

Gene expression profiles of human glioblastomas are associated with both tumor cytogenetics and histopathology

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Despite the increasing knowledge about the genetic alterations and molecular pathways involved in gliomas, few studies have investigated the association between the gene expression profiles (GEP) and both cytogenetics and histopathology of gliomas. Here, we analyzed the GEP (U133Plus2.0 chip) of 40 gliomas (35 astrocytic tumors, 3 oligodendrogliomas, and 2 mixed tumors) and their association with tumor cytogenetics and histopathology. Unsupervised and supervised analyses showed significantly different GEP in low- vs high-grade gliomas, the most discriminating genes including genes involved in the regulation of cell proliferation, apoptosis, DNA repair, and signal transduction. In turn, among glioblastoma multiforme (GBM), 3 subgroups of tumors were identified according to their GEP, which were closely associated with the cytogenetic profile of their ancestral tumor cell clones: (i) EGFR amplification, (ii) isolated trisomy 7, and (iii) more complex karyotypes. In summary, our results show a clear association between the GEP of gliomas

and tumor histopathology; additionally, among grade IV astrocytoma, GEP are significantly associated with the cytogenetic profile of the ancestral tumor cell clone. Further studies in larger series of patients are necessary to confirm our observations.

Keywords: cytogenetics, FISH, gene expression, glioblastoma, glioma, histopathology, microarray.

In the last decades an increasingly high amount of information has accumulated about the cytogenetic characteristics of human gliomas.^{1–3} Although different and unique cytogenetic profiles have been described in low- vs high-grade gliomas, no clearly defined cytogenetic subgroups have been further defined among the most frequent tumor subtypes—eg, glioblastoma multiforme (GBM). Because of this, more recent attempts have been made to search for new relevant molecular markers that could be of help for a better subclassification of low- and high-grade gliomas and the identification of putative new therapeutic targets.^{4–6} Until now, such efforts have mainly concentrated on genes that are specifically altered in a significant fraction of all gliomas, particularly among high-grade tumors (eg, *EGFR*, *PTEN*, *TP53*, *CDKN2A*, *MDM2*, and *PDGFA* genes) and their potential association with specific intracellular signaling pathways such as those involving *STAT3*, *MAPK*, and *AKT*;^{7–10} however, these studies have only been partially successful.^{11–13}

In recent years, the availability of large-scale genomic approaches and new bioinformatic tools has fostered the

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identification of molecular profiles that could be characteristic of human gliomas and define specific subgroups of GBM. Thus, it has been shown that the patterns of gene expression of gliomas correlate with tumor histopathology^{14–18} and patient survival^{19,20} and gene expression profiles (GEP) emerge in some studies as an independent prognostic factor among high-grade gliomas.²⁰ According to these studies, most informative genes include genes involved in cell proliferation, RNA processing, signal transduction, and the proteasome functional activity.¹⁸ However, no clear association between the GEP of gliomas and tumor cytogenetics has been clearly established so far, which could help on the understanding of the various GEP described.

In a recent paper²¹ using multicolor interphase fluorescence in situ hybridization (iFISH), we showed the existence of highly variable and heterogeneous intratumoral patterns of cytogenetic abnormalities in human gliomas; such cytogenetic profiles are typically characterized by complex karyotypes consisting of combined gains of chromosome 7, del(9p21)/null 9p, and/or del(10q23) in association with a variable number of other more heterogeneous cytogenetic changes; interestingly, the acquisition of these cytogenetic lesions followed different clonal pathways.²¹ In this regard, it should be noted that different pathways of intratumoral cytogenetic evolution were found in low- vs high-grade gliomas, which further suggests that the latter tumors might not always correspond to more advanced stages of histologically low-grade gliomas. Even more interestingly, different recent pathways of intratumoral clonal evolution were defined within GBM.²¹ On the basis of these results, the potential impact of these unique cytogenetic pathways in tumor behavior through the expression of different GEP remains to be elucidated.

The aim of the present study was to identify whether the reported cytogenetic heterogeneity of human gliomas—particularly that of high-grade tumors—translates into the existence of different GEP, which could contribute to a better understanding of the behavior of the tumor and patient outcome. Overall, our results based on a pilot series of 40 gliomas confirm the existence of a clear association between GEP of gliomas and tumor histopathology; in addition, for the first time we show that among grade IV astrocytomas, GEP are significantly associated with the cytogenetic profile of the ancestral tumor cell clone, further supporting its potential impact on the outcome of the disease.

Materials and Methods

Patients and Samples

A total of 40 consecutive adult patients—22 males (55%) and 18 females (45%)—diagnosed with primary gliomas were studied. Mean age at diagnosis was 57 ± 17 years (range: 21–84 years). In all cases, resected tumor samples were obtained by conventional surgical procedures, after informed consent was given by the patient, according to the recommendations of the local

Ethics Committee of the University Hospital of Coimbra (Coimbra, Portugal). For each case, the most relevant histopathological, clinical, and biological features of the disease were recorded, including patient age, gender, performance status (Karnofsky index), and tumor localization (Table 1). At the moment of closing this study, 24 patients (60%) had died and 7 had relapsed after a median follow-up of 15 and 18 months, respectively.

Tumor Histopathology

Histopathological studies included assessment of both the histological and morphological features of the disease evaluated at the moment of the diagnosis by an expert neuropathologist (eg, type of cell of origin, atypia, variation in nuclear shape or size accompanying hyperchromasia, cell size, cell shape, cellular organization, endothelial proliferation, mitotic activity, overall [eg, increased] cellularity, necrosis, and/or microvascular proliferation).

Histological diagnosis and classification of the tumors were performed according to the WHO criteria²² and patients were distributed as follows: 35 cases (87.5%) were astrocytomas (grade I pilocytic astrocytomas, 3 cases; grade II diffuse astrocytomas, 2; grade III anaplastic astrocytomas, 2 patients; grade IV glioblastomas, 26, and grade IV gliosarcomas, 2 cases); 3 patients (7.5%) had oligodendrogliomas (1 grade II oligodendroglioma, and 2 grade III anaplastic oligodendrogliomas), and the remaining 2 cases (5%) corresponded to mixed oligoastrocytomas (both grade III anaplastic oligoastrocytomas).

