PDGF autocrine stimulation dedifferentiates cultured astrocytes and induces oligodendrogliomas and oligoastrocytomas from neural progenitors and astrocytes in vivo

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We present evidence that some low-grade oligodendrogliomas may be comprised of proliferating glial progenitor cells that are blocked in their ability to differentiate, whereas malignant gliomas have additionally acquired other mutations such as disruption of cell cycle arrest pathways by loss of Ink4a–Arf. We have modeled these effects in cell culture and in mice by generating autocrine stimulation of glia through the platelet-derived growth factor receptor (PDGFR). In cell culture, PDGF signaling induces proliferation of glial precursors and blocks their differentiation into oligodendrocytes and astrocytes. In addition, coexpression of PDGF and PDGF receptors has been demonstrated in human gliomas, implying that autocrine stimulation may be involved in glioma formation. In this study, using somatic cell type-specific gene transfer we investigated the functions of PDGF autocrine signaling in gliomagenesis by transferring the overexpression of PDGF-B into either nestin-expressing neural progenitors or glial fibrillary acidic protein (GFAP)-expressing astrocytes both in cell culture and in vivo. In cultured astrocytes, overexpression of PDGF-B caused significant increase in proliferation rate of both astrocytes and neural progenitors. Furthermore, PDGF gene transfer converted cultured astrocytes into cells with morphologic and gene expression characteristics of glial precursors. In vivo, gene transfer of PDGF to neural progenitors induced the formation of oligodendrogliomas in about 60% of mice by 12 wk of age; PDGF transfer to astrocytes induced the formation of either oligodendrogliomas or mixed oligoastrocytomas in about 40% of mice in the same time period. Loss of Ink4a–Arf, a mutation frequently found in high-grade human gliomas, resulted in shortened latency and enhanced malignancy of gliomas. The highest percentage of PDGF-induced malignant gliomas arose from of Ink4a–Arf null progenitor cells. These data suggest that chronic autocrine PDGF signaling can promote a proliferating population of glial precursors and is potentially sufficient to induce gliomagenesis. Loss of Ink4a–Arf is not required for PDGF-induced glioma formation but promotes tumor progression toward a more malignant phenotype.

[Key Words: PDGF; oligodendroglioma; INK4a–ARF; glial differentiation]

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The PDGF receptor has two isoforms (α and β) and belongs to the receptor tyrosine kinase family (Claesson-Welsh et al. 1989). Upon binding PDGF, these receptors activate a number of downstream signal transduction pathways, including PI3 kinase/AKT, RAS/MAP kinases, and PLC/PKC pathways (de Vries-Smits et al. 1992, Franke et al. 1995, Moriya et al. 1996). Accumulated evidence has suggested that PDGF signaling may play an important role in both normal development and tumorigenesis of the central nervous system (CNS). In cell culture, PDGF functions to block differentiation and promote proliferation of the O2A glial progenitor that gives rise to either oligodendrocytes or type-2 astrocytes (Raff et al. 1983; Noble et al. 1988; Richardson et al. 1988). Although it is known that PDGFR, especially the α isoform, is expressed at high levels in glial progenitors, mature astrocytes also express low levels of PDGFR (Hutchkins 1995; Fruttinger et al. 1996). PDGFRα and PDGF receptors are frequently coexpressed in human glioma cell lines as well as gliomas including oligodendrogliomas (Nister et al. 1988; Di Rocco et al. 1998; Robinson et al. 2001), and amplification of PDGFRα is found in some high-grade oligodendrogliomas (Shoshan et al. 1994, Smith et al. 2000), suggesting the possible existence of an autocrine loop comprising of PDGF and its receptors (Guha et al. 1995). Recently, PDGF-B chain, encoded in a replication competent retroviral vector system, was shown to induce primary brain tumors in mice with multiple histopathological appearances (Uhrbom et al. 1998). These data indicate that abnormal PDGF signaling can contribute to the etiology of brain tumors, but whether additional mutations are required for gliomagenesis in this system is not known. The variable histology seen in the above experiment may be due to the variety of cell types infected by this vector system. Dissecting the correlation between the histology of PDGF-induced tumors and their cell of origin requires cell type-specific genetic transfer.

We have achieved cell type-specific gene transfer of PDGF using the replication competent ALV splice acceptor (RCAS)/tv-a system (Holland and Varmus 1998; Holland et al. 1998a,b). This system consists of the avian leukosis virus (ALV)-based RCAS viral vectors and transgenic mice that express the RCAS receptor TVA from cell type-specific promoters. One mouse line [Gtv-a] expresses the TVA receptor from the astrocyte-specific glial fibrillary acidic protein (GFAP) promoter; a second mouse line [Ntv-a] expresses the TVA from the nestin promoter, as nestin is expressed in the CNS progenitor cells. Comparing the effects of gene transfer to Gtv-a and Ntv-a mice enables us to determine the influence of differentiation status on the ability of a cell to respond to a genetic stimulus. For example, the combined gene transfer of active forms of K-RAS and AKT induces GBM formation from Ntv-a mice but not Gtv-a mice (Holland et al. 2000a), implying that neural progenitors are more sensitive to the oncogenic effects of these particular signaling pathways than differentiated astrocytes.

