Living-cell imaging of transgenic rat anterior pituitary cells in primary culture reveals novel characteristics of folliculo-stellate cells

Kotaro Horiguchi, Motoshi Kikuchi, Kenji Kusumoto, Ken Fujiwara, Tom Kouki, Kotaro Kawanishi and Takashi Yashiro

Division of Histology and Cell Biology, Department of Anatomy, Jichi Medical University School of Medicine, 3311-1 Yakushiji, Shimotsuke, Tochigi 329-0498, Japan

(Correspondence should be addressed to K Horiguchi; Email: kota@jichi.ac.jp; T Yashiro; Email: tyashiro@jichi.ac.jp)

Abstract

Folliculo-stellate (FS) cells in the anterior pituitary gland appear to possess multifunctional properties. Recently, the development of transgenic rats (S100b–green fluorescent protein (GFP) rats) that express GFP specifically in FS cells in the anterior pituitary gland has allowed us to distinguish and observe living FS cells in other kinds of pituitary cells. We used S100b–GFP rats to investigate the topographic affinity of FS cells for other pituitary cells. We observed living FS cells in enzymatically dispersed anterior pituitary cells of S100b–GFP rats under a fluorescent microscope, and noted that FS cells markedly extended and contracted cytoplasmic processes and formed interconnections with neighboring FS cells. In addition, FS cells adhered to small clusters of GFP-negative cells, which were primarily hormone-producing cells, and these clusters further aggregated during the course of cytoplasmic contraction. In the presence of laminin, fibronectin, and varying types of collagen, FS cells showed marked changes in shape and specific proliferative activity; however, GFP-negative cells did not. On reverse transcription-PCR analysis and immunohistochemistry, FS cells were shown to express integrin subunits, which are the cell surface receptors for extracellular matrix (ECM). In the anterior pituitary gland, FS cells and the various types of hormone-producing cells generate a unique topography in the presence of basement membrane components and interstitial collagens. The novel characteristics of FS cells observed in the present study suggest that in the anterior pituitary gland, FS cells play important roles in determining and/or maintaining local cellular arrangement in the presence of ECM components. Journal of Endocrinology (2010) 204, 115–123

Introduction

The anterior pituitary gland is composed of five types of hormone-producing cells and the folliculo-stellate cells (FS cells), which do not produce the classical anterior pituitary hormones. In vivo, these cells are surrounded by different types of extracellular matrices (ECMs), which provide the mechanical integrity, rigidity, and elasticity that are essential for these cells to perform their respective roles (Gon et al. 1987, Soji & Herbert 1989, Paez-Pereda et al. 2005). It has been shown that the lobular structures that are surrounded by ECM are responsible for organizing the functional unit in the anterior pituitary gland (Gon et al. 1987, Soji & Herbert 1989, Inoue et al. 1999, Shirasawa et al. 2004). Knowledge of the topographic features of FS cells is essential for understanding this lobular structure, the location of FS cells in the core of the lobular structure, and the characteristic enveloping of hormone-producing cells by their cytoplasmic processes, the tips of which sometimes attach to the ECM (Soji & Herbert 1989, Inoue et al. 1999, Shirasawa et al. 2004).

In addition, the anterior pituitary cells construct a unique topography by means of homophilic and heterophilic affinities between cell types (Noda et al. 2001), and by differential affinities to the ECM types (Denduchis et al. 1994, Diaz et al. 2002). The functions of hormone-producing cells are mainly regulated by endocrine, autocrine, and paracrine signaling. In addition to this humoral regulation, their functions are believed to be modified by juxtacrine signaling, by which cells receive components of the plasma membrane from adjacent cells as signals (Vankelecom & Denef 1997), and matricrine signaling, by which cells receive components of the ECM as a signal (Paez-Pereda et al. 2005).

