Brain injury expands the numbers of neural stem cells and progenitors in the SVZ by enhancing their responsiveness to EGF

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ABSTRACT

There is an increase in the numbers of neural precursors in the SVZ (subventricular zone) after moderate ischaemic injuries, but the extent of stem cell expansion and the resultant cell regeneration is modest. Therefore our studies have focused on understanding the signals that regulate these processes towards achieving a more robust amplification of the stem/progenitor cell pool. The goal of the present study was to evaluate the role of the EGFR [EGF (epidermal growth factor) receptor] in the regenerative response of the neonatal SVZ to hypoxic/ischaemic injury. We show that injury recruits quiescent cells in the SVZ to proliferate, that they divide more rapidly and that there is increased EGFR expression on both putative stem cells and progenitors. With the amplification of the precursors in the SVZ after injury there is enhanced sensitivity to EGF, but not to FGF (fibroblast growth factor)-2. EGF-dependent SVZ precursor expansion, as measured using the neurosphere assay, is lost when the EGFR is pharmacologically inhibited, and forced expression of a constitutively active EGFR is sufficient to recapitulate the exaggerated proliferation of the neural stem/progenitors that is induced by hypoxic/ischaemic brain injury. Cumulatively, our results reveal that increased EGFR signalling precedes that increase in the abundance of the putative neural stem cells and our studies implicate the EGFR as a key regulator of the expansion of SVZ precursors in response to brain injury. Thus modulating EGFR signalling represents a potential target for therapies to enhance brain repair from endogenous neural precursors following hypoxic/ischaemic and other brain injuries.

Key words: brain injury, epidermal growth factor receptor (EGFR), neural stem cell, neurosphere, subventricular zone.

INTRODUCTION

A number of studies have demonstrated that the brain possesses a limited regenerative capacity following injury (Magavi et al., 2000; Arvidsson et al., 2002; Parent et al., 2002; Goings et al., 2004; Ohab et al., 2006). These investigations have predominantly evaluated neuronal production in the adult brain following injury (Jin et al., 2001; Zhang et al., 2001; Plane et al., 2004). Although encouraging, the extent of cell replacement is limited. However, by understanding the adaptive mechanisms that occur in the SVZ (subventricular zone) subsequent to injury, therapeutic strategies can be designed to achieve a more significant level of regeneration after CNS (central nervous system) injury.

Brain injury, as a consequence of cardio-respiratory insufficiency, is the most common cause of morbidity in premature and term infants. In the US alone, it has been estimated that one in every eight babies is born premature, and that over 80 000 infants per year are born very pre-term. Up to 40% of the infants born pre-term and weighing less...
than 1500 g will develop neurological and psychological problems ranging from classic cerebral palsy to mild learning deficits that may not appear until later in life (Volpe, 2001). There is an even larger cohort of infants that acquire developmental brain injuries as a consequence of insults such as intrauterine infection or asphyxia at birth. Presently, there are limited treatments available to prevent brain damage from injuries sustained in the perinatal period and there are no approved therapeutic regimens to stimulate CNS regeneration.

EGFRs [EGF (epidermal growth factor) receptors] are expressed in the infant, juvenile, young adult and adult human SVZs (Weickert et al., 2000), as well as on actively dividing cells in the postnatal rodent SVZ (Seroogy et al., 1995; Weickert and Blum, 1995). EGFR and its ligands maintain embryonic and adult neural stem cells (Doetsch et al., 1999; Junier, 2000; Yarden, 2001; Doetsch et al., 2002) and promote the diversification of neural precursors during embryonic cortical development (Burrows et al., 1997; Kornblum et al., 1997; Burrows et al., 2000; Caric et al., 2001). Postnatally, EGFR confers a motile phenotype to non-migratory cortical neuronal progenitors (Aguirre et al., 2005). Surprisingly, there are very few studies on the role of EGFR in the regenerative responses of the neural stem/progenitor cells following brain damage (Justicia and Planas, 1999; Teramoto et al., 2003).

Using a rodent model of neonatal brain injury, in the present study we demonstrate that there is a robust increase in EGFR signalling in both neural stem cells and progenitors. We also show that the regenerative response can be recapitulated in vitro by transfecting NSPs (neural stem progenitors) with a constitutively active EGFR, and that pharmacologically inhibiting EGFR in vitro abrogates the expansion of the NSPs observed subsequent to injury. The results of the present study suggest that enhanced EGFR expression is critical for NSP expansion.

MATERIALS AND METHODS

Neonatal H/I (hypoxia/ischaemia) model

All animal work was approved by the institutional animal care and use committee guidelines of the New Jersey Medical School. Timed pregnant Wistar rats (Charles River Laboratories) were housed on a 12 h light/dark cycle at 25°C with Purina rodent chow #5001. Following a normal delivery, the litter size was culled to 12 pups per litter. Neonatal H/I was induced on postnatal day 6 (P6) rat pups (PO as day of birth) by a permanent right CCA (common carotid artery) cauterization followed by systemic hypoxia. Briefly, P6 rat pups were anaesthetized with isoflurane (5% induction, 3% maintenance) following which a midline neck incision was made. The right CCA was separated from the vagus nerve and cauterized using a bipolar cauterizer (Malis bipolar cauterize and bipolar cutter; Codman) at 10 V. The skin incision was sutured using 4-0 silk and the pups were returned to the dam for 2 h. For mouse neonatal H/I, the litter size was culled to 12 pups per litter and permanent CCA cauterization was performed on P10 pups as explained above for rats. Prior to hypoxia, the pups were pre-warmed in jars partially submerged in a 37°C water bath for 20 min after which rat pups were subjected to systemic hypoxia in jars at 8% O2/92% N2 for 75 min and mouse pups at 10% O2/92% N2 for 65 min. Sham-operated control animals had their right CCA separated from the vagus nerve, but it was not coagulated. They were subjected to hypoxia. Following hypoxia, the animals were left in the jars for 15 min at normoxia, after which they were returned to their cages.