In this study, we used an RCAS vector that encodes PDGF-B chain to infect Gtv-a and Ntv-a cells both in cell culture and in vivo. In culture, infection with this RCAS vector caused a significant increase in the proliferation rate of both GFAP-expressing astrocytes and nestin-expressing neural progenitors. Additionally, the dramatic changes in morphology and in the expression pattern of cellular differentiation markers are reversibly induced by overexpressing PDGF. These data indicate that signaling from this receptor appears to induce a conversion from differentiated astrocytes to glial precursor-like cells. In vivo, gene transfer of PDGF-B alone induced oligodendroglioma and oligoastrocytoma formation from nestin-expressing neural progenitors and GFAP-expressing astrocytes, respectively. Experimental loss of Ink4a–Arf was not required for PDGF-induced glioma formation; however, this additional mutation did result in shortened tumor latency and enhanced tumor malignancy. The highest percentage of malignant gliomas arose from the combination of neural progenitor origin and loss of Ink4a–Arf.

Results

Construction of RCAS vectors

An RCAS vector containing a bicistronic recombinant construct was generated by connecting the PDGF-B/c-sis and eGFP (green fluorescence protein) coding sequences together with an internal ribosome entry site (IRES), which results in expression of both gene products and allows detection of infection of cells. This vector is referred to as RCAS–PBIG (RCAS–PDGF-B–IRES–EGFP). A second RCAS vector was also constructed that contains only the PDGF-B sequence and is referred to as RCAS–PB. The vectors, along with an RCAS vector carrying the gene encoding β-galactosidase (lacZ) (gift of Yi Li, Memorial Sloan-Kettering Cancer Center, New York, NY) and another RCAS vector carrying pozyma virus middle T antigen (MTA) (Holland et al. 2000b), were propagated in the chicken fibroblast cell line DF-1.

PDGF-B overexpression promotes proliferation of both GFAP-expressing astrocytes and nestin-expressing CNS progenitors in cell culture

To test the mitogenic functions of PDGF-B in specific cell types, we infected primary brain cell cultures from both Gtv-a (astrocytes) and Ntv-a (glial progenitors) transgenic mice with RCAS–PB, RCAS–PBIG, and RCAS–lacZ vectors respectively. After infection, these cells were passaged in culture for 40 d and their proliferative capacity was measured by growth curves (Holland et al. 1998b). Infection of either Gtv-a (Fig. 1A) or...
Ntv-a cell cultures (data not shown) with PDGF-encoding viruses resulted in a substantial increase in the growth rate of these cells relative to infection with the control LacZ-encoding vector. A similar growth rate was seen for both RCAS–PBIG and RCAS–PB indicating that the expression of GFP did not affect the growth potential of these cells. Cells infected with RCAS–PBIG were visualized by green fluorescence secondary to the expression of eGFP [Fig. 1B].

**PDGF/PDGFRe autocrine loop stimulation dedifferentiates GFAP-expressing astrocytes to glial precursor-like cells in culture**

In addition to the enhanced proliferative capacity induced by PDGF signaling, PDGF also induced morphological and gene expression characteristics of undifferentiated cells. These changes were most striking in the astrocytes from Gtv-a mice. After 40 d in culture, the control RCAS–lacZ-infected Gtv-a cells showed a large, flat morphology expected for cultured astrocytes. In contrast, both RCAS–PB and RCAS–PBIG infected Gtv-a cells showed a small, elongated, bipolar or simple multipolar morphology, characteristics of glial progenitor cells [Fig. 1B].

Western blot and immunocytochemical analyses showed gene expression patterns similar to that of glial precursors [Fig. 1C]. Expression of myelin proteolipid protein (PLP) [Asakura et al. 1998; Spassky et al. 1998], PDGFRα [Hart et al. 1989; Hall et al. 1996] and β, A2B5 [Williams et al. 1985; Dubois-Dalcq 1987; data not shown], all known as markers for glial progenitors, were up-regulated in cells infected with PDGF-encoding vectors. Moreover, these cells also showed elevated expression of Id4, an HLH gene whose expression is shown to correlate inversely with differentiation status of oligodendrocyte precursors [Andres-Barquín et al. 1999; Konedo and Raff 2000; Nogueira et al. 2000]. By contrast, these cells did not express O4 antigen, which is expressed in premature and mature oligodendrocytes [Sommmer and Schachner 1981], implying that they are not differentiating to oligodendrocytes. Of note, NG2, which is expressed in bipotential O2A progenitors [Stallcup and Beasley 1987], was not expressed in these cells, nor was the expression of neuronal markers such as neurofilament (NF) or synaptophysin [data not shown]. Therefore, the expression pattern of cellular markers, along with the morphology and the proliferation capacity, suggest an early glial precursor stage for these cells, similar but not identical to the well-known O-2A progenitors (Raff et al. 1983). The potential for autocrine loop stimulation of PDGF and PDGFRe is demonstrated by the elevated expression of both PDGFRα and β compared with the RCAS–lacZ-infected cells.

