a single tumor type.

Table 3: Top recurrent associations for each class of variables

Variable	Metabolism	Number of associations	Tumors associated
miRNA: hsa-mir-222	INOSITOL PHOSPHATE	12	BLCA, BRCA, CESC, HNSC, KIRC, LGG, LIHC, LUAD, SARC, STAD, TGCT, THCA
Methylation	GLYCEROLIPID	10	BLCA, CESC, COAD, HNSC, LGG, PCPG, SARC, TGCT, THCA, THYM
Methylation	INOSITOL PHOSPHATE	10	BLCA, COAD, KIRP, LGG, LIHC, LUAD, PCPG, TGCT, THCA, THYM
Methylation	PHENYLALANINE	10	BLCA, CESC, COAD, LGG, LIHC, LUAD, OV, PCPG, SARC, TGCT
Methylation	PURINE	10	BLCA, CESC, KIRP, LIHC, LUAD, PCPG, SARC, TGCT, THCA, THYM
Clinical: histological type	TYROSINE	8	BRCA, ESCA, LGG, PCPG, SARC, THCA, THYM, UCEC
RPPA: CCNB1 Cyclin B1	PURINE	7	BLCA, BRCA, KIRC, KIRP, LUAD, TGCT, THYM
CNA: chr17p12	FATTY ACID	6	COAD, KIRP, LGG, OV, PAAD, TGCT
CNA: chr17q11	GLYCEROLIPID	6	BRCA, LGG, LUSC, SARC, TGCT, THYM
CNA: chr17q21	GLYCEROLIPID	6	BRCA, HNSC, LGG, LUSC, TGCT, THYM
CNA: chr3p	INOSITOL PHOSPHATE	6	BLCA, BRCA, ESCA, KIRC, LUSC, UVM
CNA: chr3p14	INOSITOL PHOSPHATE	6	BLCA, BRCA, ESCA, KIRC, LUSC, UVM
CNA: chr3p21	ARGININE AND PROLINE	6	BRCA, ESCA, KIRC, PRAD, SARC, THYM
CNA: chr3p21	INOSITOL PHOSPHATE	6	BLCA, BRCA, ESCA, KIRC, LUSC, UVM
CNA: chr3q	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q24	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q25	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q26	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q27	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q28	FATTY ACID	6	BRCA, ESCA, HNSC, LGG, LUAD, LUSC
CNA: chr3q28	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
CNA: chr3q29	FATTY ACID	6	BRCA, ESCA, HNSC, LGG, LUAD, LUSC

Variable	Metabolism	Number of associations	Tumors associated
CNA: chr3q29	PURINE	6	BRCA, ESCA, HNSC, LUSC, UCEC, UVM
OS	PROPANOATE	2	KIRC, LGG
OS	PURINE	2	ACC, KIRC
OS	PYRIMIDINE	2	ACC, LGG
OS	SPHINGOLIPID	2	KIRC, LGG

Figure 5 shows the expression of microRNA miR-22 in the clusters of breast cancer and lung adenocarcinoma patients obtained with the genes associated to inositol phosphate metabolism.

Specific examples

In the following we show some examples of associations, some confirming known results and some suggesting new lines of investigation, especially in less widely studied tumor types.

TP53 mutations and glucose metabolism

Mutant TP53 has been shown to promote the Warburg effect, possibly the most notorious metabolic hallmark of cancer, in breast and lung cancer cell lines [11]. Indeed clustering breast and lung tumors using the genes associated to glucose metabolism according to REACTOME shows, in both cancer types, the appearance of a cluster characterized by overexpression of several Warburg effect signature genes (such as GAPDH, PGK1, and PKM2) and by an enrichment in TP53 mutations (Fig. 6).

Pyruvate metabolism in thyroid cancer

The clustering of thyroid cancer samples according to the genes associated by KEGG to pyruvate metabolism is shown in Fig. 7A. These 2 clusters show many significant differences in both clinical and molecular parameters. In particular (Fig. 7B,C), the prevalence of BRAF and NRAS-mutated samples is strikingly different in the two clusters, with *all* NRAS-mutated samples found in cluster 2 and the large majority of BRAF-mutated samples found in cluster 1. The two clusters also strongly differ in the respective prevalence of histological types.

Fatty acid metabolism in thymoma

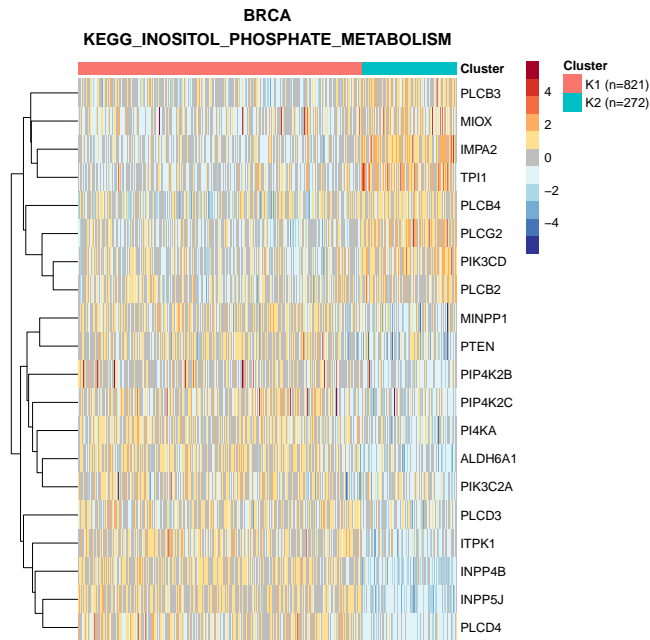
Clustering of thymoma samples using the genes associated to fatty acid metabolism divides the samples in two clusters (Fig. 8A) which largely overlap the known histological types (Fig. 8B) and differ in the prevalence of a recurrent GTF2I mutation [12].

