

A Workflow Utilizing R/Bioconductor, GSEA, and TIMER 2.0 to explore the role of the Vav Protein Family in Cutaneous Melanoma.

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Melanoma and Vav proteins

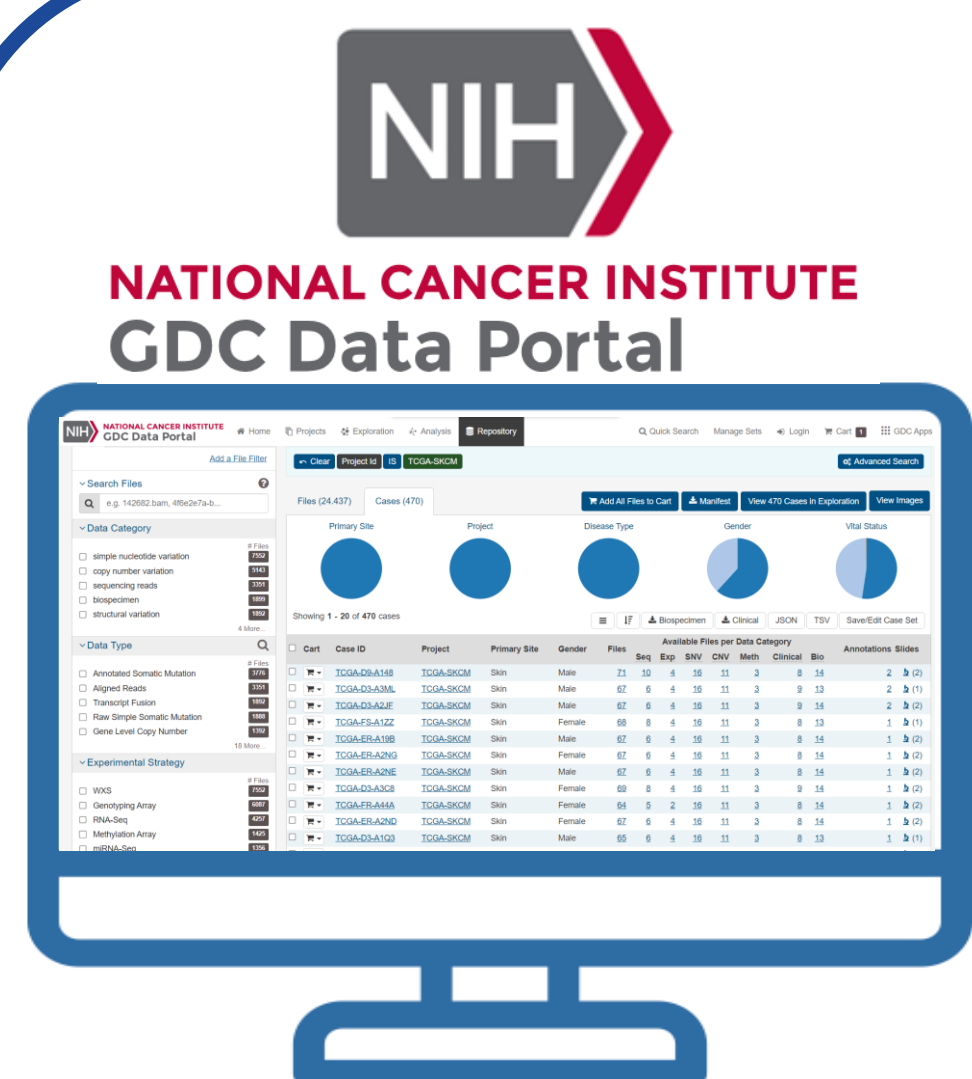
Melanoma represents the most aggressive manifestation of skin cancer, arising from the malignant transformation of cutaneous melanocytes. Its global incidence is rising, and it stands out as one of the most highly metastatic cancer types with limited treatment options available.

