

Longitudinal data exploratory analysis

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

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, Assay and longitudinal.

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>
longitudinal_palette_file <- file.path(root_dir, "figures", "palettes", "longitudinal_color_palette.tsv</pre>
# File path to `plots` directory
plots_dir <- file.path(analysis_dir, "plots")</pre>
figures_plots_dir <- file.path(plots_dir, "figures-longitudinal")</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
assays_number <- length(df$SJUID)
samples_number <- length(unique(df$SJUID))</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

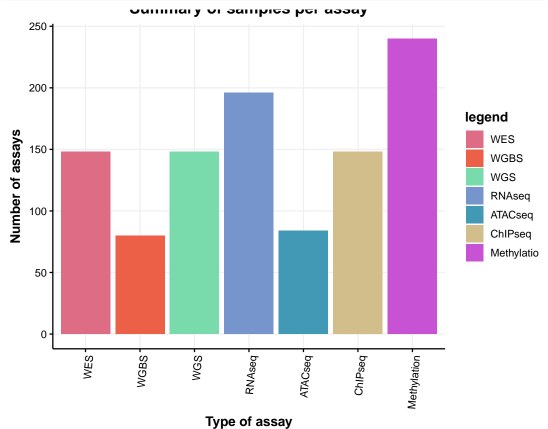
# Read color palette for longitudinal
longitudinal_palette_df <- readr::read_tsv(longitudinal_palette_file, guess_max = 100000, show_col_type)

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

5 Number of samples with assay information

Table 1: Summary of samples per assay

Assay	n
WES	148
WGBS	80
WGS	148
RNAseq	196
ATACseq	84
ChIPseq	148
Methylation	240



```
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 1044 assays in total.

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

cancer_type_brain	n
Ependymoma	480
HGG	112
LGG	408
Medulloblastoma	44

ounniary or samples and assays per brain cancer type 500 400 Number of assays legend 300 Ependymoma HGG LGG 200 Medulloblastom: 100 0 Ependymoma • .997 **Brain tumor type**

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

pdf ## 2

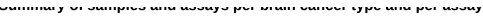
6.2 Per assay

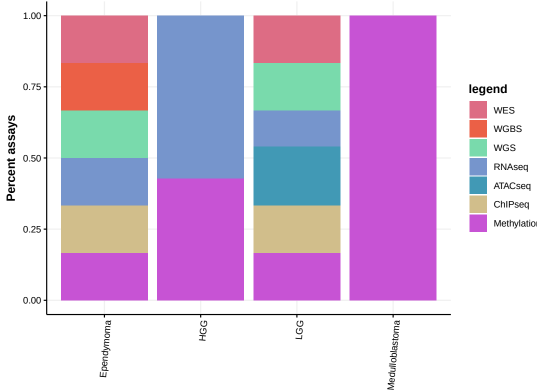
Table 3: Summary of samples and assays per brain cancer type and per assay

cancer_type_brain	WES	WGBS	WGS	RNAseq	ChIPseq	Methylation	ATACseq
Ependymoma	80	80	80	80	80	80	0

cancer_type_brain	WES	WGBS	WGS	RNAseq	ChIPseq	Methylation	ATACseq
HGG	0	0	0	64	0	48	0
LGG	68	0	68	52	68	68	84
Medulloblastoma	0	0	0	0	0	44	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-longitudinal.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()
```

6.3 Per disease stage

Table 4: Summary of samples and assays per brain cancer type, per disease stage and per assay $\,$

can-								ATAC-
cer_type_brain	longitudinal	WES	WGBS	WGS	RNAseq	ChIPseq	Methylation	seq
Ependymoma	Diagnosis	20	20	20	20	20	20	0
Ependymoma	Remission	20	20	20	20	20	20	0
Ependymoma	Relapse	20	20	20	20	20	20	0
Ependymoma	Deceased	20	20	20	20	20	20	0
HGG	Diagnosis	0	0	0	16	0	12	0
HGG	Remission	0	0	0	16	0	12	0
HGG	Relapse	0	0	0	16	0	12	0
HGG	Deceased	0	0	0	16	0	12	0
LGG	Diagnosis	17	0	17	13	17	17	21
LGG	Remission	17	0	17	13	17	17	21
LGG	Relapse	17	0	17	13	17	17	21
LGG	Deceased	17	0	17	13	17	17	21
Medulloblastoma	Diagnosis	0	0	0	0	0	11	0
Medulloblastoma	Remission	0	0	0	0	0	11	0
Medulloblastoma	Relapse	0	0	0	0	0	11	0
Medulloblastoma	Deceased	0	0	0	0	0	11	0

