Plasticity of Scarpa’s ganglion neurons as a possible basis for functional restoration within vestibular endorgans

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In a previous study, we observed spontaneous restoration of vestibular function in young adult rodents following excitotoxic injury of the neuronal connections within vestibular endorgans. The functional restoration was supported by a repair of synaptic contacts between hair cells and primary vestibular neurons. This process was observed in 2/3 of the animals studied and occurred within 5 days following the synaptic damage. To assess whether repair capacity is a fundamental trait of vestibular endorgans and to decipher the cellular mechanisms supporting such a repair process, we studied the neuronal regeneration and synaptogenesis in co-cultures of vestibular epithelia and Scarpa’s ganglion from young and adult rodents. We demonstrate that, under specific culture conditions, primary vestibular neurons from young mice or rats exhibit robust ability to regenerate nervous processes. When co-cultured with vestibular epithelia, primary vestibular neurons were able to establish de novo contacts with hair cells. Under the present paradigm, these contacts displayed morphological features of immature synaptic contacts. Preliminary observations using co-cultures of adult rodents suggest that this reparative capacity remained in older mice although to a lesser extent. Identifying the basic mechanisms underlying the repair process may provide a basis for novel therapeutic strategies to restore mature and functional vestibular synaptic contacts following damage or loss.

Keywords: vestibule, primary neurons, sensory epithelia, synapse, plasticity, repair

INTRODUCTION
Impairment of synaptic contacts between cochlear hair cells and primary auditory neurons has been demonstrated to underlie hearing loss following acoustic trauma (Pujol and Puel, 1999; Ruel et al., 2007). Current debate focuses on whether or not the acute loss of afferent terminals is reversible or permanent (Kujawa and Liberman, 2009; Lin et al., 2011). In the vestibule, impairment of synaptic contacts between vestibular hair cells and their afferent nerve terminals arising from Scarpa’s ganglion neurons may also cause several types of vestibular deficits. Although current methods of clinical evaluation preclude direct characterization of the state of synapses in the vestibule following inner ear impairment, indirect measurements of neural activity through video-nystagmography has previously demonstrated a correlation between alteration of nerve activity and vestibular deficits (Freyss et al., 1973; Haid and Mirsberger, 1985). Moreover, recent development of animal models of vestibular deficits confirmed an impact on primary vestibular synapses regardless of the origin of the impairments. Swelling of the post-synaptic terminals, often followed by synapse uncoupling, were reported upon ototoxic (Seoane et al., 2001; Vignaux et al., 2012) or excitotoxic (Liu, 1999; Shimogori and Yamashita, 2004; Brugeaud et al., 2007) vestibular insult in adult mammals. Excitotoxic damage results from the overactivation of glutamate receptors by excessive release of neurotransmitter (Olney, 1969). This leads to a massive entry of calcium and sodium into the neurons that induce both swelling of nerve terminals and subsequent activation of apoptosis. Such pathological conditions are believed to occur following local ischemia and temporal bone trauma. As in the cochlea, questions remain on the transitory or permanent nature of the synaptic impairments. Recent clinical observations revealed that recovery of vestibular activity occurs within days or weeks following an acute impairment: partial recovery of the vestibular activity determined by VOR measurement was reported in 35% of patients with vestibular neuritis (Choi et al., 2007). It is therefore of interest to define whether this functional recovery effectively relies on endogenous repair of peripheral sensory synapses. Central compensation is assumed to also support part of the functional recovery.

Spontaneous regeneration of vestibular receptors and their innervating afferents is well documented in birds, reptiles and, amphibians (Hernandez et al., 1998; Zakir and Dickman, 2006; Haque et al., 2009), but it has long been thought that this repair capacity was restricted to non-mammals. However, we recently reported the first evidence of spontaneous vestibular synap tic repair in adult mammal following mild excitotoxic damage (Brugeaud et al., 2007). Although local delivery of kainic acid in the middle ear of young adult rats evoked acute swelling of nerve terminals at synapses with hair cells, endogenous repair occurred in the days following the lesion. This synaptic repair was proposed to support, at least in part, spontaneous restoration of the vestibular function taking place during the same time frame.
The present study was designed to further investigate and extend understanding of the capacity of primary vestibular neurons for re-afferentation of hair cells following synaptic damage and loss. Here we achieved complete vestibular de-afferentation by isolating both vestibular epithelia and Scarpa’s ganglions from rodents and studied the ability of primary vestibular neurons to grow de novo neurites and re-innervate hair cells under co-culture conditions. Most experiments were performed using neonate rodents to take advantage of the highly plastic young tissue and to investigate the effect of different factors such as Brain Derived Neurotrophic Factor (BDNF) and Protein Kinase C (PKC) on such re-innervation potential. We also report preliminary experiments with adult tissue that extend these plastic capacities to mature rodents and bring new perspective on repair within vestibular sensory organs of adult mammals.