In all cases, additional tumor samples were obtained at diagnostic surgery, cut into several pieces, and freshly frozen to be used afterwards for genetic and genomic studies. Only those parts of the tumor showing both macroscopical and microscopical infiltration were used for iFISH analyses as well as to extract RNA. The remaining tumor sample was fixed in formalin and embedded in paraffin. Infiltration by tumor cells in tissue areas mirror-cut to those used for GEP was constantly >65%, as assessed by microscopical analysis of hematoxylin and eosin-stained tissue specimens.

iFISH Studies

In all cases, iFISH analyses were performed on freshly frozen single tumor cell suspensions obtained by mechanical disaggregation of tumor samples, after fixation in methanol/acetic 3/1 (v/v) (Panreac). For iFISH analyses the following commercially available dual color probes (DCP)—all obtained from Vysis, Inc, except the 7p12/alphasatellite 7 DNA DCP that was obtained from Q-BIOgene—were systematically used in double-stainings, for the detection of numerical abnormalities of 16 different chromosome regions that have been reported as frequently altered in gliomas (Supplementary Material, Table S1): (i) for chromosome 1, the LSI 1p36/LSI 1q25 DCP was used; (ii) for

Table 1. Clinicobiological characteristics of glioma patients and their corresponding tumors analyzed by both interphase FISH and DNA oligonucleotide expression arrays ($n = 40$)

Case ID	Histology	WHO grade	Ancestral tumor cell clone	Age	Gender	Karnofsky index (%)	Patient status	Overall survival (months) ^b
1	Pilocytic astrocytoma	I	Tetraploidy	21	F	80	Alive	24
2	Pilocytic astrocytoma	I	Tetraploidy	50	F	90	Alive	22
3	Pilocytic astrocytoma	I	+7+10q	34	F	90	Alive	10
4	Difuse astrocytoma	II	+7p	67	F	70	Dead	11
5	Difuse astrocytoma	II	Tetraploidy	38	M	90	Alive	27
6	Anaplastic astrocytoma	III	+7+10q	30	M	80	Dead	4
7	Anaplastic astrocytoma	III	Tetraploidy	27	M	90	Alive	28
8	GBM	IV	Amp 7p	67	F	60	Dead	15
9	GBM	IV	Amp 7p	50	F	80	Dead	13
10	GBM	IV	Amp 7p	71	F	60	Dead	8
11	GBM	IV	Amp 7p	70	M	80	Alive	18
12 ^a	GBM	IV	Amp 7p	70	F	70	Alive	17
13 ^a	GBM	IV	Amp 7p	45	F	60	Alive	15
14	GBM	IV	Amp 7p	48	M	90	Dead	13
15	GBM	IV	Amp 7p	74	M	60	Alive	8
16	GBM	IV	+7	69	F	60	Dead	1
17	GBM	IV	+7	79	M	80	Dead	4
18	GBM	IV	+7	62	M	60	Dead	3
19	GBM	IV	+7	67	M	60	Dead	2
20 ^a	GBM	IV	+7q	70	F	80	Dead	18
21	GBM	IV	+7q	35	F	80	Dead	14
22 ^a	GBM	IV	+7q	39	F	80	Dead	19
23 ^a	GBM	IV	+7p	30	F	80	Alive	62
24	GBM	IV	+7p	69	M	60	Dead	5
25	GBM	IV	-9p	67	F	70	Dead	9
26	GBM	IV	-9p	50	F	50	Dead	2
27	GBM	IV	-9p	73	M	70	Dead	2
28	GBM	IV	+7p-9p	66	M	80	Dead	25
29	GBM	IV	+7p-9p	74	M	70	Dead	1
30	GBM	IV	+7p-9p	84	M	70	Alive	10
31	GBM	IV	+7p-9p	56	M	80	Alive	8
32	GBM	IV	-10q	76	F	60	Alive	11
33	GBM	IV	-10q	60	M	60	Dead	2
34	Gliosarcoma	IV	+17p+19q	59	M	70	Dead	8
35 ^a	Gliosarcoma	IV	+7+10q+17p	39	M	80	Dead	21
36	OL	II	-1p-19q	37	M	90	Alive	11
37	AO	III	-9	64	M	70	Dead	5
38	AO	III	+7p-9p	53	M	70	Alive	11
39	Mixed AOA	III	+7-9p	81	F	70	Dead	3
40 ^a	Mixed AOA	III	-9p	65	M	70	Alive	17

Abbreviations: FISH, fluorescence in situ hybridization; F, female; M, male; GBM, glioblastoma multiforme; OL, oligodendroglioma; AO, anaplastic oligodendroglioma; AOA, anaplastic oligoastrocytoma.

^aRecurrent tumor.

^bOverall survival defined from diagnosis to the last visit or date of death.

chromosome 19, the LSI 19q13/LSI 19p13 DCP was employed; (iii) for chromosome 7, the LSI *ELN*/LSI 7q31 DCP was applied; (iv) for chromosomes 9 and 22, the LSI *bcr/abl* ES DCP was used; (v) in addition, for chromosome 9, the LSI 9p21/CEP-9 DCP was also employed, and (vi) for chromosome 10, the LSI

PTEN/CEP-10 DCP was applied. For chromosomes 13 and 17, the LSI *Rb1* and LSI *p53* probes conjugated with spectrum orange were used, respectively.

