# The Open Pediatric Cancer Project

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### In Brief

# **Highlights**

# **Summary**

# **Keywords**

## Introduction

# **Results**

# **Discussion**

# **Acknowledgments**

# **Author Contributions**

| Author             | Contributions  |
|--------------------|--|
| Eric Wafula        | Formal analysis, Software  |
| Krutika S. Gaonkar | Data curation, Formal analysis, Investigation,<br>Methodology, Software, Writing – Original draft, Writing –<br>Review and editing |
| Run Jin            | Data curation, Formal analysis, Visualization, Writing –<br>Original draft, Writing - Review and editing                           |
| Komal S. Rathi     | Formal analysis, Investigation, Methodology, Writing –<br>Original draft   |

| Author               | Contributions  |
|----------------------|--|
| Yuankun Zhu          | Data curation, Formal analysis, Investigation,<br>Methodology, Supervision   |
| Bailey K. Farrow     | Data curation, Software  |
| Daniel P. Miller     | Formal analysis  |
| Mariarita Santi      | Investigation, Validation, Writing - Review and editing  |
| Adam A. Kraya        | Methodology  |
| Xiaoyan Huang        | Formal analysis  |
| Bo Zhang             | Data curation, Formal analysis   |
| Brian M. Ennis       | Data curation, Formal analysis   |
| Ryan J. Corbett      | Formal analysis  |
| Sharon J. Diskin     | Investigation, Supervision, Validation, Funding acquisition, Writing - Review and editing  |
| Nicholas Van Kuren   | Data curation, Software  |
| Noel Coleman         | Data curation  |
| Christopher Blackden | Resources  |
| Jennifer L. Mason    | Supervision  |
| Miguel A. Brown      | Data curation, Methodology, Formal analysis  |
| Adam C. Resnick      | Conceptualization, Funding acquisition, Resources, Supervision   |
| Jo Lynne Rokita^     | Conceptualization, Data curation, Formal analysis,<br>Funding acquisition, Investigation, Methodology,<br>Software, Supervision, Writing – Original draft, Writing -<br>Review and editing |

# **Declarations of Interest**

# **Figure Titles and Legends**

# **Table Titles and Legends**

# **OPENPEDCAN METHODS**

## **RESOURCE AVAILABILITY**

### **Lead contact**

Requests for access to OpenPedCan raw data and/or specimens may be directed to, and will be fulfilled by Jo Lynne Rokita (rokita@chop.edu).

# **Materials availability**

This study did not create new, unique reagents.

## Data and code availability

Merged summary files for OpenPedCan v12 are openly accessible in <u>CAVATICA</u> or via download script in the <a href="https://github.com/PediatricOpenTargets/OpenPedCan-analysis">https://github.com/PediatricOpenTargets/OpenPedCan-analysis</a> repository. Cancer group summary data are visible within the NCI's pediatric <a href="Molecular Targets Platform">Molecular Targets Platform</a> and cohort, cancer group, and individual data are visible within <a href="PedcBioPortal">PedcBioPortal</a>

OpenPedCan analysis modules were developed within OpenPBTA [1], modified based on OpenPBTA, or newly created and can be found within the following publicly available repositories. OpenPBTA module analyses can be found at <a href="https://github.com/AlexsLemonade/OpenPBTA-analysis">https://github.com/AlexsLemonade/OpenPBTA-analysis</a>. OpenPedCan module analyses can be found at

https://github.com/PediatricOpenTargets/OpenPedCan-analysis. OpenPedCan api code can be found at https://github.com/PediatricOpenTargets/OpenPedCan-api.

Software versions are documented in **Table XX**.

### **Data releases**

### EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

### **METHOD DETAILS**

## Nucleic acids extraction and library preparation

### **Data generation**

## **DNA WGS Alignment**

Please refer to the OpenPBTA manuscript for details [1].

# **Quality Control of Sequencing Data**

Please refer to the OpenPBTA manuscript for details [1].

### SNP calling for B-allele Frequency (BAF) generation

Please refer to the OpenPBTA manuscript for details [1].

# **Somatic Mutation Calling**

### SNV and indel calling

Please refer to the OpenPBTA manuscript for details [1].

### **VCF** annotation and MAF creation

**Gather SNV and INDEL Hotspots** 

### **Consensus SNV Calling**

**Somatic Copy Number Variant Calling (WGS samples only)** 

### **Consensus CNV Calling**

# **Somatic Structural Variant Calling (WGS samples only)**

Please refer to the OpenPBTA manuscript for details [1].

**Methylation Analysis** 

**Gene Expression** 

**Abundance Estimation** 

**Gene Expression Matrices with Unique HUGO Symbols** 

Gene fusion detection

## **QUANTIFICATION AND STATISTICAL ANALYSIS**

Focal Copy Number Calling (focal-cn-file-preparation analysis module)

Please refer to the OpenPBTA manuscript for details on assignment of copy number status values to CNV segments, cytobands, and genes [1]. We applied criteria to resolve instances of multiple conflicting status calls for the same gene and sample, which are described in detail in the <u>focal-cn-file-preparation</u> module. Briefly, we prioritized 1) non-neutral status calls, 2) calls made from dominant segments with respect to gene overlap, and 3) amplification and deep deletion status calls over gain and loss calls, respectively, when selecting a dominant status call per gene and sample. These methods resolved >99% of duplicated gene-level status calls.

Gene Set Variation Analysis (gene-set-enrichment-analysis analysis module)

Please refer to the OpenPBTA manuscript for details [1].

Fusion prioritization (fusion\_filtering analysis module)

Mutational Signatures (mutational-signatures analysis module)

Tumor Mutation Burden (snv-callers analysis module)

**Clinical Data Harmonization** 

**WHO Classification of Disease Types** 

**Molecular Subtyping** 

Here, we build upon the molecular subtyping performed in OpenPBTA [1].

High-grade gliomas...

Atypical teratoid rhabdoid tumors..

Neuroblastoma tumors...

### Integration of brain tumor methylation classifications

# TP53 Alteration Annotation (tp53\_nf1\_score analysis module)

Please refer to the OpenPBTA manuscript for details [1].

## Prediction of participants' genetic sex

Please refer to the OpenPBTA manuscript for details [1].

## Selection of independent samples (independent-samples analysis module)

For analyses that require all input biospecimens to be independent, we use the OpenPedCan-analysis <u>independent-samples</u> module to select only one biospecimen from each input participant. For each input participant of an analysis, the independent biospecimen is selected based on the analysis-specific filters and preferences for the biospecimen metadata, such as experimental strategy, cancer group, and tumor descriptor.

# **Supplemental Information Titles and Legends**

### Consortia

# References

## 1. OpenPBTA: The Open Pediatric Brain Tumor Atlas

Joshua A Shapiro, Krutika S Gaonkar, Stephanie J Spielman, Candace L Savonen, Chante J Bethell, Run Jin, Komal S Rathi, Yuankun Zhu, Laura E Egolf, Bailey K Farrow, ... Jaclyn N Taroni *Cell Genomics* (2023-05) <a href="https://doi.org/gr92p6">https://doi.org/gr92p6</a>

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