Integrated Molecular Meta-Analysis of 1,000 Pediatric High-Grade and Diffuse Intrinsic Pontine Glioma

Highlights

- Pediatric HGG and DIPG comprise a diverse set of clinical and biological subgroups
- Somatic coding mutations per tumor range from none to among the highest seen in human cancer
- Histone mutations co-segregate with distinct alterations and downstream pathways
- H3/IDH1 WT tumors may resemble low-grade lesions and have targetable alterations

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In Brief

Mackay et al. perform an integrated analysis of >1,000 cases of pediatric high-grade glioma and diffuse intrinsic pontine glioma. They identify co-segregating mutations in histone-mutant subgroups and show that histone wild-type subgroups are molecularly more similar to lower-grade tumors.
Integrated Molecular Meta-Analysis of 1,000 Pediatric High-Grade and Diffuse Intrinsic Pontine Glioma

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SUMMARY

We collated data from 157 unpublished cases of pediatric high-grade glioma and diffuse intrinsic pontine glioma and 20 publicly available datasets in an integrated analysis of >1,000 cases. We identified co-segregating mutations in histone-mutant subgroups including loss of FBXW7 in H3.3G34R/V, TOP3A rearrangements in H3.3K27M, and BCOR mutations in H3.1K27M. Histone wild-type subgroups are refined by the presence of key oncogenic events or methylation profiles more closely resembling lower-grade tumors. Genomic aberrations increase with age, highlighting the infant population as biologically and clinically distinct. Uncommon pathway dysregulation is seen in small subsets of tumors, further defining the molecular diversity of the disease, opening up avenues for biological study and providing a basis for functionally defined future treatment stratification.

INTRODUCTION

Pediatric glioblastoma (pGBM) and diffuse intrinsic pontine glioma (DIPG) are high-grade glial tumors of children with a median overall survival of 9–15 months, a figure that has remained unmoved for decades (Jones et al., 2012). Although relatively rare in this age group (1.78 per 100,000 population), taken together, gliomas are nonetheless the most common malignant...
brain tumors in children, and represent the greatest cause of cancer-related deaths under the age of 19 years (Ostrom et al., 2016). Unlike histologically similar lesions in adults, which tend to be restricted to the cerebral hemispheres, diffuse high-grade gliomas in childhood (pHGG) can occur throughout the CNS, with around half occurring in midline locations, in particular the thalamus and the pons (Jones and Baker, 2014), where the lack of available surgical options confers an especially poor prognosis (Kramm et al., 2011). Numerous clinical trials of chemotherapeutics and targeted agents extrapolated from adult GBM studies have failed to show a survival benefit, and more rationally derived approaches based upon an understanding of the childhood diseases are urgently needed (Jones et al., 2016).

It has become increasingly apparent that pHGG differ from their adult counterparts, with molecular profiling studies carried out over the last 6–7 years having incrementally identified key genetic and epigenetic differences in pHGG associated with distinct ages of onset, anatomical distribution, clinical outcome, and histopathological and radiological features (Jones and Baker, 2014; Sturm et al., 2014). In particular, the identification of unique recurrent mutations in genes encoding histones H3.3 and H3.1 (Schwartzentruber et al., 2012; Wu et al., 2012) have demonstrated the distinctiveness of the pediatric disease, with the G34R/V and K27M variants appearing to represent different clinicopathological and biological subgroups. This has been recognized by the World Health Organization (WHO) classification of CNS tumors, with the latest version including the novel entity, diffuse midline glioma with H3K27 mutation (Louis et al., 2016). Further refinements incorporating other clearly delineated subsets of the disease in future iterations appear likely and might prove clinically useful.
In addition to these uniquely defining histone mutations, detailed molecular profiling has served to identify numerous targets for therapeutic interventions. These include known oncogenes in adult glioma and other tumors with an elevated frequency in the childhood setting (e.g., **PDGFRA** (Paugh et al., 2013; Puget et al., 2012) or certain rare histological variants (e.g., **BRAF** V600E) (Nicolaides et al., 2011; Schiffman et al., 2010), as well as others seemingly unique to DIPG (e.g., **ACVR1** (Buczkwicz et al., 2014; Fontebasso et al., 2014; Taylor et al., 2014a; Wu et al., 2014)). Future trials will need to exploit these targets, but also incorporate innovative designs that allow for selection of the patient populations within the wide spectrum of disease who are most likely to benefit from any novel agent (Jones et al., 2016).

Despite these advances, driven by the efforts of several international collaborative groups to collect and profile these rare tumors, individual publications remain necessarily modestly sized, involving a range of different platforms and analytical techniques. This leaves certain subgroups poorly represented across studies, widely differing individual gene frequencies in different cohorts, and an inability to draw robust conclusions across the whole spectrum of the disease. We have therefore gathered together publicly available data, supplemented with 157 new cases, in order to provide a statistically robust, manually annotated resource cohort of >1,000 such tumors for interrogation.

**RESULTS**

**Sample Cohort**

In total, we obtained data from clinically annotated high-grade glioma (WHO, 2007, grade III or IV) or DIPG (radiologically diagnosed, grades II–IV) in 1,067 unique cases (Figure S1A). These were predominantly from children but also included young adults, in order to capture more HF3A G34R/V mutations, as well as to explore an otherwise under-studied population. There was a median age at diagnosis of 9.8 years, and 982 cases aged 21 years or younger (Table S1). These included 910 taken from 20 published series (Barrow et al., 2011; Bax et al., 2010; Buczkwicz et al., 2014; Carvalho et al., 2014; Castel et al., 2015; Fontebasso et al., 2013, 2014; Grasso et al., 2015; International Cancer Genome Consortium PedBrain Tumor Project, 2016; Khuong-Quang et al., 2012; Korshunov et al., 2015; Paugh et al., 2010, 2011; Puget et al., 2012; Schwartzentruber et al., 2012; Sturm et al., 2012; Taylor et al., 2014a; Wu et al., 2012; Wu et al., 2014; Zargooshi et al., 2010) and 157 unpublished cases. The vast majority of samples were obtained pre-treatment (biopsy or resection, n = 913), as opposed to post-therapy (relapse or autopsy, n = 146). Samples were classified as occurring within the cerebral hemispheres (n = 482), brainstem (n = 323 in pons, of which 322 were DIPG; three additional cases were in the midbrain and one in the medulla) or other non-brainstem midline locations (n = 224, predominantly thalamus, but also cerebellum, spinal cord, ventricles, and others; referred to as “midline” for simplicity throughout) (Figure 1A). There was a significant association of anatomical location with age of diagnosis, with medians of 13.0 years for hemispheric, 10.0 years for midline, and 6.5 years for DIPG, respectively (p < 0.0001, ANOVA; all pairwise comparisons adjusted p < 0.0001, t test) (Figure 1B), in addition to clinical outcome, with a median overall survival of 18.0 months for hemispheric tumors (2 year overall survival 32%), 13.5 months for midline (2 year overall survival 21.4%), and 10.8 months for DIPG (2 year overall survival 5.2%; p < 0.0001 for all pairwise comparisons, log rank test) (Figure 1C). Children 3 years of age and younger had a markedly improved clinical outcome (p = 0.0028, log rank test), although this benefit was largely restricted to children 1 year and under (n = 40, 2 year survival 61%, p < 0.0001, log rank test), with this association significant in all anatomical locations (p = 0.0402, hemispheric; p < 0.0001, midline; p = 0.00286, pons, log rank test). There were, however, proportionally fewer midline and pontine tumors in <1-year-olds compared with 1- to 3-year-olds (12/40, 30.0% versus 46/85, 45.9%, p = 0.0131 Fisher’s exact test) (Figures S1B and S1C).

