Transplanted Oligodendrocyte Progenitor Cells Expressing a Dominant-Negative FGF Receptor Transgene Fail to Migrate In Vivo

Donna J. Osterhout,1 Sylvie Ebner,1 Jingsong Xu,2 David M. Ornitz,2 George A. Zazanis,1 and Randall D. McKinnon1

1Division of Neurosurgery, Department of Surgery, University of Medicine and Dentistry of New Jersey, Robert Wood Johnson Medical School, Piscataway, New Jersey 08854, and 2Department of Molecular Biology and Pharmacology, Washington University Medical School, St. Louis, Missouri 63110

The proliferation, migration, survival, and differentiation of oligodendrocyte progenitor cells, precursors to myelin-forming oligodendrocytes in the CNS, are controlled by a number of polypeptide growth factors in vitro. The requirement and roles for individual factors in vivo, however, are primarily unknown. We have used a cell transplantation approach to examine the role of fibroblast growth factor (FGF) in oligodendrocyte development in vivo. A dominant-negative version of the FGF receptor-1 transgene was introduced into oligodendrocyte progenitors in vitro, generating cells that were nonresponsive to FGF but responsive to other mitogens. When transplanted into the brains of neonatal rats, mutant cells were unable to migrate and remained within the ventricles. These results suggest a role for FGF signaling in establishing a motile phenotype for oligodendrocyte progenitor cell migration in vivo and illustrate the utility of a somatic cell mutagenesis approach for the study of gene function during CNS development in vivo.

Key words: CNS development; myelin; oligodendrocyte; O-2A progenitor; transplantation; migration; fibroblast growth factor receptor; dominant-negative

Oligodendrocytes, the myelinating cells of the CNS, are generated from progenitor cells that arise in the subventricular zone and then migrate through the brain parenchyma into axonal tracts and gray matter (Hardy and Reynolds, 1991; Levison et al., 1993; Price, 1994). These migratory precursors arise at embryonic day 12–14 in the rat spinal cord (Noll and Miller, 1993; Timis et al., 1995) from neuroectodermal cells positioned throughout the rostral–caudal neural axis (Hardy and Friedrich, 1996) and are thought to originate in the ventral–lateral portion of the neural tube (Pringle and Richardson, 1993; Miller, 1996). In vitro studies have identified distinct stages of oligodendrocyte progenitor cell maturation (Pfeiffer et al., 1993) from preprogenitors (Grinspan et al., 1990; Hardy and Reynolds, 1991) to motile progenitor cells (Raff et al., 1983), to nonmotile late progenitor cells (Dubois-Dalcq, 1987), and finally to postmitotic oligodendrocytes that form myelin internodes on contact with neuronal axons. The motile progenitors, first characterized from the rat optic nerve, generate either oligodendrocytes or a type of astrocyte under different culture conditions and are termed O-2A (oligodendrocyte-type-2 astrocyte) progenitor cells (Raff et al., 1983).

A number of polypeptide growth factors have been identified that affect oligodendrocyte progenitor cell development in vitro, including factors that affect progenitor cell proliferation (Noble et al., 1988; Raff et al., 1988; Bogler et al., 1990; McKinnon et al., 1990; Barres et al., 1994a; Canoll et al., 1996), migration (Armstrong et al., 1990; Milner et al., 1997), survival (Barres et al., 1993; Mayer et al., 1994; Gard et al., 1995; Yasuda et al., 1995), and differentiation (McMorris et al., 1986; McMorris and Dubois-Dalcq, 1988; McKinnon et al., 1993b; Barres et al., 1994b; Noll and Miller, 1994) and the synthesis of myelin (McMorris and Dubois-Dalcq, 1988). The most extensively characterized of these, platelet-derived growth factor (PDGF), promotes cell survival (Barres et al., 1992) and limited cell division (Noble et al., 1988; Raff et al., 1988) and induces a phenotype characterized by a bipolar morphology (Gard and Pfeiffer, 1993; McKinnon et al., 1993a) that is specifically associated with cell migration (Small et al., 1987). To date, PDGF is the only cytokine known to be chemotactic for oligodendrocyte progenitor cells in vitro (Armstrong et al., 1990). The fibroblast growth factors FGF-1 and FGF-2 are also mitogenic for oligodendrocyte progenitors (Besnard et al., 1989; McKinnon et al., 1990). In contrast to PDGF, FGF-2 promotes unlimited division and prevents oligodendrocyte progenitors from entering terminal differentiation (McKinnon et al., 1990). FGF also induces the expression of PDGF α-receptors (PDGFRα) on oligodendrocyte progenitor cells, increasing their sensitivity to PDGF (McKinnon et al., 1990). Thus, although FGF is not chemotactic (Armstrong et al., 1990), it primes oligodendrocyte progenitors to respond to PDGF and thereby contributes to their ability to adopt the migratory phenotype (McKinnon et al., 1993a).
FGF stimulates the proliferation of progenitor cells \textit{in vitro} in both neural and non-neural systems (for review, see Baird, 1994). FGF-1 and FGF-2 are expressed in both developing and adult brain (Ernfors et al., 1990; Gonzalez et al., 1990; Kalcheim and Neufeld, 1990; Schnurich and Risau, 1991), and FGF-1 is associated with enhanced myelination after a demyelinating lesion (Tourbah et al., 1992). Although these studies are consistent with a role for FGF in CNS myelination, this has yet to be demonstrated \textit{in vivo}. To date, a genetic approach to study the role of FGF in oligodendrocyte development \textit{in vivo} has not been informative, because targeted disruption of the murine FGF receptor gene \textit{fgfr} results in aborted development before or during gastrulation (Deng et al., 1994; Yamaguchi et al., 1994) and precludes an examination of the consequences of null mutations on later-emerging tissues.

