Introduction

In most brain regions, the number and relative density of neurons of each type vary remarkably little among individuals of the same animal strain (Williams and Herrup, 1988). This tight regulation controls the size of the different brain regions and allows the formation of proper neural connections. Cell death plays a key role in regulating the number of neurons in the brain from its earliest stages of development. Death has been described among proliferating neuroblasts, early postmitotic cells and in many populations of neurons at the time they form synaptic connections (reviewed by Davies, 2003; de la Rosa and de Pablo, 2000; Lossi and Merighi, 2003; Oppenheim, 1991; Pettmann and Henderson, 1998). A long established idea is that the survival of developing neurons depends on trophic factors produced in limited amounts, but recent studies have shown that neuronal death may also partly be due to the activation of cytotoxic signaling mechanisms, through cell receptors such as FasL, TNF-R and P75 (reviewed in Davies, 2003; Dechant and Barde, 2002; Raoul et al., 2000).

A conserved mechanism of cell death activation has been described in non-neuronal cells (Falzoni et al., 1995; Girolomoni et al., 1993; Koshlukova et al., 1999), whereby extracellular ATP (e-ATP) triggers cell death by binding the P2X\textsubscript{7} receptors. Upon sustained or repeated activation, the P2X\textsubscript{7} receptors induce large non-selective membrane pores, which eventually lead to cell death (Di Virgilio et al., 1998; North, 2002).

P2X\textsubscript{7} receptors have been shown to be expressed in regions of the nervous system (North, 2002) and, therefore investigations were undertaken to determine whether endogenous e-ATP is involved in controlling the number and density of cells in specific neural populations. The retina is an ideal region to test this, since most of its neurons are arrayed with precisely controlled density, and there is a prominent expression of P2X\textsubscript{7}. Among the developing retinal neurons, the cholinergic neurons seemed particularly relevant, because they are regularly spaced from early in development (Galli-Resta et al., 1997), and play a key role in normal development by triggering the waves of spontaneous neuronal impulse activity controlling the refinement of retinal projections to the brain (Wong, 1999).

We found that the retinal cholinergic cells express the P2X\textsubscript{7} receptors, and could be sources of endogenous e-ATP. Degrading e-ATP in vivo, or blocking the P2X\textsubscript{7} receptors increases the density of the cholinergic neurons in the developing retina by preventing their naturally occurring death. This was also confirmed by directly monitoring e-ATP-induced death of individual cholinergic neurons in isolated retinas. Death induced by e-ATP in the retina is specific to the cholinergic neurons, normally removing cholinergic cells too close to one another, and thereby contributing to the regular density and spacing of these neurons.

Summary

The precise assembly of neuronal circuits requires that the correct number of pre- and postsynaptic neurons form synaptic connections. Neuronal cell number is thus tightly controlled by cell death during development. Investigating the regulation of cell number in the retina we found an ATP gated mechanism of neuronal death control. By degrading endogenous extracellular ATP or blocking the P2X\textsubscript{7} ATP receptors we found that endogenous extracellular ATP triggers the death of retinal cholinergic neurons during normal development. ATP-induced death eliminates cholinergic cells too close to one another, thereby controlling the total number, the local density and the regular spacing of these neurons.

Key words: Retina, Cell death, ATP, P2X\textsubscript{7}, Amacrine cells, Mosaics, Rat

Materials and methods

Animal and tissue treatment

Experiments were conducted on Long Evans hooded rats in accordance with national legislation and ARVO regulations. Animals were bred in the laboratory. Intraocular injections in newborns anaesthetized with ether were performed as described previously (Galli-Resta et al., 2002). Intraocular concentrations were determined assuming a 10 µl vitreal volume at P1, as previously estimated (Galli-Resta et al., 2002). Bromodeoxyuridine (BrdU), which is incorporated during DNA synthesis (Gratzner, 1982), was injected intraperitoneally (i.p.; 50 mg/kg body weight) every 6 hours starting immediately before the treatment in order to label all the cells entering S-phase after treatment. The S-phase in the retina is 12-18 hours at this
stage (Alexiades and Cepko, 1996). Animals were sacrificed by decapitation. Eye dissection, fixation, retinal flat-mounting or sectioning, and immunocytochemistry were performed as described previously (Galli-Resta and Ensini, 1996).