Analysis of cancer cell lines

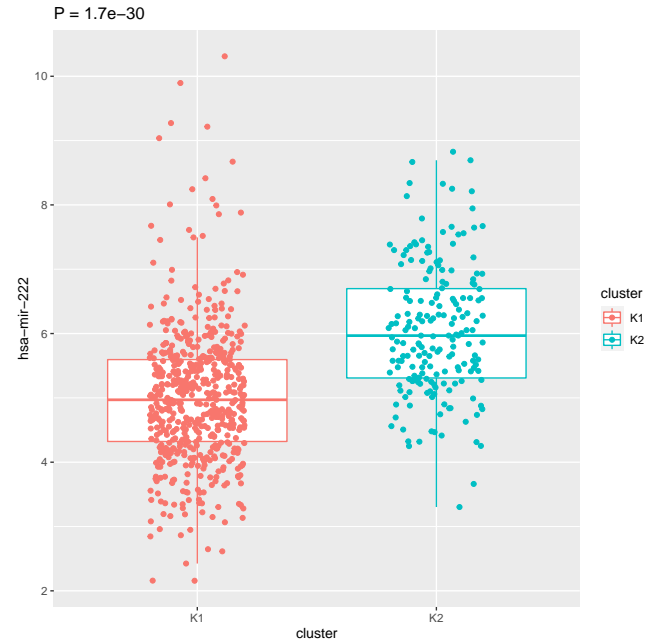
We performed the same analysis on the cancer cell lines included in the cancer cell line encyclopedia [13]. These cell lines are classified in terms of TCGA tumor types (22 types are represented by a total of 723 CCLE cell lines). Within each tumor type, cell lines were clustered based on their transcriptomic data, using the expression of the genes in the same metabolic gene sets used to cluster TCGA samples. The clusters were then correlated with molecular and phenotypic data, including copy number alterations, specific point mutations, gene-level mutations, protein expression, microRNA expression, and drug response data (IC50 values).

Since the number of cell lines in the CCLE is about one order of magnitude smaller than the number of TCGA samples, the power to detect significant associations is also much smaller. Nevertheless, we found 485 significant associations at a Bonferroni-corrected significance level of 0.05. For example, KEAP1 mutations are associated with xenobiotic metabolism in both TCGA samples and CCLE cell lines (Fig. 9), in agreement with the known role of the NRF2/KEAP1 pathway in the regulation of xenobiotic response [14]. Note that KEAP1 mutations appear to be associated to the activation of the pathway, as observed in [15].