An Axioscope fluorescence microscope equipped with a 100 \times oil objective (Zeiss) was used to count the number of hybridization spots per nuclei ($n \geq 200$) for

each slide. Only those spots with a similar size, intensity, and shape in nonoverlapping nuclei were evaluated; doublet signals were considered as single spots. The extensive description of the iFISH techniques and the cut-offs used to define the presence of numerical abnormalities, for each individual chromosome, have been previously described in detail.²³ The definition of the major tumor cell clones present in each sample was arbitrarily based on the presence of $\geq 10\%$ nuclei carrying identical numbers of hybridization spots for all probes analyzed in that tumor sample.^{23,24} On the basis of iFISH results a clonal hierarchy was defined at the intratumoral cell level within each tumor. Accordingly, for each tumor the ancestral tumor cell clone was defined as being formed by those cells carrying cytogenetic changes shared by all tumor cells in the sample. In turn, secondary tumor cell clones were identified as those carrying genetic abnormalities, which were only present in a fraction of the tumor cells in the same sample, as previously defined.²¹

Isolation of Tumoral RNA and Microarray Analysis of GEP

Frozen tissue samples stored at -80°C were used for GEP studies. After thawing, tumor tissue samples were homogenized using a Potter-'S'-Elvehjem homogenizer (Uniform). Total RNA was isolated in 2 steps using TRIzol (Invitrogen Life Technologies) and the RNeasy Mini Kit (QIAGEN). The integrity and purity of the extracted RNA were determined using a microfluidic electrophoretic system (Agilent 2100 Bioanalyzer, Agilent Technologies). GEP were analyzed with the Gene Chip Human Genome U133Plus2.0 Array (Affymetrix Inc.) according to the manufacturer's instructions, using the 1-cycle cDNA synthesis kit and the Poly-A RNA gene chip control kit (Affymetrix, Inc.).

Analysis of Microarray Data

Owing to the relatively limited number of tumors analyzed, a carefully defined statistical plan was established prior to data analysis. First, the whole set of probes contained in the array was used in a prospective unsupervised exploratory analysis of all tumor samples to identify the most significant differences in GEP among the distinct subtypes of gliomas. Subsequently, a classification approach was attempted, aimed at identifying the best combination of genes to define the specific GEP subgroups identified in the first step. Special emphasis was placed in the investigation of the associations existing between the GEP and both tumor histopathology and cytogenetics.

Briefly, data files containing expression levels for the 40 tumors—obtained through the analysis of the Affymetrix oligonucleotide arrays—were normalized using the Robust Multi-array Average (RMA) expression measure²⁵ and analyzed using the R (version 2.7.1; <http://www.r-project.org>) and Bioconductor (<http://www.bioconductor.org>) software tools. In a first step, a prospective unsupervised

exploratory analysis was performed to identify the structure of the data at the gene expression level, using all probe sets contained in the array, for which RNA expression was quantified, except the Affymetrix control sequences (total of 54 613 out of 54 675 probe sets); then, a heatmap with a hierarchical cluster was built using euclidean distances and complete linkage. A multidimensional scaling (MDS) analysis—a powerful statistical tool for the multivariate identification of those variables associated with the highest contribution in explaining differences (separability) among individual samples or groups of samples—including those 6000 probes showing the highest level of variation among samples was performed to identify the most relevant differences between groups of tumors.

In a second step, a classification process was performed in order to derive a quantitative measure of class predictability and select the most relevant genes related to WHO tumor grade. An iterative classification process based on random forest algorithms was used²⁶ to rank gene classification power. This approach was selected because it is particularly well-suited for the analysis of a relatively limited number of cases (as it applies for the 40 tumors analyzed in our study); all cases were used for the initial test set. In brief, the first iteration started with the whole collection of probe sets, and a classifier was built taking all of them into consideration. In each of the subsequent iterations, those 90% genes showing the greatest classification power in the previous iteration were used to build the new classifier. In each of the following steps, in order to identify the family of genes with the greatest classification power, 100 different iterations with a data set containing all cases were performed, and a random forest was built and tested on them; all except the 2 gliosarcoma cases were considered in this part of the analysis ($n = 38$ tumors). The percentage of error obtained in the 3800 (38 testing cases \times 100 iterations) classification trials was used to measure the overall classification power of each family of genes. The most informative genes were chosen to be those from the selected family of genes associated with the lowest error (false discovery rate, FDR). The process was only stopped when there were no more genes available. In order to assure the reproducibility of the classification obtained, only those genes with very low FDR ($P < 0.001$) were selected into the model.

A similar unsupervised analysis to that described above was specifically repeated for the 26 GBM tumors and new significance and class prediction analyses were performed using the R, SAM, and PAM software programs (Significance Analysis of Microarrays software version 2.23 and Prediction Analysis of Microarrays software version 2.1, Tibshirani Lab, Department of Statistics, University of Stanford), respectively.

Other Statistical Methods

In order to assess the statistical significance of differences observed between groups of patients, the

Student's *t*-test and the Mann–Whitney *U* test were used for parametric and nonparametric (continuous) variables, respectively; for qualitative variables, the χ^2 test was applied (SPSS software, SPSS 15.0, SPSS Inc.). Survival curves were plotted according to the method of Kaplan and Meier and the log-rank test was used to assess the statistical significance of differences observed in survival between distinct groups of patients (SPSS software). *P*-values ≤ 0.05 were considered to be associated with statistical significance.

Results

GEP of Gliomas

GEP of gliomas as measured by DNA oligonucleotide microarrays were highly heterogeneous and were associated with tumor grade and histopathology (Fig. 1). Unsupervised hierarchical clustering of all tumors analyzed ($n = 40$) according to their GEP, showed that gliosarcomas ($n = 2$) were those tumors that more significantly differed from all other tumors; in turn, the latter clustered into 2 major groups, one of which included all GBM (Fig. 1A and B). Among those 72 genes showing the highest discriminating power between gliosarcomas and other gliomas, 18 were overexpressed in the former group, and 54 were underexpressed. Of note, overexpressed genes included *KRT80* gene (keratin 80) coded at 12q13.13 and the *BAIAP2L1* gene (BAI1-associated protein 2-like 1) coded at 7q21.3, whereas the *PTPRZ1* gene (protein tyrosine phosphatase, receptor-type, Z polypeptide 1) coded at 7q31.3 and the *GFAP* gene (glial fibrillary acidic protein) coded at 17q21 were among the most relevant underexpressed genes. A more detailed description of these and other relevant genes is shown in Supplementary Material, Table S2.