**Blocking PDGF autocrine signaling reverses the PDGF-induced phenotype in vitro**

The PDGF-induced alterations described above could either be direct effects of elevated PDGF signaling or a result of selection for growth advantage in culture. To address this issue, we treated long-term cultured PDGF-transformed astrocytes [from Gtv-a mice] with a tyrosine kinase inhibitor [PTK787/ZK222584] (Wood et al. 2000) that blocks the activation of PDGF receptors. At 1 µM
drug concentration, the proliferation rate of the PDGF-expressing cells was dramatically inhibited (Fig. 2A,B) and the progenitor-like morphology reverted back to a large, flat, astrocyte-like morphology (Fig. 2C). Western blot analyses of RCAS–PBIG-infected astrocytes showed that the expression levels of PLP and Id4 were downregulated by treatment with PTK787 implying a direct effect of PDGF signaling in control of these genes. In contrast, the elevated PDGFRα and PDGFRβ levels seen in the PDGF overexpressing astrocytes were not reduced by PTK787, suggesting that the elevated expression of PDGFRs may be due to selection during extended culture for cells with high level of PDGFR expression that can maximally respond to overexpression of PDGF. The expression level of eGFP pre- and post-drug treatment was similar, indicating that the effects of PTK787 were not due to decreased expression of virally encoded PDGF-B (Fig. 2D). We demonstrated that this effect of PTK787 was not a general property of this drug by its effect on astrocytes infected with an RCAS vector carrying the coding sequence for polyoma middle T antigen (RCAS–MTA). MTA expression can activate similar signal transduction pathways as PDGF and infection with RCAS–MTA causes increased proliferation rate of astrocytes in cell culture and induces oligoastrocytomas from astrocytes in vivo (Holland et al. 2000b). At the same concentration of PTK787 that reversed the phenotype of RCAS–PBIG-infected cells, there was no effect on the proliferation capacity or morphology of RCAS–MTA-infected cells, implying specificity of the PTK787 effect for signaling through PDGFR in these glial cells.

**Gene transfer of PDGF-B into nestin-expressing neural progenitors induces oligodendroglioma formation in mice**

To investigate the effect of PDGF autocrine loop stimulation on these same cell types in vivo, we infected Ntv-a and Gtv-a transgenic mice with RCAS–PBIG by a single intracranial injection of about 10⁴ DF-1 cells producing this vector. All the mice were sacrificed at the end of week 12, or earlier if they showed signs or symptoms of intracranial disease such as macrocephaly or lethargy. The brains were fixed and sections were analyzed by standard hematoxylin and eosin (H&E) staining. Previous experiments have demonstrated that gene transfer of the marker gene alkaline phosphatase, activated Ras, or activated Akt alone to either of these mouse lines by this procedure does not induce tumors or histologic abnormalities (Holland and Varmus 1998; Holland et al. 2000a).

Initially, we directly infected a total of 34 newborn

![Figure 2](image_url)

**Figure 2.** Reversion of PDGF-induced phenotypic effects by blocking PDGF receptor kinase. Astrocytes (1 × 10⁵) were plated and grown in culture either with or without 1 µM PTK787 that inhibits the PDGF receptor kinase. The cells were counted after 14 d. (A) The proliferation of RCAS–PBIG-infected cultured astrocytes was inhibited by 1 µM PTK787 treatment. In contrast, the proliferation of RCAS–MTA-infected astrocytes was not inhibited by PTK787 (B). (MTA) Polyoma virus middle T antigen. (C) Giemsa staining shows the morphology of RCAS–PBIG-infected astrocytes with and without 1 µM PTK787 for 14 d. 100× magnification; bars, 100 µm. (D) Western blot analyses show the changes in expression pattern of molecular markers after blocking autocrine PDGF stimulation. (PLP) Myelin proteolipid protein; (PDGFR) platelet-derived growth factor receptor; (GFP) green fluorescent protein; (Id4) inhibitor of binding 4, a dominant negative HLH protein. α-Tubulin was used as loading control.
Ntv-a transgenic mice. Over 60% of this population of mice harbored gliomas. Histologically, these tumors were diffusely infiltrating neoplasms composed of small tumor cells with monotonous, regular round nuclei surrounded by cleared cytoplasm (“perinuclear halos”) (Fig. 3B). These morphologic features are identical to those seen in human oligodendrogliomas. The tumors did not express neuronal differentiation markers, such as NeuN and synaptophysin, as demonstrated by immunohistochemistry (data not shown). Further, also similar to human oligodendrogliomas, except for scattered entrapped reactive astrocytes the majority of tumor cells were immunonegative for GFAP, suggesting the absence of astrocytic differentiation (Fig. 3C). In addition to the morphologic and immunohistochemical similarity of the individual tumor cells to human oligodendrogliomas, the infiltrative behavior and interactions with normal host cellular constituents also closely resembled that seen in human gliomas. The murine gliomas displayed all of the classical morphologic formations seen in infiltrating human oligodendrogliomas referred to as secondary structures of Scherer (1938), including intrafascicular queuing in white matter tracts of the corpus callosum and subpial infiltration, perivascular satellitosis, and perineuronal satellitosis in areas of cortical invasion (Fig. 4). The expression of exogenous PDGF in these tumors was verified by positive immunostaining for eGFP expression (data not shown).