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

Table 5: Summary of cohort per brain cancer type, SJUID, disease stage, and assay $\,$

		11:						Nf - + 11 -	ATAC-
can- cer_type_brain	SJUID	longitudi- nal	WES	WGBS	WGS	RNAseq	ChIPsea	Methyla- tion	seq
Ependymoma	SJH0H5WYR	-	1	1	1	1	1	1	0
Ependymoma	SJH0H5WYR		1	1	1	1	1	1	0
Ependymoma	SJH0H5WYR	-	1	1	1	1	1	1	0
Ependymoma	SJH0H5WYR		1	1	1	1	1	1	0
Ependymoma	SJH2HPPEE	-	1	1	1	1	1	1	0
Ependymoma	SJH2HPPEE		1	1	1	1	1	1	0
Ependymoma	SJH2HPPEE	-	1	1	1	1	1	1	0
Ependymoma	SJH2HPPEE		1	1	1	1	1	1	0
Ependymoma	SJH51B396IV	-	1	1	1	1	1	1	0
Ependymoma	SJH51B396IV		1	1	1	1	1	1	0
Ependymoma	SJH51B396IV		1	1	1	1	1	1	0
Ependymoma	SJH51B396IV		1	1	1	1	1	1	0
Ependymoma	SJH5HCKPC		1	1	1	1	1	1	0
Ependymoma	SJH5HCKPC		1	1	1	1	1	1	0
Ependymoma	SJH5HCKPC	-	1	1	1	1	1	1	0
Ependymoma	SJH5HCKPC		1	1	1	1	1	1	0
Ependymoma	SJH98QNKIU		1	1	1	1	1	1	0
Ependymoma	SJH98QNKIU		1	1	1	1	1	1	0
Ependymoma	SJH98QNKIU	-	1	1	1	1	1	1	0
Ependymoma	SJH98QNKIU		1	1	1	1	1	1	0
Ependymoma	SJH9YML-	Diagnosis	1	1	1	1	1	1	0
	FOTI								
Ependymoma	SJH9YML-	Remission	1	1	1	1	1	1	0
	FOTI								
Ependymoma	SJH9YML-	Relapse	1	1	1	1	1	1	0
	FOTI								
Ependymoma	SJH9YML-	Deceased	1	1	1	1	1	1	0
	FOTI								
Ependymoma	SJHA6KC56J	-	1	1	1	1	1	1	0
Ependymoma	SJHA6KC56J		1	1	1	1	1	1	0
Ependymoma	SJHA6KC56J		1	1	1	1	1	1	0
Ependymoma	SJHA6KC56J		1	1	1	1	1	1	0
Ependymoma	SJH-	Diagnosis	1	1	1	1	1	1	0
	BKS7QSFO								
Ependymoma	SJH-	Remission	1	1	1	1	1	1	0
	BKS7QSFO								
Ependymoma	SJH-	Relapse	1	1	1	1	1	1	0
	BKS7QSFO								
Ependymoma	SJH-	Deceased	1	1	1	1	1	1	0
	BKS7QSFO								
Ependymoma	SJHC70DZR.	JSDiagnosis	1	1	1	1	1	1	0
Ependymoma	SJHC70DZR.	J:Remission	1	1	1	1	1	1	0
Ependymoma	SJHC70DZR.	JS Relapse	1	1	1	1	1	1	0
Ependymoma	SJHC70DZR.	-	1	1	1	1	1	1	0
Ependymoma	SJHCN03RC		1	1	1	1	1	1	0
- •		~							

