Somatic copy number alterations are associated with EGFR amplification and shortened survival in patients with primary glioblastoma

Abstract

Glioblastoma (GBM) is the most common malignant primary tumor of the central nervous system. With no effective therapy, the prognosis for patients is terrible poor. It is highly heterogeneous and EGFR amplification is its most frequent molecular alteration. In this light, we aimed to examine the genetic heterogeneity of GBM and to correlate it with the clinical characteristics of the patients. For that purpose, we analyzed the status of EGFR and the somatic copy number alterations (CNAs) of a set of tumor suppressor genes and oncogenes. Thus, we found GBMs with high level of EGFR amplification, low level and with no EGFR amplification. Highly amplified tumors showed histological features of aggressiveness. Interestingly, accumulation of CNAs, as a measure of tumor mutational burden, was frequent and significantly associated to shortened survival. EGFR-amplified GBMs displayed both a higher number of concrete CNAs and a higher global tumor mutational burden than their no EGFR-amplified counterparts. In addition to genetic changes previously described in GBM, we found PARK2 and LARGE1 CNAs associated to EGFR amplification. The set of genes analyzed allowed us to explore relevant signaling pathways on GBM. Both PARK2 and LARGE1 are related to receptor tyrosine kinase/P13K/PTEN/AKT/mTOR-signaling pathway. Finally, we found an association between the molecular pathways altered, EGFR amplification and a poor outcome. Our results underline the potential interest of categorizing GBM according to their EGFR amplification level and the usefulness of assessing the tumor mutational burden. These approaches would open new knowledge possibilities related to GBM biology and therapy.

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**Introduction**

Glioblastoma, \(IDH\)-wildtype (GBM) is the most frequent malignant neoplasm of the human central nervous system (CNS). Although scientists and clinicians all over the world have made countless efforts over the past decades to improve the therapies for GBM patients, tumor tends to spread rapidly and to return after treatment, resulting in a very short median survival. The 2016-revised \(WHO\) classification has incorporated valuable molecular features for brain tumor classification \[1\]. However, while detection of \(IDH1\) mutation is now used in diagnosis for its prognostic meaning in glioma, the prognostic value of other common genetic characteristics, as \(EGFR\) amplification in GBM, remains unclear \[1,2\]. Recent research demonstrated an enhanced migratory behavior of cells within \(EGFR\)-amplified tumors, \[2\] supporting a relationship between \(EGFR\) status and the clinical course. This fact underlines the interest of deeper in the role of \(EGFR\) considering also the existence of different levels of \(EGFR\) amplification, as previous works have delineated \[3—6\].

GBM is characterized by both, inter- and intra-tumor heterogeneity, with great variations at the histological and the molecular levels \[1,7\]. This heterogeneity is responsible in part of drug resistance and treatment failure \[8,9\]. GBM heterogeneity reach levels that, even regarding \(EGFR\), several variants have been described; among them, variant III (\(EGFRvIII\)) promotes cell proliferation, angiogenesis and invasion in different models, \[10—12\] making worthy to give it special attention. The identification of differential targets among GBM \(IDH\)-wildtype genetic-subgroups could lead to reach better approaches to GBM management.

Among the multiple signaling pathways deregulated in cancer, GBM stands out by alterations in the receptor tyrosine kinase/\(PI3K/RPTEN/ AKT/mTOR\)-signaling pathway. In addition, both the \(CDKN2A/Cdk4/6\)/retinoblastoma and the \(p53/MDM2/p14ARF\) molecular pathways are also widely affected \[1,13—15\]. Those signaling alterations seem to be a core requirement for GBM pathogenesis and they are associated with poor prognosis \[16,17\]. Many groups have used high-throughput techniques for the genomic analysis of these pathways in GBM \[18—20\]. However, the enormous complexity of the results (in part because of tumor heterogeneity) usually leads to sum the data in relation to the chromosomal \(loci\) affected, more than gene-by-gene detail with exemption of a little number of well-known genes \[1,18\]. A novel approach to better understand the genetic results in cancer, considers the global extent of somatic copy number alterations (CNAs), introducing the term of tumor mutational burden \[21—23\] or CNV-load \[24\]. Despite different definitions according to the experimental design, this concept may be important in GBM, as it is a genetic feature that in several tumor types correlates with response to immune-checkpoint inhibitors \[21,23,24\]. Multiplex ligation-dependent probe amplification (MLPA) seems to be appropriate to explore concrete genetic changes but also the accumulation of alterations per case, as tumor mutational burden \[25,26\].

The aim of the present work is to characterize in a semi-guided way the genetic landscape of fresh primary GBM, \(IDH\)-wildtype, with different \(EGFR\) amplification status; we want to identify potential biological targets differentially distributed according to \(EGFR\), in order to improve, in the near future, the prognostic and therapy of this heterogeneous tumor.

**Material and methods**

**Patient samples and histological analysis**

This work included 46 tumor samples from patients that underwent surgery in the \(Hospital Clínico Universitario\) in Valencia. The study was reviewed and approved by the clinical investigation ethics committee at the \(Hospital Clínico Universitario\) (CEIC). Tumor samples were fixed in neutral-buffered formalin, embedded in paraffin, sectioned, and stained with hematoxylin-eosin. They were diagnosed according to the WHO classification criteria \[1\] as primary GBM by two different neuropathologists. Immunohistochemistry analysis (IHC) was performed on paraffin-embedded sections using the avidin-biotin peroxidase method. IHC was carried out using antibodies directed against glial fibrillary acidic protein (GFAP), Ki-67 (MIB1) and \(EGFR\) -clone H11 (all from Dako, Glostrup, Denmark). The proliferation rate was calculated as the percentage of MIB1 immunopositive nuclei. GFAP and \(EGFR\) expression were scored according to the staining intensity and the number of stained cells using previously described criteria: 0, no staining; 1, light or focal staining; 2, moderate staining present in 50% to 75% of the sample and 3, strong staining, present in more than 75% of the sample. For \(EGFR\) IHC analysis, 0—1 were defined as non-overexpression and 2—3 were considered overexpression of \(EGFR\) \[27\].