Recently, Itakura et al. (2007) succeeded in producing transgenic rats (S100b–green fluorescent protein (GFP) rats) that express GFP specifically in FS cells in the anterior pituitary gland. The existence of S100b–GFP rats allows us to distinguish living FS cells and to separate them from the hormone-producing cells. Cell-to-cell interactions between FS cells and between FS cells and hormone-producing cells have been studied and described by several research groups (Rinehart & Farquhar 1953, Vankelecom & Denef 1997, ...
cells in the anterior pituitary gland. However, by utilizing S100b–GFP rats, we are now able to develop new approaches to investigate these interactions.

In a recent study using these transgenic rats, we found that FS cells of the S100b–GFP rats encircled hormone-producing cells with their cytoplasmic processes to form cell clusters (i.e. FS cells aggregated homophilically) both in vivo and in vitro (Horiguchi et al. 2008). It is likely that these histological features provide a suitable environment for cell-to-cell communication in the anterior pituitary gland. However, there is a limited understanding of the means by which FS cells reconstruct these histological features in primary culture, and the mechanisms underlying the interaction between FS cells and ECM. In the present study, we used living FS cells of the S100b–GFP rats to investigate both the behavior of FS cells during cluster formation and the morphological and functional responses of FS cells to the different ECM components in primary culture. We believe that the use of the S100b–GFP rats, in conjunction with living-cell imaging, may be the only method available to reveal these salient characteristics of FS cells in the anterior pituitary gland.

Materials and Methods

Animals

Transgenic S100b–GFP rats that express GFP under the control of the promoter of the S100b protein gene – a marker of FS cells – were donated by Prof. K Inoue of Saitama University and were bred in our laboratory (Itakura et al. 2007). Eight- to 10-week-old male rats weighing 250–300 g were given access to food and water ad libitum, and were housed under conditions of 12 h light:12 h darkness. Rats were killed by exsanguination from the right atrium under deep Nembutal anesthesia, and were then perfused with Ca²⁺- and Mg²⁺-free Hanks’ solution for primary culture or with 4% paraformaldehyde in 50 mM cacodylate buffer (pH 7.4) for immunohistochemistry. All animals were treated in accordance with the Guidelines for Animal Experimentation of Jichi Medical University. These guidelines are based on the NIH Guidelines for the Care and Use of Laboratory Animals.

Cell culture

Anterior pituitary cells of S100b–GFP male rats were dispersed as described previously (Horiguchi et al. 2008). Dispersed cells were plated onto eight-well glass chamber slides (1 cm²/well; Nalge Nunc International, Rochester, NY, USA) with or without a coating of ECM substrates – 3 μg/cm² of collagen type I, type III, or type IV (Nitta Gelatin, Osaka, Japan), or 10 μg/cm² of laminin (Millipore, Bedford, MA, USA) – at a density of 1×10⁵ cells/cm² in 400 μl of Medium 199 with Earle’s salts (Invitrogen) supplemented with 10% fetal bovine serum (Sigma–Aldrich Corp.), 0.5 U/ml penicillin, and 0.5 μg/ml streptomycin (Invitrogen). Cells were also plated onto a fibronectin-coated dish (9 cm²/well; IWAKI, Tokyo, Japan) under the same conditions described above. Cells were then cultured for 72 h at 37°C in a humidified atmosphere of 5% CO₂ and 95% air.

Time-lapse observation of primary culture

The cells were cultured in a CO₂ gas culture chamber (Sankei Corp., Tokyo, Japan) with a thermostat (Kokensha Engineering Corp., Tokyo, Japan) on a fluorescence-inverted microscope (IX71; Olympus Corp., Tokyo, Japan). Culture conditions were the same as described above. We time-lapse recorded the cells by using a digital camera (ORCA-ER; Hamamatsu Photonics, Shizuoka, Japan) and MetaMorph software (Molecular Devices Corp., Downingtown, PA, USA). Each observation was performed in triplicate.