To control apyrase effectiveness in reducing e-ATP levels in vivo, we collected 5 µl fluid samples from the posterior eye chamber, reasoning that changes in ATP levels in this fluid would parallel those in the adjacent retina. P4 rats were anesthetized with avertin (10 ml/kg body weight, i.p.; 3.3% tri-bromo-ethanol, 2% tertiary amyl-alcohol in saline) 3 hours after apyrase or vehicle intraocular injection.

Membrane permeability was assessed by adding propidium iodide (PI; 1.5 µM, for 1 minute) to the ACSF. Exposure of phosphatidylserine on the outer membrane leaflet was assessed with Cy3-annexin V (3 µg/ml, for 20 minutes). Images were acquired with the FITC (488 nm) filter setting for Oregon-Green-488-dextran, with the TRITC filter setting (568 nm) for PI, Cy3-annexin V and Alexa-Fluo-568-dextran.

Quinacrine (1 µM) was incubated for 10-20 minutes at room temperature and imaged using a Leica TCNS confocal microscope, keeping the laser at minimal power (5-10%). Since quinacrine easily bleaches, a maximum of two cells were scanned per retina, setting the scanning depth for each cell under the TRITC filter (displaying dextran labeling), then scanning with the FITC filter (quinacrine), and finally with the TRITC filter.

Data acquisition and analysis

The cholinergic cell arrays were sampled using a Leica TCNS confocal microscope. Four 400×400 µm² samples of either cholinergic cell arrays were taken in each whole-mount retina at mid-eccentricity along four perpendicular axes. The density of cholinergic cells does not vary with eccentricity at these ages (Galli-Resta et al., 2002). The same proved true in a dedicated analysis of the treated cases (eight samples per retina taken at two different eccentricities along four perpendicular axes, two retinas per treatment; not shown).

Since in each case the objective was positioned midway between the papilla and the retinal margin without prior examination of the cell distribution, each sampling was unbiased. Sampled fields and retinal images were examined using an Image analyzer (Imaging Ontario, Canada) to determine cell density, cell positioning and retinal area. Cell counts and coordinates were obtained by feeding each sample field to the Imaging system and using an automatic cell counter based on intensity threshold and size exclusion criteria to eliminate noise. Total numbers of cells were estimated multiplying average cell density and retinal area. To investigate death among the cholinergic neurons, the sampled fields were screened for cellular debris retaining immunoreactivity for ChAT. ChAT-positive (ChAT+) debris within 15 µm of one another were counted as a single occurrence. Pycnotic cells in the GCL were counted in four samples (250×250 µm²) per retina, after staining with propidium iodide. To sample non-cholinergic cells in the ganglion cell layer (GCL), whole-mount retinas were immunostained for ChAT (FITC; green), then labeled with PI (TRITC). The entire thickness of the GCL was scanned with a confocal microscope under the FITC+TRITC filter setting, taking eight samples (250×250 µm²) per retina at two regularly spaced eccentricities along four perpendicular axes. All the PI cells that were not labeled for ChAT were then counted using Metamorph. Horizontal cells (tau-immunoreactive), dopaminergic amacrine (inner nuclear layer (INL) cells immunoreactive for tyrosine hydroxylase (TH)) and AII amacrine cells (Disabled 1 immunoreactive) were also counted: horizontal cells were sampled in four (400×400 µm²) samples. TH cells were sampled in eight regularly spaced (400×400 µm²) samples limited to the dorso-temporal retina where these cells are restricted early in life (Wu and Cepko, 1993); AII cells were sampled in four mid-eccentricity (250×250 µm²) fields along four perpendicular axes.

Plots, frequency histograms and statistical analysis were made using Origin 7.0 (Microcal). Custom made programs were used to compute the autocorrelation, the density recovery profile (DRP), and the frequency of cells closer than 15 µm to one of their neighbors. The DRP and the exclusion radius (ER) are computed as follows. Concentric circles are traced at constant distances (here 2.5 µm) from one another in the autocorrelation, and a histogram (DRP) is obtained of the density of autocorrelation counts in the annuli delimited by two consecutive circles. The DRP values are very low close to the center of the autocorrelation (reflecting the central hole), and then rise to a final constant density in the bins far from the origin. The ER is obtained as the radius of the first circle (FC) where the histogram reaches or exceeds its average final density minus a correction weighing the autocorrelation counts found in the annuli contained
within FC. This correction is computed as the density of counts within the FC circle divided by the average DRP plateau value reached away from the origin, and multiplied for the histogram bin size (Rodieck, 1991). To evaluate the frequency of cell pairs with an intercellular distance below 15 µm, data from retinas treated with apyrase and oxidized ATP (oATP) were pooled together. The frequency of cell pairs was computed as the percentage of the total number of cell pairs in the field. In a field with n cells there are \( n^2(n-1)/2 \) pairs of cells.