**DNA extraction and DNA sequencing for \(IDH1/R2\) and \(TP53\) mutations**

Genomic DNA was extracted from fresh tissue samples using a \(QIAamp DNA Mini Kit\) (Qiagen, Inc., Valencia, CA, USA) according to the manufacturers instructions. We analyzed by direct sequencing the genomic regions spanning wild-type R132 of \(IDH1\) and wild-type R172 of \(IDH2\). We also carried out \(TP53\) sequencing in four different PCR amplification reactions to analyze exons 5—8. PCR was performed using standard buffer conditions, 200 ng of DNA and an AmpliTaq Gold Master Mix (Thermo Fisher Scientific). PCR products were purified with \(Centricon\) columns (Amicon, Beverly, MA, USA) and they were analyzed on an \(ABI 3100 \text{ Sequence} \) (Applied Biosystems, Foster City, CA, USA). Primer sequences forward (fw) and reverse (rv) were as follow: \(IDH1\) fw 5-ACCAATGCGACCACTAGCA, \(IDH1\) rv 5-TCAACA TTATTGCG-CAACATGACTT, \(IDH2\) fw 5-CCAATGGAACTATCCCGGAAC, \(IDH2\) rv 5-CCTCTCCAACCTGGGC CTAC, \(TP53\) (e5) fw 5-CAGCCCGTGTCTCCTCAGG, \(TP53\) (e5) rv 5-TTCAACTCTG TCTCCTTCT, \(TP53\) (e6) fw 5-GTCCGGCCCCTTCC AGCAT, \(TP53\) (e6) rv 5-GTCTGCGCCCCTTCCAGCAT, \(TP53\) (e7) fw 5-CTCATCTTG GGCCTGTGTTA, \(TP53\) (e7) rv, 5-AGTTGCGAGGGTGCGCAATGT, \(TP53\) (e8) fw 5-ACCTGATTTCC TTACTGCTTTGTC and \(TP53\) (e8) rv 5-GTCCCTGCTGTTCCTAC TC.

**Fluorescence in situ hybridization analysis (FISH)**

To evaluate the \(EGFR\) gene amplification status in this series, we used FISH on tissue microarrays (TMAs) that included representative samples from de 46 tumors studied. Firstly, we build eight TMAs using a \(QIAamp DNA Mini Kit\) (Qiagen, Inc., Valencia, CA, USA) according to the manufacturers instructions and the nuclei were counterstained with DAPI. Fluorescent hybridizations (FISH) were performed using the \(LSR-EGFP\) Spectrum Orange/\(CEP-7\) Spectrum Green Probe from \(Vysis\) (Abbott Laboratories, Downers Grove, IL, USA) according to the manufacturer’s instructions and the nuclei were counterstained with DAPI. Fluorescent signals were detected using a \(Leica DM4000B\) photomicroscope equipped with an appropriate filter set (Leica microsystems AG, Wetzlar, Alemania). Signals were counted in 100—150 non-overlapping tumor cell nuclei and their values were used to calculate the \(EGFR/CEP-7\) signal ratio. The \(EGFR\) gene was considered to be amplified when the \(EGFR/CEP-7\) signal was elevated.
ratio was > 2. The exact ratio was not calculated in cases with high amplification levels [28]. Cases were subclassified according to previous descriptions as H-amp GBMs when more than 20% of the cells showed more than 20 copies of EGFR; L-amp GBMs included cases with 5—20% of cells with 3—12 copies of EGFR and cases with 2 copies of EGFR composed the N-amp group of GBMs [3].

**Multiplex ligation-dependent probe amplification analysis (MLPA).**

MLPA was performed to determine somatic CNAs of multiple oncogenes and tumor-suppressor genes simultaneously. Among them, probes for all the exons of EGFR were included, allowing us to determine a wildtype status (wt) or the presence of the mutant form EGFRvIII, which losses exons 2—7, [29] in addition to CNAs. Two different MLPA kits (Salsa MLPA kit P105-C1 and Salsa MLPA KIT ME001-C2) were used following the manufacturers instructions (MRC-Holland, Amsterdam, Netherlands). Both kits are approved for investigational-use only. Amplification fragments were separated by capillary electrophoresis in an ABI 310 Sequencer (Applied Biosystems) and data analysis was performed using the Coffalyser excel-based MLPA analysis software (MRC-Holland).

The thresholds established to classify losses and gains of genetic material were set at 0.75 and 1.3, respectively. The set of probes included in these assays covered the following genes: APC, ATM, BCL2, BRCAM, BRCAM2, CASP8, CASR, CD27, CD44, CDH1, CDH13, CDK4, CDK6, CDKN1B, CDKN2A, CDKN2B, CHD7, CHFR, CREM, CTNNB1, DAPK1, EGFR, ER1, FHTT, GSTP1, HIC1, HRIP3, IGSF4, IL4, KLK3, LARGE1, MDM2, MIR26, MLH1, MLH3, MYBC3, MYOSB, NFKB1A, OCA2, PAX, PARK2, PDGFRα, PHD1, PTTEN, RAB7, RARB, RASSF1, RYBP, SAMHD1, SCNA1, TIMPD3, TP53, TSC2 and VHL. The collection of genes included allowed us to estimate the implication of the receptor tyrosine kinase/P3K/PTEN/AKT/mTOR-signaling pathway, the CDKN2A/CDKN2B/retinoblastoma-signaling pathway and the p53/MDM2/p14ARF signaling pathway. Considering MLPA results we have established different thresholds to assess the tumor genetic burden as accumulation of CNAs. In order to be able to take into account the dispersion of the data within the three groups formed on the basis of FISH-EGFR status, we have determined thresholds for losses, gains and global CNAs. This formula was the product of the maximum value and the average value, divided by the product of the variance and the asymmetry.