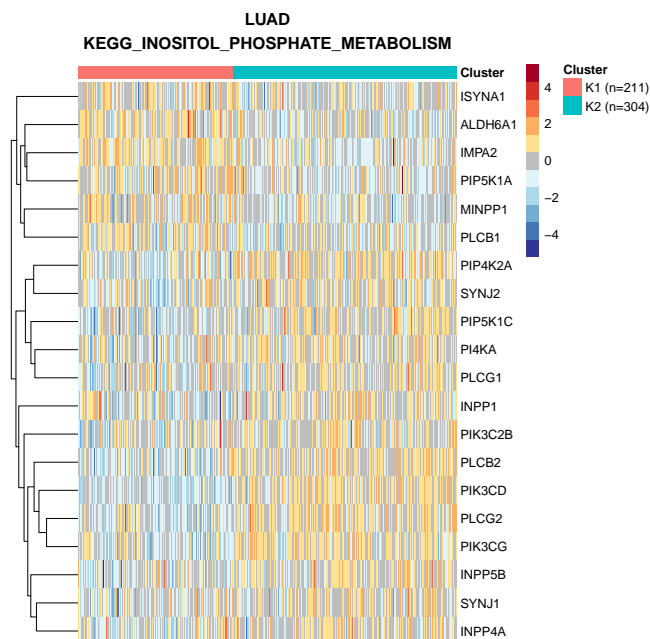
A



B



C



D

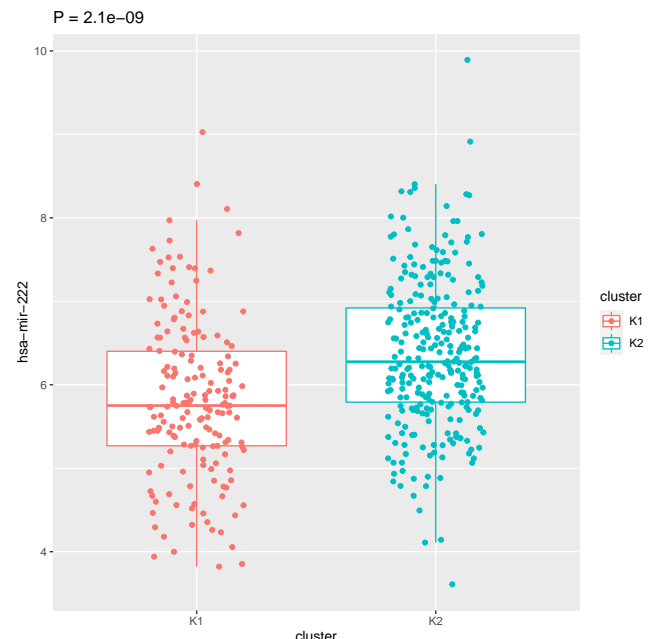
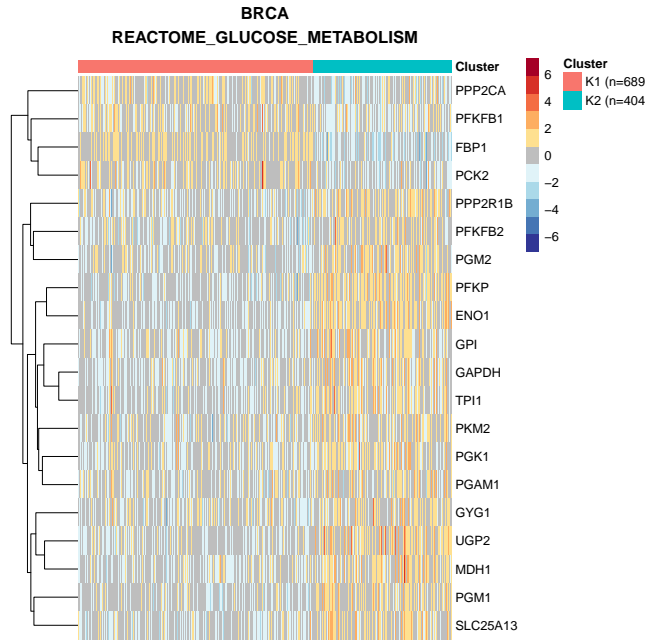
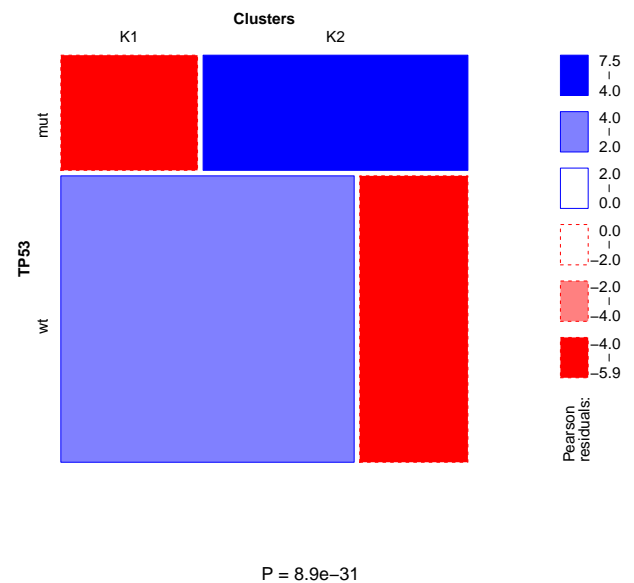


Figure 5: (A) Clustering of breast cancer samples using the genes associated to inositol phosphate metabolism according to KEGG (B) Expression of miR-222 in the two clusters. (C, D) Same for lung adenocarcinoma samples