Reagents
Fluorescent conjugated dextran (10 kDa, conjugated with Alexa Fluo 488 or 568 or Oregon Green 488), and fluorescent conjugated secondary antibodies were from Molecular Probes, Eugene, OR. Suramine was from Calbiochem. Antibodies to choline acetyltransferase (ChAT) and tyrosine hydroxylase (TH) were from Chemicon. The BrdU monoclonal antibody was from Roche. The P2X7 polyclonal has been described previously (Ferrari et al., 1997); the A8 polyclonal to Islet 1/2 was a kind gift from T. Jessell; the antibody to Disabled 1 was a kind gift from B. Howell. All other chemicals were from Sigma.

Results
The rat retina has two arrays of cholinergic neurons, positioned in two separate layers (GCL and INL). To test whether e-ATP controls the death of these neurons during normal development, we injected postnatal day 1 (P1) rats intraocularly with blockers of the P2X purinergic receptors or with an ATP degrading enzyme, and analyzed the number of retinal cholinergic neurons in these animals 24 hours later (P2). Irreversible or slowly reversible agents were used because alterations in the rate of cell death must occur for some time to affect the total number of cells.

Extracellular ATP participates in the developmental control of the density and number of cholinergic amacrine cells
Neonatal rats subjected on P1 to intraocular injection of oxidized-ATP (oATP, 300 µM), an irreversible blocker of the P2X purinergic receptors (Murgia et al., 1993; North, 2002), displayed on P2 (i.e. 24 hours after the injection) a significant increase in the density of retinal cholinergic neurons (Fig. 1A,B). This was about 23% higher than normal for the cholinergic cells in the GCL, and 22% for the cholinergic cells in the INL (n=8 retinas; Fig. 1C,D). A similar density increase (+26% in the GCL, +28% in the INL; n=8 retinas) was observed 24 hours after intraocular injection of suramine (150 µM; Fig. 1C,D), a generic blocker of the purinergic P2 receptors (Ralevic and Burnstock, 1998), as well as after injections of apyrase (30 U/ml, +30% GCL, +21% INL; n=8 retinas; Fig. 1C,D), an enzyme that hydrolyses e-ATP (Komoszynski and Wojtczak, 1996). In all cases the increase in the density of cholinergic neurons was statistically significant (P<0.0001, t-test).

An increased cell density was not a generalized effect following e-ATP blockade, since we did not observe any change in retinal area (not shown), in the thickness of the retinal layers, or in the density of a number of cell populations that we analyzed as a control. These populations include the non-cholinergic cells in the GCL [mostly retinal ganglion cells at this age (Perry et al., 1983; Rabacchi et al., 1994)], the horizontal cells, the long-range dopaminergic amacrine cells and the short-range AII amacrine cells (Table 1; Fig. S1 in supplementary material).

Since retinal areas were not affected by treatments blocking e-ATP signaling, the observed increase in the density of cholinergic neurons reflected true increments in the total number of these cells in both cholinergic arrays (Fig. 2A,B). An increase in the number of cholinergic cells was also observed 24 hours after treatments administered on P2 or P3 but not on P5 (Fig. 2A,B).

e-ATP causes death of the cholinergic neurons in normal retinal development
Since the cholinergic neurons found in treated retinas outnumbered the maximal number of these cells found in normal development (Fig. 2A,B), an accelerated migration of the cholinergic neurons to their layers could not account for the observed effects. New cell genesis was also excluded, since bromodeoxyuridine (BrdU), administered to label all the progenitor cells synthesizing DNA in the interval between treatment and analysis, was not detected in any of the 3000
cholinergic cells analyzed in sections from five treated retinas (Fig. 2C).

