equations

Brian M. Schilder

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Tutorials

- https://lpembleton.rbind.io/posts/annotate-equations/
- https://github.com/st--/annotate-equations/blob/main/annotate-equations.pdf

Install latex

Only need to do this once. Instructions using homebrew.

brew install texlive

Methods

Human Phenotype Ontology

Let us denote:

- D as the set of d diseases.
- p as a phenotype.
- g as a gene.

The final evidence-weighted gene x phenotype matrix ${\cal M}_g, p$ can be expressed as:

$$M_{g,p} = \frac{\sum_{d \in D} \mathbf{R}(g,p,d) \times \mathbf{E}(g,d)}{\sum_{d \in D} \mathbf{R}(g,p,d)}$$

(1=relationship, 0=no relationship)

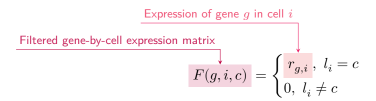
Single-cell transcriptomic atlases

Let us denote:

- g as a gene.
- c as a cell type.
- i as a single cell.

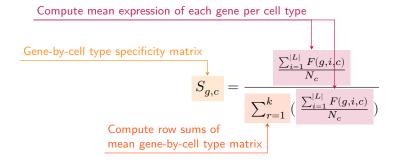
Genes with very no expression across any cell types were considered to be uninformative and were therefore removed before computing the specificity matrix.

$$F(g,i,c) = \begin{cases} r_{g,i}, \ l_i = c \\ 0, \ l_i \neq c \end{cases}$$

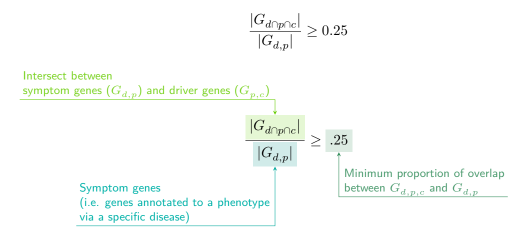


The gene expression specificity matrix construction can be defined as the following:

$$S_{g,c} = \frac{\frac{\sum_{i=1}^{|L|} F(g,i,c)}{N_c}}{\sum_{r=1}^{k} (\frac{\sum_{i=1}^{|L|} F(g,i,c)}{N_c})}$$

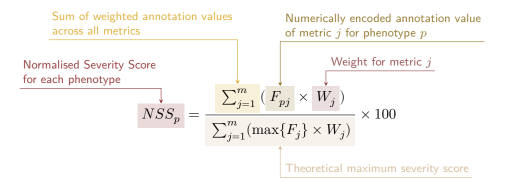


Symptom-cell type associations



Annotation of phenotypes using generative large language models

$$NSS_p = \frac{\sum_{j=1}^m (F_{pj} \times W_j)}{\sum_{j=1}^m (\max\{F_j\} \times W_j)} \times 100$$



Session Info

```
utils::sessionInfo()
R version 4.3.1 (2023-06-16)
Platform: aarch64-apple-darwin20 (64-bit)
Running under: macOS Sonoma 14.3.1
Matrix products: default
       /Library/Frameworks/R.framework/Versions/4.3-arm64/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/4.3-arm64/Resources/lib/libRlapack.dylib;
locale:
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
time zone: Europe/London
tzcode source: internal
attached base packages:
[1] stats
              graphics grDevices utils
                                            datasets methods
                                                                base
loaded via a namespace (and not attached):
 [1] compiler_4.3.1
                       fastmap_1.1.1
                                         cli_3.6.2
                                                           tools_4.3.1
                                                           rmarkdown_2.25
 [5] htmltools_0.5.7
                       rstudioapi_0.15.0 yaml_2.3.8
 [9] knitr_1.45
                       jsonlite_1.8.8
                                         xfun_0.42
                                                           digest_0.6.34
[13] rlang_1.1.3
                       evaluate_0.23
```