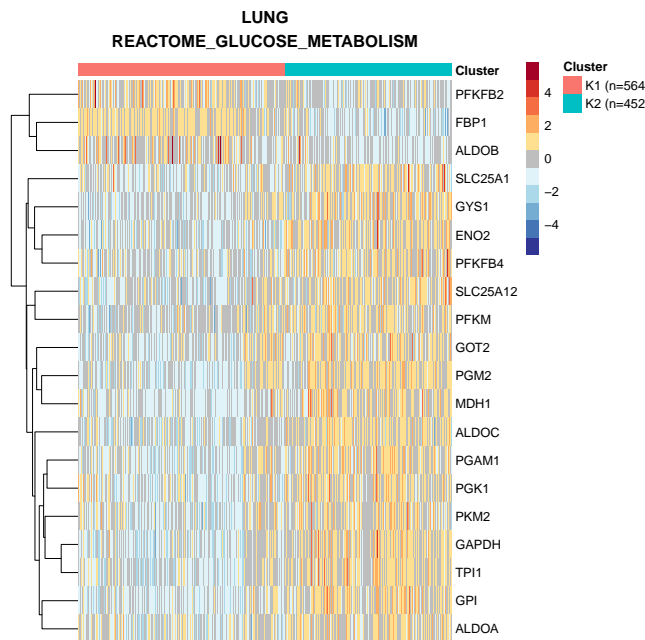
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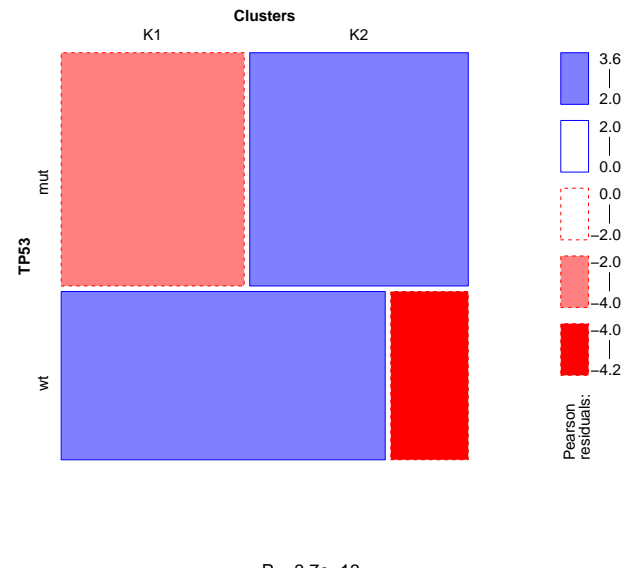
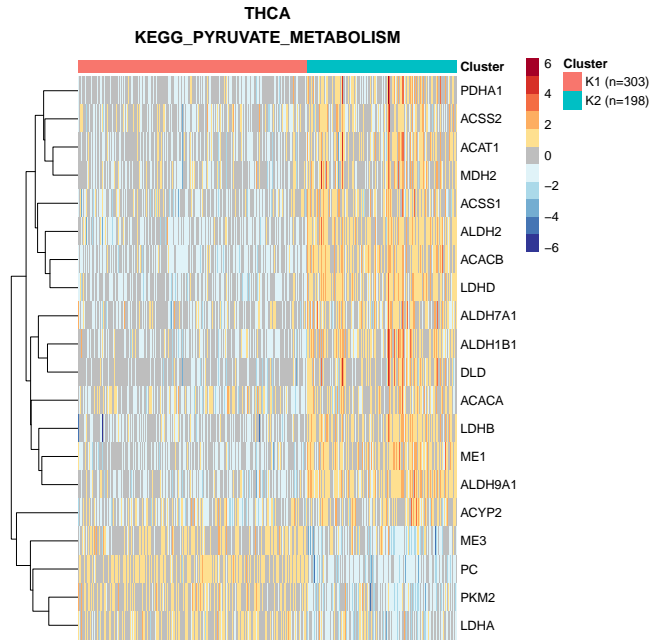
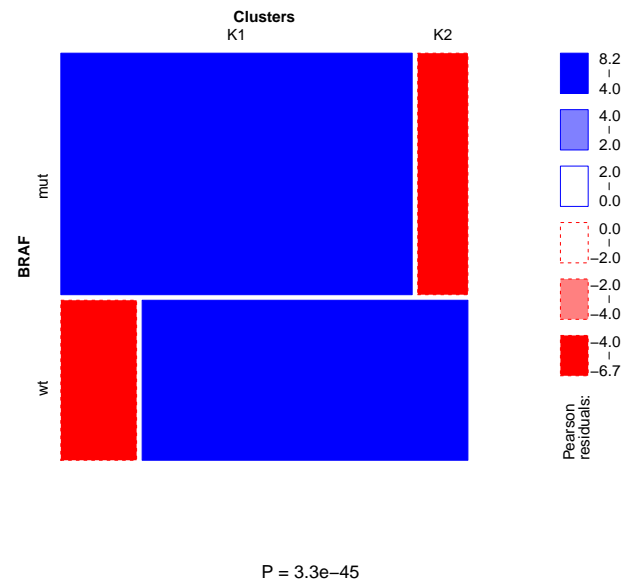


Figure 6: (A) Clustering of breast cancer samples using the genes associated to glucose metabolism according to REACTOME. Cluster K2 shows upregulation of Warburg effect genes such as GAPDH, PGK1, and PKM2 (B) The prevalence of TP53-mutated samples is significantly higher in cluster K2. (C, D) Same for lung cancer samples

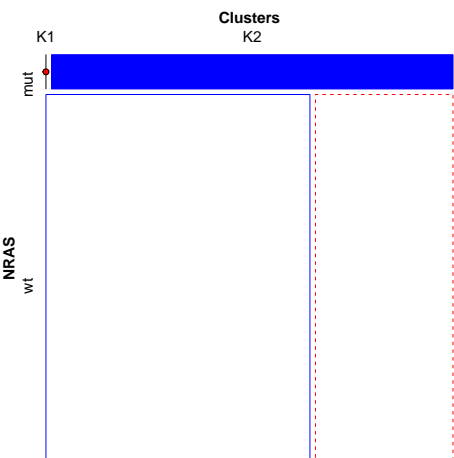
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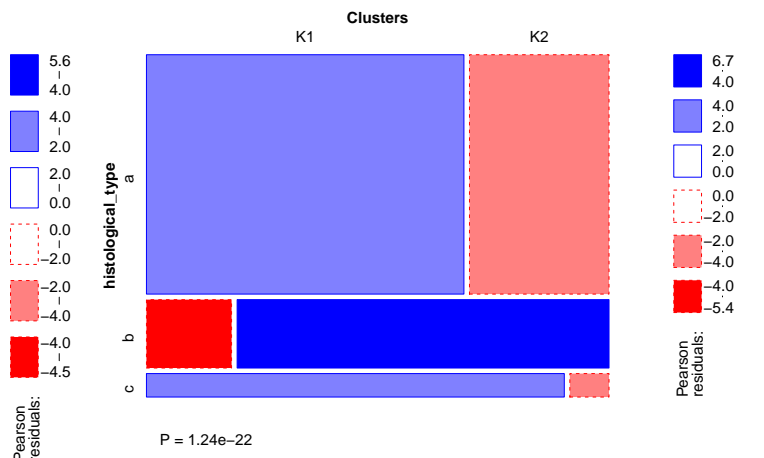


Figure 7: (A) Clustering of thyroid cancer samples using the genes associated by KEGG to pyruvate metabolism. (B) Most BRAF-mutated samples fall into cluster 1 (C) All NRAS-mutated samples fall into cluster 2. (D) The two clusters significantly differ in the prevalence of histological types.

