

1 **Tumor evolution of glioma intrinsic gene expression subtype associates with**  
2 **immunological changes in the microenvironment**

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29 mesenchymal transition, gene expression profiling, tumor microenvironment,  
30 macrophages/microglia, immune cells.

31 **ABSTRACT**

32 We leveraged IDH wild type glioblastomas (GBM) and derivative neurosphere models to  
33 define the tumor-intrinsic mRNA transcription phenotype. We found that intratumoral  
34 heterogeneity is reflected in the transcriptome and associated with increased tumor  
35 microenvironment presence. We performed in silico cell sorting to demonstrate that M2  
36 macrophages/microglia are the most frequent type of immune cells in the glioma  
37 microenvironment, followed by CD4+ T lymphocytes. Longitudinal transcriptome  
38 analysis of 124 pairs of primary and recurrent glioma pairs showed expression subtype  
39 is retained in 53% of cases with no proneural to mesenchymal transition being apparent.  
40 Inference of the tumor microenvironment through gene signatures revealed a decrease  
41 in invading monocytes but a subtype dependent increase in M2 macrophages/microglia  
42 cells after disease recurrence. Our longitudinal expression dataset can be accessed at  
43 <http://ackbar.cnio.es:3838/RecuR/>. Our study provides a comprehensive transcriptional  
44 and cellular landscape of IDH wild type GBM during treatment modulated tumor  
45 evolution.

## 46 INTRODUCTION

47 The intrinsic capacity of glioblastoma (GBM) tumor cells to infiltrate normal brain  
48 impedes surgical eradication and predictably results in high rates of early recurrence.  
49 To better understand determinants of GBM tumor evolution and treatment resistance,  
50 The Cancer Genome Atlas Consortium (TCGA) performed high dimensional profiling  
51 and molecular classification of nearly 600 GBM tumors<sup>1-5</sup>. In addition to revealing  
52 common mutations in genes such as *TP53*, *EGFR*, *IDH1*, and *PTEN*, as well as the  
53 frequent and concurrent presence of abnormalities in the p53, RB and receptor tyrosine  
54 kinase pathways. Unsupervised transcriptome analysis identified four clusters, referred  
55 to as classical, mesenchymal, neural and proneural, that were tightly associated with  
56 genomic abnormalities<sup>6</sup>. The proneural and the mesenchymal expression subtypes  
57 have been most consistently described in literature with proneural relating to a more  
58 favorable outcome and mesenchymal to unfavorable survival<sup>7-9</sup>, but these findings were  
59 affected by the relatively favorable outcome of IDH-mutant glioblastoma which are  
60 consistently classified as proneural<sup>4,6</sup>. Proneural to mesenchymal switching upon  
61 disease recurrence has been described as a source for treatment resistance in GBM  
62 relapse<sup>7,10-12</sup>, but the relevance of this phenomenon in glioma progress remains  
63 ambiguous.

64 GBM tumor cells along with the tumor microenvironment create a complex milieu  
65 that ultimately promotes tumor cell plasticity and disease progression<sup>13</sup>. The presence  
66 of tumor-associated stroma results in a mesenchymal tumor gene signature and poor  
67 prognosis in colon cancers<sup>14</sup>. Furthermore, the association between a mesenchymal  
68 gene expression signature and reduced tumor purity has been identified as a common

69 theme across cancer <sup>15,16</sup>. Tumor-associated macrophages/microglia in GBM have  
70 been proposed as regulators of proneural-to-mesenchymal transition through NF-κB  
71 activation <sup>10</sup> and may provide growth factor mediated proliferative signals, which could  
72 be therapeutically targeted <sup>17-19</sup>.

73 Here, we explored the properties of the microenvironment in different GBM gene  
74 expression subtypes and characterized the transition between molecular subtypes  
75 before and after therapeutic intervention. In doing so, we improved the robustness of  
76 gene expression subtype classification through revised gene signatures and proposed  
77 analytical methodology. Our results suggested that the tumor microenvironment  
78 interferes with expression based classification of GBM, both at the primary disease  
79 stage as well as at disease recurrence, and suggest a role for the  
80 macrophage/microglia in treatment response.

81 **RESULTS**

82 ***Harnessing glioma sphere-forming cells identifies GBM specific intertumoral***  
83 ***transcriptional heterogeneity***

84 We set out to elucidate the tumor-intrinsic and tumor microenvironment independent  
85 transcriptional heterogeneity of GBMs. We performed a pairwise gene expression  
86 comparison of independent set of GBMs and the derivative glioma sphere-forming cells  
87 (GSCs) (n = 37)<sup>20</sup>. In total, 5,334 genes were found to be significantly higher  
88 expressed in parental GBMs relative to derived GSCs that could be attributed by the  
89 tumor associated GBM microenvironment (**Figure 1A**). To focus the analysis on the  
90 tumor-intrinsic transcriptome, these genes were filtered from further analysis. GBMs  
91 with IDH mutations have distinct biological properties and favorable clinical outcomes  
92 compared to IDH wild-type GBMs<sup>1,4,5,21</sup>. Using the filtered gene set, we performed  
93 consensus non-negative matrix factorization clustering to identify three distinct  
94 subgroups amongst 369 IDH wild type GBMs (**Figure 1B; Figure 1C**). When comparing  
95 the clustering result with the previously defined proneural (PN), neural (NE), classical  
96 (CL) and mesenchymal (MES) classification<sup>1,3</sup>, three subgroups were strongly enriched  
97 for CL, MES and PN GBMs, respectively (**Figure S1**). Consequently, we labeled the  
98 groups as CL, MES and PN. None of the three subgroups was enriched for the NE  
99 class, suggesting its neural phenotype is non-tumor specific. The NE group has  
100 previously been related to the tumor margin where normal neural tissue is more likely to  
101 be present<sup>22</sup> and such contamination might explain why the neural subtype was the  
102 only subtype to lack characteristic gene abnormalities<sup>1,23</sup>. To be able to classify  
103 external GBM samples, we implemented a single sample gene set enrichment analysis

104 (ssGSEA) based equivalent distribution resampling classification strategy using 70-gene  
105 signatures for each subgroup (**Table S1**)(**Figure 1D**), to assign each sample three  
106 empirical classification p-values by which we determined the significantly activated  
107 subtype(s) in the samples. We prepared an R-library to facilitate others to evaluate our  
108 approach (**Supplementary File**). Using this method we found that the stability of cluster  
109 assignments of 144 TCGA GBM samples profiled using both RNA sequencing and  
110 Affymetrix U133A microarrays was 95% concordance (**Figure S2, Table S2**). This was  
111 an improvement over the 77% subtype concordance determined using previously  
112 reported methods <sup>3</sup>. We evaluated the distribution of somatic variants across all the  
113 three molecular subtypes (**Figure 1E**)(**Figure S3**) and confirmed the strong  
114 associations between subtypes and genomic abnormalities in previously reported driver  
115 genes <sup>1,3</sup>.

116

### 117 ***Multi-activation of subtype signatures associated with intratumoral heterogeneity***

118 We observed that 34/369 (9.2%) samples showed significant enrichment of multiple  
119 ssGSEA scores (empirical classification p-value<0.05), suggesting these cases activate  
120 more than one transcriptional subtype (**Figure 2A**). To quantify this phenomenon, a  
121 score ranging from 0 to 1 was defined to quantitatively evaluate the simplicity of subtype  
122 activation based on order statistics of ssGSEA score. Samples with high simplicity  
123 scores activated a single subtype and those with lowest simplicity scores activated  
124 multiple subtypes. All multi-subtype TCGA samples showed simplicity scores of less  
125 than 0.1 (**Figure 2A**). To determine whether transcriptional heterogeneity associated  
126 with genomic intratumoral heterogeneity, we correlated simplicity scores, total mutation

127 rates and subclonal mutation rates. Included in the analysis were 224 TCGA GBMs with  
128 available whole exome sequencing data <sup>24</sup> and ABSOLUTE <sup>25</sup> determined high tumor  
129 purity (> 0.8) to equalize the mutation detection sensitivity <sup>26</sup>. Although not significant  
130 (Wilcoxon rank test p-value=0.143), the total mutation rate was less in the bottom 30%  
131 with lowest simplicity scores versus the top 30% samples with highest simplicity scores.  
132 The subclonal mutation rate was significantly higher (p-value=0.024) in samples with  
133 lowest simplicity scores (**Figure 2B; Table S3**), suggesting that increased intratumoral  
134 heterogeneity associates with increased transcriptional heterogeneity.

135 We compared outcomes amongst the three transcriptional groups and observed  
136 no significant differences (**Figure S4**). However, when restricting the analysis to  
137 samples with high simplicity scores, a clear trend of MES showing worst survival and  
138 PN the most favorable outcome became visible. For example, Kaplan-Meier analysis of  
139 88 samples with simplicity scores >0.99 showed a median survival of 11.4, 14.7 and  
140 16.7 months were detected in MES, CL and PN, respectively, which was significantly  
141 different (log rank test, p=0.048)(**Figure 2C**)(**Figure S4**)(**Table S4**). Higher simplicity  
142 scores correlated with relative favorable outcome within the PN set, non-significant in  
143 the CL subtype, and correlated with relatively unfavorable survival in the MES class  
144 (**Figure S5**).

145 Single GBM cell RNA sequencing recently suggested that GBMs are comprised  
146 of a mixture of tumor cells with variable GBM subtype footprints <sup>18</sup>. We used this data to  
147 classify 502 single GBM cells in addition to the bulk tumor derived from five primary  
148 glioblastomas (**Table S5**). All bulk tumor samples showed simplicity scores less than  
149 0.05 suggesting high transcriptional heterogeneity compared to 45 of 369 TCGA GBM

150 samples with simplicity scores below 0.05 (**Figure 2D**). In four of five cases (MGH26,  
151 MGH28, MGH29 and MGH30), the bulk tumor samples were classified in the same  
152 primary subtype as the majority of their single cells (**Figure 2D**). Our analysis suggests  
153 that the heterogeneity observed at the single cell level is captured in the expression  
154 profile of the bulk tumor, and that the five GBM samples studied at the single cell level  
155 represented samples with relatively high transcriptional heterogeneity.