e-ATP can activate cell death in different non-neural cell populations (Di Virgilio et al., 1998). Investigations were therefore carried out to determine whether blocking e-ATP increased the number of cholinergic cells by preventing cell death among these neurons. In line with this hypothesis we found that cellular debris expressing cholinergic markers (ChAT+) had an average frequency of 3.7±1.4 every 1000 cholinergic cells in normal retinas (n=8 retinas), and decreased after e-ATP blockade, being 0.8±0.6 in oATP-treated retinas (n=8) and 0.7±0.6 in apyrase-treated retinas (n=8). This decrease was statistically significant (P<0.001, t-test) in both cases. Conversely, we observed up to a sixfold increase in the occurrence of ChAT+ debris (20±10 per 1000 cholinergic cells; n=3 retinas) 90 minutes after injection of 5 mM ATP into the eye. These changes in the frequency of ChAT+ cellular debris are consistent with the hypothesis that eATP regulates the basal level of cell death among the cholinergic neurons. e-ATP blockade did not affect the global occurrence of pycnotic cells in the GCL which was 6.5±2 every 1000 GCL cells in normal retinas (n=4) and 7.0±2 in oATP-treated retinas (n=4).

To test directly whether ATP kills the cholinergic neurons, we monitored the fate of individually labeled cholinergic cells after the application of ATP or other agents in isolated neonatal retinas aged between P2 and P8. Since experiments measuring endogenous e-ATP release in the rat retina have detected local eATP concentrations around 0.1 mM (Newman, 2001), we challenged cholinergic cells with this or higher concentrations of ATP. Developing cholinergic cells displayed dendritic and soma blebbing within minutes of ATP application (20/20 cells with 0.1 mM ATP; 5/5 cells with 0.5 mM ATP; 25/25 cells with 1 mM ATP; Fig. 3A-E). P2X7 receptor activation induces membrane blebbing, membrane permeabilization to large cations such as propidium iodide (PI), and eventually leads to cell death (Di Virgilio et al., 1998; North, 2002). ATP-induced permeability to PI was observed in all individually identified cholinergic cells we tested (33/33 cells; Fig. 3F). Furthermore, Cy3-annexin V revealed phosphatidylserine exposure on the external leaflet of the cytoplasmic membrane (10/10 cells; right inset in Fig. 3C), a typical early indicator of the activation of a death process (Reutelingsperger et al., 2002).

Within 2-12 hours after ATP application the cell soma shrunk (8/8 cells; Fig. 3G). In most cases dendritic blebbing increased with time (e.g. Fig. 3A-C), then persisted for hours (3-12 hours) after ATP application. In a few cases, the cells shed their blebs and became very faint. 2,3′-benzoyl-4-benzoyl-ATP (BzATP, 100 μM), a potent agonist at the P2X7 receptors induced the same effects as did ATP, but with a faster dynamics, so that after extensive blebbing, all blebs were shed 30 minutes after application, leaving faintly labeled cells displaying intense PI nuclear staining (6/6 cells; not shown).

The effects of ATP application on the cholinergic cells were the same throughout the age range we tested (P2-P8; e.g. Fig. 2).
ATP-gated neural death in developing retinas

Any detectable effect of ATP (1 mM) was prevented by the simultaneous incubation with apyrase (30 U/ml; 16/16 cells; Fig. 4C,D). Similarly, 2 hours of pre-incubation with the irreversible P2X antagonist oATP (300 µM) prevented blebbing and PI permeability (ATP 1 mM; 11/11 cells; Fig. 4E,F), even for ATP incubations as long as 2 hours. Finally, 30 minutes to 1 hour pre-incubation with Brilliant Blue G [BBG; a selective antagonist of rat P2X7 when used below the micromolar range (North, 2002)], prevented blebbing and membrane permeability induced by ATP (1 mM...
for 30 minutes) in 95% of the tested cells (10/10 cells BBG 0.5 μM; 9/10 cells BBG 0.2 μM; Fig. 4G,H). The protective effects of BBG were reversible, since 1 mM ATP application after BBG washing induced membrane blebbing and permeabilization (5/5 cells BBG 0.5 μM; 5/5 cells BBG 0.2 μM; 10-minute washing followed by 1 mM ATP for 30 minutes; Fig. 4I). Membrane blebbing, loss of membrane integrity or annexin V labeling were never observed in 30 individually labeled cholinergic control cells, subjected to the same light exposure as the treated cells.

**The cholinergic neurons express the P2X7 receptor and are potential sources of e-ATP**

The previous findings suggest that the P2X7 receptors are a major pathway through which extracellular ATP triggers cholinergic cell death. Double immunostaining showed that the cholinergic cells express these receptors from P0 (Fig. 5A).