can-	longitudi-						Methyla-	ATAC-
cer_type_brain	SJUID nal	WES	WGBS	WGS	RNAseq	ChIPseq	tion	seq
Ependymoma	SJHCN03RCVRemission	1	1	1	1	1	1	0
Ependymoma	SJHCN03RCVDRelapse	1	1	1	1	1	1	0
Ependymoma	${\bf SJHCN03RCV}\textbf{\textbf{D}}eceased$	1	1	1	1	1	1	0
Ependymoma	SJHD1KI- Diagnosis UMM6	1	1	1	1	1	1	0
Ependymoma	SJHD1KI- Remission UMM6	1	1	1	1	1	1	0
Ependymoma	SJHD1KI- Relapse UMM6	1	1	1	1	1	1	0
Ependymoma	SJHD1KI- Deceased UMM6	1	1	1	1	1	1	0
Ependymoma	SJHFG- Diagnosis GJRHYO	1	1	1	1	1	1	0
Ependymoma	SJHFG- Remission GJRHYO	1	1	1	1	1	1	0
Ependymoma	SJHFG- Relapse GJRHYO	1	1	1	1	1	1	0
Ependymoma	SJHFG- Deceased GJRHYO	1	1	1	1	1	1	0
Ependymoma	SJH- Diagnosis HZH67WMF	1	1	1	1	1	1	0
Ependymoma	SJH- Remission HZH67WMF	1	1	1	1	1	1	0
Ependymoma	SJH- Relapse HZH67WMF	1	1	1	1	1	1	0
Ependymoma	SJH- Deceased HZH67WMF	1	1	1	1	1	1	0
Ependymoma	SJHI8CC7QT8Diagnosis	1	1	1	1	1	1	0
Ependymoma	SJHI8CC7QT8Remission	1	1	1	1	1	1	0
Ependymoma	SJHI8CC7QT8 Relapse	1	1	1	1	1	1	0
Ependymoma	SJHI8CC7QT8Deceased	1	1	1	1	1	1	0
Ependymoma	SJHIQT- Diagnosis NAYIF	1	1	1	1	1	1	0
Ependymoma	SJHIQT- Remission NAYIF	1	1	1	1	1	1	0
Ependymoma	SJHIQT- Relapse NAYIF	1	1	1	1	1	1	0
Ependymoma	SJHIQT- Deceased NAYIF	1	1	1	1	1	1	0
Ependymoma	SJHKVQE- Diagnosis BOCP	1	1	1	1	1	1	0
Ependymoma	SJHKVQE- Remission BOCP	1	1	1	1	1	1	0
Ependymoma	SJHKVQE- Relapse BOCP	1	1	1	1	1	1	0
Ependymoma	SJHKVQE- Deceased BOCP	1	1	1	1	1	1	0
Ependymoma	SJHKYUIYAMDiagnosis	1	1	1	1	1	1	0
Ependymoma	SJHKYUIYAMæmission	1	1	1	1	1	1	0
Ependymoma	${\bf SJHKYUIYAM {\bf I\!R} elapse}$	1	1	1	1	1	1	0
Ependymoma	SJHKYUIYAMDeceased	1	1	1	1	1	1	0

can-		longitudi-						Methyla-	ATAC-
cer_type_brain	SJUID	$_{\mathrm{nal}}$	WES	WGBS	WGS	RNAseq	ChIPseq	tion	seq
Ependymoma	SJH-	Diagnosis	1	1	1	1	1	1	0
- •	PVXLA-								
	CLM								
Ependymoma	SJH-	Remission	1	1	1	1	1	1	0
- •	PVXLA-								
	CLM								
Ependymoma	SJH-	Relapse	1	1	1	1	1	1	0
	PVXLA-								
	CLM								
Ependymoma	SJH-	Deceased	1	1	1	1	1	1	0
1 0	PVXLA-								
	CLM								
Ependymoma	SJHUMKP2I	L6 D iagnosis	1	1	1	1	1	1	0
Ependymoma	SJHUMKP2I		1	1	1	1	1	1	0
Ependymoma	SJHUMKP2I	L6WRelapse	1	1	1	1	1	1	0
Ependymoma	SJHUMKP2I	-	1	1	1	1	1	1	0
Ependymoma	SJHXTBTQ		1	1	1	1	1	1	0
Ependymoma	SJHXTBTQ		1	1	1	1	1	1	0
Ependymoma	SJHXTBTQ		1	1	1	1	1	1	0
Ependymoma	SJHXTBTQ	-	1	1	1	1	1	1	0
HGG	SJH5QHZM4		0	0	0	1	0	1	0
HGG	SJH5QHZM4	~	0	0	0	1	0	1	0
HGG	SJH5QHZM4		0	0	0	1	0	1	0
HGG	SJH5QHZM4	-	0	0	0	1	0	1	0
HGG	SJHADL5OJ		0	0	0	1	0	1	0
HGG	SJHADL5OJ		0	0	0	1	0	1	0
HGG	SJHADL5OJ		0	0	0	1	0	1	0
HGG	SJHADL5OJ	-	0	0	0	1	0	1	0
HGG	SJH-	Diagnosis	0	0	0	1	0	1	0
1166	BISK5KBU	Diagnosis	U	U	U	1	U	1	U
HGG	SJH-	Remission	0	0	0	1	0	1	0
IIGG	BISK5KBU	Reimssion	U	U	U	1	U	1	U
HGG	SJH-	Dolongo	0	0	0	1	0	1	0
DDII		Relapse	0	U	0	1	0	1	0
HCC	BISK5KBU	Deceased	0	0	0	1	0	1	0
HGG	SJH- BISK5KBU	Deceased	0	0	0	1	0	1	0
HCC		DW:	0	0	0	1	0	1	0
HGG	SJHBL7CDY		0	0	0	1	0	1	0
HGG	SJHBL7CDY		0	0	0	1	0	1	0
HGG	SJHBL7CDY		0	0	0	1	0	1	0
HGG	SJHBL7CDY		0	0	0	1	0	1	0
HGG	SJHD-	Diagnosis	0	0	0	1	0	0	0
шаа	DTE0SYL	ъ	0	0	0	4	0	^	0
HGG	SJHD-	Remission	0	0	0	1	0	0	0
HÃÔ	DTE0SYL	ъ.	•	_	0		_	_	
HGG	SJHD-	Relapse	0	0	0	1	0	0	0
	DTE0SYL		_	_			_	_	_
HGG	SJHD-	Deceased	0	0	0	1	0	0	0
	DTE0SYL								
HGG	SJHEOVURI		0	0	0	1	0	1	0
TICC	SJHEOVURI	BWRAmission	0	0	0	1	0	1	0
HGG HGG	SJHEOVURI		-		0	1	0	1	0

