Identification of multipotent progenitors in the embryonic mouse kidney by a novel colony-forming assay

Kenji Osafune\textsuperscript{1,2,3}, Minoru Takasato\textsuperscript{2,4}, Andreas Kispert\textsuperscript{5}, Makoto Asashima\textsuperscript{2,3} and Ryuichi Nishinakamura\textsuperscript{1,4,6,*}

Renal stem or progenitor cells with a multilineage differentiation potential remain to be isolated, and the differentiation mechanism of these cells types in kidney development or regeneration processes is unknown. In an attempt to resolve this issue, we set up an in vitro culture system using NIH3T3 cells stably expressing Wnt4 (3T3Wnt4) as a feeder layer, in which a single renal progenitor in the metanephric mesenchyme forms colonies consisting of several types of epithelial cells that exist in glomeruli and renal tubules. We found that only cells strongly expressing SalI (SalI-GFP\textsuperscript{high} cells), a zinc-finger nuclear factor essential for kidney development, form colonies, and that they reconstitute a three-dimensional kidney structure in an organ culture setting. We also found that Rac- and JNK-dependent planar cell polarity (PCP) pathways downstream of Wnt4 positively regulate the colony size, and that the JNK pathway is also involved in mesenchymal-to-epithelial transformation of colony-forming progenitors. Thus our colony-forming assay, which identifies multipotent progenitors in the embryonic mouse kidney, can be used for examining mechanisms of renal progenitor differentiation.

KEY WORDS: Progenitor, Kidney, Colony-forming assay, SalI, Wnt, PCP, JNK, Rho, Mouse

INTRODUCTION

Mammalian adult kidney, metanephros, is formed by reciprocally inductive interaction between two precursor tissues derived from the intermediate mesoderm, the metanephric mesenchyme and the ureteric bud. The ureteric bud induces the metanephric mesenchyme to differentiate into the epithelia of glomeruli and renal tubules, endothelial and stromal cells (Saxen, 1987). Inductive signals have been vigorously investigated, and several factors have been elucidated that trigger epithelialization of metanephric mesenchyme in explant culture system; the members of Wnt family (Herzlinger et al., 1994; Kispert et al., 1998), leukemia inhibitory factor (LIF) (Barasch et al., 1999; Plisov et al., 2001), and transforming growth factor \( \beta2 \) (TGF\( \beta2 \)) (Plisov et al., 2001). These studies have also suggested the presence of clonal cells in mesenchymal rudiments, which sequentially form renal condensation, comma (C)- and S-shaped bodies, and terminally epithelia of glomeruli and renal tubules, and the existence of single epithelial precursors responding to LIF was demonstrated in mesenchyme (Barasch et al., 1999). One previous report suggested retrospectively the presence of multipotent cells in embryonic kidneys, demonstrating that in several portions of nephron were derived from a single stem cell using \( \text{lacZ} \) gene transduction with retrovirus into a single cell of mesenchyme (Herzlinger et al., 1992). However, none has isolated prospectively the renal progenitor cells with a multilineage differentiation potential from the embryonic kidney, and none has examined their differentiation mechanisms in a single cell culture. There has been a lack of assay systems that specifically identify renal progenitors, as in cases of the neurosphere method for neural stem cells (Reynolds et al., 1992) and the colony assay for hematopoietic progenitors (Pluznik and Sachs, 1965; Bradley and Metcalf, 1966).

We previously generated mice in which the green fluorescence protein gene (GFP) was knocked into the locus of SalI (SalI-GFP mice), a zinc finger nuclear factor that is expressed in the metanephric mesenchyme and that is essential for kidney development (Nishinakamura et al., 2001; Takasato et al., 2004). SalI is also expressed in the subventricular zone of the central nervous system and progress zones of limb buds, where neural and mesenchymal stem cells reside, respectively, leading to speculation that SalI might have some association with stem cells in several organs, including the kidney.

Targeted disruption of Wnt4 results in kidney agenesis and impairs mesenchymal-to-epithelial transformation (Stark et al., 1994), and co-culture with 3T3Wnt4 induces tubulogenesis in the mesenchyme rudiment in organ culture (Kispert et al., 1998), suggesting both essential and sufficient roles of Wnt4 for epithelial differentiation of metanephric mesenchyme. Recently, Wnt9b expressed in the ureteric bud was shown to function upstream of Wnt4 (Carroll et al., 2005). Thus, we attempted to set up assay systems that can identify and characterize the progenitor cells with multipotent differentiation potential from uninduced metanephric mesenchyme using Wnt4 signal. Wnt genes are known to regulate multiple cellular functions using at least three intracellular signaling branches: the \( \beta \)-catenin pathway (canonical pathway), in which stabilized \( \beta \)-catenin interacts with members of the lymphoid enhancer factor/T cell factor (LEF/TCF) family of transcription factors and activates gene expression in the nucleus (Wodarz and Nusse, 1998; Miller et al., 1999); the planar cell polarity (PCP) pathway, which involves Jun N-terminal kinase (JNK) and the Rho family of small guanosine triphosphatases (GTPases) and which directs cytoskeletal rearrangements, coordinated polarization within the plane of epithelial sheets, and morphogenetic movements during development (Veevan et al., 2003; Wallingford et al., 2002); and the Wnt/Ca\( ^{2+} \) pathway, which leads to release of intracellular calcium and is implicated in Xenopus ventralization and in the

\textsuperscript{1}Division of Stem Cell Regulation, The Institute of Medical Science, The University of Tokyo, Tokyo 108-8639, Japan. \textsuperscript{2}Department of Life Sciences (Biology), Graduate School of Arts and Sciences, The University of Tokyo, Tokyo 153-8902, Japan. \textsuperscript{3}ICORP, JST, Satama 332-0012, Japan. \textsuperscript{4}Division of Molecular Embryology and Genetics, Kumamoto University, Kumamoto 860-0811, Japan. \textsuperscript{5}Institut für Molekularbiologie, Medizinische Hochschule Hannover, 30625 Hannover, Germany. \textsuperscript{6}PRESTO, JST, Satama 332-0012, Japan.