**Molecular Subgrouping**

Hotspot mutation data for the genes encoding histone H3 were available or newly generated for 903 cases. At minimum, this included Sanger sequencing for **H3F3A** (H3.3) and **HIST1H3B** (H3.1); however, the absence of wider screening or next-generation sequencing data for 310 cases annotated as H3 wild-type (WT) means we cannot rule out rare variants in other H3.1 or H3.2 genes in those cases. In total, the cohort comprised 67 H3.3G34R/V (n = 63 G34R, n = 4 G34V), 316 H3.3K27M, 66 H3.1/3.2K27M (n = 62 HIST1H3B, n = 2 HIST1H3C, n = 2 HIST2H3C), and 454 WT. There were profound distinctions in anatomical location (p < 0.0001, Fisher’s exact test) (Figure 1D), age at diagnosis (p < 0.0001, ANOVA; all pairwise comparisons adjusted p < 0.0001, t test) (Figure 1E), and overall survival (p < 0.0001, log rank test) (Figure 1F). H3.3G34R/V tumors were almost entirely restricted to the cerebral hemispheres (accounting for 16.2% total in this location, particularly parietal and temporal lobes), were found predominantly in adolescents and young adults (median 15.0 years), and had a longer overall survival compared with other H3 mutant groups (median 18.0 months, 2 year overall survival 27.3%; p < 0.0001 versus H3.3K27M, p = 0.00209 versus H3.1H27M, log rank test). H3.3K27M were spread throughout the midline and pons, where they account for 63.0% DIPG and 59.7% non-brainstem midline tumors. In all locations (including ten cases reported to present in the cortex), these mutations conferred a significantly shorter time to death from disease (overall median 11 months, 2 year overall survival 4.7%) (Figures S1D–S1F). H3.1/3.2K27M were highly specific to the pons (21.4% total) where they represent a younger age group (median 5.0 years) with a significantly longer overall survival (median 15.0 months) than H3.3K27M (p = 0.00017, log rank test) (Figure S1F). In multivariate analysis incorporating the histone mutations alongside age, WHO grade, and gender, K27M mutations in both H3.3 and H3.1 are independently associated with shorter survival (p < 0.0001, Cox proportional hazards model).

**BRAF** V600E status was available for 535 cases, with mutant cases (n = 32, 6.0%) present only in midline and hemispheric locations, and conferring a significantly improved prognosis (2 year survival 67%, p < 0.0001, log rank test) (Figures S1G–S1I). There was additional annotation for **IDH1** R132 mutation status in 640 cases (n = 40, 6.25%), representing a forebrain-restricted, significantly older group of patients (median 17.0
Figure 1. Clinicopathological and Molecular Subgroups of pHGG/DIPG
(A) Anatomical location of all high-grade glioma cases included in this study, taken from original publications (n = 1,033). Left, sagittal section showing internal structures; right, external view highlighting cerebral lobes. Hemispheric, dark red; non-brainstem midline structures, red; pons, pink. Radius of circle is proportional to the number of cases. Lighter shaded circles represent a non-specific designation of hemispheric, midline, or brainstem.
(B) Boxplot showing age at diagnosis of included cases, separated by anatomical location (n = 1,011). The thick line within the box is the median, the lower and upper limits of the boxes represent the first and third quartiles, and the whiskers 1.5 x the interquartile range. ***Adjusted p < 0.0001 for all pairwise comparisons, t test.
(C) Kaplan-Meier plot of overall survival of cases separated by anatomical location, p value calculated by the log rank test (n = 811).
(D) Anatomical location of all cases separated by histone mutation (top, n = 441) and histone WT (bottom, n = 314). Left, sagittal section showing internal structures; right, external view highlighting cerebral lobes. Blue, H3.3G34R/V; green, H3.3K27M; dark green, H3.1K27M. Radius of circle is proportional to the number of cases. Lighter shaded circles represent a non-specific designation of hemispheric, midline, or brainstem.
(E) Boxplot showing age at diagnosis of included cases, separated by histone mutation (n = 753). The thick line within the box is the median, the lower and upper limits of the boxes represent the first and third quartiles, and the whiskers 1.5 x the interquartile range. ***Adjusted p < 0.0001 for all pairwise comparisons, t test.
(F) Kaplan-Meier plot of overall survival of cases separated by histone mutation, p value calculated by the log rank test (n = 693). See also Figure S1 and Table S1.
years, p < 0.0001, t test) with longer overall survival (2 year survival 59%, p < 0.0001, log rank test) (Figures S1J–S1L).

For 441 cases, Illumina 450k methylation BeadArray data was available, which provides robust classification into clinically meaningful epigenetic subgroups marked by recurrent genetic alterations (Korshunov et al., 2015, 2017). We used the Heidelberg brain tumor classifier to assign tumors into following subgroups: H3G34R/V (n = 51), H3K27M (n = 119), HGG WT (n = 156), IDH1 (n = 36), low-grade glioma (LGG)-like (n = 27), pleomorpho-xanthoastrocytoma (PXА)-like (n = 43), and “other” (n = 9) (Figure S2A), visualized by hierarchical clustering (Figure 2A) (Table S2). As reported previously, these subgroups have profound differences in anatomical location (Figure 2B), age at diagnosis (p < 0.00001 ANOVA) (Figure 2C), and overall survival (p < 0.0001, log rank test) (Figure 2D), with LGG-like group representing a younger cohort (median 4.0 years, 10/16 infant cases under 1 year, p < 0.0001 Fisher’s exact test) with excellent prognosis (2 year survival 74%, p < 0.0001 versus WT, log rank test), while the PXА-like group are enriched for BRAF V600E mutations (19/34, 56%) and carry an intermediate risk (median 38 months, 2 year survival 56%, p = 0.00423 versus WT, log rank test). After removing the PXА- and LGG-like groups, the remaining histone H3/IDH1 WT tumors had a 2 year survival of 23.5% (median overall survival 17.2 months). MGMT promoter methylation was significantly enriched in the H3G34R/V (65.1%, globally hypomethylated) and IDH1 (78.1%, globally hypermethylated)-like groups, and largely absent from H3K27M tumors (4.5%, all tests versus rest, p < 0.0001 Fisher’s exact test) (Figure S2B). Total methylation was lowest in H3G34R/V (median beta value 0.452), and highest in the IDH subgroup (median beta value 0.502), as reported previously (Sturm et al., 2012); however it was also found to be significantly elevated in PXА-like tumors (median beta value 0.501, p < 0.0001 t test) (Figure S2C).

DNA Copy Number

High-quality DNA copy-number profiles were obtained from 834 unique cases of pHGG/DIPG, taken from BAC and oligonucleotide arrays (n = 112), SNP arrays (n = 128), 450k methylation arrays (n = 428), and whole-genome or exome sequencing (n = 325) (Table S3). Clustering on the basis of segmented log2 ratios highlighted some of the defining chromosomal features of the pediatric disease, including recurrent gains of chromosome 1q, and losses of chromosomes 13q and 14q (Figure 3A). There are also a significant proportion of tumors (n = 147, 17.6%) with few if any DNA copy-number changes, with no bias toward lower-resolution platforms (p = 0.134, Fisher’s exact test), and the presence of other molecular markers obviating concerns of a substantial normal tissue contamination. These cases were found throughout the CNS, were younger at diagnosis (7.0 versus 10.3 years, p < 0.0001, t test) and had a longer overall survival (median 18.0 versus 14.0 months, p = 0.0107 log rank test) (Figure S3B). Common large-scale chromosomal alterations with prognostic significance included loss of 17p (n = 156), which targets TP53 at 17p13.1 and confers a shorter overall survival in tumors of all locations and all subgroups (Figure S3B), and gains of 9q (n = 108), more broadly encompassing a region of structural rearrangement on 9q34 in medulloblastoma (Northcott et al., 2014), and correlating with shorter overall survival in multiple pHGG/DIPG subgroups (Figure S3C).