MATERIALS AND METHODS

ANIMALS

Experiments were performed using newborn, postnatal day 2–9 (P2–9) and young adult (1 month old), and adult (3 month old) wild type rodents (Wistar rats and Swiss mice, CERJ, Le Genest, France) in accordance with the French Ministry of Agriculture regulations and European Community Council Directive no. 86/609/EEC, OJL 358. All efforts were made to minimize the number of animals used and their suffering.

REAGENTS

Phenol-red free Matrigel matrix was purchase from Becton Dickin-son (Pont de Claux, France). Leibovitz medium, N2 supplement, Dulbecco’s Modified Eagle medium with Ham’s F12 nutrient (DMEM/F12), and 15 mM Hepes were purchased from Invitrogen (Cergy Pontoise, France). BDNF was a Peprotech product purchased from TEBU-bio (Le Perray en Yvelines, France). Phorbol 12-myristate 13-acetate (PMA), laminin, fish gelatin, and protease inhibitor cocktail were purchased from Sigma-Aldrich (Saint Quentin Fallavier, France). Normal swine serum was purchase from Jackson Immuno Research Europe (Cambridge, UK).

3D ORGANOFTYPC CO-CULTURES

The protocol for co-cultures of vestibular sensory epithelia and primary ganglia was derived from the 3D cell culture protocol previously detailed (Gaboyard et al., 2005). Briefly, vestibular epithelia and ganglia were aseptically removed from animals and carefully dissected in Leibovitz medium. Explants were put in a drop of phenol-red free Matrigel (15 μL) placed on laminin (10 μg/mL) coated glass coverslips with utricle, crista, and ganglion carefully oriented and positioned close to each other before jelling of the coated glass coverslips with utricle, crista, and ganglion carefully dissected in Leibovitz medium. Explants were put in a drop of phenol-red free Matrigel (15 μL) placed on laminin (10 μg/mL) coated glass coverslips with utricle, crista, and ganglion carefully oriented and positioned close to each other before jelling of the extracellular matrix. These preparations were incubated for 30 min at 37°C in a 95%/5% O2/CO2 atmosphere, at saturating moisture. Then, co-cultures were fed with culture medium DMEM/F12 15 mM Hepes supplemented with N2 cocktail (2%). Treatments in some cases with BDNF (10 ng/mL) or PMA (0.1 μM) were performed during the entire culturing period: the drugs were added to the culture feeding medium from seeding to fixation. The feeding medium was renewed every 3 days. BDNF experiments were performed using explants from both mice and rats while PMA experiments only used rat tissue. For each age and condition, cultures from 3 to 12 L were initiated and fixed over time to follow the time course of re-innervation and explore newly formed synaptic contacts. Figures are illustrative of several independent and repeated experiments, of which the total number is indicated in the Section “Results.”

ANTIBODIES

Different combinations of primary antibodies were used. In most cases, a mouse monoclonal anti-neurofilament 200 kDa (1:500; clone N52, Sigma-Aldrich) and rabbit serum anti-calretinin (1:3500; Swant, Marly, Switzerland) or for few experiments, rabbit serum anti-S100β (1:200; Swant) were used. In the remaining experiments, mouse monoclonal anti-CtBP2 to label the protein ribeye, a molecular constituent of ribbon dense core (1:350; BD Transduction Laboratories, Franklin Lakes, NJ, USA), rabbit serum anti-synaptophysin (1:500; Dako, Glostrup, Denmark) and goat serum anti-calretinin (1:3500; Swant) were mixed. Fluorescent secondary antibodies were donkey sera conjugated to Alexa-488 anti-mouse, Alexa-594 anti-rabbit, and Alexa-647 anti-goat IgG, respectively (1:700; Fisher Scientific).