156

### 157 ***Transcriptional subtypes differentially activate the immune microenvironment***

158 Despite restricting the cluster analysis to genes exclusively expressed by GBM cells, we  
159 found that tumor purity predictions based on ABSOLUTE were significantly reduced in  
160 GBM classified as MES (Student T-test p-value < 10e-14; **Figure 3A**). This was  
161 corroborated by gene expression based predictions of tumor purity using the  
162 ESTIMATE method (Student T-test p-value < 10e-32; **Figure 3B**)<sup>15</sup>. The ESTIMATE  
163 method has been optimized to quantify tumor-associated fibroblasts and immune cells  
164 <sup>15</sup> and the convergence of a decreased ABSOLUTE and decreased ESTIMATE tumor  
165 purity confirmed previous suggestions on the increased presence of microglial and  
166 neuroglial cells in mesenchymal GBM <sup>12,27,28</sup>. The mean simplicity score of samples  
167 classified as MES was 0.53 which was significantly lower than in PN (Wilcoxon rank test  
168 p-value<0.019) and CL subtypes (Wilcoxon rank test p-value<0.0001), confirming  
169 increased transcriptional heterogeneity. In order to identify genomic determinants of  
170 macrophage/microglia chemoattraction, we compared genomic alterations between  
171 mesenchymal class samples with high (n=51) and low (n=51) ABSOLUTE based tumor  
172 purity. GBM carrying hemizygous loss of *NF1* or somatic mutations in *NF1* showed

173 reduced tumor purity compared to GBM with wild type *NF1* (Wilcoxon rank test p-  
174 value=0.0007) and this association was similarly detected when limiting the analysis to  
175 MES samples (Wilcoxon rank test p-value=0.017)(**Figure S6**). Formation of dermal  
176 neurofibromas in the context of *Nf1* loss of heterozygosity has been reported to be  
177 context and microenvironment dependent <sup>29</sup>. Functional studies may clarify whether  
178 *NF1* deficient GBM are able to recruit cells that provide them with a proliferative  
179 advantage, or whether *NF1* loss provides that proliferative advantage in a specific  
180 tumor-associated microenvironment context.

181 To determine the cellular components of the tumor microenvironment across  
182 different transcriptional subtypes, we used the CIBERSORT in silico cytometry <sup>30</sup>  
183 method to evaluate absolute immune cell fractions. We evaluated 22 different immune  
184 cell types in 69 PN, 137 CL and 96 MES samples, after filtering samples with  
185 classification simplicity scores less than 0.1 (**Table S6**). Microglia are the resident  
186 macrophages in the central nervous system. Peripheral blood monocytes also give rise  
187 to tumor associated macrophages. These innate immune cells can be broadly classified  
188 as the proinflammatory M1 type and the alternative tumor promoting M2 type<sup>31</sup>. The M2  
189 macrophage gene signature showed a greater association with the MES subtype  
190 (13.4%) relative to the PN (4.6%) and CL (6.0%)(**Figure 3C**), consistent with previous  
191 analysis of the TCGA database<sup>28,32</sup>. In addition to the M2 macrophage gene signature,  
192 there was also a significantly higher fraction of MES samples that expressed M1  
193 macrophage (Student T-test p-value 3.20E-5) and neutrophil (Student T-test p-value  
194 1.30E-9) gene signatures. In contrast, the activated natural killer T-cell gene signature  
195 (Student T-test p-value 4.91E-2) was significantly reduced in the MES subtype and

196 resting memory CD4+ T cells (Student T-test p-value 5.40E-7) were less frequently  
197 expressed in the PN subtype.

198 To confirm the association of macrophages/microglia with the MES GBM subtype,  
199 we assessed protein expression levels of the ITGAM (alternatively known as CD11B)  
200 and IBA1 (also known as AIF1) macrophage/microglial markers in a set of 18 GBM for  
201 which we also characterized the expression subtype (**Figure 3D**) as well as through  
202 immunohistochemistry (**Figure S7**). We confirmed the microenvironment as the main  
203 source for *ITGAM/IBA1* transcription by comparing transcriptional levels in 37 GBM-  
204 neurosphere pairs used for gene filtering, which showed that neurospheres do not  
205 express *ITGAM/IBA1* (**Figure 3E**). The association of the MES GBM subtype with  
206 increased level of M2 microglia/macrophages may suggest that in particular MES GBM  
207 are candidates for therapies directed against tumor-associated macrophages<sup>17</sup>.  
208 Activated dendritic cell signatures (Student T-test p-value 7.36E-3)(**Figure 3C**) were  
209 significantly higher in the CL subtype, suggesting this subtype may benefit from  
210 dendritic cell vaccines<sup>33</sup>. Dendritic cells may require an activated phenotype in order to  
211 direct the immune system. A previous study suggested that MES GBM patients treated  
212 with dendritic cells were more likely to benefit<sup>34</sup>.

213

### 214 ***Phenotypic plasticity upon GBM recurrence***

215 Glioblastoma has long been hypothesized to progress along a proneural to  
216 mesenchymal axis<sup>7</sup>. To determine the relevance of this transition process in IDH  
217 wildtype glioma evolution, we performed a longitudinal analysis of the subtype  
218 classification and tumor-associated microenvironment in sample pairs obtained at

219 diagnosis and first disease recurrence from 124 glioma patients. The cohort included 96  
220 initial GBM and first recurrence, eight pairs of primary low grade glioma and matching  
221 secondary GBM, and 20 pairs of primary and recurrent low grade glioma. Gene  
222 expression profiles of 78 tumor pairs were analyzed through transcriptome sequencing,  
223 and remaining pairs were generated using Affymetrix (n = 31) and Illumina (n = 15)  
224 microarray, respectively. To facilitate exploration of this dataset we have made it  
225 available through the GlioVis portal <http://ackbar.cnio.es:3838/RecuR/>. We used a gene  
226 expression signature<sup>35</sup> to determine that 33 of 124 cases were IDH-mutant/GCIMP at  
227 presentation and recurrence (**Table S7**). We used the renewed gene signatures and  
228 classification method to determine molecular subtype of the 91 pairs of IDH wild type  
229 cases and found that expression class remained consistent after disease recurrence for  
230 48 of 91 IDH-wildtype cases (52%)(**Figure 4A**). The MES subtype was most stable  
231 (64%) while the CL (47%) and PN (43%) phenotypes were less frequently retained.  
232 Nine, sixteen and eighteen post-treatment tumors switched subtypes to become CL,  
233 MES and PN at disease recurrence, respectively, indicating that PN and MES increased  
234 in frequency after recurrence while the CL subtype was least frequently found (**Figure**  
235 **4A**). The CL expression class was previously found to be most sensitive to intensive  
236 therapy and it is possible that therapy provides a competitive advantage for non-CL  
237 cells, which could explain the reduced post-treatment incidence<sup>3</sup>. Our results did not  
238 identify enrichment for proneural to mesenchymal transitions.

239 We observed a significant difference in transcriptional simplicity between primary  
240 GBM retaining their expression class, versus those that switched to a different  
241 phenotype (**Figure 4B**). GBMs with a primary tumor simplicity score greater than 0.5,

242 indicated lower transcriptional heterogeneity, were classified as the same subtype in 31  
243 of 48 (64.5%) cases, compared to 15 of 41 (36.6%) cases with primary tumor simplicity  
244 scores less than 0.5 (Fisher exact test p-value=0.01).

245

### 246 ***Microenvironment transitions upon GBM recurrence***

247 Debulking surgery, radiotherapy and chemotherapy provide therapeutic barriers but  
248 nonetheless induce tumor evolution, including influences on the tumor  
249 microenvironment. We explored this possibility by comparing the tumor associated  
250 microenvironment in primary and recurrent GBMs using CIBERSORT (**Table S8**)<sup>30</sup>. A  
251 comparison between 91 primary and recurrent IDH-wild type tumors revealed a  
252 decrease in monocyte gene signature expression at recurrence, suggesting relative  
253 depletion of circulation derived monocytes (**Figure 5A**). Next, we dissected  
254 microenvironment fluctuations between diagnosis and recurrent tumors across different  
255 subtype combinations. Primary non-MES (CL or PN) tumors showed relatively high  
256 tumor purity and consequently, recurrent tumors classified as non-MES demonstrated a  
257 relatively global decrease of immune cells while cases transitioning to MES at  
258 recurrence represented a trend towards increased immune cell fractions (**Figure 5B**).  
259 Gene signatures of immunosuppressive regulatory T cells showed an increase in gene  
260 expression at recurrence across several primary-recurrence subtype combinations  
261 although the inferred cellular fractions are relatively small (**Figure 5B**). In contrast to the  
262 trend of monocyte depletion, the imputed M2 macrophage frequency was significantly  
263 higher at recurrence in cases transitioning to MES (**Figure 5C**). This observation  
264 converges with the higher fraction of M2 macrophages in primary MES GBM relative to

265 primary non-MES GBM. M1 macrophages and neutrophils also correlated with primary  
266 MES GBM, but these associations were not confirmed for recurrent GBM. We validated  
267 the increase in macrophages using immunostaining of IBA1 expression in two primary-  
268 recurrent GBM pairs which were classified as CL to MES (**Figure 5D**). IBA1  
269 immunoexpression was restricted to macrophages/microglia, cells exhibiting either  
270 globular or filamentous/spidery morphology, with no expression in glioma tumor cells  
271 (**Figure 5D**). These findings further solidify the association between MES GBM and  
272 macrophage/microglia and extend this mutual relationship to disease recurrence. MES  
273 tumors at recurrence compared to primary MES tumors showed an increase in  
274 transcriptional activity associated with non-polarized M0 macrophages, which has been  
275 previously described <sup>28</sup>, but also dendritic cells which is potentially motivated by the  
276 increased levels of neoantigens at disease recurrence <sup>24</sup>. In contrast, primary PN GBM  
277 were found to contain significantly higher fractions of five immune cell categories  
278 compared to recurrent PN GBM, indicating a relative absence of immune infiltration in  
279 PN GBM upon recurrence.

280 We evaluated the effect of transcriptional class on patient survival. The analysis  
281 was restricted to 50 cases for whom annotation on overall survival (OS) time and time to  
282 disease progression (PFS) were available and with high simplicity scores, indicating low  
283 transcriptional heterogeneity. We confirmed the worse prognosis for patients whose  
284 primary tumor was classified as MES on overall survival (logrank test  $p=0.029$  with  
285  $HR=1.97$ )(**Figure 6A; Figure 6B**). This pattern was retained in patients whose  
286 secondary glioma was classified as MES (logrank test  $p=0.09$  with  $HR=1.71$ )(**Figure 6C;**  
287 **Figure 6D**). Consequently, cases for whom both primary and recurrent tumor were

288 classified as MES subtype showed the least favorable outcome, suggesting an additive  
289 effect of transcriptional class at different time points (**Figure 6E, Figure 6F**)(**Figure S8**).

