Targeted Neocortical Cell Death in Adult Mice Guides Migration and Differentiation of Transplanted Embryonic Neurons

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Local expression of cellular and molecular signals is required for normal neuronal migration and differentiation during neocortical development and during periods of plasticity in the adult brain. We have previously shown in neonatal and juvenile mice that induction of apoptotic degeneration in neocortical pyramidal neurons by targeted photolysis provides an altered environment that directs migration and differentiation of transplanted embryonic neurons. Here we employ the same paradigm in adult mice to test whether targeted photolysis induces the reexpression in the mature brain of developmental signals that control migration, differentiation and integration of embryonic neurons. We examined both the time course of migration and the morphologic and immunocytochemical differentiation of embryonic neurons transplanted into regions of targeted photolytic cell death.

Pyramidal neurons in neocortical lamina II/III underwent photolytically induced apoptosis after retrograde incorporation of the photoactive chromophore chlorin e, and transdural exposure to 674 nm near-infrared laser energy. Embryonic day 17 neocortical neurons were prelabeled with fluorescent nanospheres and the lipophilic dye PKH26, transplanted into regions of ongoing neuronal degeneration in adult mice, and examined histologically and immunocytochemically. Transplanted neurons began migration into regions of neuronal death within 3 days and differentiated into large pyramidal neurons similar to those degenerating. In contrast, neurons transplanted into intact cortex did not migrate, and they differentiate into small presumptive interneurons. Migration up to 430 μm in experimental mice was complete by 2 weeks; approximately 45% of the donor neurons migrated greater than 3 SDs beyond the mean for neurons transplanted into intact neocortex of age-matched adult hosts. Following migration, dendrites and axons of many donor neurons were properly oriented toward the pial surface and corpus callosum, indicating integration into the host parenchyma. Neurofilament and neuron-specific enolase staining further support appropriate differentiation and integration.

These results indicate that signals guiding neuronal migration and differentiation in neocortex are reexpressed in adult mice well beyond the period of corticogenesis within regions of targeted photolytic cell death. Elucidating the molecular mechanisms underlying these events by comparison with adjacent unperturbed regions will contribute to efforts toward future therapeutic transplantation and control over endogenous plasticity.

[Key words: adult mouse, apoptosis, differentiation, integration, laser, migration, neocortex, neuronal degeneration, targeted photolysis, transplantation]

Control over neuronal migration and differentiation depends in part upon local expression of neuronal, glial, and extracellular signals during development or during periods of plasticity in the adult (Hyues et al., 1986; Hyues et al., 1989; Ghosh et al., 1990; Good et al., 1991; Ferrari et al., 1993; Goodman and Shatz, 1993). In neocortex, neurogenesis is limited to a brief embryonic period, and neuronal migration continues only during the first postnatal week (Angevine and Sidman, 1961; Austin and Cepko, 1990; Bayer and Altman, 1991). Disappearance of a highly ordered temporal and spatial pattern of molecular and activity-dependent signals during maturation is thought to underlie the relative lack of plasticity observed in the mature mammalian CNS. Understanding the signals controlling initial neuronal migration and differentiation during neocortical development may allow future therapeutic manipulation of these regulatory molecules.

Disruption of cell-cell and cell-matrix interactions by selective photolytic degeneration of neocortical pyramidal neurons with callosal projections produces a local environment that regulates the migration and differentiation of embryonic neocortical neurons transplanted into the neocortex of neonatal or juvenile host mice (Macklis, 1993). Targeted neuronal death is produced in a controlled, synchronous manner in vivo while maintaining complex three-dimensional relationships between cells. This new microenvironment activates migration by donor neurons into only affected regions in lamina II/III, differentiation to replace the degenerated pyramidal neurons, formation of anatomically correct axons and dendrites, and ultrastructural integration with afferent synapse formation. These phenomena were not observed with control transplants into intact cortex or cortex lesioned by kainic acid with resultant necrotic and inflammatory neuronal and glial injury. Additional experiments demonstrated the localization of signal alterations to the regions of neuronal...
Degeneration in vivo can be effected within lamina II/III by targeted photolytic cell death (Macklis, 1993; Madison and Macklis, 1993). We have recently demonstrated that neuronal degeneration initiated following targeted photolysis occurs via apoptosis (Sheen et al., 1992; Sheen and Macklis, 1994). Degeneration results from the photoactivation of retrogradely transported nanospheres containing the chromophore chlorin e6. Chlorin e6 produces the cytotoxic molecule singlet oxygen when excited at near-infrared wavelengths; light at these wavelengths (produced by a specific 674 nm laser and beam controlling optics) penetrates deeply through nervous system tissue without absorption or cellular injury to nontargeted cells. Thus, intermixed neurons, glia, axons, and connective tissue are entirely spared, and only neurons selectively labeled with the chromophore undergo degeneration which is progressive and noninflammatory. Nanospheres containing chlorin e6 are localized to lysosomal granules, and photolytic activation of singlet oxygen leads to lipid peroxidation of lysosomal membranes and release of proteolytic contents. The cell undergoes an internal autolysis by singlet oxygen and proteases, leading to a slowly progressive cell death with a loss of calcium homeostasis through L-type calcium channel activation and an increase in membrane permeability. Disruption of cytoplasmic proteins and cytoskeletal components with preservation of membrane bound organelles is accompanied by the appearance of apoptotic membrane bodies and internucleosomal DNA fragmentation. As with other initiating events leading to apoptosis, these targeted cells degenerate and are phagocyted in vivo without the inflammation or scar formation observed with necrotic cell death. It is possible that the slow, non-necrotic process of targeted neuronal cell death following photolytic activation of singlet oxygen in vivo produces many of the same physiological cues that are activated by programmed cell death during normal development and during organizational refinement in the adult vertebrate nervous system. The similarity in the mechanisms of cell death may underlie similar compensatory mechanisms resulting from a loss of cell-cell and/or cell-matrix interactions.