Primary neurosphere assay

After 3 days of recovery from H/I, P9 rat pups were decapitated and their brains removed. Under aseptic conditions, a cut was made 2 mm from the anterior pole of the brain. A second cut was made approx. 3 mm posterior to the first cut. The hippocampus, corpus callosum and the meninges were removed under the microscope. Using forceps, 12 o’clock and 3 o’clock incisions were made and the region enclosed between the cortex and the ventricle containing the SVZ was removed and placed in fresh PGM [1 mM MgCl2 and 0.6% dextrose in PBS (pH 7.3)]. The tissue was mechanically minced using forceps and then enzymatically dissociated using 0.05% trypsin/EDTA at 37°C for 7 min. The trypsin was inactivated by the addition of an equal volume of newborn calf serum and the tissue was resuspended in ProN media [DMEM (Dulbecco’s modified Eagle’s medium)/F12 (1:1) media containing 10 ng/ml D-biotin, 25 μg/ml insulin, 20 nM progesterone, 20 mM Heps, 100 μM putrescine, 5 ng/ml selenium, 50 μg/ml apo-transferrin and 50 μg/ml gentamycin] and triturated in ProN. The triturated suspension was passed through a 40 μM Nitex screen and the cells were collected by centrifugation at 200 g for 2 min and washed with ProN. The cells were counted with 0.1% Trypan Blue dye under a haemocytometer and plated at 5 × 10⁴ cells/ml in ProN medium supplemented with 2 ng/ml EGF or 1 ng/ml FGF (fibroblast growth factor)–2 or as stated. The cells were cultured at 37°C in 5% CO2 incubators and fed every 2 days by removing approx. half of the medium and replenishing with fresh medium.

Neurosphere quantification

A neurosphere was defined as a free-floating cluster of at least 25 μm diameter. Prior to counting the spheres, the plates were shaken to ensure uniform distribution of the spheres in the well. The number of neurospheres in five random fields under 4 x magnification was determined for each well. The total number of neurosphere-producing cells, the neural stem/progenitor cells, in the population was
extrapolated from the average number of spheres per field, the area of the field and the area of the well. For neurosphere volumetric measurements, digital photographs were captured from at least ten neurospheres from each condition (n = 6 SVZ per condition) and volumes were calculated using IP Lab 3.6 software.

Immunofluorescence

Cryostat brain sections were collected from animals killed at 48 h of recovery. Sections were stained using a rabbit anti-EGFR antibody (Santa Cruz Biotechnology, catalogue number 1005; used at a 1:100 dilution). Secondary antibodies were LRSC (lissamine rhodamine sulfonyl chloride)-conjugated goat anti-rabbit IgG (Jackson Immunoresearch). Sections were thawed in ice-cold PBS and blocked with PGB superblock (10% BSA and 1% normal goat serum in PBS) for 1 h at room temperature (20°C). The superblock was aspirated and primary antibodies were applied in a 1:5 dilution of PGB superblock and incubated overnight at 4°C. The slides were rinsed extensively with PBS and then incubated in secondary antibodies for 1 h at 37°C. The sections were washed, counterstained with DAPI (4',6-diamidino-2-phenylindole), and coverslipped with Gel Mount (Biomeda). To control for non-specific staining, the primary antibody was replaced with a 1:200 dilution of normal rabbit serum. Images were obtained using a Zeiss LSM410 confocal laser-scanning microscope using a 40× objective. The exposure time for capturing the images was identical. Images were cropped and assembled into a montage using Photoshop 9.0.

Flow cytometry

The SVZ was dissected out of 3 day H/I recovery and sham-operated control P9 Wistar rat pups and dissociated as described above. Approx. 2 × 10⁵ cells were used for each analysis. The cells were incubated with a rabbit polyclonal anti-EGFR antibody (Ab 15669, Abcam; used at a 1:50 dilution) together with either a mouse monoclonal anti-NG2 (neuro-glial protein 2; Millipore; used at a 1:100 dilution) or a mouse monoclonal IgM anti-PSA-NCAM (poly-sialated neural cell adhesion molecule; Millipore; used at a 1:500 dilution) for 1 h on ice. After several washes, the cells were incubated with appropriate secondary antibodies conjugated with fluoro-chromes for 1 h on ice protected from light. To analyse EGFR-expressing cells in mice, subsequent to neonatal H/I, approx. 2 × 10⁶ cells were stained using a mouse IgM monoclonal anti-SSEA-1 (stage-specific embryonic antigen-1)/LeX antibody (Zymed Laboratories). Following several washes, the cells were incubated with biotin-conjugated anti-rabbit antibody and then streptavidin-conjugated Alexa Fluor® 488 for EGFR detection, and RedX-conjugated anti-mouse IgM for LeX detection for 1 h each on ice and protected from direct light. The cells were analysed using a BD FACS Calibur®. A population of unstained rat SVZ cells was used to calibrate the BD FACS Calibur® machine prior to analysis. Negative controls used included replacing the primary antibody with normal rabbit serum (Jackson Immunoresearch; used at a 1:3000 dilution) and normal mouse serum (Jackson Immunoresearch; used at a 1:3000 dilution) followed by secondary and tertiary antibodies as described above. The data were analysed using FlowJo version 4.4.4 software.