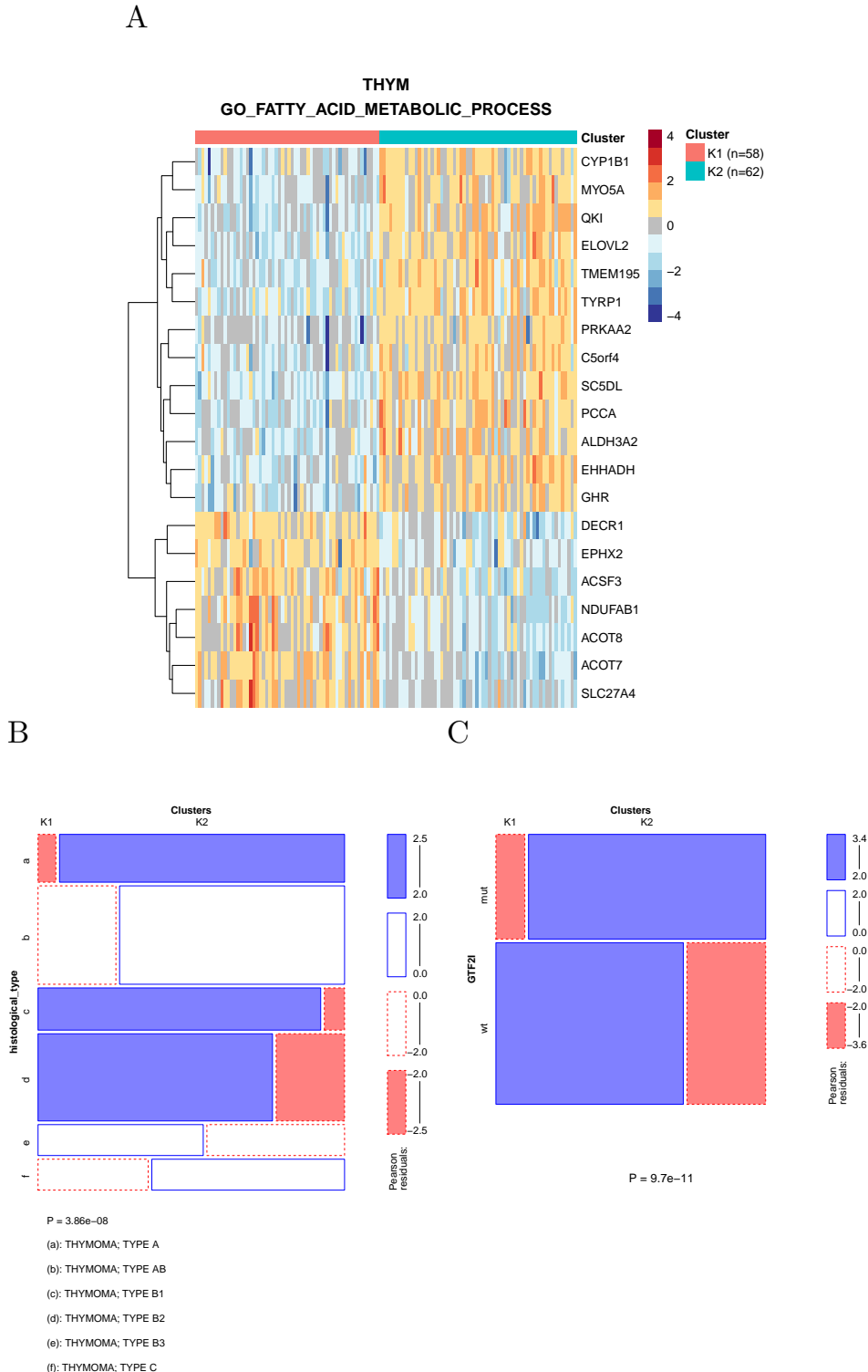


Figure 8: (A) Clustering of thymoma samples using the genes associated by KEGG to fatty acid metabolism. (B) The clusters significantly overlap the known histological types (C) A specific GTF2I mutation significantly segregates between the two clusters