can-	longitudi-						Methyla-	ATAC-
cer_type_brain	SJUID nal	WES	WGBS	WGS	RNAseq	ChIPseq	tion	seq
HGG	SJHEOVURBMDeceased	0	0	0	1	0	1	0
HGG	SJHEQG3P4F\(\mathbb{E}\) iagnosis	0	0	0	1	0	1	0
HGG	SJHEQG3P4FRemission	0	0	0	1	0	1	0
$_{ m HGG}$	${\bf SJHEQG3P4FKRelapse}$	0	0	0	1	0	1	0
HGG	SJHEQG3P4FIDeceased	0	0	0	1	0	1	0
HGG	SJHHPN- Diagnosis	0	0	0	1	0	1	0
	QERSQ							
HGG	SJHHPN- Remission	0	0	0	1	0	1	0
	QERSQ							
HGG	SJHHPN- Relapse	0	0	0	1	0	1	0
	QERSQ							
HGG	SJHHPN- Deceased	0	0	0	1	0	1	0
	QERSQ							
HGG	SJHHW23UJ9PDiagnosis	0	0	0	1	0	1	0
HGG	SJHHW23UJ9Remission	0	0	0	1	0	1	0
HGG	SJHHW23UJ9PRelapse	0	0	0	1	0	1	0
HGG	SJHHW23UJ9Deceased	0	0	0	1	0	1	0
HGG	SJHJ59RLHSUDiagnosis	0	0	0	1	0	1	0
HGG	SJHJ59RLHSURemission	0	0	0	1	0	1	0
HGG	SJHJ59RLHSU Relapse	0	0	0	1	0	1	0
HGG	SJHJ59RLHSUDeceased	0	0	0	1	0	1	0
HGG	SJHKKD- Diagnosis	0	0	0	1	0	1	0
шаа	DXOYH	0	0	0	1	0	1	0
HGG	SJHKKD- Remission	0	0	0	1	0	1	0
HOO	DXOYH	0	0	0	1	0	1	0
HGG	SJHKKD- Relapse DXOYH	0	0	0	1	0	1	0
HGG	SJHKKD- Deceased	0	0	0	1	0	1	0
1166	DXOYH	U	U	U	1	U	1	U
HGG	SJHMI643DMIDiagnosis	0	0	0	1	0	1	0
HGG	SJHMI643DMRemission	0	0	0	1	0	1	0
HGG	SJHMI643DMDRelapse	0	0	0	1	0	1	0
HGG	SJHMI643DMIDeceased	0	0	0	1	0	1	0
HGG	SJHQBOPS9F D iagnosis	0	0	0	1	0	0	0
HGG	SJHQBOPS9FRemission	0	0	0	1	0	0	0
HGG	SJHQBOPS9FDRelapse	0	0	0	1	0	0	0
HGG	SJHQBOPS9FDeceased	0	0	0	1	0	0	0
HGG	SJHRO- Diagnosis	0	0	0	1	0	1	0
	JUQZAP							
HGG	SJHRO- Remission	0	0	0	1	0	1	0
	JUQZAP							
HGG	SJHRO- Relapse	0	0	0	1	0	1	0
	JUQZAP							
HGG	SJHRO- Deceased	0	0	0	1	0	1	0
	JUQZAP							
HGG	SJHUT- Diagnosis	0	0	0	1	0	0	0
	PISXFQ							
HGG	SJHUT- Remission	0	0	0	1	0	0	0
	PISXFQ							
HGG	SJHUT- Relapse	0	0	0	1	0	0	0
	PISXFQ							