We have taken an alternative approach to examine the role of FGF signaling in CNS myelination \textit{in vivo}. Isolated progenitor cells were rendered nonresponsive to FGF \textit{in vitro} using a dominant-negative FGF receptor 1 (FGFR1) transgene; then their development was examined after transplantation into neonatal rodent brain. We demonstrate that oligodendrocyte progenitors with impaired FGF signaling are unable to migrate into parenchyma and persist within the ventricles of recipient brain. These studies thus indicate that oligodendrocyte progenitor cells require FGF signaling to acquire their migratory phenotype \textit{in vivo} and demonstrate the utility of a somatic cell mutagenesis approach for the study of gene function during CNS development.

**MATERIALS AND METHODS**

\textbf{Recombinant DNA.} The construct pMo.FGFRx.iresNeo (pMoFRx) was used in this study (Fig. 1A) contains a cDNA copy of the murine FGF (mFGF) receptor-1 (Ornitz and Leder, 1992; Benvenisty and Ornitz, 1995) and the neo resistance (G418 \textsuperscript{8}) gene (Southern and Berg, 1982), under transcriptional regulation of the Moloney MuLV retroviral long terminal repeat (Mo-LTR) promoter. The cDNA insert FGFRx encodes the three Ig domain forms of FGFR1 (splice form c), generating a protein that binds both FGF-1 and FGF-2 (Ornitz and Leder, 1992; Ornitz et al., 1994). A syntenic construct was placed at position 418 by an internal ribosome entry sequence (IRES) derived from FGFRx, insertion mutagenesis (Benvenisty and Ornitz, 1995). The encoded protein product contains extracellular and transmembrane domains of FGFR1 but lacks a cytoplasmic (catalytic protein tyrosine kinase) domain. This construct produces a biecostic transcript with mFGFR1 and neo separated by an internal ribosome entry sequence (IRES) derived from the IRES-containing region of the neomycin phosphotransferase II gene (Ghattas et al., 1991). Because neo lies downstream (3\textsuperscript{9}) from FGFR1, G418 resistant cells should express both transgenes. Control constructs encoding neomycin phosphotransferase II (NPTII; G418 \textsuperscript{R}) were generated (Benvenisty and Ornitz, 1995).

\textbf{Primary cell culture.} Primary glial cultures were established from 2-d-old Sprague Dawley rats (Taconic Farms, Germantown, NY), and A2B5-immunoreactive oligodendrocyte progenitor cells were isolated from these cultures by immunoselection as described previously (McKinnon et al., 1990). Purified cells were plated on Falcon culture dishes (Becton Dickinson, Rutherford, NJ) or glass coverslips (Bellco Glass, Cherry Hill, NJ) in saline, followed by 4% paraformaldehyde in phosphate-buffered saline (PBS), frozen in OCT compound (TissueTek; Baxter Scientific, Boston, MA), and serially sectioned (20 \mu m) for microscopy.

\textbf{DNA transfections.} DNA transfections were performed using the calcium technique (Graham and Van der Eb, 1973) with cells plated at a density of 2 x 10\textsuperscript{5} per 60 mm dishes in medium containing 10\% FBS for 72 hr as described previously (McKinnon and Zazanis, 1996). DNA precipitates were prepared by combining 1 \mu g of plasmid DNA purified by cesium chloride gradient centrifugation with 14 \mu g of high molecular weight rat genomic DNA in 10 mm Tris, pH 7.5, and 1 mm EDTA, followed by the addition of 0.1 x volume of 2.5 M CaCl\textsubscript{2}. This was added dropwise to an equal volume of 2 x HEPEPS-buffered saline, pH 7.05, with mixing, generating a CaPO\textsubscript{4}–DNA coprecipitate that was added directly to the culture medium. The cultures were refed after 72 hr and every 4 d thereafter with defined medium containing 20% B104-cm plus 100 \mu g/ml genitin (G418; Sigma). B104-cm was replenished every 48 hr. Colonies were routinely visible after 10 d of culture, and individual colonies were isolated in cloning cylinders (Bellco Glass) and expanded as subclones in medium containing 200 \mu g/ml G418.

\textbf{In vitro analysis.} Mitogen-stimulated cell proliferation assays were performed in 96 well Falcon plates in culture medium supplemented with recombinant human PDGF-AA or FGF-2 (R & D Systems, Minneapolis, MN). Cells were plated at 3-4 x 10\textsuperscript{4} cells/well in triplicate wells, and the cells were treated with various concentrations of mitogens for 72 hr, and DNA synthesis was measured by measuring \textit{in vitro} characterization.

\textbf{Cell transplantation.} Transfected cells expressing recombinant constructs encoding neomycin phosphotransferase II (NPTII; G418 \textsuperscript{8}) were maintained in medium lacking G418 for one passage before transplantation. All cells were labeled \textit{in vitro} with fluorescent cell markers (either PKH2 (fluorescein) or PKH26 (rhodamine) optics; Sigma) according to the manufacturer’s directions, then resuspended in PBS (10 mm sodium phosphate, 150 mm NaCl, pH 7.2), and transplanted into the right thalamus of neonatal rat recipients at postnatal day 2 (P2) or day 6 (P6). The animals were anesthetized with halothane (Halocarbon Labs, River Edge, NJ) before transplant. A Hamilton 701 \mu l syringe was used to inject 100 \mu l (5,000 \(\times\) transfected cells) into the brain 1 mm rostral from bregma and 1 mm right of midline, to a depth of 3 mm (P2 rat) or 5 mm (P6 rat). The animals were returned to their mothers after the procedure. At the indicated times after transplantation, the animals were anesthetized with sodium pentobarbital (40 mg/kg, i.p.; Abbott Labs, Irving, TX) and perfused with 1 U/ml heparin (Elkins-Sinn, Cherry Hill, NJ) in saline, followed by 4\% paraformaldehyde in phosphate buffer, pH 7.4. The brains were removed, post-fixed in 4\% paraformaldehyde overnight, equilibrated in 10\% followed by 25\% sucrose in PBS, frozen in OCT compound (TissueTek; Baxter Scientific, Boston, MA), and serially sectioned (20 \mu m) sections on a Jung Frigocut cryostat. Transplanted cells were visualized in the sections using a Zeiss Axiosvert 100 TV with a 40\times LD Acroplan objective.