Within the intricate landscape of cancer biology, the Vav family of proteins assumes a significant role as activators of Rho GTPases, which are implicated in pro-oncogenic processes when their activity is deregulated.

This family consists of three members, which typically exhibit functional redundancy and are associated with proactive functions in cancer. However, their role in melanoma remains largely unexplored.

Our aim was to establish a systematic approach, utilizing bioinformatic techniques, to investigate the role of each member of the Vav family in melanoma.

Search, Download, and Data Cleaning



```
library("SummarizedExperiment")
library("TCGAbiolinks")
library("RTCGA.clinical")
library("edgeR")
```

```
query.exp <- GDCquery(
  project = "TCGA-SKCM",
  data.category = "Transcriptome Profiling",
  data.type = "Gene Expression Quantification",
  workflow.type = "STAR - Counts")
```

```
GDCdownload(
  query = query.exp,
  files.per.chunk = 100)
```

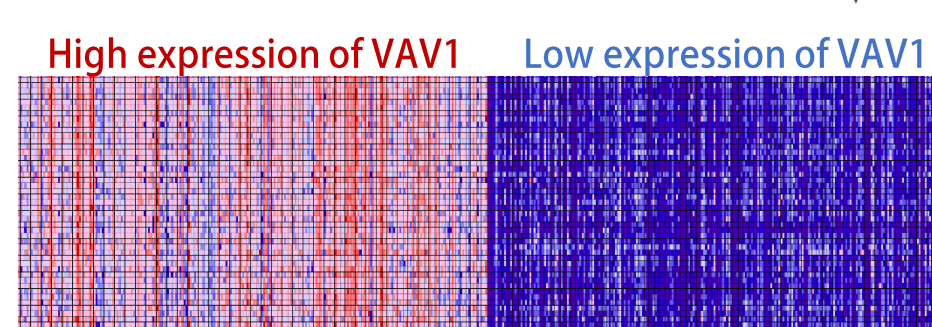
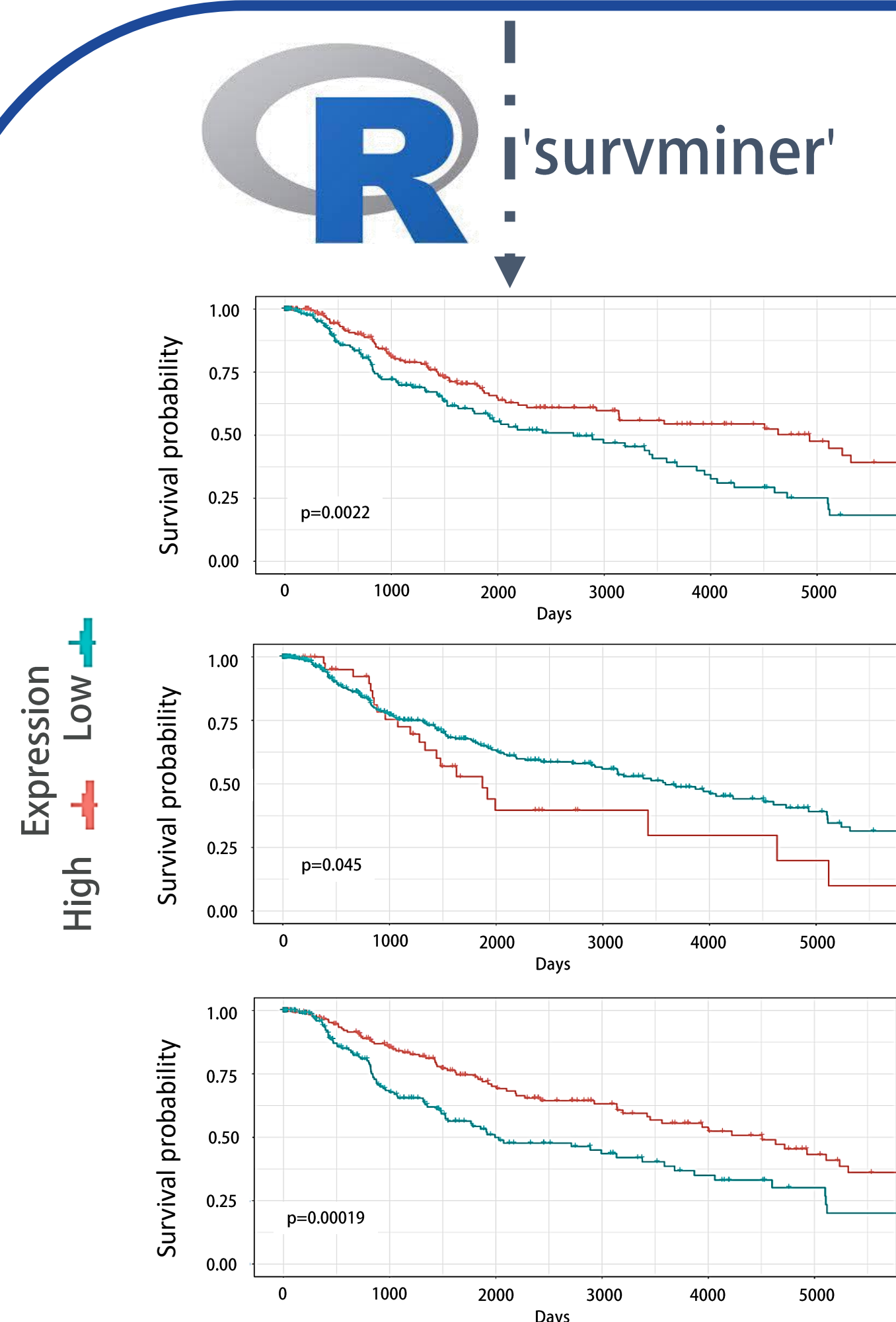
```
skcm.exp <- GDCprepare(
  query = query.exp,
  save = TRUE,
  save.filename = "skcmExp.rda")
```