In the mammalian CNS, plasticity and developmental signals influencing neuronal migration and differentiation are thought to decrease with maturity. The age of the 2 week old host mice used in our prior experiments was chosen to be beyond the period of normal neuronal migration, which ends by 5–10 d postnatally, but still during a period of relative plasticity observed in juvenile animals. Recent experiments using this transplantation paradigm with a clonal donor population demonstrated that the phenotypic fate of multipotent precursor cells placed into regions of targeted photolytic neuron death is influenced by the alteration of microenvironmental signals such that differentiation into pyramidal neurons occurs in postnatal experimental mice, but no neuronal differentiation occurs in intact mice or mice with nonspecific kainic acid lesions (Macklis et al., 1994; E. Y. Snyder, C. H. Yoon, J. D. Flax, and J. D. Macklis, unpublished observations). Together, these results suggest that developmental signals in the microenvironment have been either upregulated or disinhibited to effect both neuronal migration and directed differentiation. Study of the temporal and spatial patterns of signal alterations within these regions of targeted apoptosis would be most productive within mature hosts in whom normal developmental signal patterns have long disappeared.

The present experiments involve an examination in adult mice of the time course of migration and the differentiation and integration of embryonic day 17 (E17) neurons transplanted through regions of targeted photolytic neuronal degeneration. These experiments evaluate whether the developmentally appropriate events seen in juvenile mice (Macklis, 1993) can occur in mature mice in whom decreased plasticity might be expected, allowing examination of mechanisms underlying neocortical migration and differentiation in vivo. We provide evidence at three levels of analysis that the directed migration and differentiation of transplanted embryonic neurons within regions of targeted photolytic cell death in neocortex also occur in adults. We observed the initiation of cellular migration by 3 d after transplantation and the repopulation of regions following photolytic degeneration in the adult hosts. Neurons transplanted within regions of neuron deficiency assumed morphologies and immunocytochemical staining characteristic of the previously targeted neocortical pyramidal neurons, often displaying appropriately radially oriented axons and dendrites. These findings suggest that signals regulating neocortical development may be reexpressed in a temporally and spatially correct manner within the regions of targeted photolytic cell death in fully adult mice in which most developmentally regulated molecules are normally downregulated.

A preliminary report of some of these experiments has been presented previously (Macklis and Sheen, 1992).

Materials and Methods

This study is based on data from 156 adult C57B/6J mice. Fifty-nine host mice and 14 separate embryonic dissections were used for studies of neuronal migration and differentiation after transplantation. Each dissection contained cells from six to nine mouse embryos at E17. Sixty mice were used for analysis of neuronal targeting and degeneration in experimental hosts. An additional 37 mice were used to assess neuronal degeneration by silver staining and DNA end terminal labeling following initiation of targeted cell death by noninvasive exposure to laser energy.

The experimental paradigm (Fig. 1a) requires targeting of callosally projecting pyramidal neurons by unilateral injection of fluorescent latex nanospheres conjugated with chlorin e6 into P1 to P16 host mice. After 2 weeks to allow for retrograde transport of latex nanospheres, neuronal degeneration was initiated in the experimental mice by targeted laser photolysis. Cellular transplantation, using dissociated cell suspensions of embryonic neocortex labeled with rhodamine nanospheres that label neurons relatively selectively (Macklis et al., 1990; Macklis, 1993) was performed 2 weeks later in 4 or 6 week old experimental mice and in age-matched intact controls. Targeted neurons underwent a gradual and progressive degeneration extending over several weeks such that neurons in the process of cell death are still present at the time of transplantation within the region of cortex exposed to laser energy. Initial analysis was performed on littermate mice 4–5 d following cortical injection of nanospheres to ensure successful transport to pyramidal neurons in contralateral cortex. One day to 2 weeks following targeted laser photolysis, other mice were analyzed to ensure induction of neuronal degeneration. Experimental mice underwent light microscopic (LM) analysis of serial sections alternately processed for fluorescence and routine histology. Intact control adult hosts underwent LM analysis by fluorescence and routine histology 1 week (n = 7), 3 weeks (n = 2), and 4 weeks (n = 4) after transplantation. Intact control adult hosts underwent LM analysis by fluorescence and routine histology 1 week (n = 6), 2 weeks (n = 7), 3 weeks (n = 2), and 4 weeks (n = 4) following transplantation of E17 neurons.
Figure 1. Experimental paradigm for transplantation of embryonic neocortical neurons into regions of targeted photolytic neuronal death in adult mice. a, Schematic representation of the sequential experimental steps. The photoactive chromophore chlorin e₆ is injected unilaterally into motor cortex of host mice. After uptake by axonal terminals and retrograde labeling of callosally projecting pyramidal neurons (dotted pyramids), mice are transdurally exposed to near-infrared 674 nm laser energy via optics that delimit the geographic region of uniform exposure. Photoactivation of chlorin e₆ initiates a progressive, apoptotic degeneration within labeled neurons in lamina II/III (dotted, irregular pyramids). Embryonic donor neurons (solid circles) with multiple fluorescent labels are transplanted across the thickness of neocortex. The extent of donor neuron migration and differentiation is assessed histologically and immunocytochemically from 3 d to 6 weeks following transplantation. b, Appearance under fluorescein fluorescence of retrogradely labeled pyramidal neurons in lamina II/III of motor cortex 3 d after injection of nanospheres carrying chlorin e₆. c, Phase contrast photomicrograph of the same field, with neocortical laminae indicated. d, Low power fluorescence photomicrograph of neocortex 4 weeks following laser exposure reveals by the loss of neuronal fluorescence that most of the labeled neurons within regions of targeted neuronal degeneration (arrowheads) have undergone cell death and phagocytosis. Remaining fluorescence is often observed within phagocytic microglia, most frequently in the corpus callosum (cc) and occasionally within the parenchyma of neocortex. Regions not exposed to light activation (open arrow) are not affected. e and f, Higher magnification fluorescence and brightfield photomicrographs of microglia (arrows)
Additional control transplants into host mice receiving conjugated nanosphere injections, but no laser exposure, were similarly examined 2 weeks after transplantation \((n = 3)\).

**Targeting of callosal projection neurons.** Degeneration of neuronal subpopulations in vivo requires labeling with the photoactive chromophore chlorin \(e_6\) that generates singlet oxygen upon noninvasive exposure to near-infrared 674 nm light. Retrograde labeling of contralateral callosal projection neurons with chlorin \(e_6\) was performed with slight modifications of previously described methods (Macklis, 1993). The C3/B6D1 mice (P13–14 or P14-16) were deeply anesthetized through hypothermia or with Avertin, respectively. The skull was exposed by a 5–7 mm dorsal midline incision, and a rectangular craniotomy approximately 2 mm by 4 mm was opened to expose the dura overlying motor cortex. A series of 8–10 microneedles of fluorescein latex nanospheres (Luminaloe) conjugated with chlorin \(e_6\) was performed unilaterally at equally spaced intervals across the length and width of motor cortex to label the callosal neurons. After all injections were performed, the overlying skin was closed with sutures and the pups were returned to maternal care. Consistent with previous results (Macklis and Quatraro, 1991; Macklis, 1993), the latex nanospheres remained localized to the injection site, thereby providing specificity of labeling, and pyramidal neurons in laminae II/III and V were labeled through retrograde transport of nanospheres conjugated with chlorin \(e_6\) (Fig. 1b,c).