\textbf{Immunohistochemistry.} Cells growing on 12 mm glass coverslips (Bellco Glass) were washed in PBS, fixed for 5 min in 2\% paraformaldehyde, and then exposed to antibody solutions for 30 min at room temperature in a humid chamber as described previously (McKinnon et al., 1990). Primary antibody dilutions in PBS were: monoclonals A2B5, O1, and O4, 1:10 dilution of tissue culture supernatants; anti-myelin basic protein (MBP) serum, 1:500 dilution; and anti-GFAP, anti-\textit{5-bromo-2′-deoxyuridine} (BrdU), and anti-neurofilament 160 kD (anti-SMI-31) antibody (Oncogene Research Products, Cambridge, MA), as recommended by the manufacturer. Fluorescent-conjugated secondary antibodies (Pierce, Rockford, IL) were used at 25 \mu g/ml. For MBP immunoreactivity, the coverslips were first treated with Bouin’s fixative.
(5 min) and then 10% normal goat serum (10 min) before incubating with antiserum (60 min), and immunoreactivity was detected by incubating with biotinylated anti-antibody for 30 min, washing in 0.1 M NaHCO3, and 0.15 M NaCl, pH 8.4, and then incubating with fluorescein avidin (10 µg/ml; 30 min; Vector Laboratories, Burlingame, CA). The coverslips were mounted on glass slides with Fluoromount-G (Southern Biotechnology, Birmingham, AL) and then viewed and photographed with Kodak TMAX 400 ASA print or Elite 110 color slide film.

FGF protein was detected in cultured cells in vitro, and in transplanted cells in vivo, using biotinylated anti-NPTII (5 Prime–3 Prime Inc., Boulder, CO). Cultured cells were fixed in acid:alcohol (5:95) at −20°C for 5 min, followed by incubation with anti-NPTII antibody (1:100 in PBS) for 2 hr at room temperature (RT). Cryostat sections of paraformaldehyde-fixed tissue samples (grafted brain) were permeabilized with 0.5% Triton X-100 in PBS for 5 min and then incubated first with 10% normal goat serum for 30 min at RT followed by anti-NPTII antibody (1:100 in PBS) overnight at 4°C. The sections were rinsed and incubated in ABC reagent according to the manufacturer’s instructions (Vectorstain Elite kit; Vector Laboratories), and NPTII immunoreactivity was detected with the DAB substrate kit (Vector Laboratories).

The proliferation of grafted cells was determined by immunohistochemical analysis of BrdU (Sigma) incorporation (Nowakowski et al., 1989). Transplant recipients received six intraperitoneal injections (each injection, 50 µg/gm) over the course of 9 hr and then were perfused as described above 30 min after the last injection. The brains were embedded in paraformaldehyde and serially sectioned (8 µm sections). Before staining, the sections were deparaffinized and permeabilized using 0.1% trypsin followed by 2 N HCl treatment, then incubated with anti-BrdU antibody (1:75 dilution; Chemicon), and visualized with the Vectorstain Elite and DAB substrates (Vector Laboratories) according to the manufacturer’s instructions.

Biological techniques. For Northern blot analysis, total RNA was isolated from cultured oligodendrocyte progenitor cells as described by Ansel et al. (1988). Cells in monolayer culture were lysed in 4 M guanidinium isothiocyanate, 5 mM sodium citrate, pH 7.0, 0.1 M β-mercaptoethanol, and 0.1% Sarkosyl; and total RNA was purified by CsCl gradient centrifugation, extracted in 4:1 chloroform/butanol, and precipitated in ethanol. The RNA precipitate was resuspended in RNase-m, mercaptoethanol, and 0.1% Sarkosyl, and then RNA was purified by CsCl gradient centrifugation, extracted in 4:1 chloroform/butanol, and precipitated in ethanol. The RNA precipitate was resuspended in RNase-free water and quantitated by spectrophotometry. Total RNA (10 µg) was separated on a 1.5% agarose gel containing 2.2 M formaldehyde, and in transplanted cells in vivo, using biotinylated anti-NPTII (5 Prime–3 Prime Inc., Boulder, CO). Cultured cells were fixed in acid:alcohol (5:95) at −20°C for 5 min, followed by incubation with anti-NPTII antibody (1:100 in PBS) for 2 hr at room temperature (RT). Cryostat sections of paraformaldehyde-fixed tissue samples (grafted brain) were permeabilized with 0.5% Triton X-100 in PBS for 5 min and then incubated first with 10% normal goat serum for 30 min at RT followed by anti-NPTII antibody (1:100 in PBS) overnight at 4°C. The sections were rinsed and incubated in ABC reagent according to the manufacturer’s instructions (Vectorstain Elite kit; Vector Laboratories), and NPTII immunoreactivity was detected with the DAB substrate kit (Vector Laboratories).

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RESULTS

Inactivation of FGF signaling in oligodendrocyte progenitors expressing a truncated FGFR1 transgene

To disrupt FGF signaling in oligodendrocyte progenitor cells, we introduced a recombinant plasmid encoding a modified version of the wild-type murine FGFR1 (Fig. 1A) into cultured oligodendrocyte progenitors by DNA-mediated gene transfer (McKinnon and Zazapis, 1996). The cDNA insert of this construct contains a stop codon introduced at amino acid position 418 of the fgfr1 sequence and encodes a truncated form of the receptor (denoted FGFRx) that lacks the cytoplasmic signaling domain of the protein. When expressed in transfected cells, the truncated FGFRx receptor inhibits signal transduction by multiple types of endogenous wild-type FGFRs (Ueno et al., 1992; Benvenisty and Ornitz, 1995) in a dominant-negative manner (Herskowitz, 1987). This expression vector also encodes the NPTII gene (Fig. 1A), which confers resistance to the antibiotic geneticin (G418) and provides both a selectable (Southern and Berg, 1982) and immunohistochemically detectable marker for cells that express the transgene. Both transgenes are expressed on a bicistronic RNA transcript linked by the IRES, with NPTII encoded downstream of the soma. Scale bar, 25 µm. C, 125I-FGF binding proteins present in extracts of nontransfected parental progenitor cells (lane 1; wt) and progenitor cells transfected with pMoFRx (lane 2; FRx). Sizes of electrophoretic markers are indicated in kilodaltons, and arrows indicate a prominent FGF-binding protein migrating at the expected size for wild-type FGFR in both lanes and a protein with the predicted size of FGFRx (FRx; 100 kDa) in transfected cells.