A supervised hierarchical clustering analysis (Fig. 1C) with a classifier built on those 71 genes that would mostly contribute to discriminate between low-grade—grade I and II—vs high-grade—grade III and IV—gliomas (after excluding gliosarcomas), showed a high classification power with only 6% of the cases being misclassified (Supplementary Material, Table S3). Interestingly, one-fourth of the 71 selected genes corresponded to genes coded in chromosome 10 (21/71), 7/71 genes were coded in chromosome 17, and 3 genes were coded in chromosomes 7 and 22, each. The remaining 36 genes were spread among the other chromosomes, with the exception of chromosomes 2, 3, 16, 18, and 21, which did not contain informative genes differentially expressed in the 2 groups of patients. The 6 most informative genes included the *GRAMD1B* gene (GRAM domain containing 1B) coded at 11q24.1, the *PITPNC1* gene (phosphatidylinositol transfer protein, cytoplasmic 1) localized at chromosome 17q24.2, the *C1orf21* gene (chromosome 1 open reading frame 21) at 1q25, the *VPS26A* gene (vacuolar protein sorting 26 homolog A) at 10q21.1, the *MSRB2* gene (methionine sulfoxide reductase B2) at 10p12, and the *MGEA5*

(meningioma expressed antigen 5) coded at 10q24.1–q24.3. Interestingly, 19 of these 71 genes had been either previously specifically related to gliomas or generally involved in carcinogenesis through their participation in the regulation of cell proliferation, apoptosis, DNA repair, transcription, and signal transduction: *EXO1* (1q42–q43), *CBX8* (17q25.3), *NCOA4* (10q11.2), *SGMS1* (10q11.2), *ZYX* (7q32), *TK1* (17q23.2–q25), *SCAPER* (15q24), *CENPE* (4q24–q25), *BLOC1S2* (10q24.31), *DLG7* (14q22.3), *CDC45L* (22q11.21), *AURKB* (17p13.1), *CDCA5* (11q12.11), *CHAF1A* (19p13.3), *RECQL4* (8q24.3), *CCNA2* (4q25–q31), *BIRC5* (17q25), *GTSE1* (22q13.2–q13), and *UBE2C* (20q13.12) genes.

The clinico-biological and cytogenetic characteristics of the 2 groups of gliomas distributed according to tumor grade are summarized in Table 2. As displayed in it, low-grade gliomas more frequently occurred in younger adults (< 40 years) ($P = 0.01$) and they showed both a higher Karnofsky index ($P = 0.002$) and a better outcome ($P = 0.03$); in addition, they displayed a lower frequency of both multilobular and temporal tumors ($P < 0.001$), lower number of neoplastic cell clones per tumor ($P = 0.04$), and a higher frequency of tetraploid ancestral tumor cell clones ($P = 0.008$).

GEP of Grade IV Astrocytomas

Owing to the relatively high heterogeneity of the GEP of grade IV astrocytic tumors (Fig. 1B and C), we further investigated the potential association between the gene expression signatures and the genetic, clinical, and biological features of the disease, specifically among grade IV astrocytomas/GBM. Unsupervised hierarchical clustering analysis of the GEP of the 26 GBM showed 3 clearly distinct groups of tumors for the 54 613 probes analyzed (Fig. 2A).

Overall, 47 genes were identified, which allowed for a clear discrimination among the 3 groups of GBM showing different GEP (Fig. 2B and Table 3). Four of these 47 genes have been previously shown to be involved in the pathogenesis of gliomas acting either as tumor suppressor genes, or being involved in signal transduction: *EGFR* (7p12), *ENPP2* (8q24.1), *GAS1* (9q21.3–q22), and *FTL* (19q13.3–q13). Further analyses searching for the best combination of genes to predict for the classification of GBM into the three GEP groups identified a combination of 79 informative genes (Fig. 3A and Supplementary Material, Table S4). These genes were coded in 16 different chromosomes: 8 genes in chromosomes 7 and 5 each; 7 in chromosome 4; 6 in chromosomes 1, 3, and 12; 5 in chromosome 8; 4 genes in chromosomes 10 and 11, and either 3 or 2 genes localized in chromosomes 2, 6, 9, 13, 14, 15, 17, 18, 19, 20, and 22. Once again, among the most discriminating genes, the *EGFR* (epidermal growth factor receptor) gene was selected, together with the *SOCS2* (suppressor of cytokine signaling-2) (both overexpressed in GEP 1 cases), *SPATA18* (spermatogenesis/