## 290 **DISCUSSION**

291 Transcriptome profiling of tumor samples is a commonly used modality for interrogating  
292 pathway functionality and phenotype based patient classification. The transcriptional  
293 footprint left by the tumor microenvironment, which may constitute 10-80% of cells in a  
294 tumor biopsy<sup>15</sup>, can obscure the true activity of the signaling network<sup>14,36</sup>. Here, we  
295 employed in silico methods to integrate mRNA expression profiles from glioma samples  
296 and glioma cell culture models to provide insights into glioma-intrinsic pathway activities  
297 and classification, and to deconvolute the glioma associated stroma into its  
298 immunological cellular components.

299 GBM expression subtype classification has emerged as an important concept to  
300 better understand the biology of this devastating disease<sup>8,37,38</sup>. Robust classification of  
301 new GBM tumors is therefore critical to ensure consistency in reporting between  
302 different studies. The transcriptional glioma subtypes we discovered using tumor-  
303 intrinsic gene expression values strongly overlapped with the proneural, classical and  
304 mesenchymal subtypes but identified the neural class as normal neural lineage  
305 contamination. Our updated methods, released through a R-library, were found to be  
306 highly robust and provide the community with a standardized strategy for classification  
307 of gliomas.

308 Through re-classification of primary GBM samples from TCGA and despite using  
309 tumor-only transcripts, we observed that the mesenchymal GBM subtype associated  
310 with the presence of tumor-associated glial and microglial cells. Mesenchymal glioma  
311 cell differentiation status has been found to correlate with enrichment of  
312 macrophages/microglia<sup>10,39</sup>. Through in silico cell type identification we additionally

313 detected enrichment of various adaptive immunity cell types, including CD4+ T  
314 lymphocytes.

315 Longitudinal analysis of tumor samples is complicated by the lack of tissue  
316 collections including such pairs. Through aggregation of existing and novel datasets we  
317 compiled a cohort of 124 glioma pairs, including 91 pairs of IDH wild type tumors.  
318 Comparison of pairs of initial gliomas and first disease recurrence did not identify the  
319 trend of proneural GBM transitioning to a mesenchymal phenotype that has often been  
320 suspected <sup>7</sup>. Mesenchymal subtype at diagnosis and at disease recurrent correlated  
321 with relatively poor outcome. The recurrent IDH wild type GBM immuno environment  
322 showed fewer blood derived monocytes which may reflect lower penetration through the  
323 blood brain barrier. While the frequency of M2 macrophage/microglia was increased in  
324 recurrent mesenchymal GBM compared to primary non-mesenchymal GBM, the overall  
325 fraction of M2 macrophage/microglia remained stable. This possibly suggests that the  
326 majority of these cells are derived from resident CNS macrophages than through active  
327 recruitment from the circulation.

328 In summary, our study defines a new strategy to determine transcriptional  
329 subtype, and associated expression classes to the tumor-associated immuno-  
330 environment. Our findings may aid in the implementation of immunotherapy approaches  
331 <sup>40</sup> in a disease type with very limited treatment options. Collectively, our results have  
332 improved our understanding of determinants of GBM subtype classification, the critical  
333 impact of the tumor microenvironment, and provide new handles on the interpretation of  
334 transcriptional profiling of glioma.

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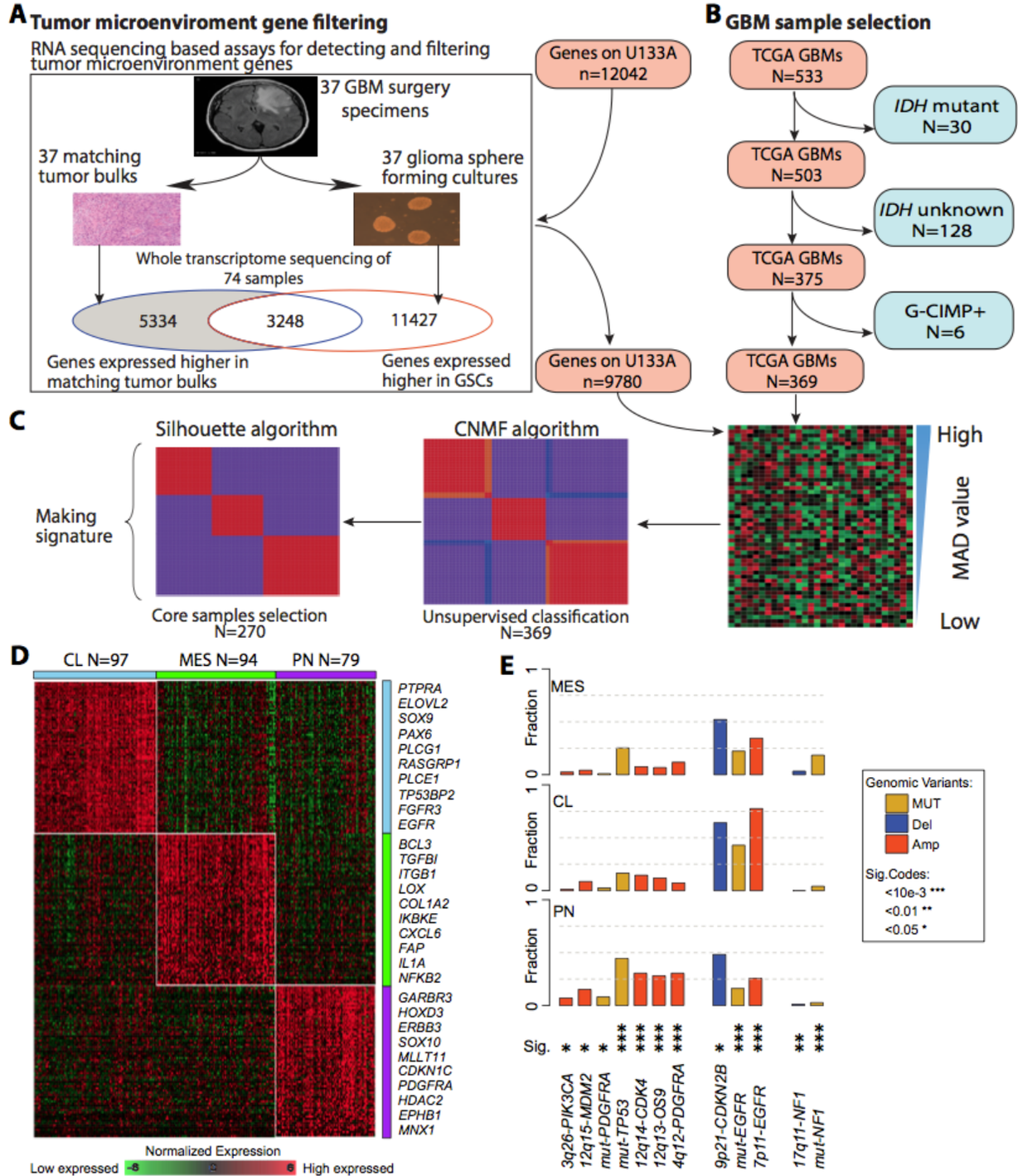
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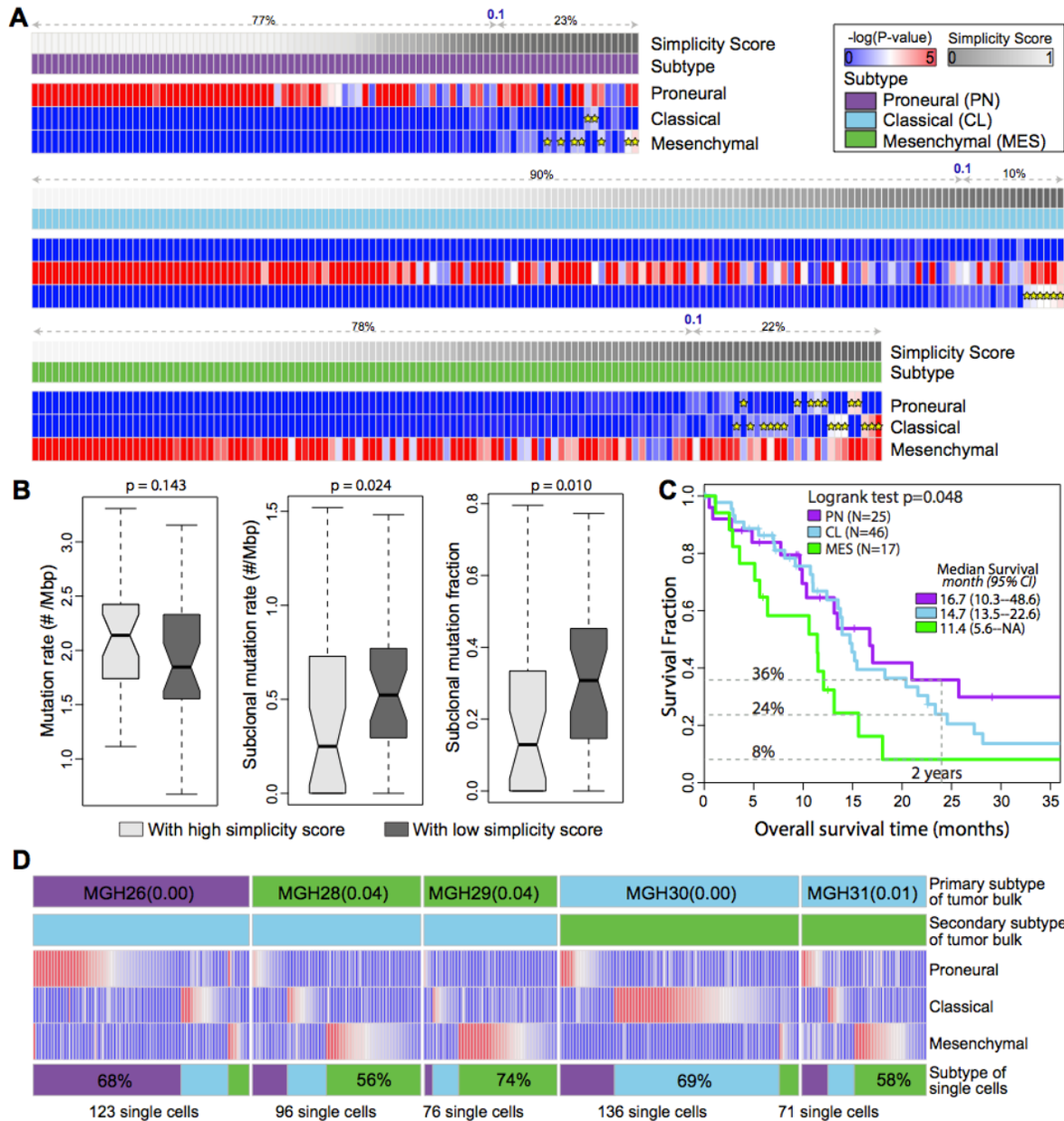
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444 **Figures and Legends**