Figure 1. Expression of a dominant-negative FGF receptor transgene in oligodendrocyte progenitor cells. A, Schematic representation of transfection vector pMo.FGFRx.iresNeo (pMoFRx), encoding (left to right) the Moloney LTR transcriptional regulatory sequence (Mo-LTR; arrow indicates polarity of transcription), a truncated form of the murine FGFR1 receptor (mFGFRx), internal ribosome entry sequence (IRES), and the neomycin phosphotransferase (NPTII) gene. The vertical arrow indicates the location of a stop codon introduced in FGFR1 by XbaI linker mutagenesis (Benvenisty and Ornitz, 1995) and causing premature termination of translation 22 amino acids after the transmembrane domain (tm) and upstream from the cytoplasmic tyrosine kinase (tk) domain. B, Anti-NPTII immunoreactivity in oligodendrocyte progenitor cells transfected with pMoFRx. Transfected cells had prominent staining of the soma. Scale bar, 25 µm. C, 125I-FGF binding proteins present in extracts of nontransfected parental progenitor cells (lane 1; wt) and progenitor cells transfected with pMoFRx (lane 2; FRx). Sizes of electrophoretic markers are indicated in kilodaltons, and arrows indicate a prominent FGF-binding protein migrating at the expected size for wild-type FGFR in both lanes and a protein with the predicted size of FGFRx (FRx; 100 kDa) in transfected cells.
including the wild-type FGFR1 receptor, PDGFα, or fms/ PDGFα chimera or with the truncated FGFRx transgene and clonally selected in medium containing G418. The fms/PDGFα chimera transgene encodes the ectodomain of human CSF1 receptor, which recognizes human but not rodent CSF1 and therefore would not be expected to respond to endogenous ligand when transfected cells are transplanted into the rodent CNS. Individual transformants from independent transfection experiments were isolated and expanded as independent cell strains (O2A FRx, O2A fms, and O2A FRx cells, respectively). The frequency of G418 resistant colonies obtained with the FGFRx construct was comparable with that obtained with control constructs (Table 1), indicating that the presence of transgenes encoding the truncated FGFRx did not impair the ability of oligodendrocyte progenitor cells to survive or proliferate under the culture conditions used in this study. Analysis of one clonal derivative (O2A FRx subclone cells) demonstrated the presence of both fgfr.nptII transcripts (data not shown), and NPTII immunoreactivity (Fig. 1B), and cross-linking studies (Fig. 1C) revealed an FGF-binding protein of M, 100 kDa, expected for the truncated FGFRx receptor, that was expressed at a level at least fivefold higher than that of wild-type receptors. Additional bands detected in this analysis (Fig. 1C, lane 2) may represent multimeric forms of FGF-binding proteins.

To characterize the transfectants further, we compared the phenotypes of untransfected “parental” and O2A FRx progenitor cells under different culture conditions in vitro. When maintained in the presence of B104-conditioned medium, both parental (data not shown) and O2A FRx (Fig. 1B) cells were multipolar with thin branched processes and were immunopositive with antibodies A2B5 (50.5 and 74.1%, O2A FRx, O2A fms, and O2A FRx cells, respectively).

Table 1. Transfection of oligodendrocyte progenitor cells with pMo.FGFRx.iresNeo constructs

<table>
<thead>
<tr>
<th>Vector</th>
<th>No. of colonies</th>
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<tr>
<td>no DNA</td>
<td>0</td>
</tr>
<tr>
<td>PDGFα</td>
<td>&gt;76</td>
</tr>
<tr>
<td>FGFR1</td>
<td>56</td>
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<tr>
<td>FGFRx</td>
<td>58</td>
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The frequency of G418 colonies obtained after transfection of oligodendrocyte progenitor cells with an expression vector pMo.FGFRx.iresNeo (Fig. 1) encoding cDNA versions of human PDGFα, murine FGFR1, or a truncated (dominant-negative) version, FGFRx. Values represent the number of colonies/2 × 10⁵ transfected cells/5 μg of plasmid DNA.

dependent increase in [³H]thymidine incorporation in response to PDGF-AA (Fig. 2A). Parental O-2A progenitor cells responded maximally with <5 ng/ml PDGF, whereas two independently derived O2A FRx lines required somewhat higher (5–10 ng/ml) PDGF concentrations for maximal response. Wild-type oligodendrocyte progenitors also responded to FGF-2 (Fig. 2B), with maximal response at 1–2 ng/ml. In contrast, O2A FRx cells failed to respond to FGF-2 at concentrations ranging from 1 to 25 ng/ml (Fig. 2B). One clonal derivative (O2A FRx subclone cells) responded weakly to FGF-2 at much higher (>50 μg/ml) concentrations, whereas a second subclone (O2A FRx subclone) was nonresponsive at this higher dose. This pattern of growth factor-induced thymidine incorporation was observed in four independently isolated O2A FRx clones tested, suggesting that impaired FGF signaling was not caused by random clonal variation and that the truncated FGFRx transgene acts in a dominant-negative manner to interrupt FGF signaling in these cells. These observations also imply that FGF signaling is not essential for the proliferation of oligodendrocyte progenitor cells under the culture conditions used (in the presence of B104-conditioned medium).