Fluorescent cytochemistry of actin filaments

To observe cellular localization of actin filaments, cells were fixed with 4% paraformaldehyde in 25 mM phosphate buffer (pH 7.4) for 20 min, and were permeabilized with 0.4% Triton X-100 in PBS for 5 min at room temperature. Fixed cells were incubated in PBS containing 2% normal goat serum for 20 min at 30°C, incubated with 0.1 μM rhodamine–phalloidin (Cytokeleton Inc., Denver, CO, USA) for 30 min at room temperature, and washed with PBS. Cells were scanned using a confocal laser microscope (FV1000; Olympus).

Immunohistochemistry

Pituitary glands were carefully excised and fixed overnight in 4% paraformaldehyde in 50 mM cacodylate buffer (pH 7.4) at 4°C. Tissues were then immersed in cacodylate buffer (pH 7.2) containing 30% sucrose for 2 days at 4°C, embedded in Tissue-Tek OCT compound (Sakura Finetechical, Tokyo, Japan), and frozen rapidly. Frozen frontal sections of 8 μm thickness were obtained using a cryostat (CM3000; Leica Microsystems, Wetzlar, Germany) and were mounted on slide glasses. Sections were incubated in PBS containing 2% normal goat serum for 20 min at 30°C, and were then incubated overnight with mouse anti-rat integrin β1 monoclonal antibody (12.5 ng/ml; BD Biosciences, Erembodegem, Belgium) at room temperature. After washing with PBS, sections were incubated in PBS with Alexa Fluor 568-conjugated goat anti-mouse IgG (Invitrogen) diluted to 1:200 and were washed with PBS again. Sections were scanned using a confocal laser microscope.
Proliferation assay

To visualize the proliferative activities of cells, the nucleotide analog 5-bromo-2’-deoxyuridine (BrdU, Sigma–Aldrich Corp.) was added to primary culture for 24 h at a concentration of 3 μg/ml. Cells were fixed in 4% paraformaldehyde in 25 mM phosphate buffer (pH 7.4) for 20 min at room temperature, and were then treated with 4 M HCl in PBS for 10 min. Cells were incubated in PBS containing 2% normal goat serum for 1 h at 30 °C, and were then incubated with anti-rat BrdU mouse monoclonal antibody (1:5 μg/ml, Sigma–Aldrich Corp.) for 90 min at 30 °C. After washing with PBS, cells were incubated in PBS with Alexa Fluor 568-conjugated goat anti-mouse IgG (Invitrogen) diluted to 1:200. The absence of an observable nonspecific reaction was confirmed using normal mouse serum. Thirty random fields were imaged per well using a confocal laser microscope with a 60-fold objective lens. The percentage of immunoreactive cells was calculated by counting at least 1500 cells per well. The observations were done three times for each experimental group.

Reverse transcription-PCR analysis of integrin subunit gene expression

Total RNA fractions were prepared with Trizol reagent (Invitrogen) from anterior pituitary glands, anterior pituitary cell primary culture, and the GFP-positive cell fraction of S100b–GFP male rats. They were then sorted by a MoFlo XDP (Beckman Coulter, Inc., Fullerton, CA, USA) and were incubated with RNase-free DNase I (1 U/tube; Promega Corp). As positive controls of integrin subunit gene expression, total RNA fractions were also prepared from the adrenal gland (Otis et al. 2007) and whole fetus of S100b–GFP male rats. After inactivation of DNase I by heating for 10 min at 65 °C, cDNA was synthesized using a Superscript III reverse transcription kit with oligo-(dT)20 primer (Invitrogen). For PCR, 1 μl of the reverse transcription (RT) reaction product was added to 9 μl of PCR buffer containing 1 μl deoxy-NTPs (2 mM), 0.1 μl KOD Dash DNA polymerase (2.5 U/μl; TOYOBO, Osaka, Japan), and 0.1 μl of each oligonucleotide primer listed below (10 μM).