Western blot analysis

SVZ tissue dissected from H/I and sham-operated P9 rat pups was homogenized with a tissue/lysis buffer ratio of 0.5 g of tissue/2 ml of lysis buffer in a glass homogenizer (Kontes glass homogenizer with Pestle ‘A’). The HES lysis buffer used contained 10 mM Hepes, 1 mM EDTA and 250 mM sucrose (pH 7.4) with 2 mM sodium orthovanadate and one tablet of protease inhibitor (Complete Mini PI cocktail tablets; Roche diagnostics) for 10 ml of buffer, added just prior to use. The homogenate was spun at 1000 g for 10 min in a table top Beckman centrifuge to pellet nuclei. The supernatant then spun at 150 000 g in a Beckman Ultracentrifuge with the titanium SW55Ti swinging bucket rotor. This membrane-enriched fraction was resuspended in HES buffer and the protein concentration was estimated using the BCA (bicinchoninic acid) assay (Pierce Biotech). Total protein (30 μg) was mixed with 4 × NuPage LDS sample buffer and 10 × NuPage reducing agent and heated for 10 min at 90°C and loaded on to 4–12% NuPage Bis-Tris pre-cast gels (Invitrogen). Approx. 2 μl of MagicMark XP (Invitrogen) was loaded for standard molecular mass markers. Following electrophoresis, the proteins were transferred on to a nitrocellulose membrane and stained with Ponceau S to evaluate equal protein loading into the wells. The blots were blocked with 5% non-fat dried skimmed milk/1% BSA in PBS-Tween followed by incubation with primary antibody in 1% BSA in PBS-Tween overnight at 4°C with gentle rocking. The primary antibodies used were rabbit polyclonal anti-EGFR antibody (Ab 15669, Abcam; used at a 1:250 dilution) or mouse monoclonal anti-phospho-tyrosine-100 (Cell Signalling Technologies; used at a 1:2000 dilution). Following three rinses with PBS-Tween the following day, the blots were incubated with the corresponding secondary antibodies such as DAR-HRP (horseradish peroxidase)-conjugated (Jackson Laboratories; used at a 1:10000 dilution) or DAM-HRP conjugated (Jackson laboratories; used at a 1:5000 dilution) antibodies for 2 h at room temperature. The blots were washed and the signal developed with Western Lightning chemiluminescence reagent (PerkinElmer) according to the manufacturer’s protocol. The bands were visualized using UVP EpiChem® and processed with Labworks 4.0 digital quantification software (UVP).

RNA isolation

SVZs were microdissected as outlined above and snap frozen in 0.5 ml of TRIzol® (Invitrogen) per SVZ in a slush of dry ice/ethanol and stored at −80°C. Just prior to extraction, the
tissues were thawed and homogenized using a hand-held tissue homogenizer. Chloroform (100 µl) was added per sample and mixed to obtain a cloudy suspension that was centrifuged at 800 g for 5 min, following which the aqueous phase was transferred to a new tube. Approx. 250 µl of 70% ethanol was added and the mixture was applied to an RNeasy Mini-spin column (Qiagen) and the manufacturer’s protocol was followed to remove contaminants. The purity and amount of total RNA obtained was determined from the absorbance values measured using a UV–visible spectrophotometer (Becton Dickinson). The total RNA concentration was adjusted to 200 ng/µl and stored at −80 °C until further use.

**Thymidine incorporation assay**

At 72 h, post-H/I and sham-operated control rat pups were decapitated and the SVZs dissected out as described above. The SVZ cells were dissociated and cultures were set up at 3 x 10^6 cells/ml in 20 ng/ml EGF in ProN overnight. The following morning the cells were spiked with 8 µCi of [³H]thymidine. Starting every 4 h, the cells were collected on centrifugation at 800 g for 5 min, following which the aqueous phase was transferred to a new tube. Approx. 250 µl of 70% ethanol was added and the mixture was applied to an RNeasy Mini-spin column (Qiagen) and the manufacturer’s protocol was followed to remove contaminants. The purity and amount of total RNA obtained was determined from the absorbance values measured using a UV–visible spectrophotometer (Becton Dickinson). The total RNA concentration was adjusted to 200 ng/µl and stored at −80 °C until further use.

**EGFR gain-of-function**

Plasmid pEF2-Fc–γR1–IRESGFP [where IRES is internal ribosome entry site and EGFP is enhanced GFP (green fluorescent protein)] was digested with NheI and Spel followed by heat inactivation of the restriction enzymes. The digested ends were dephosphorylated with shrimp alkaline phosphatase to avoid vector re-ligation. Plasmid pEF2–FLAF–EGFR was digested with NheI and Xbal to obtain the EGFR fragment. This fragment was ligated to the vector to obtain pEF2–FLAF–EGFR–IRESGFP. The resulting plasmid was transfected into COS cells and the transfection resulted in increased levels of phosphorylated EGFR as a proof of concept (results not shown).

Primary neurospheres were generated from rat pup SVZs and dissociated with accutase to yield single cells. Approx. 2 x 10^6 cells/condition were used for pEGFR_IRES_EGFP and pEGFP plasmid using the rat nucleofector kit (Lonza, catalogue number VPG-1005). Mock transfection controls had cells that were pulsed using the Amaxa system, but with no added DNA. Fluorescence in the transfected cells was readily evident by 24 h. At this time, experiments were performed to assess cell division. Cells were diluted to 5 x 10^4 cells/well in neurosphere growth medium, cultured 7 days in vitro. Some cells were incubated with 10 µM BrdU (bromodeoxyuridine) and samples were collected 6 and 12 h post-BrdU addition and processed for BrdU incorporation analysis using the BD Pharmingen BrdU Flow Kit.

**Statistical analysis**

Data were analysed using one-way ANOVA followed by Tukey’s post-hoc test or Student’s t test to detect significant differences between the means with P<0.05. Analyses were performed using SAS (Cary). For real-time PCR data, the REST program was used to detect significant differences between the means of treatment groups and untreated controls based on P values of 0.05.
RESULTS

Neurosphere number and size increase following H/I in the presence of EGF, but not FGF-2

The neurosphere has remained a standard assay for measuring numbers of neural stem cells in the SVZ, therefore we isolated SVZ cells to determine whether there was an increase in the number of sphere-forming cells after neonatal H/I. Neurosphere cultures of rat SVZs were established from ipsilateral, contralateral and sham-operated control animals 72 h post-H/I. Spheres were propagated in a chemically defined serum-free medium (ProN) supplemented with either EGF (2 ng/ml) or FGF-2 (1 ng/ml). After 7 DIV, the number of neurospheres obtained from the ipsilateral SVZ following injury was 5-fold higher in medium supplemented with EGF as compared with FGF-2 [1381.7 ± 105.2 compared with 261.5 ± 32.4 (means ± S.E.M.), n=6 brains] (Figure 1B). There was a significant difference in the number of neurospheres generated from the ipsilateral SVZ compared with the contralateral SVZ in EGF-supplemented media (1381.7 ± 105.2 compared with 511.5 ± 90.3, n=6 brains), but no significant difference in the number of spheres obtained from the ipsilateral SVZ compared with the contralateral SVZ when cultured in FGF-2-supplemented media (261.5 ± 32.4 compared with 223.10 ± 24, n=6 brains). There was no difference in the number of neurospheres obtained from sham-operated control SVZs in medium supplemented with EGF as compared with FGF-2 (114.8 ± 42.2 compared with 82.9 ± 33.3, n=6 brains). The addition of FGF-2 to the EGF-containing medium had no effect on the outcome of this assay (Figure 1A).

