Breakout Session 8: Track A

Applying Gerchberg-Saxton Algorithm on Biomedical Data to Mitigate Sampling Bias on Under-Represented Populations

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Developing Unbiased AI/Deep Learning Pipelines to Address Lung Cancer Health Disparities Research

Applying Gerchberg-Saxton Algorithm on Biomedical Data to Mitigate Sampling Bias on Underrepresented Populations

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Agenda

- 1. Project Motivation
- 2. Gerchberg-Saxton Algorithm
 - Previous Studies and Results
- 3. Project Plan
- 4. Current Phase and Initial Results
- 5. Future Directions and Expected Outcomes

Project Motivation - Challenges in Medical Al

- PHI data regulations poses challenges on data distributions.
- Upholding data integrity and privacy is paramount yet challenging.
- There is a noticeable scarcity in the availability of public medical datasets.



Project Motivation - Data Quality and Representation

- The imperative to improve the quality of data generated by individual institutions is clear.
- The distribution among population groups in the US is uneven, which is mirrored in medical data, potentially biasing Al model predictions.
- Existing Al-based solutions, such as synthetic data generation, face challenges like **inadequate source data quality or quantity**.

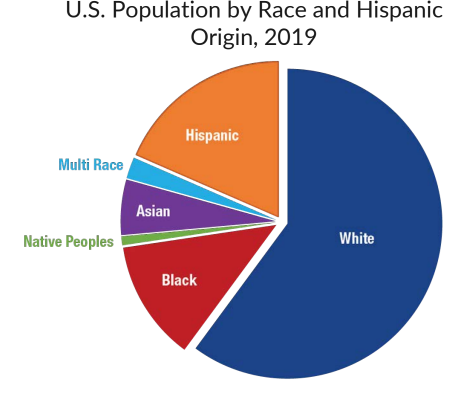


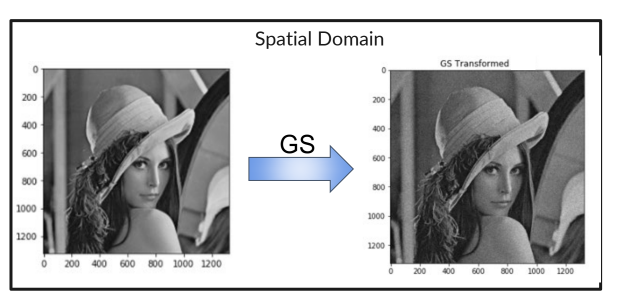
Image Source: U.S Census Bureau, Population Estimates,

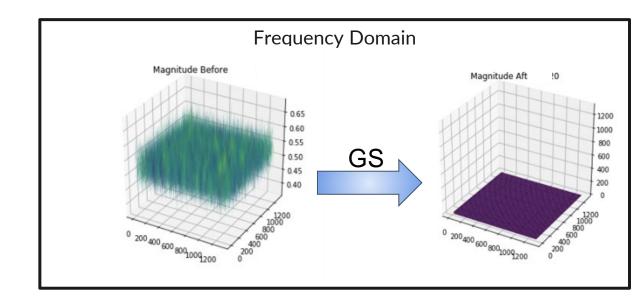
Project Motivation - Innovative Solution

- We propose an innovative data transformation approach utilizing the Gerchberg-Saxton algorithm to address data quality and representation issues.
- This algorithm transforms data in the frequency domain, balancing intensity components while preserving phase information, thereby enhancing data uniformity.
- The transformed data will more uniformly represent each population group, aiming for fairer results in machine learning applications.

