

Data exploratory analysis

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| ## ## | The following object is masked _byGlobalEnv: | |
| ## | | |

Project: Comprehensive Omics Catalogue for Hartwell

St. Jude Children's Research Hospital Bio
Hackathon Team ${\bf 1}$

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1 Information about this notebook

This is an exploratory analysis of the data availability in terms of assays in the Comprehensive Omics Catalogue for Hartwell. This is critical for mitigating duplicate sequencing requests and efforts through Hartwell. This notebook aims to showcase: (1) which samples have already been sequenced by Hartwell, and (2) what omics data are available per sample.

For demo purposes, we use dummy data cohort and subset by human brain tumor samples. Finally, we investigate the number of samples per cancer_type_brain and Assay.

2 Set up

```
suppressPackageStartupMessages({
  library(tidyverse)
})
```

3 Directories and paths to file Inputs/Outputs

```
attach(params)
## The following object is masked _by_ .GlobalEnv:
##
##
       root_dir
analysis_dir <- file.path(root_dir, "analyses", "data-exploratory-analysis")
input_dir <- file.path(analysis_dir, "input")</pre>
# We will first read in metadata file as we need to define sample_name
metadata_file <- file.path(input_dir, input_file) # metadata input file</pre>
palette_file <- file.path(root_dir, "figures", "palettes", "assay_color_palette.tsv")</pre>
tumor_palette_file <- file.path(root_dir, "figures", "palettes", "tumor_color_palette.tsv")</pre>
# File path to `plots` directory
plots_dir <- file.path(analysis_dir, "plots")</pre>
if (!dir.exists(plots_dir)) {
  dir.create(plots_dir)}
figures_plots_dir <- file.path(plots_dir, "figures")</pre>
if (!dir.exists(figures_plots_dir)) {
  dir.create(figures_plots_dir)}
source(paste0(analysis_dir, "/util/function-create-barplot.R"))
source(paste0(root_dir, "/figures/scripts/theme_plot.R"))
```

4 Read metadata file

We will subset by human brain tumor samples.

```
# Read metadata
df <- read.csv(metadata_file, stringsAsFactors=FALSE)
# Number of samples per cancer_type_brain</pre>
```

4.1 Color palette for plotting

```
# Read color palette
palette_df <- readr::read_tsv(palette_file, guess_max = 100000, show_col_types = FALSE)

# Define and order palette
palette <- palette_df$hex_codes
names(palette) <- palette_df$color_names

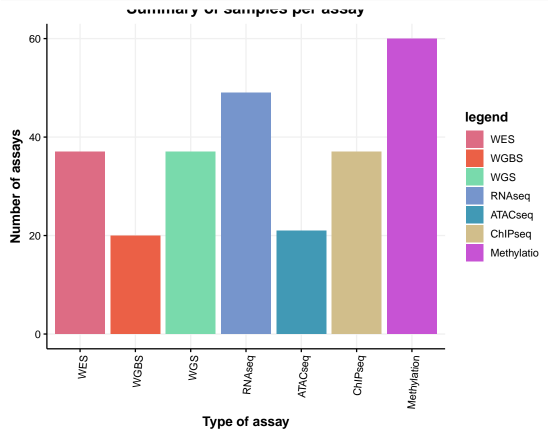
# Read color palette for tumor type
tumor_palette_df <- readr::read_tsv(tumor_palette_file, guess_max = 100000, show_col_types = FALSE)

# Define and order palette
tumor_palette <- tumor_palette_df$hex_codes
names(tumor_palette) <- tumor_palette_df$color_names</pre>
```

5 Number of samples with assay information

Table 1: Summary of samples per assay

| Assay | n |
|-------------|----|
| WES | 37 |
| WGBS | 20 |
| WGS | 37 |
| RNAseq | 49 |
| ATACseq | 21 |
| ChIPseq | 37 |
| Methylation | 60 |
| | |



```
pdf(file = fname, width = 6, height = 5)
print(p)
dev.off()
```

pdf ## 2

6 Number of samples per brain cancer type and assay

6.1 Overall assays

There are 60 brain tumor samples with 261 assays in total.

Table 2: Summary of samples and assays per brain cancer type

| cancer_type_brain | n |
|-------------------|-----|
| Ependymoma | 120 |
| HGG | 28 |
| LGG | 102 |
| Medulloblastoma | 11 |

ounniary or samples and assays per brain cancer type 125 100 Number of assays legend 75 Ependymoma HGG LGG 50 Medulloblastom: 25 0 Ependymoma • .997 **Brain tumor type**

```
pdf(file = fname, width = 6, height = 5)
print(p)
dev.off()
```

pdf ## 2

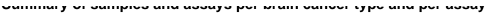
6.2 Per assay

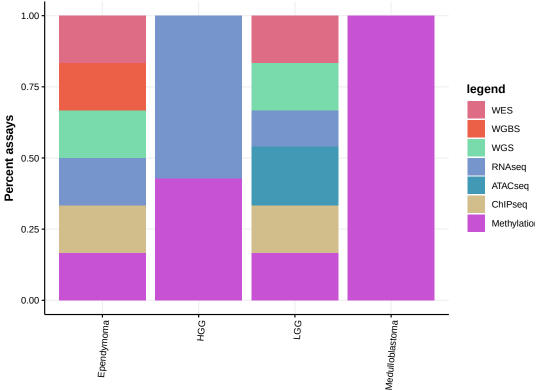
Table 3: Summary of samples and assays per brain cancer type and per assay

