

RAPID COMMUNICATION

Data-driven identification of core tumor-secreted factors associated with cachexia prevalence

Secreted components from the tumor microenvironment actively drive the development and progression of cancer cachexia. These cancer cachexia tumor factors (CCTFs) trigger systemic inflammation, metabolic alterations, skeletal muscle wasting, and adipose tissue lipolysis. Many of these factors are members of the cytokine–cytokine receptor interaction (CCRI) pathway, which is particularly enriched in tumor types highly associated with cachexia. For example, pancreatic ductal adenocarcinoma (PDAC) has the highest prevalence of cachexia and exhibits the highest number of overexpressed secreted genes. Our previous pan-cancer study demonstrated that secretory genes (tumor secretome) belonging to the CCRI pathway are up-regulated in tumors highly associated with cachexia.¹ PDAC expressed the highest number of secreted genes from the CCRI pathway. Considering the potential of this pathway to have other CCTFs not previously associated with cachexia, it is crucial to determine a core of genes representative of the pathway in the context of cancer cachexia. Moreover, we expect to distinguish the primary source of CCTF by analyzing tumor transcriptomes (bulk and single-cell). Therefore, we aimed to investigate the secretome genes related to the CCRI pathway in 12 cancer types with different prevalences of cachexia, thereby generating what we called the core of cancer cachexia tumor factors (core-CCTF).

Using a pan-cancer approach, we evaluated secreted CCTF genes in 12 cancer types with varying cachexia prevalence (4651 TCGA samples from BRCA: invasive breast carcinoma; COAD: colon adenocarcinoma; ESCA: esophageal carcinoma; HNSC: head and neck squamous cell carcinoma; LAML: acute myeloid leukemia; LIHC: liver hepatocellular carcinoma; LUAD: lung adenocarcinoma; LUSC lung squamous cell carcinoma; PAAD: pancreatic

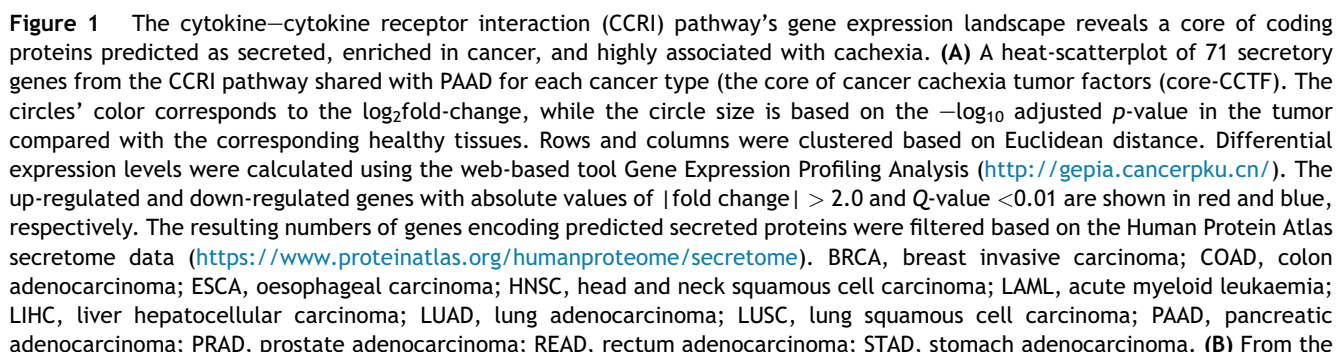
adenocarcinoma; PRAD: prostate adenocarcinoma; READ: rectal adenocarcinoma; STAD: stomach adenocarcinoma) compared with matched normal tissues (2737 GTEx samples). First, we selected 263 genes from the CCRI pathway and predicted as secreted by The Human Protein Atlas. We identified 71 genes up-regulated in PDAC and shared with at least one cancer type. Most of these genes are highly expressed in cachexia-associated cancers and absent/down-regulated in cancer types where cachexia is not prevalent (Fig. 1A). We denominated this gene set as a core-CCTF. Through Pearson correlation-based clustering analysis, we identified a distinct co-expression module comprising CCTF with the highest positive correlations, which also shows increased expression in tumor types with higher cachexia prevalence (Fig. S1), suggesting a coordinated transcriptional program potentially associated with cachexia-related mechanisms. To understand how these molecules interact, we constructed a protein–protein interaction network with 26 core-CCTF genes up-regulated in PAAD, STAD, READ, COAD, ESCA, and HNSC (cancer types highly associated with the syndrome). This indicates a potential use of the described genes as a panel of biomarkers for cachexia risk stratification or potential therapeutic targets. Indeed, C-X-C motif chemokine ligand 8 (CXCL8), also known as interleukin-8 (IL-8), has the highest centrality and interacts with other genes related to the chemokine interleukin-8-like superfamily (Fig. 1B). IL-8 is a tumor-derived factor directly associated with muscle wasting in lung and pancreatic cancers, and its neutralization with blocking antibodies prevented *in vitro* muscle atrophy (refer to the supplementary materials for references).