IMMUNOCYTOCHEMISTRY

Co-cultures were fixed in 4% paraformaldehyde (PFA) for 2 h, followed by methanol for 30 min at –20°C. Non-specific sites were blocked and permeabilization was performed by immersing the samples overnight at 4°C in a Tris/NaCl solution (Tris 100 mM with NaCl 1.5%, pH:7.4) containing 10% swine serum, 0.5% fish gelatin, 0.2% Tween-20, 0.1% Triton X-100, and 0.1% protease inhibitors. Samples were then incubated with primary antibodies for 48 h at 4°C diluted in Tris/NaCl solution with 5% swine serum, 0.1% fish gelatin, 0.02% Tween-20, 0.01% Triton X-100, and 0.1% protease inhibitors. Specific labeling was revealed with fluorescent secondary antibodies (1:1000) in Tris/NaCl solution eventually combined with ToPro (1:1000) staining overnight at 4°C. Samples were observed with a Zeiss 5 live duo laser scanning confocal microscope (MIR INM, Montpellier). Final image processing was done with Adobe Photoshop software (San Jose, CA, USA).

TRANSMISSION ELECTRON MICROSCOPY

Co-cultures (n = 2; 18 DIV from P6 rats) were fixed with 1.2% PFA and 0.25% glutaraldehyde directly added into the medium for 48 h at 4°C. A post-fixation in PBS with 1% PFA, 2.5% glutaraldehyde, 1% picric acid, and 0.1% glucose followed for 3 h at room temperature. Then, samples were postfixed in 0.5% OsO4, dehydrated, and embedded in Embed 812- DER 736 (Delta Microscopies, Aigues Vive, France). Silver-gold ultrathin sections (80–90 nm) were collected, stained with uranyl acetate (RHEM INM, Montpellier, France) then viewed and digitized in a Hitachi H7100 electron microscope (CRIC, Montpellier, France).

RESULTS

PRIMARY VESTIBULAR NEURONS PREFERENTIALLY ORIENT NEURITE OUTGROWTH TOWARD SENSORY EPITHElia

Once removed from the inner ear of neonate mice or rats, isolated vestibular endorgans were cultured as 3D organotypic explants as previously reported (Gaboyard et al., 2005). Under these conditions hair cells rapidly recovered from the isolation procedure while an endolymphatic compartment containing enriched K

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liquid regenerated (Bartolami et al., 2011). In the present study, vestibular (Scarpa’s) ganglia were also dissected and co-cultured in the vicinity (within a distance of 0.5–1 mm) of the sensory epithelia (utricles and cristae ampullaris). During the first 3 days in vitro (DIV), both the proximal (not shown) and distal (Figure 2A) nerve processes of primary vestibular neurons degenerated after being severed from their targets or cell bodies respectively. Subsequently, de novo neurite outgrowth was observed. Bundles of fibers arising from Scarpa’s ganglion neurons with preferential orientation toward the sensory epithelia were clearly observable in all co-cultures after 6 DIV (mice \(n = 12\); rats \(n = 10\); Figures 1A,B).

As BDNF was previously demonstrated to play a critical role during developmental innervation of sensory epithelia by vestibular primary neurons (Ernfors et al., 1995), we used the co-culture model from young rodents to examine the effect of BDNF addition on de novo neurite outgrowth. When BDNF (10 ng/mL; mice \(n = 4\); rats \(n = 4\)) was added to the culture medium, the orientation of de novo neurite outgrowth was significantly altered (Figure 1C), forming a radiating pattern from Scarpa’s ganglia, and only occasionally still innervating the sensory epithelia. This observation confirms the role of BDNF in the guidance of neuritis during the innervation of sensory epithelia. It agrees with studies that show controlled release of BDNF by the targeted cells or their immediate environment is required for innervation of sensory epithelia by primary vestibular neurons (Brugeaud et al., 2007). Schwann cells have also been shown to play an important role in the guidance of neuronal processes toward their targets during peripheral nerve regeneration (Zhang and Yannas, 2005). Using immunostaining of the S100\(\beta\) protein, specifically expressed in vestibular glial cells (Bartolami et al., 2003), we confirmed that Schwann cells were part of the re-innervation process in the co-culture preparation. Indeed, the neuritic outgrowth of nerve fibers was never observed without accompanying Schwann cells (rats, \(n = 4\); Figure 1D).