\*Author for correspondence (e-mail: ryuichi@kaiju.medic.kumamoto-u.ac.jp)

Accepted 24 October 2005
regulation of embryonic cell movements (Miller et al., 1999; Veeman et al., 2003; Wallingford et al., 2002). Mechanisms by which Wnt pathways mediate cellular effects in kidney development are poorly understood.

In this study, we established a novel colony-forming assay system using 3T3-expressing Wnt4 to identify renal progenitors in the metanephric mesenchyme. Combining our colony-forming assay with flow cytometry, we found that these progenitors could be enriched by using Sall1 as a marker. We also examined the effects of Wnt downstream branches on the renal progenitors.

MATERIALS AND METHODS

In vitro colony-forming assay

Metanephric mesenchyme of embryonic day (E) 11.5 mice was isolated surgically from embryonic kidney rudiment. The mesenchyme was incubated in 0.05% trypsin-EDTA at 37°C for 10 minutes and then transferred into DMEM with 10% fetal calf serum. Mesenchymal cells were then mechanically dissociated by gentle aspiration through repeated pipetting. Metanephros of E14.5 and 17.5 embryos was incubated in 1 mg/ml Dispase (Invitrogen) at 37°C for 30 minutes then mechanically dissociated by repeated pipetting. NIH3T3 cells stably expressing Wnt3a, Wnt4 and lacZ (Kispert et al., 1998) were mitotically inactivated with mitomycin C before use. Single mesenchymal cells were sorted by FACS Vantage (Becton Dickinson) and plated on these feeder cells at a low density (5 × 10^3 cells/well of 6-well plates), then cultured in DMEM/F12 with 5% knockout serum replacement (Invitrogen), 10 μg/ml insulin, 6.7 μg/l sodium selenite, 5.5 μg/ml transferrin, 1 × 10^-7 mol/dexamethasone, 10 mmol/l nicotinamide, 2 mmol/l L-glutamine, 50 mmol/l β-mercaptoethanol, 5 mmol/l HEPES and penicillin/streptomycin.

RT-PCR

Primers used for PCR were as follows:

- Pax2, 5′-AGGCCATCCTGCCTAGATGAC-3′ and 5′-CTCGGCTTCTCCTCTCCTAC-3′
- Lim1 (Lhx1 – Mouse Genome Informatics), 5′-TGGACCTGTTCCTCCCTTGAAC-3′ and 5′-TGGTCTTCTTTGCCACACTG-3′
- Eya1, 5′-CCTCTCAGGAGTTTCTCCAGATT-3′ and 5′-AATCCTCTGGCATGAGACAGTAC-3′
- Slit1, 5′-CTCTCAGGAGTTTCTCCAGATT-3′ and 5′-GTACAGCTTCTCCTCAGAC-3′
- Wt1, 5′-ACCACCGCCTGCAATAAAGAAG-3′ and 5′-GCTGAAACGCTTCTCCTTGTT-3′
- Hoxa11, 5′-GGATTTTGATGACCGCTGTCT-3′ and 5′-GAGTAGTCAGTGCGCCAGATT-3′
- glial cell lineage derived neurotrophic factor (Gdnf), 5′-CCCGAAGATATATCCTGGACCA-3′ and 5′-TAGCCCAACACAAAGCTGAT-3′
- integrin α8, 5′-GGCGAAAGGTGCAGTCTTAAA-3′ and 5′-GAAGGAGAACATTCCATGATG-3′
- integrin α3, 5′-CCGGCTTGTCATCAATATCCT-3′ and 5′-CGCAATTTGTCATCAGAC-3′.

Cell cell adhesion molecule (Ncam), 5′-ACGTCCGCTTTCA-TAGTTCCCTG-3′ and 5′-CTATGGTCTCCATCTCCTTT-3′

E-cadherin (cadherin 1 – Mouse Genome Informatics), 5′-GAC-CTTCTCTCTCTGTGCTCTC-3′ and 5′-GTGGACCTGGCTCTTACAGTAT-3′

K-cadherin (cadherin 6 – Mouse Genome Informatics), 5′-CTAGTGGCTCTCCACAGAAG-3′ and 5′-CGTACTGGTGACACACCTAATG-3′

Rt, 5′-GGCTTACCCAGGAGATGTAAGAAG-3′ and 5′-CATCAGGGAAACAGTGGTAG-3′

Hoxb7, 5′-TTCCGGCAACAAACTCTTCTG-3′ and 5′-CGGAGAGGTTCTGTCTAAAG-3′

α-actinin-4, 5′-TGGTGAACATCTCTACTTCTG-3′ and 5′-CGGACGCTGGCTACTA-3′

CD2-associated protein (CD2-AP), 5′-AGGAATCAGGCCACCAT-3′ and 5′-TGGCTACGGCTTTGAGTTC-3′

P-cadherin (cadherin 3 – Mouse Genome Informatics), 5′-CAC-A CGACCTTGATGTACACC-3′ and 5′-GAATGTGGCCCTCCATCA-3′

Podoplanin, 5′-TCTACTGGCAAGGCACATCCT-3′ and 5′-GCTCTT TAGGGCCGAGCCTT-3′

Podocalyxin-like, 5′-ACTAATGCGGCTCTCACC-3′ and 5′-AAAT- TCTGACGTCCTGTGGA3′

Aquaporin 1 (Aqp1), 5′-CTCTCCAGGACATGCTTCTC-3′ and 5′-CAGTGCGCTTCTGACCTT-3′

Chloride channel 5 (Clec5), 5′-TGGCCTCTTGTTGCTCTTT-3′ and 5′-GGCCAAGAAAGAGCCACATG-3′

Cubulin (intrinsic factor coabsorptive receptor, 5′-CAACCTT GCCGGTTGTCTTAC-3′ and 5′-GCTCTAGTACGTGCTGTTGA3′