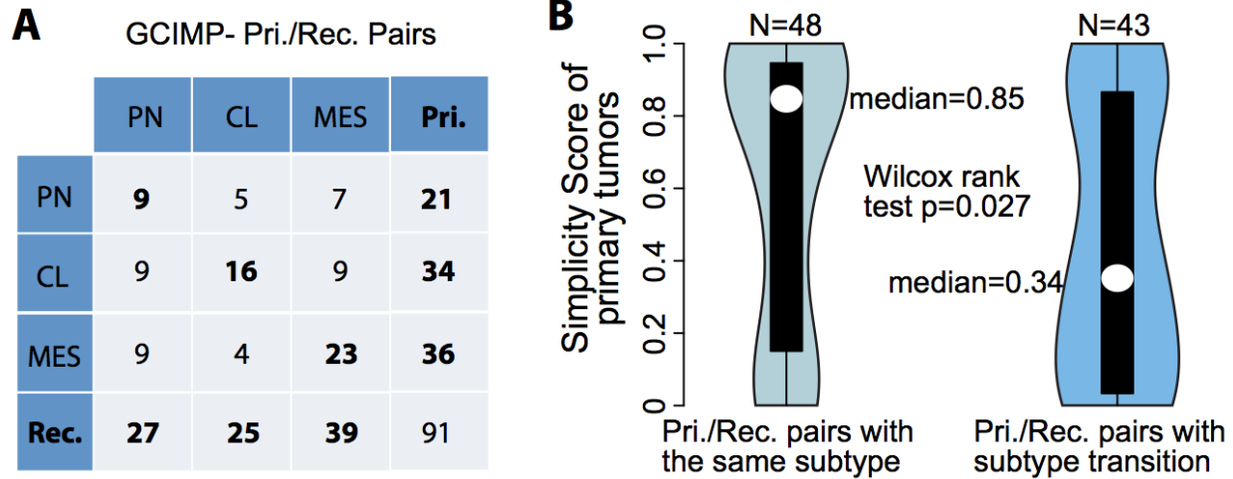
445

446 **Figure 1. Molecular classification of IDHwt GBMs.** (A) Filtering tumor associated  
 447 microenvironment genes. (B) Discarding IDH mutation related GBMs. (C) Overview of NMF  
 448 clustering. (D) Heatmap of 70-gene signatures by gene expression subtype were developed  
 449 based on 270 GBMs. Top ten genes are shown for each subtype. (E) Frequency of subtype  
 450 related somatic genomic alterations.



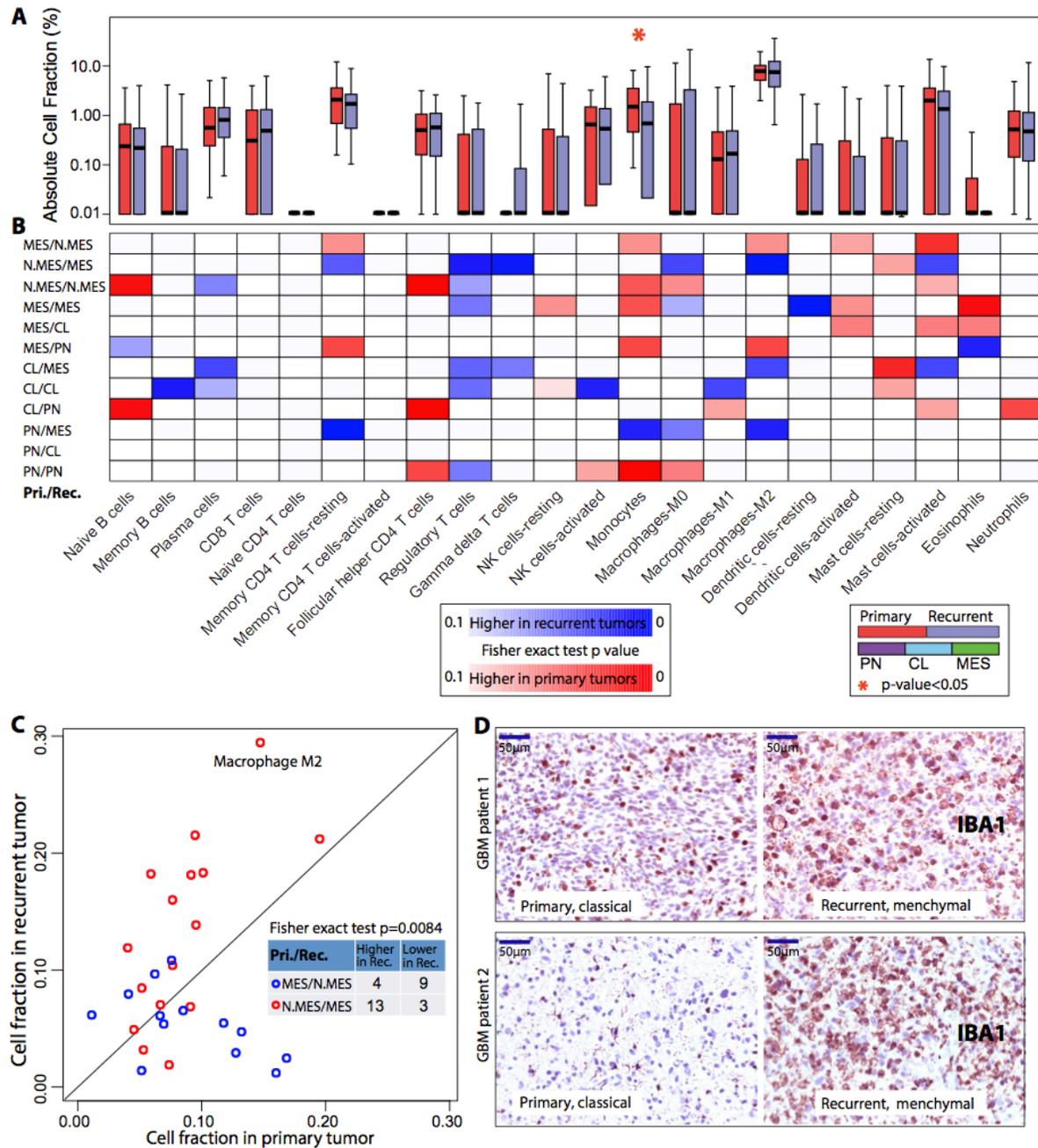
451  
 452 **Figure 2. Multi activation of transcriptional subtypes associated with intratumoral**  
 453 **heterogeneity.** (A) The expression profiles of 369 IDHwt GBMs were analyzed using Affymetrix  
 454 U133A. The empirical  $-\log(P\text{-value})$  of raw ssGSEA enrichment scores at each signature are  
 455 shown as heatmaps, with dark blue representing no activation and bright red as highly activated.  
 456 Yellow star indicates the secondary activated subtype (empirical  $p\text{-value} < 0.05$ ). For each panel,  
 457 the first row shows simplicity score, and the second row indicates transcriptional subtype. (B)  
 458 Comparison of mutation rate, subclonal mutation rate and subclonal mutation fraction between  
 459 IDHwt GBMs with high and low simplicity scores. P-values were calculated using Wilcoxon rank  
 460 test and shown at the top of each panel. (C) Kaplan-Meier survival curve by subtype. (D)  
 461 Transcriptome classification of five bulk tumor samples and 502 single GBM cells derived from  
 462 them. The top two row of each panel show the dominant and secondary subtype of the GBM  
 463 tumor bulk. The heatmap of each panel shows the empirical  $-\log(P\text{-value})$  of the ssGSEA  
 464 scores of the derived single GBM cells on each of the three subtype signatures. The bottom row  
 465 shows the subtype distribution of derived single GBM cells within the same GBM tumor of origin.



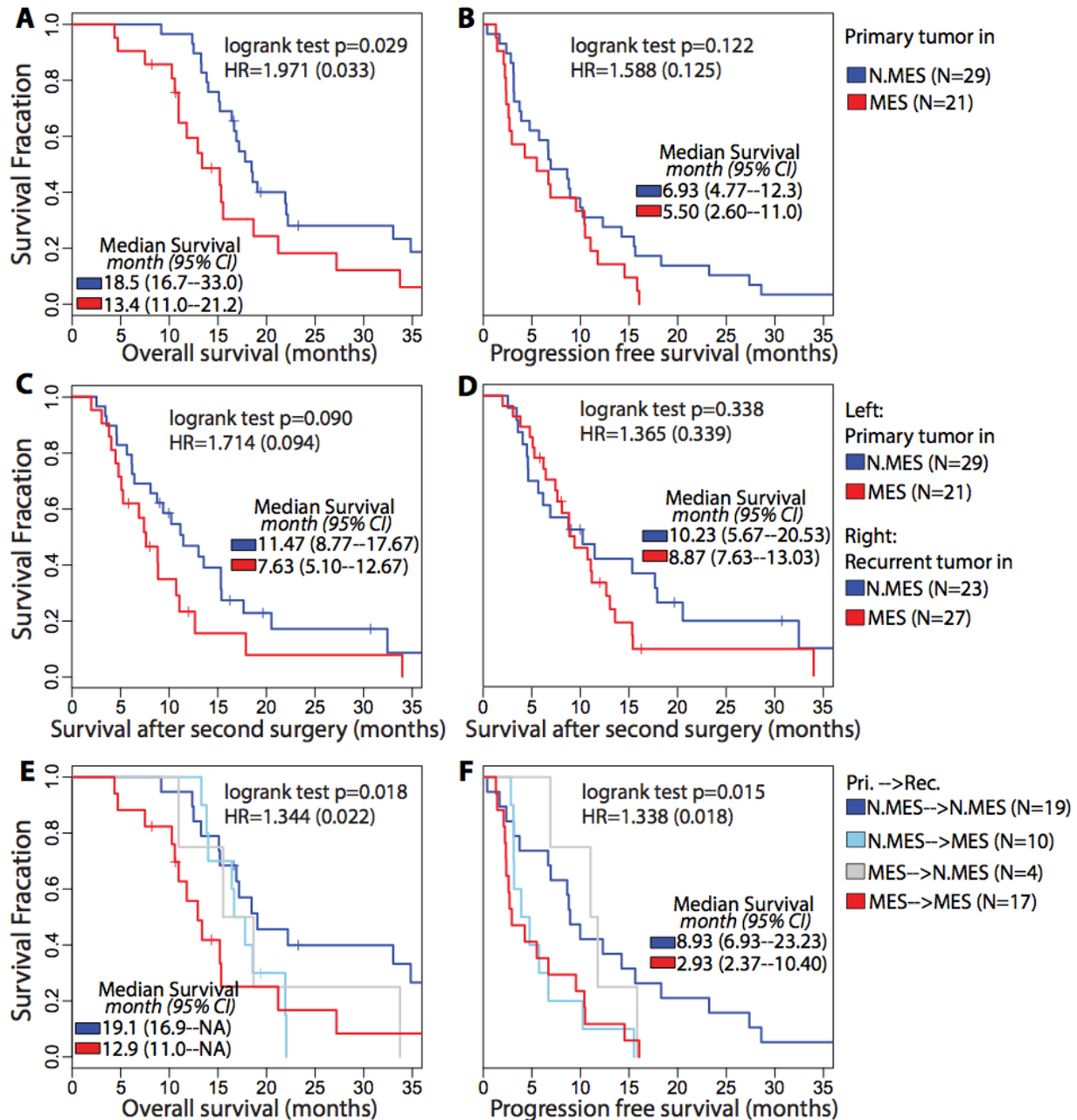


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**Figure 4. Comparison between transcriptional subtype of primary and paired recurrent tumors.** (A) Rows and columns of the cross table represents subtype distribution frequency of primary and paired recurrent tumors, respectively. (B) Violin plots show the distribution of simplicity scores of pairs with (left) and without (right) subtype transition.