We used GISTIC genomic identification of significant targets in cancer) in order to determine subgroup-specific copy-number drivers based on focality, amplitude, and recurrence of alterations. The most common focal events were the previously described high-level amplifications (Figure 3B) at 4q12 (PDGFRA/KIT/KDR, n = 77), 2p24.3 (MYCN/ID2, n = 42), chromosome 7 (7p11.2 (EGFR, n = 32), 7q21.2 (CDK6, n = 14), and 7q31.2 (MET, n = 19) (Figures S3D–S3F), as well as focal deletions (Figure 3C) at 9p21.3 (CDKN2A/CDKN2B, n = 102) (Figure S3G). Amplifications conferred a shorter overall survival, and CDKN2A/CDKN2B deletion a better prognosis, either across the whole cohort or selected subgroups (Figures S3B–S3G). In addition, the aggregated data identified less-frequent alterations, recurrent across multiple studies, identifying pHGG/DIPG candidates including NF1 (nuclear factor I B, n = 77), 9p23-p22.3, n = 4), GAB2 (GRB2-associated binding protein 2, 11q14.1, n = 4), SMARCE1 (SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily e, member 1, 17q21.2, n = 4), and others (Figures 3B and 3C).

Subgroup-Specific Alterations

When IDH1-mutant tumors were removed and the cohort restricted to those cases for which histone H3 status was available, we were able to investigate subgroup-specific DNA copy-number changes in 705 pHGG/DIPG (Figure 4A). Applying GISTIC within these case sets revealed specific focal events enriched within individual subgroups, including AKT1 amplifications in H3G34R/V, MYC and CCND2 amplification in H3K27M, and MYCN/ID2, MDM4/PK3C2B, and KRAS amplification in H3 WT (Figure 4B) (Table S4). These latter events were generally restricted to hemispheric tumors, while MYCN/ID2 were enriched in H3 WT DIPG (Figure S4A). H3.1K27M tumors generally lacked amplifications/deletions, but were instead characterized by frequent gains of 1q and the whole of chromosome 1q.

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Figure 2. Methylation-based Subclassification of pHGG/DIPG
(A) Unsupervised hierarchical clustering and heatmap representation of (i) values for 441 samples profiled on the Illumina 450k BeadArray platform (red, high; blue, low). Samples are arranged in columns clustered by most variable 1,381 classifier probes. Age at diagnosis is provided below. Clinicopathological and molecular annotations are provided as bars according to the included key.
(B) Anatomical location of methylation-defined PXА-like (n = 43) and LGG-like (n = 27) cases. Left, sagittal section showing internal structures; right, external view highlighting cerebral lobes. Dark gold, PXА-like; tan, LGG-like. Radius of circle is proportional to the number of cases. Lighter shaded circles represent a non-specific designation of hemispheric, midline, or brainstem.
(C) Boxplot showing age at diagnosis of included cases, separated by simplified methylation subclass (n = 440). The thick line within the box is the median, the lower and upper limits of the boxes represent the first and third quartiles, and the whiskers 1.5 x the interquartile range. **Adjusted p < 0.0001 for all H3 G34R/V pairwise comparisons, t test; **adjusted p < 0.01 for LGG-like versus WT, t test.
(D) Kaplan-Meier plot of overall survival of cases separated by simplified methylation subclass, p value calculated by the log rank test (n = 307). See also Figure S2 and Table S2.
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2, and the loss of 16q (Figure 4C). PXA-like tumors had frequent CDKN2A/B deletions and a unique loss at 1q, associated with shorter overall survival within this group (Figure S4B).

Whole-arm losses were also enriched in H3.3G34R/V tumors, specifically 3q, 4q, 5q, and 18q, where smallest regions of overlap were in some instances able to narrow the common region to a handful of candidate genes (Figure S4C). On chromosome 4q this appeared to target FBXW7 at 4q31.3, also aligning with the GISTIC data (Figure 5A). Across three independent platforms, gene expression over the whole arm was significantly lower when 4q was lost (Agilent, p = 0.00231; Affymetrix, p = 0.000102; RNA sequencing (RNA-seq), p = 0.0398; Wilcoxon signed-rank test) (Figure S5). (Table S5).

There were also four patients with three different somatic mutations identified (below), two truncating and one missense, three of which were in hemispheric tumors, and two with H3F3A G34R (Figure 5B).

In cases with 4q loss, median FBXW7 gene expression was reduced compared with those with normal copy number (Agilent, p = 0.029; Affymetrix, p = 0.015; RNA-seq, p = 0.4; Mann-Whitney U test) (Figure 5C).

Within H3.3K27M tumors, we identified a recurrent amplification at 17p11.2 (n = 17; 170 kb to 11.96 Mb), across multiple platforms and significantly enriched in DIPGs, which appears to target TOP3A within these tumors (Figure 5D). Where available (n = 6) (Figure S5B), whole-genome sequencing data reveals this occurs via complex intra- and inter-chromosomal rearrangements (Figure 5E) leading to increased mRNA expression of TOP3A in amplified versus non-amplified cases in Agilent (n = 1), and Affymetrix and RNA-seq (p = 0.011 and p = 0.016, respectively, Mann-Whitney U test) data (Figure 5F) (Table S5).

In an integrated dataset, TOP3A was the most differentially expressed gene in the region in amplified cases (adjusted p = 0.00856 Mann-Whitney U test). We further identified a single somatic missense mutation (C685Y) in an additional case of DIPG, and, taken together, TOP3A alterations were mutually exclusive with ATRX deletion/mutations found in H3.3K27M DIPG (0/13).

Whole-Genome and Exome Sequencing

Out of 372 sequenced cases (n = 118 whole genome, n = 247 exome, 7 both), we were able to retrieve raw data from 351 for integration of somatic variant calling (Table S6) and DNA copy-number changes. Of these, RNA-seq data was available for 47, allowing for candidate fusion gene nomination in 150 cases (RNA-seq or whole-genome sequencing restricted to high-confidence nominations in relevant pathway-associated genes, Table S6). Taking a conservative approach to variant calling given the disparate sequencing coverage (median 88x, range 16–295x), capture platforms, and availability of germline data, we report a median number of somatic single nucleotide variants (SNVs) and insertion/deletions (InDels) of 12, with 97% cases in the range 0–305. DNA copy neutral cases had significantly fewer somatic mutations (median 8.37 versus 21.32 SNVs/InDels per case), with those copy neutral cases also having no detectable mutations falling into the youngest age group (median = 3.9 years).

There was only a modestly elevated mutation rate between samples taken post- compared with pre-treatment (1.5-fold, p = 0.056, t test) (Figure S6A). However, 11 cases had a vastly increased mutational burden, described as a hypermutator phenotype (median 13,735 SNVs/InDels, range 852–38,250), with distinctive mutational spectra from non-hypermutated pHGG/DIPG (Shlien et al., 2015), including three cases with plausible somatic activating POLE mutations (Figure S6B).

