### **Breakout Session 5:**

# Cloud-Based Machine Learning and Biomarker Visual Analytics for Salivary Proteomics

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"Cloud-Based Machine Learning and Biomarker Visual Analytics for Salivary Proteomics"

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Supplement grant # 3R01DE016937-18S1 (Parent grant: R01DE016937) Date: 01-18-23





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Advanced Search



Parotid glands (592 proteins) Whole saliva (2643 proteins)

- share your mass spec data
- annotate the proteins
- give your thoughts in discussions
- tell us how we can improve the site



**HSP WIKI Grant Aims:** 

Aim 1. Transfer the HSP Wiki to a cloud-based server, maintain and develop the database computing services to provide the research community with a reliable, curated, and constantly evolving (up-to-date) data platform for evaluating salivary proteomics datasets.

Aim 2. Implement a novel computational infrastructure for salivary proteomic datasets that includes visual analytics, faster processing, and efficient measures of success.

### Aim 3. Expand the HSP Wiki database to include bacterial and phage proteins.

The long-term goal of this project is to establish the resources and develop new tools to facilitate salivary research for both scientific discovery and diagnostic applications.



## Human Salivary Proteome Wiki 2.0





# **ANALYSIS PAGE**

Human Salivary Proteome

HOME BROWSE - SEARCH - ANALYZE - HELP -

### ANALYSIS

Analysis will identify proteins with differential abundance between experiments in Groups A and B based on their normalized spectral counts.

#### MULTIPLE SEQUENCE ALIGNMENT

ClustalW is a general purpose multiple sequence alignment program for DNA or proteins.

#### DIFFERENTIAL EXPRESSION

Shows the difference in protein abundance between samples in Groups A and B.

#### PROTEIN SIGNATURE SEARCH

Finds regions of sequence similarity, which will yield functional and evolutionary clues about the structure and function of your novel sequence.

#### PROTEIN SIMILARITY SEARCH

InterProScan is a tool that combines different protein signature recognition methods into one resource.



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Forsyth
Image: Construction of the university of the u



**Cloud Grant Aims:** 

Aim 1: Develop a novel proteomics data analytics platform to support explainable and optimal machine learning identification of salivary proteomic biomarkers through visual analytics and parallel computing.

Aim 2 : Develop a scalable and on-demand cloud pipeline to identify protein modification, protein structure prediction using AlphaFold2, and 3D models comparison tool using MolStar.



Specific Aim 1: Develop a novel proteomics data analytics platform to support explainable and optimal machine learning identification of salivary proteomic biomarkers through visual analytics and parallel computing.





## Goals





## **Preliminary Data**

P-value_smaller_than0.01	Gene Name
P06331	IGHV4-34
Q9Y3D6	FIS1 TTC11 CGI-135
P31949	S100A11 MLN70 S100C
P62937	PPIA CYPA
P26599	PTBP1 PTB
P02533	KRT14
P05109	S100A8 CAGA CFAG MRP8
P07737	PFN1
P13646	KRT13
P29401	TKT
P04083	ANXA1 ANX1 LPC1
A0A0B4J1X5	IGHV3-74
P28799	GRN
P61769	B2M CDABP0092 HDCMA22P
O60235	TMPRSS11D HAT
P23528	CFL1 CFL
P01624	IGKV3-15
P01037	CST1
P06702	S100A9 CAGB CFAG MRP14
P05164	MPO
P52209	PGD PGDH
P07195	LDHB



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#### High Serum Progranulin Levels in COVID-19 Patients: A Pilot Study

Inflammation Research (2022) 71:369–376 https://doi.org/10.1007/s00011-022-01545-7

Inflammation Research

**ORIGINAL RESEARCH ARTICLE** 

Check for updates

The prognostic value of S100A calcium binding protein family members in predicting severe forms of COVID-19

Rank sum test p-value < 0.01



### Infrastructure





Specific Aim 2: Develop a scalable and on-demand cloud pipeline to identify protein modification, protein structure prediction using AlphaFold2, and 3D models comparison tool using MolStar.





# **Tools Needed**

#### **Dimensionality Reduction**

 Principal component analysis (PCA)

#### **Protein Clustering**

- CD-HIT
- MMSEQ2
- kClust

#### **Differential Abundance**

- LIMMA
- Fold Change analysis Based on statistical Tests.

#### Classification

Random Forest (RF)

#### Heatmap

 pheatmap (K-means clustering)

#### Statistical Tests

- t-test (or ANOVA)
- Wilcoxon Rank Sum Test
- Wilcoxon Signed Rank Test

#### Pathway Enrichment

- TopGO
- KEGG Pathway (KEGGREST)
- GOANA and KEGGA (LIMMA)

#### Network Analysis

- Cytoscape (RCyc3)
- Ggraph, igraph
- STRING (database)



# **Future Insights: Host-Microbial Proteins**

Genes encoding the protein can be displayed in HOMD Genome Browser, with link back to WiKi





Overview of 16,745 predicted prophagelike entities in HOMD bacterial genomes (HOMD v0.1 draft). Nodes represent predicted prophage genomes; selected phages are colored by the host in which they reside (e.g. *Streptococcus*: green; *Tannerella*: orange; *Prevotella*: purple; *Aggregatibacter*: red; *Fusobacterium*: blue).

From: Dr. Kauffmm (U at Buffalo)



From: Drs. Jessica Mark Welch and George Chen (Forsyth Institute)

# **Open Source Visual Analytics (NCBI)**



Gene Expression Omnibus (GEO) Search. Download. Visualize.

Search, Download, and Visualize Human RNA-Seq Gene Expression Data in NCBI's Gene Expression Omnibus (GEO)



*Figure 1: Screenshot of GEO2R differential gene expression analysis results, including Volcano, Mean difference, Mean variance, UMAP, Venn, Boxplot, and Histogram plots.* 



Source: NCBI

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### Questions



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"Enhancing human immunity to impact global disparities"