Since PDAC was the tumor type with significant up-regulation of the 71 core-CCTF genes, we further focused on this tumor type and this gene list for the subsequent analyses. We found that gene amplification is a genomic alteration most found on core-CCTF genes, present in 217

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out of 988 PDAC patients from cBioPortal (Fig. S2A). These patients with alteration in core-CCTF genes have higher scores of tumor hypoxia (Fig. S2B) and lower disease-free survival rates (Fig. S2C; $n = 69$).

Further, we analyzed the public single-cell RNA-sequencing data from pancreatic cancer samples (CRA001160)² to characterize the expression landscape of core-CCTF genes. We found that macrophages, B cells, and T cells were most abundant in cancer tissues and reduced in the normal pancreas (Fig. S3A and S3B). T cells and macrophages were also increased in TCGA PDAC samples compared with GTEx normal pancreas, as demonstrated by the CIBERSORTx deconvolution tool (Fig. S3C). Sixty-nine core-CCTF genes were available in the single-cell analysis. Overall, malignant ductal cells (Ductal2) express 57.9% of core-CCTF genes, followed by macrophages (39.1%), fibroblasts (39.1%), and T cells (27.5%) (Fig. 1C; Fig. S3D). Normal ductal cells (Ductal1) express fewer core-CCTF genes (29%) when compared with the neoplastic cells (Fig. 1C). We verified that the majority of genes were expressed by higher percentages in malignant cells (25%–87%; Fig. S4). *CXCL5* (78.93%), C–C motif chemokine ligand 15 (*CCL15*) (85.54%), and platelet factor 4 (*PF4*) (87.03%) were particularly expressed in malignant ductal cells (Fig. S3D; Fig. S4). Liu et al have shown that macrophages potentiate pancreatic cancer-induced muscle wasting by promoting tumor necrosis factor (TNF)-like weak inducer of apoptosis (TWEAK) secretion from tumor cells, and that depletion of macrophages can reverse tumor-driven muscle degradation. Macrophages activate malignant cells through the CCL5/TNF receptor-associated factor 6 (TRAF6)/nuclear factor-kappa B (NF- κ B) pathway, leading to non-autonomous secretion of TWEAK (refer to the supplementary materials for reference). This evidence supports a model in which malignant epithelial cells and macrophages act cooperatively to sustain a pro-cachectic signaling network, consistent with our finding that both cell types are central sources of core-CCTFs in PDAC. We validated the increased expression of TNF receptor superfamily member 1A (*TNFRSF1A*), *TNFRSF6B*, *TNFRSF12A*, *TNFRSF14*, *TNFRSF25*, *FAS*, Bone morphogenetic protein 4 (*BMP4*), activin A receptor type 1 (*ACVR1*), interferon alpha and beta receptor subunit 1 (*IFNAR1*), interleukin 15 receptor

subunit alpha (*IL15RA*), interleukin 1 receptor accessory protein (*IL1RAP*), interleukin 4 receptor (*IL4R*), oncostatin M receptor (*OSMR*), leukemia inhibitory factor (*LIF*), transforming growth factor beta 1 (*TGFB1*), *TGFB2*, *CXCL1*, *CXCL8*, and *CXCL5* in pancreatic cancer cell lines from the Cancer Cell Line Encyclopedia (Fig. S5). The PDAC cell line PANC0213 presented the most enriched profile for core-CCTF genes (Fig. S5), suggesting that this cell line is the most similar to PDAC cancer cells in terms of the transcriptional profile of CCTF genes.

To infer the crosstalk between tumor cells and tissues affected by cachexia, we used the ligand-receptor consensus list by Ramilowski et al³ to verify the expression of the CCTF receptors in human primary cells from the FANTOM5 project. We observed that adipose cells (mature adipocytes), followed by liver cells (hepatocytes), express the highest number of receptor genes (85.5% and 62%, respectively) for the core-CCTFs expressed in the tumors highly associated with cachexia (Fig. 1D). Neurons and bone cells express the lowest number of core-CCTF receptor genes (31% and 18.3%, respectively). Skeletal muscle cells express 43% of core-CCTF receptor genes. This may indicate that adipose tissue is more responsive to cachexia stimulus than skeletal muscle. A previous study has shown a more pronounced transcriptomic response in the subcutaneous adipose tissue compared with the rectus abdominis muscle in PDAC cachectic patients.⁴ The receptors C–X–C motif chemokine receptor 1 (*CXCR1*), *CXCR2*, *CXCR3*, C–C motif chemokine receptor 5 (*CCR5*), and *CCR1* presented more than three ligands from the core-CCTF list, as demonstrated in the KEGG database (Fig. S6A). In a pancreatic cancer cachexia model (GSE51931),⁵ *Ccr1* and *Ccr5* were up-regulated ($\log_2$ fold-change > 1.15 and p -value < 0.05) in skeletal muscle, white adipose, and liver tissues (Fig. S6B). *Cxcr2* was only significantly overexpressed in the liver of cachectic animals (Fig. S6B).