Impaired migration of oligodendrocyte progenitors expressing a truncated FGFR1 transgene in vitro

FGF maintains oligodendrocyte progenitors at the O4-immunoreactive progenitor cell stage (Pfeiffer et al., 1993) and increases PDGF expression levels (McKinnon et al., 1990), thereby modulating the ability of progenitor cells to respond to PDGF (McKinnon et al., 1993a). Consequently, in addition to its inhibition of FGF signaling, overexpression of the dominant-negative FGFRx transgene would be predicted to alter the response of these cells to PDGF. In two subclones examined, O2A FRx cells showed a slight decrease in response to mitogenic stimulation by PDGF relative to control cells (Fig. 2A), suggesting a modulating effect of the FGFRx transgene on PDGF signaling. Because PDGF can stimulate cell migration (Armstrong et al., 1990; Milner et al., 1997) as well as proliferation, we also compared the ability of parental and O2A FRx cells to migrate in vitro (Fig. 3). Cells plated on coverslips were cultured in the

Figure 2. Impaired FGF signaling in oligodendrocyte progenitor cells expressing the truncated mFGFRx receptor. A, B. [³H]Thymidine incorporation in nontransfected parental oligodendrocyte progenitor cells (squares) and two independently isolated, clonally derived strains transfected with pMoFGRx (triangles, circles). Cells were cultured for 20 hr in the presence of the indicated concentrations of PDGF-AA (A) or FGF-2 (B), and results represent the mean ± SD of triplicate samples and are representative of a minimum of three independent assays. All cell strains responded to physiological levels of PDGF (5 ng/ml), although transfected cells were nonresponsive at physiological levels of FGF-2 (2–5 ng/ml).
presence of FGF plus PDGF, conditions under which parental cells revert from stellate progenitors to bipolar, migratory progenitors (McKinnon et al., 1993a). Under these conditions, parental cells migrated in a radial array from the coverslip onto the culture dish surface (Fig. 3, top). O2A FRx cells, in contrast, did not migrate away from the coverslip (Fig. 3, bottom). When monitored by time lapse cinematography, O2A FRx cells were also nonmotile compared with control cells over a 16 hr time period (data not shown). The lack of migration of O2A FRx cells was unlikely to be a result of the clonal selection process that generated these cells, because O-2A progenitor cells transfected with the chimeric fms/PDGFRα construct (Yu et al., 1994) migrated in response both to PDGF and to recombinant human CSF1 (via the transgene receptor) under similar conditions in vitro (S. Ebner, unpublished observations). Because the migration of O-2A progenitors in vitro is stimulated by PDGF (Armstrong et al., 1990), these results further suggest that expression of the FGFRx transgene modulated the ability of transfected cells to respond to PDGF in vitro.

Impaired migration of oligodendrocyte progenitors expressing a truncated FGFR1 transgene in vivo

The behavior of O2A FRx progenitor cells in vivo was examined by transplanting into neonatal rat brain and examining their distribution at various times after transplantation. The fate of O2A FRx cells (Figs. 4A, 5) was compared with that of wild-type (parental) oligodendrocyte progenitors (Fig. 4B) and with that of progenitors transfected with control cDNA constructs encoding wild-type versions of FGFR1, PDGFRα, and a fms/PDGFRα chimera (see Fig. 6) that is inactive in rodent brain. All cells were first labeled in vitro with fluorescent markers, either PKH2 or PKH26, and then injected into the telencephalon (thalamus) of postnatal day 4 rat brain (rostral, top) 48 hr after transplantation with parental progenitors (PKH2 label, green fluorescence) plus pMoFRx-transfected progenitors (PKH26 label, red fluorescence). A, Retention of distinct fluorescent markers by cotransplanted wild-type and mutant cells. Transplanted cells located within the lateral ventricle (LV) specifically retained cell surface-associated PKH2 or PKH26 fluorescent markers. B, Migration of parental oligodendrocyte progenitor cells in host CNS. PKH2-labeled (wild-type) progenitor cells were found distal to the site of injection, entering the corpus callosum rostral to the LV. The fluorescence label was predominantly in the cell soma, whereas processes extended in situ in the presumed leading edge of migration (arrows) were less intensely labeled. Scale bar, 35 μm.

Figure 3. Impaired migration of oligodendrocyte progenitor cells expressing the truncated mFGFRx receptor. Photomicrographs of parental oligodendrocyte progenitor cells (top) and progenitor cells transfected with pMoFRx (bottom) that were cultured for 72 hr in the presence of PDGF plus FGF-2 are shown. Parental progenitor cells showed extensive radial migration, whereas FGFRx cells did not. Short-term time lapse cinematography also indicated that transfected cells did not migrate (see Results). Scale bar, 650 μm.

Figure 4. Cotransplantation of fluorescent-labeled oligodendrocyte progenitor cells. Horizontal sections of postnatal day 4 rat brain (rostral, top) 48 hr after transplantation with parental progenitors (PKH2 label, green fluorescence) plus pMoFRx-transfected progenitors (PKH26 label, red fluorescence). A, Retention of distinct fluorescent markers by cotransplanted wild-type and mutant cells. Transplanted cells located within the lateral ventricle (LV) specifically retained cell surface-associated PKH2 or PKH26 fluorescent markers. B, Migration of parental oligodendrocyte progenitor cells in host CNS. PKH2-labeled (wild-type) progenitor cells were found distal to the site of injection, entering the corpus callosum rostral to the LV. The fluorescence label was predominantly in the cell soma, whereas processes extended in situ in the presumed leading edge of migration (arrows) were less intensely labeled. Scale bar, 35 μm.
2 recipient rats. The fluorescent cell markers were retained in the plasma membranes of labeled cells both in vitro and in vivo throughout the time points examined in this study (7 d) and had no demonstrable effects on the viability of labeled cells in vitro. The labeling became somewhat punctate at longer time points, with processes extended in situ having less label than the cell soma. The injection path was identifiable 24 hr after transplantation by a needle track penetrating the tissue (see Fig. 5A). The transplantation produced a gliotic reaction around this track, detected after 72 hr by immunoreactivity of glial fibrillary acidic protein (data not shown). When visualized under fluorescence after 24 hr, transplanted cells were found along this needle tract and within a small cyst at the site of transplant. This distribution of grafted cells was typical of all cells examined at 24 hr after transplantation in this study.