can- cer_type_	longitudi- brain SJUID nal	WES	WGBS	WGS	RNAseq	ChIPseq	Methyla- tion	ATAC- seq
HGG	SJHUT- Deceased PISXFQ	0	0	0	1	0	0	0
HGG	SJHVIS5Q8HXDiagnosis	0	0	0	1	0	0	0
HGG	SJHVIS5Q8HXRemission	0	0	0	1	0	0	0
$_{\mathrm{HGG}}$	SJHVIS5Q8HX Relapse	0	0	0	1	0	0	0
HGG	SJHVIS5Q8HXDeceased	0	0	0	1	0	0	0
LGG	SJH2W47P7D@iagnosis	1	0	1	1	1	1	1
LGG	SJH2W47P7D Remission	1	0	1	1	1	1	1
LGG	SJH2W47P7DGRelapse	1	0	1	1	1	1	1
LGG	SJH2W47P7D (Deceased	1	0	1	1	1	1	1
LGG	SJH5HCKPC9Diagnosis	0	0	0	0	0	0	1
LGG	SJH5HCKPC9Remission	0	0	0	0	0	0	1
LGG	SJH5HCKPC97Relapse	0	0	0	0	0	0	1
LGG	SJH5HCKPC97Deceased	0	0	0	0	0	0	1
LGG	SJHAY4GX4A D iagnosis	1	0	1	1	1	1	1
LGG	SJHAY4GX4ARemission	1	0	1	1	1	1	1
LGG	SJHAY4GX4ANRelapse	1	0	1	1	1	1	1
LGG	SJHAY4GX4A Deceased	1	0	1	1	1	1	1
LGG	SJHBN- Diagnosis JSZHW6	1	0	1	1	1	1	1
LGG	SJHBN- Remission JSZHW6	1	0	1	1	1	1	1
LGG	SJHBN- Relapse JSZHW6	1	0	1	1	1	1	1
LGG	SJHBN- Deceased JSZHW6	1	0	1	1	1	1	1
LGG	SJHBV3Q6UV R iagnosis	1	0	1	1	1	1	1
LGG	SJHBV3Q6UVRemission	1	0	1	1	1	1	1
LGG	SJHBV3Q6UVRelapse	1	0	1	1	1	1	1
LGG	SJHBV3Q6UV R eceased	1	0	1	1	1	1	1
LGG	SJHC70DZRJSDiagnosis	0	0	0	0	0	0	1
LGG	SJHC70DZRJSRemission	0	0	0	0	0	0	1
LGG	SJHC70DZRJS Relapse	0	0	0	0	0	0	1
LGG	SJHC70DZRJSDeceased	0	0	0	0	0	0	1
LGG	SJH- Diagnosis CLGFJTIG	1	0	1	1	1	1	1
LGG	SJH- Remission CLGFJTIG	1	0	1	1	1	1	1
LGG	SJH- Relapse CLGFJTIG	1	0	1	1	1	1	1
LGG	SJH- Deceased CLGFJTIG	1	0	1	1	1	1	1
LGG	$\mathrm{SJHCN03RCV}\mathbf{D}$ iagnosis	0	0	0	0	0	0	1
LGG	SJHCN03RCVR emission	0	0	0	0	0	0	1
LGG	SJHCN03RCVDRelapse	0	0	0	0	0	0	1
LGG	SJHCN03RCVDeceased	0	0	0	0	0	0	1
LGG	SJHD- Diagnosis DTE0SYL	1	0	1	0	1	1	1
LGG	SJHD- Remission DTE0SYL	1	0	1	0	1	1	1
LGG	SJHD- Relapse DTE0SYL	1	0	1	0	1	1	1