Measuring the volume of the neurosphere can provide insights into the proliferative capacity of the NSPs; therefore we measured the diameter of at least ten neurospheres from each culture condition (n=6 SVZs per condition). The size of the neurospheres obtained from the ipsilateral hemisphere of the injured brain cultured in 2 ng/ml EGF was 17-fold larger (P<0.05) compared with neurospheres from the ipsilateral SVZs cultured in 1 ng/ml FGF-2 (470143.9 ± 79422.2 μm$^3$ compared with 26676.2 ± 7397.6 μm$^3$, n=6 brains). Contralateral SVZ cells cultured in EGF-supplemented medium were 2.5-fold larger (P<0.05) in size compared with their counterparts cultured in FGF-2-supplemented medium (27065.2 ± 9611.3 μm$^3$ compared with 11055.8 ± 1583.7 μm$^3$, n=6 brains). There was no significant difference in sham-operated control SVZ sphere sizes between EGF- and FGF-2-supplemented media (7579.2 ± 2275.7 μm$^3$ compared with 7833.4 ± 3273.4 μm$^3$). Lastly, there was a 62-fold change in neurosphere size difference between ipsilateral SVZ- and control SVZ-generated spheres in EGF-supplemented media (ipsilateral neurosphere, 470143.7 ± 79422 μm$^3$; control neurosphere, 7579.2 ± 2275.7 μm$^3$) as compared with a 3.4-fold change in FGF-2-supplemented media (ipsilateral neurosphere, 26676.2 ± 7397.6 μm$^3$; control neurosphere, 7833.4 ± 3273.4 μm$^3$) (Figures 1A and 1C).

Pharmacologically inhibiting EGFR abrogates the increase in precursors following neonatal H/I

To test the hypothesis that an increase in EGFR signalling was responsible for the expansion of the neural progenitors during recovery from neonatal H/I, we administered a commercially available selective inhibitor of the EGFR, PD153035 (alternatively named AG1517 or Compound 32) (Bos et al., 1997; Lichtner et al., 2001). PD153035 is a competitive inhibitor of ATP binding and, hence, prevents functional activation of the receptor. This inhibitor also locks the receptors into inactive dimers and thus reduces EGFR signalling. At 72 h post-H/I, SVZs from ipsilateral and contralateral hemispheres, along with sham-operated control SVZs, were dissected out and cultured in ProN media supplemented with EGF and FGF-2 alone or with the addition of 300 nM or 1 μM PD153035. The neurosphere numbers obtained were quantified at 7 DIV. PD153035 at both 300 nM and 1 μM abrogated the increase in neurosphere number and size measured using the neurosphere assay. The number of neurospheres obtained 7 DIV from the ipsilateral SVZ at 3 day recovery from H/I cultured in growth-factor-supplemented media containing 300 nM and 1 μM PD153035 were 5.5-fold and 16-fold lower (P<0.05) respectively, compared with their counterparts cultured in the absence of the inhibitor (ipsilateral SVZ cells: growth media, 916.8 ± 53.8; +300 nM PD153035, 169.6 ± 43.2; +1 μM PD153035, 55.9 ± 12.3; n=3 with six animals per experiment) (Figure 2). Analogously, at 7 DIV, the number of neurospheres obtained from the sham-operated control SVZs at 3 days of recovery cultured in growth-factor-supplemented media containing 300 nM and 1 μM PD153035 were 3.6-fold and 6-fold lower respectively (P<0.05) compared with their counterparts cultured in growth-factor-supplemented media in the absence of the inhibitor (control SVZ cells: growth media, 324.4 ± 104.06; +300 nM PD153035, 88.5 ± 52.6; +1 μM PD153035, 54.1 ± 32.4; n=3 with six animals per experiment). The number of neurospheres obtained from ipsilateral SVZs in 300 nM PD153035 was not different from that of uninjured control SVZs cultured in growth media containing EGF and FGF-2 (ipsilateral SVZ cells in 300 nM inhibitor, 169.6 ± 43.2; control SVZ cells in growth media, 324.4 ± 104.1; n=3 with six animals per experiment). Finally, the numbers of neurospheres obtained in the presence of either doses of the inhibitor (300 nM and 1 μM PD153035) were not significantly different between injured ipsilateral SVZ and sham-operated control SVZ at 3 day recovery from H/I (at 300 nM inhibitor: ipsilateral, 169.6 ± 43.1 and control, 88.5 ± 52.6; at 1 μM inhibitor: ipsilateral, 55.9 ± 12.3 and control, 54.1 ± 32.4). Notably, FGF-2 at 1 ng/ml [the $K_d$ of FGF-2/FGFR (FGF receptor) 1 is 50 pM] was unable to recapitulate the EGF effect upon inhibition of the EGFR activity (Figure 2).

