Molecular identification of CNS NB-FOXR2, CNS EFT-CIC, CNS HGNET-MN1 and CNS HGNET-BCOR pediatric brain tumors using tumor-specific signature genes

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Abstract

Four molecular types of rare central nervous system (CNS) tumors have been recently identified by gene methylation profiling: CNS Neuroblastoma with FOXR2 activation (CNS NB-FOXR2), CNS Ewing Sarcoma Family Tumor with CIC alteration (CNS EFT-CIC), CNS high grade neuroepithelial tumor with MN1 alteration (CNS HGNET-MN1) and CNS high grade neuroepithelial tumor with BCOR alteration (CNS HGNET-BCOR). Although they are not represented in 2016 updated WHO classification of CNS tumors, their diagnostic recognition is important because of clinical consequences. We have introduced a diagnostic method based on transcription profiling of tumor specific signature genes from formalin-fixed, paraffin-embedded tumor blocks using NanoString nCounter Technology. Altogether, 14 out of 187 (7.4%) high grade pediatric brain tumors were diagnosed with either of four new CNS categories. Histopathological examination of the tumors confirmed, that they demonstrate a spectrum of morphology mimicking other CNS high grade tumors. However, they also exhibit some suggestive histopathological and immunohistochemical features that allow for a presumptive diagnosis prior to molecular assessment. Clinical characteristics of patients corroborated with the previous findings for CNS EFT-CIC, CNS NB-FOXR2 and CNS HGNET-MN1 patients, with a favorable survival rate for the latter two groups. Among six CNS HGNET-BCOR patients, three patients are long term survivors, suggesting possible heterogeneity within this molecular category of tumors. In summary, we confirmed the effectiveness of NanoString method using a single, multi-gene tumor specific signature and recommend this novel approach for identification of either one of the four newly described CNS tumor entities.

Keywords: Brain tumors, CNS EFT-CIC, CNS NB-FOXR2, CNS HGNET-MN1, CNS HGNET-BCOR, Genes expression
Introduction

High grade pediatric brain tumors are characterized by a continuing high mortality rate. Some tumors, especially central nervous system primitive neuroectodermal tumors (CNS-PNETs) were challenging to diagnose, being heterogeneous with variable neuronal, ependymal or glial differentiation [3]. Integrated use of technologies for whole genome analysis revealed molecular heterogeneity of CNS-PNETs [16] and the diagnosis of CNS-PNET was removed from the recent 4th revision of WHO Classification of Tumors of the Central Nervous System [9, 11]. Instead, a new diagnostic category of genetically defined embryonal brain tumor was introduced, namely C19MC-altered embryonal tumor with multilayered rosettes (ETMR).

The multicenter study conducted on 323 CNS-PNETs using genes methylation profiling [16] revealed that, on the molecular basis, the majority of CNS-PNETs cluster with other tumors, mainly with high grade gliomas (HGGs), ependymomas (EPNs), medulloblastomas (MBs) or ETMRs. Importantly, four previously unknown types of tumors were discovered, each characterized by the specific recurrent genetic rearrangements:

- CNS Neuroblastoma with FOXR2 activation (CNS NB-FOXR2), CNS Ewing Sarcoma Family Tumor with CIC alteration (CNS EFT-CIC), CNS high-grade neuroepithelial tumor with MN1 alteration (CNS HGNET-MN1), and CNS high-grade neuroepithelial tumor with BCOR alteration (CNS HGNET-BCOR). Although the latter four groups were discovered among previously diagnosed CNS-PNETs, subsequent analysis revealed that several tumors were also initially diagnosed as other entities, specifically CNS HGNET-MN1 tumors were classified mainly as EPNs or astroblastomas and CNS HGNET-BCOR tumors as HGGs or MBs [2, 4, 18].

Despite the fact that the newly discovered types of tumors are rare and are not represented in the recent updated edition of WHO 2016 classification, they require diagnostic recognition because of clinical importance. For example, there is a remarkable difference in survival of patients, with poor outcome for CNS HGNET-BCOR tumors and good outcome for CNS HGNET-MN1 and CNS NB-FOXR2 tumors. Following this, de-escalation of treatment for patients with CNS NB-FOXR2 tumors is being considered [6, 8]. Consequently, cIMPACT-NOW (the Consortium to Inform Molecular and Practical Approaches to CNS Tumor Taxonomy) reviewed new CNS tumor types and their nomenclature and recommended that they should be included in the future WHO classification of CNS tumors [10].

Since a diagnosis based on routine histopathology is not feasible, the only reliable and recommended so far method for detection of all four types of tumors was DNA methylation profiling. CNS HGNET-BCOR tumors were also recently diagnosed using capture-based next-generation DNA sequencing [4], nevertheless both methods are still challenging for routine diagnosis.

Importantly, the newly discovered types of tumors also display distinctive gene expression profiles [17]. We have therefore applied an alternative diagnostic method based on transcription profiling of marker genes using NanoString nCounter Technology. This approach allowed for analysis of genes expression from formalin-fixed, paraffin-embedded (FFPE) tumor blocks, routinely prepared in hospitals in everyday practice. We used tumor specific signature genes for identification of all four categories in a series of 187 high grade pediatric brain tumors. We confirmed the effectiveness of this method and performed clinical and histological characteristics of the recently identified types of tumors.

Materials and methods

Patients and tumor material

Pediatric patients diagnosed between 1998 and 2019 with high grade brain tumors in The Children’s Memorial Health Institute in Warsaw, Poland, were included in the analysis. The only criterion for inclusion was the availability of tumor material. Only samples examined previously by NanoString method and diagnosed molecularly as MBs [12, 13] were excluded from analysis. Analysis was performed on archive FFPE tumor material obtained at diagnosis and one relapsed sample. Hematoxylin-eosin-stained slides (H&E) were used for pathological evaluation and re-analysis to confirm the original diagnosis and determine the tumor tissue content. Whole preparations were scanned in Hamamatsu NanoZoomer 2.0 RS scanner at the original magnification 40x and analyzed independently by two experienced neuropathologists (W.G. and E.M.). For further investigation only samples with >70% of tumor cells (90% of samples) and >60% of tumor cells (10% of samples) were taken into account.