**Selective neuronal degeneration.** After 2 weeks to allow for sufficient uptake and retrograde transport of photoactive nanospheres, host mice were deeply anesthetized with Avertin, overlying skin was incised and bilaterally reflected, and the dura was exposed by craniotomy overlying the homologous region of neocortex, contralateral to the initial nanosphere injection site. Cortical illumination by a continuous wave solid state diode laser operating at 674 nm (Candela Laser Corp.) delivered 275–350 J/cm\(^2\) of laser energy over 10–12 minutes via a quartz optical fiber 1 mm in diameter with special optics (Candela Laser) to achieve cross sectional energy uniformity over a 2 mm diameter cortical surface. Coluromation of the laser energy provided specific targeting of pyramidal neurons labeled with nanospheres only in lamina II/III and initiated a slowly progressive neuronal degeneration within this cellular subpopulation (Macklis and Madison, 1985; Madison et al., 1988; Macklis and Madison, 1991; Macklis, 1993). Regions of selective pyramidal neuron degeneration could be defined from the loss of approximately two-thirds of the pyramidal neurons containing fluorescein fluorescence 3–4 weeks following initiation of targeted neuronal death (Fig. 1d), in agreement with previous quantitative results (Macklis, 1993). Non-neuronal cellular fluorescence was also observed within these regions and in the adjacent corpus callosum, localized within microglia which were labeled by the macrophage/macrophage-specific marker F4/80 (Fig. 1e,f).

**In situ detection of neuronal degeneration.** Neuronal degeneration following targeted laser energy. Mouse brain was assessed by both silver degeneration staining, and DNA end terminal labeling. Labeled pyramidal neurons underwent progressive cell death as early as 24 h following exposure to laser energy when assessed in situ for DNA fragmentation, whereas unlabeled pyramidal neurons receiving the same level of laser energy were unaffected (Fig. 2a–e). Silver degeneration staining revealed the same pattern of highly selective pyramidal neuron degeneration within lamina II/III of exposed regions, maximal at 8–10 d following exposure to laser energy in agreement with prior studies (Macklis and Madison, 1993).

Silver degeneration staining was performed by previously described methods (Nadler and Evenson, 1983; Macklis and Macklis, 1993). Experimental mice were assessed histologically 3–14 d following exposure to incident laser energy to assess the onset of neuronal degeneration. Additional control mice receiving laser energy without retrograde labeling by conjugated nanospheres were also processed to confirm prior findings that there is not nonspecific injury due to light alone (Macklis and Macklis, 1993). Cryosections from fresh frozen brains were fixed with 4% paraformaldehyde/0.1 M phosphate buffered saline (PBS, pH 7.4), washed extensively in PBS, and postfixed in ethanolic acid (v/v 2:1). The tissue was washed in PBS and incubated with 15% hydrogen peroxide to inactivate endogenous peroxidases. Sections were then placed in equilibration buffer, reacted with the terminal deoxynucleotise transferase (Tdt) enzyme, and rinsed in stop/wash buffer. After washing with PBS, cells were placed in anti-digoxigenin peroxidase and the complex was visualized with diamobenzidine (DAB) or DAB with cobalt and nickel chloride intensification (Pierce). Positive control samples were pretreated in DNease (1000 U/ml).

Neuronal degeneration was also seen via immunocytochemical staining specific for neurons. Four weeks after initiation of neuronal degeneration, staining for neuron-specific enolase (NSE) revealed that most of the remaining neurons were small, ovoid, presumptive interneurons; only rare neurons had projecting pyramidal neuron morphology. The pattern of neural filament labeling within regions of targeted neuronal degeneration in laminae II/III revealed a geographically defined loss of pyramidal somata and large apical dendrites, further confirming the previously reported (Macklis, 1993; Madison and Macklis, 1993) selective loss of callosally projecting neurons (Fig. 2f).

**Neocortical dissection and labeling.** Isolation and double fluorescent labeling of E17 neocortical neurons followed methods previously reported (Macklis, 1993). Timed E17 pregnant C57/6J mice were cervically dislocated, and embryos were removed and placed in calcium- and magnesium-free Hanks' buffer solution (HBSS-CMF; Gibco), supplemented with glucose (4500 mg/liter), penicillin (24 U/ml), streptomycin (24 mg/ml), and Heps buffer (25 mm) at 4°C. Hemispheres were dissected, dura was removed and the cortex was minced before placing in supplemented HBSS-CMF at 4°C. Cortical tissue was washed twice with supplemented HBSS-CMF by centrifugation at 1500 rpm and aspiration of supernatant. Trypsin solution (0.03% with 0.1% FDTA) was used to dissociate cells by gentle agitation for 30 min at 37°C. Cell suspensions were washed with the supplemented HBSS-CMF, and triturated in growth medium containing calcium- and magnesium-free Dulbecco's modified Eagle's medium (GIBCO), supplemented with glucose (4500 mg/liter), penicillin (24 U/ml), streptomycin (24 mg/ml), and Heps buffer (25 mm), supplemented calf serum (10%, Hyclone) and sodium pyruvate (1 mm) at 37°C. Cells were washed again with the supplemented HBSS-CMF at room temperature, pelleted, and labeled for 30 min with supplemented HBSS-CMF solution containing nanospheres with incorporated rhodamine 6G (12.5 μM) (Madison et al., 1990; gift of Prof. C. Thies, Washington University). These nanospheres provide labeling relatively specific for neurons as identified morphologically by phase-contrast and Nomarski DIC illumination in vitro and in vivo, with neurons displaying distinct punctate granular fluorescence and glia largely unlabeled (Madison et al., 1990; Macklis, 1993). This useful specificity is not essential to interpretation of these studies, as identities of labeled neurons were confirmed by immunocytochemical staining with anti-neuronal degeneration. The cell suspensions were washed three times with the supplemented HBSS-CMF at 4°C, counted during trypan blue exclusion to assure cell viability of greater than 75%, and plated in a sterile vial on ice until use. To assess further the morphology and local projections of donor neurons, cells were prelabelled with the red fluorescent, lipophilic cell linker PKH26-4-L (PKH26, Sigma) using recommended methods. Following transplantation, transplants were placed dorsal to the eye monitored in vitro to assess viability and differentiation of neurons (specifically somatic morphology, neurite extension, and granular compartmentalization), thus ensuring transplantation of viable cells.
Figure 2. Pyramidal neurons in regions of targeted photolysis undergo selective, apoptotic neuronal degeneration. a, Degenerating neurons are stained in situ for DNA fragmentation by 3′-OH end terminal labeling. Apoptotic DNA fragmentation is apparent within lamina II/III 24 hr after laser exposure, and neuronal degeneration continues over 2–3 weeks, restricted to lamina II/III. The boxed region is expanded in b. Neocortical laminae are indicated to the left of the photomicrograph. b, Higher magnification photomicrograph of the boxed field in a, showing pyramidal neurons undergoing apoptotic degeneration (arrowheads). c, Phase contrast photomicrograph of the field in b, indicating the same degenerating pyramidal neurons (arrowheads). d, Photomicrograph of control neocortex 24 hr after exposure to laser energy without targeting by chlorin e6 conjugated nanospheres. Pyramidal neurons do not undergo DNA fragmentation or degeneration, confirming that both chlorin e6 and light activation are necessary to induce cell death. The boxed region is expanded in e. e, Higher magnification photomicrograph of the boxed field in d, showing the absence of DNA fragmentation and degeneration of pyramidal neurons receiving light energy alone. f, Bright-field photomicrograph showing neurofilament staining of neocortex 4 weeks after laser exposure and initiation of targeted neuronal cell death. Loss of callosal projection neurons is further demonstrated by the reduction of neurofilament staining of large cell bodies and apical dendrites within the targeted region of neocortical lamina II/III (open arrows), compared with an immediately adjacent, nontargeted region (curved arrows). Scale bars: a, d, f, 200 μm; b, c, e, 50 μm.