**Analysis of TCGA dataset**

To validate the association between EGFR amplification status and somatic CNAs in the genes mentioned above, we obtained the data for GBM samples from The Cancer Genome Atlas (TCGA) by using cBioPortal for Cancer Genomics (www.cbioportal.org) [20,30,31]. We studied the Genomic Profile Putative copy number alterations from GIS-TIC for latest dataset available in cBioportal for GBM (TCGA, Provisional 604 samples). Copy number alteration data from 577 cases were obtained and further analyzed. A User-defined List including EGFR, BCL2, CDK6, CDKN1B, CDKN2B, ER1, LARGE1, MYBC3, PARK2 and PAH was entered into the Enter Gene box. Samples were classified for each gene into 5 groups according to their putative copy number variation calculated by GISTIC with default cBioportal thresholds [30]. The groups were Diploid (0), Shallow Deletion (-1), Deep deletion (-2), Gain (1) and Amplification (2). The associations between EGFR amplification and the copy number profile were analyzed using the Plots tool and retrieving the raw data. A contingency table was built for expressing the distribution among groups of all samples according to the EGFR amplification status. A Fishers exact test was used for calculating statistical significance. Statistical significance for amplification vs diploid and deletion (shallow or deep) vs diploid for each gene was calculated.

**Statistical analysis**

The statistical analysis of the different analyzed parameters was carried out according to the type of variable. Quantitative variables (age, Karnovsky index, tumor size, survival and Ki-67) were evaluated using the Kolmogorov-Smirnov and Levene tests; depending on their results and their characteristics, two-tailed Students t-test, ANOVA, Mann-Whitney U test or Kruskal-Wallis test were performed. For comparisons among categorical variables, Fisher exact test calculator was used for 2 2 tables and Chi-square test was used for tables with higher number or rows/columns. We also carried out survival analyses using the Kaplan-Meier method. The statistical significance of these survival curves was calculated using the log-rank (Mantel-Cox) test. Significance was accepted at least at p < 0.05 level. To facilitate understanding, in addition to p value, symbols were added to significant results (* when p < 0.05), very significant results (**p < 0.01) and highly significant results (***/p < 0.001). Data were analyzed with SPSS (version 22) software (IBM, Madrid, Spain).

**Results**

**Patient population and histopathological study**

We studied a series of 46 primary GBM from adult patients; 29 patients were men and 17 women, reflecting the GBM male predominance. The patients ages ranged from 24-75, with a mean of 59 years. The tumor size ranged between one and eight cm³. Tumor location affected the frontal lobe in 12 cases, temporal lobe in 18 cases, parietal lobe in 13 cases, occipital lobe in two cases and one case in callosa commissure. The patients received, in addition to the surgical treatment, chemotherapy in 2 cases, radiotherapy (RT) in 18 cases, both RT and chemotherapy in 23 cases and 3 cases only received palliative care. Five patients that did not received RT tend to be older than the other 41 patients (68.4 ± 4.0 and 57.2 ± 1.3, p = 0.058). Overall survival for these patients averaged 13.2 months, without differences between women and men. Histologically, all the tumors demonstrated features of GBM with pleomorphic, astrocytic tumor cells, prominent microvascular proliferation, and necrosis (Figure 1A—C). In every case, GFAP expression was confirmed in neoplastic cells. Ki-67 indicated a high proliferation potential with a mean value of 15% (Figure 1D). EGFR expression was variable both among tumors and among different regions of the same tumor. In 24 cases, a level 2—3 was determined and 8 cases scored level 1 (Figure 1E—F). The remaining 14 cases were negative for EGFR expression. The clinical data and the histopathological results are summarized in Table 1.

**Sequencing studies of IDH1/IDH2 and TP53**

Genomic analysis encompassing the R132 region of IDH1 and the R172 region of IDH2 revealed no mutations in any of the 46 cases studied. This fact confirmed their classification as IDH-wildtype GBMs, according to the most recent WHO classification [1]. Exons 5, 6, 7 and 8 on TP53 were also studied. TP53 mutations were found in case 29 (heterozygous substitution-missense mutation p.C141R, in exon 5) and in case 33 (heterozygous substitution-missense mutation p.R273C, in exon 8).

**EGFR characterization: amplification status is associated to EGFR protein expression and to EGFRvIII form**

EGFR copy number analysis by FISH was successfully assessed in the 46 primary GBMs and revealed EGFR amplification in 30 cases (65.2%).
Based on this analysis cases 1—20 showed a high level of $EGFR$ amplification (H-amp group, Figure 1G), cases 21—30 showed a low level of $EGFR$ amplification (L-amp group, Figure 1H) and cases 31—46 displayed no $EGFR$ amplification (N-amp group, Figure 1I). Immunohistochemical analysis of $EGFR$ protein expression is shown in Table 1. It revealed a strong statistical association between the genetic status of $EGFR$ determined by FISH and the protein levels detected by IHC ($p = 4.3E^{-6}***$). The receptor was overexpressed in every case from the H-amp group (levels 2—3). One case showed $EGFR$ protein overexpression in the L-amp group and only three cases from the N-amp group displayed $EGFR$ protein overexpression. These results agreed with previous reports that confirm the appropriateness of FISH for the classification of $EGFR$ amplification [3]. MLPA study showed the presence of the mutant form of $EGFR$ gene, $EGFR_vIII$, in 9 cases. All of them were also H-amp GBMs ($p = 0.001***$). All these data are summarized in Table 1.