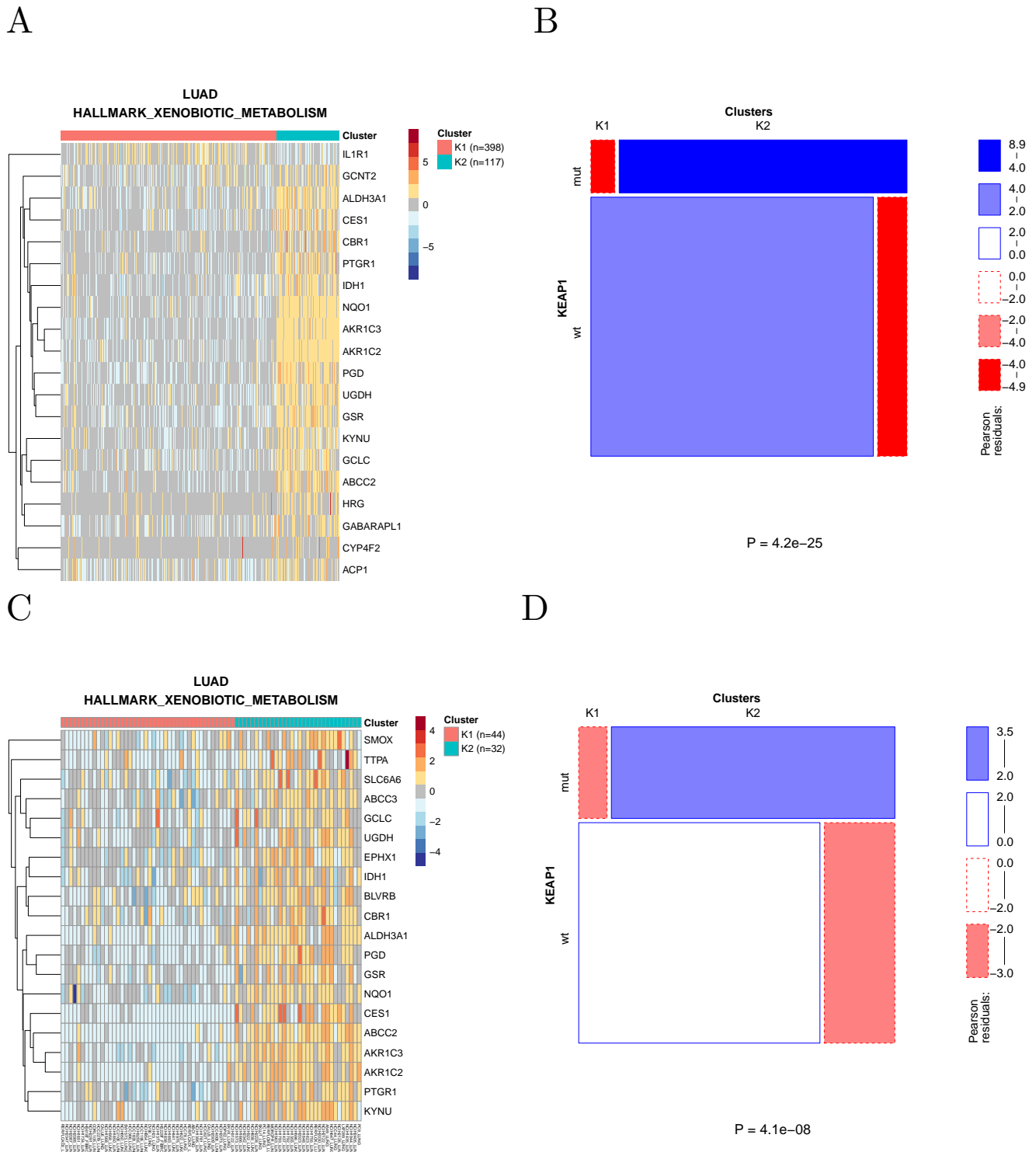


Figure 9: (A) Clustering of lung adenocarcinoma samples using the genes in the xenobiotic metabolism hallmark gene set. (B) Mutations of KEAP1 are strongly enriched in the K2 cluster where the pathway appears to be activated. (C) Clustering of lung adenocarcinoma cell lines using the same gene set. (D) Also in cell lines mutations are enriched in the cluster showing pathway activation.

Discussion

In this work we systematically analyzed cancer metabolic subtypes, defined through clustering of patients using the transcription profiles of metabolic gene sets. These subtypes are highly specific of each individual metabolism used to define the gene sets, and show a myriad of significant associations with both molecular and phenotypic features of the tumors.

We present these associations as a resource for the community, available from metaminer.unito.it which we believe will be useful in two related but distinct ways. One one hand, our results highlight the key role of metabolic profiles in classifying individual patients in a way that is potentially useful to devise precision treatments. This is particularly compelling for tumors for which no established classification exists, but in which metabolic subtypes strongly associated with phenotypic characteristics can be found.

On the other hand, our results can help formulating hypotheses on the mechanisms underlying the associations between metabolic subtypes on one hand and molecular and phenotypic features on the other, which can then be experimentally tested in cell lines or organoids. For example, while both miR-222 (see e.g. [16] for a recent review) and inositol phosphate metabolism [17,18] have been often associated to cancer progression, their association, which recurs in a large number of tumor types, has not been previously reported to the best of our knowledge, and deserves a mechanistic investigation.

Indeed the correlative nature of our results is the most obvious limitation of our results, and of all results obtained by the retrospective analysis of primary tumors.

Methods

Data

TCGA data

All TCGA data (release 2016_02_28) were obtained from the Broad TCGA GDAC site (<https://gdac.broadinstitute.org/>), by means of firehose_get, version: 0.4.1. The gene expression data clustered to generate the metabolic subtypes were the normalized gene-level TPMs obtained with RSEM.

Copy number data were obtained from the segmented data using the cghMCR [19], DNAcopy [20], and CNTTools [21] Bioconductor packages. The cghMCR package allows the calculation of segment gain or loss (SGOL) from segmented data, by means of a modified version of the GISTIC algorithm. The *segment* function of the DNAcopy package is used to segment the normalized data so that chromosome regions with the same copy number have the same segment mean values. Then, using the *getRS* function from CNTTools, the data returned by segment are organized in a matrix format. Finally the *SGOL* function of cghMCR is used to compute gene-level SGOL scores for genes by calculating the sum (parameter method) of all the positive and negative values, respectively above and below a set threshold (0.5, -0.5). Finally, SGOL scores for all genes included in each Broad Institute positional gene set were averaged to obtain the SGOL score of each region.