In the experiment reported here, both low- and high-grade tumors arose from PDGF gene transfer in vivo. The high-grade oligodendrogliomas, called anaplastic oligodendrogliomas in the current World Health Organization Classification of Tumours (WHO 2000), were characterized by dense cellularity without intervening neuropil, easily identifiable mitotic figures, cellular and nuclear pleomorphism and microvascular proliferation (Fig. 3D–F). These anaplastic oligodendrogliomas frequently were additionally characterized by the presence of foci of tumor necrosis with surrounding tumor cell pseudopalisading (Fig. 3F). The majority of the PDGF-induced gliomas, arising in either Gtv-a or Ntv-a mice having a wild-type genetic background, were low-grade tumors that lacked the features just described for high-grade oligodendrogliomas.

Loss of Ink4a–Arf enhances tumor initiation and tumor progression in Ntv-a transgenic mice

The above data suggest that autocrine stimulation of PDGF signaling, in the absence of other experimentally induced genetic alterations, could be sufficient to induce low-grade gliomas from neural progenitors. Human high-grade gliomas have additional mutations that result in disruption of cell cycle arrest pathways such as loss of Ink4a–Arf (Cairncross et al. 1998). To elucidate the roles of disruption of G1 cell cycle arrest pathway in PDGF-induced gliomagenesis, we compared the incidence and characteristics of gliomas occurring in both Ink4a–Arf+/−

![Figure 3. Histologic analysis of oligodendrogliomas induced from neural progenitors. PDGF-induced gliomas in Ntv-a mice with either Ink4a–Arf+/− (A–C) or Ink4a–Arf−/− (D–F) genetic backgrounds. (A) Low magnification of H&E-stained section illustrates the diffuse infiltrating low-grade tumor (indicated by the arrow). (B) H&E-stained section shows small, homogenous tumor cells with perinuclear “halo” appearance. (C) GFAP immunostaining shows negative oligodendroglioma cells and a few positive, trapped reactive astrocytes. (D) Low magnification of H&E-stained section illustrates a high-grade oligodendroglioma. The arrow indicates the necroses. (E) H&E-stained section shows microvascular proliferation [indicated by the arrow], a criterion for diagnosing high-grade gliomas. (F) H&E-stained section shows the palisading necrosis [indicated by the arrow], a criterion for diagnosing high-grade gliomas.](image-url)
and Ink4a–Arf−/− mice. At 12 wk of age, 61% of Ink4a–
Arf+/+ and 57% of Ink4a–Arf−/− mice developed gliomas
respectively (Fig. 5A). All of these tumors were oligoden-
drogliomas according to the above histopathological cri-
teria and as described in the WHO 2000 classification.
Although the overall glioma incidence was similar be-
tween these two groups, over half of the gliomas forming
in Ink4a–Arf+/+ mice were low-grade and asymptomatic
at the 12-wk time point whereas only about 10% of tu-
mors arising from Ink4a–arf−/− mice were low grade. By
contrast, using the same criteria these Ink4a–Arf−/− tu-
mors not only were more likely to be malignant but were
also more likely to be symptomatic, resulting in earlier
sacrifice and analysis (Fig. 5A, B). Therefore, experi-
mental disruption of the Ink4a–Arf locus appears not essen-
tial for the induction of these gliomas from nestin-ex-
pressing neural progenitors; however, it does either fa-
cilitate the progression of gliomas toward a more
malignant phenotype, or it selects for a more aggressive
tumor cell that outgrows the population in vivo.