can- cer_type_brain	longitudi- SJUID nal	WES	WCB	s wgs	RNAseq	ChIPsea	Methyla- tion	ATAC- seq
LGG	SJHD- Deceased DTE0SYL	1	0	1	0	1	1	1
LGG	SJHI52BLNWIDiagnosis	1	0	1	1	1	1	1
LGG	SJHI52BLNW R emission	1	0	1	1	1	1	1
LGG	${\it SJHI52BLNWKRelapse}$	1	0	1	1	1	1	1
LGG	SJHI52BLNWkDeceased	1	0	1	1	1	1	1
LGG	SJHIQT- Diagnosis NAYIF	0	0	0	0	0	0	1
LGG	SJHIQT- Remission NAYIF	0	0	0	0	0	0	1
LGG	SJHIQT- Relapse NAYIF	0	0	0	0	0	0	1
LGG	SJHIQT- Deceased NAYIF	0	0	0	0	0	0	1
LGG	SJHN- Diagnosis PJTQHIT	1	0	1	1	1	1	1
LGG	SJHN- Remission PJTQHIT	1	0	1	1	1	1	1
LGG	SJHN- Relapse PJTQHIT	1	0	1	1	1	1	1
LGG	SJHN- Deceased PJTQHIT	1	0	1	1	1	1	1
LGG	SJHO- Diagnosis FORRR7C	1	0	1	1	1	1	1
LGG	SJHO- Remission FORRR7C	1	0	1	1	1	1	1
LGG	SJHO- Relapse FORRR7C	1	0	1	1	1	1	1
LGG	SJHO- Deceased FORRR7C	1	0	1	1	1	1	1
LGG	SJHOY05OJJNDiagnosis	1	0	1	1	1	1	1
LGG	SJHOY05OJJNRemission	1	0	1	1	1	1	1
LGG	SJHOY05OJJN Relapse	1	0	1	1	1	1	1
LGG	SJHOY05OJJNDeceased	1	0	1	1	1	1	1
LGG	${\rm SJHQBOPS9F}\textbf{D}{\rm iagnosis}$	1	0	1	0	1	1	1
LGG	SJHQBOPS9FRemission	1	0	1	0	1	1	1
$_{ m LGG}$	SJHQBOPS9FDRelapse	1	0	1	0	1	1	1
LGG	SJHQBOPS9FDeceased	1	0	1	0	1	1	1
LGG	SJHUT- Diagnosis PISXFQ	1	0	1	0	1	1	1
LGG	SJHUT- Remission PISXFQ	1	0	1	0	1	1	1
LGG	SJHUT- Relapse PISXFQ	1	0	1	0	1	1	1
LGG	SJHUT- Deceased PISXFQ	1	0	1	0	1	1	1
LGG	SJHVIS5Q8HXDiagnosis	1	0	1	0	1	1	1
LGG	SJHVIS5Q8HXRemission	1	0	1	0	1	1	1
LGG	SJHVIS5Q8HX Relapse	1	0	1	0	1	1	1
LGG	SJHVIS5Q8HXDeceased	1	0	1	0	1	1	1
LGG	SJHWS0NRZVA iagnosis	1	0	1	1	1	1	1
_~~	2011, 701, 102 (Hangilooli)	-	V	-	-	-	-	-