To search for e-ATP sources in the neonatal retina we used quinacrine, a vital green-fluorescent dye labeling high levels of ATP bound to peptides in large granular vesicles (Bodin and Burnstock, 2001). In P2-P8 rat retinas where individual cells had been labeled with Alexa-Fluor-564-dextran (red) to allow cell identification, we observed quinacrine labeling in the astrocytes of the central retina (not shown) and in all the cholinergic cells we had labeled (10/10 cells; Fig. 5B-D), while none of the individually labeled non-cholinergic amacrine cells or RGCs (0/25 cells) contained quinacrine (not shown). These data suggest (but do not prove) that the cholinergic cells could be endogenous sources of e-ATP in developing retinas, in accordance with previous observations (Santos et al., 1999).

**Death induced by e-ATP eliminates cholinergic cells getting too close to one another**

We reasoned that if the cholinergic cells were able to release e-ATP, e-ATP-induced death should mostly affect cholinergic cells close to one another, because e-ATP spreads only short distances (<50 μm) away from its source (Newman, 2001). In agreement with this prediction, we found a higher than normal frequency of cholinergic cells very close to one another after e-ATP blockade: while in vehicle injected control retinas the frequency of cholinergic cells very close to one another after agreement with this prediction, we found a higher than normal distances (<50 cells close to one another, because e-ATP spreads only short e-ATP, e-ATP-induced death should mostly affect cholinergic cells. We reasoned that if the cholinergic cells were able to release e-ATP, e-ATP-induced death should mostly affect cholinergic cells very close to one another after agreement with this prediction, we found a higher than normal.

**Fig. 5.** The cholinergic neurons express the P2X7 receptors and accumulate ATP in granules, which suggests that they could be sources of e-ATP. (A) In the neonatal rat retina the cholinergic cells (red) express the P2X7 receptors (green), as do many retinal ganglion cells. Confocal image of a P4 retinal section. (B-D) Quinacrine, which labels high levels of ATP stored in granules, is found in the cholinergic cells. (B) Confocal image of a P8 rat cholinergic neuron labeled with Alexa Fluor 568 dextran. (C) A higher magnification of the same cell shows that its soma (red) contains quinacrine labeling (green), which is also shown in isolation in D. Scale bar: 20 μm (A); 40 μm (B); 10 μm (C,D).

**Discussion**

We have shown that degrading endogenous extracellular ATP (e-ATP) by intraretinal injections of apyrase significantly increased the density and number of retinal cholinergic neurons in neonatal rats. The same effect was obtained with the purinergic antagonist suramine, or with oATP, a selective irreversible blocker of the P2X purinergic receptors (North,
New cell genesis is excluded, since BrdU, which is incorporated into the DNA of proliferating cells, was not detected in any cholinergic neurons after e-ATP blockade. Accelerated cell migration is insufficient, as after e-ATP blockade we observed more cholinergic cells than are ever observed in the normal rat retina. Thus, e-ATP blockade must hamper a mechanism which normally reduces the number of cholinergic cells during development.

In principle the number of cholinergic cells can be reduced by inducing cell death among these neurons, or by promoting their trans-differentiation into a different type of cell, or by acting at both levels. Since e-ATP has been shown to trigger cell death in different non-neural cell populations (Di Virgilio et al., 1998) we first tested the effects of ATP on cholinergic cell death. In vivo we found that blocking e-ATP reduced the occurrence of cellular debris immunoreactive for cholinergic markers, while a pulse of exogenous ATP in the eye increased the cell debris. Furthermore, direct application of ATP in living isolated retinas rapidly induced clear signs of the activation of a death process in individually identified cholinergic neurons. These signs included membrane blebbing, loss of membrane integrity, exposure of phosphatidylserine on the external leaflet of the cytoplasmic membrane, and, finally, nuclear shrinkage. These effects were never reversible. Initially, we applied 1-0.5 mM ATP to isolated retinas, considering that the effective concentration at the tissue level was likely to be much lower, for the rapid degradation of ATP by endogenous ectonucleotidases (reviewed by Zimmermann, 1996). However, cholinergic cell death was also induced by 0.1 mM ATP, corresponding to typical concentrations observed close to the endogenous sources of ATP in the rat retina (Newman, 2001). The death-inducing effects of ATP were totally prevented by the simultaneous application of apyrase, which degrades e-ATP (Komoszynski and Wojtczak, 1996), or by preincubation with oATP, an irreversible P2X blocker (Murgia et al., 1993; North, 2002). Finally, Brilliant Blue G ensured total protection of 95% of the tested cells, at concentrations that selectively block the P2X7 receptors (North, 2002). Taken together these data indicate that e-ATP can kill the cholinergic neurons, and does so during development.