Neuronal transplantation. Experimental and control mice received transplants of E17 neurons at 4 or 6 weeks of age. Cell suspensions (approximately 1 × 10⁴ cells/μl) were slowly pressure-injected into and beneath cortical regions of photolytic degeneration or intact cortex via pulled-glass micropipettes with tip diameters of approximately 100 μm. Micropipettes were lowered directly through the neuron deficient regions to a depth of 500 μm and withdrawn over 250 μm at 50 μm intervals while injecting approximately 50 nl of cell suspension at each level (total 300 nl) to ensure a “tubular” distribution of cells across laminae II/III to V, thereby offering the donor neurons a laminar choice for potential migration and integration. After closing overlying skin, mice were returned to normal care until perfusion for histological analysis.

Histological analysis. We performed histological analysis from 3 d to 6 weeks following the transplantation of neocortical cell suspensions. Mice were deeply anesthetized with Avertin, transcardially perfused with saline and heparin (10,000 U/liter), followed with 2% paraformaldehyde in 0.1 M phosphate-buffered saline (pH 7.3–7.4), and postfixed overnight in the same fixative. The brains were blocked for coronal sectioning through regions of laser exposure and cellular transplantation, and serial sections 40–50 μm thick were cut on a vibratome. Alternate sections were processed and mounted for fluorescence and cresyl violet staining to assess cellular morphology, movement from the radially oriented transplantation site, and extent of cellular integration. Fluorescence sections were air dried on gelatin-coated slides, and subsequently either dipped briefly in Histoclear (National Diagnostics) and permanently mounted in Fluoromount (Gurr), or examined directly mounted in mineral oil. Histologic examination was performed using a microscope equipped with epifluorescence and high numerical aperture optics (Zeiss).

The extent of cellular migration and differentiation was initially assessed by examination under low magnification, then in more detail...
under fluorescence, phase contrast, and Nomarski differential interference contrast (DIC) optics at high magnification to determine neuronal identity and morphology. Highly anionic nanospheres containing rhodamine 6G were preferentially incorporated into lysosomes of neurons, providing a first level of neuronal identification in transplants (Madison et al., 1990). Further criteria of neuronal identification included axonal and dendritic extension, somatic morphology, and immunocytochemical analysis determined under phase contrast and Nomarski DIC optics.

Cellular locations were documented by camera lucida, and neuronal positions were digitized using a video-based image analysis system (Java, Jandel) following previously described methods (Macklis, 1993). Alternate, serial sections from experimental mice underwent quantitative analysis of the degree of migration from the transplantation site 3 days (n = 4), 1 week (n = 2), 2 weeks (n = 6), and 4 weeks (n = 2) after transplantation of dissociated embryonic cell suspensions (n = 4069 labeled donor neurons). Of those experimental mice analyzed 2 weeks following transplantation, n = 2 hosts were transplanted at 4 weeks of age, and n = 4 were transplanted at 6 weeks of age. Positions of transplanted neurons within intact, control mice were digitized from serial sections 1 week (n = 3 mice, n = 6 transplants), 2 weeks (n = 3 mice, n = 5 transplants), and 4 weeks (n = 20) after transplantation to compare cellular dispersion and morphologic differentiation (n = 1282 labeled donor neurons). Additional control transplants assessed positions of donor neurons within host mice receiving conjugated nanosphere injections without laser energy exposure two weeks after transplantation (n = 3 mice; n = 292 labeled donor neurons). These controls evaluated whether injection of photoactive nanospheres or surgical manipulations influenced the migration and differentiation of transplanted immature neurons, reproducing prior controls for these variables (Macklis, 1993).

Statistical analysis. The distributions of neuronal distance from the site of injection could be approximated by curves fit to a one-sided γ function \( \gamma(x, \alpha, n) = \frac{\alpha^n}{\Gamma(n)} x^{n-1} e^{-\alpha x} \) where x distance from transplantation track, and \( \alpha, n \) is optimized. Optima for \( \alpha \) and \( n \) were calculated with the maximum likelihood estimator \( L(\alpha, n) \) for the γ function with \( L(\alpha, n) = \frac{\alpha^n}{\Gamma(n)} \) . \( \alpha \) and \( n \) were obtained by the differentiation of \( L(\alpha, n) \) with respect to \( \alpha \) and \( n \) and determining \( \alpha \) and \( n \) for \( L(\alpha, n) = 0 \) and \( L(\alpha, n) = 0 \) for the respective distributions (B. Nalebuff, personal communication).