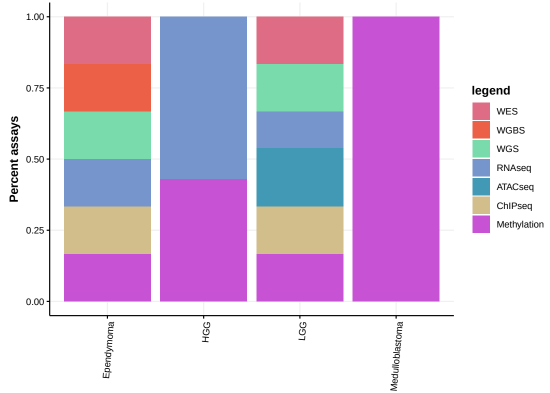
can- cer_type_brain	longitudi- SJUID nal	WES	WGBS	s wgs	RNAseq	ChIPseq	Methyla- tion	ATAC- seq
LGG	SJHWS0NRZWAemission	1	0	1	1	1	1	1
LGG	SJHWS0NRZVARelapse	1	0	1	1	1	1	1
LGG	SJHWS0NRZV A eceased	1	0	1	1	1	1	1
LGG	SJHXIKCWNKDiagnosis	1	0	1	1	1	1	1
LGG	SJHXIKCWN Remission	1	0	1	1	1	1	1
LGG	SJHXIKCWNK K elapse	1	0	1	1	1	1	1
LGG	SJHXIKCWNK Meceased	1	0	1	1	1	1	1
LGG	SJHYP- Diagnosis	1	0	1	1	1	1	1
Luu	KTG3P5	1	U	1	1	1	1	1
LGG	SJHYP- Remission KTG3P5	1	0	1	1	1	1	1
LGG	SJHYP- Relapse KTG3P5	1	0	1	1	1	1	1
LGG	SJHYP- Deceased KTG3P5	1	0	1	1	1	1	1
LGG	SJHZW7GYEFP9iagnosis	1	0	1	1	1	1	1
LGG	SJHZW7GYEF@mission	1	0	1	1	1	1	1
LGG	SJHZW7GYEFRelapse	1	0	1	1	1	1	1
LGG	SJHZW7GYEF P eceased	1	0	1	1	1	1	1
Medulloblas-	SJH77NRD- Diagnosis	0	0	0	0	0	1	0
toma	WUX							
Medulloblas-	SJH77NRD- Remission	0	0	0	0	0	1	0
toma	WUX							
Medulloblas-	SJH77NRD- Relapse WUX	0	0	0	0	0	1	0
toma Medulloblas- toma	SJH77NRD- Deceased WUX	0	0	0	0	0	1	0
Medulloblas- toma	SJH8HIE3P0P Diagnosis	0	0	0	0	0	1	0
Medulloblas- toma	SJH8HIE3P0PRemission	0	0	0	0	0	1	0
Medulloblas- toma	SJH8HIE3P0P Relapse	0	0	0	0	0	1	0
Medulloblas-	SJH8HIE3P0P Deceased	0	0	0	0	0	1	0
toma Medulloblas- toma	SJHAF- Diagnosis TIOMPQ	0	0	0	0	0	1	0
Medulloblas-	SJHAF- Remission	0	0	0	0	0	1	0
toma	TIOMPQ							
Medulloblas-	SJHAF- Relapse	0	0	0	0	0	1	0
toma	TIOMPQ							
Medulloblas-	SJHAF- Deceased	0	0	0	0	0	1	0
toma	TIOMPQ							
Medulloblas- toma	SJHC1QST5GRDiagnosis	0	0	0	0	0	1	0
Medulloblas- toma	${\rm SJHC1QST5G} \textbf{R} emission$	0	0	0	0	0	1	0
Medulloblas- toma	${\bf SJHC1QST5GRRelapse}$	0	0	0	0	0	1	0
Medulloblas- toma	SJHC1QST5GRDeceased	0	0	0	0	0	1	0