Figure 5. Role of Ink4a–Arf loss in PDGF-induced gliomagenesis from neural progenitors. (A) Tumor-free survival curve indicating percent of mice developing symptomatic tumors over time. All mice were sacrificed at 12 wk, clinically occult tumors are indicated by the vertical line at 12 wk. The total number of mice without detectable tumors at 12 wk was ~40% in both animal groups as indicated by the arrows. (B) Ink4a–Arf loss significantly enhanced tumor malignancy. The grading criteria are described in the text. (O) Oligodendroglioma (WHO classification); (AO) anaplastic oligodendroglioma (WHO classification). AO tumors having the histo-
logic feature of pseudopalisading necrosis are listed as “anaplastic with necrosis.”
Gene transfer of PDGF-B into GFAP-expressing astrocytes induces either oligodendrogliomas or mixed oligoastrocytomas in mice

The experiments described thus far were performed in Ntv-a mice, that is, gene transfer to nestin-expressing neural progenitors. To test the influence of cell of origin on PDGF-induced gliomagenesis, we infected Gtv-a transgenic mice (astrocytes) with RCAS–PBIG. The total incidence of glioma formation was somewhat lower (∼40%), and the tumors were either oligodendrogliomas or gliomas of mixed histology containing both oligodendroglioma and astrocytoma components (oligoastrocytoma). Figure 6A shows a representative example of an oligoastrocytoma in which distinct oligodendroglioma and astrocytoma components are present. The oligodendroglioma component exhibits morphologic features identical to those seen in human oligodendrogliomas, including monotonous, uniform round nuclei and scant-to-nondetectable cytoplasm. By contrast, in the astrocytoma component the constituent tumor cells are more pleomorphic and atypical, with irregular nuclei and prominent eosinophilic cytoplasm that is strongly immunoreactive for GFAP (Fig. 6B,C). In addition, multinucleated giant cells are intimately associated with the astrocytoma component (data not shown), as is sometimes seen in human astrocytomas. These gliomas arising in Gtv-a mice also occurred in low-grade and malignant forms. However, over 70% of these tumors were low grade in mice with a wild-type Ink4a–Arf genetic background.

In Gtv-a mice, loss of Ink4a–Arf increases tumor incidence and tumor malignancy

To test the effects of Ink4a–Arf loss on the formation of the mixed gliomas arising from astrocytes in Gtv-a mice, we compared PDGF-induced glioma formation in Gtv-a, Ink4a–Arf+/− and Gtv-a, Ink4a–Arf−/− mice. Similar to the Ntv-a mice, gliomas arising from Gtv-a mice with Ink4a–Arf deficiency were of higher grade and shorter latency than those in mice with an Ink4a–Arf wild-type background (Fig. 7A). In contrast to Ntv-a mice, in which a higher percentage of animals developed gliomas and no increase in tumor incidence was seen with Ink4a–Arf loss, loss of Ink4a–Arf in Gtv-a mice resulted in a nearly twofold increase in the incidence of PDGF-induced gliomas at 12 wk of age (Fig. 7A). The most malignant gliomas with the shortest latency were seen with the combination of both neural progenitor cell of origin and loss of Ink4a–Arf (Fig. 7B).

Deletions of 1p, 19q, and 10q are not present in PDGF-induced gliomas

It is not clear whether PDGF signaling is sufficient for the induction of oligodendrogliomas or whether additional chromosomal abnormalities during tumor initiation or progression are required to cooperate with PDGF signaling in gliomagenesis. We investigated this question by analyzing these tumors for allelic loss of the mouse chromosomes syntenic to regions that are frequently lost in human gliomas. Approximately half of human oligodendrogliomas demonstrate deletions of chromosomes 1p and 19q, and malignant gliomas often show loss of chromosome 10q (Ransom et al. 1992; Reifenberger et al. 1994; von Deimling et al. 1992, Bello et al. 1995; Maier et al. 1997). Such regions would be likely candidates for additional mutations that might be required for PDGF-induced glioma formation. We addressed this issue by using fluorescence in situ hybridization (FISH) for murine loci that are syntenic to the candidate tumor suppressor regions on human chromosomes 1p, 19q, and 10q. Murine chromosomal arms (murine 3, 84.9 cM and murine 4, 46.6 and 81.5 cM, for human 1p; murine 7, 4–5.5 and 23.0 cM for human 19q; and murine 19, 26.0 cM for human 10q), and corresponding centromeres were studied in 21 PDGF-induced gliomas.

Figure 6. Histology of mixed gliomas induced from GFAP-expressing astrocytes. (A) Low magnification of GFAP-immunostained section illustrates the coexistence of both positive astrocytoma cells and negative oligodendroglioma cells. (B) H&E-stained section shows astrocytic tumor cells with irregular nuclei and eosinophilic cytoplasm. (C) GFAP immunostaining illustrates the strongly positive astrocytoma cells; the staining pattern distinguishes them from reactive astrocytes.
mas of various grades. For each case at each locus, the chromosomal counts ranged from 1.71 and 2.08, satisfying the criteria for two chromosomal copies (Fig. 8). Thus, in no case was there evidence for loss of any of these regions. Therefore, those common mutations implicated in human oligodendrogliomas appear not to be required for PDGF-induced glioma formation. These data may imply that PDGF signaling achieves the same biologic effect as 1p and 19q loss and that there is no selective pressure for generating these mutations, or that this model may simulate those human oligodendrogliomas without such mutations.