Altogether, 95 cases located in the infratentorial region included 50 EPNs, 37 MBs, 4 choroid plexus carcinomas (CPCs), 3 atypical teratoid rhabdoid tumors (ATRTs) and one glioblastoma (GBM).

Second group of 92 cases located in the supratentorial region included 28 HGGs, 23 PNETs, 20 EPNs, 10 CPCs, 7 ATRTs, 2 ETMRs and 2 tumors classified as CNS embryonal tumors “not otherwise specified” (NOS). Five PNETs were previously analyzed by DNA methylation profiling and presented in the already published series [17]. They included one CNS EFT-CIC, one CNS HGNET-MN1, one CNS HGNET-BCOR and two CNS NB-FOXR2 tumors, therefore we labelled them as the reference samples.

Informed consent was obtained to use tumor material according to the procedures outlined by The Bioethics Committee at the Children’s Memorial Health Institute’s, Warsaw, Poland.
Identification of marker genes for molecular classification by microarrays in silico analysis

The CEL files deposited in database Gene Expression Omnibus (GEO) were re-analyzed. We selected only pediatric cases (patients <18 years old) with high grade brain tumors, all analyzed on the platform Affymetrix Human Genome U133 Plus 2.0.

The following datasets were investigated: GSE73038 [17] which included CNS NB-FOX2, CNS EFT-CIC, CNS HGNET-MN1, CNS HGNET-BCOR, ATRTs, ETMR, HGGs and MBs; GSE70678 [7] which included ATRTs and GSE64415 [15] which included EPNs. CEL files were uploaded to R environment and subsequently normalized by MA55 method using “affy” library. Next quantile normalization was done separately on infra- and supratentorial tumors. Data were log2 transformed and affy probes with the lowest variance (variance< 0.25) were filtered out. In supervised approach, affy probes were selected by Student t-test. A set of samples with histologic diagnosis or a molecular subtype was compared 100 times to a set of the same size of samples drawn from patient’s samples with different histologic diagnosis or molecular subtype. A mean p-value and mean fold change from 100 t-tests were used as measures of good marker candidate. Random selection of samples and repeating t-test 100 times was done to assure robustness of marker selection.

Detection of molecular subtypes of tumors at the RNA level

NanoString nCounter system analysis (NanoString Technologies, Seattle, USA) was applied for identification of four molecular groups (CNS NB-FOX2, CNS EFT-CIC, CNS HGNET-MN1 and CNS HGNET-BCOR) in a series of altogether 187 tumors. Total RNA was extracted from FFPE tumors using RNeasy kits (Qiagen). RNA integrity was assessed using an Agilent 2100 Bioanalyzer.

For group assignment a custom NanoString CodeSet was applied, which consisted of marker genes and three housekeeping genes (ACTB, GAPDH and LDHA).

The sequences of the probes which were designed to target the regions of the marker genes are presented in Additional File 1.

Hybridization of the probes to tumor RNA samples was performed in the Clinical Research Centre, Medical University of Bialystok, Poland, according to NanoString Technologies procedures for hybridization, detection and scanning. Raw counts for each gene underwent technical and biological normalization using nSolver software. Clustering of the samples was performed with Euclidean distance metrics and average settings.

Area under curve (AUC) analysis of the selected marker probes

Nanostring data was uploaded to R and ROCR library was used. Prediction of classes was based on the expression of their marker probes/genes. Performance of the prediction was visualized by the ROC curves, where True positive rate (TPR) is on the y-axis and False positive rate (FPR) on the x-axis.

Detection of internal tandem duplication in BCOR gene

Genomic DNA was extracted from 6 tumor samples, which showed positive HGNET-BCOR signature, using the QIAamp DNA FFPE Tissue Kit (Qiagen). The duplicated region in exon 15 of BCOR was detected by targeted PCR using the following primers, as described by Appay et al., 2017 [1]. BCOR_15F: TCCTCCCGCATATTCTCGTG and BCOR_15R: ACACACTGTACATGGTGGGTCC (35 cycles of 98 °C 10s, 60 °C 30s, 72 °C 120 s). Bidirectional sequencing was performed using a 3500 Genetic Analyser (Applied Biosystems, Foster City, CA, USA). The sequences were determined on both DNA strands from at least two independent PCR products. The analyzed sequence fragments were compared with the BCOR cDNA (Gen-Bank RefSeq: NM_001123385.1) sequence using Mutation Surveyor software version 3.30 (Soft Genetics, LLC, State Collage, PA, USA).

Detection of MN1 fusion by targeted next-generation sequencing

Targeted cancer panel sequencing – the Ampliseq Childhood Cancer Panel for Illumina was used to detect MN1 gene fusions in four tumors. Prior to the library preparation, total RNA was extracted from FFPE tumor samples using RNeasy Mini Kit (Qiagen) and quantified with QuantiFlor RNA system (Promega). The percentage of RNA fragments > 200 nucleotides using Agilent 2100 Bioanalyzer was calculated (DV200). About 20 ng of RNA was used for library construction according to the manufacturer’s protocol. Each library was qualified using Agilent 2100 Bioanalyzer, and quantified using the QuantiFlor RNA system. The obtained library was then amplified using universal primers targeting the paired-end adapters. Clusters were generated and sequenced on the MiniSeq instrument (Illumina) using the MiniSeq High Output Kit (2 × 150 cycles). Alignment was performed using the Burrows Wheeler Aligner. The BaseSpace RNA Amplicon workflow (Illumina) was used to determine the fusion presence in MN1 gene.