The possibility that e-ATP could also control the cholinergic cell number by inducing a transdifferentiation of postmitotic cholinergic neurons into non-cholinergic cells was discounted. To our knowledge transdifferentiation of postmitotic neurons has never been observed in the retina, where cell fate appears to be determined around the time of the last cell division (Livesey and Cepko, 2001). Furthermore, we never observed any indication of mix-phenotypes (e.g. cells immunopositive for cholinergic and non-cholinergic markers) in the retina of e-ATP-blocked eyes.
for ChAT but not displaying features typical of the cholinergic amacrine cells such as soma size, dendritic arrangement, planar organization, Islet immunoreactivity, or displaying features typical of non cholinergic cells) in vivo in normal retinas or after eATP blockade. Finally, direct application of ATP in concentrations normally observed in the rat retina (Newman, 2001) never induced anything but typical cell death indicators in individually labeled cholinergic cells in isolated retinas.

Future studies will be necessary to investigate the death process activated by e-ATP in the cholinergic neurons. Strong evidence points to P2X7 as the most likely P2 receptor subtype responsible for the e-ATP cytotoxic effect, as suggested by the potent killing activity of BzATP, the protective effect of oATP and Brilliant Blue G, and the high level of expression of this receptor in the cholinergic neurons. This does not exclude the possibility that other P2X receptors for which a pore-forming ability has been demonstrated (e.g. P2X2 or P2X4) (Khakh et al., 1999; Virginio et al., 1999) might also be involved, although a death-inducing activity for P2X subtypes other than the P2X7 has not been shown (North, 2002).

P2X7 receptors have been shown to trigger both apoptosis and necrosis, depending on the cell type (Di Virgilio et al., 2001). In the retina we have been unable to label cholinergic cells using TUNEL or ISEL staining to detect fragmented DNA but these negative results may simply reflect the loss of cholinergic markers before these neurons reach advanced stages of death. More interestingly, we found that broad caspase inhibitors were unable to increase the number of cholinergic cells (not shown), suggesting that e-ATP induced a caspase-independent death.

**Endogenous e-ATP as a local, possibly cell-type-specific cytotoxic agent for retinal cholinergic cells**

Death induced by e-ATP in the developing retina appears specific to the cholinergic neurons. Blocking e-ATP signaling increased the number of cholinergic cells, but did not induce any generalized effect on retinal area, thickness or layering. Furthermore, treatments affecting e-ATP signaling did not alter cell density in a number of control cell populations. These include the horizontal cells, the short range amacrine cells AII, the long range dopaminergic amacrine cells, as well as all the non-cholinergic cells in the GCL (mostly retinal ganglion cells at these ages), notwithstanding most of these latter cells express the P2X7 receptors (Fig. 5A) (Wheeler-Schilling et al., 2001).

These results indicate that P2X7 expression is not enough to make cells vulnerable to e-ATP-induced death, in line with previous results showing that susceptibility to ATP may depend on the level of expression of the P2X7 receptors and the potential coupling of these receptors to cytoplasmic effectors (North, 2002). In addition, local e-ATP degradation by endogenous ecto-ATPases may selectively protect specific cells (Zimmermann, 1996), keeping endogenous e-ATP below the P2X7 activation threshold around these cells. A similar reasoning may explain why we found that endogenous e-ATP controlled the death of cholinergic neurons only before P5, while direct ATP application could kill the cholinergic neurons also at later stages. Our preliminary data on the retinal distribution of the major ecto-ATPase (CD39) (Zimmermann, 1996) are consistent with this hypothesis, showing a concentration of CD39 around the ganglion cells between P0 and P5, and a much broader expression pattern afterwards (not shown).

Muller cells and astrocytes are sources of e-ATP in the adult retina (Newman, 2001; Newman, 2003), but in the first postnatal days Muller glia has not been generated yet (Wong and Godinho, 2003), and astrocytes are only found in the central retina (Ling et al., 1989). Therefore, we searched for e-ATP sources in the neonatal retina. Using quinacrine, a vital stain that binds ATP-containing vesicles, we found that the cholinergic cells store ATP in granules, which suggests that they are potential sources of e-ATP in the developing retina. Proving that the cholinergic cells do release ATP however, will require future studies.