**Experimental loss of p53 does not promote PDGF-induced gliomagenesis**

The above data indicate that some of the common genetic alterations found in human gliomas do not occur in our PDGF-induced model for gliomas, potentially implying that other mutations are not required for PDGF-induced gliomagenesis. To further investigate the possibility that additional mutations might enhance the formation of PDGF-induced gliomas, we crossed the Gtv-a mouse to a p53−/− background. Loss of p53 has been shown to induce chromosomal instability in some tumor model systems and cultured astrocytes (Donehower et al. 1995; Yahanda et al. 1995; Bertrand et al. 1997). Intriguingly, mice with a p53−/− genetic background showed neither shortened tumor latency nor increased incidence of PDGF-induced gliomas relative to wild type mice (Fig. 7A). In fact, there appeared to be a somewhat lower tumor incidence in these p53−/− mice. This result is consistent with the finding that TP53 mutations rarely occur in human oligodendrogliomas (Ohgaki et al. 1993; Reifenberger et al. 1994; Bigner et al. 1999). In contrast, p53 loss cooperates with combined activation of Ras and Akt pathways in Gtv-a mice to induce glioblastomas (E.C. Holland et al., unpubl.), which is also consistent

**Figure 7.** Role of Ink4a–Arf loss, p53 loss in PDGF-induced gliomagenesis from Gtv-a mice. (A) Ink4a–Arf loss, but not p53 loss, resulted in a shortened tumor latency and an increased total tumor incidence (70% vs. 39%); (B) Ink4a–Arf loss also significantly enhanced the malignancy of gliomas induced from Gtv-a mice. In summary, the most malignant gliomas arose from the combination of both neural progenitor cell of origin and loss of Ink4a–Arf. (O) Oligodendroglioma (WHO classification); [AO] anaplastic oligodendroglioma (WHO classification). Final tumor incidence for each genetic background is indicated by the arrows.

**Figure 8.** Fluorescence in situ hybridization analysis of PDGF-induced gliomas. (A) Two copies of murine chromosome 4 syntenic to human chromosome 1p. Green and red signals indicate the locus of murine chromosome 4, centromere and 81.5 cM, respectively. (B) Two copies of murine chromosome 7 syntenic to human chromosome 19q. Green signals are for the centromere of murine chromosome 7, and red for chromosome 7, 23 cM.
with the high frequency of TP53 mutations in human astrocytic tumors (Louis et al. 1993; Wu et al. 1993; Weber et al. 1996). Therefore, although it remains formally possible that additional mutations are required for PDGF-induced glioma formation in mice, if such mutations exist they are not the common ones found in human oligodendrogliomas nor is their occurrence enhanced by loss of p53.

Discussion

Previous studies indicate that PDGF signaling functions during glial cell development by maintaining glial precursors in an undifferentiated and proliferating state. By forcing the overexpression of PDGF we have induced an autocrine signaling loop in cells that also express the PDGF receptor. In our experiments, the alterations in proliferation rate, cell morphology, and gene expression pattern in cultured astrocytes infected with RCAS–PBIG indicate that PDGF autocrine stimulation can dedifferentiate GFAP-expressing astrocytes to glial progenitor-like cells. This process can be reversed by blocking the PDGF receptor kinase, indicating that these PDGF-driven cells continue to depend on signaling through the PDGF receptor in culture, and have not genetically evolved to be independent of these signaling pathways.

The in vivo equivalent of this dedifferentiation effect of PDGF autocrine stimulation, and maintenance of glial progenitors in a proliferative capacity, appears to be the formation of low-grade gliomas. Depending on the cell of origin, these tumors are either predominantly oligodendroglomas from glial progenitors or oligoastrocytomas from astrocytes. Similar to human gliomas, the low-grade mouse gliomas do not have Ink4a–Arf loss. Additional Ink4a–Arf loss in the mouse model results in increased malignancy but only a modest increase in tumor incidence. All of these data are consistent with the hypothesis that low-grade gliomas may represent collections of glial progenitors that are trapped in an undifferentiated and proliferating state. There could well be numerous mechanisms that lead to glial progenitors being unable to differentiate in vivo. PDGF autocrine stimulation models this phenomenon in mice and may be responsible for a subset of human oligodendrogliomas. The conversion of these tumors to high-grade neoplasms could be due to acquisition of additional mutations, such as Ink4a–Arf loss, and result in an outgrowth of tumors with increased malignancy. This hypothesis implies that low-grade gliomas may respond well to drugs that promote differentiation such as histone deacetylase inhibitors and retinoic acid derivatives, and in some cases potentially PDGFR kinase inhibitors. High-grade gliomas that have additional mutations may be more refractory to this strategy alone.

The evidence presented in this paper indicates that PDGF signaling alone may be sufficient to induce glioma formation. The reversion of PDGF-induced phenotypes in vitro by blocking PDGF signaling strongly implies that PDGF signaling is necessary and sufficient to maint-
ground including contribution from C57Bl6, 129, Balb/C, and FVB/N.