**Figure 1.** Histopathology, immunohistochemistry and fluorescence in situ hybridization (FISH) analysis of $EGFR$ in primary GBM. (A) Pleomorphic, astrocytic tumor cells, (B) prominent microvascular proliferation and (C) presence of necrosis. (D) Ki-67 positive cells ratio indicated a higher proliferation range. (E) $EGFR$ overexpression with a level 3 and (F) with a level 2 was present. (G) Tumor cells showing a high level of $EGFR$ amplification, (H) cells with a low level of $EGFR$ amplification and (I) cells without $EGFR$ amplification, determined by FISH.

$EGFR$ genetic status correlates with morphological and histological features of aggressiveness

The frequency of men was higher in both L-amp (8 men/ 2 women) and N-amp groups (11 men/ 5 women) whereas that difference disappeared in the H-amp group (10 men/ 10 women). The average size of the tumors was the greatest in the H-amp group (4.2 cm$^3$), followed by the L-amp group (3.7 cm$^3$) and the N-amp group (3.2 cm$^3$) but these differences did not reach statistical significance (supplementary file 1). $EGFR_vIII$ cases showed an average of 3.1 cm$^3$, which was significantly higher than 3.4 cm$^3$ in $EGFR$ wildtype (wt) cases ($p = 0.013^*$) but these differences disappear if we consider exclusively variant III and wild-type cases among $EGFR$ H-amp GBMs. Overall survival of the patients was similar among the different groups (12—14 months). Regarding Ki-67, the H-amp group presented a mean value of 20.3%. It was statistically higher than the average Ki-67 of the L-amp group and the N-amp group, which showed 14.4% and 8.7%, respectively ($p = 0.020^*$). In addition, $EGFR_vIII$ cases showed 27.8% Ki-67 compared to 11.8% in $EGFR_{wt}$ cases ($p = 0.013^*$).

Somatic copy number alterations are frequent in GBM and their accumulation is associated to shortened survival

Somatic copy number alteration (CNA) analysis showed losses and/or gains in, at least, one of the regions explored in all the tumor samples assayed (Table 2). CNAs were frequent in our series showing an average of 12.6 8.9 CNAs per patient. CNAs were statistically higher in the 41 patients that received RT than in the other 5 patients (5.8 0.8 and 2.8 0.9, respectively, $p = 0.025$). From the 54 genes studied, 12 presented losses in more than 20% of cases: $CDKN2A$ (54.3%), $CDKN2B$ (54.3%),
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PTEN (41.3%), CDH1 (37.0%), TP53 (34.8%), LARGE1 (34.8%), CREM (32.6%), MYBPC3 (32.6%), VHL (32.6%), GSTP1 (28.3%), SAMHD1 (28.3%) and PAH (23.9%). The losses for CDKN2A and CDKN2B were mostly in homozygosis, and they were confirmed by probes included into the two kits assayed. Likewise, six genes showed gains that frequency: EGFR (60.9%), CDK6 (60.9%), CHD7 (39.1%), RARB (28.3%), CDKN1B (26.1%), HIC1 (23.9%) and PARK2 (21.7%). These alterations were located on chromosomes 3, 6, 7, 8, 9, 10, 11, 12, 16, 17, 20 and 22 (Figure 2).

Tumors displaying CDK6 gains were significantly bigger, with a mean value of 4.2 cm³, compared to 3.0 cm³ in CDK6wt tumors (p = 0.038*). Furthermore, patients tumors with losses in LARGE1 tended to present statistically higher Ki-67, with an average of 24.0% compared to 10.3% (p = 0.038*). On the whole, cases under 7 CNAs showed 556.0 days while from that threshold it fell to 308.9 (p = 0.015*) although considering together losses and gains seems to lose biological perspective. Kaplan-Meier survival analysis showed significant associations through Long Rank (Mantel-Cox) statistic for both, >4 gains/patient (p = 0.036*) and >10 losses / patient (p = 0.009**) (Figure 3A, B). The statistical strength reduced when we consider together losses and gains, and Kaplan-Meier survival analysis offered significance through Breslow analysis. It is remarkable that all the cases above 10 copy number losses exhibited simultaneously more than 4 copy number gains among different loci.

EGFR amplified GBMs display a high number of concrete genetic copy number alterations

We analyzed the distribution of CNAs among the three EGFR amplification groups. Five genes presented significant CNA changes among them (Figure 3C). CDKN2A (p = 0.030*), CDKN2B (p = 0.030*), LARGE1 (p = 0.009**), PARK2 (p = 0.046*) and CDK6 (p = 0.046*). All these genes were statistically more affected in the H-amp group or both.