The behavior of parental and O2A<sub>FRx</sub> progenitor cells was directly compared by cotransplanting both cell types into the same host (Fig. 4A), after mixing cells labeled in vitro with the distinct fluorescence markers PKH2 (parental) and PKH26 (O2A<sub>FRx</sub>). At 48 hr after transplantation, many PKH2-labeled parental cells had moved from the site of injection and were observed both within the lateral ventricle (Fig. 4A) and within axonal tracts including the corpus callosum (Fig. 4B). In contrast, PKH26-labeled O2A<sub>FRx</sub> cells remained either at the site of injection or within the neighboring ventricles (Fig. 4A). A similar distribution was seen at 7 d after transplantation. The fluorescent cell markers specifically reported the location of individual grafted cells, and not phagocytic cells, because the fluorescent dyes were cell surface associated and mutually exclusive (Fig. 4A). Because control cells migrated under these conditions, the lack of migration of O2A<sub>FRx</sub> cells was not attributable to some aspect of the transplantation procedure. The cotransplantation studies thus demonstrated a specific lack of migration of O2A<sub>FRx</sub> cells into brain after transplantation.

The distribution of transplanted O2A<sub>FRx</sub> mutant cells in vivo was distinct from that of control transplants, including both nontransfected parental cells and progenitor cells transfected with control cDNA constructs. At all time points examined, O2A<sub>FRx</sub> cells were located primarily along the surface of the ventricles (Fig. 5B,C) and adjacent to the choroid plexus (Fig. 5D) and were rarely found within the host brain parenchyma. This distribution pattern was visible as early as 48 hr after transplantation, did not change over the time points examined (up to 7 d), and was observed with two independently isolated O2A<sub>FRx</sub> clonal lines. At 72 hr after transplantation, the morphology of O2A<sub>FRx</sub> cells resembled that of the parental transplanted cells, extending short processes and interacting with adjacent cells in the choroid plexus and ependymal layer of the ventricles (Fig. 5C, D). The inability to migrate was specifically associated with the dominant-negative FGFRx transgene (see below), suggesting that impaired FGFR signaling may be responsible for their failure to migrate in vivo.

**Migration of control oligodendrocyte progenitor cells in vivo**

In contrast to O2A<sub>FRx</sub> cells, the distribution of PKH2-labeled parental cells changed significantly after transplantation (Fig. 4B). After 48 hr, O2A<sup>WT</sup> cells were dispersed in brain parenchyma, and after 72 hr, grafted cells could be found within axonal fiber tracts including the corpus callosum (Fig. 4B), internal capsule, and hippocampal fimbria. Parental cells were also observed along the needle tract and at the site of injection, although...
the number of these cells declined over time (see below). A similar distribution of control transfected cells was observed with transplants of O2A<sup>imm</sup> (Fig. 6), O2A<sup>FR</sup><sub>x</sub> (Fig. 7), and O2A<sup>FR1</sup> (data not shown) cells. The majority of cells that had moved away from the injection site extended several processes within host brain parenchyma (Figs. 4B, 6C). The age of the host at the time of transplantation (postnatal day 2 or 6) had no apparent effect on the distribution of grafted cells within the brain. Because the expression of other transgenes had no apparent effect on the dispersion and distribution of transplanted cells in the brain, the lack of migration observed with O2A<sup>FR</sup><sub>x</sub> cells was likely because of the presence of the dominant-negative FGFRx transgene.

**Maintenance of transgene expression in vivo**

Maintenance of the dominant-negative effects of FGFRx would require the stable and continued expression of the fgfr<sub>x</sub>.nptII transgene in transplanted O2A<sup>FR</sup><sub>x</sub> cells in the absence of G418 selection in vivo. When cultured for 7 d in the absence of the G418 in vitro, O2A<sup>FR</sup><sub>x</sub> cells continued to maintain high levels of NPTII immunoreactivity, indicating that the transgene was stably integrated and expressed in these cells. These cells also stably expressed the FGFRx transgene in vitro, as determined both by steady-state RNA blot analysis and ligand cross-linking analysis (Fig. 1C). Because reagents to specifically detect murine FGFR1 in rat brain are unavailable, we used anti-NPTII immunohistochemistry to examine transgene expression in O2A<sup>FR</sup><sub>x</sub> cells. In the expression vector used in this study, the fgfr<sub>x</sub>.nptII transgene is bicistronic with and encoded upstream from nptII (Fig. 1A); thus, expression of NPTII from the single promoter in this vector ensures coexpression of FGFRx. At 7 d after transplantation, PKH26-labeled O2A<sup>FR</sup><sub>x</sub> cells were identifiable with anti-NPTII antibodies directly lining the ventricles, adjacent to the choroid plexus, and in the ependymal layer as determined by immunohistochemistry (Fig. 5D). Thus the fgfr<sub>x</sub>.nptII transgene expression was maintained in transplanted O2A<sup>FR</sup><sub>x</sub> cells in vivo, suggesting that the phenotype associated with the dominant-negative FGFRx transgene persisted in these grafted cells over the time frame analyzed in this study.