Immunohistochemistry

Immunohistochemical reactions were performed on 4 μm thick FFPE sections using the Ventana BenchMark ULTRA IHC/ISH autostaining system. The following
Ventana primary antibodies were applied: for Glial Fibrillary Acidic Protein (GFAP), anti-GFAP [clone EP672Y], Rabbit Monoclonal Antibody; for Ki-67, anti-Ki-67 [clone 30–9], Rabbit Monoclonal Antibody, for Olig2, anti-Olig2 [clone EP112], Rabbit Monoclonal Antibody and for Synaptophysin, anti-Synaptophysin [clone MRQ-40], Rabbit Monoclonal Antibody. Antigen retrieval was performed in CC1 buffer followed by detection with the Ultra View HRP system (Roche/Ventana).

Expression of BCOR was detected using commercially available antibody (sc-514,576; Santa Cruz, Dallas, TX), [clone C10] at a dilution of 1:400. Antigen retrieval was performed using Target Retrieval Solution, High pH, (DAKO, Glostrup, Denmark) for 30 min in 99.5 °C. The specificity of BCOR immunoreexpression was tested in classic MB and normal testis. Whole slides were scanned in Hamamatsu NanoZoomer 2.0 RS scanner (Hamamatsu Photonics, Hamamatsu, Japan) at the original magnification 40 ×.

Results
Identification of marker genes for CNS NB-FOXR2, CNS EFT-CIC, CNS HGNET-MN1 and CNS HGNET-BCOR tumors
In the first step, we prepared the list of the top 100 significant genes for one of the four new CNS categories of tumors from the supplementary Table S12 published by Sturm et al., 2016 [17]. Subsequently, from every list we selected the candidates, which were both significantly up-regulated and showed the highest expression level as compared with the all other analyzed types of CNS high grade tumors.

Since >10% of the cohort analyzed by Sturm et al., 2016 [17] included adult cases, we performed a supplementary analysis of microarray data from pediatric cases only. They included, in addition to the four new CNS tumor entities, also 41 EPNs (GSE64415), 29 HGGs (GSE73038), 12 ATRTs (GSE73038 and GSE70678) and 6 ETMRs, as described in the Material and Methods section. Only supratentorial tumors were analyzed, since CNS NB-FOX2R and CNS EFT-CIC were not described in the infratentorial region.

Again, we prepare the list of the top 100 significant genes for each category, which were up-regulated at least 10 fold, as compared to all other types of tumors. Of note, among the top 100 significant genes for CNS NB-FOX2R, more than 30% genes were down-regulated, therefore fewer candidate markers for this group could be identified. Following this, a mean p-value and mean fold change from 200 t-tests were used as a measure of good marker candidates for this group. ECHS1 and ETV4 genes were not present among the top 100 significant genes from our analysis, nevertheless, we included them to examine their NanoString performance.

Based on the two lists of genes, we finally selected candidate marker genes for each group as presented in Table 1.

NanoString probes hybridization performance
Marker genes were analyzed separately in supratentorial and infratentorial locations since the distribution (e.g. MBs, CNS NB-FOX2R and CNS EFT-CIC) or molecular characteristics (e.g. EPNs) of pediatric CNS tumors is different in these compartments. Therefore, the marker genes for CNS HGNET-BCOR and CNS HGNET-MN1 tumors were analyzed in both supratentorial and infratentorial location, as opposed to CNS NB-FOX2R and CNS EFT-CIC tumors, which were described in the supratentorial region only.

Good hybridization was considered when level of expression for the probe in the reference sample was higher (as expected) than mean level of all remaining samples. It ranged between 3.9 fold (for MYOD1) and 92.7 fold (for FOXR2) in our series.

For CNS HGNET-BCOR detection three probes (for SHISA8, MNX1 and MMP15) showed good hybridization in both compartments and additional two probes (for PRDM6 and EMX1) were chosen for detection of supratentorial tumors only, since these genes were over-expressed in several posterior fossa EPNs or MBs. Instead, probe for RTN4RL1 gene was added to the custom set for detection of posterior fossa CNS HGNET-BCOR tumors.

For CNS HGNET-MN1 detection all six probes in supratentorial tumors (for BEND2, MUM1, SHOX, APCDD1L, FAM3B and ECHS1) showed good hybridization. Only three probes were used in the posterior fossa tumors (for BEND2, SHOX and APCDD1L) and tested positive. Two additional probes examined in the latter compartment (for RBP4 and PRRX2) showed poor hybridization and these genes were excluded from further analysis.

All probes for detection of supratentorial CNS NB-FOX2R and CNS EFT-CIC tumors showed the good hybridization result.

Clustering of tumors according to the expression level of marker genes analyzed by NanoString method
Unsupervised hierarchical clustering was performed separately, as mentioned before, for 92 supratentorial and 95 infratentorial tumors (plus two reference samples) with the selected marker genes using Euclidean distance metrics and average settings. The results from supratentorial tumors analysis showed only one reference sample with CNS EFT-CIC signature, three samples with CNS NB-FOX2R signature (including two reference samples), four samples with CNS HGNET-MN1 signature (including one reference sample) and five samples with CNS...
HGNET-BCOR signature (including one reference sample), (Fig. 1a). The results from infratentorial tumors analysis showed only one reference sample with CNS HGNET-MN1 signature and three samples with CNS HGNET-BCOR signature (including one reference sample), (Fig. 1b).

