Validation and use of a computer-assisted counting procedure to quantify BrdU-labeled proliferating cells in the early postnatal mouse hippocampus

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**ABSTRACT**

The dentate gyrus is one of the few brain regions that show proliferation of neuronal precursors postnatally and in adult life. Proliferation in the dentate gyrus has been shown to be influenced by exercise, stress and drugs such as antidepressants. Traditionally, proliferation studies rely on the time consuming and subjective manual count of labeled cells. Here we adapted the Metamorph software to automatically count cells labeled in the S phase in the developing dentate gyrus of mice. The validity of the computer-assisted method was established by showing an outcome similar to that obtained with the established manual counting procedure. In addition, by using a genetically modified mouse line with increased proliferation, the ability of the computer-assisted method to detect changes in proliferation was demonstrated.

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1. Introduction

The hippocampus has long been known to be involved in spatial learning and memory and recent studies also identified its role in anxiety (Bannerman et al., 2004; Squire et al., 2004). The dentate gyrus is the only hippocampal subregion that develops postnatally and which shows neurogenesis in the adult brain (Kempermann et al., 1997, 2004). During early postnatal development, neuronal progenitors proliferate in the tertiary matrix located in the hilus and in the subgranular zone (SGZ) of the dentate gyrus (Altman and Das, 1965). Both the hilar and SGZ cells migrate outward to form the granule cell layer (GCL). The SGZ continues to give rise to new neurons during adult life (Kempermann et al., 2004). Interestingly, adult neurogenesis is increased by exercise (Cotman and Berchtold, 2002) and following chronic antidepressant treatment in mammals (Dulawa et al., 2004; Malberg et al., 2000). Stress on the other hand reduces neurogenesis (Duman, 2003; Malberg and Duman, 2003).

Proliferation in the dentate gyrus is assessed by counting cells pulse labeled by bromodeoxyuridine (BrdU) followed by the visualization of labeled cells using anti-BrdU antibodies in histological sections. Traditionally cells are manually counted. However this procedure is labor intensive limiting the scope of studies. Manual counting is a daunting task especially when the number of immunopositive cells is relatively high in the area of interest such as in the developing dentate gyrus. Image analysis software with interactive definition of the region of interest and threshold are available and have been used to count cells in various biological systems (Cunnane et al., 1999; el-Salhy et al., 1997; Goedkoop et al., 2005; Markey et al., 2003; Taylor and Levenson, 2006; Went et al., 2006). Here we tested the Metamorph software and its cell counting module for counting S-labeled nuclei in the developing dentate gyrus of mice in various experimental paradigms. The computer-assisted counting method is fast but still has the accuracy of the traditional manual counting method.

2. Materials and methods

2.1. Animals

Four groups of animals were used in these studies: Wild-type (WT), serotonin (5-HT) 1A receptor (5-HT1A R) homozygote knock-out (KO) (Parks et al., 1998), Ink4a homozygote KO (Kim and Sharpless, 2006) and 5-HT1AR/Ink4a double KO mice. These lines were originally used in a study that analyzed the effect of genetically inactivating either the 5-HT1AR, the Ink4a or both genes on cell proliferation (Gleason and Toth, unpublished). All mice were on the mixed (50–50%) Swiss Webster × FVB genetic background.

2.2. BrdU immunohistochemistry

Postnatal day (P) 7 pups received 1 subcutaneous 100 mg/kg injection of 10 mg/ml BrdU (Sigma, St. Louis, MO). We selected...
pups from at least 3 different litters because development during the first postnatal week in mice can show substantial individual variability within and between litters. Factors could include the time of delivery, size of the litter and postnatal maternal care. This variability may be reflected in the total number of proliferating cells as proliferation peaks during the first week of life and a slight delay or acceleration in proliferation in individuals could change the number of BrdU positive cells labeled on P7. Therefore, we tried to minimize the variability in individual development by excluding pups that deviated by more than 0.5 g from the 4.5 g average weight.