InD1H1-mutant cases were again excluded (n = 14), with genetic profiles identical to that described in adults for astrocytic tumors (13/14 TP53, 7/14 ATRX mutations), and oligodendrogial tumors (1p19q co-deletion, TERT promoter, CIC, FBUBP1 mutations) entirely absent (Figure S6C). We were thus left with an integrated dataset of 326 pHGG/DIPG, providing robust annotation of the most frequently altered genes across histone H3 subgroups and anatomical locations (Figure 6A). As well as known associations such as hemispheric H3.3G34R/V and TP53/ATRX (18/20, 90%; p = 0.0001), midline H3.3K27M and FGFR1 (8/39, 20.5%; p = 0.212, not significant), pontine H3.1K27M and ACVR1 (28/33, 84.8%; p = 0.0073), and PXA-like GBM with BRAF V600E (17/28, 60.7%; p < 0.0001), we also identified previously unrecognized co-segregating mutations including H3.3G34R/V and ARID1B (2/20, 10%; p = 0.0720), H3.3K27M DIPG and ATM and ASXL1 (5/93, 10.7%; p = 0.0473), and H3.1K27M and BCOR (6/37, 16.2%; p = 0.0022, all Fisher’s exact test) (Figure S6D). We also identified recurrent events in genes such as PTPN23 (protein tyrosine phosphatase, non-receptor type 23, n = 5), SOX10 (SRY-box 10, n = 5), SRCAp (Snf2-related CREBBP activator protein, n = 5), DEPDC5 (DEP domain-contain 5, member of GATOR complex, n = 4), SGK223 (PEAK1-related kinase activating pseudokinase, n = 4), and others (Figure 6B).

Integrated Pathway Analysis

Many of the rare variants we identified (Figure S6E) were found in genes associated with intracellular signaling pathways and processes more commonly targeted by high-frequency events, often in a mutually exclusive manner. In total, 297/326 (91.1%) of cases were found to harbor genetic alterations in one or more of nine key biological processes (Figure 7A). These included well-recognized pathways such as DNA repair (198/326, 60.7%), largely driven by TP53 mutations (n = 160), but also by common mutually exclusive (p < 0.0001, Fisher’s exact test) activating truncating alterations in PPM1D (n = 18), as well as heterozygous mutations in a diverse set of genes including those involved in homologous recombination (ATM, BRC2, BLM, ATR, PALB2, RAD50, and RAD51C) and numerous Fanconi anemia genes (BRIP1, FANCN, FANCA, and FANCG), among others (Figure S7A). Although TP53 is almost always found in concert with H3.3G34R/V in the cerebral hemispheres, these additional DNA repair pathway mutations were enriched in...
H3K27M DIPG (36/68, 52.9%). Also co-segregating with H3.3G34R/V and TP53 is ATRX, although mutations/deletions of the latter gene are also frequently found in conjunction with H3K27M (28/54, 51.8%). ATRX accounts for a large proportion of the cases harboring mutations in genes coding for chromatin modifiers (54/118, 45.8%); however, there is a diverse set of readers, writers, and erasers also targeted at lower frequency, especially in DIPG, including the previously mentioned BCR (n = 14) and ASXL1 (n = 6) in addition to SETD2 (n = 8), KDM6B (n = 6), SETD1B (n = 5), and ARID1B (n = 5) among many others (Figure S7B).

While CDKN2A/CDKN2B deletions were almost entirely absent from DIPG (1/154, 0.65%), dysregulation of the G1/S cell-cycle checkpoint was common throughout anatomical locations and subgroups (82/326, 25.2%), with amplifications of CCND2 and deletions of CDKN2C predominating in the pons (n = 5/7 and 5/5 DIPG, respectively), in contrast to recurrent homozygous RB1 deletions and CDK6 amplifications (n = 6/7 and 4/6 hemispheric) (Figure S4A) (Figure S7C).

Subgroup-specific dysregulation was also observed when considering discrete components of the RTK-PI3K-MAPK pathway. In total, 201/326 (61.7%) cases harbored alterations in any given node; however, for H3.3G34R/V this was predominantly at the RTK level (11/20, n = 9 PDGFR) (Figure S7D), whereas H3.1K27M cases were enriched for PI3K/mTOR alterations (17/37, n = 9 PIK3CA, n = 5 PIK3R1) (Figure S7E), and H3 WT cases harbored the highest frequency of MAPK alterations (mainly BRAF V600E in PXA-like, n = 5/10 plus one NF1) (Figure S7F). NRTK1-NRTK3 fusions were enriched in the infant group (4/6 fusions under 1 year old, median age 3.25 versus 8.5 years, p = 0.00033, t test) (Figure S7D). We further identified mutations in genes regulating mTOR signaling, including TSC2 (n = 3), RPTOR, and MTOR itself (both n = 2), as well as a diverse series of SNVs and fusion candidates in MAPKs across all subgroups and locations (MAP2K7, MAP3K15, MAP3K4 and others) (Figure S7F).

BMP signaling was significantly enriched in H3.1K27M DIPG due to the strong correlation with ACVR1 mutations; however, alterations in other pathway members such as amplification of ID2 (n = 10) or ID3 (n = 3) and mutations in BMP3 (n = 5), BMP2K (n = 3), and others across locations and subgroups, extends the proportion of tumors for which this pathway may be relevant (62/326, 18.7%) (Figure S7G). There was also a subset of cases harboring alterations in members of the WNT signaling pathway (16/326, 4.9%), including AMER1, APC (both n = 3), and WNT8A, WNT9A, PLAGL2, and TCF7L2 (all n = 2) (Figure S7H).

Uniquely, the accumulated data uncovered a series of additional processes involved in maintenance of DNA replication, genome integrity, or transcriptional fidelity, targeted by infrequent but mutually exclusive alterations in pHGG and DIPG. These included mutations in splicing factors (SF3A1, SF3A2, SF3A3, SF3B1, SF3B2, and SF3B3, total n = 10), sister chromatid segregation (STAG2, STAG3, and ESPL1, total n = 9), pre-mRNA processing (Dicer and Drosha, total n = 4), DNA polymerases (PolK, PolQ, and PolR1B, total n = 4), as well as genes involved in centromere (CENPB, n = 3) and telomere maintenance (PML, n = 2; TERT, n = 7) (Figure S7I). TERT promoter mutations were found in 5/326 (1.5%) cases; however, alternative lengthening of telomeres (ALT) status was only available for 26 cases, although the 5 ALT-positive samples (19.2%) were mutually exclusive with TERT alterations.

We incorporated the integrated dataset into a pathway enrichment analysis (significant gene sets, false discovery rate [FDR] < 0.05, visualized as interaction networks by Cytoscape Enrichment Map) in order to gain additional insight into dysregulated biological processes. In addition to the subgroup-specific differential targeting of distinct nodes within common signaling pathways already described (e.g., RTK, PI3K/mTOR, and MAPK), additional dysregulated processes across the diversity of the disease were identified (Figure 7B). This revealed the perhaps not unexpected dysregulation of numerous developmental and CNS-associated gene sets (various immature organ systems, neuronal communication), but also previously unrecognized areas such as nuclear transport, cell migration, and the immune response (Table S7), which may provide further insight into disease biology as well as represent potential therapeutic strategies targeting key regulators of tumor phenotype. Indeed, neuronal communication with pGBM and DIPG cells is a recently demonstrated microenvironmental driver of pediatric glioma growth (Qin et al., 2017; Venkatesh et al., 2015).