Megalin (low density lipoprotein receptor-related protein 2 – Mouse Genome Informatics), 5′-CAGGGAGCTCTCTGAGAAG-3′ and 5′-CTCCCTCTTTGGAGAC-3′

Sodium glucose transporter 1 (Sglt1, Slc5a1 – Mouse Genome Informatics), 5′-GCCATCCTTTCTCTTAGCAT-3′ and 5′-ACCAGCTTGCTTACCAG-3′

Clek2 (Clekkb – Mouse Genome Informatics), 5′-CCTCTCCT CGCTCTGTTGCT-3′ and 5′-AAGAAGTCCGCGCTGTGTA-3′

Polycystin 2, 5′-GTGGTGGTTGCAAACTAGAACT-3′ and 5′-TCTCCTGACATACAG-3′

Renal outer medulla K channel 2 (Romk2), 5′-TGGTCTCCCACAAAGTGAGAAAG-3′ and 5′-ATGGCACCACACATGAAGA3′

Epithelial Na channel (EnaC, Scnn1g – Mouse Genome Informatics), 5′-GCCCTACGTCTTTCAAGGAC-3′ and 5′-CCAAGTGGAAAAT-GCTA-3′

Na/Ca exchanger, 5′-TGTTTTTACGTGGTCTTGAC-3′ and 5′-TGGAAAGTGTGCCTGCCTC-3′

Polycystin 1, 5′-CCTCTGTCGCCCTTCTGAGTC-3′ and 5′-TGATCC ATCTTCCATGAC-3′

Fosx1, 5′-CTCGGTGACAGCCCTCTTCATT-3′ and 5′-GCCGTTGCCTCCTCTTCG-3′

Slc12a1, 5′-CTCGGTGACAGCCCTCTTCATT-3′ and 5′-GCCGTTGCCTCCTCTTCG-3′

Slc5a1, 5′-CTCGGTGACAGCCCTCTTCATT-3′ and 5′-GCCGTTGCCTCCTCTTCG-3′

CD45 (Ppcre – Mouse Genome Informatics), 5′-CCACACGGACACT GAACTT-3′ and 5′-TAGCACTGAGTGGCTTTCG-3′

Glyceraldehyde-3-phosphate dehydrogenase (Gapdh), 5′-TGATG ACAAGAGGTGGTGTAGA3′ and 5′-TCTCGTGGAGGCACATGTA GGCCAT-3′

PCR cycles were as follows: Gapdh, initial denaturation at 94°C for 2.5 minutes, followed by 22 cycles of 94°C for 30 seconds, 60°C for 30 seconds, 72°C for 30 seconds, and final extension at 72°C for 10 minutes; other genes, initial denaturation at 94°C for 2.5 minutes, followed by 28-33 cycles of 94°C for 30 seconds, 58°C for 1 minute, 72°C for 30 seconds, and final extension at 72°C for 10 minutes.

Organ culture

In order to examine the in vitro differentiating potential of cell populations included in the metanephric mesenchyme, each cell population was separated by flow cytometry and was pelleted down by low-speed centrifugation (380 g). The resultant cell pellet (1 × 10^6 cells per pellet) was cultured on 3T3Wnt4 cells at air-fluid interface on a polycarbonate filter (0.4 μm, Nucleopore) supplied with DMEM plus 10% fetal calf serum at 37°C, 5% carbon dioxide. 3T3Wnt4 cells (50,000 cells in 50 μl medium) were seeded on the filter 24 hours before the experiments, as described (Kispert et al., 1998). To examine the influence of reagents on tubulogenesis, two metanephris or mesenchyme rudiments from E11.5 embryos were cultured on a polycarbonate filter. For the culture of mesenchyme rudiments, 3T3Wnt4 cells were used as described above.
Retroviral infection
The cDNA clones of the active mutant form of β-catenin (pUC-EF-1α-β-catenin ΔN-HA) (Miyaishi et al., 2000), the full length of rat axin (pRSKS
rAxin) (Ikeda et al., 1998), and both constitutively-active and dominant-negative mutant forms of human Rac1 and RhōA with N-terminus flag tag (pCAGIP-flag-Rac1 (Val), pCAGIP-flag-Rac1 (Asn), pCAGIP-flag-RhōA (Val), pCAGIP-flag-RhōA (Asn)) were subcloned into retroviral vector pMY-IRE-EGFP (Kitamura et al., 2003). To produce recombinant retrovirus, these plasmid vectors were transfected into the virus packaging cell line PLAT-E (Moria et al., 2000) using FuGENE (Roche), and supernatant from the transfected cells was collected to infect cells of the metanephric mesenchyme. The viral supernatant was centrifuged at 20,000 g overnight at 4°C to concentrate the virus. To infect mesenchymal cells with the retrovirus, dissociated mesenchymal cells were resuspended into the concentrated virus supernatant with adding polybrene. The suspension was centrifuged 1400 g for 4 hours at room temperature. After washing with PBS, mesenchymal cells were plated onto 3T3 feeder cells.

Immunocytochemistry and lectin staining
The colonies formed on 3T3Wnt4 feeder were fixed with 4% paraformaldehyde in PBS for 20 minutes at 4°C. After washing with PBS, PBS containing 2% skimmed milk and 0.1% Triton-X was incubated as a blocking solution for 1 hour at room temperature. The fixed dishes were incubated with primary antibodies overnight at 4°C followed by incubation with secondary antibodies for 1 hour at room temperature. The following antibodies were used: rabbit anti-Pax2 (Babco), rabbit anti-WT1 (Santa Cruz), mouse anti-E-cadherin (Becton Dickinson), rabbit anti-AQP1 (Chemicon), and rabbit anti-phosphorylated JNK1 and 2 (Biosource). Rhodamine-conjugated anti-rabbit IgG (H+L) and anti-mouse IgG (Chemicon) were used as secondary antibodies. To examine the expression of a proximal renal tubule-specific marker, fluorescein isothiocyanate (FITC)-conjugated Lotus Tetragonolobus lectin (LTL; Vector Labs) was used. After each step, the cultured cells were washed three times with PBS containing 0.1% Triton-X. For detection of Sall1, mesenchymal cells derived from Sall1-GFP heterozygote embryos were cultured on 3T3 feeder and subjected to GFP immunostaining procedure using rabbit anti-GFP (Molecular Probes). Rhodamine-conjugated peanut agglutinin (PNA; Vector Labs) staining was done as described (Gilbert et al., 1994). Organ culture tissues were fixed with 4% paraformaldehyde in PBS for 1 hour at 4°C and incubated in PBS including 0.1% saponin (Sigma) for 1 hour at 37°C, then the same staining procedure was carried out. Staining with rabbit anti-secreted frizzled-related protein 2 (sFRP2; Santa Cruz) and FITC-conjugated Dolichos biflorus agglutinin (DBA; Vector Labs) were also used on sections of paraffin-embedded explants to examine the effect of reagents on tubule formation and branching, respectively.