| Case | Age/Sex | Location | Treatment | Survival (months) | Ki-67 (%) | EGFR expression | EGFRampl. | EGFRwt/mutant |
|------|---------|----------|-----------|------------------|-----------|----------------|-----------|---------------|
| 1    | 56/F    | T        | RT + CH   | 23               | 5         | 3              | 2         | EGFRwt        |
| 2    | 72/F    | F        | RT + CH   | 8                | 20        | 3              | 2         | EGFRwt        |
| 3    | 63/F    | T        | RT + CH   | 11               | 8.5       | 3              | 3         | EGFRwt        |
| 4    | 69/M    | T        | RT        | 7                | 15        | 3              | 2         | EGFRwt        |
| 5    | 48/M    | T        | RT + CH   | 11               | 3         | 3              | 2         | EGFRwt        |
| 6    | 55/F    | CC       | CH        | 4                | 14.5      | 3              | 3         | EGFRwt        |
| 7    | 59/F    | P        | RT        | 18               | 40        | 3              | 3         | EGFRwt        |
| 8    | 58/F    | T        | RT + CH   | 23               | 35        | 3              | 2         | EGFRwt        |
| 9    | 61/M    | T        | RT + CH   | 12               | 4         | 3              | 3         | EGFRwt        |
| 10   | 59/F    | P        | RT + CH   | 20               | 30        | 3              | 3         | EGFRwt        |
| 11   | 66/M    | P        | RT + CH   | 6                | 19.7      | 2              | 3         | EGFRwt        |
| 12   | 55/M    | T        | RT + CH   | 5                | 6         | 2              | 2         | EGFRwt        |
| 13   | 66/F    | F        | RT        | 36               | 8         | 2              | 2         | EGFRwt        |
| 14   | 69/M    | T        | RT + CH   | 12               | 50        | 2              | 3         | EGFRwt        |
| 15   | 66/M    | F        | RT        | 5                | 45        | 3              | 3         | EGFRwt        |
| 16   | 61/F    | F        | RT        | 2                | 22.5      | 3              | 3         | EGFRwt        |
| 17   | 58/F    | O        | RT + CH   | 20               | 5         | 2              | 2         | EGFRwt        |
| 18   | 57/M    | F        | RT        | 17               | 12        | 2              | 3         | EGFRwt        |
| 19   | 71/M    | F        | RT + CH   | 3                | 20        | 2              | 3         | EGFRwt        |
| 20   | 63/M    | T        | RT        | 5                | 43        | 2              | 2         | EGFRwt        |
| 21   | 45/M    | T        | RT        | -                | 3         | 1              | 1         | EGFRwt        |
| 22   | 24/M    | T        | RT        | 2                | 30        | 1              | 1         | EGFRwt        |
| 23   | 67/M    | T        | RT + CH   | 7                | 43        | 0              | 1         | EGFRwt        |
| 24   | 73/M    | P        | RT        | 5                | 22        | 0              | 1         | EGFRwt        |
| 25   | 45/M    | P        | RT + CH   | 10               | 6.8       | 2              | 1         | EGFRwt        |
| 26   | 42/M    | P        | RT        | -                | 3         | 0              | 1         | EGFRwt        |
| 27   | 60/M    | P        | RT        | 5                | 2         | 1              | 1         | EGFRwt        |
| 28   | 31/F    | F        | RT        | -                | 3         | 1              | 1         | EGFRwt        |
| 29   | 73/F    | P        | RT + CH   | 26               | 1         | 1              | 1         | EGFRwt        |
| 30   | 35/M    | T        | RT        | 38               | 30        | 1              | 1         | EGFRwt        |
| 31   | 74/F    | F        | NONE      | 1                | 12        | 0              | 0         | EGFRwt        |
| 32   | 66/M    | T        | RT + CH   | 11               | 5         | 0              | 0         | EGFRwt        |
| 33   | 35/F    | P        | RT        | -                | 2         | 0              | 0         | EGFRwt        |
| 34   | 54/M    | T        | RT + CH   | 5                | 26        | 0              | 0         | EGFRwt        |
| 35   | 63/M    | F        | RT + CH   | 36               | 23        | 0              | 0         | EGFRwt        |
| 36   | 75/M    | P        | NONE      | 6                | 44        | 2              | 0         | EGFRwt        |
| 37   | 55/M    | F        | RT + CH   | 26               | 3         | 1              | 0         | EGFRwt        |
| 38   | 75/M    | O        | RT + CH   | 11               | 2         | 0              | 0         | EGFRwt        |
| 39   | 63/F    | P        | CH        | 9                | 2         | 1              | 0         | EGFRwt        |
| 40   | 67/M    | P        | RT + CH   | 21               | 7.5       | 2              | 0         | EGFRwt        |
| 41   | 75/F    | F        | NONE      | 2                | 2         | 0              | 0         | EGFRwt        |
| 42   | 60/M    | F        | RT + CH   | 2                | 1         | 0              | 0         | EGFRwt        |
| 43   | 50/M    | T        | RT        | 10               | 5         | 0              | 0         | EGFRwt        |
| 44   | 65/F    | T        | RT + CH   | 22               | 1         | 2              | 0         | EGFRwt        |
| 45   | 38/F    | P        | RT        | 30               | 1         | 0              | 0         | EGFRwt        |
| 46   | 52/M    | T        | RT        | 23               | 2         | 0              | 0         | EGFRwt        |

We analyzed the distribution of CNAs among the three EGFR amplification groups. Five genes presented significant CNA changes among them (Figure 3C). CDKN2A (p = 0.030*), CDKN2B (p = 0.030*), LARGE1 (p = 0.009**), PARK2 (p = 0.046*) and CDK6 (p = 0.046*). All these genes were statistically more affected in the H-amp group or both.

Table 1. Clinical and histopathological data and genetic status of EGFR in 46 GB cases. T: temporal; F: frontal; P: parietal; O: occipital; CC: callosa commissure; RT: radiotherapy; CH: chemotherapy; amp: amplification; wt: wild type; EGFRvIII: EGFR mutant form.
in H-amp and L-amp, compared to N-amp group (Figure 3C). CDKN2A, CDKN2B and LARGE1 displayed mainly losses, CDK6 showed gains and PARK2 exhibited both losses and gains. Taken together all the genetic changes detected, we found that somatic CNAs were lower in N-amp group, with 9.3 ± 7.6 CNAs, compared to L-amp and H-amp which accounted 14.8 ± 6.6 and 14.1 ± 10.1 CNAs, respectively (p > 0.050).