**Histone H3/IDH1 WT Subgroups**

Finally, we wanted to explore those cases absent of any histone H3 or IDH1 mutations in more depth. Using a t statistic-based stochastic neighbor embedding projection of the 450k methylation data, we identified three distinct clusters of tumors separate from the G34, K27, and IDH1 groups (Figure 8A). Consensus clustering of the H3/IDH1 WT cases alone confirmed the presence of three robust subgroups (Figure 8B), which were also recapitulated by unsupervised hierarchical clustering of the 10,000 probe classifier subset (Figure 8C). These groups included a largely hemispheric set of tumors containing, but not restricted to, the PXA- and LGG-like subgroups (WT-A). These tumors were driven by BRAF V600E, NF1 mutations, or fusions in RTKs including MET, FGFR2, and NTRK2,3 (Figure 8D). Although including many younger patients, the ages varied widely (Figure 8E). Regardless, this group had the best overall survival (median = 63 months, p < 0.0001 versus rest, log rank test) (Figure 8F), with the non-PXA/LGG-like tumors within this group themselves having an extended median survival time of...
Figure 5. Alterations Targeting FBXW7 in H3.3G34R/V pHGG and TOP3A in H3.3K27M DIPG

(A) Segmented exon-level DNA copy-number heatmaps for 4q loss in H3.3G34R/V tumors (dark red, amplification; red, gain; dark blue, deletion; blue, loss; n = 28). An ideogram of chromosome 4 is provided indicating enlarged genome browser view and genes within common regions targeted across samples (gray). Clinicopathological and molecular annotations are provided as bars according to the included key.

(B) Cartoon representation of amino acid position for four somatic mutations found in FBXW7, colored by annotated functional domains and numbers provided for recurrent variants.

(C) Boxplots representing gene expression differences between FBXW7 lost/mutated cases (blue) and those with normal copy/WT (gray) in three independent gene expression platform datasets. The thick line within the box is the median, the lower and upper limits of the boxes represent the first and third quartiles, and the whiskers 1.5 times the interquartile range.

(D) Segmented exon-level DNA copy-number heatmaps for 17p11.2 amplification in predominantly H3.3K27M DIPG (dark red, amplification; red, gain; dark blue, deletion; blue, loss; n = 17). Chromosome 17 ideogram is provided indicating enlarged genome browser view and genes within common regions targeted across samples (gray). Clinicopathological and molecular annotations are provided as bars according to the included key.

(legend continued on next page)
38 months (p = 0.00928 versus other H3/IDH1 WTs, log rank test). Taking an integrated gene expression profiling dataset (Figures S8A–S8E), these tumors were found to have upregulation of gene signatures associated with cytokine signaling and cell junction organization (Figures S8F and S8G). A second group of tumors (WT-B) were found in all anatomical compartments, and were distinguished by chromosome 2 gains (Figure 8C) and, most notably, by high-level amplifications in EGFR, CDK6, and MYCN (p = 0.00033, p = 0.0299, p = 0.00037, respectively, Fisher’s exact test), with an imperfect overlay to the classifier “GBMpedRTK” and “GBM_MYCN” groups (Figure 8D). This group had strong upregulation of MYC target genes, and had the poorest overall survival (median = 14 months) (Figure 8F). The remaining cases encompassed a methylation classifier group described as “HGG_MID,” although in fact were split 80:20 hemispheric:midline (WT-C) (Figure 8C). This group was enriched for chromosome 1p and 20q loss, 17q gain (p = 0.00595, p = 0.0286, p = 0.0478, respectively, Fisher’s exact test) (Figure 8C), harbored PDGFRα and MET amplifications (p = 0.0159, Fisher’s exact test) (Figure 8D), and was strongly associated with the adult GBM-defined “Proneural” gene signature. These patients had a median survival of 18 months.

Although there remain tumors without detectable genetic alterations, we are nonetheless able to assign clinically meaningful subgroups with plausible driver alterations to the vast majority of pediatric HGG/DIPG.

DISCUSSION

Integrated molecular profiling has revolutionized the study of diffusely infiltrating high-grade glial tumors in children, providing evidence for unique mechanisms of molecular pathogenesis reflecting their distinct developmental origins (Baker et al., 2015; Jones and Baker, 2014). Although they are relatively rare, the present study accumulates 1,067 unique cases, a number similar to the aggregated analysis of the The Cancer Genome Atlas adult LGG/GBM cohorts (n = 1122, with grade III included in the “lower-grade” series) (Ceccarelli et al., 2016). Although there are clearly the usual caveats with such retrospective analyses of inconsistently annotated and treated cases, the cohort appears to represent a clinically useful approximation of the diversity of the pHGG/DIPG population.

In adults, the key distinction is between IDH1 mutant (G-CIMP/ATRX/TP53 or 1p19q co-deleted/TERT promoter mutated) and WT (classical, mesenchymal, PA-like) (Ceccarelli et al., 2016), whereas in the childhood setting IDH1 mutations were restricted to a small proportion (6.25%) of tumors mostly in adolescents (representing the tail end of an overwhelmingly adult disease), and harbored only rare examples of the common alterations seen in WT adult GBM (e.g., 4.9% EGFR mutation/amplification). Instead, most prominent among the differences between pediatric and adult studies is the frequency of hotspot mutations in genes encoding histone H3 variants: 2/820 (0.2%) in adults (Ceccarelli et al., 2016) versus 449/893 (50.3%) in the present pHGG/DIPG series.

The importance of recurrent H3 mutations in the childhood setting has become increasingly clear since their unexpected discovery in 2012 (Schwartzentruber et al., 2012; Wu et al., 2012), with clear clinicopathological differences associated with distinct variants (Jones and Baker, 2014; Jones et al., 2016; Sturm et al., 2014), and fundamental insights into mechanisms of epigenetically linked tumorigenesis (Bender et al., 2013; Bjerke et al., 2013; Chan et al., 2013; Funato et al., 2014). Despite this, precisely how we can target these mutations clinically remains elusive (Grasso et al., 2015; Henrika et al., 2017). Data from such a large series of tumors demonstrates the robustness of the histone-defined subgroups in terms of anatomical location, age of incidence, clinical outcome, methylation and gene expression profiles, copy-number changes, co-segregating somatic mutations, and pathway dysregulation. As most of the non-histone molecular alterations previously reported in pHGG/DIPG have been relatively infrequent, it is only through this accumulated dataset that we have been able to uncover subgroup-specific genes/processes that may play a role as diagnostic, prognostic, or predictive markers or drug targets in these diseases.

H3.3G34R/V-mutant tumors are restricted to the cerebral hemispheres and co-segregate with ATRX and TP53 mutations; they are also the only pediatric subgroup to harbor frequent MGMT promoter methylation (Korshunov et al., 2015). Copy-number profiling of 63 cases highlighted a significant enrichment of chromosomal arm losses at 3q, 4q, 5q, and 18q, further refined by smallest region of overlap and GISTIC analyses. At 4q31.3, this identified FBXW7 as a candidate gene target of the loss. FBXW7 encodes a member of the F box protein family and is frequently deleted/mutated in cancer, supporting its tumor-suppressive function (Davis et al., 2014); notably in relation to H3.3G34R/V it has been reported to play a role in MYC/MYC stabilization through its action as a component of the SCF-like ubiquitin ligase complex that targets MYC/MYC for proteosomal degradation (Welcker et al., 2004; Yada et al., 2004). With MYCN upregulated in H3.3G34R/V tumors through differential H3K36me3 binding (Bjerke et al., 2013), this observation adds to the mechanisms by which Myc proteins exert their influence in this subgroup, and provide further rationale for the observed effects of disrupting these interactions, such as with Aurora kinase A inhibitors which target the direct interaction between the catalytic domain of Aurora A and a site flanking Myc Box I that also binds SCF/FbxW7 (Richards et al., 2016).