| cancer_type_brain | WES | WGBS | WGS | RNAseq | ChIPseq | Methylation | ATACseq |
|-------------------|-----|------|-----|--------|---------|-------------|---------|
| Ependymoma | 20 | 20 | 20 | 20 | 20 | 20 | 0 |

| cancer_type_brain | WES | WGBS | WGS | RNAseq | ChIPseq | Methylation | ATACseq |
|-------------------|-----|------|-----|--------|---------|-------------|---------|
| HGG | 0 | 0 | 0 | 16 | 0 | 12 | 0 |
| LGG | 17 | 0 | 17 | 13 | 17 | 17 | 21 |
| Medulloblastoma | 0 | 0 | 0 | 0 | 0 | 11 | 0 |

```
tables1 <- df %>% count(cancer_type_brain, Assay) %>%
  as.data.frame() %>%
  mutate_all(funs(replace_na(.,0)))
## Warning: `funs()` was deprecated in dplyr 0.8.0.
## i Please use a list of either functions or lambdas:
##
## # Simple named list: list(mean = mean, median = median)
##
## # Auto named with `tibble::lst()`: tibble::lst(mean, median)
##
## # Using lambdas list(~ mean(., trim = .2), ~ median(., na.rm = TRUE))
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was generated.
# Plot stacked barplot
fname <- paste0(figures_plots_dir, "/", "cancer-type-brain-per-assay.pdf")</pre>
p <- create_stacked_barplot(plot_df = tables1,</pre>
                             x_value = tables1$cancer_type_brain,
                             use_palette = palette,
                            xtitle ="Brain tumor type",
                             legend = tables1$Assay,
                             title_value = caption_value)
```





Brain tumor type

```
pdf(file = fname, width = 6, height = 5)
print(p)
dev.off()
```

7 Number of samples per brain cancer type, assay, and SJUID

Table 4: Summary of samples and assays per brain cancer type, per assay and per ${\rm SJUID}$

| can- cer_type_brain | SJUID | WES | WGBS | WGS | RNAseq | ChIPseq | Methylation | ATAC- seq |
|------------------------|----------------------------|-----|------|-----|--------|---------|-------------|--------------|
| | | | | | | | • | |
| Ependymoma | SJH0H5WYREF SJH2HPPEEKM | | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | | | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH51B396IW | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH5HCKPC97 | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH98QNKIUU | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH9YML- FOTI | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHA6KC56J6 | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH- BKS7QSFO | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHC70DZRJS | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHCN03RCVD | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHD1KI- UMM6 | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHFG- GJRHYO | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJH- HZH67WMF | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHI8CC7QT8 | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHIQT- NAYIF | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHKVQE- BOCP | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHKYUIYAMI | ₹ 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHPVXLA- CLM | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHUMKP2L6V | 7 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| Ependymoma | SJHXTBTQ5ZT | | 1 | 1 | 1 | 1 | 1 | 0 |
| HGG | SJH5QHZM4US | | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHADL5OJME | | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJH- BISK5KBU | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHBL7CDYRN | 1 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHD- DTE0SYL | 0 | 0 | 0 | 1 | 0 | 0 | 0 |
| HGG | SJHEOVURBM | 10 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHEQG3P4FK | | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHHPN- QERSQ | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHHW23UJ9P | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHJ59RLHSU | 0 | 0 | 0 | 1 | | 1 | 0 |
| HGG | SJHKKDDX- | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| 1166 | ОҮН | U | U | U | 1 | 0 | 1 | U |

| can- cer_type_brain | SJUID V | VES | WGBS | WGS | RNAseq | ChIPseq | Methylation | ATAC- seq |
|------------------------|------------------|-----|------|-----|--------|---------|-------------|--------------|
| HGG | SJHMI643DMD | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHQBOPS9FD | 0 | 0 | 0 | 1 | 0 | 0 | 0 |
| HGG | SJHRO- JUQZAP | 0 | 0 | 0 | 1 | 0 | 1 | 0 |
| HGG | SJHUT- PISXFQ | 0 | 0 | 0 | 1 | 0 | 0 | 0 |
| HGG | SJHVIS5Q8HX | 0 | 0 | 0 | 1 | 0 | 0 | 0 |
| LGG | SJH2W47P7DG | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHAY4GX4AN | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHBN- JSZHW6 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHBV3Q6UVR | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJH- CLGFJTIG | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHD- DTE0SYL | 1 | 0 | 1 | 0 | 1 | 1 | 1 |
| LGG | SJHI52BLNWK | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHN- PJTQHIT | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHO- FORRR7C | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHOY05OJJN | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHQBOPS9FD | 1 | 0 | 1 | 0 | 1 | 1 | 1 |
| LGG | SJHUT- PISXFQ | 1 | 0 | 1 | 0 | 1 | 1 | 1 |
| LGG | SJHVIS5Q8HX | 1 | 0 | 1 | 0 | 1 | 1 | 1 |
| LGG | SJHWS0NRZVA | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHXIKCWNKY | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHYP- KTG3P5 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJHZW7GYEF9 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| LGG | SJH5HCKPC97 | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| LGG | SJHC70DZRJS | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| LGG | SJHCN03RCVD | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| LGG | SJHIQT- NAYIF | 0 | 0 | 0 | 0 | 0 | 0 | 1 |
| Medulloblastoma | SJH77NRD- WUX | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJH8HIE3P0P | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHAF- TIOMPQ | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHC1QST5GR | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHF- CYGKSDY | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHJAC- CGA3S | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHQS0D51KH | 0 | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHXVMEU21L | | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHY4W1ZWCY | | 0 | 0 | 0 | 0 | 1 | 0 |
| Medulloblastoma | SJHZ40PT- CYH | 0 | 0 | 0 | 0 | 0 | 1 | 0 |