Area under curve (AUC) analysis results for NanoString marker probes

AUC analysis was performed on supratentorial data where all four categories of tumors were detected. Prediction of HGNET-BCOR or HGNET-MN1 or NB-FOXR2 or EFT-CIC classes was based on the expression level of their marker probes presented in Additional File 1. AUC was close to 1 for all the tested markers for classes HGNET-BCOR (6 marker probes), HGNET-MN1 (6 marker probes), NB-FOXR2 (4 marker probes) and EFT-CIC (6 marker probes), (Additional File 2). Close to ideal performance for tested markers may be due to small number of samples of these rare tumors. Especially in case of EFT-CIC subtype only one sample was available. These results indicate that selected NanoString markers can distinguish the individual tumor classes compared to all other classes.

Further characteristics of CNS HGNET-BCOR tumors

In addition to the reference sample diagnosed previously as PNET, four newly detected CNS HGNET-BCOR tumors were originally diagnosed as HGG. Therefore, we performed further analyses to confirm that their molecular profiles differ from the original diagnosis. We included into the custom set three probes for genes expressed in glioma tumors (GFAP, OLIG2 and PMP2), in addition to the CNS HGNET-BCOR marker probes, and performed hybridization in the series of 27 tumors originally diagnosed as HGG and one HGG relapsed sample. NanoString data analysis revealed two distinct clusters and one outlier sample (probably not

| Group       | Probe     | Gene  | This study | Sturm et al., 2016 |
|-------------|-----------|-------|------------|--------------------|
|             |           |       | Top position | Fold | Top position |                  |
|             |           |       | significance |      | significance |                  |
| HGNET-BCOR  | 219433_at | BCOR  | Group marker |       | Group marker |                  |
| HGNET-BCOR  | 215264_at | EMX1  | 1 | 65.5 | 122 |
| HGNET-BCOR  | 230311_s_at | PRDM6 | 3 | 13.2 | 47 |
| HGNET-BCOR  | 230700_at | RTHN4RL1 | 22 | 12.2 | 40 |
| HGNET-BCOR  | 244467_at | SHHSA8 | 29 | 29.2 | 12 |
| HGNET-BCOR  | 214614_at | MNX1  | 37 | 27.9 | 60 |
| HGNET-BCOR  | 203365_s_at | MMP15 | 59 | 10.1 | 13 |
| HGNET-MN1   | 1553842_at | BEND2 | Group marker |       | Group marker |                  |
| HGNET-MN1   | 223347_at | MUM1  | 2 | 16 | 8 |
| HGNET-MN1   | 207570_at | SHOX  | 3 | 362 | 7 |
| HGNET-MN1   | 235548_at | APCDD1L | 5 | 185 | 18 |
| HGNET-MN1   | 219140_s_at | RBP4 | 6 | 158 | 51 |
| HGNET-MN1   | 219729_at | PRRX2 | 10 | 38.6 | 112 |
| HGNET-MN1   | 227194_at | FAM3B | 15 | 320 | 13 |
| HGNET-MN1   | 201135_at | ECHS1 | – | – | 9 |
| NB-FOXR2    | 1569669_at | FOXR2 | Group marker |       | Group marker |                  |
| NB-FOXR2    | 239499_at | DNAH2 | 45 | 34.5 | 7 |
| NB-FOXR2    | 209842_at | SOX10 | 74 | 95.9 | 23 |
| NB-FOXR2    | 222095_s_at | FAM163A | 93 | 19.8 | 24 |
| EFT-CIC     | 212338_at | MYO1D | 11 | 31.7 | 8 |
| EFT-CIC     | 209616_s_at | CES1 | 17 | 197 | 43 |
| EFT-CIC     | 235849_at | SCARA5 | 44 | 167 | 45 |
| EFT-CIC     | 215161_at | CAMK1G | 62 | 130 | 1 |
| EFT-CIC     | 235238_at | SHC4 | 80 | 37.5 | 6 |
| EFT-CIC     | 1554576_s_at | ETV4 | – | – | 91 |
glioma), (Fig. 2). Cluster 1 included six samples with expressed CNS HGNET-BCOR signature genes and cluster 2 included other samples expressing only GFAP, OLIG2 or PMP2. Two samples from the same patient, originally diagnosed as anaplastic oligodendroglioma and the relapsed sample, expressed both CNS HGNET-BCOR signature and glioma marker genes (cluster 1B). Immunohisto pathological analysis confirmed positive reaction for expression of both GFAP and OLIG2 in both samples.

Tandem duplication in exon 15 of BCOR gene was assessed in all samples diagnosed by NanoString method as CNS HGNET-BCOR. Among 6 tumor samples from cluster 1 (Fig. 2) four cases showed the presence of tandem duplication, including two samples from cluster 1B, one case showed no duplication and in one case the result was inconclusive. Additional CNS HGNET-BCOR cerebellar tumor, originally diagnosed as unclassified malignant neuroepithelial tumor, showed lack of exon 15 of BCOR gene duplication and negative reaction for GFAP and synaptophysin.

BCOR immunohistochemistry was performed on three tumors with negative or inconclusive BCOR tandem duplication results. Two tumors with negative result showed diffuse and strong BCOR nuclear immunopositivity, including tumor with unique papillary architecture originally diagnosed as unclassified malignant neuroepithelial tumor (Fig. 3a,b). The third tumor with inconclusive result revealed only focal BCOR immunopositivity of neoplastic cell nuclei (Fig. 3c). Medulloblastoma tumor, in contrast to CNS HGNET-BCOR tumors, appeared to be BCOR immunonegative (Fig. 3d.)

Detection of MN1 fusion transcript
In four tumors showing CNS HGNET-MN1 signature, we were able to perform NGS analysis in order to confirm NanoString results. In all cases the presence of MN1:BEND2 fusion transcript was detected.
Histopathological characteristics of tumors

CNS EFT-CIC
Tumor displayed features attributed to originally diagnosed CNS-PNET. It appeared as densely cellular, small cell tumor with nodular, fascicular and focally alveolar growth pattern (Fig. 4a). The nodules were pale and consisted mainly of small round cells exhibiting a neurocytic phenotype. Larger cells with visible eosinophilic cytoplasm were found only sporadically (Fig. 4b). The stromal pattern could be observed focally (Fig. 4c).