**Death induced by e-ATP controls the local density and regular spacing of the cholinergic neurons**

We have found that the frequency of neighboring cholinergic cells spaced by less than 15 µm was higher than normal after e-ATP signaling blockade. In addition, autocorrelation analysis showed that the major effect of e-ATP blockade on the spacing of the cholinergic cells was a reduced efficacy of the mechanism by which each cholinergic cell normally excludes other cholinergic neurons from a limited region surrounding its soma. Thus, death induced by e-ATP not only controls the density of the cholinergic cells, but also contributes to the regular spacing that these neurons normally display in the retina. We do not know how this occurs, but we can propose a working explanation based on the assumption that the cholinergic cells release ATP: when two cell sources of e-ATP, which are both vulnerable to e-ATP-induced death, get too close to one another, if one or both release ATP, this can kill one, or even both of these cells, thereby reducing local overcrowding.

Most neurons in the retina form arrays, or mosaics, where cells of the same type are regularly spaced (Cook and Chalupa, 2000; Galli-Resta, 2002; Wässle and Riemann, 1978). This orderly arrangement of neurons of the same type is thought to ensure an even sampling of the visual field. Previous studies have shown that lateral cell displacement plays an important role in the formation of retinal mosaics (Galli-Resta et al., 1997; Reese and Galli-Resta, 2002; Reese et al., 1995), but simulation experiments (Jeyarasasingam et al., 1998) and studies of retinal mosaics in death-suppressing transgenic mice (Raven et al., 2003) strongly suggested that cell death also participates in this process. The present study provides direct evidence of a death control mechanism contributing to regular cell spacing and density in a neural population.

The selective elimination of cells too close to one another is a potent mechanism to prevent local overcrowding. However, cell death cannot by itself create a regular distribution of cells, unless a continuous provision of new cells is ensured until the final cell density and regular cell spacing are both achieved. This process would require an enormous amount of cell genesis, as simulation experiments easily show (e.g. simulating the formation of a normal cholinergic array by generating cells in random places and eliminating them whenever they do not obey the minimal spacing rule of 15±2 µm typical of the neonatal cholinergic arrays, shows that on average as many cells should be eliminated as finally remain in the array) (see also Eglen and Willshaw, 2002). This would not be the case, however, if cells too close to one another could also move apart: this process would reduce their risk of death, contribute to regular cell spacing, and reduce the number of new cells...
necessary to achieve a final regular density. Tangential cell dispersion could in principle be enough to space cells appropriately, but we know little of the controlling mechanisms. Lateral cell migration has been shown to involve dendritic interactions (Galli-Resta et al., 2002), to be limited to a specific developmental time window, and to displace cells no more than 100-150 μm away from their clone of origin (reviewed by Reese and Galli-Resta, 2002). In several situations therefore, it might be more expedient for misplaced cells to die rather than move around till they are properly placed.

A dynamic regulation of cell number during cholinergic cell development

We have found that blocking for 24 hours the e-ATP-induced death of retinal cholinergic neurons in vivo significantly increases the total number of these cells (Fig. 2A,B). This means that many cholinergic neurons are normally dying at these ages. However, new cholinergic cells are also observed to migrate to the cholinergic arrays at these same times (Galli-Resta, 2000; Galli-Resta et al., 1997). The total number of cholinergic neurons reflects the balance between these two opposite contributions. This is not the first example of a cell population simultaneously undergoing cell death and cell replacement in the developing nervous system. This dynamic behavior has already been shown in the early development of retinal ganglion cells (Frade et al., 1997; Frade et al., 1996), in the avian ciliary ganglion (Lee et al., 2001) and in populations of retinal and cortical neuroblasts (reviewed by de la Rosa and de Pablo, 2000). The simultaneous presence of cell death and new cell addition makes it very difficult to estimate the real amount of cell death going on during development, or even to detect cell death as a decrease in the total number of cells.

Death induced by e-ATP has never been reported in the developing nervous system. However, a number of non neuronal cell populations, ranging from human macrophages (Falzoni et al., 1995) and keratynocytes (Girolomoni et al., 1993) to colonies of fungi (Koshlukova et al., 1999) undergo cell death induced by ATP through the P2X receptor, suggesting that the cholinergic neurons are a new example of an ancient mechanism of cell death control.

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Supplementary material

Supplementary material for this article is available at http://dev.biologists.org/cgi/content/full/132/12/2873/DC1

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