**Quantitative analysis of transplanted oligodendrocyte progenitor cells**

The relative distribution of transplanted O2A<sup>FR</sup><sub>x</sub> cells was also examined using immunohistochemical analysis of NPTII, at both 2 and 7 d after transplantation (Fig. 7). Serial sections from brains that received either of two independent O2A<sup>FR</sup><sub>x</sub> lines were analyzed. After 2 d, a significant number (21.5 ± 0.7%) of O2A<sup>FR</sup><sub>x</sub> cells remained at the site of injection (572 of 2664 cells counted), whereas after 7 d relatively fewer (6.0 ± 8.5%) were found at the injection site (45 of 1077 cells counted). At both time points examined, <10% of NPTII-immunoreactive O2A<sup>FR</sup><sub>x</sub> cells were found within the brain parenchyma at a minimum distance of at least two cell diameters from the ventricular space (Fig. 7). The majority of O2A<sup>FR</sup><sub>x</sub> cells identified by NPTII staining were located within the ventricles, accounting for 58 ± 6% (1521 of 2664 cells counted) and 84 ± 11% (930 of 1077 cell counted) at 2 and 7 d after transplantation, respectively (Fig. 7). The relative increase in O2A<sup>FR</sup><sub>x</sub> cells within ventricles from 2 to 7 d may represent a selective loss of NPTII-immunoreactive cells at the injection site; when these cells were excluded from the analysis, the proportion of O2A<sup>FR</sup><sub>x</sub> cells within ventricles at the 2 d time point represented 73% of NPTII-immunoreactive cells counted, comparable with that found after 7 d (Fig. 7). The relatively small decrease in O2A<sup>FR</sup><sub>x</sub> cells located within the parenchyma, from day 2 (10 ± 2.8%) to day 7 (6.0 ± 1.4%) after transplantation, suggested that there was no appreciable difference in the survival of cells located in these separate compartments. This analysis suggests that the primary defect of O2A<sup>FR</sup><sub>x</sub> cells is a lack of migration into brain parenchyma.

This distribution of transplanted O2A<sup>FR</sup><sub>x</sub> cells was in sharp contrast to the behavior of other populations of NPTII-immunoreactive oligodendrocyte progenitor cells (O2A<sup>FR1</sup>, O2A<sup>FR</sup><sub>x</sub>). The behavior of control transfectants (Fig. 7) paralleled that of the untransfected parental progenitors. At 48 hr, a significant proportion of NPTII-immunoreactive O2A<sup>FR</sup><sub>x</sub> cells (34%) had migrated away from the injection site into axonal tracts, whereas relatively few (13%) remained within the ventricles (93 of 723 NPTII-immunoreactive cells counted). The comparable pattern of migration observed with control transfectants and parental progenitor cells indicated that the clonal selection process used to generate oligodendrocyte progenitor cells in vitro did not affect their ability to migrate in vivo. Thus, the specific lack of migration of O2A<sup>FR</sup><sub>x</sub> cells was likely because of the overexpression of the truncated FGF receptor and suggests this is a result of the disruption of FGF signaling in these cells.

**DISCUSSION**

Fibroblast growth factor is one of a large number of polypeptide growth factors that, either individually or in combination, affect the proliferation, migration, survival, and differentiation of oligodendrocyte progenitor cells in vitro (McMorris and McKinnon, 1996). FGF-2 is both a mitogen (Bogler et al., 1990; McKinnon et al., 1990) and survival factor for oligodendrocyte progenitors...
(Yasuda et al., 1995) and induces mature oligodendrocytes to dedifferentiate and re-enter the cell cycle (Fressinaud et al., 1993; Grinspan et al., 1993; Muir and Compston, 1996). The present study demonstrates that a principal requirement of FGF signaling in vivo is during the migration of oligodendrocyte progenitor cells. By introducing a dominant-negative version of the murine FGF receptor FGFR1 into oligodendrocyte progenitors, clonally derived cells were generated that were unable to respond to FGF-2 in vitro. When transplanted into neonatal rodent brain, these cells failed to migrate into brain parenchyma. The failure of mutant cells to migrate was independent of clonal selection and was specific to cells expressing the dominant-negative FGF receptor transgene. Our results are therefore consistent with a model in which FGF signaling is required for oligodendrocyte progenitors to acquire a migratory-competent phenotype in vivo.

At the time of transplantation in this study, endogenous oligodendrocyte progenitors are migrating from the subventricular zone into brain parenchyma (Altman, 1966; Paterson et al., 1973; Levine and Goldman, 1988; Hardy and Reynolds, 1991). When grafted into this environment, wild-type oligodendrocyte progenitor cells are able to respond to local environmental cues including signals for migration and maturation into differentiated oligodendrocytes (Duncan and Milward, 1995; Franklin and Blakemore, 1995). Because mutant cells would be subject to these same environmental cues after transplantation, their phenotype reflects a cell autonomous defect. The inability of O2A FRx cells to migrate under these conditions (Figs. 4, 5) implies a role for an FGF ligand–receptor signaling pathway in oligodendrocyte progenitor cell migration in vivo. This is in agreement with recent observations describing a role for FGF in progenitor cell migration in vitro (Milner et al., 1997). These findings are also consistent with the observation that the FGF receptor homolog homolg breathless is essential for migration of specific midline glial cells in Drosophila (Klambt et al., 1992). FGF has not been implicated previously in the chemotactic migration of oligodendrocyte progenitor cells in vitro (Armstrong et al., 1990) but has been shown to prime these cells to respond to PDGF (McKinnon et al., 1993a). Our results thus illustrate the benefits of investigating signal transduction pathways under in vivo conditions, in which the complexity and interplay of biological responses can reveal functions that are not uncovered by in vitro analyses.

The cells examined in this study were amplified in vitro in the presence of mitogens from neuroblastoma B104-conditioned medium. Although this system is accessible to molecular genetic manipulations, it is also subject to the limits of an in vitro approach including the potential to either select for or generate immortalized cells. Because the cells we examined were isolated from neonatal rodent brain, they would be expected to have a longevity somewhat in excess of the 50 ± 10 doublings defined for presenescent adult human fibroblasts (Hayflick, 1965). In the present studies, we have focused on cultures that have been expanded for <15 passages (1:3 split ratio) and that are able to differentiate into mature oligodendrocytes both after mitogen withdrawal in vitro and after transplantation in vivo. We have found that at later passages these cell populations can acquire both a mitogen independence for cell proliferation and the inability to differentiate into mature oligodendrocytes after removal of mitogens. However, the early passage cell strains examined in this study did not exhibit these characteristics in vitro and did not have a high mitotic index in vivo, because <3% of PKH26-labeled cells could be detected by immunohistochemistry with antibodies to BrdU. The cells examined in this study thus do not meet the criteria for immortalized or tumorigenic cell lines.