RESULTS
In vitro colony formation from E11.5 metanephric mesenchyme
We cultured single cells from the metanephric mesenchyme of E11.5 embryos, using 3T3Wnt4 as a feeder layer in a serum-free condition. The metanephric mesenchyme of transgenic mice ubiquitously expressing enhanced green fluorescence protein (EGFP; Okabe et al., 1997) was used to distinguish mesenchyme-derived cells from feeder cells, and single cells sorted by flow cytometry were cultured at a low cell density on 3T3Wnt4. This culture condition resulted in the formation of sheet-like colonies not formed on 3T3lacZ (Fig. 1A, upper panels), while scattered fibroblast-like cells were observed in both conditions (Fig. 1A, lower panels, arrows). Colonies were not formed in the presence of frizzled (Fz)-Fc chimeric protein, a Wnt inhibitor, thus confirming an essential role of Wnt4 for colony formation (Fig. 1B). Colonies were not formed by culturing in the conditioned medium from 3T3Wnt4 without feeder cells (data not shown). Colonies were also formed on 3T3Wnt3a, but not in feeder-free conditions using a purified recombinant Wnt3a protein (data not shown). These data suggested the requirement of other signals from 3T3 cells, in addition to the Wnt signals for the colony formation. In the presence of serum, colonies were not formed even on 3T3Wnt4, and some factors in the serum might prevent colony formation (data not shown). When colonies on 3T3Wnt4 were dissociated and plated onto fresh feeder cells at day 10 of culture, few colonies were obtained, and maintenance of these colonies could not be achieved (data not shown). When we tried colony-formation by using polycarbonate filters, which separate mesenchymal cells from the feeder layer, colonies were formed but the number of colonies formed was much smaller than that formed by directly culturing on feeder cells (data not shown).

To characterize the molecular profiles of the colonies, genes expressed in the metanephric mesenchyme were examined by RT-PCR using RNA from the colonies together with 3T3Wnt4 (Fig. 1C). All the mesenchymal genes examined (Pax2, Lim1, Eya1, Sall1, WT1, Hoxa11, Gdnf, integrin α8, integrin α3, Ncam, E-cadherin and K-cadherin) were expressed, and the expression continued to day 20 (Fig. 1C, lanes 3-5). By contrast, when cultured on 3T3lacZ, the expression of these genes was below the detection level (lanes 7-9). The expression of ureteric bud markers (Ret and Hoxb7) were not detected in mesenchyme separated from ureteric bud, suggesting that the separation was successful (lane 1). To determine the potential for differentiation within the colonies, markers for terminally differentiated epithelia in glomeruli (podocyte), proximal or distal tubules, and the loop of Henle were also examined (glomeruli: α-actinin-4, CD2-AP, P-cadherin, podoplanin and podocalyxin; proximal tubule: Aqp1, Clc5, cubulin, megalin and Sglt1; Henle’s loop: Bmt1 and Nkcc2; Henle’s loop or distal tubule: Clc2, polycystin 2, and Romk2; distal tubule: ENaC, Na/Ca exchanger and polycystin 1. These markers encode: (1) cytoskeletal or structural proteins: α-actinin-4, CD2-AP, P-cadherin, podoplanin and podocalyxin; (2) transcription factor: Bmt1; (3) water or ion channels: Aqp1, Clc5, Clc2, Romk2, ENaC, and polycystin 1 and 2; and (4) transporters: cubulin, megalin, Sglt1, Nkcc2 and Na/Ca exchanger. As shown in Fig. 1C, almost all the genes examined were expressed at day 20 on 3T3Wnt4 (lane 5), while these markers were not expressed on 3T3lacZ (lanes 7-9). To ascertain that these genes were expressed by the colony-forming cells, colonies were formed from GFP transgenic mesenchyme, and cells expressing GFP were separated from feeder layers by using flow cytometry sorting. RT-PCR using RNA from these cells suggested that the marker genes examined were indeed expressed by colony-forming cells (lane 10). Furthermore, we made use of immunocytochemistry and found that Pax2 (Fig. 1D-F), E-cadherin (Fig. 1G-I), Sall1 (Fig. 1J,K), and Aqp1 (Fig. 1L,M) were expressed on colonies. The expression of Pax2 and E-cadherin was not detected on immunocytochemistry at day 3, and was subsequently upregulated by day 10, which was consistent with the result of RT-PCR (Fig. 1D,E,G,H). These data suggest that dissociated cells from the metanephric mesenchyme form colonies on 3T3Wnt4 feeder cells in serum-free conditions, and that these colonies contain differentiated epithelia expressing marker genes for epithelia in glomeruli (podocyte), proximal or distal tubules, and the loop of Henle.