485  
 486 **Figure 5. Microenvironment transition between primary and paired recurrent tumors. (A)**  
 487 Red and blue boxplots represent the immune cell fraction distribution of each immune cell type.  
 488 Immune cell fraction was calculated using CIBERSORT and adjusted using ESTIMATE purity  
 489 scores. Difference between cell fraction of primary and paired recurrent tumors was calculated  
 490 using Wilcoxon rank test. **(B)** The blue-to-red heatmap represents immune cell fraction changes  
 491 upon tumor recurrence per subtype transitions which were list on the left of the heatmap. **(C)**  
 492 Each dot represents a pair of primary and recurrent GBM with axes indicating M2 macrophage  
 493 cell fraction. **(D)** Immunohistochemical staining of the IBA1 microglia marker in two matched  
 494 pairs of primary and recurrent GBM. Cellular morphology in combination with the distribution of  
 495 the staining suggested exclusive microglial expression of monocyte lineage markers IBA1 in  
 496 GBM tumors.



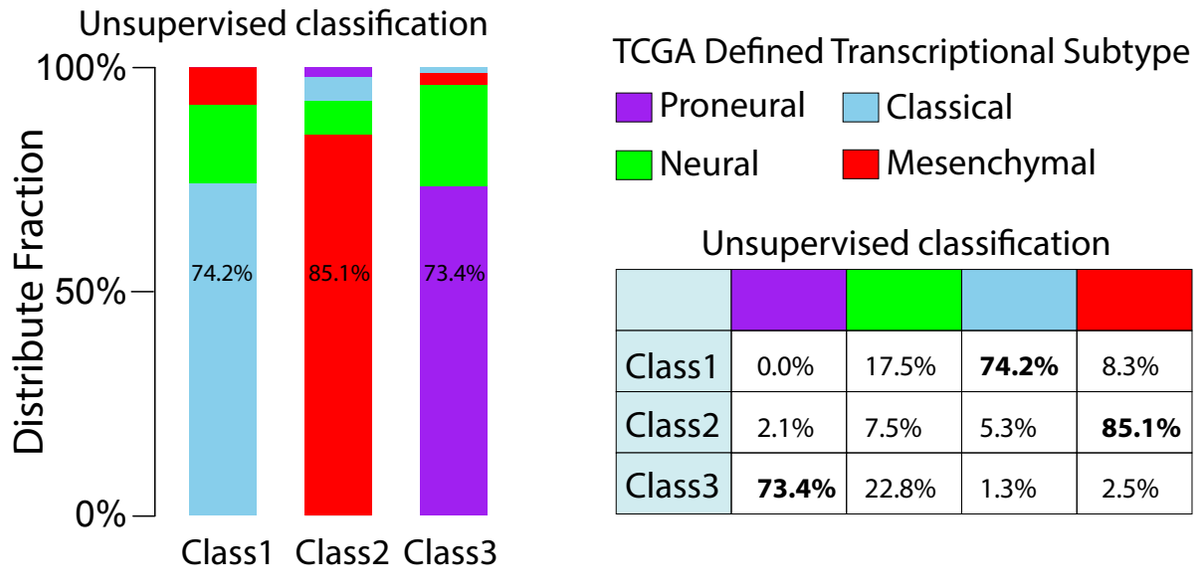
497  
 498 **Figure 6. Survival analysis of paired IDH wild type GBM.** (A, B) OS and PFS analyses  
 499 between samples with different primary subtype. (C) Difference of survival time after secondary  
 500 surgery between patients with non-MES and MES in primary tumors. (D) Survival analysis of  
 501 time after secondary surgery between patients with non-MES and MES in recurrent tumors. (E,  
 502 F) OS and PFS analyses between samples with difference recurrent subtype.

503

## Supplementary Information

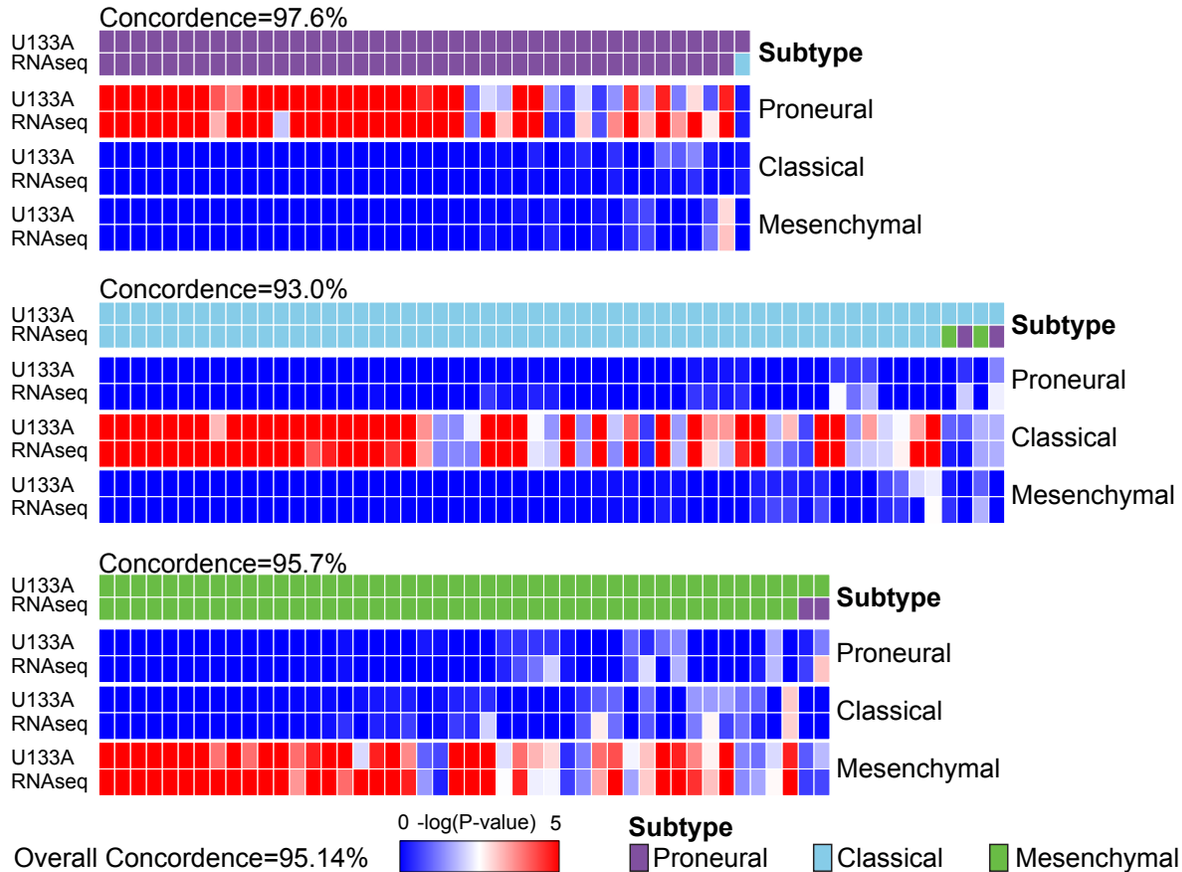
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### 505 Supplementary Figures and Legends



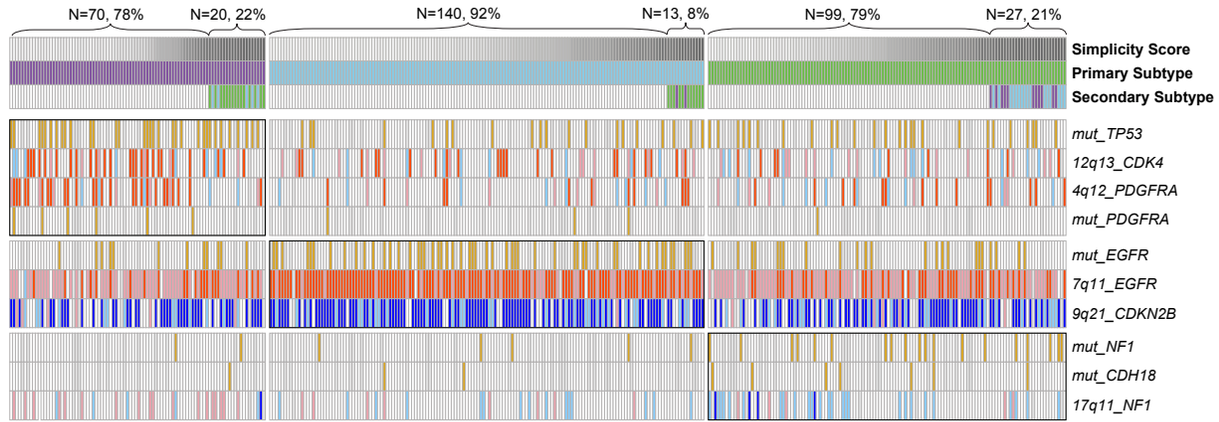
506

507 **Figure S1.** Comparison between GCIMP- GBM specific classification and TCGA defined GBM  
508 subtypes.