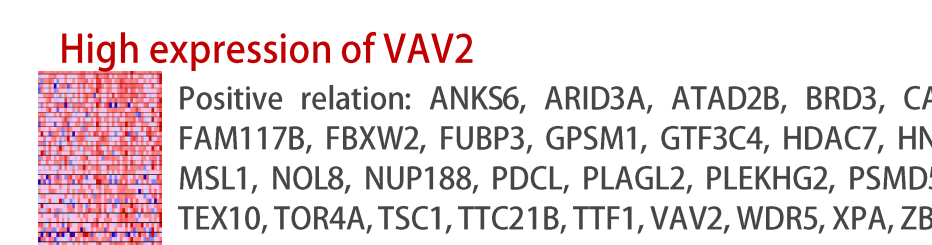
ID	times	status	VAV1	VAV2	VAV3	FenoVAV3	Ft
1	TCGA-3N-A9WB	518	1	0.7584649	15.448733	11.0975395	High
2	TCGA-3N-A9WC	1656	0	26.3088208	24.706554	51.3120911	High
3	TCGA-3N-A9WD	395	1	17.2641888	38.202245	7.7659861	Low
4	TCGA-BF-A1PU	387	0	0.7206094	30.368538	1.7353450	Low
5	TCGA-BF-A1PV	14	0	1.9078738	37.608427	0.897289	Low
6	TCGA-BF-A1PX	282	1	23.3542221	22.089916	30.0374890	High
7	TCGA-BF-A1PZ	12	0	2.9778407	1.980264	1.6973692	Low
8	TCGA-BF-A1QD	17	0	7.1820481	56.883170	4.9397655	Low
9	TCGA-BF-A3DI	464	0	9.6267260	25.806781	5.1711129	Low
10	TCGA-BF-A3DL	28	0	2.1988719	7.789224	0.5404007	Low
11	TCGA-BF-A3DM	14	0	2.4255881	14.399794	5.6027668	Low
12	TCGA-BF-A3DN	32	0	2.8289981	10.858781	13.6877786	High
13	TCGA-BF-A3EO	338	0	6.9953220	13.714625	1.5947239	Low
14	TCGA-BF-A3EP	11	0	1.5407419	26.992980	1.7808576	Low
15	TCGA-BF-A3EQ	12	0	12.6369774	30.269510	4.1859988	Low
16	TCGA-BF-A3ER	12	0	1.1346008	30.890422	0.7320005	Low