Immunocytochemistry. Immunocytochemical staining for neuron specific enolase (NSE; Zymed), neurofilament (NF, pan-neuronal, nonphosphorylated-specific; Sternerberger, and NF pan-neuronal, Zymed), and the macrophage specific marker F4/80 (gift of Dr. A. Ezekowitz) was performed on cryosections of brains fixed in 4% paraformaldehyde and cryopreserved with sucrose, or Vibratome sections of brains fixed with 2% paraformaldehyde. Sections were rinsed in PBS containing 0.25% Triton X-100 and 1% goat serum, incubated in block solution (10% goat serum in PBS), and placed into the primary antibodies (stock NSE, 1:2000 NF, 1:200 F4/80) for 2-3 h at 4°C. The tissue was then washed in blocking solution, and processed according to the peroxidase ABC kit (Vectastain). Neurofilament, NSE, and F4/80 staining were visualized with a DAB substrate alone or a DAB substrate enhanced by nickel and cobalt chloride (Pierce). Donor neurons were identified by rhodamine fluorescent nanosphere labeling within antibody stained tissue sections.

Results

Transplantation of E17 neocortical neurons into intact adult cortex

Donor E17 neurons in intact regions of adult mouse neocortex showed an extremely limited extent of migration and integration 1-4 weeks following transplantation. Donor neurons spanning laminae II/III to V remained localized to the site of injection (Fig. 3a-d). Donor neurons were located at an average distance of 23 ± 18 μm from the transplantation track with a maximal spread of 140 μm. There was no significant change in distribution of donor neurons over time. The distribution of neuronal positions was similar but slightly more restricted than in the previously reported control transplants into juvenile hosts (Macklis, 1993). This minor difference may be explained by ongoing growth of the brain in juvenile hosts. The positions of E17 neurons transplanted into neocortex of mice following injections of photoactive nanospheres without light exposure did not differ significantly from transplants into intact mice (Fig. 3e).

Morphologically, donor neurons in all controls remained small and ovoid with only localized process extension, resembling endogenous interneurons in size and morphology (Fig. 4). Donor neurons could be identified by combined immunocytochemical staining for NSE and labeling with rhodamine fluorescent nanospheres, demonstrating the diminished size and interneuron morphology of transplanted neurons compared with host pyramidal neurons (Fig. 4d,e). Nissl staining further demonstrated the nonpyramidal morphology of donor neurons near the sites of transplantation, compared with host pyramidal neurons further from the injection track (Fig. 4f).

Transplantation of E17 neocortical neurons into regions of adult cortex undergoing targeted neuronal degeneration

Embryonic (E17) donor neurons within the regions of targeted neuronal death in adult mice displayed strikingly different behavior by undergoing directed migration and differentiation 3 d to 2 weeks after transplantation, without further migration observed 4 and 6 weeks after transplantation. Three days to 1 week after transplantation into experimental host mice, clusters of donor embryonic neurons could be observed with elongated morphologies within regions of targeted cell death and in locations midway between the transplantation track and the photolytically targeted regions, suggesting that these neurons were still undergoing an active process of migration at these times (Fig. 5a,b). These included clusters oriented obliquely in deeper laminae IV and V, migrating across these layers to their final locations. One to 2 weeks following transplantation, neurons resided in lamina II/III (Fig. 5c,d). Donor neurons within regions of photolytic degeneration displayed a directed and specific migration into neocortex undergoing targeted cell death over the first 2 weeks (Fig. 6a,b). Donor neurons 3 d after transplantation had a mean distribution of 39 ± 31 μm from the transplantation track. The mean distribution of donor neurons increased to 96 ± 94 μm from the transplantation track one week following transplantation, with 42% of the cells lying beyond 3 SDs of the mean from the control cases (Fig. 6d). After 2 weeks, quantitative analysis of neuronal distance from the transplantation track of experimental mice indicated an average neuronal dispersion of 86 ± 74 μm (Fig. 6e). Neuronal positions 2 weeks after transplantation into 4 week old or 6 week old hosts did not differ significantly by 2 weeks (Fig. 6e). Four weeks after transplantation, donor neurons were positioned 102 ± 93 μm from the transplantation track with a maximal migration of 430 μm perpendicular to the track (Fig. 6f). In comparison with control populations, 45% and 48% of neurons in experimental mice 2 and 4 weeks after transplantation were distributed more than 3 SDs beyond the mean for control distributions. Donor neurons 6 weeks following transplantation displayed a qualitatively similar distribution and reintegration into neuron-deficient regions in lamina II/III.

The experimental distributions obtained for neuronal distance from the site of injection could be approximated by curves fit to a one-sided gamma function \( \gamma(x, \alpha, n) = \frac{\alpha^n}{\Gamma(n)} x^{n-1} e^{-\alpha x} \) where x is distance, \( \alpha, n \) is optimized (B. Nalebuff, personal communication). Both control and experimental cases were best fit to the gamma function \( \gamma(n = 1) \), which approximates an exponential decay. Control cases, 4 weeks following transplantation, exhibited a modified exponential density with mean of 20 μm from the transplantation site (\( \alpha = 0.05 \)), differing significantly from the experimental
Figure 3. Quantitative analysis of donor neuron positions following transplantation into control cortex of adult mice. a, Fluorescence appearance of transplanted neurons within an intact control mouse 2 weeks following transplantation. Donor neurons remain highly confined to the transplantation site (arrows). Corresponding neocortical laminae are indicated to the left. b, Representative camera lucida drawing demonstrates the positions
Figure 4. Nonpyramidal morphology of transplanted neurons within control intact neocortex of adult mouse. a and b, Fluorescence and phase contrast photomicrographs of embryonic day 17 (E17) neocortical neurons 2 weeks following transplantation into intact cortex. Neurons are small in size and show extremely limited dispersion from the transplantation track (arrow). c, Nissl stain of an adjacent section similarly demonstrates small, ovoid donor neurons, spanning laminae II/III to V and remaining highly localized to the site of transplantation. d and e, Higher magnification fluorescence and brightfield photomicrographs of donor neurons stained for neuron-specific enolase. Morphologically, donor neurons (arrows) appear small and ovoid with only localized process extension, resembling endogenous interneurons in size and morphology, compared with much larger, characteristic pyramidal neurons of the host (arrowheads). f, Higher magnification photomicrograph of the Nissl stained section in c shows that the donor neurons (arrows) remain small and ovoid, unlike the endogenous pyramidal neurons of the host (arrowheads). Scale bars: a-c, 200 μm; d-f, 50 μm.

cases, examined 2 or 4 weeks after transplantation, which follow a modified exponential with mean of 100 μm from the transplantation site (α = 0.01). The mean distance migrated by donor neurons following transplantation into regions of targeted cell death was therefore greater than and differed significantly from the mean dispersion for donor neurons transplanted into intact cortex.