The primer pairs used and putative product lengths were as follows: integrin β1 (GeneBank accession no. NM_017022), forward: 5’-CTTACTCAGGCGGAGTTTGC-3’, reverse: 5’-TCACAATGGCACACAGGTTT-3’ (598 bp); integrin α1 (NM_030994), forward: 5’-ACTGGAACGGAACTGGTCG-3’, reverse: 5’-CTTATTGGTTCAGGCTCAGC-3’.

Figure 1 Time-lapse images of S100b–GFP male rat anterior pituitary cells in primary culture. Cells were time-lapse recorded at 2-min intervals under a fluorescence-inverted microscope from 28 to 37 h after the cells were plated. The cytoplasmic processes of FS cells (arrowheads) formed interconnections (arrows) (A and B). These FS cells moved closer to each other by contracting their cytoplasmic processes. Finally, they gathered nearby small clusters of GFP-negative cells, thereby forming a large cluster (K and L). (A, C, E, G, I, and K) Phase-contrast images of single frames from time-lapse movie 1. (B, D, F, H, J, and L) Fluorescence images corresponding to A, C, E, G, I, and K from time-lapse movie 2 respectively. The time elapsed from plating of cells is shown in the lower left corner as h:mins. Scale bar, 100 μm.
(Yashpal et al. 2005); integrin α6 (XM_215984), forward: 5′-CCCAAGGAGATTAGCAATGGC-3′, reverse: 5′-CA-GTCTTTGAGGAAACACCG-3′ (452 bp) (Yashpal et al. 2005); integrin α10 (NM_001107699), forward: 5′-TGTTGAAGCCTTGGCTTCTTT-3′, reverse: 5′-GGC-AGGAGGTGAGAGGCAGTG-3′ (494 bp); integrin α11 (NM_001108156), forward: 5′-AGAAGGCCGTGCAGA-GATA-3′, reverse: 5′-CCCTGTTGTAAGACTAGCTCTT-3′ (420 bp); GH (NM_001034848), forward: 5′-CAAGAGGCTGCTTTACC-3′, reverse: 5′-CAACCACAGCTCAATCTCAGC-3′ (436 bp) (Yashpal et al. 2005); integrin α6 (NM_001107699), forward: 5′-TGTGGAAGCTTGGCTTCTTT –3′, reverse: 5′-GGCAGGAAGGTGAGAGAGTG-3′ (452 bp); integrin α11 (NM_001108156), forward: 5′-AGAAGCCCGTGCAAGAATA-3′, reverse: 5′-CCCAAGGAGATTAGCAATGGC-3′, reverse: 5′-CA-CTTCTGAGCCTCCCTTCCATT-3′ (527 bp); and glyceraldehyde 3-phosphate dehydrogenase (Gapdh; M_17701), forward: 5′-CCATCACCATCTTC-CAGGAG-3′, reverse: 5′-TTCAGCTCTGGGATGAC-CTT -3′ (427 bp). Samples were subjected for 2 min at 94 °C, 25 (GH, S100B protein, and GAPDH) or 30 (integrin-β1, -α1, -α2, -α3, -α6, -α10, and -α11) cycles of 30 s at 94 °C, 2 s at 58 °C, 30 s at 74 °C, and then for an additional 7 min at 72 °C in a GeneAmp PCR System 9700.

Figure 2. Number of cells in clusters with or without FS cells. (A, B, and C) GFP images superimposed on phase-contrast images using a confocal laser microscope (A; Intact, B; GFP-negative, and C; Mixture). (D) The average numbers of cells in a cluster. The cell number was obtained by counting at least 1500 cells per well in 30 random fields. Data are expressed as mean ± S.E.M. (n = 3). The significance of the differences between intact and GFP-negative cultures and between mixture and GFP-negative cultures was determined by the Bonferroni test (P < 0.05). Scale bar, 10 μm.