**PRIMARY VESTIBULAR NEURONS FORM CELL-TO-CELL CONTACTS WITH HAIR CELLS WHEN CO-CULTURED WITH SENSORY EPITHELIA**

At 3 DIV, no remaining severed nerve processes were observed in sensory epithelia (\(n = 8\); Figure 2A). Conversely beginning at 6 DIV, growing neuronal processes arising from vestibular primary neurons entered the sensory epithelia in all co-cultures (mice and rats; \(n = 20\)). High magnification observations indicated that the neuronal processes formed cell-to-cell contacts with the hair cells (Figures 2B,C). In most cases, growing neuronal processes established contacts with multiple hair cells. Surprisingly, at this time

![Figure 1](https://www.frontiersin.org)
PMA POTENTIATES RE-INNERRAVATION OF HAIR CELLS BY PRIMARY VESTIBULAR NEURONS

Phorbol 12-myristate 13-acetate (PMA), an activator of PKC, has been implicated in the regulation of diverse cellular functions (Giorgi et al., 2010), among them, potentiation of dendritic spine development and plasticity (Goldin and Segal, 2003; Ethell et al., 2005). We tested for potential modulatory effects of PMA on the re-innervation process in vestibular co-cultures of young rodents by examining neurite outgrowth in presence of PMA (0.1 μM; rats n = 5) in the bathing medium. Confocal imaging was used to track de novo neurite outgrowth from vestibular ganglion neurons up to the contacted hair cells in the sensory epithelia (Figures 3A–D). In our model, addition of PMA potentiated the neurite outgrowth process. Gross observation indicated that the presence of PMA increased the number of neural processes growing toward and entering the sensory epithelia (Figures 3A–D). Moreover, under these conditions calyx-type contacts could be observed using 3D reconstruction in both the utricles (Figure 3C) and cristae (Figure 3E). Table 1 summarizes the results concerning the density, type of innervation, and cell-to-cell contacts in the different culture conditions. Evaluation is ranged as a mean observation obtained from the indicated number of repeated cultures. Although the mechanism underlying PMA potentiation of hair cell re-innervation by primary vestibular neurons remains unknown, it demonstrates the potential for modulating the re-innervation of peripheral vestibular targets.

DE NOVO CELL-TO-CELL CONTACTS DISPLAY CHARACTERISTICS OF SYNAPSES

To assess whether the de novo cell-to-cell contacts corresponded to synapses between afferent terminals and hair cells, we first used immunohistochemistry to label the pre- and post-synaptic elements. Hair cells were counterstained using calretinin, a calcium binding protein used as a specific marker (Dechesne et al., 1994; Zheng and Gao, 1997). The pre-synaptic ribbons, specific structures controlling the pool of releasable synaptic vesicles, were immuno-localized in the hair cell cytosol using antibodies against ribeye, the main component of the ribbon dense core element (Schmitz et al., 2000; Zanazzi and Matthews, 2009). The post-synaptic terminal element was identified and localized within the sensory epithelia using immunolabeling of synaptophysin. This small synaptic vesicle associated protein is characteristic of vestibular afferent terminals during synaptogenesis (Scarfone et al., 1991; Dechesne et al., 1997; Gaboyard et al., 2003). The juxtaposition of ribeye (Schmitz et al., 2000; Zanazzi and Matthews, 2009) and synaptophysin labeling indicated that the de novo cell-to-cell synaptic contacts were formed correctly at the site of pre-synaptic active zones. At 11 DIV (mice, n = 4), synaptophysin immunolabeling was observed in neuronal processes of both utricles (Figure 4A) and cristae (Figure 4B). In most cases, the synaptophysin immunoreactivity strongly correlated with the ribeye labeling (anti-CtBP2 antibodies; Figure 4B). Note that vesicles spread out along the fibers indicate an immature stage of synaptogenesis.