Genome-wide DNA methylation was obtained from the gene-level summarized Human Methylome 450k data, as provided by GDAC, then summed over all genes to associate a global DNA methylation value to each sample. miRNA expression was obtained from the Illumina HiSeq data as deposited in GDAC. Specific point mutations for each sample were obtained from the “mutation packager oncotated calls” provided by GDAC, considering only non-synonymous mutations. For gene-level mutations, a sample was considered as mutated in a gene if it carried one or more specific point mutations associated to the gene.

Protein expression data (RPPA) were obtained from the corresponding files annotated with gene names from the GDAC site. Survival and other clinical data were obtained from the “Tier 1 clinical pick” files available from GDAC.

CCLE data

Gene expression data of 723 cell lines associated to one of the TCGA tumor types were obtained, as gene-level TPMs computed with RSEM, from the CCLE web site. We also retrieved from the CCLE web site the

following data to be correlated to metabolic subtypes:

- gene-level copy number data: the absolute score provided by CCLE was processed with the same steps used for the SGOL score reported for the TCGA data to obtain region-based copy number data
- miRNA expression data (which were log-transformed for display in the figures)
- somatic mutation data (also here we considered only non-synonymous mutations)
- IC50 values for the available drugs
- metabolite levels

Metabolic gene sets

Gene sets were obtained from the c2.KEGG, c2.REACTOME, c5.BP and Hallmark MSigDB v5.2 collections. Metabolic gene sets were defined as those whose name matched the string “metabol” but not the string “regul” (to include only gene sets directly involved in metabolic pathways rather than in their regulation). In this way we obtained 345 metabolic gene sets (41, 35, 265, and 4 for the c2.KEGG, c2.REACTOME, c5.BP, and Hallmark collections, respectively).

Generation of metabolic clusters

For each tumor type and each metabolic gene set, the patients were divided into metabolic subtypes using partition around medoid (PAM) clustering [9] on the Spearman rank correlation coefficient-based distance matrix obtained from the expression of the genes in the gene set. The number of clusters (the maximum being set at 10) generated by each metabolic gene set was based on optimizing the average silhouette width. The normalized mutual information between two clusterings of the same patients based on different gene sets was computed with the R package *infotheo* [22], using the entropy of the empirical probability distribution and normalizing to the geometric mean of the entropies. Genes differentially expressed among clusters were identified using a Kruskal-Wallis test: the top 20 genes by P-value are shown in the heatmaps.

Statistical tests of association

The statistical tests used for establishing the association between each clinical/molecular feature and metabolic subtypes depend on the type of variable describing the feature. The metabolic subtype is always a categorical variable indicating the cluster membership of each patient.

- For continuous variables (microRNA expression, DNA methylation, CNA SGOL scores) we used Mann-Whitney U test (Kruskal-Wallis for $k > 2$). In the case of microRNA expression, given the large number of microRNAs with expression equal to zero in each sample, the expression data were rank-transformed with random resolution of ties before the statistical test.
- For categorical variables (e.g. presence of a given mutation) we used Fisher’s exact test (χ^2 test for tables larger than 2×2 , removing levels for which the expected count was less than 5)
- For survival (RFS or OS) we used the log-rank test. The test was not considered valid when the expected number of events in any metabolic subtype was less than 5

Multiple testing

Bonferroni correction was applied separately to each variable type but to all tumor types together. Thus, for example, the Bonferroni correction for the association with miRNA expression takes into account all tests of association between all miRNAs and all metabolic gene sets across all tumor types. We chose to perform the correction separately for the feature types because these contain very different numbers of variables (hundreds of variables for miRNA expression and a single one for overall survival): Therefore an overall Bonferroni correction would unduly penalize the variable types containing few variables.

Data availability

The results of the analysis can be browsed interactively at <https://metaminer.unito.it/>.

Supplementary material

- Supplementary data 1: clustering of TCGA patients into metabolic subtypes.
- Supplementary data 2: associations between metabolic subtypes and molecular/phenotypic features in TCGA.
- Supplementary data 3: clustering of CCLE cell lines into metabolic subtypes.
- Supplementary data 4: associations between metabolic subtypes and molecular/phenotypic features in CCLE.