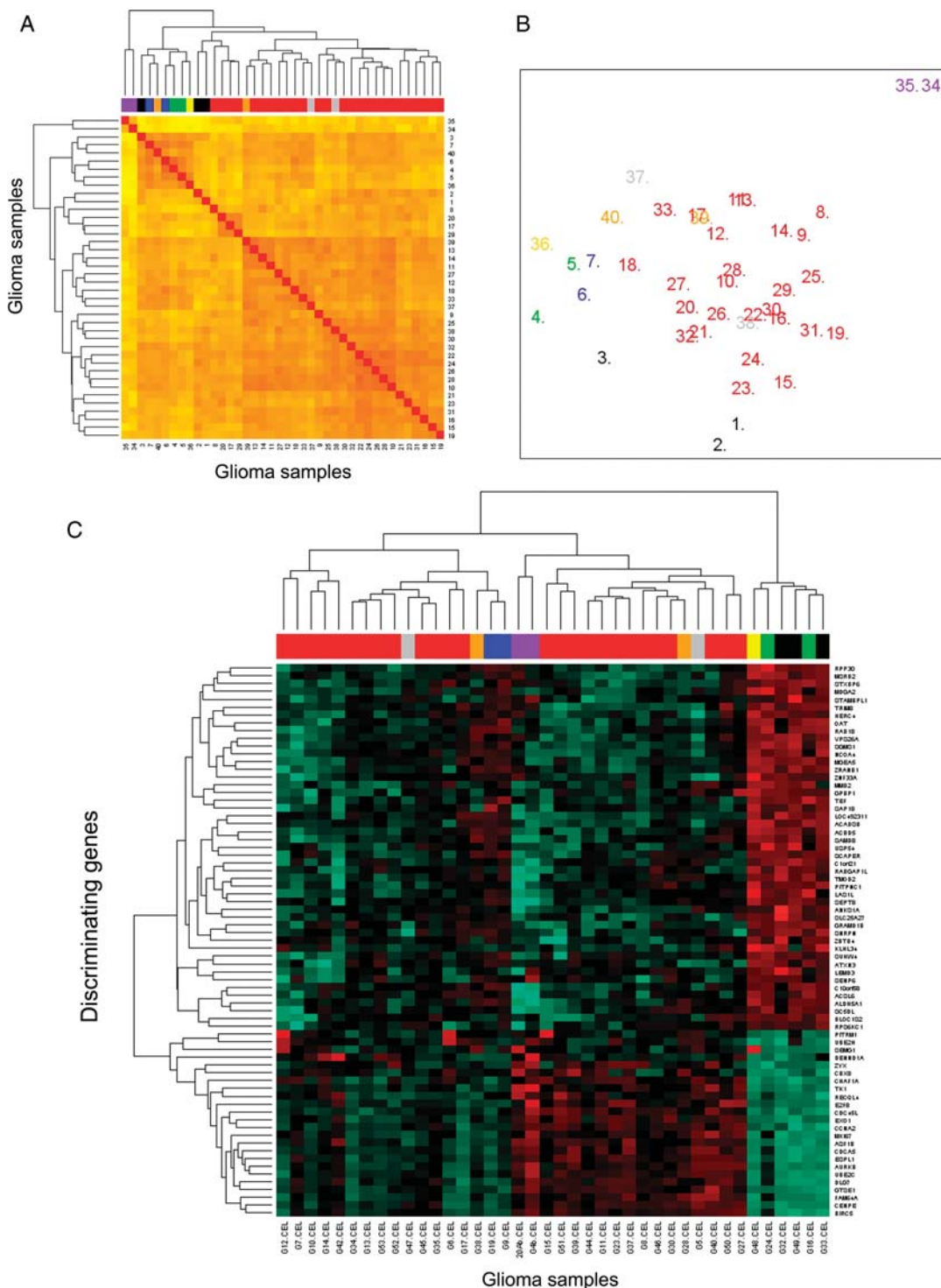


Fig. 1. Unsupervised and supervised classification of glioma samples based on their GEP. In (A) unsupervised classification of the tumors according to the 54 613 probes included in the Gene Chip Human Genome U133 Plus 2.0 array are shown; in (A) different histopathological subtypes of gliomas ($n=40$) are identified by a distinct color code: pilocytic tumors are coded black; diffuse astrocytomas are identified as green; anaplastic astrocytomas, blue; glioblastoma multiforme, red; gliosarcomas, purple; oligodendrogliomas, yellow; anaplastic oligodendrogliomas, grey, and mixed oligoastrocytomas correspond to those cases depicted as orange. In (B) an MDS plot corresponding to the same 40 gliomas (identified according to their histopathologic features with the same color code as described for [A]) classified according to those 6000 genes showing the highest variability in their expression among the different tumors. In (C) is a heat map representation of the results reflecting a hierarchical cluster analysis based on the expression of those 71 genes showing the highest classification power for tumor grade/histopathology; in (C) the different histopathological subtypes of gliomas are also identified by the same color code as described for (A).

Table 2. Clinical and cytogenetic characteristics of glioma tumors ($n = 40$) grouped according to the WHO histological grade

	Low-grade (I–II) ($n = 6$)	High-grade (III–IV) ($n = 34$)	<i>P</i> -value
Age			
<40 years	4/6	6/34	0.01
>40 years	2/6	28/34	
Gender			
Male	2/6	20/34	0.25
Female	4/6	14/34	
Karnofsky index ^a	85 ± 8 (70–90)	71 ± 10 (50–90)	0.002
Tumor localization			
Frontal	2/6	12/34	<0.001
Temporal	0/6	12/34	
Parietal	1/6	0/34	
Occipital	1/6	0/34	
Posterior fossa	1/6	0/34	
Cerebellum	1/6	0/34	
Multilobular	0/6	10/34	
Ancestral tumor cell clone			
Amp7	0/6	8/34	0.008
+7q or +7p	1/6	5/34	
+7	0/6	4/34	
–9p/–9p	0/6	5/34	
–10q	0/6	2/34	
+7p–9p	0/6	6/34	
+7+10q	1/6	2/34	
+17p+19q	0/6	1/34	
Tetraploidy	3/6	1/34	
–1p–19q	1/6	0/34	
Number of tumor cell clones ^a (by iFISH)	3 ± 1 (2–4)	5 ± 2 (2–9)	0.04
Number of relapses	0/6	7/34	0.22
Number of deaths	1/6	22/34	0.03

Abbreviation: WHO, World Health Organization. Results expressed as number (N) of cases/total cases.

^aMean ± one standard deviation (range).

phosphodiesterase), *SLC39A12* (solute carrier family 39, zinc transporter) (overexpressed in GEP 2 tumors), *ENPP2* (ectonucleotide pyrophosphatase/phosphodiesterase), and *SULF1* (sulfatase-1) (underexpressed in GEP 1 and overexpressed in both GEP 2 and GEP 3 cases) genes localized at the 7p12, 12q, 4q12, 10p12.33, 8q24.1, and 8q13.2–q13.3 chromosome regions, respectively (Fig. 3A and Supplementary Material, Table S4).

Interestingly, a significant ($P = 0.001$) association was found between the GEP and the cytogenetic patterns of the ancestral tumor cell clones of GBM (Table 4). Accordingly, GEP 1 and GEP 2 cases systematically showed isolated chromosome 7 abnormalities in the ancestral tumor cell clone: although GEP 1 tumors typically showed amplification of the *EGFR* gene in chromosome 7p, GEP 2 cases displayed partial gains of either the long- or the short arm of chromosome 7 in their

