

# From data disparity to data harmony: A comprehensive pan-cancer omic data collection

Abstract 6209

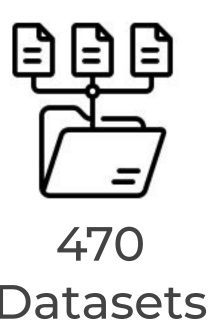
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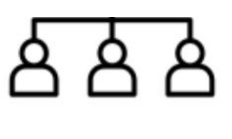
### INTRODUCTION

- The exponential growth of omics datasets offers a significant opportunity for scientific advancement in cancer research.
- However, though the lack of uniform standards, in both clinical and omic data, hinder the effective utilization of these datasets, thus impeding our understanding of cancer biology and the development of innovative therapies.
- We have created a novel collection of pan-cancer datasets with extensive clinical data harmonization and consistent omic data normalization.
- This approach enhances data quality, and is also cost-effective, offering significant advantages in the realm of cancer research.

Here, we focused on patient-derived gene expression microarray datasets from the Gene Expression Omnibus<sup>1</sup> (GEO) database.

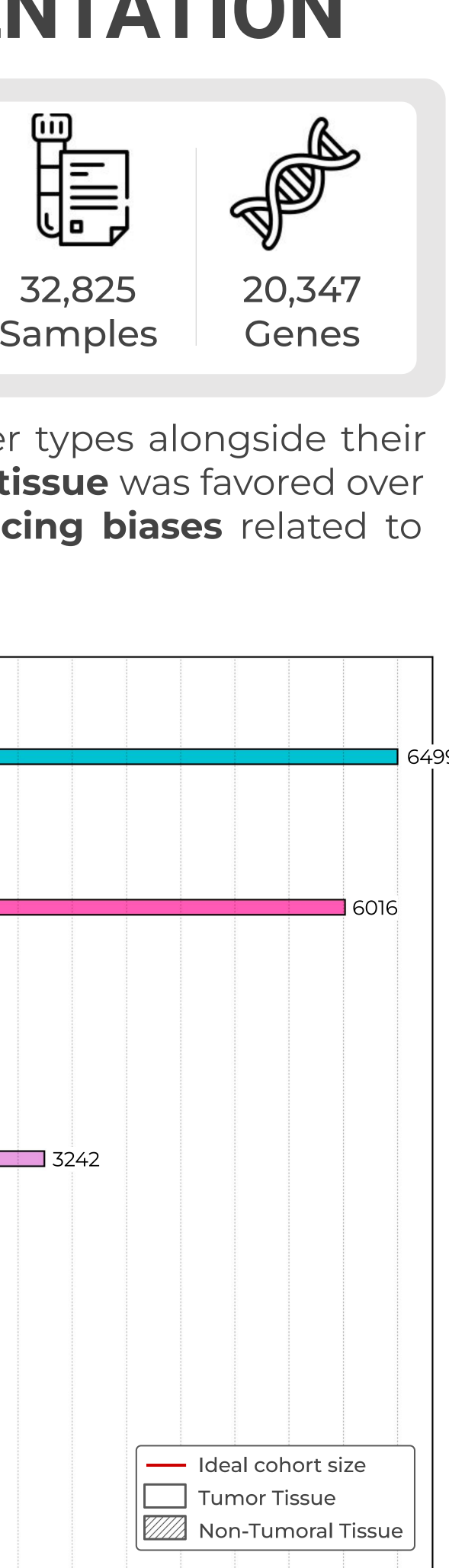
## **DATA COLLECTION PRESENTATION**











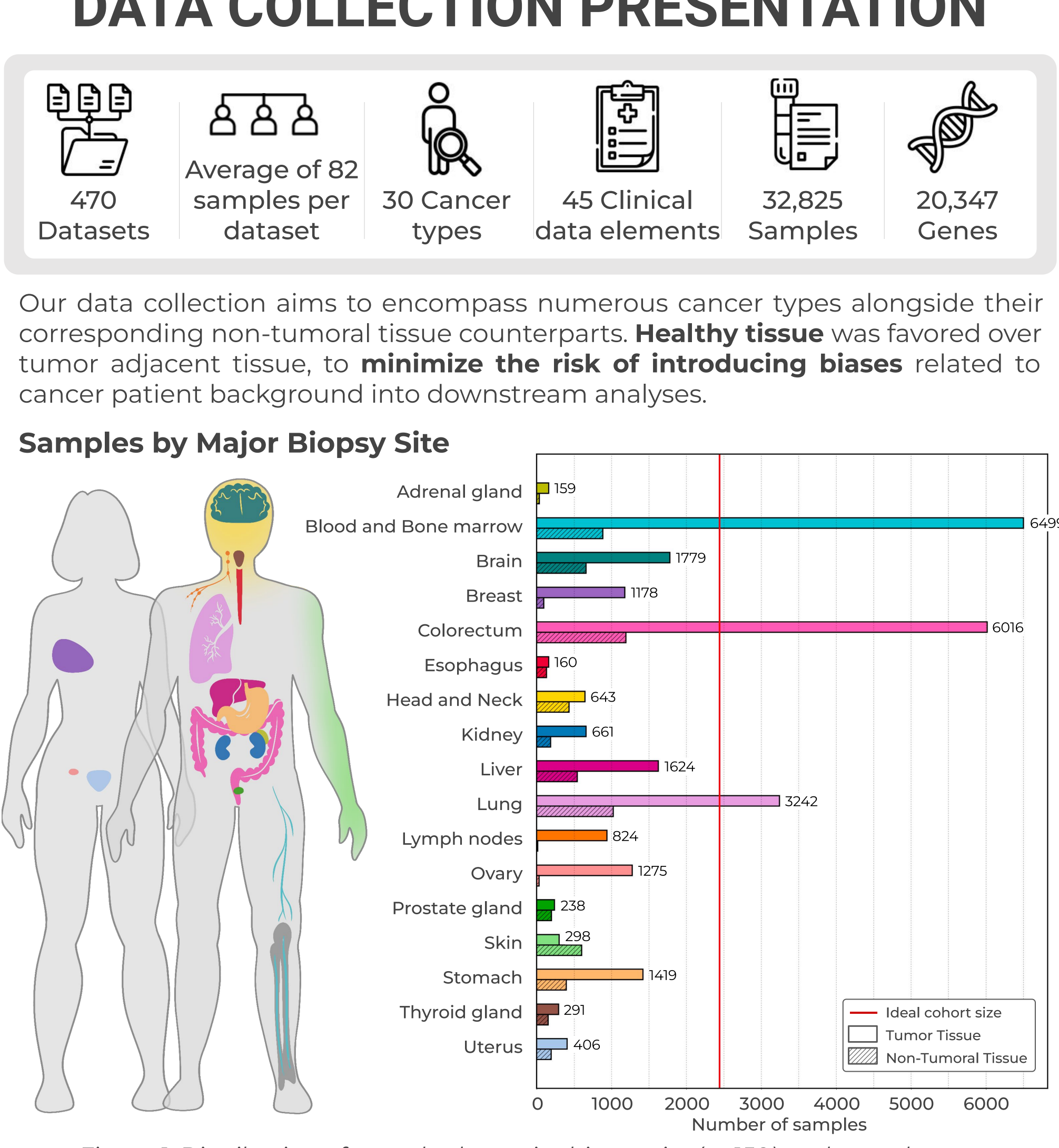
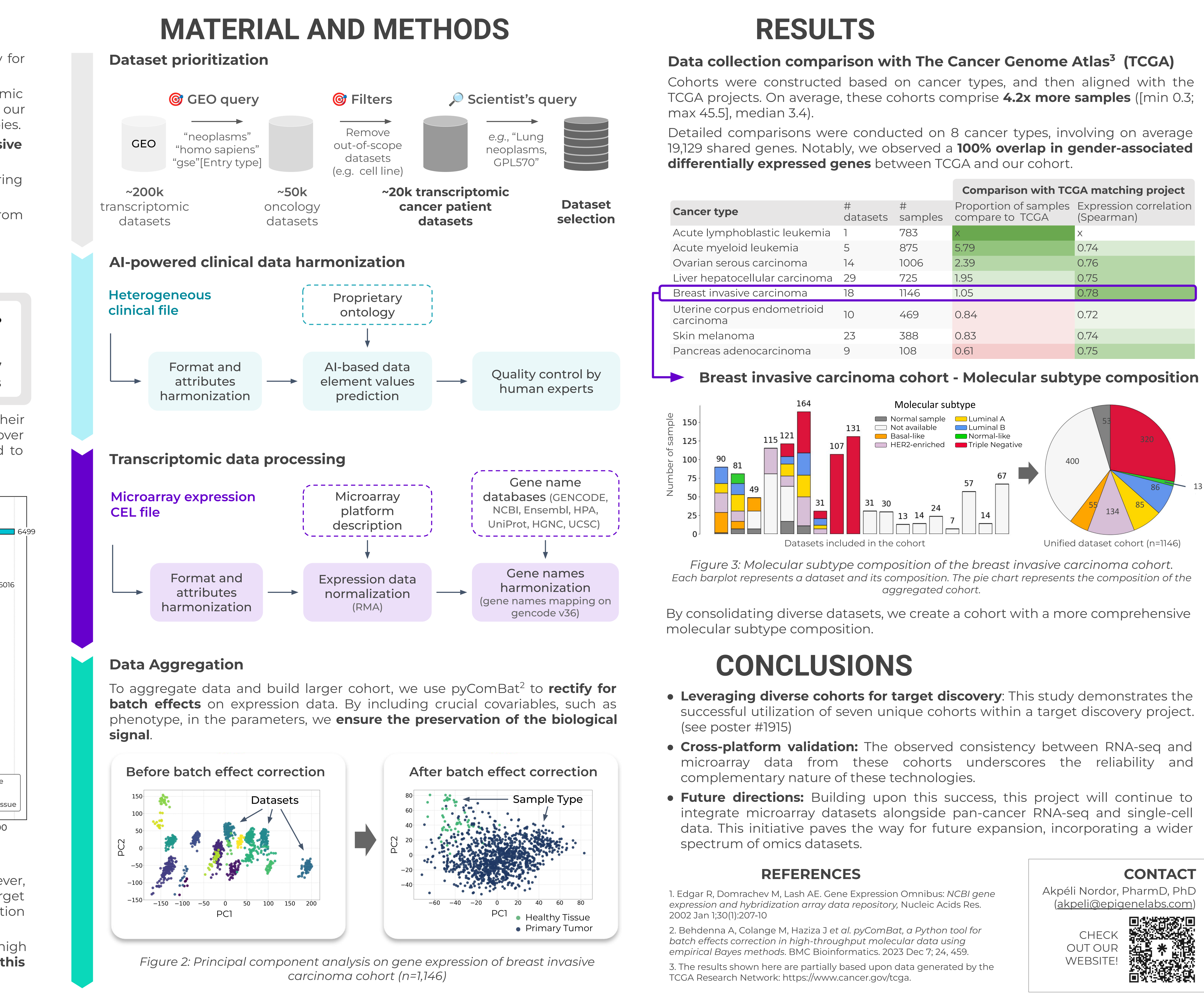