The most straightforward interpretation of this study is that oligodendrocyte progenitor cells expressing a dominant-negative FGFR1 transgene are defective in cell migration. This study does not entirely exclude a possible defect in the survival of O2A FRx mutant cells that have migrated into the host tissue. It is possible, for example, that O2A FRx cells persist within the ventricles because the choroid plexus provides sufficient quantities of factors such as IGF (Stylianopoulou et al., 1988), a potent survival factor for cells of this lineage (Barres et al., 1992, 1993). O2A FRx cells that migrated into tissue, in contrast, may fail to survive because of their inability to respond to FGF, which promotes oligodendrocyte survival in vitro (Yasuda et al., 1995). However, several observations suggest that the primary defect of O2A FRx cells may not be cell survival. First, the ability to expand mutant oligodendrocyte progenitor cells lacking FGF signal transduction indicates that FGF signaling is not essential for the survival of these cells in vitro. Second, the cotransplantation of parental and mutant cells revealed that O2A FRx mutants persisted for times comparable with that of wild-type progenitors in vivo (Fig. 4A). Third, a quantitative analysis did not reveal a selective loss of transplanted O2A FRx cells located within either ventricles or tissue (Fig. 7). These observations suggest that O2A FRx cells are not defective for cell survival and are thus consistent with the interpretation that the dominant-negative FGFRx transgene affects the ability of oligodendrocyte progenitor cells to migrate in vivo.

The mechanism by which the FGFRx transgene may affect the migration in vivo is not known. The truncated form of FGFR1 acts in a dominant-negative manner to interrupt FGF signaling in these cells (Fig. 2B), implicating altered signal transduction as the primary cause. Oligodendrocyte lineage cells express several forms of FGFRs (Bansal et al., 1996), and because a truncated form of FGFR1 may inhibit signaling by multiple types of fibroblast growth factor receptors (Ueno et al., 1992), O2A FRx cells should be nonresponsive to any of the FGFs expressed in the CNS. Signaling through the FGF receptor has also been implicated in neurite outgrowth stimulated by the adhesion molecules L1, N-CAM, and N-cadherin (Williams et al., 1994; Saffell et al., 1997). Thus, if cell motility is promoted by cell adhesion molecules acting through the FGF receptor, these interactions may be disrupted in O2A FRx cells. Cell migration can also be regulated by interactions of cells with extracellular matrix molecules via cell surface integrins (Reichardt and Tomasselli, 1991). Oligodendrocyte progenitors express a number of integrins, and the pattern of integrin subunit expression changes as progenitors differentiate into mature oligodendrocytes (Milner and Firench-Constat, 1994). Because FGF regulates the expression of other cell surface receptors on oligodendrocyte progenitors (McKinnon et al., 1990; Gallo et al., 1994), it is conceivable that integrin expression is also altered in O2A FRx cells. To date, the role of cell adhesion molecules and/or integrins in the migration of oligodendrocyte progenitor cells has not been described. Finally, although the primary focus of expressing a dominant-negative FGFRx transgene is its effects on FGF signal transduction, other possible effects such as altered nuclear translocation of ligand (Bugler et al., 1991; Mason, 1994) could contribute to the inadequacy of these cells to migrate in vivo.

The expression of PDGF by neurons during CNS development (Sasahara et al., 1991; Yeh et al., 1991) is consistent with a role for neuronal PDGF in promoting the migration of oligodendro-
cyte progenitors along axonal tracts in vivo. Although FGF does not promote chemotactic migration in vitro (Armstrong et al., 1990), one dramatic effect of FGF is the upregulation of PDGF receptors, leading to an increased sensitivity of these cells to PDGF (McKinnon et al., 1990). Thus, the absence of FGF signaling could affect the ability of these cells to acquire a PDGF-responsive state. Consistent with this, in vitro studies indicate that O2A cells have a slightly decreased mitogenic response to PDGF (Fig. 2A), and initial in vivo analysis indicated that transplanted wild-type cells have a more immature phenotype than do transplanted O2A mutant progenitors (Ebner, unpublished observations). The failure of transplanted O2A progenitors to migrate in vivo could be an indirect result of an impaired response to PDGF, resulting in an inability to adopt the more migratory phenotype in vivo. This model is thus consistent with the observation that transplanted O4-immunoreactive oligodendrocyte progenitors have a decreased migratory ability relative to A2B5-immunoreactive (O-2A) progenitors (Warrington et al., 1993).

The expression of dominant-negative forms of growth factor receptors has proven to be a useful tool for studying the role of growth factor signaling in development. In Xenopus, mesoderm formation is disrupted with dominant-negative forms of both activin (Hemmatti-Brivanlou and Melton, 1992) and FGF receptors (Amaya et al., 1991), and a dominant-negative FGFR also causes errors in Xenopus retinal ganglion cell axonal target recognition (McFarlane et al., 1996). Dominant-negative FGF receptors also affect the development of both keratinocytes (Werner et al., 1993) and cardiac myocytes (Mima et al., 1995). By combining ex vivo gene manipulations with cell transplantation, we have been able to address the role of FGF signaling in mammalian CNS development. This approach offers advantages over direct gene transfer in vivo, such as the delivery of receptor transgenes using retroviral vectors (Lillien, 1995), or in vivo transfection (McFarlane et al., 1996), in that we could confirm the dominant-negative effects of the FGFRx transgene in vitro before the analysis of transplanted cells in vivo. Our finding that transgene expression was maintained in vivo suggests that this approach may also be useful for sustained overexpression of molecules such as growth factors, which could potentially be used to study repair processes in models of neurological diseases including multiple sclerosis. Thus, ex vivo gene transfer into populations of CNS progenitor cells is an attractive method for studying many aspects of their biology.

REFERENCES


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