Gene expression data from cutaneous melanoma patients were obtained from the Cancer Genome Atlas database. Raw counts were subsequently normalized to counts per million (CPM) using the 'edgeR' package. The patient cohort (n=460) was stratified based on high or low expression levels of Vav1, Vav2, and Vav3.

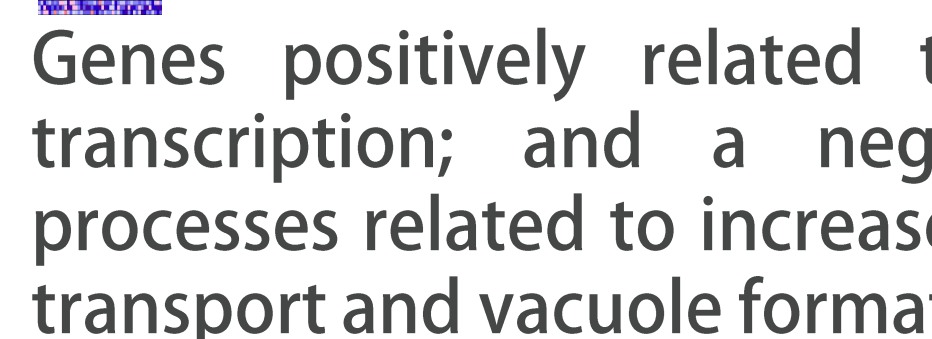
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Genes positively related with the inhibition of tumor metastasis, reduction of cell motility, prevention of damage from inflammatory processes, activation of Ras for remodeling of the cytoskeleton and processes related to immune homeostasis



Genes positively related to division cellular and transcription; and a negative relationship with processes related to increased cellular acidity, proton transport and vacuole formation, were found.