Morphology and local projections of transplanted neurons in regions of targeted photolytic neuronal death

Repopulation of the neocortex was limited to regions of photolytic degeneration within lamina II/III, with many transplanted neurons displaying extension of axons and dendrites, and integration into host neocortex (Fig. 7). Local projections of donor neurons labeled with the lipophilic dye PKH26 within regions of targeted cell death were often appropriately oriented, with apical dendrites and axons. Transplanted neurons displayed characteristic pyramidal neuron morphology under fluorescence and Nomarski DIC, with large cell somas, large nuclei, and prominent nucleoli. Dendritic branching often extended toward the pial surface, while axons could be seen to project for several hundred micrometers toward the corpus callosum. Donor neurons were distinctly identified by granular, rhodamine fluorescence, and they assumed a large, pyramidal phenotype similar of neurons from another control mouse 2 weeks after transplantation into intact neocortex. Neurons exhibit extremely limited dispersion from the site of injection (arrows). c, a, f, Neurons transplanted into intact neocortex remain highly localized to the transplantation site 1-4 weeks following transplantation. Neuronal positions were digitized from camera lucida drawings, measured perpendicularly from the injection track. Donor neurons are located an average of 23 μm from the transplantation track with a maximal spread of 140 μm. e, The positions of donor neurons transplanted into neocortex of control mice following injections of photoactive nanospheres without light exposure did not differ significantly from transplants into intact mice. Scale bars: a, 150 μm; b, 250 μm.
Figure 5. Progressive migration of transplanted embryonic neurons into regions of targeted neuronal degeneration in neocortex of adult mice. a, Camera lucida drawing for orientation, including the transplantation site (asterisks) within a region of targeted cell death, 1 week following transplantation. The boxed field is shown at higher magnification in the adjacent fluorescence photomicrograph b. CC, Corpus callosum; LV, lateral ventricle; neocortical laminae are indicated. c, Higher magnification fluorescence photomicrograph of the boxed field in a. In mice with targeted neuronal degeneration in lamina II/III, 1 week following transplantation, clusters of immature neurons are seen migrating from the injection track (asterisks). These include clusters oriented obliquely in layer IV (arrows), migrating across this layer to their final locations in lamina II/III. Camera lucida drawing for orientation, including the transplantation site within a region of targeted cell death, 2 weeks following transplantation. The boxed field is shown at higher magnification in the adjacent fluorescence photomicrograph d. CC, corpus callosum; LV, lateral ventricle; HC, hippocampus; neocortical laminae are indicated). d, Higher magnification fluorescence photomicrograph of the boxed field in c. Two weeks after transplantation, donor neurons assume positions within neuron-deficient regions in lamina II/III and differentiate into pyramidal neurons. e, Nissl stain of the same field in an adjacent section demonstrates donor neurons within superficial lamina II/III with large pyramid morphology (large arrows), laterally dispersed from the injection track. Donor neurons with pyramidal neuron morphology are interspersed with endogenous host interneurons (small arrows). Scale bars: a and c, 500 μm; b, 50 μm; d and e, 40 μm.

to remaining, endogenous pyramidal neurons when examined under Nomarski DIC optics. Large cell diameters ranging from 12–25 μm, and fluorescent labeling at the periphery of the cell soma and into the proximal processes, indicated neuronal identity. In contrast, neurons remaining at the site of transplantation, or transplants into intact neocortex, typically remained small and ovoid with presumptive interneuron morphology. The frequent observations of donor neurons in experimental mice with typical pyramidal neuron morphology and appropriate local projections support a directed reintegration and repopulation of neuron-deficient regions.

Expression of neuron-specific markers by donor neurons within regions of targeted cell death

The integration and differentiation of donor neurons were further confirmed by antibody labeling with the neuron-specific markers.
REGIONS OF PHOTOLYTIC DEGENERATION (3 DAYS)

REGIONS OF PHOTOLYTIC DEGENERATION (2 WEEKS)

REGIONS OF PHOTOLYTIC DEGENERATION (1 WEEK)

REGIONS OF PHOTOLYTIC DEGENERATION (4 WEEKS)

NEURONAL MIGRATION FROM INJECTION SITE (µm)

NEURONAL FREQUENCY (% of TOTAL)

NEURONAL MIGRATION FROM INJECTION SITE (µm)

NEURONAL FREQUENCY (% of TOTAL)

NEURONAL MIGRATION FROM INJECTION SITE (µm)

NEURONAL FREQUENCY (% of TOTAL)

NEURONAL MIGRATION FROM INJECTION SITE (µm)

NEURONAL FREQUENCY (% of TOTAL)
Figure 7. Pyramidal morphology and local projections of donor neurons within regions of targeted photolytic neuronal cell death. a and b, Fluorescence photomicrographs of transplanted neurons, labeled with the lipophilic dye PKH26, within the neuron-deficient regions. The neurons (open arrows) display characteristic pyramidal neuron morphology including axons (large arrows) extending ventrally to the corpus callosum, and apical dendrites (small arrows) oriented dorsally toward the pial surface. c, Fluorescence image of a donor neuron (large arrow) labeled with PKH26, revealing the apical dendrite (small arrows). d, Appearance of the same donor neuron (large arrow) under Nomarski DIC optics reveals a large soma, large nucleus, and prominent nucleoli (arrowhead). The base of a large apical dendrite is outlined (small arrows). e and f, Fluorescence appearance of a transplanted neuron (open arrow) in two focal planes, illustrating the ventral extension of the axon (large arrows) toward the corpus callosum and dorsal extension of the dendrite (small arrows) toward the pial surface. Scale bars: a and b, 25 μm; c and d, 15 μm; e and f, 25 μm.

NSE and NF. Donor neurons had cell body diameters characteristic of differentiated pyramidal neurons, displayed prominent axons and dendrites, and were widely dispersed within regions of targeted neuronal degeneration (Fig. 8). Neuron-specific enolase displayed most intense staining within the neuronal cytoplasm, leaving the nucleus and processes largely unlabeled. Staining by NSE within cells containing granular nanosphere fluorescence supported the morphologic criteria used to identify transplanted neurons, determined by large cell somas, nuclei and prominent nucleoli (Fig. 8c,d). Neurofilament staining exhibited a characteristic fibrillar pattern in neuronal processes with intense staining in the soma. Neurons of donor origin showed both fluorescent nanosphere labeling and NF staining in processes which were often oriented radially toward the pial surface and corpus callosum (Fig. 8e,f). The pattern of neuronal immunocytochemical labeling within donor neurons was indistinguishable from endogenous pyramidal neurons, further indicating appropriate integration and directed differentiation into pyramidal neuron phenotype within regions of targeted neuronal cell death.