Figure 3. Time-lapse images of FS cell adhesion. Cells were time-lapse recorded at 3-min intervals under a fluorescence-inverted microscope from 24 to 26 h after the cells were plated. FS cells extended their cytoplasmic processes, which resulted in adherence (arrowheads). FS cells can be distinguished from fibroblasts by the presence of GFP in the former. (A, C, and E) Single frames from a time-lapse movie. (B, D, and F) Fluorescence images corresponding to A, C, and E respectively. The time elapsed from plating of cells is shown in the lower left corner as h:min:s. Scale bar, 10 μm.
The amplified products were analyzed on 1.5% agarose gels and were visualized by ethidium bromide staining. Negative controls were subjected to RT-PCR omitting RT, cDNA, or KOD Dash DNA polymerase, and they showed no reaction bands.

Statistical analyses

The data are presented as the mean ± S.E.M. for at least three rats in each group. The significance of differences between control and test values was determined by Dunnett’s test and the two-tailed multiple t-test with Bonferroni correction (three comparisons in three groups). Differences were considered significant when $P$ was $<0.05$.

Results

Living-cell imaging of FS cells in primary culture

We observed the behavior of living FS cells in primary culture of S100b–GFP male rat anterior pituitary cells. The cells were cultured in a CO$_2$ gas culture chamber on a fluorescence-inverted microscope and were photographed every 2 or 3 min for a period of 72 h. We observed that FS cells were remarkably motile (Fig. 1, Supplementary movies 1 and 2, see section on supplementary data given at the end of this article) as compared to GFP-negative cells, which were primarily hormone-producing cells. At 28 h of primary culture, cytoplasmic processes of some FS cells had extended ∼50 μm, and FS cells were interconnected by these processes (Fig. 1B). Thereafter, the FS cells contracted their cytoplasmic processes, aggregated (Fig. 1D), and gathered adjacent GFP-negative cells into a cluster. Between 31 and 37 h of culture, FS cells (Fig. 1F, H, and J) continued the extension and contraction of their cytoplasmic processes, thereby assembling ever-larger clusters. FS cells moved at an average rate of 6 μm/min during a 9-h observation. During the course of observation, the cytoplasmic processes of some FS cells encircled GFP-negative cells (Fig. 1L). After 72 h, the morphology of cells did not differ from that observed in conventional primary culture. Next, we counted the number of cells in clusters with or without FS cells. Dispersed cells were separated into GFP-positive and GFP-negative cells by a cell sorter (MoFlo XDP; Beckman Coulter, Inc). GFP-positive and GFP-negative cells amounted to 5 and 95% of dispersed cells respectively. The percentage of GFP-positive cells among all dispersed cells was nearly equal to that noted in a previous report (Itakura et al. 2007). Unsorted cells (Fig. 2A), GFP-negative cells (Fig. 2B), and a mixture of 5% GFP-positive/95% GFP-negative cells (Fig. 2C) were cultured at a density of $1 \times 10^5$ cells/cm$^2$ for 72 h on the uncoated surface of eight-well glass chamber slides. The number of cells among the GFP-negative fraction cluster was lower than that among the unsorted cells (Fig. 2D). When GFP-positive cells were added to GFP-negative cells, the number of cells in a cluster was similar to that observed among the unsorted cells (Fig. 2D).

Figure 3 shows the cytoplasmic process of an FS cell attaching to another FS cell at ∼24 h of primary culture. This extension of the cytoplasmic process from one FS cell to another was quite precise (Fig. 3B, D, and F).

Actin arrangement in FS cells

We observed the actin arrangement of motile FS cells. FS cells were photographed during the course of cell binding under a fluorescence-inverted microscope at ∼24 h after they were plated (Fig. 4A). They were then fixed with 4% paraformaldehyde and were subjected to fluorescent cytochemistry for actin. Actin filaments were detected along the elongated cytoplasmic processes (Fig. 4B).