Colonies are derived from a single multipotent renal progenitor
To confirm that these colonies were derived from a single cell, each single cell sorted from the EGFP transgenic mesenchyme was cultured in an individual well of 96-well plates coated with 3T3Wnt4. The sheet-like colony was found in 166 wells out of a total of 1632 (10.2%) from three independent experiments (Fig. 2A).
To examine the multilineage differentiation of single cell-derived colonies, RT-PCR was done for 22 independent wells containing a colony at day 20. The representative data from three colonies are shown in Fig. 2B (lanes 1-3). Although variation existed between colonies, all the colonies expressed markers for each of the three segments: glomerular podocytes, proximal and distal tubules, and the loop of Henle. Double staining using PNA and LTL, specific to glomerular podocytes and the proximal renal tubule, respectively, showed that adult kidney (8 weeks old) contained three kinds of cells; single-positive for PNA (those in the glomerulus); single-positive for LTL (those in the proximal renal tubule); and double-negative for LTL or PNA (Fig. 2C, left panel). Similarly, a single cell-derived colony at day 20 contained these three kinds of cells (Fig. 2C, right panel). With a combination of LTL and E-cadherin, at least three cell types were observed in adult kidney (Fig. 2D, left panel) and in a single cell-derived colony (right panel): cells strongly expressing only E-cadherin characteristic of distal renal tubules (Fig. 2D, arrows), and LTL-positive or -negative cells, with a faint expression of E-cadherin in the cell boundary. These results suggest that a colony was derived from a single progenitor, with multipotent differentiating capacity into epithelial cells in glomeruli, proximal and distal tubule, and the loop of Henle.

Colony-forming progenitors exist in the Sall1-GFP<sup>high</sup> subpopulation of the metanephros

We next attempted to identify prospectively the renal progenitor cells using Sall1-GFP<sup>high</sup> knock-in mice (Takasato et al., 2004). As Sall1 is expressed in mesenchyme-derived tissues, GFP was detected in the mesenchyme around the ureteric bud at E11.5 in the Sall1-GFP<sup>heterozygous</sup> mouse (Fig. 3A, arrows). At E17.5, GFP-expressing cells were observed in the mesenchyme near the surface, as well as in C- or S-shaped bodies, and parts of renal tubules (Fig. 3B). By flow-cytometrical analysis, three subpopulations were fractionated based on the expression of Sall1-GFP: Sall1-GFP<sup>high</sup>
Sall1-GFP<sup>low</sup> and Sall1-GFP<sup>negative</sup> (Fig. 3C), and cells in these subpopulations were separated by flow cytometry sorting to be characterized using RT-PCR. As shown in Fig. 3D, Sall1-GFP<sup>high</sup> cells expressed Sall1 and Pax2. Sall1-GFP<sup>low</sup> cells expressed markers of stroma (Foxd1, previously known as BF2), endothelia (Flk1 and VE-cadherin), and blood cell (Cd45), in addition to Sall1 and Pax2. Sall1-GFP<sup>negative</sup> cells expressed Flk1 and Cd45. The markers of fully differentiated renal epithelia were not expressed in these three populations. These data suggested that cells of stromal lineage were included in cell populations weakly expressing Sall1 and that those of hemangiogenic lineage were included in both Sall1-GFP<sup>low</sup> and Sall1-GFP<sup>negative</sup> populations. Then, the numbers of the colony-forming progenitors in each subpopulation were examined using the low-density culture on 3T3Wnt4 (Fig. 3E). At E11.5, colonies were formed exclusively from the Sall1-GFP<sup>high</sup> population, and not from Sall1-GFP<sup>low</sup> or Sall1-GFP<sup>negative</sup> populations. At E14.5 and 17.5, colonies were also formed only from Sall1-GFP<sup>high</sup> subpopulations, but the frequency of colony-forming progenitors decreased as gestation proceeded. These results indicate that renal progenitors with multipotent differentiating capacity are included in cell populations strongly expressing Sall1 throughout gestation periods.

**Sall1-GFP<sup>high</sup> mesenchyme reconstitutes a three-dimensional structure in organ culture**

We next examined the in vitro differentiation capacity of three subpopulations in E11.5 mesenchyme by modifying organ culture of mesenchyme rudiments (Grostein, 1953; Kispert et al., 1998). Sall1-GFP<sup>high</sup>, Sall1-GFP<sup>low</sup> and Sall1-GFP<sup>negative</sup> cells were separated by flow cytometry, aggregated to form a cell pellet by centrifugation and cultured on 3T3Wnt4 feeder cells in an organ culture setting. Starting from day 3 in culture, tubulogenesis was observed only in the aggregate of the Sall1-GFP<sup>high</sup> population (Fig. 4A, upper panels), while that from Sall1-GFP<sup>low</sup> or Sall1-GFP<sup>negative</sup> did not differentiate and disappeared by day 7 (Fig. 4A, lower panels; data of Sall1-GFP<sup>negative</sup>, not shown). In sections of the Sall1-GFP<sup>high</sup> aggregate (Fig. 4B), many tubule- (t) and glomerulus-like structures (g) were observed, and the expression of markers for glomerular podocyte (Wt1, Fig. 1C, red) and proximal tubule (LTL, green) was confirmed by confocal microscopy. These data suggest that only Sall1-GFP<sup>high</sup> cells differentiate into renal epithelia in vitro in a three-dimensional setting, in addition to forming colonies.

**Colonies size is affected by the absence of Sall1**

To investigate the role of Sall1 in colony formation, mesenchymal cells from Sall1<sup>+/+</sup>, Sall1<sup>+/–</sup> and Sall1<sup>–/–</sup> embryos at E11.5, which were obtained from intercrosses of Sall1-GFP mouse, were plated on 3T3Wnt4 feeder cells at a low density. Ten days after culture, double immunostaining using anti-GFP and anti-E-cadherin antibodies was done to strengthen the green fluorescence and to examine the expression of E-cadherin, respectively (Fig. 5). The numbers of colonies formed were not significantly different among wild-type, heterozygous and homozygous mesenchyme, suggesting that colony-forming progenitors do exist and are not decreased in the absence of Sall1 (data not shown). Colonies derived from Sall1<sup>+/+</sup> wild-type mesenchyme were not stained with GFP (Fig. 5A), while Sall1<sup>+/–</sup> and Sall1<sup>–/–</sup> colonies were positive for GFP (Fig. 5C,E, green), indicating that Sall1 itself is not required for Sall1 promoter activity. Colonies from all three groups (Sall1<sup>+/+</sup>, Sall1<sup>+/–</sup> and Sall1<sup>–/–</sup>) were also positive for E-cadherin (Fig. 5B,D,F), suggesting that differentiation (mesenchymal-to-epithelial transformation) may not be impaired in the absence of Sall1. Indeed, marker gene expression for terminally differentiated epithelia in glomeruli and renal tubules was not changed among Sall1<sup>+/+</sup>, Sall1<sup>+/–</sup> and Sall1<sup>–/–</sup> colonies on RT-PCR analyses (data not shown). By contrast, the size of Sall1<sup>–/–</sup> colonies (Fig. 5E,F) was significantly smaller than Sall1<sup>+/+</sup> and Sall1<sup>+/–</sup> colonies (Fig. 5B-D), and this was confirmed statistically (Table 1). Thus, Sall1 is not