CNS NB-FOXR2
All three tumors, originally diagnosed as PNETs, displayed embryonal-like morphology of a highly cellular tumor composed of small, round cells with hyperchromatic nuclei surrounded by a clear halo (Fig. 5a). Occasionally neuroblastic-like rosette formations (Fig. 5b) and neuropil islands (Fig. 5c) could be seen. The tumors were supported by a rich capillary network with presence of pseudorosette arrangement of neoplastic cells. Foci of myxoid degeneration (Fig. 5d) and microvascular

Fig. 2 Clustering of tumors with HGG original diagnosis using NanoString method. Cluster 1 represents the samples with high expression of CNS HGNET-BCOR markers. Cluster 2 represents the samples with high expression of glioma markers. Outlier sample is probably not a glioma tumor. Red arrow indicates a location of the CNS HGNET-BCOR reference sample. Colors represent log2 gene expression differences.

Fig. 3 BCOR immunohistochemistry. a, b A strong, widespread nuclear BCOR immunoreactivity and c focal tumor BCOR nuclei staining, in contrast to d negative immunostaining of medulloblastoma. The scale bars: a–c – 50 μm.
proliferation was found (Fig. 5e). The tumor cells exhibited extensive Olig2 immunoexpression (Fig. 5f).

**CNS HGNET-MN1**

Four tumors were originally diagnosed as anaplastic EPN (2 cases), CPC (1 case) and PNET (1 case). Most of them exhibited ependymoma-like or astroblastoma-type perivascular pseudorosettes composed of elongated cells with abundant eosinophilic cytoplasm and clear processes radiating towards central blood vessels (Fig. 6a). Pseudorosettes with hyaline sclerosis were commonly observed (Fig. 6b). Perivascular formations formed a focal ribbon-like pattern (Fig. 6c). The tumor initially identified as CPC showed areas with a typical papillary structure (Fig. 6d). A characteristic feature was the extensive hyalinization of the fibro-vascular stroma (Fig. 6e,f). The solid growth pattern with papillary formation (Fig. 6g) and microcystic changes was also seen (Fig. 6h). Only one tumor diagnosed as PNET consisted of densely packed, round to ovoid cells with scanty cytoplasm and high mitotic activity (Fig. 6i). In this tumor, perivascular pseudorosettes have been observed only sporadically.

**CNS-HGNT BCOR**

Three tumors previously identified as GBM exhibited similar morphology with a solid, highly cellular, mostly monomorphic growth pattern (Fig. 7a), accompanied by a rich, thin-walled capillary network. The majority of neoplastic cells were characterized by ovoid or oval nuclei with fine chromatin and scant eosinophilic cytoplasm with delicate fibrillary processes. Two tumors previously identified as GBM showed an additional pleomorphic cellular component with the appearance of astrocytes or gemistocytes with round to oval nuclei and abundant eosinophilic cytoplasm (Fig. 7b). In one case, the cellular elements contained oligodendroglial-like cells with round nuclei surrounded by clear cytoplasm (Fig. 7c). One tumor diagnosed as pediatric anaplastic oligodendroglioma displayed uniform cells with round nuclei and perinuclear halo, chicken-wire pattern of vessels, numerous persistent neurons with perinuclear satellitosis and microcalcifications (Fig. 7d).

Another tumor, initially identified as PNET, showed embryonal-like cytological features with a predominance of densely-packed, small, round, hyperchromatic cells with brisk mitotic activity (Fig. 7e). The most characteristic
feature of all BCOR tumors was a distinctive branching network of delicate capillary blood vessels with the dominant chicken-wire appearance (Figs. 7f, g). Perivascular ependymoma-like pseudorosettes were rare (Fig. 7h). Only in the tumor originally considered as PNET, both in the primary tumor and relapse, the small vessels surrounded by perivascular eosinophilic, occasionally microcystic zones (Fig. 7i) and lacking fibrillary processes were more often found. In some areas, the hyalinized vessels (Fig. 7j) or thin-walled, round vessels in the spongy matrix (Fig. 7k) could be observed. The PNET relapse displayed the same morphology as the original tumor. Two cases revealed focal microcystic or cystic changes or areas composed of stellate cells within myxoid background (Fig. 7l). In all cases, micronecrosis (Fig. 8a) and areas of ischemic necrosis were present, sometimes surrounded by pseudopalisading. However, none of the cases identified as HGG showed microvascular proliferation, a key histologic feature of GBM.

The only tumor, initially diagnosed as unclassified malignant neuroepithelial tumor, had a completely different papillary structure (Fig. 8b) with uniform, small cells similar to neurocytes arranged around eosinophilic acellular cores, rarely around blood vessels (Fig. 8c). In some parts of the tumor, the columnar arrangement was seen and the background was partially microcystic. Tumors demonstrated brisk mitotic activity and foci of necrosis.

Immunohistochemically, the majority of CNS HGNET-T-BCOR tumors did not reveal expression of GFAP, except of some reactive astrocytes (Fig. 8d). Only one tumor, initially diagnosed as pediatric oligodendroglioma, showed strong GFAP immunoreactivity related to numerous entrapped reactive astroglial cells and dense network of their processes in the tumor background (Fig. 8e). Especially intense GFAP positivity was observed around blood vessels (Fig. 8f). However, some areas of tumor were immunonegative for GFAP (Fig. 8g), except a few tiny glial processes (Fig. 8h). In all CNS HGNET-BCOR tumors immunoexpression of synaptophysin was absent and Olig2 immunoreactivity was detected only focally. The Ki67 proliferation index was high in originally diagnosed GBMs and PNET, and also in unusual papillary, neuroepithelial tumor (Fig. 8i).