Figure 1: Distribution of samples by major biopsy site (n>150) and sample type

On average, GEO individual datasets typically hold around 60 samples. However, by adjusting the expected outcome of a Kolmogorov-Smirnov test to a target p-value of 5%, we estimated the ideal cohort size to study sub-population composition (theoretically set at 5) to be 2,441 samples (Fig. 1).

Surpassing the size of popular databases, 3 biopsy sites in our data meet the high cohort size limit. With ongoing data integration, we anticipate surpassing this limit for various biopsy sites, enhancing the robustness of our analyses.







|          |               |              | Comparison with TCGA matching project |                                      |
|----------|---------------|--------------|---------------------------------------|--------------------------------------|
|          | #<br>datasets | #<br>samples | Proportion of samples compare to TCGA | Expression correlation<br>(Spearman) |
| eukemia  | 1             | 783          | X                                     | X                                    |
| nia      | 5             | 875          | 5.79                                  | 0.74                                 |
| oma      | 14            | 1006         | 2.39                                  | 0.76                                 |
| arcinoma | 29            | 725          | 1.95                                  | 0.75                                 |
| oma      | 18            | 1146         | 1.05                                  | 0.78                                 |
| netrioid | 10            | 469          | 0.84                                  | 0.72                                 |
|          | 23            | 388          | 0.83                                  | 0.74                                 |
| oma      | 9             | 108          | 0.61                                  | 0.75                                 |

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