H3.3K27M tumors are found in two-thirds of DIPG and non-brainstem midline pHGG alike, where they are associated with a shorter overall survival in both locations, as well as in the small number of cases reported in the cortex. Although presumably reflecting a common or overlapping origin, the pattern of co-segregating mutations differ, e.g., PdgfRa alterations predominating.
Figure 6. Somatic Mutations in pHGG/DIPG

(A) Oncoprint representation of an integrated annotation of somatic mutations and DNA copy-number changes for the 30 most frequently altered genes in 326 pHGG/DIPG (n > 6, frequency barplot on the right). Selected common fusion events are also shown where available. Samples are arranged in columns with genes labeled along rows. Age at diagnosis is provided below. Underneath, barplots are provided on a log 10 scale for numbers of copy-number aberrations and somatic mutations per case. Clinicopathological and molecular annotations are provided as bars according to the included key.

(B) Barplot of all recurrent somatic mutations across all 326 cases, in order of frequency, and colored independently by both anatomical location and histone mutation. See also Figure S6 and Table S6.
in the pons, and FGFR1 variants being largely restricted to the thalamus (Fontebasso et al., 2014). Our analysis of more than 300 cases further identifies differential amplification of CCND2 (DIPG) and CDK4 (non-brainstem midline), and, most strikingly, an amplification at 17p11.2 involving TOP3A in H3.3K27M DIPG. This complex rearrangement often involves loss of the more distal part of 17p involving TP53, along with intra- or inter-chromosomal translocations to deliver an increase in TOP3A copy number and gene expression. TOP3A encodes DNA topoisomerase III alpha, which forms a complex with BLM (Wu et al., 2000), has an important role in homologous recombination (Yang et al., 2010), and has been implicated in maintenance of the ALT phenotype (Temime-Smaali et al., 2009). Notably, TOP3A amplification/mutation was found to be mutually exclusive with ATRX mutation in H3.3K27M DIPG, with depletion by small interfering RNA reducing ALT cell survival (Temime-Smaali et al., 2008), and therefore represents a potential therapeutic target in this subgroup.
Figure 8. Integrated Analysis of H3/IDH1 WT pHGG/DIPG

(A) t-Statistic-based stochastic neighbor embedding (t-SNE) projection of the combined 450k methylation dataset (n = 441). The first three projections are plotted in the x, y, and z axes, with samples represented by dots colored by histone H3G34 (blue), H3K27 (green), IDH1 (red), PXA-like (dark gold), LGG-like (tan), and “others” (gray).

(B) K means consensus clustering on the H3/IDH1 WT cases highlights three stable clusters (left, black/brown [WT-A], gray/pink [WT-B], and dark cyan [WT-C]) as the most robust subdivision of the data (right, area under the curve analysis for different cluster numbers).

(C) Unsupervised hierarchical clustering and attendant heatmap of the H3/IDH1 WT cases (n = 219). Samples are arranged in columns clustered by the most variable 1,521 classifier probes. Age at diagnosis is provided below. Clinicopathological and molecular annotations are provided as bars according to the included key.

(D) Oncoprint representation of an integrated annotation of somatic mutations and DNA copy-number changes for the H3/IDH1 WT cases (n = 50). Samples are arranged in columns with genes labeled along rows. Age at diagnosis is provided below. Clinicopathological and molecular annotations are provided as bars according to the included key.

(E) Boxplot showing age at diagnosis of H3/IDH1 WT subgroups, separated by anatomical location (n = 190). The thick line within the box is the median, the lower and upper limits of the boxes represent the first and third quartiles, and the whiskers 1.5× the interquartile range.

(F) Kaplan-Meier plot of overall survival of H3/IDH1 WT subgroups separated by anatomical location, p value calculated by the log rank test (n = 150). See also Figure S8 and Table S8.
H3.1K27M tumors by contrast are restricted to the pons, patients are younger and with a slightly longer survival (Castel et al., 2015), and are largely defined at the copy-number level by whole chromosomal arm gains and losses (Taylor et al., 2014a). They have the well-recognized association with ACVR1 mutation (Taylor et al., 2014b); however, we also identify an enrichment of downstream PI3K pathway mutations (PIK3CA and PIK3R1) in comparison with the largely upstream RTK alterations present in H3.1K27M DIPGs, important in designing stratified trials and combinatorial therapies. Further association with mutations of the BCL6 repressor gene BCO2, commonly altered in medulloblastomas, neuroepithelial tumors, and sarcomas, highlights a further avenue for intervention study through its regulation of the SHH pathway (Tiberi et al., 2014).

In H3/IDH1 WT cases, methylation profiling refines the heterogeneous collection of tumors, particularly identifying two predominantly hemispheric intermediate risk subgroups that classify alongside other entities (PXA- and LGG-like) in a larger series of better outcome tumors (WT-A). These had already been strongly linked with dysregulation of the MAPK pathway (BRAF V600E) (Korshunov et al., 2015) along with CDKN2A/CDKN2B deletion (Nicolaides et al., 2011). However, with molecular markers such as losses at 1q and 17p appearing to confer a worse outcome there may be more than one subgroup within this entity, and a co-clustering group of H3/IDH1 WT tumors appeared distinctly driven by somatic NF1 mutation. The LGG-like tumors generally occur in very young patients, where the appearance of few genetic alterations and a significantly better prognosis is shared by the majority of infant HGG. Gene fusion events, including those targeting NTRK1-NTRK3, are common in this age range. Notably this enhanced survival is restricted to patients diagnosed under 12 months of age, and is not recapitulated in the 1–3 year age group, although this is the common clinical definition of “infants” in many centers.

Excluding these morphologically high-grade but biologically and clinically low-grade tumors, the remaining H3/IDH1 WT cases can be further split into two poor-outcome groups driven by EGFR/MYC/CDK6 (WT-B) or PDGFRα/MEK (WT-C) or amplifications. These groups overlap with other methylation-based classification groups (PDGFRα versus EGFR versus MYCN (Korshunov et al., 2017); “GBM_pedRTK” versus “GBM_MYCN” versus “HGG_MID” (moleculareuropathology.org/mpn), however, are uniquely defined here spanning anatomical locations and integrated with sequencing data. Further exploration of these heterogeneous subgroups in order to refine integrated molecular diagnostics to prioritize patient subpopulations for stratified treatment remains a priority.

The remarkable biological diversity spanning pediatric malignant glioma is finally demonstrated by the <5% tumors with a hypermutator phenotype, some of the greatest mutational burdens in all human cancer, and candidates for immune checkpoint inhibitors (Bouffet et al., 2016). Previously unrecognized processes altered in small subsets of tumors identified through this meta-analysis, such as the splicing machinery, mRNA regulation, and the WNT pathway offer further areas for exploration. The thorough cataloging of dysregulated molecular pathways across the whole spectrum of pediatric diffusely infiltrating gliomas in the present study provides the basis for novel therapeutic development.

**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental Information includes eight figures and eight tables and can be found with this article online at http://dx.doi.org/10.1016/j.ccell.2017.08.017.

**AUTHOR CONTRIBUTIONS**


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the human genome. In this study, we aimed to identify novel genetic markers and therapeutic targets in pediatric glioblastoma.