Overall, we identified a core of secreted factors associated with the CCRI pathway, specifically enriched in tumor types with a high prevalence of cachexia. By analyzing bulk and single-cell data from PDAC samples, the cancer type with the highest risk of inducing cachexia, we found increased immune cellularity (number and diversity) compared with normal samples. We also found that

up-regulated genes identified in (A), protein–protein interactions (PPI) were identified in STRING of secreted proteins coded by 26 up-regulated genes shared by six cancer types highly associated with cachexia (PAAD, ESCA, STAD, HNSC, COAD, and READ). Edge thickness indicates confidence. Interacting proteins from the CCRI pathway are represented in blue and correspond to a gene set of 26 out of the 263 genes constituting this pathway. We also observed a central subcluster in this PPI pathway enriched by 16 proteins of the chemokine interleukin-8-like superfamily pathway, represented in green. Only proteins significant in at least four tissue sites are represented in the PPI pathway. PPI was created using STRING tool v.11 (<https://string-db.org/>) and Cytoscape (v3.7.2). The arrow indicates the up-regulated genes used to generate the PPI. (C) Dotplot with expression values of core-CCTF genes from (A) in 57,530 pancreatic cells using single-cell RNA sequencing data (GSA: CRA001160) from PDAC and normal-like pancreatic tissues ($n = 27$ and $n = 14$, respectively). (D) Inference of the crosstalk between tumor cells and tissue cells affected by cachexia using core-CCTF gene expression data. Left: Alluvial diagram connecting the ligands from the core of factors (CCTF) into its matched receptors (data from Ramilowski et al). Right: Heat map of the expression levels (z-score normalized Transcripts Per Million - TPM) of predicted receptors in human primary cell lineages, previously described by Ramilowski et al³ using the FANTOM5 project. Bottom: The scatter plot showing the percentage of core-CCTF receptor genes expressed in each tissue. Muscle: 31 out of 71 receptors (43.7%); Adipose tissue: 60 out of 71 receptors (85.5%); Liver: 44 out of 71 receptors (62%); Brain: 22 out of 71 receptors (31%); Bone: 13 out of 71 receptors (18.3%). Adipocyte1, breast adipocyte; Adipocyte2, omental adipocyte; Adipocyte3, perirenal adipocyte; Preadipocyte1, breast preadipocyte; Preadipocyte2, omental preadipocyte; Preadipocyte3, perirenal preadipocyte; Preadipocyte4, subcutaneous preadipocyte; Preadipocyte5, visceral preadipocyte.

malignant ductal cells were responsible for the expression of most genes belonging to the core, followed by stromal and immune cells. Moreover, we found that the core-CCTF genes coded proteins able to interact with muscle cells, hepatocytes, and adipocytes via specific receptors expressed in these cells, especially in mature adipocytes. These results show novel potential pro-cachectic mediators that can trigger wasting in the targeted tissues most affected in cancer cachexia.

CRediT authorship contribution statement

Sarah Santiloni Cury: Writing – review & editing, Writing – original draft, Validation, Methodology, Investigation, Formal analysis, Data curation. **Paula Paccielli Freire:** Writing – review & editing, Visualization, Methodology, Investigation, Formal analysis, Data curation. **Robson Francisco Carvalho:** Writing – review & editing, Writing – original draft, Supervision, Resources, Project administration, Methodology, Investigation, Funding acquisition, Conceptualization.

Data availability

The results shown here are whole based upon data generated by the TCGA Research Network (<http://cancergenome.nih.gov/>) and by the Genotype-Tissue Expression project (GTEx) (<https://gtexportal.org/>). The results are also partly based on publicly available microarray data from the Gene Expression Omnibus (GEO) and single-cell RNA-Seq dataset from the Genome Sequence Archive (GSA): GSE51931 and CRA001160, respectively.

Declaration of generative AI and AI-assisted technologies in the writing process

During the preparation of this work the author(s) used Grammarly and ChatGPT 4 in order to improve language and readability. After using this tool/service, the author(s) reviewed and edited the content as needed and take(s) full responsibility for the content of the publication.

Conflict of interests

The authors have read the journal's authorship agreement and declare no conflict of interests.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.gendis.2025.101883>.

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