Subcellular characteristics of the newly formed cell-to-cell contacts were determined using electron microscopy (rats, n = 3). Bouton-type synaptic contacts were observed in cristae (Figures 5A,C,D). Afferent terminals full of mitochondria often faced multiple ribbons located in hair cells, surrounded by clusters of mitochondria. Post-synaptic densities (PSD) were observed on membranes of afferent fibers facing pre-synaptic ribbons, confirming that newly formed synaptic contacts were established correctly. Calyx-type terminals were observed entering the sensory cell layer and enveloping type I hair cells (Figures 5B,E–J). Single and multiple ribbons were apposed to the calyx terminal membrane; some facing PSDs were observed on the nerve terminal membrane. Both bouton- and calyx-type terminals were full of mitochondria and vesicles. Two main types of vesicles were identified (Figures 5C,D,E–J): typical small synaptic vesicles and large granular vesicles (almost 100 nm length) filling both bouton-like terminal endings on type II hair cells and calyx-type climbing terminal afferents along type I hair cells. In summary, in this co-culture model, nerve endings arising from primary vestibular neurons entered vestibular sensory epithelia and contacted both type I and type II hair cells forming calyx- and bouton-type synaptic connections respectively. Several features of these newly formed synaptic contacts (expression of synaptophysin, presence of multiple ribbons, and post-synaptic accumulation of vesicles) suggest that at least up to 18 DIV, these contacts remained immature.
FIGURE 3 | PMA potentiates vestibular synapse formation in vitro.
Eighteen DIV co-cultures of P6 rat ganglion (G), utricle (U), and crista (C) in presence of PMA (A–E) illustrates the potentiation of neurite outgrowth and synapse formation in vitro. Insets show higher magnification of utricles (B,C) and cristae (D,E). Multiple neurites were observed inside the utricles and detailed observation (inset, B) on nerve endings reconstructed in 3D (C) showed the additional formation of calyceal innervations in utricles. In cristae, PMA increased the number of neurites entering the sensory epithelia (D), and the number of calyces as shown in the 3D reconstruction (E). Neurofilaments stained in green and nuclei in red. Scale bar 100 μm (A).

CAPACITY FOR RE-INNERRATION PERSISTS IN MATURE TISSUE
Since the previously described co-culture experiments were performed using tissue explanted from neonatal rodents with ongoing synaptogenesis, we undertook preliminary experiments to test whether re-innervation capacity remained in mature tissue. Scarpa’s ganglia and vestibular epithelia from young adult (Figures 6A,B; 4 weeks old; n = 4) and mature adult (Figures 6C,D; 3 months old; n = 3) rats were co-cultured with PMA added to potentiate the re-innervation and re-connection process. With both young adult and mature adult tissue (Table 1), we observed oriented neurite re-growth toward sensory epithelia, utricles, and cristae (Figures 6A,C). At higher magnification, nerve terminals were seen penetrating the sensory cell layer, weaving between hair cells and eventually establishing bouton- and calyx-type contacts (Figures 6B,D). Note that the lack of calretinin labeling at the calyx-type connection supports the hypothesis that this typical synaptic connection was properly formed with mature type I hair cells, as expression of this calcium binding protein is lost after synapse formation. These preliminary experiments suggest that the re-innervation and re-connection capacity was still present in mature tissue following de-afferentation.