Genetic landscape of L-amp GBMs partially overlaps with both H-amp GBMs and N-amp GBMs

Then, we made an overview of the most relevant genes based on the frequency they were altered. We considered the five aforementioned genes and others that showed CNAs affecting at least 40% of the cases within an EGFR amplification group although they did not show statistical differences among all the three subgroups (MYBPC3, PTEN, BCL2, PAH, CREM, CDH1, CHD7, CDKN1B, RAR8, PARK2 and HIC1). H-amp group exhibited five genes reaching the 40% of cases with loss, L-amp group presented eight genes and N-amp only one gene (Figure 4A). These results reflect an increasing affectation from N-amp to H-amp and an overlapping profile between L-amp and H-amp. Regarding to the gains, H-amp group presented two genes reaching this threshold, L-amp group showed six genes and N-amp two genes (Figure 4B). The analysis of TCGA putative copy number alterations data for GBM showed that there are statistically significant associations between EGFR amplification and CDK6 amplification, CDKN2A and CDKN2B deletion, MYBPC3 deletion and PAH deletion (Table 3). CDKN1B, ESR1, PARK2 or LARGE1

| Table 2. Somatic copy number alteration analysis. Individual results (part I). |
|-----------------------------------|-----------------|-----------------|-----------------|
| **Gene**                          | **H-AMP CASES** | **L-AMP CASES** | **N-AMP CASES** |
| CDKN1                           | 09q21.3         |                 |                 |
| EGFRR                            | 07q11.2         |                 |                 |
| CNGT7                            | 08q12.2         |                 |                 |
| RASB                             | 05q24.2         |                 |                 |
| MYO1B                            | 18q11.1         |                 |                 |
| RYBP                             | 03q13           |                 |                 |
| MLH1                             | 03q22.2         |                 |                 |
| CDKN1B                           | 12p13.1         |                 |                 |
| HIC1                             | 17p13.3         |                 |                 |
| SCN1A                            | 02q34.3         |                 |                 |
| PDGFRB                           | 04q12           |                 |                 |
| IGSB4                            | 11q23.3         |                 |                 |
| MIR26                            | 12q14.15        |                 |                 |
| CTHB1G1                          | 03q22.1         |                 |                 |
| CASR                             | 03q21           |                 |                 |
| PKHD1                            | 09q12.3         |                 |                 |
| ATM                              | 11q22.3         |                 |                 |
| TSG5                            | 16p13.3         |                 |                 |
| BRCC1A                           | 17q21.31        |                 |                 |
| IL4                              | 19q13.1         |                 |                 |
| RAB7A                            | 03q21.3         |                 |                 |
| HRPR2                            | 16q11.2         |                 |                 |
| KL13                             | 19q13.3         |                 |                 |
| RASSF1                           | 09q31.21        |                 |                 |
| CCK       | 12q14.15        |                 |                 |
| CASP8                            | 02q33           |                 |                 |
| TIMP3                            | 10p12.1         |                 |                 |
| DAPK1                            | 09q21.33        |                 |                 |
| ESR1                             | 18q23.1         |                 |                 |
| MLH3                             | 14q24.3         |                 |                 |

The copy number alterations are color-coded. Gene losses are shown in red. Gene gains are shown in green. Abbreviation: H-AMP, high-EGFR amplification level; L-AMP, low-EGFR amplification level; N-AMP, no EGFR amplification.
copy number alterations are published in the TCGA collection but their association to EGFR amplification were not statistically significant.

Relevant signaling pathways on GBM are altered in EGFR amplified tumors and related to a poor outcome

From the 46 GBM studied cases, 84% presented CNAs in genes involved in the receptor tyrosine kinase\(\text{PI3K/PIK3/PI3K/PIK3/PI3K/PTEN}}\) signaling pathway (TRK-pat), 43.5% had gene alterations in genes from p53-pathway (p53-pat) and 65% in genes associated with RB-signaling pathway (RB-pat). CNAs in genes from TRK-pat were detected in all the samples belonging to H-amp group and in 90% of the samples in L-amp group, but only in 37.5% of N-amp cases \((p = 0.001^{**})\). Similar frequencies were observed for CNAs in genes from RB-Pat, but with closer percentages (90% in H-amp, 80% in L-amp and 56.3% in N-amp, \(p = 0.059\)). Affectionation of genes from RB-pat, was similar among the three EGFR amplification groups, with values ranging 30—50%.

Regarding the clinical characteristics of the cases, we found that patients that showed alterations on genes in TRK-pat and RB-pat presented bigger tumors: tumor size was 2.7 cm\(^3\) and 2.5 cm\(^3\) when TRK-pat or RB-pat were not altered and it was 4.1 cm\(^3\) when any of them were affected \((p = 0.019^{*} \text{ and } p = 0.013^{*}, \text{ respectively})\). In addition, Kaplan-Meier analysis showed that survival was significantly shortened when we found alterations in genes belonging to the three different pathways simultaneously in a case: it was 584 days in cases which alterations belong to only one pathway, 411 when two pathways were altered and 234 days when genes from the three pathways were implicated \((p = 0.016^{*})\). Although it did not reach significant meaning, 45% of H-amp GBMs showed the three pathways altered compared with 20.0% of L-amp and 18.8% of N-amp.