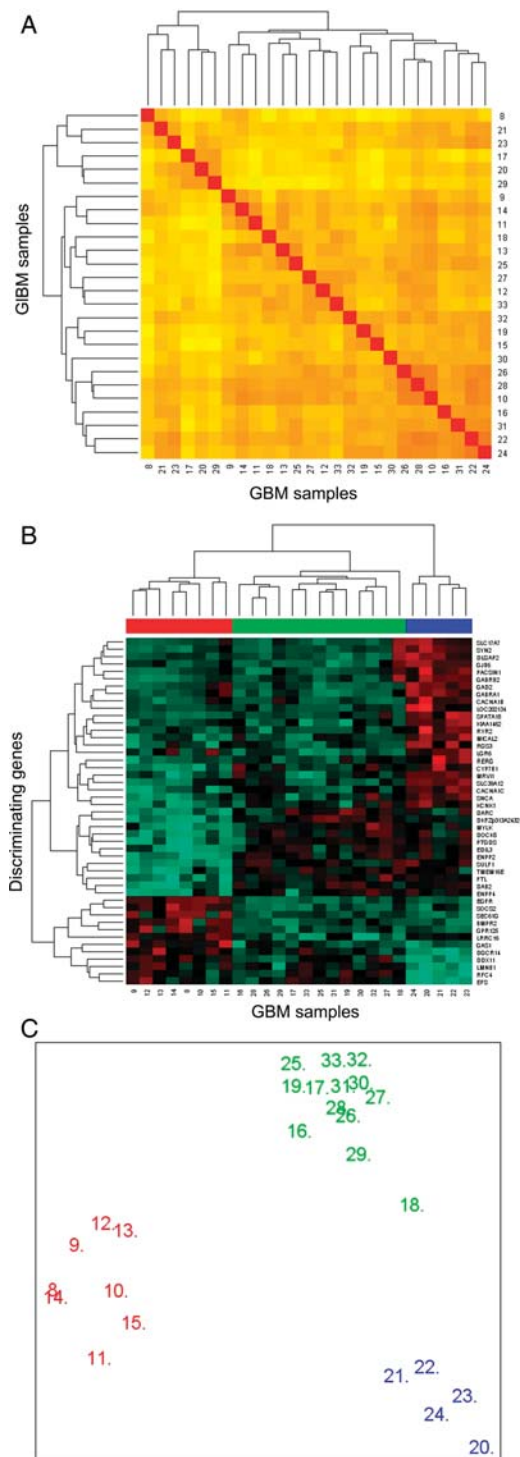


Fig. 2. Unsupervised and supervised classification of GBM ($n = 26$) according to their GEP. In (A) unsupervised classification based on the 54,613 probes included in the GeneChip U133A plus 2.0 array is shown. In turn, in (B) hierarchical clustering based on those 47 genes with the highest discrimination power as identified by significance microarray analysis (SAM) shows 3 groups of tumors with distinct GEP (GEP groups: GEP 1, GEP 2, and GEP 3 are colored red, blue, and green, respectively) is displayed. In (C) a MST plot corresponding to the same tumors classified according to their GEP and identified with the same color code as described above.

Table 3. Genes (*n* = 47) differentially expressed in the three groups of GBM identified by their GEP

Gene name	Gene symbol	Chromosomal localization	GEP tumor groups		
			GEP 1	GEP 2	GEP 3
Duffy blood group, chemokine receptor	DARC	1q21–q22	–	+	+
Leucine-rich repeat containing G protein-coupled receptor 6	LGR6	1q32.1	–	+	–
Ryanodine receptor 2 (cardiac)	RYR2	1q42.1–q43	–	+	–
Potassium channel, subfamily K, member 1	KCNK1	1q42–q43	–	+	–
Bone morphogenetic protein receptor, type II	BMPR2	2q33–q34	+	–	–
Synapsin II	SYN2	3p25	–	+	–
Myosin, light chain kinase	MYLK	3q21	–	+	+
Replication factor C (activator 1) 4, 37 kDa	RFC4	3q27	+	–	+
G protein-coupled receptor 125	GPR125	4p15.31	+	–	–
Spermatogenesis associated 18 homolog	SPATA18	4q12	–	+	–
Synuclein, alpha	SNCA	4q21	–	+	–
Disabled homolog 2, mitogen-responsive phosphoprotein	DAB2	5p13	–	+	+
EGF-like repeats and discoidin I-like domains 3	EDIL3	5q14	–	+	+
Lamin B1	LMNB1	5q23.3–q31.1	+	–	+
Gamma-aminobutyric acid (GABA) A receptor, beta 2	GABRB2	5q34	–	+	–
Gamma-aminobutyric acid (GABA) A receptor, alpha 1	GABRA1	5q34–q35	–	+	–
Hypothetical protein LOC202134	LOC202134	5q35.2	–	+	–
Ectonucleotide pyrophosphatase/phosphodiesterase 4	ENPP4	6p12.3	–	+	+
Protein kinase C and casein kinase substrate in neurons 1	PACSIN1	6p21.3	–	+	–
Leucine rich repeat containing 16	LRRC16	6p22.2	+	+	–
Sec61 gamma subunit	SEC61G	7p11.2	+	–	–
Epidermal growth factor receptor	EGFR	7p12	+	–	–
Dedicator of cytokinesis 5	DOCK5	8p21.2	–	+	+
Discs, large (Drosophila) homolog-associated protein 2	DLGAP2	8p23	–	+	–
Sulfatase 1	SULF1	8q13.2–q13.3	–	+	+
Cytochrome P450, family 7, subfamily B, polypeptide 1	CYP7B1	8q21.3	–	+	+
Ectonucleotide pyrophosphatase/phosphodiesterase 2	ENPP2	8q24.1	–	+	+
Growth arrest-specific 1	GAS1	9q21.3–q22	+	–	–
Regulator of G protein signaling 3	RGS3	9q32	–	+	–
Calcium channel, voltage-dependent, N type, alpha 1B subunit	CACNA1B	9q34	–	+	–
Prostaglandin D2 synthase	PTGDS	9q34.2–q34.3	–	+	+
Glutamate decarboxylase 2	GAD2	10p11.23	–	+	–
KIAA1462	KIAA1462	10p11.23	–	+	–
Solute carrier family 39 (zinc transporter), member 12	SLC39A12	10p12.33	–	+	–
Hypothetical protein DKFZp313A2432	DKFZp313A2432	11p14.2	–	+	+
Transmembrane protein 16E	TMEM16E	11p14.3	–	+	+
Murine retrovirus integration site 1 homolog	MRV1	11p15	–	+	–
Microtubule associated monooxygenase, calponin and LIM 2	MICAL2	11p15.3	–	+	–
DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 11	DDX11	12p11	+	–	+
RAS-like, estrogen-regulated, growth inhibitor	RERG	12p12.3	–	+	–
Calcium channel, voltage-dependent, L type, alpha 1C	CACNA1C	12p13.3	–	+	–
Suppressor of cytokine signaling 2	SOCS2	12q	+	–	–
Gap junction protein, beta 6	GJB6	13q11–q12.1	–	+	–
Embryonal Fyn-associated substrate	EFS	14q11.2–q12	+	–	+
Solute carrier family 17, member 7	SLC17A7	19q13	–	+	–
Ferritin, light polypeptide	FTL	19q13.3–q13.4	–	+	+
DiGeorge syndrome critical region gene 14	DGCR14	22q11.21	+	–	+