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# STAR METHODS

## KEY RESOURCES TABLE

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CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Chris Jones (chris.jones@icr.ac.uk).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Patient Samples
All new patient material was collected after informed consent and subject to local research ethics committee approval. We collated and profiled 157 unpublished cases of HGG in children and young adults up to the age of 30 years at diagnosis obtained from the Royal Marsden, St Georges and Kings College Hospitals, (n=39, all London, UK), Chinese University of Hong Kong (n=24,
Hong Kong, China), Qilu University Hospital (n=23, Jinan, China), Farhad Hatched Hospital (n=14, Sousse, Tunisia), Federal University of São Paolo (n=14, São Paulo, Brazil), Morozov Children’s and Dmitri Rogachev Hospitals (n=12, Moscow, Russia), Queensland Children’s Tumor Bank (n=8, Brisbane, Australia), Hospital San Joan de Déu (n=6, Barcelona, Spain), City Hospital #31 (n=6, St Petersburg, Russia), Barretos Cancer Hospital (n=4, Barretos, Brazil), Centre Hospitalier Régional et Universitaire Hautepierre (n=3, Strasbourg, France), and Our Lady Children’s Hospital Crumlin (n=2, Dublin, Ireland). A full description of the samples included are provided in Table S1.

METHOD DETAILS

Nucleic Acid Extraction
DNA was extracted from frozen tissue by homogenisation prior to following the DNeasy Blood & Tissue kit protocol (Qiagen, Crawley, UK). DNA was extracted from formalin-fixed, paraffin-embedded (FFPE) pathology blocks after manual macrodissection using the QIAamp DNA FFPE tissue kit protocol (Qiagen). Matched normal DNA was extracted from blood samples using the DNeasy Blood & Tissue kit (Qiagen, Crawley, UK). Concentrations were measured using a Qubit fluorometer (Life Technologies, Paisley, UK). RNA was extracted by following the RNaseasy Mini Kit protocol (Qiagen), and quantified on a 2100 Bioanalyzer (Agilent Technologies).

Sanger Sequencing of H3F3A / HIST1H3B
PCR for H3F3A and HIST1H3B was carried out using primers obtained from Life Technologies (Paisley, UK). Products were purified using the QIAquick PCR purification kit (Qiagen), subjected to bidirectional sequencing using BigDye Terminator mix 3.1 (Applied Biosystems, Foster City, CA, USA), with capillary sequencing was done on an ABI 3100 genetic analyzer (Applied Biosystems, Foster City, CA, USA). Sequences were analysed using Mutation Surveyor (SoftGenetics, PN, USA) and manually with 4Peaks (Nucleobytes, Aalsmeer, Netherlands).

Methylation Profiling
50-500 ng DNA was bisulphite-modified and analyzed for genome-wide methylation patterns using the Illumina HumanMethylation450 BeadArray (450k) platform at either the DKFZ or the University College London Genomics Centre, according the manufacturer’s instructions. All samples were checked for expected and unexpected genotype matches by pairwise correlation of the 65 genotyping probes on the 450k array.

Exome and RNA Sequencing
50-500 ng DNA was sequenced at the Tumor Profiling Unit, ICR, London, UK using the SureSelect Human All Exon capture sets V4 or V5 (Agilent, Santa Clara, CA, USA), and paired-end-sequenced on an Illumina HiSeq2000 (Illumina, San Diego, CA, USA) with a 100 bp read length. Coverage ranged from 29-295x (median=105x). RNA was sequenced at the ICR Tumor Profiling Unit after SureSelect RNA capture on an Illumina HiSeq2500 with a 125 bp read length.

QUANTIFICATION AND STATISTICAL ANALYSIS

Published Data Sources
These data were combined with those obtained directly from the authors or from public data repositories representing 20 published studies (Barrow et al., 2011; Bax et al., 2010; Buczkowicz et al., 2014; Carvalho et al., 2014; Castel et al., 2015; Fontebasso et al., 2013, 2014; Grasso et al., 2015; International Cancer Genome Consortium PedBrain Tumor Project, 2016; Khuong-Quang et al., 2012; Korshunov et al., 2015; Paugh et al., 2010, 2011; Puget et al., 2012; Schwartzentruber et al., 2012; Sturm et al., 2012; Taylor et al., 2014a; Wu et al., 2012; Wu et al., 2014; Zarhoo et al., 2010) with the following accession numbers: EGA - EGAS00001000226, EGAS0000100192, EGAS00001000575, EGAS00001000720, EGAS00001001139; the Gene Expression Omnibus (www.ncbi.nml.nih.gov/geo/) - GSE19578, GSE26576, GSE21420, GSE34824, GSE36245, GSE36278 GSE50022, GSE50021, GSE50024, GSE55712; ArrayExpress - E-TABM-857, E-TABM-1107. The full cohort included a total of 1254 molecular profiles from 955 samples across 12 platforms, which after quality control and manual annotation to remove duplicates, and supplemented with targeted sequencing of an additional 158 cases, resulted in a total dataset comprised of 1067 individual patients. The full dataset comprises genomic profiles from DNA copy number arrays (Agilent 44K, n=127; Affymetrix 500K, n=100; Affymetrix SNP6.0, n=78; 32k BAC, n=61), Illumina 450K methylation arrays (n=441), whole exome (n=254), genome (n=125), targeted (n=212) and RNA sequencing (n=82), as well as gene expression from Affymetrix U133Plus2 (n=102) and Agilent WG2.5 (n=67) platforms.

DNA Copy Number
DNA copy number data was obtained as array CGH (Agilent 44k and 32K BAC), SNP arrays (Affymetrix 500K and SNP6.0), 450k methylation arrays (Illumina) and/or sequencing data (whole genome and exome). Two color CGH data was read and normalized using the R packages limma and marray. Log intensity data from Affymetrix SNP arrays was derived using the aroma.affymetrix and aroma.cn package. Combined log2 intensity data from Illumina 450K methylation arrays was processed using the R packages minfi and conumee. For sequenced samples, coverage of aligned reads was binned into known genes and exons with BEDTools and
log₂ ratios of median coverage in tumor and normal sequences were processed with in-house scripts. To combine copy number platforms, median log₂ ratios were recovered within all known genes and exons and normalized such that the median displacement of X in male:female comparisons was rescaled to an average of -1. Exon-level median log ratios and smoothed values were then combined across platforms and thresholded to call gains and losses above and below log₂ ratios of ±0.3 with a contiguous of ~1MB and amplifications and deletions above and below a threshold of ±1.5 with a minimum of 3 contiguous exons.

CBS binary segmentation from the DNAcopy package was applied to each dataset to provide smoothed log₂ ratios. Genes within common CNVs in normal individuals were excluded from further analysis with reference to the CNV map of the human genome. DNA copy number data was clustered based upon categorical states (deep deletion, loss, no change, gain and amplification) based upon the Euclidean distance method with a Ward algorithm. Gains and losses in chromosomal arms were called based upon contiguous regions covering more than one third of the exonic regions within each arm. For regions of focal copy number change cases carrying copy number alterations were ranked according to the length of the largest CNA in each case and are plotted as heatmaps aligned to precise genomic coordinates alongside genomic tracks based upon hg19 made with the R package gviz. Minimal regions of copy number alteration were assigned based on the frequency of categorical states within each region. Focal amplifications and deletions were identified in CBS segmented data using the GISTIC algorithm in MATLAB on the exon-level data, with thresholds for gain and loss of 0.3 and gene-level filters to remove regions of common copy number variation in normal individuals based on the CNV map of the human genome.