---

**Table 1. Colonies derived from Sall1-mutant metanephric mesenchyme**

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Embryos&lt;sup&gt;*&lt;/sup&gt;</th>
<th>Area at day 10 (mean±sd) (μm&lt;sup&gt;2&lt;/sup&gt;) (n=60)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>+/+</td>
<td>2</td>
<td>16,118±7219</td>
<td></td>
</tr>
<tr>
<td>+/-</td>
<td>2</td>
<td>16,318±7473</td>
<td>0.44</td>
</tr>
<tr>
<td>–/–</td>
<td>2</td>
<td>51,40±2071</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

<sup>*</sup>Number of embryos examined. P values were analyzed against wild type (+/+). *|t*-test. Embryos of a total of four litters were analyzed in this way. Representative data from one experiment are shown.
required for generation or differentiation of renal progenitors, but the colony size is affected by Sall1 absence. This is consistent with our previous report that Sall1-deficient mesenchyme is competent with respect to epithelial differentiation tested by spinal cord recombination (Nishinakamura et al., 2001). In the spinal cord recombination experiments, Sall1-deficient mesenchyme was consistently smaller than wild-type mesenchyme, but this could be due to differences in the initial size of the mesenchyme. Using the colony-forming assay starting from a single cell, we now show that Sall1 is indeed required for the colony from the mesenchyme to develop into a normal size.

The PCP pathway regulates colony size and the differentiation of colony-forming cells

By combining the colony-forming assay set up in this study and gene transfer using retroviral vector pMY-IRES-EGFP (Morita et al., 2000; Kitamura et al., 2003), we observed EGFP expression in 12.9% of colony-forming progenitor cells (116 colonies expressing green fluorescence per total of 896 colonies formed from three independent experiments). Thus, the colony-forming assay in this study enables us to investigate direct effects of reagents and gene transduction on colony-forming progenitor cells, allowing us to examine the roles of Wnt and its downstream branches in kidney development. Positive immunostaining of the colonies for activated JNK1 and 2 indicated that the JNK branch of the PCP pathways (Boutros et al., 1998) may be activated downstream of Wnt4 (Fig. 6A). Indeed, the addition of two kinds of JNK inhibitor (JNKI1 and JNKI2) (Bonny et al., 2001; Bennett et al., 2001) gave rise to smaller colonies than did the control without reagents (Fig. 6B,C, colonies from EGFP transgenic mesenchyme, Table 2). The result of control experiments using the HIV-TAT peptide excluded the possibility that the effects of JNKI1 were derived from non-specific toxicity of the peptide constituting the inhibitor (Fig. 6B). We then investigated effects of both activation and inactivation of Rac1, one of the Rho family GTPases implicated in PCP pathways (Habas et al., 2003), on colony formation. Cells from wild-type E11.5 mesenchyme were transduced with both constitutively active (CA) and dominant-negative (DN) forms of Rac1 using the retroviral vector pMY-IRES-EGFP. Colonies consisting of cells expressing both EGFP and CA-Rac1 were larger than those transduced with pMY-IRES-EGFP controls (Fig. 6D, Table 3), suggesting positive effects on colony size. By contrast, the transduction of DN-Rac1 gave rise to smaller colonies.

Table 2. Effects of reagents on the area of colony

<table>
<thead>
<tr>
<th>Reagent</th>
<th>n</th>
<th>Area at day 10 (mean±s.d.) (µm²)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (without reagents)</td>
<td>20</td>
<td>35,429±15,132</td>
<td>0.25</td>
</tr>
<tr>
<td>HIV-TAT peptide</td>
<td>20</td>
<td>38,198±11,357</td>
<td></td>
</tr>
<tr>
<td>JNK inhibitor 1</td>
<td>20</td>
<td>777±14,520</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Control (without reagents)</td>
<td>30</td>
<td>36,330±15,065</td>
<td></td>
</tr>
<tr>
<td>JNK inhibitor 2</td>
<td>20</td>
<td>668±23,834</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Y27,632</td>
<td>20</td>
<td>92,359±24,768</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LiCl</td>
<td>20</td>
<td>568±23,535</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>BIO</td>
<td>30</td>
<td>743±4897</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Dkk-1</td>
<td>20</td>
<td>31,998±12,566</td>
<td>0.147</td>
</tr>
</tbody>
</table>

n, number of colonies measured. P values were analyzed against control using a t-test. For each reagent, more than three independent experiments were performed. Representative data from one experiment are shown.
colonies than did the controls (Fig. 6D, Table 3). The numbers of colonies formed were not significantly changed either with the addition of inhibitors or with gene transduction (data not shown). These data indicate that Rac and JNK pathways positively regulate colony size.