DISCUSSION
SPONTANEOUS SYNAPTIC REPAIR VERSUS DEVELOPMENTAL SYNAPTGENESIS
Based on the knowledge of developmental synaptogenesis in mammal vestibular endorgans gathered over the last decades, it can be assumed that the process we report in the present study relies on postlesional repair of damaged vestibular synapses. Indeed, it has been well documented through morphological and functional studies (for review see Eatock and Hurley, 2003) that most afferent fibers arising from Scarpa’s ganglion neurons in rodents enter the sensory epithelia a few days before birth, with the first immature synaptic contacts already identifiable at birth. Synapses acquire
their mature phenotype in the first postnatal days (Desmadryl and Sans, 1990; Dechesne et al., 1994), giving rise to the first recordable synaptic activity (Curthoys, 1982; Desmadryl et al., 1986). During this period, hair cells gradually acquire their mature biophysical properties (Rusch et al., 1998). In the presently described co-cultures prepared from neonatal rodents, the observed ability of primary neurons to restore synaptic contacts with vestibular hair cells may be supported or enhanced by developmental properties that allow formation of native synapses during the first postnatal week. The influence of such developmental properties is particularly noticeable in cultures of rodent utricles (Gaboyard et al., 2005), which only retain the remarkable capacity to regenerate a sealed vestibular cyst when removed from their native environment during the first postnatal week (Bartolami et al., 2011). The presently reported process of synaptic repair is in several ways comparable to the regenerative properties in co-cultures of spiral ganglia with denervated organ of Corti from neonate mice (Martinez-Monedero et al., 2006). In this preparation, auditory neurons removed from neonate mice were demonstrated to regenerate processes that grew toward and entered the sensory epithelia, eventually re-innervating hair cells in organs of Corti. Immunocytochemical examinations revealed that the newly formed contacts expressed the synaptic vesicle markers synapsin and SV2 at points of contact with hair cells. The presently observed sustained capacity for neurite regeneration and vestibular organ re-innervation in adult tissue suggests that this repair ability extends beyond the period of neonatal synaptogenesis. It is likely that this property supports the recovery we previously described for adult rats subjected to excitotoxic vestibular insult in vivo (Brugeaud et al., 2007). Further experiments will be required to examine whether the re-innervation process results in functional synapses and whether the mechanism underlying the regenerative process is identical during the neonate and adult periods.

### Table 1 | Regeneration of vestibular neurons in different culture conditions. Semi-quantitative comparison of innervation and synaptic contact re-connections evaluated over several experiments (n).

| Type of culture                | n  | Innervation | Synaptic contacts |
|-------------------------------|----|-------------|------------------|
|                               |    | Crista      | Utricle Crista   | Utricle |
| Mouse (P2–9)                  | 12 | ++          | + +              | + +     |
| Rat (P2–9)                    | 10 | + +         | + +              | + +     |
| Mouse or rat (P2–9) + BDNF (10 ng/mL) | 8  | ±           | ± +              | + +     |
| Rats (P2–9) + PMA (0.1 μM)    | 5  | ++++        | ++++             | ++++    |
| Rats (>4 weeks) + PMA (0.1 μM) | 7  | ++          | ++ +             | ++ +    |

**Innervation:** ± no preferential direction; ++ orientated; +++ orientated and dense.

**Synaptic contacts:** + boutons, no calyx; ++ boutons and calyces; ++++ numerous boutons and calyces.

### FIGURE 4 | Maturing synaptic contacts. In utricles (A) and cristae (B) of 11 DIV co-cultures of P4 rats, synaptic contacts were stained and observed between hair cells (blue) and neurites. Anti-CtBP2 antibodies used to label the ribeye protein, major molecular component of synaptic ribbons (green, small bright dots) and nuclei of hair cells, while anti-synaptophysin antibodies labeled small synaptic vesicles (red) contained in maturing synaptic afferents. Terminals of innervating fibers (arrows) entering sensory epithelia were observed. Co-labeling with ribeye shows close apposition of these synaptic terminals with ribbons in hair cells (arrow heads) in both cristae (A) and utricles (B). Scale bars 50 μm.
FIGURE 5 | Details of synaptic contacts. In crista, electron microscopy highlights the different synaptic contacts formed de novo between hair cells and vestibular afferent terminals in 18 DIV co-cultures from P6 rats. Some fibers (green) contact multiple hair cells ([A], stars), while others contact only one hair cell forming a terminal that ensheathes the pre-synaptic type I hair cell ([B], I). Bouton terminals ([C,D]) contact hair cells. Ribbons (white arrows) in hair cells face synaptic densities (white arrow heads). The fibers ensheathing the type I-like hair cells ([E–J]) display typical features of mature and immature calyceal terminals. Fibers climb along the hair cell membrane facing ribbons ([E–G]), presenting some post-synaptic densities ([F,G]). Contacts with multiple ([E,F]) or single ([G]) ribbons were found. In these growing afferent fibers ([H–J]), typical small synaptic vesicles of immature calyceal innervations were observed in the newly formed terminals (black arrows) ([I,J]). Large granular vesicles (black arrow head) typical of peptidergic synaptic vesicles were also present in these growing calyceal terminals ([I,J]). Scale bars 5 μm in ([A,B]), 500 nm in ([C,E,H]), and 100 nm in ([D,F,G,I,J]).