Discussion
Our results that targeted photolytic neuronal death in adult mice produces directed migration and differentiation of transplanted embryonic neurons provide evidence for selectively altered expression of molecular signals that are usually developmentally regulated. The transplanted embryonic neurons in areas undergoing targeted apoptosis underwent a sequence of migratory, differentiation, and integration events appropriate to the initial development of the superficial lamina II/III. These events did not occur in control mice. Clusters of embryonic neurons began migrating away from the transplantation track within 3 d after transplantation, and migration up to 430 μm was largely complete by 2 weeks. The subpopulation of neurons repopulating the regions of photolytic degeneration underwent directed differentiation into pyramidal neurons when compared to neurons remaining at the transplantation site or in control mice, which resembled endogenous interneurons. The finding of extremely limited migration by donor neurons in intact adult mice is consistent with previous studies in which embryonic neurons transplanted into intact juvenile mice were maximally dispersed and morphologically differentiated by 2–3 weeks following transplantation (Macklis, 1993), and with results from the extensive literature involving grafts into the parenchyma of neocortex in postmigrational and adult hosts. Together, these findings suggest that alterations in the host microenvironment, possibly due to loss of cell-cell and/or cell-matrix interactions, can be elicited
in neocortex of fully adult mice and in turn, influence the mig-
ration, differentiation, and integration of transplanted immature
neurons.

The extent of migration and differentiation by donor neurons
within regions of targeted photolytic neuronal death in adult
hosts was similar to that seen in juvenile mice (Macklis, 1993).
This supports the hypothesis that the specific loss of neurons by
photolytically induced apoptosis can activate intercellular signal
changes leading to plasticity independent of host age. Both stud-
ies demonstrate a preferential migration and assumption of py-
ramidal neuron morphology, limited to the region undergoing
targeted photolytic neuronal degeneration in lamina II/III. Donor
neurons partially restored normal cortical architecture by repa-
pulating neuron-deficient regions. Neurons placed adjacent to
regions of photolytic degeneration in juvenile mice migrated up
to 780 μm perpendicularly from the transplantation track. Neu-
rons centrally located in regions of photolytic degeneration in
adult mice migrated up to 430 μm perpendicularly from the
transplantation track. These perpendicular measurements may be
underestimates of the true migration distances; neurons with
oblique trajectories would be expected to migrate over consid-
erably longer distances (approximately 600–700 μm). The dif-
cerence observed between adolescent and adult hosts may also
simply be due to the change in experimental paradigm, since
cell suspensions in the present experiments were transplanted
through the center of the regions of neuronal degeneration rather
than adjacent to these zones. The modification effectively halves
the lateral dimension of regions of degeneration into which neu-
rons may migrate, while increasing the probability of successful
engraftment into a region of targeted cell death. The results of
these experiments in adult mice provides an opportunity to iso-
late the molecular alterations in adults with less potential con-
fusion by unrelated late elements of the normal developmental
timetable that could be present in younger mice.

Several observations demonstrate that the labeled neurons are
indeed the donor neurons. Evidence from the current experi-
ments argues in three ways for specific identification of neurons
of donor origin and against nonspecific labeling of host neurons
by accumulation of free label following transplantation or from
release of label by dying or lysed donor cells. (1) Neurons la-

Figure 8. Expression of neuron-specific antigens by donor neurons within regions of targeted neuronal degeneration. Fluorescence photomicrograph
demonstrates donor neurons labeled with rhodamine fluorescent nanospheres (arrows), laterally dispersed from the transplantation site (asterisks) 2
weeks after transplantation. Unlabeled, remaining endogenous neurons are indicated (arrowheads). b, The same donor neurons (arrows) assume a
large pyramidal morphology indistinguishable from remaining, endogenous pyramidal neurons (arrowheads) when examined under Nomarski DIC
optics. c and d, A transplanted neuron (large arrow), identified by intracellular fluorescence in c, is double stained for neuron-specific enolase
(NSE) in d. Neuron-specific enolase displays most intense staining within the cytoplasm, leaving the nucleus and processes largely unlabeled. The
pattern of NSE staining observed in donor neurons within regions of neuronal degeneration is identical to that of the remaining endogenous
pyramidal neurons (small arrow). e and f, Fluorescence and bright field images of a transplanted neuron (large arrow) immunocytochemically
stained for neurofilament. Neurofilament staining is localized within the periphery of the soma and within axons (arrows) and dendrites (small
arrowheads), properly oriented toward the corpus callosum and pial surface, respectively. Donor neurons resemble endogenous, host neurons (large
arrowhead), assuming large pyramidal morphology within regions of targeted neuronal degeneration. Scale bars: a and b, 50 μm; d-f, 20 μm.
beled with the neuron-selective, rhodamine fluorescent nanospheres or the lipophilic dye PKH26 are located distant from the transplantation site selectively within regions of targeted cell death in superficial lamina II/III while labeled neurons in deeper cortical layers remain extremely close to the site of injection. Nonspecific uptake of either label by host neurons would not show laminar preference. (2) The intensity of labeling following direct application of fluorescent nanospheres or PKH26 in vitro leads to distinct and more pronounced labeling of donor neurons than is observed following nonspecific uptake and labeling of host neurons at a nanosphere injection site in vivo. (3) Neurons transplanted into regions of targeted cell death morphologically assume a pyramidal phenotype. Nonspecific uptake of the fluorescent labels would be expected to occur in both interneurons and pyramidal neurons far from the transplantation site. Further evidence derives from prior control studies of labeled, hypoosmotically lysed neocortical neurons and neuroblastoma cells, viable neocortical neurons, and viable cerebellar neurons transplanted into intact cortex (Macklis, 1993); these experiments did not exhibit lamina-specific migration and differentiation of donor neurons. Similarly, labeled neocortical neurons transplanted into kainic acid lesioned hosts did not display directed migration or differentiation by donor neurons. When lysed neurons did release their contents within the host brain, non-neuronal phagocytes engulfed and cleared the debris, as observed by fluorescence and electron microscopy. Taken together, such findings argue strongly against the possibility of widespread, nonspecific uptake of fluorescent nanospheres or PKH26 by host neurons.