Morphological changes of FS cells on ECM

We observed the behavior of FS cells in primary culture on various ECM-coated surfaces. As shown in Fig. 5, many FS cells, with distinct morphological changes, were present on type III collagen after 72 h of culture; GFP-negative cells
retained their round shape. FS cells had flattened and markedly extended their cytoplasmic processes. Furthermore, the cytoplasmic processes of a large number of FS cells had interconnected. Similar morphological changes in FS cells were observed on the other ECM surfaces, i.e. collagen type I, collagen type IV, laminin, and fibronectin (data not shown). However, a small number of FS cells retained their round shape, even on ECM.

**Proliferation of FS cells on ECM**

In primary culture on ECM-coated surfaces, proliferation of FS cells was observed frequently from 48 to 72 h after the cells were plated. Figure 6A–D and Supplementary movie 3, see section on supplementary data given at the end of this article show the division of an FS cell over a period of 1 h. Observation of BrdU incorporation revealed that only FS cells with cytoplasmic processes showed proliferative activity; round FS cells did not show proliferative activity (Fig. 6E and F). The percentage of BrdU-positive cells among all FS cells was much higher on the type I collagen-coated surface (22.6 ± 1.8%, mean ± s.e.m., n = 3) than on the uncoated surface (0.1 ± 0.1%, mean ± s.e.m., n = 3). Next, we counted the number of FS cells before and after primary culture. The number of FS cells did not differ significantly before and after culture for 72 h on uncoated glass (Fig. 6G). In contrast, the number of FS cells increased by more than 50% on all the examined ECM-coated surfaces (Fig. 6G). These differences in the number of FS cells on uncoated and ECM-coated surfaces were statistically significant (P < 0.01, Dunnett’s test).

**Expression of integrin subunit genes in FS cells**

We examined the expression of integrin subunit genes in FS cells by RT-PCR. Integrin-β1, -α1, -α3, and -α6 were detected in anterior pituitary cells both in vivo and in vitro; integrin-α2, -α10, and -α11 were not detected (Fig. 7A). Analysis of GFP-positive and GFP-negative fractions in a cell sorter resulted in identical results, with the exception of GH and S100β protein gene, which were not detected in the fractions of GFP-positive and GFP-negative cells respectively (Fig. 7A). Furthermore, immunohistochemical techniques revealed that integrin β1 was expressed in GFP-positive and GFP-negative cells (Fig. 7B).

**Figure 6** Proliferation of FS cells on collagen type III in primary culture. Cells in culture were time-lapse recorded at 3-min intervals under a fluorescence-inverted microscope from 60 to 61 h after the cells were plated. During a 1-h period, an FS cell (A, arrowhead) divided on a surface coated with collagen type III in primary culture (A–D). (A–D) Single frames from time-lapse movie 3. (E) Confocal image of BrdU incorporation (green, FS cells and red, BrdU). (F) Phase-contrast image of the same field shown in E. (G) Number of FS cells before and after primary culture on an uncoated glass chamber (before and after), and after primary culture on glass chambers coated with collagen type I (Col I), collagen type III (Col III), collagen type IV (Col IV), laminin (Lami), and fibronectin (Fibro). Cells were retrieved by incubation in 0.3% EDTA and were then counted with a hemocytometer. The time elapsed from plating of cells is shown as h:mins in the lower left corner of panels A, B, C, and D. Scale bars, 10 μm. Data are expressed as mean ± s.e.m. (n = 5). Differences in the numbers of FS cells on uncoated and ECM-coated surfaces were statistically significant (P < 0.01, Dunnett’s test).
GFP-negative cells. Scale bar, 10 μm. was observed in both GFP-positive cells (FS cells, green) and pituitary gland. Immunoreactivity of the integrin α1 subunit in the anterior pituitary tissue, primary culture (primary culture), the GFP-positive cell fraction (GFP+), and the GFP-negative cell fraction (GFP−) were analyzed by RT-PCR for mRNA of integrin-β1, -α2, -α3, -α6, -α10, and -α11 subunits; GH; S100β protein (S100β); and GAPDH. (B) Immunohistochemistry for integrin β1 subunit in the anterior pituitary gland. Immunoreactivity of the integrin β1 subunit (red) was observed in both GFP-positive cells (FS cells, green) and GFP-negative cells. Scale bar, 10 μm.