Clinical characteristics of patients
In our series, 14 out of 187 (7.4%) patients were diagnosed with either of four new CNS categories of tumors, Table 2. All but two tumors were detected in the supratentorial region.

Only one patient presented with metastases. Patients were treated according to the Polish Pediatric Neurooncology Group (PPNG) protocols. All patients underwent
gross total resection of tumors. The specific protocols for particular patients are listed in Table 2.

Only one infant girl was diagnosed with CNS EFT-CIC (the reference sample). She was treated without radiotherapy during primary treatment, relapsed and died within 1 year from diagnosis.

Three CNS NB-FOXR2 patients (age 4.5, 5, 7 years old) had the previous original diagnosis as PNETs and one patient had metastases to the spinal cord. Because of the same original diagnosis and the similar age, patients were treated according to the same MB/PNET protocol with the cerebrospinal irradiation (CSI). None of the patients relapsed and all are alive at least 10 years since diagnosis.

Four CNS HGNET-MN1 female patients (age 2–11 years old) had different original diagnoses and therefore were treated according to different protocols. However, none of them relapsed and all are long term survivors, with at least 11 years of observation time.

Six CNS HGNET-BCORs patients had various tumor location (four tumors in supra-tentorial region, one in the pons and one in the cerebellar hemisphere). They had also diverse original diagnoses, age (1.5–9 years) and survival rate. Three patients survived at least 5 years without relapse. Notably, two of them, who were treated according to the HGG I version protocol (including Ifosfamide, Etoposide and Adriamycin) are long term survivors, with at least 12 years of observation time.

**Discussion**

New molecular categories of tumors discovered by DNA methylation profiling, CNS NB-FOXR2, CNS EFT-CIC, CNS HGNET-MN1 and CNS HGNET-BCOR, are rare pediatric brain malignancies that have not yet been

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**Fig. 7** Representative histopathology of CNS-HGNT BCOR tumors. a) Solid, highly cellular, monomorphic growth pattern with rich, thin-walled capillary network. b) Pleomorphic cells of astroglial-like appearance. c) Oligodendroglial-like monomorphic cells with round nuclei surrounded by clear cytoplasm accompanied by chicken-wire pattern of vessels. d) Infiltration of uniform neoplastic cells with round nuclei and microcalcifications. e) Densely-packed, small, hyperchromatic cells and thin-walled capillary blood vessels. f, g) Network of capillary blood vessels with chicken-wire appearance. h) Perivascular ependymoma-like pseudorosette. i) Small vessels surrounded by perivascular eosinophilic zones. j) Hyalinized vessels or k) thin-walled, round vessels in the spongy matrix. l) Area composed of stellate cells within myxoid background. The scale bars: a, b, d-e, h, i – 50 μm; f – 40 μm; c, g-l – 30 μm.
included in the recent WHO 2016 classification. Therefore, reliable identification of new cases diagnosed either retrospectively or prospectively is crucial for a better description of biological and morphological features, as well as clinical characteristics of patients.

In this study, we applied a novel approach for the diagnosis of all four new entities using the NanoString nCounter Analysis System and multi-genes tumor-specific signatures. This method allows for analysis of degraded RNA, is therefore compatible with FFPE specimens and was previously successfully tested for detection of molecular subtypes in MBs [12–14].

We applied a multi-gene detection method with a limited but specific number of probes, what makes such approach both reliable and cost effective. The signatures include group specific marker genes (BCOR, BEND2 and FOXR2) which are highly expressed in the relevant groups. All reference RNA samples previously analyzed by DNA methylation profiling clustered within the expected groups, thus validating our NanoString gene expression based diagnosis.

CNS HGNET-MN1 genes (BEND2, MUM1, SHOX, APCDD1L, FAM3B and ECHS1) showed robust and clear expression signature. Since no specific immunohistochemical markers were described in this group of tumors, our detection method seems to be of particular diagnostic value. In all four cases analyzed by NGS, the presence of MN1:BEND2 fusion transcript was detected, again validating NanoString results.

For CNS HGNET-BCOR detection, three probes for SHISA8, MNX1 and MMP15 showed diagnostic usefulness regardless of tumor location. Multi-gene detection method in this type of tumors can be supplemented by immunohistochemical analysis of BCOR protein. Nevertheless, BCOR can be also expressed in other brain tumors [1, 4, 5], especially in CNS HGNET-MN1, what is visible at the RNA level in our series as well (Fig. 1a).

We should underline, that the described method is based on the open system, where more genes/probes can be added to the custom set. This may be useful for further investigation, especially of CNS EFT-CIC tumors, since only four microarrays profiles were available for identification of marker genes for this group. Also, more genes may be used for better characterization of CNS HGNET-BCOR patients, including potential prognostic markers.

It is important to mention, that the reference samples from the reliable, previously diagnosed tumors should be included in the analysis of new cases. Data from such samples can be stored as RCC files and used for diagnostic comparison. Clustering analysis using freely available nSolver software is rapid and straightforward. In our experience whole diagnostic process from RNA extraction,
hybridization and RCC file analysis can be accomplished within 3 days.

We detected altogether 14 samples (7.4%), which showed tumor-specific expression signatures representative for new molecular categories of tumors. Only one CNS EFT-CIC (reference sample) was detected in supratentorial location, confirming very rare occurrence of this type of tumor. This tumor displayed features attributed to originally diagnosed CNS-PNET, as reported in tumors of this molecular group. The cells showed a neurocytic phenotype but lacked immunoexpression of neuronal differentiation.