Discussion

EGFR amplification has been identified as a genetic hallmark of primary GBM and occurs in approximately 40—60% of primary GBMs, but rarely in secondary GBMs; \([28,32,33]\) this frequency is similar to the one detected in our series. EGFR genetic status was studied by two complementary techniques: FISH and MLPA. MLPA is an excellent tool to study EGFR amplification but also it has been used to demonstrate that EGFR\(\text{vIII}\) affects up to 67% of the GBMs with EGFR amplification \([13,34,35]\). However, it is not able to discriminate low levels of EGFR amplification, because of trisomies and polysomies of chromosome 7 \([5]\). For that reason, FISH was chosen to distinguish L-amp GBMs. A previous report from our group showed that high amplification presented as double minutes whereas low amplification presented as insertions into different loci on chromosome 7 \([3]\). EGFR genetic status use to correlate to the EGFR protein expression level \([28,33,36]\), fact that is broadly supported by our results with a high statistical meaning. In our study, all the H-amp GBMs presented overexpression of the protein. EGFR variable expression among L-amp GBMs indicates that there are cases closer to the H-amp group and others closer to the N-amp group. The presence of a little subgroup of N-amp cases with high EGFR protein levels suggests the existence of mechanisms for its expression, which would be independent to DNA amplification, as previous works have investigated and underlines the important role of EGFR in GBM \([37,38]\).

The use of MLPA and FISH allowed as to confirm the high frequency of EGFR amplification, both as L-amp and as H-amp, and the presence of EGFR\(\text{vIII}\) limited to H-amp GBMs, in agreement with previous reports \([11,12]\). We found statistically increased proliferation and tumor size in patients that presented EGFR\(\text{vIII}\) in concordance with previous works \([10—12]\); our association among H-amp/EGFR\(\text{vIII}/\text{tumor size/Ki-67}\) goes in favor of descriptions that propose a direct role of EGFR\(\text{vIII}\) on aggressiveness \([11,12,29,39]\).

In the present work, we show a semi-guided genetic analysis of many TSGs and oncogenes that are important in GBM, and we correlate these analyses to the EGFR amplification status. Although it would be desirable to confirm our results in larger series, the genetic study of fresh tumor specimens provides high quality data with low background making easy to analyze the information obtained by this technical approach.

CNAs were, as previously described \([18,40]\), frequent in our series supporting the classical idea of a high chromosomal and genetic instability, and they are also increased in response to RT \([7,14,41]\). The global inci-
The occurrence of CNAs was higher in both groups of tumors with EGFR amplification but even higher in EGFR H-amp GBMs than in L-amp tumors. In addition, the high frequency of concrete genes altered in the EGFR amplified GBMs, and especially in the EGFR H-amp group, indicated they would be directly involved in the genesis and progression of GBM.

CDK6 gains and LARGE1 losses were significantly associated to tumor size and Ki-67. Both genes, plus CDKN2A/B were statistically more altered in H-amp GBMs and all of them were validated with TCGA cohort data [30]. CDK6 and CDKN2A/B are well-known genes in GBM. It is necessary to remark the potential influence of LARGE1 and PARK2 in GBM related pathways (Figure 5). There is little described about them on GBM literature but both resulted significantly altered in EGFR amplified GBMs in this study. LARGE1 encodes a glycosyltransferase that participates in glycosylation of alpha-dystroglycan. Mutations in this gene cause diverse forms of congenital muscular dystrophies which include brain damage among their characteristics [42,43]. Dystroglycan alterations have also been related to different cancers, as RMS, prostate, colon or breast cancer [42,44,45]. Dystroglycan requires a correct function of LARGE1 gene and interacts with MEK and ERK components of the MAP kinase cascade, and thus, with TRK-pat [46]. PARK2 has also recently demonstrated a role in this pathway and it is one of the most frequently altered tumor suppressor genes in cancer [47]. Its depletion leads to PTEN inactivation and the co-occurrence of PARK2 and PTEN losses promotes tumorigenesis in vivo [47]. All these facts highlight the potential role of these genes in
Figure 4. Genes analyzed above the established threshold of somatic CNAs. The figures show the proportion of cases with alterations in each gene within each EGFR amplification group. (A) With respect to the losses, H-amp group exhibited five genes reaching the 40% of cases with losses, L-amp group presented eight genes and N-amp only one gene. (B) With respect to the gains, H-amp group presented two genes reaching the 40% of cases with losses, L-amp group presented six and N-amp two. Definitions: CNAs, copy number alterations; H-amp, glioblastomas with a high number of extra copies of EGFR; L-amp, glioblastomas with a low number of additional copies of EGFR; N-amp, glioblastomas without EGFR amplification.

Table 3. Associations between EGFR amplification and SCNAs in other genes (TCGA analysis).

| Gene   | EGFR diploid | EGFR amplification |
|--------|--------------|---------------------|
|        | Dip | Amp | Del | Dip | Amp | Del | p-value (1) | p-value (2) |
| CDK6   | 56  | 2   | 1   | 25  | 15  | 0   | 1.5E-05**** | 1.0E + 00   |
| CDKN1B | 47  | 1   | 9   | 0   | 1   | 26  | 3.4E-01     | 3.6E-01     |
| CDKN2A | 22  | 1   | 39  | 207 | 1   | 26  | 4.0E-01     | 1.8E-04***  |
| CDKN2B | 22  | 1   | 39  | 0   | 35  | 215 | 4.0E-01     | 1.8E-04***  |
| ESR1   | 46  | 0   | 14  | 0   | 0   | 72  | 1.0E + 00   | 4.2E-01     |
| LARGE1 | 41  | 0   | 19  | 179 | 0   | 69  | 1.0E + 00   | 6.3E-01     |
| MYBPC3 | 46  | 0   | 17  | 216 | 0   | 26  | 1.0E + 00   | 2.0E-03**   |
| PAH    | 50  | 1   | 11  | 222 | 0   | 14  | 1.9E-01     | 7.2E-03**   |
| PARK2  | 45  | 4   | 15  | 155 | 7   | 91  | 2.9E-01     | 9.6E-02     |