H/I recruits more SVZ cells into the cell cycle and they divide more rapidly in response to EGF

To determine whether the increase in neurosphere size obtained from the ipsilateral SVZ in EGF-supplemented...
medium was the result of altered cell-cycle kinetics, we performed a cumulative [3H]thymidine incorporation assay. Cells isolated from ipsilateral, contralateral and sham-operated control SVZs at 3 days of recovery from H/I were cultured overnight in EGF-supplemented medium spiked with 8 μCi of [3H]thymidine. The DNA was acid precipitated and the radioactivity was quantified at 4 h intervals. This assay revealed that a significantly greater number of cells in the ipsilateral hemisphere achieved a higher level of [3H]thymidine incorporation compared with cells from the contralateral and control hemispheres. By 16 h, the slope of the rate of [3H]thymidine from the ipsilateral SVZ began to plateau with maximum incorporation of radioactivity reaching approx. 2.5-fold and 10-fold higher levels compared with contralateral and control respectively (at 16 h: ipsilateral, 2935.2 d.p.m.; contralateral, 1233 d.p.m.; control, 307.7 d.p.m.) (Figure 3). Analysing the slopes of the curves, ipsilateral cells showed a 2.5-fold and 18-fold greater slope between 4 and 16 h compared with contralateral and sham-operated control SVZ cells respectively, indicating that the cells in the ipsilateral SVZ cells have reduced cell-cycle time following injury compared with sham-operated controls (slope of ipsilateral, 182.58; slope of contralateral, 77.41; slope of control, 10.09; between 4–16 h) (Figure 3).

**Neonatal H/I increases the expression of EGFR mRNA and protein levels, but not EGFR ligands**

To determine whether the observed increase in neural stem/progenitor numbers and alteration in cell-cycle kinetics subsequent to injury could be attributed to increased EGF responsiveness, we stained brain sections for EGFR at 48 h after H/I and found that cells in the most medial aspect of the

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**Figure 1** Neurospheres grow larger in the presence of EGF, but not FGF-2

(A) SVZ cells were dissociated from ipsilateral (a and d), contralateral (b and e) and control (c and f) hemispheres at 3 days recovery from H/I and cultured for 7 DIV in ProN medium supplemented with either 2 ng/ml EGF (a–c) or 1 ng/ml FGF-2 (d–f). Phase-contrast images of representative neurospheres were captured for quantification. (B) Quantification of the average number of neurospheres obtained per 50,000 SVZ cells. (C) Quantification of the average neurosphere size from at least 30 spheres. *P < 0.05 using ANOVA and Tukey’s post-hoc tests. Values are the mean number/size of neurospheres ± S.E.M. of three experiments with six animals per experiment. IL, ipsilateral; CL, contralateral; CTL, control.
SVZ, immediately subjacent to the ependymal cells, had the pronounced increase in EGFR expression. Increased EGFR expression, however, was also seen within most of the cells of the dorsolateral SVZ. We also analysed EGFR mRNA expression between 12 and 72 h of recovery. At 12 h post-H/I there was a 1.5-fold increase \( (P<0.05) \) in EGFR mRNA in both the ipsilateral and contralateral SVZs compared with sham-operated controls (Figure 4). At 3 days recovery from H/I there was a 3-fold \( (P<0.05) \) increase in EGFR mRNA in the ipsilateral hemisphere compared with controls (Supplementary Figure S1 at http://www.asnneuro.org/an/001/an001e009add.htm).

Increased EGF responsiveness of neural stem cells after injury

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To establish whether the observed increase in EGFR mRNA translated into greater EGFR protein expression, we performed Western blot analysis on subfractionated SVZ homogenates isolated from ipsilateral hemispheres 3 days post-H/I and sham-operated control rat pups. The SVZ tissue was homogenized and fractionated into cytosolic, nuclear and membrane fractions. An analysis of the membrane fractions revealed a 2.8-fold increase ($P < 0.05$) in phosphorylated EGFR in the ipsilateral hemisphere relative to sham-operated control SVZs at 72 h post-injury (Figure 5).

Injury increases EGFR expression in SVZ stem/progenitor cells.

To investigate EGFR expression levels on stem and progenitor cells, we performed flow cytometry for EGFR expression on progenitors (PSA-NCAM and NG2 for rats) and stem cells (LeX for mice). We used anti-NG2 and anti-PSA-NCAM antibodies as they label several types of progenitors within the SVZ. At 3 days recovery from H/I, there was a 20-fold increase ($P < 0.05$) in EGFR$^{+}$/NG2$^+$ cells and a 17-fold increase ($P < 0.05$) in the EGFR$^{+}$/PSA-NCAM$^+$ cells in the ipsilateral SVZ as compared with controls (Figures 6A, 6B, 7A and 7B). There was also a 7-fold increase ($P < 0.05$) in EGFR$^{+}$/NG2$^-$ and a 1.5-fold increase ($P < 0.05$) in EGFR$^{+}$/PSA-NCAM$^-$ cells in the damaged SVZ as compared with controls (Figures 6C and 7C). No significant change was found in the double-positive EGFR/NG2 cells or EGFR/PSA-NCAM cells between the 72 h post-H/I contra-lateral hemisphere and sham-operated control SVZ cells (Figures 6B and 7B). Previously we found that progenitors, expressing PSA-NCAM, are deleted early after H/I injury (Romanko et al., 2004). The increase in the number of PSA-NCAM$^+$/EGFR$^+$ cells probably reflects the effort to replace these progenitors by augmenting their sensitivity to EGFR ligands.
To evaluate whether EGFR is induced on neural stem cells, we used a mouse model of neonatal H/I wherein we could identify putative mouse stem cells using the LeX/SSEA-1 marker. LeX is a fucose-containing trisaccharide that has been reported to mark SVZ cells in the mouse brain that possess stem cell properties (Capela and Temple, 2002). Studies from our laboratory have shown that the murine SVZ is similarly affected by H/I (Brazel et al., 2004), and that there is an increase in murine neural stem/progenitor cells following injury as observed by an increase in the numbers