Three tumors were identified as CNS NB-FOXR2, all originally diagnosed as PNETs, what corroborates with the previous findings [17]. They displayed the same embryonal-like morphology with neuroblastic-like rosettes and pseudorosettes. Foci of myxoid degeneration and microvascular proliferation could be found and the tumor cells exhibited extensive Olig2 immunoexpression. These features correspond to those described so far in CNS NB-FOXR2 tumors. In line with the previous data, all three our patients survived at least 10 years without relapses, what provides further evidence for proposed de-escalation of treatment for this molecular group [6].

Four tumors were identified as CNS HGNET-MN1, all located in the supratentorial region. This suggests that CNS HGNET-MN1 tumors are very rare in the posterior fossa location, in line with the earlier findings, where such location was described in < 10% of CNS HGNET-

| ID | Age yrs | Sex | NanoString diagnosis | Orthogonal method | Original diagnosis | Tumor location | Meta- | Relapse mts/location | DOD mts | ADF mts | Primary treatment PPNG protocols
|----|---------|-----|---------------------|------------------|-------------------|---------------|------|---------------------|---------|---------|------------------------|
| CIC 1 ref | 0.5 | f | EFT-CIC | Methylation | PNET | fronto-temporal right | no | 5/distant | 9 | – | none MB/PNET |
| BCOR 1 ref | 1.5 | m | HGNET-BCOR | Methylation, BCOR tandem dup | PNET | frontal right | no | 8/local | 35 | – | none MB/PNET |
| BCOR 2 | 9 | m | HGNET-BCOR | BCOR immuno | GBM | pons | no | 29/local | 31 | – | local HGG II version |
| BCOR 3 | 6 | f | HGNET-BCOR | BCOR immuno | GBM | frontal right | no | no | no | 144 | local HGG I version |
| BCOR 4 | 6 | f | HGNET-BCOR | BCOR tandem dup | GBM | parietal left | no | no | no | 168 | local HGG I version |
| BCOR 5 | 2.5 | f | HGNET-BCOR | BCOR immuno | Unclassified malignant neuroepithelial tumor | cerebellar hemisphere | no | no | no | 66 | CSI MB SR |
| BCOR 6 | 8 | m | HGNET-BCOR | BCOR tandem dup | Anaplastic oligodendrogloma | frontal lobe left | no | 22/local | 24<sup>foo</sup> | – | local HGG I version |
| MN1 ref | 2 | f | HGNET-MN1 | Methylation, MN1:BEND2 fusion | PNET | parieto-temporal right | no | no | no | 144 | none MB/PNET |
| MN2 | 9 | f | HGNET-MN1 | MN1:BEND2 fusion | EPN | frontal lobe right | no | no | no | 204 | local HGG I version |
| MN3 | 11 | f | HGNET-MN1 | MN1:BEND2 fusion | Papillary EPN | parietal left | no | no | no | 132 | local none |
| MN4 | 6 | f | HGNET-MN1 | MN1:BEND2 fusion | CPC | parietal right | no | no | no | 180 | local CPC |
| FOXR2 1 ref | 5 | f | NB-FOXR2 | Methylation | PNET | parietal left | yes | no | no | 120 | CSI MB/PNET |
| FOXR2 2 ref | 4.5 | m | NB-FOXR2 | Methylation | PNET | parieto-occipital left | no | no | no | 137 | CSI MB/PNET |
| FOXR2 3 | 7 | m | NB-FOXR2 | nd | PNET | frontal right | no | no | no | 240 | CSI MB/PNET |

<sup>foo</sup> reference patients, mts months, DOD Died of disease, ADF Alive disease free, 24<sup>foo</sup> Death from chemotherapy-related toxicity after 24 months, RT Radiotherapy, CSI Cerebrospinal irradiation, PPNG The Polish Pediatric Neurooncology Group, MB/PNET Medulloblastoma/PNET protocol, HGG High grade glioma protocol, MB SR Medulloblastoma standard risk protocol, CPC Choroid plexus carcinoma protocol, dup Duplication, BCOR immuno BCOR immunohistochemistry, nd Not done.
MN1 tumors by Sturm et al., 2016 [17] and in none out of 8 tumors analyzed by Lehman et al., 2019 [8]. The predominant histologic pattern of HGNET-MN1 tumors, including ependymoma-like or astroblastoma-type perivascular pseudorosettes and hyalinization, is consistent with features previously described in MN1 altered tumors, which were most often identified as astroblastoma. Only one tumor, initially recognized as PNET, consisted of densely packed, small cells with high mitotic activity and inconspicuous perivascular pseudo-rosettes.

Despite different treatment protocols, all four female patients are long term survivors. One patient received local radiotherapy only and another patient chemotherapy only. Since existing data point towards better overall survival rate [6, 9, 17], less intensive therapeutic regimen should be considered for patients with CNS HGNET-MN1 tumors.

Altogether, we have identified 6 CNS HGNET-BCOR cases, four in supratentorial and two in infratentorial location. Three supratentorial tumors harboured BCOR exon 15 tandem duplication and in one tumor the result was inconclusive. Two tumors from cluster 1A (Fig. 2), (BCOR2 and BCOR5) lack the presence of duplication despite displaying clear CNS HGNET-BCOR expression signature. An absence of duplication in CNS HGNET-BCOR tumors identified by methylation profiling was described in some cases by Sturm et al., 2016 [17], where an intragenic in-frame deletion in BCOR gene or BCOR frameshift mutations have been found. Most likely, other rearrangement than BCOR exon 15 tandem duplication, are present in BCOR2 and BCOR5 samples.