Abbreviations: GBM, glioblastoma multiforme; GEP, gene expression profiles.

ancestral tumor cell clone. In turn, cases included in the GEP 3 group had more variable cytogenetic profiles in their ancestral tumor cell clones, mainly consisting of

isolated trisomy 7 (31%), del(10q) (15%), and del(9p)/null 9p alone (23%) or in combination with gains of chromosome 7 (31%).

Table 4. Chromosomal aberrations of GBM grouped according to their GEP

Tumor ID	GEP	Ancestral tumor cell clone		Additional chromosomal aberrations ^b											
				Chromosome losses					Chromosome gains						
8	1	Amp 7p	-1p	-9p	nul9p	-10q	-13q	-22q	+1p	+7+9q	+13q	+17p	+19q	+22q	
13 ^a	1	Amp 7p	-1p	-9p	nul9p	-10q	-13q	-17p	+1p	+7+9	+10q	+13q	+17p	+19q	+22q
10	1	Amp 7p	-1p	-9p	nul9p	-10q	-13q			+7+9q			+19q	+22q	
9	1	Amp 7p	-1p	-9p	nul9p	-10q			+1p	+7+9q	+13q	+17p	+19q	+22q	
11	1	Amp 7p			nul9p	-10q	-13q		+1p	+7+9q			+19q		
15	1	Amp 7p		-9p		-10q			+1p	+7+9	+10q	+13q	+17p	+19q	+22q
14	1	Amp 7p				-10q			+1p	+7+9	+13q	+17p	+19q	+22q	
12 ^a	1	Amp 7p				-10q			+1p	+7+9	+13q	+17p	+19q	+22q	
21	2		+7q			-10q	-17p	-22q		+7+9	+13q		+19q		
22 ^a	2		+7q			-10q		-22q	+1p	+7+9	+13q	+17p	+19q	+22q	
20 ^a	2		+7q		nul9p	-10q			+1p	+7+9q	+13q	+17p	+19q	+22q	
23 ^a	2		+7p		-9p	nul9p			+1p	+7+9	+10q	+13q	+17p	+19q	+22q
24	2		+7p					-22q	+1p	+7+9	+10q	+13q	+17p	+19q	+22q
19	3		+7	-1p-7q		-10q		-19q-22q	+1p	+7+9	+10q	+13q	+17p		
16	3		+7		-9p	-10q		-19q-22q	+1p	+7+9q		+13q	+17p	+19q	+22q
17	3		+7						+1p	+7+9	+10q		+17p	+19q	+22q
18	3		+7			-10q	-13q	-17p	-22q	+1p	+7+9	+13q	+17p	+22q	
25	3		nul9p		-9p	nul9p	-10q	-13q	-17p	+7+9q	+10q		+17p	+19q	+22q
27	3		-9p	-1p	-9p	-10q	-13q	-19q-22q	+1p	+7+9	+10q	+13q	+17p	+19q	+22q
26	3		-9p		-9p	nul9p	-10q		+1p	+7+9q	+13q	+17p	+19q		
28	3		+7p-9p	-1p	-9p	-10q			+1p	+7+9q	+13q	+17p	+19q	+22q	
31	3		+7p-9p		-9p	nul9p	-10q	-19q	+1p	+7+9q	+13q	+17p	+19q	+22q	
29	3		+7p-9p		-9p	nul9p		-13q	+1p	+7+9q	+10q	+17p	+19q	+22q	
30	3		+7p-9p		-9p	nul9p			+1p	+7+9q	+10q	+13q	+17p	+22q	
32	3		-10q	-1p	-9p	nul9p	-10q	nul10q	-13q	-19q-22q		+7+9q	+17p	+22q	
33	3		-10q			-10q	-17p		+1p	+7+9	+13q	+17p	+19q	+22q	

Abbreviations: GBM, glioblastoma multiforme; GEP, gene expression profiles.

^aRecurrent tumors.

^bPresent in tumor cell clones other than the ancestral tumor cell clone (ie, not common to all altered tumor cells in the sample).

membrane receptors and chromosome ORF genes. These findings point out the expression of unique structural and functional marker profiles in gliosarcomas that in addition to GFAP²⁸ could contribute to a more refined histopathological classification of gliosarcomas vs other glioblastomas. However, these results require further confirmation in larger series of patients from brain tumor trial consortia.

Interestingly, upon excluding gliosarcomas, hierarchical clustering analysis of GEP provided a classifier for low- vs high-grade gliomas based on the expression of 71 different genes; the most informative genes included the *GRAMD1B*, *PITPNC1*, *C1orf21*, *VPS26A*, *MSRB2*, and *MGEA5* genes, together with another 23 genes that had been previously related to gliomas and/or involved in oncogenetic events in gliomas and other tumors, DNA repair, regulation of transcription, cell division, cell proliferation, tumor cell growth, and metastasis.²⁹⁻³⁴ Overall, high-grade tumors were associated with GEP characterized by greater expression of genes associated with increased cell proliferation and survival.^{31,34,35} Among other genes these included the *DLG7*, *BIRC5* (survivin), and *UBE2C* (or *UBCH10*) genes coded at 14q22.3, 17q25, and 20q13.12, which have been involved in cell cycle regulation and stem cellness (eg, *DLG7*), cell division and suppression of apoptosis (eg, survivin), and regulation of cell cycle progression and the ontogenesis of astrocytic tumors (eg, *UBEC2C*).^{29,35-37} Of note, the *STXBP6* (syntaxin

binding protein 6-amisyn) gene encoded at 14q13 was the individual gene showing the highest differential expression between high- vs low-grade gliomas; despite the fact that the protein encoded by this gene is known to be involved in vesicle-mediated intracellular transport, the precise functional role and pathways related to this protein remain largely unknown.