By contrast, inactivation of the Rho/Rho-associated protein kinase (ROCK) pathway, another branch of PCP (Strutt et al., 1997; Winter et al., 2001; Habas et al., 2001; Habas et al., 2003), with the addition of ROCK inhibitor, Y27,632 (Uehata et al., 1997) (Fig. 6E, colonies from EGFP transgenic mesenchyme, Table 2), or the transduction of DN-RhoA increased the colony size (Fig. 6F, Table 3), while the activation with CA-RhoA decreased it (Fig. 6F, Table 3). Activation of β-catenin signaling both with the addition of two kinds of glycogen synthetase kinase (GSK)-3 inhibitors, lithium chloride (LiCl) (Klein and Melton, 1996) (Fig. 6G, colonies from EGFP transgenic mesenchyme, Table 2) and (2’H11032Z, 3’H11032E)-6-Bromoindirubin-3-oxime (BIO) (Sato et al., 2004) (Table 2) and with the transduction of the active form of β-catenin (Fig. 6H, Table 3) gave rise to smaller colonies. However, inactivation of the β-catenin pathway with the addition of recombinant dickkopf homolog 1 (Dkk-1), a specific inhibitor of the β-catenin pathway (Glinka et al., 1998) and with the transduction of axin (Zeng et al., 1997) exerted no significant effects on colony formation (Tables 2, 3). These data suggest inhibitory roles of Rho/ROCK and β-catenin pathways in regulating colony size.

RT-PCR analysis showed that the expression of marker genes (E-cadherin, P-cadherin, podocalyxin, AQP1, CLC5, Brn1, Nkcc2, ENaC and Clck2) was inhibited with the addition of JNK inhibitors (Fig. 6I). Although the addition of LiCl and JNK inhibitors resulted in a decreased size of colonies to the same extent, immunostaining confirmed that E-cadherin was lost with JNK inhibitors 1 and 2, but not with LiCl (Fig. 6J, colonies from EGFP transgenic mesenchyme). Thus, the JNK pathway is likely not only to regulate colony size but also to be involved in epithelialization (mesenchymal-to-epithelial transformation) of colony-forming progenitors.

Table 3. Effects of gene transduction on the area of colony

<table>
<thead>
<tr>
<th>Gene transduced</th>
<th>n</th>
<th>Area at day 20 (mean±s.d.) (µm²)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (vector)</td>
<td>23</td>
<td>49,666±32,111</td>
<td></td>
</tr>
<tr>
<td>CA-Rac1</td>
<td>12</td>
<td>83,990±52,619</td>
<td>0.01</td>
</tr>
<tr>
<td>DN-Rac1</td>
<td>24</td>
<td>23,045±22,791</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>CA-RhoA</td>
<td>12</td>
<td>25,658±19,205</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td>DN-RhoA</td>
<td>21</td>
<td>89,723±49,989</td>
<td>0.001</td>
</tr>
<tr>
<td>Control (vector)</td>
<td>38</td>
<td>62,013±31,212</td>
<td></td>
</tr>
<tr>
<td>Active-β-catenin</td>
<td>20</td>
<td>23,241±21,685</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Axin</td>
<td>22</td>
<td>65,352±27,675</td>
<td>0.34</td>
</tr>
</tbody>
</table>

n, number of colonies measured. P values were analyzed against control using a t-test. Data from three independent experiments is shown.
The PCP pathway is involved in tubulogenesis in organ culture

To examine if the results described above were consistent with kidney formation in vivo, we tested the effect of the reagents on whole metanephroi (Fig. 7A-E) and mesenchyme rudiments (F-J) in an organ culture setting. After 7 days of culture, the size of kidney structures was measured. As compared with the control explants cultured without reagents (Fig. 7A,F), the addition of JNK inhibitor 1 (Fig. 7B,G) and JNK inhibitor 2 (Fig. 7C,H) and LiCl (Fig. 7E,J) resulted in a decrease in the size of kidney structures developed, while the addition of ROCK inhibitor Y27,632 (Fig. 7D,I) gave rise to larger ones. These findings were observed both in whole kidney and in mesenchyme rudiments, and were confirmed statistically (Table 4).

We also evaluated the effect of the reagents on tubule formation and branching of ureteric bud by staining with an antibody against secreted frizzled-related protein 5 (FRZB), which is a marker for ureteric bud branching. The addition of JNK inhibitors or LiCl resulted in a decrease in the size of kidney structures developed, while the addition of ROCK inhibitor Y27,632 gave rise to larger ones. These findings were observed both in whole kidney and in mesenchyme rudiments, and were confirmed statistically (Table 4).

Table 4. Effects of reagents on the area of organ culture

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Whole metanephroi</th>
<th>Mesenchymal rudiments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Area at day 7</td>
<td>Area at day 7</td>
</tr>
<tr>
<td></td>
<td>(mean ± s.d.) (mm²)</td>
<td>(mean ± s.d.) (mm²)</td>
</tr>
<tr>
<td>Control (without reagents)</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>HIV-TAT peptide</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>JNK inhibitor 1</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>JNK inhibitor 2</td>
<td>6</td>
<td>7</td>
</tr>
<tr>
<td>Y27,632</td>
<td>5</td>
<td>7</td>
</tr>
<tr>
<td>LiCl</td>
<td>5</td>
<td>7</td>
</tr>
</tbody>
</table>

n, number of explants measured. P values were analyzed against control by using a t-test. Data from five independent experiments each for whole metanephroi and mesenchyme rudiments are shown.
protein 2 (sFRP2; Fig. 7K-N, red), the gene expressed only in newly formed tubular epithelia (Lescher et al., 1998), and DBA (Fig. 7K,L, green), respectively. While some tubules expressing sFRP2 were found in explants treated with control HIV-TAT peptide (Fig. 7K,M, arrows), they were lost with JNK inhibitor 1 both in whole metanephroi (Fig. 7L) and in mesenchyme explants (Fig. 7N), suggesting the involvement of JNK pathways in mesenchymal-to-epithelial transformation in organ culture. By contrast, branching of ureteric bud was proportional to the size of explants, and there were no specific effects of the reagents observed on the ureteric bud itself (data not shown). These findings were consistent with the results observed in the colony-forming assay (Fig. 6, Table 2). Thus our colony-forming assay, which enables analysis at a single cell level, could be used for examining mechanisms of three-dimensional kidney development.