DNA Methylation
Methylation data from the Illumina Infinium HumanMethylation450 BeadChip was preprocessed using the minfi package in R. DNA copy number was recovered from combined intensities using the conumee package with reference to methylation profiles from normal individuals provided in the CopyNumber450kData package. We have used the Heidelberg brain tumor classifier (molecularneuropathology.org) to assign subtype scores for each tumor compared to 91 different brain tumor entities using a training set built from more than 2000 tumors implemented in the MNP R package. Simplified methylation subgroup assignments were then made to incorporate cases carrying G34RV or K27M mutations in H3 histones, IDH1 mutation at R132, low grade glioma-like profiles (predominantly diffuse infantile gangliogioma and pilocytic astrocytoma) and those similar to pleomorphic xanthoastrocytoma (PXA). Wild-type HGG encompassed many other methylation subgroups and were simply assigned by exclusion with the groups above. Clustering of beta values from methylation arrays was performed using the 10K probe set from the Heidelberg classifier based upon Euclidean distance with a ward algorithm. Methylation heatmaps show only the most variable probes of the classifier between simplified methylation subgroups. Overall methylation was calculated as the mean of the 10K classifier probe set for each subgroup and MGMT promoter methylation was calculated based upon the MGMT-SPT27 model implemented in the MNP package. t- stochastic neighbor embedding (tSNE) was used to project the methylation clustering in three dimensions using the Rtsne package. A Pearson correlation matrix of the 10K probe set was subjected to tSNE using a theta value of zero over 10,000 iterations as previously described and plotted using the rgl package.

mRNA Expression
Gene expression data was obtained from Agilent WG2.5 Affymetrix U133Plus2.0 or RNA sequencing platforms. Gene expression was preprocessed from two color Agilent microarrays using the R packages marray and limma and from single channel Affymetrix arrays using the affy package. Differential expression was assigned for microarray data using the limma package based upon a false discovery rate of 5%. RNaseq was aligned with Bowtie2 and TopHat and summarized as gene level fragments per kilobase per million reads sequenced using BEDTools and cufflinks/cuffnorm. Following rlog transformation and normalization, differential expression was assigned with DESeq2. Known Ensembl genes were further filtered to remove low abundance genes in all three datasets whose maximal expression was within the lowest 20% of all expression values based upon probe intensities or read depth. Replicate probes/features for each gene were removed by selecting those with the greatest median absolute deviation (MAD) in each dataset. Following centering within each dataset, log-transformed expression measures were combined and further normalized using pairwise loess normalization. Gene Set Enrichment Analysis was performed using the GSEA java application based upon pairwise comparisons of the major subgroups in the merged dataset. Heatmaps of gene expression across chromosomal arms were made using centered expression values rescaled across each chromosomal arm based upon the median absolute deviation of each probe. Differential expression analysis of TOP3A and FBXW7 was based on a Mann-Whitney U test of centred expression values between cases with and without losses and amplifications respectively in each case.

Sequence Analysis
Sequencing data was available as whole genome and/or whole exome (predominantly using Agilent’s SureSelect whole exome capture sets v4 and v5). Short read sequences from whole exome or whole genome sequencing were aligned to the hg19 assembly of the human genome using bwa. Following duplicate removal with Picard tools variants were called using the Genome Analysis toolkit according to standard Best Practices (Broad) including local re-alignment around Indels, down sampling and variant calling with the Unified Genotyper. Variants were annotated with the variant Effect predictor v74 from Ensembl tools and ANNOVAR to include annotations for variant allele frequency in 1000 genomes dbSNP v132 and the ExAc database as well as functional annotation tools SIFT and Polyphen). Depth of coverage varied from 16-295x (median 88x), with the greatest variation unsurprisingly in the exome data (whole genome range 50-150x, median=85x). Somatic variants were identified in regions covered by at least 10 reads in normal and
tumor sequences carrying at least 3 variant reads in the tumor and less than 2 in normal sequences. Hotspot TERT promoter mutations C228T and C250T were incidentally captured by the various exome platforms as they are located only 114 and 146 bp upstream of the translation start site, and were called even if only covered by a few reads. Mutation signatures were ascertained by grouping somatic substitutions on the basis of their 3’ and 5’ bases into 96 possible trinucleotide categories.

**Candidate Fusion Gene Nomination**

Structural variants were called from whole genome data using Breakdancer (breakdancer.sourceforge.net) filtered to remove commonly multi-mapped regions to identify somatic breakpoints separated by a minimum of 10 kbp involving at least one Ensembl gene. Fusion transcripts were detected from RNAseq data using chimerascan version 0.4.5a filtered to remove common false positives. To minimize unverified false positives, reporting of nominated fusions was restricted to genes within the core functional pathways and processes identified through integrated DNA copy number and somatic variant calling.

**Inferred Tumor Purity**

We used determined the somatic allele-specific copy number profiles using read depth from whole genome / exome sequencing, and used ASCAT (rick.ac.uk/peter-van-loo/software/ASCAT) to provide for an estimate of the non-neoplastic cell contamination of the sample as well as the overall ploidy of the tumor. Values ranged from 36-100%, with a median of 83%.

**Integrated Analysis of Driver Events**

Somatic non-synonymous coding mutations were filtered to remove common passenger mutations, polymorphisms and false positives in exome sequencing. Data were integrated with focal DNA copy number calls by GISTIC to provide gene-level binary alteration calls which were further selected for putative drive status on the basis of functional annotation. Oncoprint representations of integrated mutations, gene-level copy number alterations and fusion events were made using the online tool available at cBioportal (cbiportal.org). For the most commonly mutated genes mutations were mapped to the canonical transcript and plotted according to their predicted protein position using the Protein Painter (pecan.stjude.org). Integrated views of copy number alterations, structural variants and somatic mutations were made using CIRCOS (circos.ca) and rearrangements within TOP3A amplified regions in whole genome sequenced cases were identified using Breakdancer and aligned with copy number breakpoints in R.

**Pathway Analysis**

Pathway assignments were made for all genes carrying copy number alterations, structural variations or somatic mutations based on pathways in the MSigDB molecular signatures databases (Broad) as well as Gene Ontologies for Biological Processes and Molecular Functions (Gene Ontology consortium) and canonical pathways from KEGG, NetPath and Reactome. Genes within known CNVs and common false positives in exomic sequencing were excluded with reference to large scale genome profiling studies (CNVmap, ExAc, BCBio) Pathway analysis of genes carrying mutations, gene fusions and copy number aberrations was based on the pathways defined by these combined databases and subjected to enrichment analysis using the EnrichmentMap module within CytoScape.

**Statistical Analysis**

Statistical analysis was carried out using R 3.3.1 (www.r-project.org). Categorical comparisons of counts were carried out using Fishers exact test, comparisons between groups of continuous variables employed Student’s t-test, Wilcoxon signed –rank test, ANOVA or Mann-Whitney U test. Differences in survival were analysed by the Kaplan-Meier method and significance determined by the log-rank test. All tests were two-sided and a p value of less than 0.05 was considered significant. Multiple testing was accounted for using false discovery rate q values or the Bonferroni adjustment.

**DATA AND SOFTWARE AVAILABILITY**

All newly generated data have been deposited in the European Genome-phenome Archive (www.ebi.ac.uk/ega) with accession number EGAS00001002314 (sequencing) or ArrayExpress (www.ebi.ac.uk/arrayexpress/) with accession number E-MTAB-5528 (450k methylation).

**ADDITIONAL RESOURCES**

Processed copy number profiles are hosted as a disease-specific project within the Progenetix framework for annotated genomic analyses (dipg.progenetix.org) (Cai et al., 2014), and represented in the arrayMap resource (arraymap.org) (Cai et al., 2012). Curated gene-level copy number and mutation data are provided as part of the pediatric-specific implementation of the cBioPortal genomic data visualisation portal (pedcbioportal.org). Newly-generated raw data files are housed alongside published datasets made available to the Cavatica NIH-integrated cloud platform (www.cavatica.org).