Interestingly, low- vs high-grade tumors did not only show different GEP but they were also associated with different cytogenetic patterns. Accordingly, high-grade gliomas displayed a greater intratumoral cytogenetic heterogeneity (eg, they had a higher number of tumor cell clones), which probably reflects a greater genetic instability associated with a higher rate of accumulation of genetic aberrations.³⁸ This could be related at least to a certain extent to the differential expression in low- vs high-grade gliomas of genes involved in DNA repair (*CHAF1A* and *RECQL4*),³⁹ cell cycle and cell division (*TK1*, *CENPE*, *CDC45L*, *AURKB*, *CDC45*, *CCNA2*, and *GTSE1*)^{32,40} and apoptosis (*BLOC1S2*, *SGSM1*, and *BIRC5*).^{34,35,41} Alternatively, this could be also associated with the pattern of cytogenetic abnormalities detected in the ancestral tumor cell clone since some genetic abnormalities leading to tumor progression (eg, amplification of *EGFR*) were exclusively found among high-grade tumors whereas tetraploidy, a genetic abnormality typically associated with a better outcome,^{42,43} was restricted to low-grade astrocytomas.

Table 5. Clinical and cytogenetic characteristics of glioblastoma multiforme (GBM) patients grouped according to their tumoral gene expression profiles (GEP)

	GEP group			P-value
	GEP 1 (n = 8)	GEP 2 (n = 5)	GEP 3 (n = 13)	
Age				
<40 years	0/8	3/5	0/13	0.001
>40 years	8/8	2/5	13/13	
Gender				
Male	3/8	1/5	9/13	0.12
Female	5/8	4/5	4/13	
Karnofsky index ^a	70 ± 12 (60–90)	76 ± 9 (60–80)	67 ± 10 (50–80)	0.26
Tumor localization				
Frontal	3/8	1/5	4/13	0.50
Temporal	1/8	3/5	5/13	
Multilobular	4/8	1/5	4/13	
Ancestral tumor cell clone				
Amp7	8/8	0/0	0/0	<0.001
+7q or +7p	0/0	5/5	0/0	
+7	0/0	0/0	4/13	
–9p/–9p	0/0	0/0	3/13	
–10q	0/0	0/0	2/13	
+7p–9p	0/0	0/0	4/13	
Number of tumor cell clones ^a (by iFISH)	6 ± 2 (4–9)	4 ± 1 (3–5)	4 ± 1 (3–7)	
Median OS (months) ^b	13 ± 4 (8–18)	16 ± 8 (5–25)	6 ± 7 (0–25)	0.05
Number of relapses	2/8	3/5	0/13	0.01
Number of deaths	3/8	4/5	10/13	0.14

Abbreviation: OS, overall survival. Results expressed as number (N) of cases/total cases.

^aMean ± one standard deviation (SD).

^bMedian and range between brackets.

In line with this latter hypothesis, 3 subgroups of GBM could be established according to their GEP, which in turn were tightly associated with the cytogenetic profile of their ancestral tumor cell clones. Accordingly, EGFR amplification and overexpression emerged as typically associated with one of these GEP (GEP1) detected among GBM.^{44–46} In addition, GEP1 GBM patients also showed significantly higher expression of *SOCS2*, a gene whose methylation has been involved in the pathogenesis of GBM where it is associated with resistance to conventional therapy and patient outcome.⁴⁷

In contrast to GEP1 GBM, cases classified as GEP 2 typically showed isolated gains of one arm of chromosome 7 in their ancestral tumor cell clones in association with overexpression of the *SPATA18*, *SLC39A12*, and *ENPP2* genes. Although the former 2 genes are poorly characterized genes, they have been associated with cell differentiation and metal ion transport, respectively. In turn, *ENPP2* encodes for an autotaxin reported to be overexpressed in GBM⁴⁸ and promote tumor invasion⁴⁹ and was highly expressed in both GEP2 and GEP3 gliomas. Upon comparison with GEP1 and GEP2 tumors, GEP3 GBM typically showed more heterogeneous cytogenetic profiles in their ancestral tumor cell clones, which could contribute to explain their relatively more variable GEP. Of note, further cytogenetic evolution of tumor cell clones was frequently associated in all these GEP

subgroups of GBM with further gains of chromosome 7, del(9p), and del(10q) suggesting the simultaneous occurrence of activation of wild-type and/or mutated *EGFR* and loss of both *Ink4A/Arf* and *PTEN* tumor suppressor genes, which has been recently shown to induce a fully penetrant and rapid-onset, high-grade malignant glioma phenotype with prominent pathological and molecular resemblance to human GBM, in the CNS of adult mice.⁴⁶ Altogether, these observations suggest that the sequence of appearance of different genetic abnormalities in GBM could have a relevant impact on the GEP and the behavior of the tumor, independently of the specific combination of genetic abnormalities accumulated in secondary tumor cell clones. In other words, the sequence of appearance in a tumor of different genetic abnormalities will have a specific influence on the GEP of tumor cells, over the end result.

Interestingly, GEP of GBM also showed a clear association with patient age, the number of different tumor cell clones defined by iFISH, and survival, which points out the potential impact of the GEP of GBM in determining tumor behavior at both the cytogenetic and clinical levels.

In conclusion, in this pilot study, we show that GEP of human gliomas are significantly associated with tumor histopathology, unique GEP being found in low- vs high-grade gliomas. Most interestingly, among

GBM 3, distinct GEP were found that appear to be associated with early genetic tumor changes. Further studies in independent and larger series of patients are required to confirm these results.

Supplementary Material

Supplementary Material is available at *Neuro-Oncology* online.

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