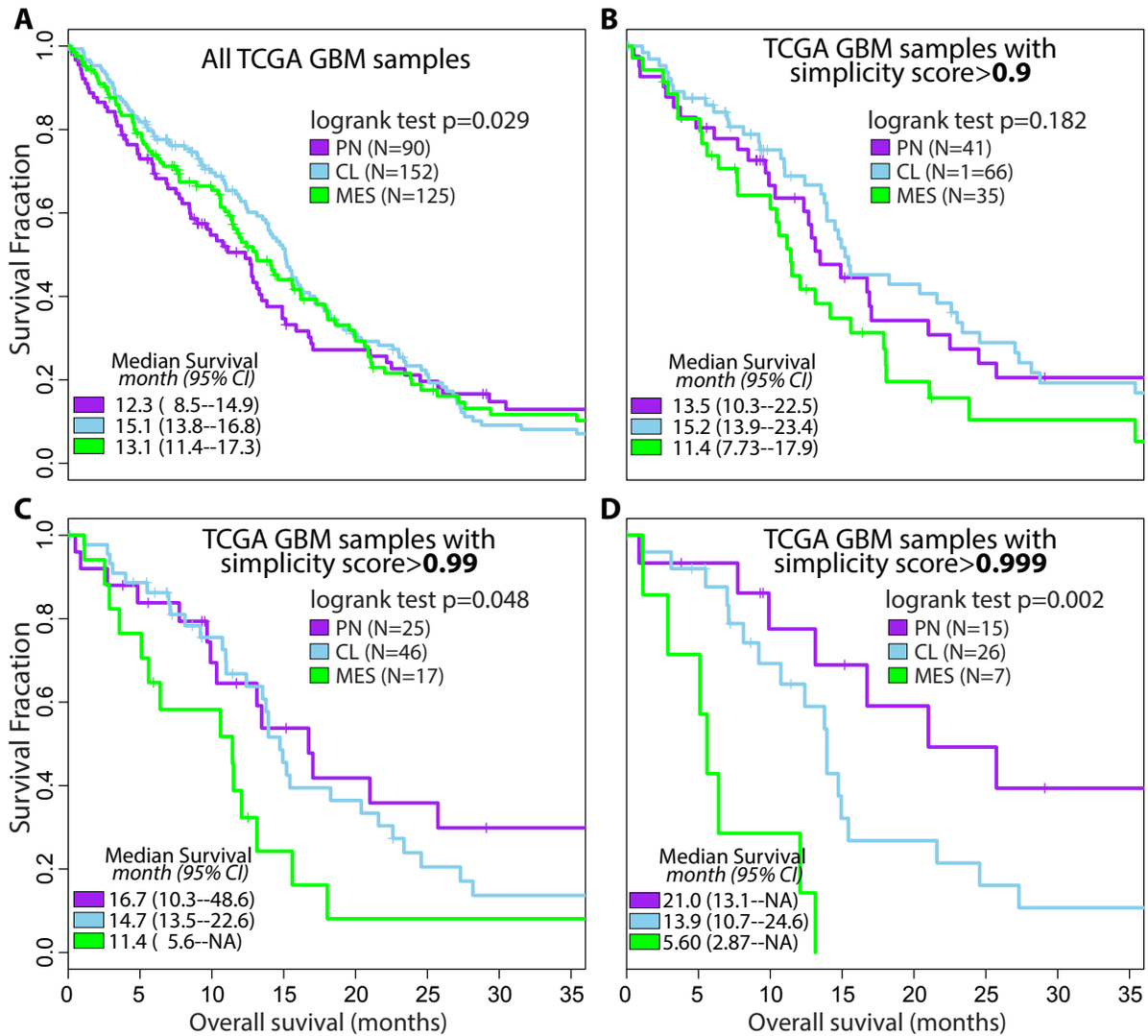
509

510 **Figure S2. Concordance of transcriptional classification of GBMs cross multiple**  
511 **platforms.** Through TCGA, the expression profiles of 144 GBM were analyzed using both  
512 Affymetrix U133A gene expression arrays and RNA sequencing. The empirical  $-\log(\text{P-value})$  of  
513 raw ssGSEA enrichment scores at each signature are shown as heatmaps, with dark blue  
514 representing no activation and bright red as highly activated. For each panel, the first row shows  
515 U133A based classification, and the second row indicates RNA-seq subtype classification.



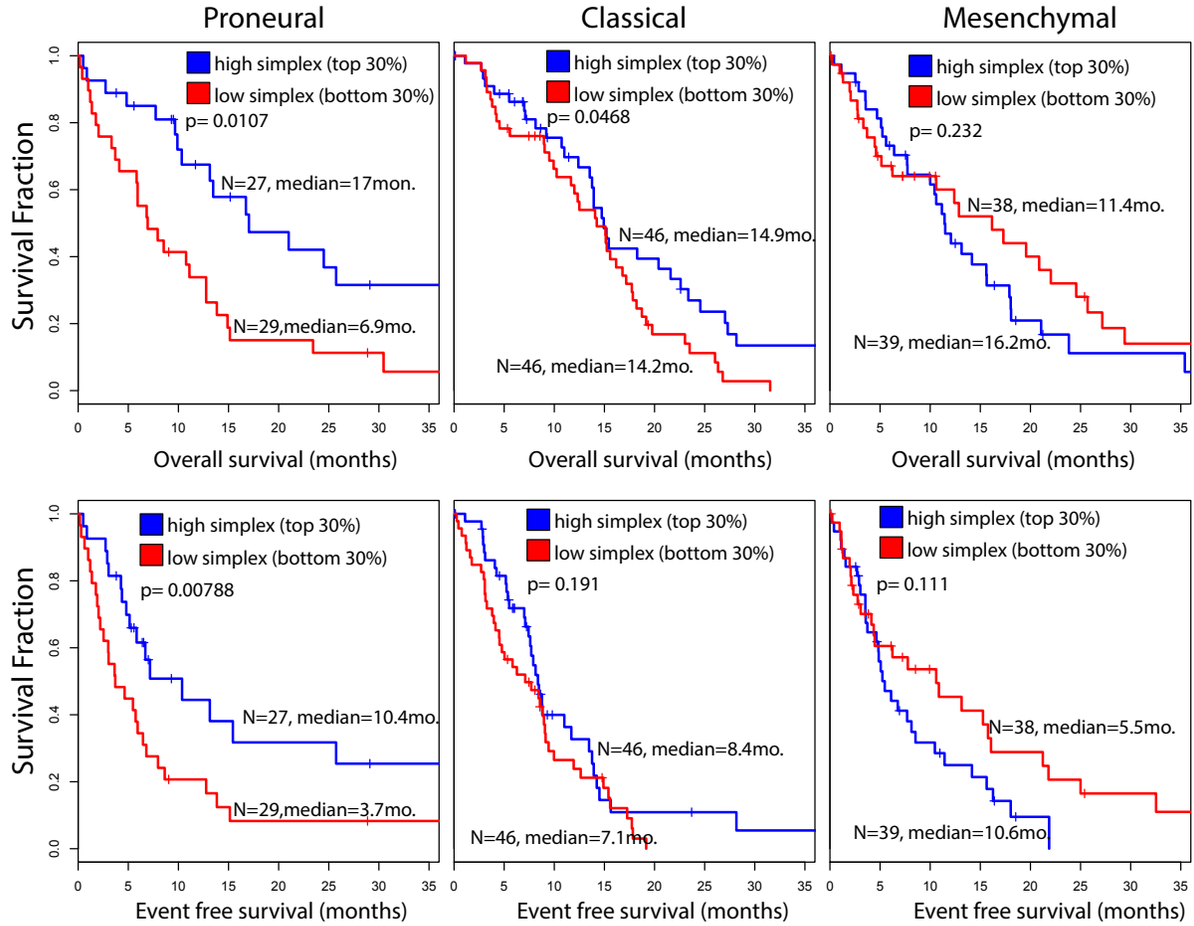
516

517 **Figure S3. Genomic alteration patterns for each subtype.** Ten prominent GBM somatic  
518 events were manually selected. Simplicity score, primary and secondary subtype were shown  
519 by the first three rows.



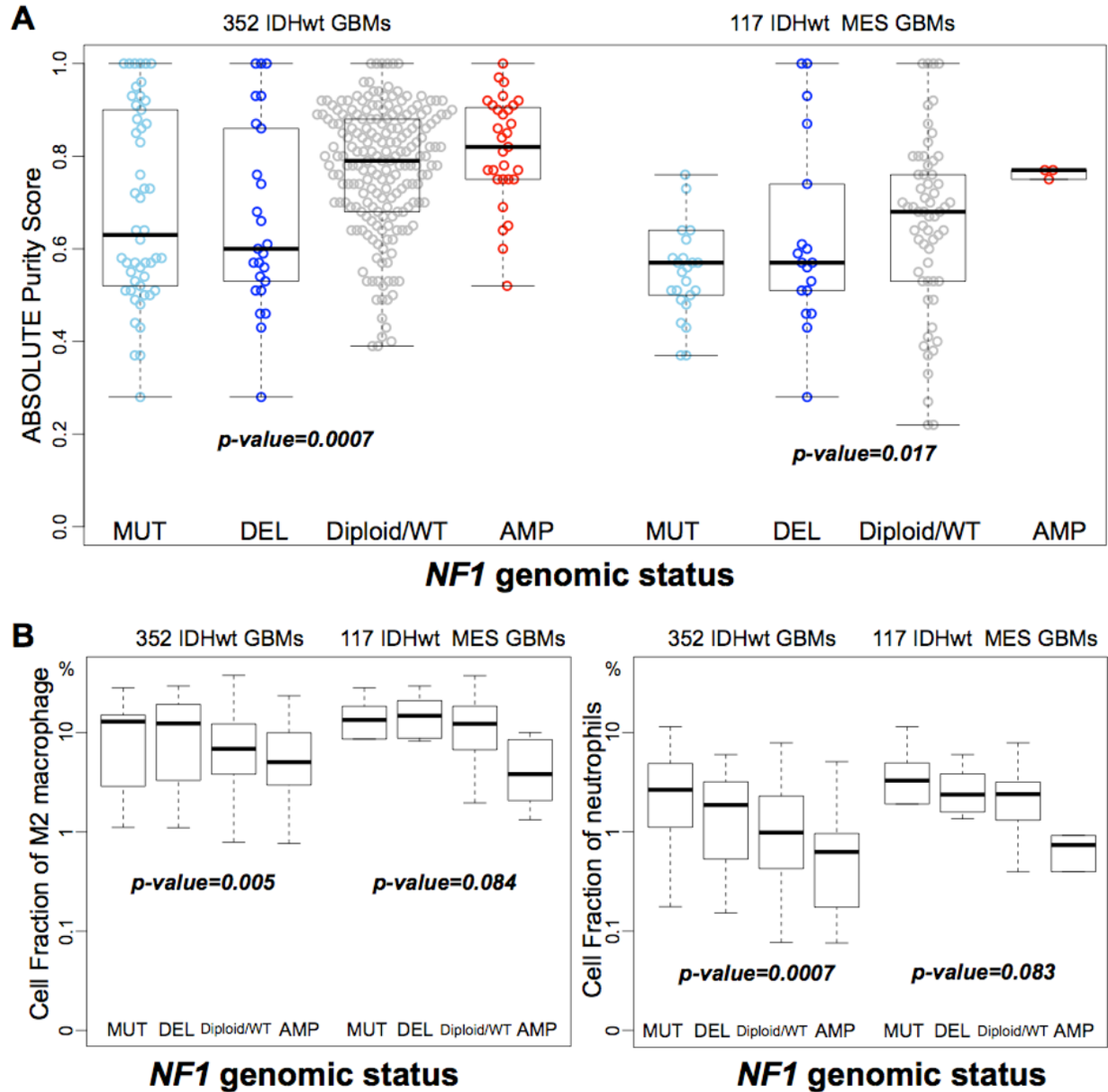
520

521 **Figure S4. Patient survival differences between transcriptional subtypes.** Samples were  
522 filtered using increased simplicity score as threshold from panel A to D.