Our results add additional support to the concept of microenvironmental regulation of neuronal migration and differentiation. The directed behavior of donor neurons is sharply limited to the regions of targeted degeneration, their development is not autonomous, and local alterations can support their migration and differentiation even in adult mice. Our results are consistent with results seen using other systems of neuronal injury within the central nervous system. Embryonic cerebellar Purkinje cells, transplanted into cerebellum of adult Purkinje cell degeneration (PCD) mutant mice, display elements of appropriate positional and temporal specificity in repopulating neuronally depleted regions (Sotelo and Alvarado-Mallart, 1986, 1987, 1988; Gardette et al., 1988; Sotelo et al., 1994). Many anatomic lesioning experiments reveal similar but more limited elements of plasticity within the host environment when appropriate embryonic neurons are transplanted into the affected sites (Itakura et al., 1990; Stromberg et al., 1992; Takeuchi et al., 1992; Cenci et al., 1993; Groves et al., 1993). Directed differentiation is observed in the goldfish retina system (Braisted and Raymond, 1992, 1993), where laser ablation of retinal cone photoreceptors leads to specific regeneration of cone photoreceptors and repopulation of the lesioned area (Braisted et al., 1993).

Microenvironmental control over neural cell fate has become increasingly supported by experimental evidence from recent transplantation studies of both primary neurons and genetically modified precursor cell lines. Primary cerebellar precursor cells can shift fate by differentiating into neurons with hippocampal phenotype upon transplantation into neonatal dentate gyrus (Vicario et al., 1993), while striatal precursors transplanted into the embryonic cortical ventricular zone generate neurons with cortical phenotypes (Fishell, 1993). Similarly, immortalized multipotent precursor cells undergo differentiation appropriate for the site of implantation within the neonatal hippocampus, neonatal cerebellum, and adult striatum (Renfranz et al., 1991; Snyder et al., 1992; Sheen et al., 1993, and unpublished observations). Studies of the immortalized C17-2 precursors (Snyder et al., 1992) derived from neonatal cerebellum implanted into intact neocortex, and along the neuraxis, of mice at various times during development and postnatal life indicate that differentiation into neuronal phenotype is limited to the period of neurogenesis within a particular CNS region (Snyder et al., 1993 and unpublished observations). Transplants of C17-2 cells into adult intact and kainic acid lesioned neocortex did not receive the necessary local cues to initiate any neuronal differentiation, but they underwent directed differentiation into both neurons and glia following transplantation into regions of targeted photolytic neuronal death (Macklis et al., 1994; E. Y. Snyder, C. H. Yoon, J. D. Flax, and J. D. Macklis, unpublished observations). The plasticity observed following targeted photolytic neuronal death cannot be attributed to a generalized response to the degeneration. These results support the hypothesis that directed perturbation of the microenvironmental by targeted apoptosis induces a localized and specific temporal and spatial sequence of signals necessary to achieve repopulation by pyramidal neurons, and that these signals can provide an additional level of hierarchical control over immature neural precursors.

Several factors could contribute to the changes we observe following transplantation into these selectively perturbed host environments. The directed migration could potentially be supported by reemergence of structural elements such as radial glia that usually transform into astrocytes when neuronal migration is complete (Rosen et al., 1992, Sheen et al., 1992, Hatton and Hoi, 1993; Hunter and Hatten, 1993; Hernt-Grant and Macklis, 1994), or it could be due to guidance provided by the extracellular matrix or diffusible trophic factors (Politis and Miller, 1985a; Stewart and Pearlman, 1987; Chun and Shatz, 1988; Liesi and Silver, 1989; Joosten et al., 1994). The appropriate differentiation could result from the altered expression of a variety of neurotrophins and growth factors, in addition to local alterations in synaptic activity (Gardette et al., 1991; Hickmott and Constantine-Paton, 1993; Kalb and Agostini, 1993; Lohof et al., 1993; Allendoerfer et al., 1994). Trophic factors centrally influence long term survival and differentiation of neurons (Cheng and Mattson, 1991; Jacobson, 1991; Sieber-Blum, 1991; Zhang et al., 1993). Recent evidence demonstrates that basic fibroblast growth factor (bFGF) acts both as a mitogen and as a trophic factor (Reynolds and Weiss, 1992; Kilpatrick and Bartlett, 1995; Ray et al., 1995; Kitchens et al., 1994), and that it contributes to the regenerative response seen in retina and optic tract following phototoxicity or retinal dystrophy (Kostyk et al., 1994) and following traumatic cortical injury (Finkelstein et al., 1988; Lin and Chen, 1994). Similarly, expression of neurotrophins such as NGF or brain-derived neurotrophic factor (BDNF), and neurotrophin-induced receptor tyrosine kinase (Trk) phosphorylation, correlate temporally with developmental periods of neuronal migration and early differentiation (Zafra et al., 1990; Komuro and Kukic, 1993; Knusel et al., 1994). Brain derived neurotrophic factor, in particular, is implicated in normal cortical neuron survival and differentiation, and expression increases following subtle injury due to hypoxia or excitotoxicity, but only in regions where cells remain partially viable and not necrotic (Ballarin et al., 1991; Comelli et al., 1992; Ghosh et al., 1994). Determination of the temporal and spatial expression of pivotal regulatory factors, and their roles within regions of photolytic apoptosis, will elucidate the mechanisms that drive
cellular migration and differentiation during normal corticogenesis.

In conclusion, these studies show that selective neuronal degeneration following targeted photolysis induces a specific sequence of events that guide migration and differentiation of transplanted embryonic neurons, potentially recapitulating signaling sequences seen during development, and leading to the restoration of elements of normal cortical cytoarchitecture. Neuronal migration extends over a 2 week period following transplantation, with donor neurons assuming characteristic pyramidal phenotype in these neuron-deficient regions. This response can occur in fully adult mice, further suggesting that the potential to guide cellular migration and differentiation through alteration of microenvironmental cues remains possible despite host ages well beyond normal development. It will be of interest to investigate the cellular and molecular alterations in the regions of targeted neuronal cell death against a background in control adult cortex in which most developmentally regulated molecules will have been downregulated. Controlled perturbations of this system and these signals may allow definition of spatial and temporal patterns of developmentally regulated molecules permissive or required for directed neuronal migration and differentiation within neocortex.

References


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