Discussion

In the present study, we observed living FS cells in primary culture of S100b–GFP rat anterior pituitary cells and noted that FS cells markedly extend and contract their cytoplasmic processes by rearrangement of their actin cytoskeleton. Toral et al. (2003) reported that in primary culture of anterior pituitary cells, changes in the morphology of hormone-producing cells of infant rats were facilitated by alteration of the arrangement of actin filaments, a process that was not observed in adults. In the present study, hormone-producing cells in adults remained round, as described in Toral et al. (2003); however, in our study, FS cells in adults behaved in a manner similar to that observed in infants. After the interconnection of cytoplasmic processes, FS cells pull toward each other by contracting these processes (Fig. 1 and Supplementary movies 1 and 2) and small clusters of GFP-negative cells attach to the FS cells, which results in the formation of a larger cluster. We also noted that the number of cells in a cluster was significantly higher in the presence of FS cells (Fig. 2). These findings suggest that this cluster formation in anterior pituitary cells partly depends on the activity of FS cells.

FS cells formed interconnections with neighboring FS cells in primary culture. A similar phenomenon was observed in a culture experiment using TtT/GF cells line, i.e. cloned FS cells (Yamasaki et al. 1997). Interestingly, some FS cells precisely extend their cytoplasmic processes toward other cells, as shown in Fig. 3. This suggests the presence of guidance factors similar to those observed in axon growth cones. Axons respond to certain chemoattractants and chemorepellents, which function as permissive and non-permissive guidance signals (Nakamura et al. 2000, Round & Stein 2007). The results of the present experiment indicate that a similar mechanism may be involved in establishing interconnections between FS cells. At the tips of the cytoplasmic processes of TtT/GF cells, annexin I, which is known to be a paracrine factor in the gland, is immunohistochemically localized, and may function as a paracrine guidance substance, as suggested by Chapman et al. (2002).

FS cells in anterior pituitary gland connect to each other not only by mechanical means, but also by gap junction-mediated biochemical communication (Soji & Herbert 1989). This system of connections is referred to as the FS cell network. In vivo electrophysiological studies have shown that FS cells form networks that transmit stimulation to neighboring FS cells as Ca2+ waves through gap junction channels (Fauquier et al. 2001, Sato et al. 2005). It has also been shown that FS cells in vitro can transfer small molecules to neighboring FS cells through gap junction coupling (Morand et al. 1996, Kabir et al. 2005). In addition, we recently reported that FS cells reconstructed gap junctions in primary culture of S100b–GFP rat anterior pituitary cells (Horiguchi et al. 2008). Indeed, the living-cell image shown in Fig. 3 may show the initiation of FS cell network reconstruction.

In our first attempt to investigate the interaction of FS cells and ECM, we examined whether FS cells respond to ECM components of the basement membrane (collagen type IV, laminin, and fibronectin) and to interstitial collagens (collagen type I and III). We found that FS cells with long cytoplasmic processes, but not in round FS cells (Fig. 4B), we believe that FS cells extend and contract their cytoplasmic processes by rearrangement of their actin cytoskeleton.
types I and III). We observed that FS cells first become flatter and markedly extend their cytoplasmic processes to form numerous interconnections with neighboring FS cells in the presence of ECM (Fig. 5). ECM affects FS cells not only morphologically, but also functionally. On the surface of ECM, FS cells exhibited marked proliferation and GFP-negative cells showed almost no proliferation (Fig. 6 and Supplementary movie 3). This difference in proliferation was confirmed by the BrdU incorporation experiment (Fig. 6E and F). Wilson (1986) examined tritiated thymidine uptake and suggested that, as compared to all types of hormone-producing cells, FS cells may proliferate more frequently in adult anterior pituitary gland in vivo. However, to our knowledge, the present study is the first to directly demonstrate proliferation of FS cells.