Primary brain cell cultures

Newborn tv-a transgenic mice were sacrificed and the whole brains were mechanically dissociated into small pieces in sterile PBS, Ca²⁺, Mg²⁺ free (pH 7.4), followed by digestion with 1 mL of 0.25% Trypsin–1 mM EDTA in HBSS (GIBCO BRL) in sterile tubes and incubation in 37°C water bath for 15 min with gentle shaking. After incubation, fresh medium was added to terminate Trypsin digestion and large debris was settled. The single cells were pelleted, resuspended in DMEM with 10% fetal calf serum (GIBCO BRL), and plated.

Infection of primary brain cell culture and growth rate analysis

The supernatants containing various RCAS vectors from DF-1 cell cultures transfected with various RCAS vectors were collected using sterile syringes and filtered through 0.22 μm filters, followed by transferring into 70%–80% confluent primary brain cell cultures that had been plated and grown in DMEM with 10% fetal calf serum. Infections were repeated 3 times with 12-h intervals. After infection, the proliferation rates were measured. Briefly, the cells were harvested by trypsin digestion and counted using a hemocytometer. Total 1 × 10⁶ cells were re-plated and grown in DMEM with 10% fetal calf serum until the cells reached confluence; the same procedure was repeated over time. Finally, the total accumulated cell numbers were calculated and plotted as growth curves.

Fluorescence microscopy

DF-1 cells transfected with RCAS–PBIG and primary brain cell cultures infected with the same vector were seeded and grown on sterilized glass coverslips. First, the cells were fixed by immersing the coverslips in 2 mL of 4% paraformaldehyde in PBS (pH 7.4) for 30 min at room temperature. After washing with PBS three times, the coverslips were mounted on slides using Fluoromount G (EMS) and visualized under fluorescence microscope (Nikon Eclipse) using conventional FITC filter sets (Nikon B-1E, EX470–490, DM505, BA520–560) for green fluorescence. The red fluorescence of brain sections stained with anti-GFP antibody (Santa Cruz Biotech) was visualized by using Texas Red filter set (Nikon G-1B, EX540/10, DM580, BA590).

Giemsa staining of cultured cells

The medium was aspirated from cells at 80% confluence and the cells were washed with PBS (pH 7.4), followed by fixing in methanol at room temperature for 10 min. Then, 1 mL of Giemsa stain (LabChem Inc) was applied for 5 min. The stain was discarded and the cells were washed with PBS several times and visualized under a bright field microscope.

Immunocytochemistry

The cells infected with RCAS–PBIG or RCAS–lacZ were fixed in either 4% paraformaldehyde in PBS (pH 7.4) or cold methanol. The dishes were blocked by using 5% normal horse serum diluted in PBS (pH 7.4) for 1 h at room temperature with shaking. Monoclonal anti-A2B5 antibody (Chemicon), polyclonal anti-NG2 antibody (Chemicon), monoclonal anti-O4 antibody (Chemicon), monoclonal anti-GFAP antibody (Boehringer Mannheim), monoclonal anti-NeuN antibody (Chemicon), and monoclonal anti-neurofilament antibody (Boehringer Mannheim) were used as primary antibodies. Appropriate biotinylated secondary antibodies and avidin-conjugated peroxidase were purchased from Vector Lab. The signal was visualized by using DAB substrate kit (Vector Lab) and examined under microscopy.

Blocking of PDGF signaling

The cells infected with RCAS–PBIG were cultured in DMEM supplemented with 10% FCS. Cells [1 × 10⁵] were plated and the experiment groups were treated with PTK787 (stock solution dissolved in DMSO) with the final concentration of 1 μM for 2 wk. The control groups were treated with the same volume of DMSO. After two wk, the cell numbers were counted, and the morphology was visualized by Giemsa staining.

Western analysis

Whole-cell protein extracts were prepared by using cold lysis buffer consisting of: 100 mM NaCl, 30 mM Tris-HCl (pH 7.6), 1% NP-40, 30 mM sodium fluoride, 1 mM EDTA, 1 mM sodium vanadate, 0.5 mM PMSF, and protease inhibitor cocktail tablets (Boehringer Mannheim). Samples were incubated on ice for 30 min and supernatants were recovered by centrifuging at 14,000 rpm at 4°C for 20 min. Protein concentrations were determined by BCA method (Pierce Biochemical). Proteins were separated on 10% SDS-PAGE and transferred to nitrocellulose membrane (Osmonics). Blocking reagent was 5% non-fat dry milk in TBS (pH 7.4). Washing buffer was TBS (pH 7.4) with 0.1% Tween-20. Polyclonal rabbit anti-PLP (Oncogene Research), polyclonal rabbit anti-PDGFRα and β (Upstate Biotech), polyclonal goat anti-vimentin (Chemicon), polyclonal rabbit anti-Ih4 (Santa Cruz Biotech), polyclonal rabbit anti-GFAP (Santa Cruz Biotech), and monoclonal anti-α-tubulin (Sigma) were used as primary antibodies. Respective HRP-conjugated secondary antibodies were purchased from Boehringer Mannheim. Signals were visualized by using ECL chemiluminescence (Amersham) and Kodak X-Omat films.