Figure 5  H/I increases the levels of activated EGFR protein
(A–C) Cryostat sections (12 μm) from animals killed 48 h after H/I by intracardiac perfusion were stained for EGFR and counterstained with DAPI. Panels are representative of the ipsilateral hemisphere (A), contralateral hemisphere (B), and an immunostaining control (C). The scale bar represents 40 μm. To evaluate EGFR activation, H/I and control SVZs were dissected after 72 h recovery and the membranes were subfractionated (see the Materials and methods section) and 30 μg of each membrane fraction was analysed by Western blot for total EGFR and for phosphorylated EGFR (D). The Western blots were quantified and normalized to total receptor levels (E). Results are averaged from three independent experiments (n=6). *P<0.05 (as measured using a Student’s t test).
of neurospheres generated from the damaged hemisphere compared with controls (Alagappan, 2008). Therefore cells acutely dissociated from P12 mouse SVZ were stained for EGFR and LeX and analysed by flow cytometry. This analysis revealed a 27-fold increase ($P < 0.05$) in the number of EGFR$^+$/LeX$^+$ cells in ipsilateral SVZs compared with contralateral and sham-operated control SVZs (Figure 8) (5.59 ± 1.4 compared with 0.2 ± 0.04; 5.59 ± 1.4 compared with 0.2 ± 0.1; $n = 6$ brains). There was no statistical difference in the EGFR$^+$/LeX$^+$ population between the contralateral and sham-operated control SVZ cells (contralateral, 0.2 ± 0.04; control, 0.2 ± 0.1; $n = 6$ brains).

**Recovery from H/I increases EGFR intensity on SVZ stem/progenitors.**

To determine whether H/I also increases the amount of EGFR expressed per cell, we measured the geometric mean fluorescence intensity of EGFR labelling in the progenitor marker-positive as well as the LeX-positive stem cell population in the ipsilateral and contralateral hemisphere SVZs at 72 h post-H/I as compared with sham-operated control SVZs in rats and mice. In rats, EGFR$^+/progenitor^+$ ipsilateral SVZ cells expressed 3-fold greater ($P < 0.05$) EGFR intensity compared with contralateral and sham-operated control SVZ [ipsilateral, 46.3 ± 6.6 a.u. (arbitrary units); contralateral, 15.9 ± 2.2 a.u.; control, 9.9 ± 1.4 a.u.; $n = 6$ brains] (Figures 9A and 9B). In mice, EGFR$^+/LeX^+$ ipsilateral SVZ cells expressed 11-fold greater ($P < 0.05$) EGFR intensity compared with contralateral and sham-operated control SVZ (ipsilateral, 21.7 ± 3.4 a.u.; contralateral, 1.9 ± 0.5 a.u.; control, 1.3 ± 0.2 a.u.; $n = 6$ brains) (Figures 9C and 9D). Not surprisingly, in rats, EGFR$^+/progenitor^+$ cells in the ipsilateral SVZ showed a 1.5-fold increase ($P < 0.05$) in the EGFR intensity compared with the contralateral hemisphere and sham-operated controls (ipsilateral, 8.5 ± 0.9 a.u.; contralateral, 5.3 ± 0.5 a.u.; control, 3.9 ± 0.3 a.u.; $n = 6$ brains) (Figure 9C). In addition, LeX intensity was also affected by H/I. EGFR$^+/LeX^+$ ipsilateral SVZ cells expressed 17-fold greater ($P < 0.05$) LeX intensity compared with their counterparts in the contralateral and sham-operated control SVZs (ipsilateral, 6.8 ± 1.3 a.u.; contralateral, 0.4 ± 0.1 a.u.; control, 0.2 ± 0.02 a.u.).

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**Figure 6** H/I increases the number of EGFR-positive NG2-expressing neural precursors in the rat SVZ

(A) Dot plot of EGFR$^+/NG2^+$ cells from the ipsilateral, contralateral and control SVZs after 72 h recovery from H/I. (B) Quantification of the EGFR$^+/NG2^+$ neural precursors in the SVZ. (C) Quantification of EGFR$^+/NG2^+$ population, encompassing neural stem cells, in the SVZ after injury. Values are the mean number ± S.E.M. and are from one representative experiment ($n = 6$ animals per experiment). *$P < 0.05$, measured using one-way ANOVA and Tukey’s post-hoc tests. IL, ipsilateral; CL, contralateral; CTL, control.
There was no significant difference in LeX intensity or EGFR intensity between contralateral hemisphere and sham-operated control hemispheres at 3 day recovery from H/I (LeX intensity: contralateral, 0.4±0.1 a.u.; control, 0.2±0.02 a.u.; EGFR intensity: contralateral, 1.9±0.5 a.u.; control, 1.3±0.2 a.u.; n=6 brains) (Figures 8D and 8F).

EGFR overexpression is sufficient to recruit cells into the cell cycle and decrease their cell-cycle time.

To establish whether overexpressing EGFR is sufficient to recruit cells into the cell cycle and to increase the rate at which they traverse through the cell cycle, we compared BrdU incorporation into SVZ stem/progenitor cells transfected with a plasmid containing a constitutively active EGFR or a control vector. The constitutively active EGFR plasmid increased the number of cells in S-phase by 1.65-fold compared with controls (pEGFR-ires_EGFP, 16.6±1.6; pEGFP, 10.1±0.4; n=3 independent experiments) following a 6 h pulse with BrdU (Figure 10E). EGFR overexpression also increased the number of cells in G2/M-phase by 1.3-fold compared with control cells (pEGFR-ires_EGFP, 64.3±1.4; pEGFP, 49.3±4.3; n=3 independent experiments) in a 12 h period (Figure 10F).

A volumetric analysis of neurosphere size was performed by measuring the diameter of at least 30 neurospheres from each transfection condition per experiment. The size of the neurospheres obtained from the constitutively active EGFR-transfected cells were 5-fold and 32-fold larger (P<0.05) compared with EGFR-transfected cells after 3 days and 7 days in culture post-transfection respectively (3 day: 3160.9±2820.4 μm³; 7 day: 2154.9±2566.8 μm³; n=6 brains) (Figures 10C and 10D).