The majority of our CNS-HGNT BCOR tumors displayed distinctive histological features, including high cellularity, rich capillary network of chicken-wire appearance and palisading necrosis. Most neoplastic cells revealed a glial morphology with eosinophilic cytoplasm and delicate fibrillar processes, less often oligodendrogial-like phenotype with numerous persistent neurons and microcalcifications. The immunophenotype was similar to the previously reported CNS HGNET-BCOR tumors [4]. Majority of currently studied tumors showed only focal positive GFAP staining limited to reactive astrocytes. Even the very strong GFAP expression in the tumor of classic oligodendrogial-like histology was associated with the abundant reactive component of stromal astroglia.

Only the case initially diagnosed as unclassified malignant neuroepithelial tumor (BCOR 5) exhibited a completely different morphology, which may indicate some heterogeneity within this molecular group. Because of high Ki67 proliferation index and cerebellar location, this 2.5-year old girl was treated according to the standard risk protocol for medulloblastoma (MB SR) with cerebro-spinal irradiation (CSI), despite young age. She has already survived 5 years without relapse of the disease. By contrast, another young patient (BCOR 1) who did not receive radiotherapy, relapsed within 1 year since diagnosis and subsequently died.

One tumor (BCOR 2) originally diagnosed as GBM was located in the pons and this is only the second CNS HGNET-BCOR described in such location so far.

Two patients also originally diagnosed as GBM (BCOR3 and BCOR4 with confirmed BCOR exon 15 tandem duplication) are of particular interest since they did not relapse and are alive at least 12 years since diagnosis. They were treated according to the PPNG HGG I version protocol (including Ifosfamide, Etoposide and Adriamycin). Taking into account the diverse outcome in our patients it seems that CNS HGNET-BCOR group may be clinically and biologically heterogeneous. Indeed, NGS analysis revealed that 40% of analyzed tumors, in addition to BCOR exon 15 internal tandem duplication, harbor other likely pathogenic genetic alterations, what may have an impact on tumor behavior [4]. Altogether, CNS HGNET-MN1 category requires further investigation both in terms of detection method and underlying or additional genetic aberrations. It is likely, that methods based on gene methylation or expression profiling detect more CNS HGNET-BCOR tumors than e.g. diagnosis based on detection of BCOR exon 15 tandem duplication only.

Conclusions

We propose a novel approach for identification of either one of four newly described CNS tumor entities using a single multi-gene signature. This method allows also for analysis of FFPE tumor samples routinely preserved in diagnostic laboratories. Our results confirm that the recently proposed molecular classification of malignant pediatric brain tumors demonstrates a spectrum of morphology that mimic other CNS high grade tumors. Nevertheless, they exhibit some suggestive histopathological features and immunohistochemical profile that allow to make a presumptive diagnosis prior to molecular assessment. Further validation and collection of new cases should improve our understanding of the biology of these rare tumors and recommend better therapeutic protocols for patients.

Supplementary information

Supplementary information accompanies this paper at https://doi.org/10.1186/s40478-020-00984-9.

Additional file 1. The target regions of the marker genes.

Additional file 2. Area under curve (AUC) analysis of the marker probes for four category of tumors. Performance of the prediction was visualized by the ROC curves, where True positive rate (TPR) is on the y-axis and False positive rate (FPR) on the x-axis. AUC was close to 1 for all markers tested for HGNET-BCOR (6 marker probes), HGNET-MN1 (6 marker probes), NB-FOXR2 (4 marker probes) and EFT-CIC (6 marker probes) classes.
Abbreviations
ATRTs: Atypical teratoid rhabdoid tumors; CNS: Central nervous system; CNS EFT-CIC: CNS Ewing Sarcoma Family Tumor with CIC alteration; CNS HGNET-MN1: CNS high-grade neuroepithelial tumor with MN1 alteration; CNS HGNE T-BCOR: CNS high-grade neuroepithelial tumor with BCOR alteration; CNS NB-FOKR2: CNS Neuroblastoma with FOKR2 activation; CNS-PNETs: Nervous system primitive neuroectodermal tumors; CPCs: Choroid plexus carcinomas; CST: Cerebrospinal irradiation; EPNs: Ependymomas; ETMR: Embryonal tumor with multilayered rosettes; FFPE: Formalin-fixed, paraffin-embedded; GBM: Glioblastoma; GFAP: Glial Fibrillary Acidic Protein; GSO: Gene Expression Omnibus; H&E: Hematoxylin-eosin-stained slides; HGGs: Grade gliomas; MBs: Medulloblastomas; NOS: Not otherwise specified

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Authors’ contributions
Maria Lastowska, – designed the study, interpreted the results and written the manuscript. Joanna Trubicka - performed laboratory and bioinformatics work, interpreted the results. Anna Sobocińska – performed laboratory and bioinformatics work, interpreted the results. Bartosz Wojtas – performed bioinformatics analyses of microarray data. Magdalena Niemira - performed laboratory work using NanoString technique. Anna Szalkowska - performed laboratory work using NanoString technique. Adam Krętówski – contributed to implementation of the project based on NanoString technology. Agnieszka Karczkowska-Więcławowska - performed immunohistopathological analyses. Magdalena Kaleta - performed scanning of preparations and immunohistopathological analyses. Maria Ejmont – performed mutational analyses. Marta Perek-Polnik - collected and interpreted clinical data of patients. Bożenna Dembovska-Bagińska - collected and interpreted clinical data of patients. Wiesława Grajkońska - performed histopathological examination of tumors and interpreted the results. Ewa Matyja - performed the histopathological examination of tumors, interpreted the results, prepared histopathological figures and was a major contributor in writing the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials
All data generated or analysed during this study are included in this published article [and its supplementary information files].

Ethics approval and consent to participate
Informed consent was obtained to use tumour material according to the procedures outlined by The Bioethics Committee at the Children’s Memorial Health Institute’s, Warsaw, Poland, approval No 1 /KBEB/2017.

Consent for publication
“Not applicable”.

Competing interests
The authors declare that they have no competing interests.

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