Infection of tv-a-transgenic mice

DF-1 cells producing various RCAS viruses were trypsinized and pelleted by centrifugation; the pellets were resuspended in ~50 μL of DMEM medium and placed on ice before injection. Using a 10 μL gas-tight Hamilton syringe, a single intracranial injection of 1 μL of cell suspension (~10⁵ cells) was made in the right frontal region of newborn mice, with the tip of the needle just touching the skull base.

Brain sectioning, H&E staining, and immunohistochemistry

Mice were sacrificed before (because of early symptoms) or at 12 wk of age and the whole brains were fixed in 4% formaldehyde in PBS for at least 36 h with shaking. Five sections of each brain were cut and embedded in paraffin, 5-micron sections were cut with a Leica microtome. The sections were stained with H&E. Immunostaining was performed using ABC kits (Vector Lab). Briefly, deparaffinized slides were first treated with antigen unmasking reagent (Vector Lab) with heating in a steamer for 30 min, followed by immersing in 10% hydrogen peroxide in methanol for 20 min to inactivate the endogenous peroxidases. Then the sections were blocked with 1.5% normal horse serum in PBS (pH 7.4) for 1 h at room temperature. Monoclonal anti-GFAP (Boehringer Mannheim), monoclonal anti-NeuN (Chemicon), polyclonal rabbit anti-synaptophysin (Biomeda), and poly-
clonal rabbit anti-GFP [Santa Cruz Biotech] were diluted in ChemMate antibody dilution buffer [Ventana Medical Systems] and incubated with sections at 37°C for 1 h. After washing with PBS/0.05% Tween 20, appropriate biotinylated secondary antibodies (Vector Lab) diluted in the same antibody dilution buffer were incubated with sections at 37°C for 30 min. Then, after washing, avidin-conjugated peroxidase or alkaline phosphatase (Vector Lab) diluted in PBS containing 1.5% normal horse serum was incubated with sections for 30 min at room temperature. Finally, after exclusive PBS-T washing, DAB or alkaline phosphatase red substrate [Vector Lab] was added to develop the color. After terminating the staining reaction, the sections were counterstained with hematoxylin and mounted. The negative controls were included with the same procedure except replacing primary antibodies with antibody dilution buffer.

**FISH**

The following BACs [Research Genetics, Huntsville, AL] were used to assay for chromosomal loss at murine regions syntenic to human glioma tumor suppressor regions [human/mouse homology maps are available at http://www.ncbi.nlm.nih.gov/ Homology/index.html and http://greengenes.llnl.gov/mouse/]; for regions syntenic to human 1p; BAC 432-H-2 [murine chromosome 3, 84.9 cM] [Korenberg et al. 1999]; BAC 150-L-24 [murine chromosome 4, 46.6 cM] [Chua et al. 1996], and BAC 362-D-3 [murine chromosome 4, 81.5 cM] [Korenberg et al. 1999]; for regions syntenic to human 19q13.3, BACs 305-J-2 [murine chromosome 7, 4–5.5 cM] and 419-I-21 [murine chromosome 7, 23.0 cM]; for the Pten locus, BAC 371-O-8 [murine chromosome 19, 26 cM] [Hansen and Justice 1999]. Centromeric BACs were also used to confirm copy numbers for chromosomes 3, 4, 7, and 19 (BAC 52-H-9, chromosome 3; BAC 39-B-18, chromosome 4; BAC 43-A-19, chromosome 7; and BAC 26-B-5, chromosome 19) [Korenberg et al. 1999]. Purified BAC DNA was fluorescently labeled with SpectrumOrange or SpectrumGreen [Vysis Inc., Downers Grove, IL] by nick translation and applied to slide-mounted tissue sections that had been dewaxed, rehydrated, and treated in 10 mM sodium citrate for 30 min at 80°C and pepsin [4 mg/mL in saliné, pH adjusted to 1.5 with HCl] for 15 min. Following denaturation for 10 min at 80°C, the sections were incubated overnight at 37°C. After washing in buffers at 45°C (2× 15 min in 1.5 M urea/0.1× SSC, 5 min in 2× SSC, 5 min in 2× SSC/0.1% NP-40), the slides were counterstained with 4,6-diamidino-2-phenyindol (DAPI), covered with coverslips and incubated with sections at 37°C for 1 h. After washing with PBS/0.05% Tween 20 and incubation with primary antibodies, an appropriate biotinylated secondary antibody (Vector Lab) diluted in PBS/0.05% Tween 20, was added and incubated with sections overnight at room temperature. After washing, sections were counterstained with hematoxylin and mounted. The negative controls were included with the same procedure except replacing primary antibodies with antibody dilution buffer.

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