(1) Statistical significance for the comparison between the ratio of amplifications for the target gene in EGFR amplified samples vs EGFR diploid samples. ****p < 0.0001.
(2) Statistical significance for the comparison between the ratio of deletions for the target gene in EGFR amplified samples vs EGFR diploid samples. **p < 0.01, ***p < 0.001.
GBM biopathology and particularly in EGFR-amplified tumors. In addition, PARK2 is located on 6q, near to MAP3K4, reinforcing the influence of MAP kinases in GBM biopathology. As we showed in a previous work, MAP kinase cascade is intensely activated through ERK1/2 in GBM that display EGFR amplification [48]. Conversely, QKI is another gene in that region which acts as a target of miR200c. Recently another work demonstrated that miR200c is under expressed on EGFR region which acts as a target of miR200. Recently another work demonstrated that miR200c is under expressed on EGFR-amplified tumors [49]. This fact would mean that in 2/3 of our samples QKI could not develop its role. In line with these considerations of chromosomal location more than concrete gene, it is to mention that CCND2 is a cyclin frequently amplified in gliomas and in about 3% of GBM. Although we did not studied it, near, in 12p13 we can find CDKN1B, which showed similar frequency of CNA in our series. Thus, both 6q and 12p segments seem to contain genes whose knowledge is necessary to better understand GBM aggressiveness. Interestingly, tumor mutational burden resulted more predictive than any individual alteration: high values of both losses and/or gains statistically correlated to shortened survival in GBM patients. This fact agrees with recent published work on gliomas IDH-mutant and other neoplasia as lung or breast cancer and supports the potential usefulness of immunotherapy for GBM patients [41,53].

The genetic landscape of GBMs was widely varied in our series, in line with previous studies that demonstrated the coexistence of multiple clones in a manner that resemble admixtures of molecular distinct tumors rather than a uniform disease [41,53]. Our results showed also common genetic alterations that are independent to EGFR status. This fact could explain the varied intrinsic biological strategies related to tumor progression that make such complicated to achieve therapeutic improvements; however, our data also reflected the presence of specific differential cell clones in H-amp GBMs or in N-amp GBMs. Interestingly, L-amp GBM genetics partially overlapped with that of H-amp and L-amp groups. These findings imply that the therapeutic actions on these three groups would not cause the same clinical results and remarks the importance of establishing genetic GBM subgroups based on EGFR amplification levels.

The deregulation of molecular signaling pathways has impact on cell proliferation (Figure 5) and it is associated with a bad prognosis in GBM [17,54]. The proportion of patients in this work with alterations in the RTK-pat agrees with previous studies that indicate ranges between 50—88% [40,55]. Interestingly, in the H-amp group we found a number of PTEN alterations that is two-fold compared with L-amp. It is worth mentioning that PTEN is currently subject of therapeutic interventions in GBM [38]. Related to P53-pat, TP53 and MDM2 showed similar alterations to previous descriptions although the pathway global affection was lower than the previously published data [1,17,40,55]. This fact may be explained because this pathway could be more EGFR-independent than the others; it also suggests that N-amp GBMs could genetically resemble secondary GB, in which p53 alterations are more frequent [1]. Regarding RB-pat, the frequency of changes is concordant with the published frequencies too [17,40,55]. Our results show that the genetic burden affecting this pathway increases progressively from N-amp to H-amp. Interestingly, all the cases with simultaneous alteration of CDKN2A/B and CDK6 belonged to H-amp group. Globally, our series showed that, p53-pat and RB-pat deregulation in GBM happens more frequently because of MDM2 and CDKN2A changes respectively, than because of TP53 and RB1 alterations, in line with previous approaches [1,13,14].

In GBM, near 80% of the cases show deregulation of more than one pathway simultaneously [56], similarly to our data. The simultaneous alteration of genes from the three pathways was higher in H-amp GBMs, and interestingly, it was statistically associated to a shortened survival, which suggests, in an indirect manner, a poorer outcome when EGFR is highly amplified although larger series need to confirm these findings.

Figure 5. Relevant signaling pathways deregulated in glioblastoma: receptor tyrosine kinase/PI3K/AKT/mTOR pathway is shown in blue, CDKN2A/CDK4/6/retinoblastoma is shown in green and p53/MDM2/p14ARF signaling pathway is shown in red. Targets for LARGE1 and PARK2 novel alterations are indicated. Modified from Ohgaki et al. [15].

Conclusions

The accumulation of genomic alterations in the studied GBMs remarks the high genomic instability of this tumor. It has been shown an interesting relationship between high CNAs in GBM and patients survival, more prominent in EGFR H-amp and L-amp tumors, which may indicate the convenience of classifying GBMs depending on EGFR status in order to better understand the behavior of this highly heterogeneous entity. The
association of high CNAs with patients survival underlines the influence of the tumor mutational burden in the progression of GBM. The current study highlights the prognostic strength of the combination of MLPA/FISH, which are technologies completely implemented in most of the hospitals. For sure, further studies are necessary to follow improving the FISH, which are technologies completely implemented in most of the hospitals. The current association of high CNAs with patients survival underlines the influence of the tumor mutational burden in the progression of GBM.

Frequent amplification of the epidermal growth factor receptor (EGFR) has been identified as crucial for glioblastoma proliferation, metastasis, and resistance to therapy. EGFR amplification and somatic CNAs in glioblastoma L. Muoz-Hidalgo et al. Neoplasia Vol. 22, No.1, 2020

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.neo.2019.09.001.

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