the epithelia, and re-innervated hair cells at pre-synaptic active zones. This is substantiated by electron microscopy analysis and demonstration of the juxtaposition of synaptophysin immunolabeling, characteristic of connected neuronal processes (Scarfone et al., 1991; Dechesne et al., 1997; Gaboyard et al., 2003) and CtBP2 immunoreactivity characteristic of ribeye protein present at pre-synaptic active zones in hair cells (Schmitz et al., 2000; Zanazzi and Matthews, 2009). It is possible that the reported regenerative and plastic capacities observed for part of the neuronal population in the co-culture model in vitro may be more robust in native tissue. Indeed, in vivo, several endogenous factors have been demonstrated to play a critical role in the guidance of fibers toward their targets and in the synaptogenesis process. Using BDNF deficient mice, it was previously demonstrated that BDNF controls the guidance of nerve fibers arising from Scarpa’s ganglion neurons toward the sensory epithelia (Ernfors et al., 1995). In the present study, BDNF application in the culture medium altered the preferred orientation of neurite outgrowth toward sensory epithelia. This confirms that (at least during the developmental process of synaptogenesis) controlled release of BDNF by the targeted cells...
or their immediate environment is required for directional control of neurite outgrowth (Bruggeaud et al., 2007). In this study, we also reported the involvement of Schwann cells in supporting and/or accompanying the re-innervation process, and we observed the potentiating effect of PMA on neurite outgrowth. Numerous other factors in vivo may play essential roles in this process. In particular, the role of efferent terminals in providing feedback input from the brain stem may participate in the maturation of synapses between hair cells and primary vestibular neurons (Favre and Sans, 1977; Desmadryl et al., 1992; Dememes et al., 2001). In the present co-culture model, the lack of efferent fibers may contribute to the immaturity of the newly formed synaptic contacts, characterized by the presence of multi ribbons in hair cells, as well as the increased expression of synaptophysin and accumulation of vesicles in afferent terminals. However, immature expression of synaptophysin may also suggest that the reperative de novo synaptogenesis recapitulates stages of developmental synaptogenesis. In support of this latter hypothesis, we previously demonstrated that hair cells of adult rat utricles in vivo control the re-formation of synaptic contacts with afferents after excitotoxic damage by re-expression of a voltage-gated sodium conductance (Bruggeaud et al., 2007) otherwise restricted to the developmental synaptogenesis stage (Chabbert et al., 2003). Whether primary vestibular neurons re-innervating hair cells re-establish pre-lesional branching patterns is an important question that needs to be addressed to substantiate the functional consequences of the re-innervation process. In the adult pigeon, elegant experiments have demonstrated that calyceal, dimorphic, and bouton afferents all regenerated with normal epithelial topographical distribution following complete vestibular sensory receptor loss and epithelial denervation, albeit following a slower time course. However, it was noted that regenerated afferent terminals were significantly smaller, had reduced morphological complexity and innervated fewer sensory cells in reduced epithelial areas than those developed through normal morphogenesis (Haque et al., 2009).

CONCLUSION

Whether newly formed neurite to hair cell contacts lead to functional synapses can only be answered by monitoring recovery of synaptic activity in co-cultures following complete denervation of the sensory epithelia. An adequate paradigm to record such activity is currently under development in our lab. Nevertheless, the observations presented in this study further emphasize the remarkable capacity of primary vestibular neurons to restore synaptic contacts with available targets, assuming sufficient proximity and/or appropriate guidance. Identification of the molecular and cellular mechanisms underlying such plastic capacities is a prerequisite for the development of therapeutic strategies to protect and repair vestibular sensory synapses. Such approaches could directly facilitate the treatment lasting functional impairments of peripheral origin encountered following, e.g., vestibular neuritis or vertebrobasilar stroke. Moreover, such knowledge could open new therapeutic opportunities to the pharmacological control of neuronal regeneration and synaptic repair at a time where significant progress exists in vestibular prosthesis (Della Santina et al., 2011; van de Berg et al., 2011).

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