DISCUSSION

Renal progenitors defined by colony-forming assay

In this study, we provide evidence, using in vitro clonal analysis combined with flow cytometry, for the presence of progenitor cells in the fetal mouse kidney. Results of staining with PNA, LTL and E-cadherin, and of RT-PCR showed the differentiating capacity of a single Sall1-GFPhigh cell into glomerular epithelia (podocyte), proximal and distal tubule, respectively. In addition to lineage-marker expression, both glomerulus- and tubule-like structures were reconstituted by Sall1-GFPhigh cells, supporting their differentiation ability. A multipotent renal stem cell line has been isolated from E11.5 mesenchyme utilizing immortalization with T antigen of SV40 virus (Oliver et al., 2002). The cell line expresses marker genes of endothelia and smooth muscle cells with treatment of TGFβ1, in addition to gene expression of mesenchymal and renal epithelia. It has not been known, however, whether they normally resided in the fetal kidney or accidentally emerged by the influence of the process with immortalization. In our colony-forming system, gene expression of endothelia and smooth muscle cells was not observed, and expression of Foxd1 (BF2), a marker gene specific to stroma, a third cell population included in metanephros was not found (data not shown). Thus, it remains to be elucidated whether embryonic kidney contains stem cells that can differentiate into endothelium, smooth muscle or stroma in addition to epithelia of glomerulus and renal tubules.

The renal progenitors defined by our colony-forming assay are included in cell populations strongly expressing Sall1 throughout gestation periods, and they might continue to reside in the outer layer of embryonic kidney, where undifferentiated metanephric mesenchyme resides and strongly expresses Sall1 (Fig. 3B). As shown in Table 5, the total cell numbers of metanephros increased and the frequency of colony-forming Sall1-GFPhigh cells decreased as gestation proceeded. Interestingly, the calculated numbers of the colony-forming cells remained almost constant throughout gestation periods (400-800 cells/embryonic kidney). The amplification of these progenitors might not occur in the embryonic kidney. One interesting question is whether they continue to remain in the adult kidney. From 8-week-old mice, however, colonies were not formed under the same culture conditions (data not shown). Renal progenitors defined by our colony-forming assay might be lost by the time kidney development is complete.

Table 5. Calculated number of colony-forming progenitors in embryonic kidney

<table>
<thead>
<tr>
<th>E11.5</th>
<th>E14.5</th>
<th>E17.5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total cell number of kidney (×10⁵)</td>
<td>0.77±0.24</td>
<td>2.4±2.3</td>
</tr>
<tr>
<td>Sall1-GFPhigh cells in kidney (%)</td>
<td>23.6±1.7</td>
<td>46.3±2.1</td>
</tr>
<tr>
<td>Colony formation in Sall1-GFPhigh cells (%)</td>
<td>32.2±2.8</td>
<td>0.79±0.18</td>
</tr>
<tr>
<td>Calculated numbers of colony-forming cells*</td>
<td>585.1</td>
<td>782.7</td>
</tr>
</tbody>
</table>

Means±s.d. (from five independent experiments each for E11.5, E14.5, and E17.5).

*Numbers of colony-forming cells were calculated by multiplying the means of the three values above.
Analysis of gene function in kidney development by colony-forming assay

The knowledge of gene function in kidney development has mainly been obtained from analyses using knockout mice, while experimental systems that investigate gene function in individual cells of metanephros have been lacking. By setting up a novel system combining colony formation from a single cell and gene transduction using a retroviral vector, our culture system enables the direct observation of effects of reagents and gene transduction on colony-forming progenitor cells. As similar results were obtained from organ culture experiments (Fig. 7), it is less likely that the cellular behavior observed in our colony-assay system might be artifactual.

Mice lacking the constituent genes involved in downstream branches of Wnt signaling pathways often show early embryonic lethality, such as Rac1 (Sugihara et al., 1998), Jnk1 and Jnk2 (Kuan et al., 1999), β-catenin (Haegele et al., 1995), axin (Zeng et al., 1997), and their functions in kidney morphogenesis remain largely unknown. Using our culture system, functions of these genes in metanephros development were elucidated. Furthermore, experiments for colony formation from mesenchyme of Sall1-mutant embryos demonstrated the roles of Sall1 for the colony size. Thus, the colony-assay system set up in this study can also be applied to the analysis of genetic mouse models.

Roles of PCP pathway in kidney development

Among downstream branches of Wnt4 signal, we found that Rac- and JNK-dependent PCP pathways positively regulated the colony size and the differentiation of colony-forming cells. This result is compatible with several previous reports (Du et al., 1995; Ungar et al., 1995; Maretto et al., 2003). In frogs and fish, the Wnt4 family might antagonistically regulate the growth of progenitors in kidney development. Recently the roles of the JNK pathway in epithelialization, as well as in regulation of colony size. Common mechanisms regulating epithelial morphogenesis might underlie these processes. The PCP pathways, including the Rho family of small GTPases and JNK, control several developmental processes, mainly by regulating cell cytoskeletons, such as the polarity of hairs on the epidermal cells of Drosophila wings, the arrangement of ommatidial cells of Drosophila eyes, the polarity of stereocilia in the inner ears of mammals, and convergent extension in Xenopus and zebrafish (Veenman et al., 2003; Wallingford et al., 2002). In addition to these processes, we provide a novel hypothesis of the involvement of the PCP pathways in kidney development.

In summary, we set up a novel colony-forming assay by which we demonstrated the presence and the frequency of multipotent progenitor cells in embryonic kidneys. This assay would serve as a useful tool for analyzing differentiation mechanisms in the kidney at a single cell level, taking advantage of the facility of gene transfer.

We thank Dr M. Okabe for providing EGFP transgenic mice, Dr A. Kikuchi for pBSKS+-Axin, Dr A. Nagauchi for pUC-EGF-4x-β-catenin55-3HA, Dr H. Koido for pCAGIP-Flag-Rac1 and RhoA, Dr T. Kitamura for pMY/IRE-EGFP and PLAT-E, Y. Morita for technical support for FACS, and Dr C. Kobayashi for critically reading the manuscript. This work was partly supported by the Ministry of Health, Labor, and Welfare of Japan.

References