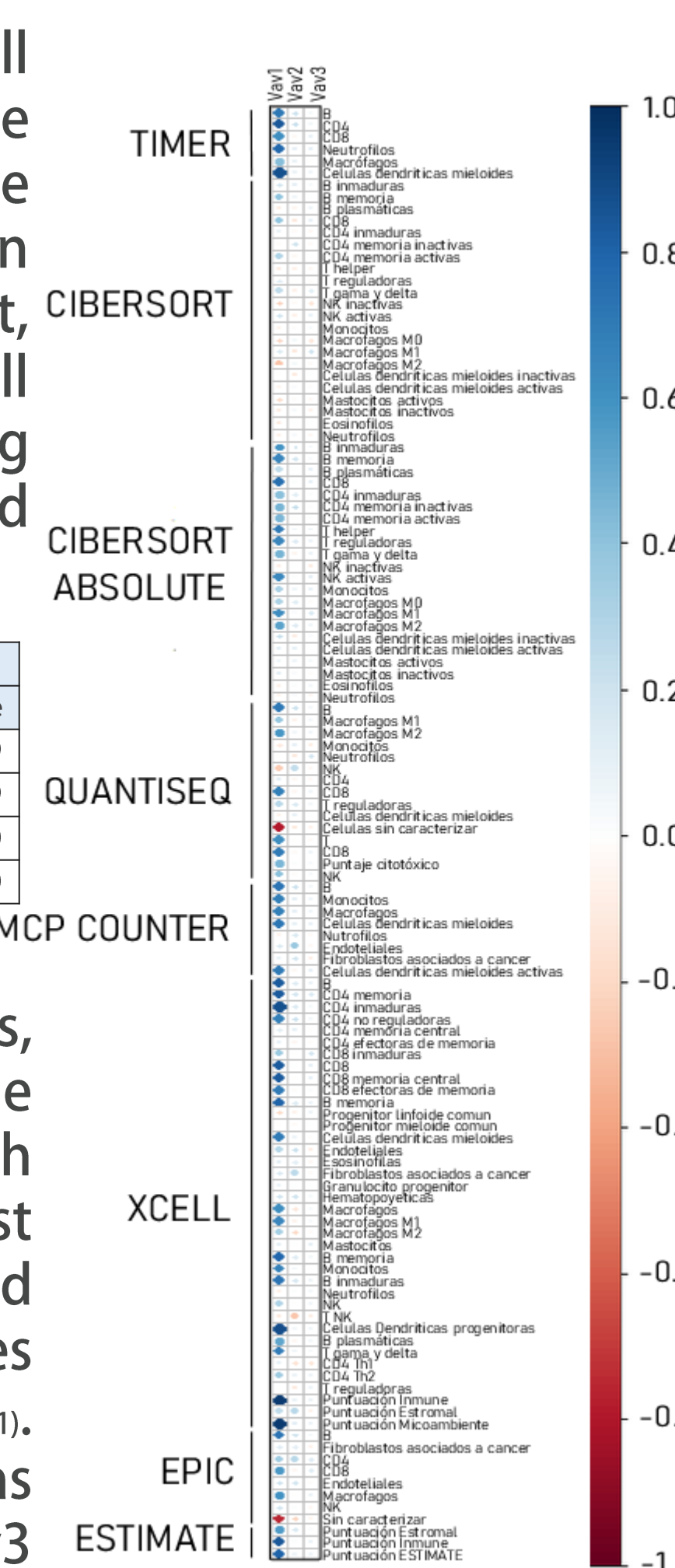
The gene set analysis did not suggest significant and characteristic differences for each phenotype.



To assess immune and stromal cell infiltration in tumor tissues, Immune Score and Microenvironment Score were calculated based on gene expression profiles of the tumor microenvironment, employing the ESTIMATE and xCell algorithms. Both Scores showed a strong and positive association with Vav1 and Vav3 expressions ($p < 0.001$).

Score		VAV1		VAV2		VAV3	
		Score	p value	Score	p value	Score	p value
Immune	xCell	0.923	2.2E-16	-0.035	0.5149	0.278	1.7E-09
	estimate	0.951	2.2E-16	-0.002	0.9696	0.275	2.6E-09
Microenvironment	xCell	0.935	2.2E-16	0.038	0.4229	0.288	3.9E-10
	estimate	0.931	2.2E-16	0.078	0.0937	0.289	2.5E-10

Then, using eight different algorithms, with the 'estimate' package and the TIMER2.0 application, correlation with some cell types was evaluated. A robust positive correlation was identified between Vav1 expression and some types of immune cell signatures ($p < 0.001$). Conversely, no significant correlation was observed between Vav2 or Vav3 expression and cell types.



Conclusions

Our findings suggest that a favorable prognosis in melanoma is linked to elevated expressions of Vav1 and Vav3, coupled with reduced Vav2 expression. This prognosis may arise from Vav1's impact on intercellular communication within the tumor microenvironment, while heightened Vav3 expression could regulate the activation of tumor cell signaling pathways, thereby promoting greater immunogenicity.

Our study presents a comprehensive pipeline that could serve to explore the implications of other proteins in diverse disease contexts.

References

Colaprico et al., 2015 - Gentleman et al., 2004 - Kassambara, 2016 - Mounir et al., 2019 - Robinson & Oshlack, 2010 - Silva et al. 2016 - Wickman & Grolemd, 2018