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References

1. Hanahan D, Weinberg RA. Hallmarks of cancer: The next generation. *Cell*. 2011;144: 646–674. doi:[10.1016/j.cell.2011.02.013](https://doi.org/10.1016/j.cell.2011.02.013)
2. Graham NA, Minasyan A, Lomova A, Cass A, Balanis NG, Friedman M, et al. Recurrent patterns of DNA copy number alterations in tumors reflect metabolic selection pressures. *Mol Syst Biol*. 2017;13: 914. doi:[10.1525/msb.20167159](https://doi.org/10.1525/msb.20167159)
3. Gaude E, Frezza C. Tissue-specific and convergent metabolic transformation of cancer correlates with metastatic potential and patient survival. *Nat Commun*. 2016;7: 13041. doi:[10.1038/ncomms13041](https://doi.org/10.1038/ncomms13041)
4. Kanehisa M, Goto S. KEGG: Kyoto encyclopedia of genes and genomes. *Nucleic Acids Res*. 2000;28: 27–30. doi:[10.1093/nar/28.1.27](https://doi.org/10.1093/nar/28.1.27)
5. Kanehisa M, Sato Y, Furumichi M, Morishima K, Tanabe M. New approach for understanding genome variations in KEGG. *Nucleic Acids Res*. 2019;47: D590–D595. doi:[10.1093/nar/gky962](https://doi.org/10.1093/nar/gky962)
6. Fabregat A, Jupe S, Matthews L, Sidiropoulos K, Gillespie M, Garapati P, et al. The Reactome Pathway Knowledgebase. *Nucleic Acids Res*. 2018;46: D649–D655. doi:[10.1093/nar/gkx1132](https://doi.org/10.1093/nar/gkx1132)
7. Ashburner M, Ball CA, Blake JA, Botstein D, Butler H, Cherry JM, et al. Gene Ontology: Tool for the unification of biology. *Nat Genet*. 2000;25: 25–29. doi:[10.1038/75556](https://doi.org/10.1038/75556)
8. The Gene Ontology Consortium. The Gene Ontology Resource: 20 years and still GOing strong. *Nucleic Acids Res*. 2019;47: D330–D338. doi:[10.1093/nar/gky1055](https://doi.org/10.1093/nar/gky1055)
9. Kaufman L, Rousseeuw P. Clustering by means of Medoids. In: Dodge Y, editor. *Statistical Data Analysis Based on the L1 Norm and Related Methods*. North-Holland; 1987. pp. 405–416.
10. Arbin N, Suhaimi NS, Mokhtar NZ, Othman Z. Comparative Analysis between K-Means and K-Medoids for Statistical Clustering. 2015 3rd International Conference on Artificial Intelligence, Modelling and Simulation (AIMS). IEEE; 2015. pp. 117–121. doi:[10.1109/AIMS.2015.82](https://doi.org/10.1109/AIMS.2015.82)
11. Zhang C, Liu J, Liang Y, Wu R, Zhao Y, Hong X, et al. Tumour-associated mutant p53 drives the Warburg effect. *Nat Commun*. 2013;4: 2935. doi:[10.1038/ncomms3935](https://doi.org/10.1038/ncomms3935)

12. Petrini I, Meltzer PS, Kim I-K, Lucchi M, Park K-S, Fontanini G, et al. A specific missense mutation in GTF2I occurs at high frequency in thymic epithelial tumors. *Nat Genet.* 2014;46: 844–849. doi:[10.1038/ng.3016](https://doi.org/10.1038/ng.3016)
13. Ghandi M, Huang FW, Jané-Valbuena J, Kryukov GV, Lo CC, McDonald ER, et al. Next-generation characterization of the Cancer Cell Line Encyclopedia. *Nature.* 2019;569: 503–508. doi:[10.1038/s41586-019-1186-3](https://doi.org/10.1038/s41586-019-1186-3)
14. Motohashi H, Yamamoto M. Nrf2-Keap1 defines a physiologically important stress response mechanism. *Trends Mol Med.* 2004;10: 549–557. doi:[10.1016/j.molmed.2004.09.003](https://doi.org/10.1016/j.molmed.2004.09.003)
15. Gong M, Li Y, Ye X, Zhang L, Wang Z, Xu X, et al. Loss-of-function mutations in KEAP1 drive lung cancer progression via KEAP1/NRF2 pathway activation. *Cell Commun Signal.* 2020;18: 98. doi:[10.1186/s12964-020-00568-z](https://doi.org/10.1186/s12964-020-00568-z)
16. Ravegnini G, Cargnini S, Sammarini G, Zanotti F, Bermejo JL, Hrelia P, et al. Prognostic Role of miR-221 and miR-222 Expression in Cancer Patients: A Systematic Review and Meta-Analysis. *Cancers (Basel).* 2019;11. doi:[10.3390/cancers11070970](https://doi.org/10.3390/cancers11070970)
17. Hakim S, Bertucci MC, Conduit SE, Vuong DL, Mitchell CA. Inositol polyphosphate phosphatases in human disease. *Curr Top Microbiol Immunol.* 2012;362: 247–314. doi:[10.1007/978-94-007-5025-8_12](https://doi.org/10.1007/978-94-007-5025-8_12)
18. Benjamin DI, Louie SM, Mulvihill MM, Kohnz RA, Li DS, Chan LG, et al. Inositol phosphate recycling regulates glycolytic and lipid metabolism that drives cancer aggressiveness. *ACS Chem Biol.* 2014;9: 1340–1350. doi:[10.1021/cb5001907](https://doi.org/10.1021/cb5001907)
19. Zhang J, Feng B. cghMCR: Find chromosome regions showing common gains/losses. Bioconductor; 2020. Available: <https://bioconductor.org/packages/release/bioc/html/cghMCR.html>
20. Seshan VE, Olshen A. DNAcopy: DNA copy number data analysis. Bioconductor; 2020. Available: <https://bioconductor.org/packages/release/bioc/html/DNAcopy.html>
21. Zhang J. CNTTools: Convert segment data into a region by sample matrix to allow for other high level computational analyses. Bioconductor; 2020. Available: <https://bioconductor.org/packages/release/bioc/html/CNTTools.html>
22. Meyer PE. Infotheo: Information-Theoretic Measures. R package version 1.2.0: CRAN; 2014.