Three hours after the administration of BrdU mice were transcardially perfused under deep anesthesia with heparinized saline followed by 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer. Brains were dissected out, postfixed overnight in fresh 4% PFA, cryoprotected in 30% sucrose in 0.1 M phosphate buffer, and sectioned in the coronal plane at 40 μm on a freezing sliding microtome. Every fourth section throughout the hippocampus was processed for BrdU immunohistochemistry (Pham et al., 2005). Briefly, sections were mounted on slides and microwaved in preheated 0.1 M citrate buffer. Next, sections were incubated in 0.6% H2O2 in phosphate-buffered saline for twenty minutes, permeabilized in 0.1 M Tris buffer containing 0.1% trypsin with 0.1% CaCl2 and finally treated with 2 N HCl in phosphate-buffered saline. Then, sections were blocked with 3% normal horse serum and 0.3% Triton X-100 and incubated at 4 °C with mouse monoclonal anti-BrdU (1:200; Novocastra, Newcastle upon Tyne, UK) overnight. The next day, sections were incubated at room temperature with the secondary antibody (biotinylated horse anti-mouse; Vector Laboratories, Burlingame, CA) for 1 h, incubated at room temperature with an avidin–biotin complex (Vector Laboratories) for 1 h, and then visualized with 3,3′-diaminobenzidine (Sigma, St. Louis, MO). Finally, slides were Nissl-stained with cresyl violet, dehydrated with an ethanol series, cleared in xylene, and coverslipped using Cytoseal mounting medium.

### 2.3. Computer-assisted counting

Sections were derived from a total of 16 mice corresponding to 4 different genotypes (see above). In some experiments 44 sections (2–3 sections per animal) were used by randomly assigning them to 3 groups of 16, 16, and 12 sections. These series were counted on separate days. In other experiments, BrdU positive cell number was analyzed by the genotype. In this case, the total number of positive cells per animals was measured by counting cells in every fourth section and then multiplying the result by 4.

Images were captured by a blinded observer (at 10× magnification) by using a Nikon TE200 inverted fluorescence microscope with an attached Nikon Coolpix 995 digital camera. The ideal aperture was determined to be 5.0 mm, and the best exposure time was determined to be 1/30 of a second. These settings were fixed and maintained throughout the duration of the image-taking process. Since the size of the dentate gyrus increases from the dorsal to ventral axis, some of the ventral sections required multiple images to be taken. These images were merged using Adobe Photoshop. Images were imported into the MetaMorph software program (Molecular Devices, Downingtown, PA). Since the SGZ in P7 mice is not yet completely restricted to the innermost 1–2 cell layers of the GCL, BrdU positive cells were counted in the entire GCL. The GCL was marked by a line outlining this area. Then the threshold tool was used to isolate immunopositive cells. The average area for single cells was 50 pixels and we limited the counting to objects larger than 20 pixels. An upper limit was imposed as well; an object was not included in the cell count if its standard area count was more than 7 cells, or 350 pixels. Although peroxidase was used to reduce nonspecific staining, blood vessels occasionally showed staining (Fig. 1). However, these objects were larger than 350 pixels, the upper limit of our counting, and were automatically excluded. Once the images were thresholded the number and size of positive objects and the corresponding cell numbers (size of the cluster in pixel/50 pixel) were automatically registered and logged to an Excel spreadsheet.

### 2.4. Manual counting with stereological analysis

BrdU-labeled cells were manually counted in each section using a Nikon microscope with the Stereoinvestigator software (Microbrightfield Inc., Colchester, VT) by a blinded observer as described previously (West and Gundersen, 1990) but instead of using the “fractionator” function, all cells were counted in the GCL. A 40× objective lens was used for all counting.

### 3. Results

While Metamorph, a software with computer-assisted quantification capabilities, has been used in numerous immunological studies (Schacker et al., 2002; Zhang et al., 1998) (see Section 1), its potential as a program for counting S phase cells in the brain has not been explored. First, we tested if the computer-assisted counting of BrdU positive cells in the developing dentate gyrus of mice has accuracy comparable to that of the manual counting technique (cross-method correlation and inter-trial accuracy).

#### 3.1. Cross-method correlation

Forty-four dentate gyrus sections from P7 mice (Fig. 1), previously counted manually, were randomly assigned to three groups of 16, 16, and 12 sections, each (see Section 2). The smaller group size was chosen to test the computer-assisted method with a group size more realistic in experimental settings. Metamorph was used to determine the number of cells in the GCL (ANOVA showed no significant difference between the three groups; $F_{2,41} = 0.946, P = 0.40$). Scatter plot analysis of the manual and computer-aided counts showed a significant correlation...
in all three groups ($P=0.0042$, $0.00019$, and $0.0019$ for groups 1–3, respectively, Fig. 2A–C). Furthermore, the standard deviation and error of all measurements (groups 1–3) by the manual and computer-assisted techniques were similar (62.61 ± 4.36 and 68.55 ± 4.41; mean ± S.D., respectively) indicating that variability in the two techniques is comparable. These data validate the computer-assisted counting as comparable to the manual counting method.