Introduction to Gerchberg-Saxton

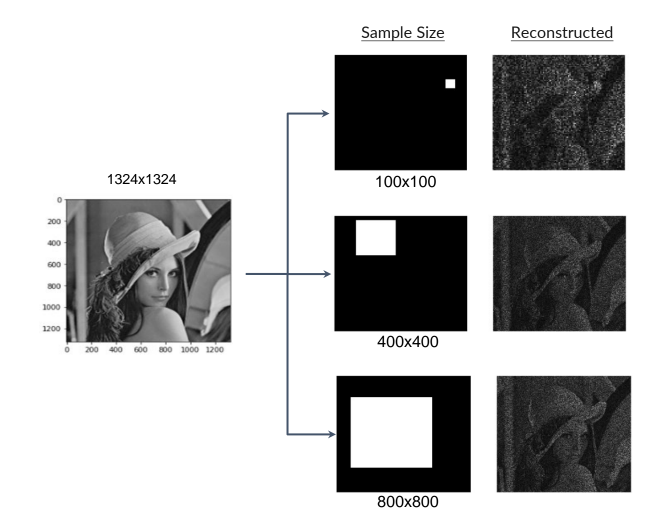
- The Gerchberg-Saxton algorithm is a cornerstone in holographic imaging, enabling the creation of holographic representations of images in digital environments.
- We leverage the Gerchberg-Saxton algorithm to utilize two crucial characteristics of holographic images to enhance medical data analysis.
 - Holographic Divisibility
 - Information Distribution





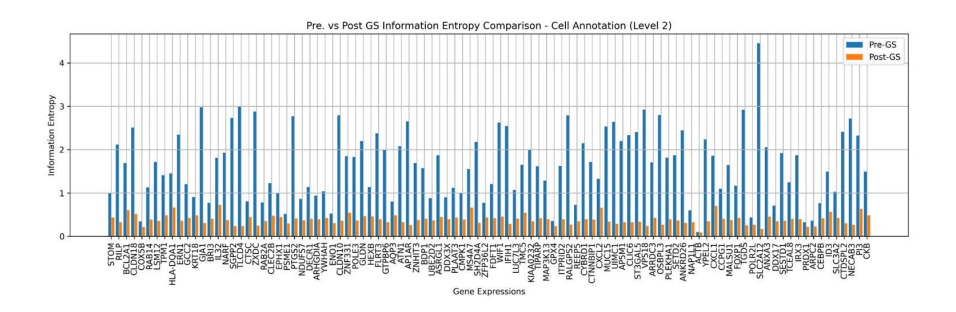
Holographic Divisibility

- Each data sample encapsulates information representative of the entire dataset, echoing the holistic nature of holograms.
 - o Small Sample, Whole Dataset
- Equitable insights and analysis across diverse patient data.



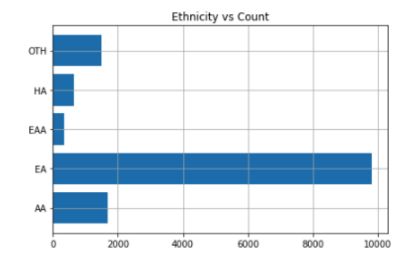
Uniform Information Distribution

- The Gerchberg-Saxton algorithm works in the frequency domain, **balancing intensity components** and preserving phase information during data transformation.
- Such transformation ensures a more uniform distribution of information across datasets.
- This uniformity is critical for obtaining fairer results from machine learning models and simplifying the identification of feature model contributions.



Previous Study - Mortality Prediction

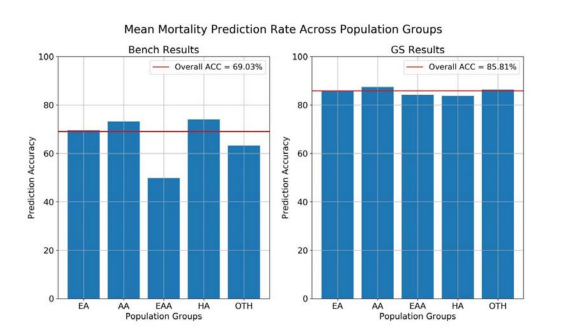
- Study Focus
 - Evaluation of mortality rates among ICU patients, with a particular emphasis on detecting and correcting bias across diverse population groups.
- Database Selection
 - o The MIMIC-III database was chosen for its comprehensive data
 - Unbalanced population distribution across different racial groups.