can- cer_type_brain	SJUID	longitudi- nal	WES	WGBS	WGS	RNAseq	ChIPseq	Methyla- tion	ATAC- seq
Medulloblas-	SJHF-	Diagnosis	0	0	0	0	0	1	0
toma	CYGKSDY								
Medulloblas-	SJHF-	Remission	0	0	0	0	0	1	0
toma	CYGKSDY								
Medulloblas-	SJHF-	Relapse	0	0	0	0	0	1	0
toma	CYGKSDY								
Medulloblas-	SJHF-	Deceased	0	0	0	0	0	1	0
toma	CYGKSDY								
Medulloblas-	SJHJAC-	Diagnosis	0	0	0	0	0	1	0
toma	CGA3S	.						_	
Medulloblas-	SJHJAC-	Remission	0	0	0	0	0	1	0
toma	CGA3S	D 1	0	0	0	0	0	1	0
Medulloblas-	SJHJAC-	Relapse	0	0	0	0	0	1	0
toma Medulloblas-	CGA3S SJHJAC-	Daggard	0	0	0	0	0	1	0
toma	CGA3S	Deceased	U	U	U	0	U	1	U
Medulloblas-	SJHQS0D511	V IDio emogia	0	0	0	0	0	1	0
toma	231162012311	KIDiagilosis	U	U	U	U	U	1	U
Medulloblas-	SJHQS0D511	KIRomission	0	0	0	0	0	1	0
toma	231162012311	1XIII(eIIIISSIOII	U	U	U	U	U	1	U
Medulloblas-	SJHQS0D511	KHRelanse	0	0	0	0	0	1	0
toma	501105502511	rifficiapse	O	O	O	O	O	1	O
Medulloblas-	SJHQS0D511	KHDeceased	0	0	0	0	0	1	0
toma	33-1-40-3-3-3			, ,	•		, and the second	_	ŭ
Medulloblas-	SJHXVMEU	21Diagnosis	0	0	0	0	0	1	0
toma		3 3 3 3							
Medulloblas-	SJHXVMEU	21Remission	0	0	0	0	0	1	0
toma									
Medulloblas-	SJHXVMEU	21IRelapse	0	0	0	0	0	1	0
toma		_							
Medulloblas-	SJHXVMEU	21Deceased	0	0	0	0	0	1	0
toma									
Medulloblas-	SJHY4W1ZV	W D Magnosis	0	0	0	0	0	1	0
toma									
Medulloblas-	SJHY4W1ZV	V E mission	0	0	0	0	0	1	0
toma									
Medulloblas-	SJHY4W1ZV	VC K elapse	0	0	0	0	0	1	0
toma	~								
Medulloblas-	SJHY4W1ZV	WCM'eceased	0	0	0	0	0	1	0
toma	G 777 F 40 D FF	. .						_	
Medulloblas-	SJHZ40PT-	Diagnosis	0	0	0	0	0	1	0
toma	СҮН	ъ	0	0	0	0	0	-1	0
Medulloblas-	SJHZ40PT-	Remission	0	0	0	0	0	1	0
toma	CYH	D 1	0	0	0	0	0	1	0
Medulloblas-	SJHZ40PT- CYH	Relapse	0	0	0	0	0	1	0
toma Medulloblas-	SJHZ40PT-	Deceased	0	0	0	0	0	1	0
toma	CYH	Deceased	U	U	U	U	U	1	U
toma Medulloblas-	SJHZZLR-	Diagnosis	0	0	0	0	0	1	0
toma	CGJ6	Diagnosis	U	U	U	U	U	1	U
Medulloblas-	SJHZZLR-	Remission	0	0	0	0	0	1	0
toma	CGJ6	101111031011	U	U	U	U	U	1	U
JOIII0	2000								

can- cer_type_brain	SJUID	longitudi- nal	WES	WGBS	WGS	RNAseq	ChIPseq	Methyla- tion	ATAC- seq
Medulloblas-	SJHZZLR-	Relapse	0	0	0	0	0	1	0
toma Medulloblas- toma	CGJ6 SJHZZLR- CGJ6	Deceased	0	0	0	0	0	1	0

```
tables1 <- df %>% count(cancer_type_brain, SJUID, longitudinal, Assay) %>%
  as.data.frame() %>%
  mutate_all(funs(replace_na(.,0))) %>%
  mutate(match_id_brain = paste(cancer_type_brain, SJUID, sep = "_"))
## 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.
# Run function
fname <- paste0(figures_plots_dir, "/", "cancer-type-brain-patient-assay.pdf")</pre>
p <- create_stacked_barplot(plot_df = tables1,</pre>
                            x_value = tables1$match_id_brain,
                            use_palette = palette,
                            xtitle ="Brain tumor type and patient case",
                            legend = tables1$Assay,
                            title_value = caption_value)
```

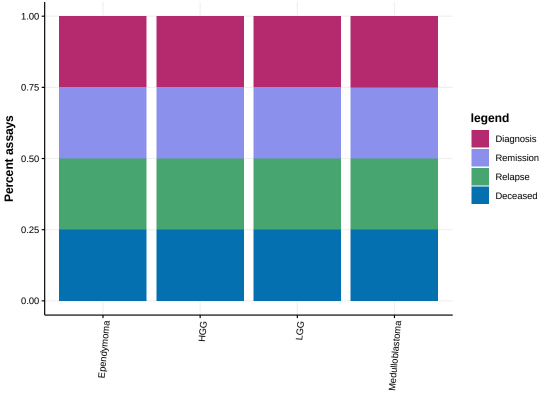
iry or samples and assays per brain valloer type, per disease stage and per assa



Brain tumor type and patient case

```
pdf(file = fname, width = 12, height = 6)
print(p)
dev.off()
```

ary or samples and assays per brain cancer type, per disease stage and per assa



Brain tumor type and patient case

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

pdf ## 2

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
```