3.2. Inter-trial accuracy of the computer-assisted counting

Seventeen images were randomly selected and the number of BrdU positive cells in the GCL was counted in two separate trials by the computer-aided method. Scatter plot analysis showed a high level of correlation between the data collected in two different days ($P=0.00022$; Fig. 3).

3.3. Detection of genotype-dependent differences in cell number by the computer-assisted counting method

The computer-assisted method was further tested by using genetically manipulated mouse lines. Since genotype differences were studied in these experiments, the total number of BrdU positive cells per animal (rather than the number of cells per section) was compared in the GCL (see Section 2). ANOVA of the manual counts of the four groups showed a genotype effect (ANOVA: $F_{3,12}=11.76$, $P=0.0007$, $N=4$ per group). LSD post hoc analysis showed that 5-HT$_{1A}$R/Ink4a double KO mice had more BrdU positive cells than WT, 5-HT$_{1A}$R KO and Ink4a KO mice (Fig. 4A). Then, the same sections were counted with the computer-assisted techniques which yielded a similar result (ANOVA $F_{3,12}=4.92$, $P=0.018$, $N=4$ per group) with significant differences between the 5-HT$_{1A}$R/Ink4a double KO mice and WT and Ink4a KO mice (Fig. 4B). The difference between the double KO and 5-HT$_{1A}$R KO mice was only a trend indicating that the computer-assisted method may not have the sensitivity of the manual counting technique to detect genotype-dependent differences. Nevertheless, the computer-assisted method correctly indicated the overall difference in the sample and the differences between groups.

4. Discussion

Traditionally, BrdU pulse-labeled neuronal precursors are counted manually. Since this procedure is time consuming, it is important to develop and validate techniques which maintain a similar accuracy as manual quantification while significantly reducing the amount of time involved. Here we show that MetaMorph can be used to count BrdU positive cells in a time-efficient way. Indeed, at least six 4–5 h sessions are necessary to complete a stereological analysis that involves two groups, each consisting of 4–6 animals, each sampled by three sections through the hippocampus (approximately 45 min are required for a single section). The same series can be analyzed in one 4–5 h session as image acquisition takes only a few minutes each and highlighting of the area of interest and threshold setting, once the images are uploaded to the MetaMorph software, take another few minutes. The difference in time between the two methods can be even bigger because normally 4–6 sections are analyzed per animal and because in some experiments more than 4–6 animals/group are used.

The computer-assisted method is particularly useful when a large number of cells are present such as during develop-
ment because the computer-aided technique allows obtaining total cell counts directly circumventing the need to extrapolate from sample areas commonly used in manual counting (i.e., with the fractionator function of StereoInvestigator). The computer-aided quantification method is less prone to subjectivity because it selects and highlights cells automatically based on red, green, and blue color intensities and size. Our data showed a good correlation between the manual and computer-assisted counting methods and was able to detect genotype-dependent differences between groups of animals in BrdU positive cell number.

Although the computer-assisted method is significantly faster than the manual technique, it has limitations. First, although typical stereology utilizes a top and bottom guard zone of the section to avoid double counting of cells present on two adjacent sections (West and Gundersen, 1990), this is difficult to implement for automated counting. Instead, we limited the size of objects to be counted to at least 20 pixels while a typical nucleus is 50 pixels. This resulted in registering cells with nuclei of at least 20 pixels but excluded cells with less than 20 pixels on both surfaces resulting in a combined effect of an overestimation of cell numbers. Considering the thickness of the slides and the diameter of cell nuclei and assuming an even distribution of cells through the rostro-caudal axis within individual sections, the overcounting was estimated to be 5.83% by the computer-assisted method. However this bias does not invalidate our results because the main question in our experiments, like in most experimental designs, was the relative difference in cell number between the experimental and control groups rather than the absolute number of cells in individual groups. If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups). If absolute count is required, the computer-assisted count could be normalized (in absolute number of cells in individual groups).

Second, the cell cluster size has to be adjusted according to the experimental conditions when using the automated method. Since a single pulse of BrdU labeling was used in our experiments, we did not encounter the problem of cell clusters bigger than seven cells (Fig. 1). In fact the size of the biggest clusters was usually 3–4 cells in our experiments because multiple cell divisions of a single-labeled progenitor will dilute the BrdU label in daughter cells below the detection limit. However, if multiple BrdU injections are used, especially if these injections extend over several cell divisions, larger than seven cell clusters may be formed. In this case either the labeling should be limited to a shorter period or the upper limit of cluster size should be increased. Despite of these limitations, we demonstrate here that the computer-aided technique, if conditions are selected correctly, can significantly expedite quantification of BrdU positive cells in neuronal tissue.

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