| can- cer_type_brain | SJUID | WES | WGBS | WGS | RNAseq | ChIPseq | Methylation | ATAC- seq |
|------------------------|------------------|-----|------|-----|--------|---------|-------------|--------------|
| Medulloblastoma | SJHZZLR- CGJ6 | 0 | 0 | 0 | 0 | 0 | 1 | 0 |

8 Future directions

The current exploratory data analysis module can be expanded by investigating samples with paired assays. Moreover, if other metadata are available, e.g., disease_stage, treatment, this will build large, longitudinal cohorts with multi-omic sequencing data. Such an analysis permits consideration of samples according to the condition(s) of the experiment and research aims. In addition, it can be used to refine research questions and/or generate new ones.

This will facilitate collaboration across departments at St. Jude, expedite discoveries, and find cures for children with cancer and other catastrophic diseases.

9 Session Info

```
## R version 4.4.0 (2024-04-24)
## Platform: x86_64-pc-linux-gnu
## Running under: Red Hat Enterprise Linux 8.8 (Ootpa)
##
## Matrix products: default
           /research/rgs01/applications/hpcf/authorized_apps/rhel8_apps/lapack/3.10.1/install/lib64/lib
## LAPACK: /research/rgs01/applications/hpcf/authorized_apps/rhel8_apps/lapack/3.10.1/install/lib64/lib
## locale:
##
  [1] LC_CTYPE=en_US.UTF-8
                                   LC_NUMERIC=C
  [3] LC_TIME=en_US.UTF-8
                                   LC_COLLATE=en_US.UTF-8
   [5] LC_MONETARY=en_US.UTF-8
                                   LC_MESSAGES=en_US.UTF-8
   [7] LC_PAPER=en_US.UTF-8
                                   LC_NAME=C
##
  [9] LC_ADDRESS=C
                                   LC_TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
##
## time zone: America/Chicago
## tzcode source: system (glibc)
## attached base packages:
## [1] grid
                 stats
                           graphics grDevices utils
                                                          datasets methods
## [8] base
## other attached packages:
  [1] ggthemes_5.1.0
                        lubridate_1.9.3 forcats_1.0.0
##
                                                         stringr_1.5.1
   [5] dplyr_1.1.4
                        purrr_1.0.2
                                        readr_2.1.5
                                                         tidyr_1.3.1
## [9] tibble_3.2.1
                        ggplot2_3.5.1
                                        tidyverse_2.0.0
## loaded via a namespace (and not attached):
## [1] sass_0.4.9
                          utf8_1.2.4
                                             generics_0.1.3
                                                               stringi_1.8.4
## [5] hms_1.1.3
                          digest_0.6.37
                                             magrittr_2.0.3
                                                               evaluate_0.24.0
## [9] timechange_0.3.0 fastmap_1.2.0
                                             jsonlite_1.8.8
                                                               tinytex_0.52
## [13] fansi_1.0.6
                          scales_1.3.0
                                             jquerylib_0.1.4
                                                               cli_3.6.3
## [17] rlang_1.1.4
                          crayon_1.5.3
                                             bit64_4.0.5
                                                               munsell_0.5.1
## [21] withr_3.0.1
                                             yaml_2.3.10
                                                               tools_4.4.0
                          cachem_1.1.0
## [25] parallel_4.4.0
                          tzdb_0.4.0
                                             colorspace_2.1-1
                                                               vctrs_0.6.5
## [29] R6_2.5.1
                          mime_0.12
                                             lifecycle_1.0.4
                                                               bit_4.0.5
## [33] vroom_1.6.5
                                             pillar_1.9.0
                          pkgconfig_2.0.3
                                                               bslib_0.8.0
## [37] gtable_0.3.5
                          glue_1.7.0
                                             xfun 0.47
                                                               tidyselect 1.2.1
                                                               htmltools_0.5.8.1
## [41] highr_0.11
                          knitr_1.48
                                             farver_2.1.2
## [45] rmarkdown_2.28
                          labeling_0.4.3
                                             compiler_4.4.0
```