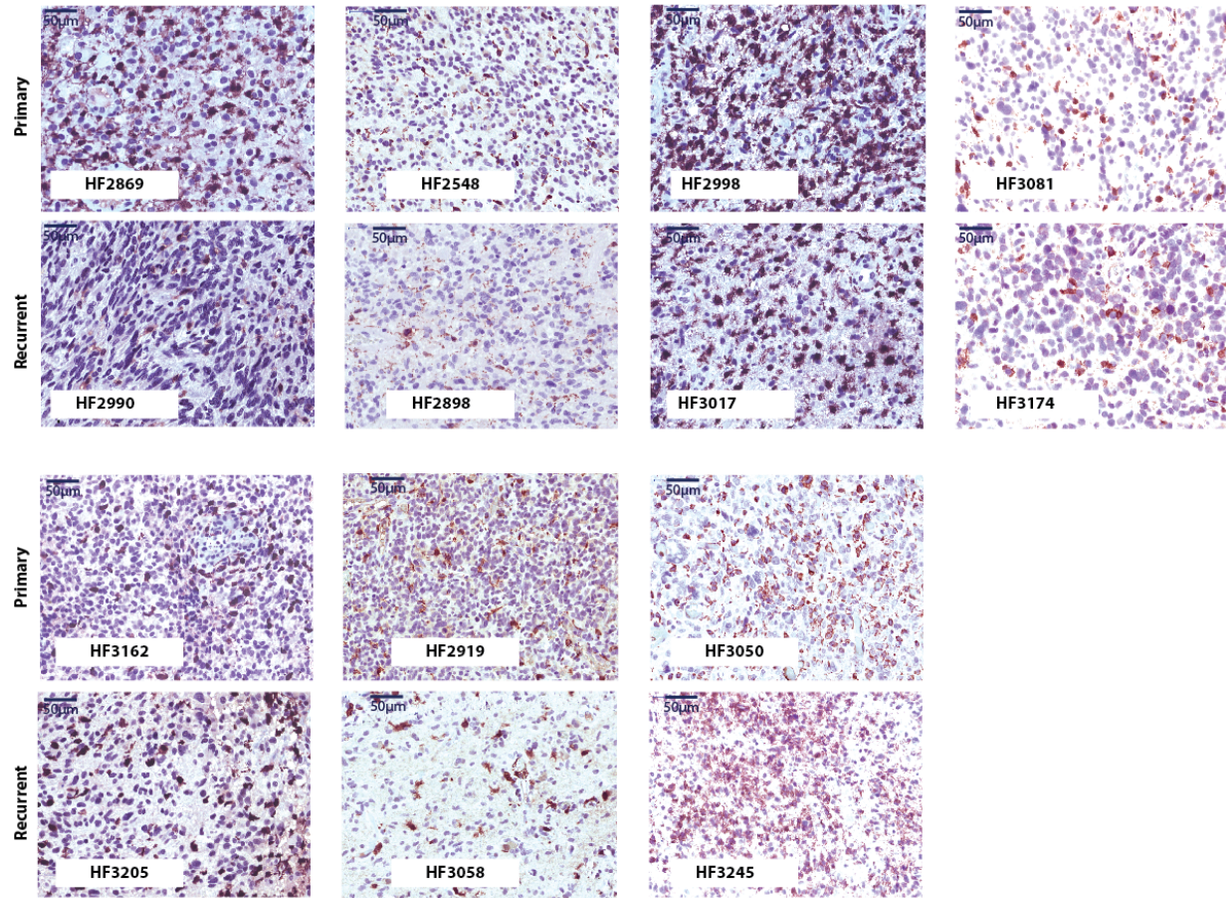
523

524 **Figure S5.** Overall and event free survival analysis comparison between samples with high and  
525 low simplicity scores in each subtype.



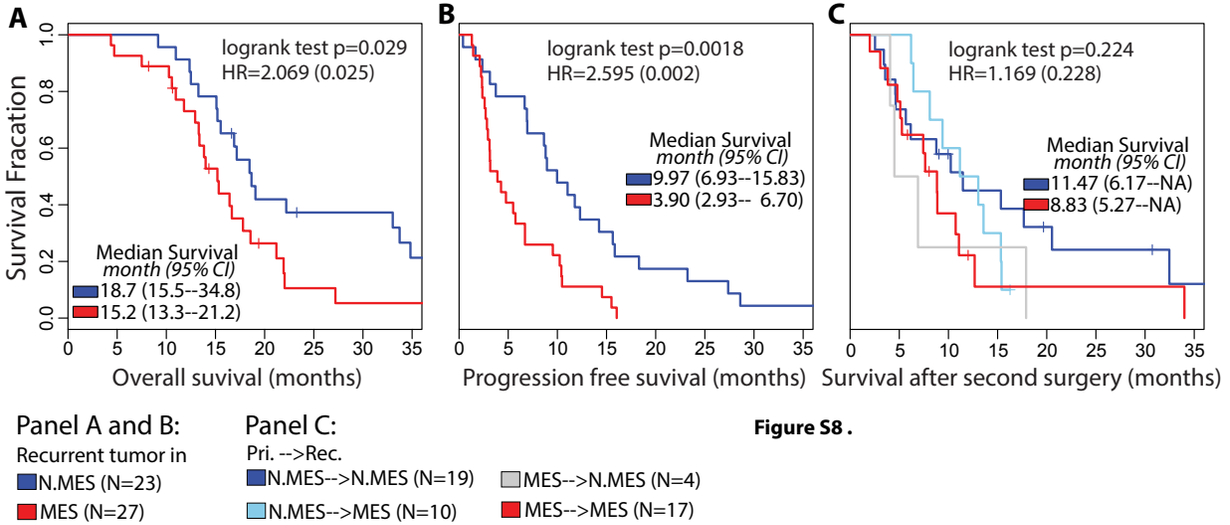
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**Figure S6. Comparison of tumor purity and immune cell fraction between GBMs with different *NF1* genomic status.** P-values were calculated using Wilcoxon rank test between samples carrying *NF1* deletion/mutation and others.



531

532 **Figure S7.** Immunohistochemistry staining of the IBA1 microglia marker.



533

534 **Figure S8. (A, B)** Overall and progression free survival analysis between samples in different  
 535 recurrent subtypes. **(C)** Survival after secondary surgery comparison between different  
 536 transition types.

537 **Supplementary Methods and Experimental Procedures**

538 **Collection of pairs of primary and recurrent glioma samples**

539 U133A array profiles for 543 primary GBM, and RNA-Seq data for 166 primary and 13  
540 recurrent GBM were obtained from the TCGA portal <https://tcga-data.nci.nih.gov/tcga/>.  
541 Mutation calls and DNA copy number profiles were obtained for all samples, where  
542 available. Tissues from 20 additional initial GBM and matched recurrent tumor were  
543 obtained from Henry Ford Hospital (n = 9) in accordance with institutional policies and  
544 all patients provided written consent, with approval from the Institutional Review Boards  
545 (Henry Ford Hospital IRB protocol #402). All RNA samples tested were obtained from  
546 frozen specimens. All of the recurrent GBMs had been previously treated with  
547 chemotherapy and radiation. Three cases had a history of lower grade astrocytoma  
548 prior to the first GBM (HF-2869/HF-3081/HF-3162). Tumors were selected solely on the  
549 basis of availability. RNA-Seq libraries were generated using RNA Truseq reagents  
550 (Illumina, San Diego, CA, USA) and paired-end sequenced using standard Illumina  
551 protocols. Read length was 76 base pairs for cases sequenced by TCGA and from  
552 Henry Ford (processed at MD Anderson). RNA-Seq data on frozen tissue from 44  
553 patients with initial and recurrent GBM that received resection at Samsung Medical  
554 Center and Seoul National University Hospital were provided by Dr. Nam's lab. Surgery  
555 specimens were obtained in accordance to the Institutional Review Board (IRB) of the  
556 Samsung Medical Center (No. 2010-04-004) and Seoul National University Hospital (No.  
557 C-1404-056-572). Affymetrix CEL files of 39 pairs of initial and recurrent glioma were  
558 retrieved from the Gene Expression Omnibus (GEO accession GSE4271, GSE42670,  
559 GSE62153)<sup>1-3</sup>. The expression profiles of the 23 pairs from GSE4271 were determined

560 using Affymetrix HG-U133 GeneChips, the 1 pairs from GSE42670 were analyzed using  
561 the Affmetrix HuGene-1-0-st platform, the 15 pairs from GSE62153 were analyzed  
562 using Illumina Human HT-12 V4.0 expression BeadChip.

563 Genome wide DNA copy number profiling and exome sequencing on thirteen  
564 TCGA tumor pairs and nine of ten Henry Ford tumor pairs were performed and data  
565 was analyzed using standard protocols and pipelines as previously described <sup>4</sup>.

566

### 567 **Data for multiplatform classification comparison**

568 RNA sequencing data was available for 162 primary GBMs<sup>5</sup> for which an Affymetrix HT-  
569 U133A gene expression profile was also available. We observed a low Pearson  
570 Correlation Coefficient ( $< 0.15$ ) between RNA sequencing based reads per kilo base of  
571 transcript per million reads (RPKM) and Affymetrix HT-U133A profiles in eighteen cases  
572 and these were removed from further analysis. In summary, in order to assess the  
573 concordance between classification results of the new 70-gene signatures and  
574 previously published 210-gene signatures <sup>6</sup>, 144 GBMs which were profiled in both RNA  
575 sequencing and Affymetrix U133A platforms were used in our further analyses.