We investigated the expressions of possible receptor molecules for the ECM components, i.e. integrin subunits, in FS cells by RT-PCR using GFP-positive cell fractions. We are the first to note that FS cells expressed integrin-α1, -α3, -α6, and -β1 (Fig. 7A); it was previously reported that integrin α5 was expressed in anterior pituitary cells in vivo and in vitro (Horacek et al. 1994, Kikuchi et al. 2005). Integrins comprise an α- and a β-subunit, which form a heterodimer. In mammals, 18 types of α-subunits and 8 types of β-subunits are known; their various combinations give rise to 24 integrin heterodimers, which differ in ligand specificity (Hynes 2002). With respect to these ligand specificities, we believe that FS cells bind collagen, laminin, and fibronectin through integrin-α1β1, -α3β1, and/or -α6β1, and integrin α5β1 respectively in the anterior pituitary gland. Immunohistochemistry confirmed that the common integrin β1 subunit was expressed in GFP-positive and GFP-negative cells (Fig. 7B). However, a number of FS cells did not extend cytoplasmic processes and maintained a round shape throughout 72 h of culture. There are two possible explanations for this difference in morphology. The difference may be due to heterogeneity in FS cells. FS cells are classified into several types based on their content, i.e. keratin, globular fibrillary acidic protein, or major histocompatibility complex class II antigen (Höfler et al. 1984, Tachibana & Yamashima 1988, Allaerts et al. 1996). The variation in response to ECM may be due to differences between these FS cell types. The morphological difference may also be due to variation in the intracellular signal pathways. In addition to its role as a molecule that facilitates mechanical adhesion to ECM, integrin is known to act as a starting point for various signaling pathways that regulate cell migration and proliferation (Martin et al. 2002). Integrin may associate differently with the intracellular signaling pathways in FS cells that do and do not extend cytoplasmic processes. If so, this would constitute matricrine regulation of FS cells.

Accumulating evidence indicates that FS cells have numerous possible functions. In the postnatal development of the anterior pituitary gland, FS cells are reported to act as stem and proliferative progenitor cells (Yoshimura et al. 1977, Horvath & Kovacs 2002, Inoue et al. 2002). They also possess phagocytic activity (Horvath et al. 1974, Ogawa et al. 1997). Furthermore, FS cells regulate hormone-producing cells by secreting paracrine factors such as basic fibroblast growth factor (Ferrara et al. 1987), vascular endothelial growth factor (Gospodarowicz & Lau 1989), IL-6 (Vankelecom et al. 1989), and nitrogen oxide (Chen et al. 2000). Finally, the present study revealed that FS cells, under the influence of ECM components, may play important functional roles in determining and/or maintaining local cellular arrangement in the anterior pituitary gland.

Supplementary data
This is linked to the online version of the paper at http://dx.doi.org/10.1677/JOE-09-0333.

Declaration of interest
The authors declare that there is no conflict of interest that would prejudice the impartiality of this research.

Funding
This work was partly supported by a Grant-in-Aid for Scientific Research (C) (19590194) and (21570067) from the Ministry of Education, Culture, Sports, Science, and Technology of Japan.

Acknowledgements
We thank Prof. K Inoue (Satama University, Japan) for supplying the transgenic rats. We are grateful to M Yatabe for her excellent technical assistance. We also thank David Kipler, ELS, of Supernatant Communications for revising the language of the manuscript.

References


Paez-Pereda M, Kuchenbauer F, Arzt E & Stalla GK 2005 Regulation of pituitary hormones and cell proliferation by components of the extracellular matrix. Brazilian Journal of Medical and Biological Research 38 1487–1494.


Vankelecom H & Denef C 1997 Paracrine communication in the anterior pituitary as studied in reaggregate cell cultures. Microscopy Research and Technique 15 150–156.


Received in final form 14 October 2009 Accepted 9 November 2009 Made available online as an Accepted Preprint 9 November 2009