**DISCUSSION**

Many factors including TGF-α, FGF-2, insulin-like growth factor 1, platelet-derived growth factor, neurotrophins, ephrins, LIFR–CNTFR–gp130 complex, Notch 1, Sonic Hedgehog and Wnts regulate the formation of the SVZ from the ventricular zone in the developing brain (Carson et al.,...
1993; Vescovi et al., 1993; Weickert and Blum, 1995; Craig et al., 1996; Johe et al., 1996; Kuhn et al., 1997; Shimazaki et al., 2001; Doetsch et al., 2002; Giuliani et al., 2004; Hsieh et al., 2004; Israsena et al., 2004; Palma et al., 2005; Felling et al., 2006). These factors also regulate the proliferation and self-renewal of SVZ stem/progenitors and control the production of neurons and macroglia. Currently it has not been established which factor(s) among this pool of molecules is/are critically involved in the expansion and proliferation of the endogenous SVZ stem/progenitors for brain repair following injury. In the present study we show that following neonatal H/I: (i) quiescent SVZ cells enter the cell cycle; (ii) the increased proliferation of these precursors occurs in response to EGF, but not FGF-2; (iii) EGFR mRNA and protein expression increases in SVZ; (iv) EGFR expression increases on progenitors and stem cells within the SVZ; (v) EGFR acquisition increases the rate that SVZ cells traverse the cell cycle; and (vi) pharmacologically inhibiting EGFR, even in the presence of FGF-2, reduces the in vitro stem/progenitor expansion observed following injury, confirming that FGF-2 stimulation cannot compensate for the EGF-mediated regenerative response.

Both EGF and FGF-2 are mitogenic for adult neural stem/progenitors. SVZ cells express FGF-2, FGFRs, EGF and EGFRs, and signal transduction through these receptors is important for their proliferation and self-renewal during embryonic and postnatal development (Richards et al., 1992; Gritti et al., 1996; Johe et al., 1996; Kuhn et al., 1997; Doetsch et al., 1999; Gritti et al., 1999; Doetsch et al., 2002; Anton et al., 2004; Fox and Kornblum, 2005). Intraventricular injection of EGF expands the numbers of nestin$^+$ precursors in the SVZ (Craig et al., 1996) and forced EGFR acquisition confers migratory capacity to non-migratory postnatal neural progenitors (Aguirre et al., 2005). In parallel, studies on FGF-2 signalling have shown that FGF-2 is mitogenic for multipotential adult murine forebrain precursors, maintains the number of slowly dividing stem cells within the anterior SVZ and FGF-2 knockouts have reduced numbers of dividing neural stem cells and fewer olfactory bulb neurons (Gritti et
al., 1996; Zheng et al., 2004). Similar to the results of the present study, the expression of the EGFR increases within nestin+ cells located adjacent to ependymal cells during recovery from middle cerebral artery occlusion in adult rats, with peak expression occurring at 3 days post-ischaemia. Furthermore, infusing EGF for 7 days into the cerebral ventricles beginning 2 days after inducing ischaemia stimulated striatal neurogenesis 100-fold compared with vehicle-infused animals (Teramoto et al., 2003). Robust induction of FGF-2 and FGFRs has been documented following chronic hypoxia and this has been correlated with increased neurogenesis (Ganat et al., 2002; Fagel et al., 2006). However, we did not observe any increased cellular responses to FGF-2 using the neurosphere assay, nor did FGF-2 infusion stimulate enhanced neurogenesis after ischaemia in adult rats (Teramoto et al., 2003) Thus the precise role and requirement for FGF signalling in SVZ cellular responses to injury remain incomplete.

In the present study we evaluated whether functional activation of the EGFR is necessary and sufficient for the proliferative response initiated by stem/progenitor cells of the SVZ subsequent to brain injury. Studies from our laboratory, as well as others, have documented a regenerative response from the SVZ following neonatal H/I (Plane et al., 2004; Felling et al., 2006). During the first week of recovery from a moderate H/I insult, there is increased proliferation of cells in the most medial aspect of the SVZ, as revealed by increased BrdU uptake, as well as by increased numbers of nestin+/PCNA+ (proliferating-cell nuclear antigen) cells in the most medial aspect of the dorsolateral SVZ. This increase in cells in S-phase within the region of the SVZ that harbour the stem cells correlates with a doubling of the number of tripotential,

Figure 9 H/I increases the expression of EGFR-positive LeX-expressing putative stem cells in murine SVZ (A) Dot plot of EGFR+/LeX- cells from the ipsilateral, contralateral and control SVZs after 72 h recovery from H/I. (B) Quantification of the EGFR+/LeX- neural stem population in the SVZ after 72 h recovery from H/I. (C) Histogram plot of EGFR intensity within the LeX-positive cells from the ipsilateral, contralateral and control SVZs after 72 h recovery from H/I. (D) Quantification of the EGFR intensity (in a.u.) within the LeX-positive stem cell population in the SVZ after injury. (E) Histogram plot of LeX intensity from the ipsilateral, contralateral and control SVZs after 72 h recovery from H/I. (F) Quantification of the LeX intensity (in a.u.) of SVZ cells after injury. Values represent the mean number ± S.E.M. of three independent experiments with n=6 animals per experiment. *P<0.05, measured using one-way ANOVA and Tukey’s post-hoc tests. IL, ipsilateral; CL, contralateral; CTL, control.
self-renewing NSPs as measured using the neurosphere assay. Fate-mapping studies have shown that there is a subsequent emigration of neuronal and glial progenitors from the SVZ to repopulate subcortical and neocortical structures (Plane et al., 2004; Ong et al., 2005; Felling et al., 2006; Yang and Levison, 2006; Yang et al., 2007; Yang and Levison, 2007). However, the extent of stem cell expansion and the resulting regeneration following injury is modest. Therefore our most recent studies have focused on understanding the signals that regulate these processes towards achieving a more robust amplification of the stem/progenitor cell pool and more significant repair.

In a key experiment we abolished the in vitro expansion of SVZ NSPs by pharmacologically inhibiting functional EGFR activation in the presence of FGF-2. We used the quinazoline PD153035, a reversible tyrosine kinase inhibitor specific to EGFR. In vitro, PD153035 inhibits EGF-dependent cellular processes completely at doses ranging up to 1 μM whereas PDGF-, FGF-2-, and serum-dependent mitogenesis is inhibited only at concentrations higher than 5 mM (Fry et al., 1994). PD153035 is the parent compound of several 4-anilooquinazolines in the clinic such as Iressa (Astra-Zeneca), Tarceva (Roche/Genentech), GW 2016 (GlaxoSmithKline) and CI-1033.