576

### 577 **Transcriptome Data processing**

578 The latest version custom CDF files (Version19, <http://brainarray.mbni.med.umich.edu>)  
579 <sup>7,8</sup> were used to map probes from the Affymetrix HG-U133A and HuGene-1\_0-st  
580 GeneChip platforms to the Ensemble transcript database, combined in one probe set  
581 per gene and normalized using the AROMA package with default parameters, resulting  
582 in RMA normalized and log transformed gene expression values <sup>9</sup>. All RNA sequencing

583 data was processed by the PRADA pipeline <sup>10</sup>. Briefly, reads were aligned using BWA  
584 against the genome and transcriptome. After initial mapping, the aligned reads were  
585 filtered out if their best placements are only mapped to unique genomic coordinates.  
586 Quality scores are recalibrated using the Genome Analysis Toolkit (GATK), and  
587 duplicate reads are flagged using Picard. Mapped features were quantified and  
588 normalized per kilo base of transcript per million reads (RPKM) and were converted to a  
589 log<sub>2</sub> scale to represent a gene expression level. RPKM values measuring the same  
590 gene that mapped to the Ensemble transcript with longest size were selected to obtain  
591 one expression value per gene and sample. RPKM values were converted to a log<sub>2</sub>  
592 scale to represent gene expression level. The statistical environment R was used to  
593 perform all the statistical analysis and graph plots.

594

### 595 **Deriving new gene signatures**

596 A pair-wise gene expression analysis identified 5,334 genes which are significant higher  
597 expressed in glioma bulk samples compared to their derivative GSCs. These genes  
598 were discarded from the gene list for developing tumor-specific molecular subtypes.  
599 Consensus non-negative matrix factorization (CNMF) clustering method identified three  
600 distinct subgroups among the 369 IDHwt primary GBMs. A set of 270 GBMs was  
601 recognized as core samples based on a positive silhouette width. The gene expression  
602 values of each subtype were compared with those from the other two subtypes  
603 combined <sup>6</sup>. Signature genes per cluster were selected on the basis of differences in  
604 gene expression level and were considered significant if they reached the cut-off value  
605 with t-test p-value < 1E-3 for higher expressed in this class, while also showing a

606 significant lower expression with t-test p-value<1E-3 in the other two classes. In the  
607 original gene signatures, genes could be either down-regulated or up-regulated, while  
608 only up-regulated genes (n=70 per gene signature) were selected for revised gene  
609 signatures. Only genes measured on both RNAseq and U133A platforms were  
610 considered, and the U133A data from 162 GBM samples measured on both platforms  
611 (which included the 144 cases used to compare U133A and RNAseq results) was used  
612 in the final comparative analysis.

613

#### 614 **Molecular classifications based on ssGSEA enrichment scores**

615 Single sample gene set enrichment analysis was performed as follows. For a given  
616 GBM sample, gene expression values were rank-normalized and rank-ordered. The  
617 Empirical Cumulative Distribution Functions (ECDF) of the signature genes and the  
618 remaining genes were calculated. A statistic was calculated by integration of the  
619 difference between the ECDFs, which is similar to the one used in GSEA but is based  
620 on absolute expression rather than differential expression<sup>11</sup>. Since the ssGSEA test is  
621 based on the ranking of genes by expression level, the uncentered and log-transformed  
622 U133A and RPKM expression levels were used as input for ssGSEA. Since the scores  
623 of the three signatures were not directly comparable, we performed a resampling  
624 procedure to generate null distributions for each of the four subtypes. First we  
625 generated a large number of virtual samples in which each gene obtains its expression  
626 level by randomly selecting an expression value of the same gene in the remainder of  
627 the samples. Then, the three ssGSEA scores for each signature were calculated.  
628 Following this procedure we generated a large number (>1,000,000) of random ssGSEA

629 scores for each subtype, to build the null distribution and to give empirical p-values for  
630 the raw ssGSEA scores of each sample. By testing on multiple datasets with different  
631 sample sizes, we found the resampling generated distribution could be replaced with  
632 Student-T distribution (sample size>30) or Normal distribution (sample size>50) for  
633 getting very similar results. R-library with the code and expression matrices used is  
634 provided upon request.

635

### 636 **Evaluate the simplicity of subtype activation**

637 For a single sample, we decreased rank the empirical p-values for each subtype to  
638 generate order statistics as  $R_{N-1}, R_{N-2} \dots R_1, R_0$ . In particular,  $R_0$  equals to the minimum  
639 empirical p-value and points to the dominant subtype, i.e., the most significantly  
640 activated subtype. The accumulative distance to the dominant subtype (ADDS) was  
641 defined as:

$$ADDS = \sum_{i=1}^{N-1} (R_i - R_0)$$

642 Similarly, the accumulative distance between non-dominant subtypes (ADNS) as:

$$ADNS = \sum_{j>i>0} (R_j - R_i)$$

643 Obviously, the ADDS and ADNS are positive and negative correlated with single  
644 activation, respectively. Hence, we defined the simplicity score by combing ADDS and  
645 ADNS together and corrected with a constant  $\frac{(R_{N-1}-R_0)}{N-1}$  as follows:

$$Simplicity\ score = [ADDS - ADNS] \times \frac{(R_{N-1} - R_0)}{N - 1}$$

646 In a three subtypes case, the simplicity score of subtype activation can be simplified as:

647 
$$\text{Simplicity score} = (\text{median-min}) \times (\text{max-min}).$$

648

### 649 **Tumor purity assessment**

650 The ESTIMATE package was used to evaluate tumor purity on the basis of the  
651 expression level of marker genes in stromal and immune cells <sup>12</sup>, where the fraction of  
652 stromal cells and immune cells in each sample were represented by stromal score and  
653 immune score respectively, and the mixed fraction of both stromal and immune cells  
654 was represented by estimate scores. The ABSOLUTE package was used to confirm the  
655 tumor purity on the basis of chromosome copy number and allele fraction ratios on  
656 samples for which single nucleotide polymorphism array data were available <sup>13</sup>.

657

### 658 **Sample collection and Neurosphere Cultures**

659 After obtaining approval from the institutional review board of The University of Texas  
660 M.D. Anderson Cancer Center, glioblastoma tumor tissues were collected and named in  
661 the order that they were acquired. Each tissue was enzymatically and mechanically  
662 dissociated into single cells and grown in DMEM/F12 media supplemented with B27  
663 (Invitrogen), EGF (20 ng/ml), and bFGF (20 ng/ml), resulting in neurosphere growth. All  
664 cell lines were tested to exclude the presence of Mycoplasma infection. To minimize  
665 any batch effect the downstream molecular analyses were performed on identical cell  
666 culture batches. Total RNA from formalin fixed, paraffin embedded tumor tissues and  
667 matching neurospheres was prepared using the Masterpure complete DNA and RNA  
668 isolation kit (Epicenter) after proteinase K digestion per to the instructions from the  
669 manufacturer. Paired-end Illumina HiSeq sequencing assays were performed resulting

670 in a medium number of 50 million 75bp paired end reads per sample. We employed the  
671 PRADA pipeline to process the RNA sequencing data<sup>10</sup>. In short, Burroughs-Wheeler  
672 alignment, Samtools, and Genome Analysis Toolkit were used to map short reads to the  
673 human genome (hg19) and transcriptome (Ensembl 64) and RPKM gene expression  
674 values were generated for each of the 135,994 transcripts of 21,165 protein coding  
675 genes in Ensembl database.

676

### 677 **Western blotting**

678 Lysates were prepared from fresh frozen sections using RPPA lysis buffer (1% Triton X-  
679 100 50mM HEPES pH 7.4, 150mM NaCl, 1.5mM MgCl<sub>2</sub>, 1mM EGTA, 100mM NaF,  
680 10mM Na pyrophosphate, 1mM Na<sub>3</sub>VO<sub>4</sub>, 10% glycerol, plus protease and phosphatase  
681 inhibitors cocktails from Roche Applied Science #05056489001 and 04906837001), with  
682 sonication and clearing by centrifugation at 10,000g. Protein concentration was  
683 measured using the BCA kit (Thermo Scientific - Pierce #23225). SDS-PAGE and  
684 western blotting was performed using Midi gel system (Life Technologies - #WR0100)  
685 and NuPage-Novex 4-12% Bis-Tris Midi (20-well) Protein Gels (Life Technologies -  
686 #WG1402) using the following antibodies: ITGAM (CD11B) (Sigma Aldrich –  
687 #HPA002274), IBA1 (AIF1) (Sigma Aldrich – #HPA049234), GFAP (Cell Signalling –  
688 #3670), YKL40 (CHI3L1) YKL40 (CHI3L1, Santa Cruz Biotechnology - #sc-30465) a-  
689 actinin (Sigma Aldrich A5044) and Tubulin (Sigma Aldrich T9026).

690

### 691 **Immunohistochemistry**

692 Formalin-fixed, paraffin-embedded tissue sections (4  $\mu$ m thick) were collected on  
693 Superfrost plus slides. Briefly, tissue sections were deparaffinized with xylene and  
694 ethanol and re-hydrated with 95, 70 and 50% ethanol. Sections were antigen unmasked  
695 using citrate buffer (Vector Labs #H-3300) and heating. Peroxidase block was  
696 conducted with 3% H<sub>2</sub>O<sub>2</sub> and blocking was with 5% goat serum (Vector Labs #S-1000).  
697 Primary rabbit polyclonal antibody against IBA1 (AIF1)(WAKO #016-20001) at 1:400  
698 was used overnight. Secondary antibody was done using with the Rabbit-on-Rodent  
699 HRP-Polymer (Biocare #RMR622L) for 1 hr at room temperature. The slides were  
700 developed with Nova-red (Vector Labs #SK-4800) and counterstained with  
701 haematoxylin, mounted and scanned with Pannoramic 250 slide scanner (Caliper Life  
702 Sciences).

703

#### 704 **Analysis of the Human Protein Atlas database**

705 We systematically queried the human protein atlas database ([www.proteinatlas.org](http://www.proteinatlas.org),  
706 Version 12) for immunohistochemistry of specific markers of macrophage/microglial  
707 lineage<sup>14</sup>. Typically, each antibody was used on a set of tissue microarrays covering  
708 healthy controls and set of cancers of diverse sites, typically 10-15 individual cases for  
709 each cancer. Only antibodies with high validation scores (concordant RNA seq/IHC  
710 staining, western blot bands and antigen profile) were retained. ITGAM (Antibody HPA  
711 002274), IBA1 (Antibody HPA 049234) and CD163 (Antibody HPA 002432) met these  
712 criteria, as well as control protein vinculin (Antibody CAB002453).

## 713 **Supplementary References**

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