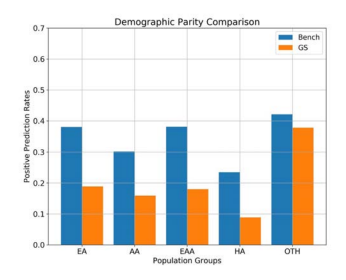


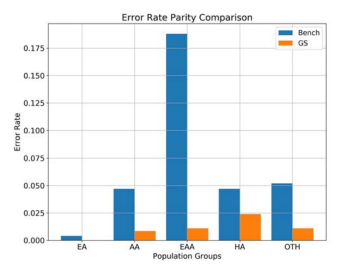


Previous Study - Results

- Implementation of GS transformations on the dataset **significantly reduced the bias**, enhancing model predictions across different population groups.
- The improvements were quantitatively supported by demographic parity and error rate parity, demonstrating more uniform model prediction rates across demographics.





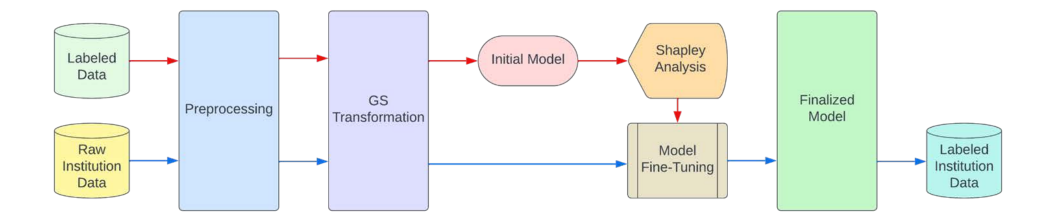


Project Objectives and Approach

- Goal: Enable fair data representations for single-cell cancer studies, improving healthcare decision-making.
- **Strategy:** Develop a pipeline to transform sc-RNA sequencing data into an AI-ready format, ensuring fair and uniform data representation through GS transformations.
 - Data Preprocessing and GS transformation
 - Model Development
 - Testing and Implementation

Pipeline

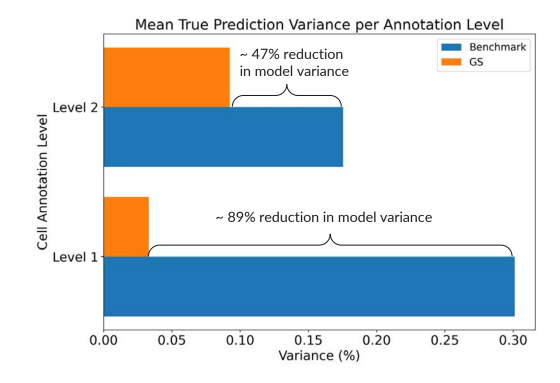
- Raw data **preprocessing** and **GS transformation** for fairness.
- Train models, then refine using **Shapley analysis** to identify and retain only **features that significantly contribute to model performance**.
- Validate model with labeled public data subsets.
- Apply to institutional data for enhanced decision-making in healthcare.



Initial Results - Model Prediction Uniformity

- Human Lung Cell Atlas, featuring **over 2 million single cell profiles from lung tissue**, with detailed cell annotations.
- Selected 20,000 cells **across diverse populations**, focusing on annotation levels 1 **(39 cell types)** and 2 **(61 cell types)**.

	Pre-GS Accuracy	Post-GS Accuracy
Level 1 (39 cell types)	90.62	89.48
Level 2 (61 cell types)	89.67 %	88.72 %



Future Directions and Expected Outcomes

- Pipeline Enhancement: Integrating a secondary model for tumor stage prediction to distinguish between normal and cancer cells, facilitating tumor microenvironment analysis.
- **Comprehensive Analysis:** Combination of models will **improve understanding** of gene regulatory networks, cell-to-cell interactions, and therapeutic pathways.
- Impact on Healthcare: Aims for enhanced, equitable decision-making in healthcare through better insights into cancer biology and treatment pathways.

Thank you

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