**Figure 10** EGFR overexpression is sufficient to increase cell-cycle kinetics of SVZ NSPs. (A and B) pEF2-ATG-FL-EGFP and pEF2-FL-EGFR-IRES-EGFP plasmids. (C) Neurospheres generated at 3 days and 7 days following transfection with pEGFP and pEGFR-IRES-EGFP into SVZ NSPs. (D) Quantification of neurosphere size obtained from (C). (E) Percentage of cells in S-phase after a 6 h BrdU pulse and in G2/M-phase (F). Values represent the mean number ± S.E.M. of three independent experiments with n=6 animals per experiment. *P<0.05, measured using a Student's t test.

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Increased EGF responsiveness of neural stem cells after injury

(Pfizer) (Boschelli, 2002). In the present study, we used 300 nM PD153035 to block EGFR activation during stimulation with FGF-2. At this dose, we found that the number of neurospheres obtained in the presence of the inhibitor at 72 h post-H/I from the damaged ipsilateral hemisphere was comparable with sham-operated control SVZs indicating that FGF-2 signalling cannot compensate for the loss of EGFR signalling, and EGFR activation is necessary for the observed reactive expansion of the NSPs subsequent to an ischaemic insult. The above compounds gain access to the CNS, whereas PD153035 is excluded. It would have been informative to use these inhibitors for in vivo experiments; however, the manufacturers of these drugs denied our requests for these reagents.

Complementary studies to the loss-of-function experiment were performed using a novel technology to introduce a constitutively active EGFR into the NSPs. Acquisition of this functionally active EGFR enhanced the number of cells traversing the cell cycle. This was manifested as an increase in neurosphere size obtained from EGFR-overexpressing SVZ NSPs compared with GFP-transfected cells. Thus augmenting EGFR signalling is sufficient to recapitulate in vitro the increased regenerative NSP response observed ex vivo after injury.

A subset of tripotential neural progenitors in the postnatal brain expressing the NG2 proteoglycan can migrate to the subcortical white matter and to the olfactory bulb via the rostral migratory stream. Several studies have shown that acquisition of EGFR by neural progenitors correlates with their ability to proliferate and migrate (Aguiirre and Gallo, 2004; Aguiirre et al., 2005; Ivkovic et al., 2008). SVZ cells migrate into damaged white and grey matter after H/I as we, and others, have shown. These studies have implicated chemokines such as MCP-1 (monocyte chemoattractant protein-1) and SDF-1α (stromal-cell-derived factor-1α) as stimulators of progenitor cell emigration post-stroke (Miller et al., 2005; Yang et al., 2007). In light of the above studies substantiating a role for the EGFR in the migration of glial progenitors in remyelination, our results suggest a previously unanticipated role for the EGFR in progenitor cell migration post-ischaemic injury.

To establish whether EGFR acquisition following injury is conserved across species, and to determine whether it is induced on neural stem cells, we evaluated the expression of EGFR on LeX+/SVZ cells. LeX/SSEA-1 is a surface carbohydrate moiety expressed by a small population of mouse SVZ cells that retain sustained proliferation and self-renewal capabilities characteristic of SVZ stem cells (Capela and Temple, 2002; Maric et al., 2003; Imura et al., 2006). Upon injury, the numbers of EGFR+/LeX- cells increased significantly in the regenerating SVZ. We have also confirmed in the rat that the EGFR increases in the progenitor marker-negative population using the strategy developed by Maric et al. (2003) and have observed a similar increase of EGFR+ cells that are AldefluorTM+ in the neonatal SVZ after H/I (Alagappan, 2008). These results are consistent with our previous in situ and neurosphere analyses that indicate that the brain initiates an active expansion of SVZ stem cells, which is not simply an activation of pre-existing quiescent stem cells for repair.

We employed a cumulative thymidine incorporation assay to evaluate cell-cycle kinetics acutely following injury. This analysis affirmed an alteration in stem/progenitor behaviour upon EGF treatment. In this assay, the height at which the curve plateaus is a measure of the number of cells recruited into cell cycle and the slope of the curve provides an index of the cell-cycle time, with a steeper slope indicating a shorter cell-cycle time (Yu et al., 1996; Khanna, 2005; Khanna and Mehra, 2006). We specifically adopted cumulative thymidine labelling compared with pulse labelling because serum starvation, typically used to synchronize cells prior to cell-cycle analysis, is not technically feasible with NSPs. Averaging across experiments, approx. ten times as many cells from the injured brain entered the cell cycle and they divided significantly faster compared with uninjured controls in the presence of EGF. SVZ cells from the contralateral SVZ also divided more rapidly, analogous to previously reported neurosphere assay data (Felling et al., 2006). Taken together, these results suggest that acquisition of EGFR by the SVZ cells following injury not only expands the numbers of stem cells, but that it recruits quiescent cells to enter the cell cycle and stimulates these precursors to transit through the cell cycle faster compared with uninjured controls. These results are reminiscent of published studies in cancer biology that show that activation of EGFR destabilizes cell-cycle inhibitor proteins to enable quiescent cells to enter the cell cycle (Lenferink et al., 2000; Boerner et al., 2005).

In conclusion, the present study demonstrates that the EGFR is a key mediator of the regenerative response elicited from the stem cell and progenitor cell pools within the SVZ subsequent to brain injury. EGFR expression is elevated in SVZ cells, which recruits relatively quiescent cells to cycle more rapidly. The in vitro response of the SVZ stem/progenitors is manifested by an increase in neurosphere size and number, which can be abolished by a pharmacological inhibitor specific to EGFR in the presence of FGF-2 and is mimicked by overexpressing the EGFR. The increase in EGFR sensitivity acquired will promote their expansion and hence can participate in their migration into pallial and subpallial structures to aid repair. We also show that EGFR levels return to control levels within a few days, therefore strategies to augment EGFR signalling may represent important avenues to pursue to initiate a robust response from the resident SVZ NSPs following injury for functional brain recovery.

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