Small Cytoskeleton-Associated Molecule, Fibroblast Growth Factor Receptor 1 Oncogene Partner 2/Wound Inducible Transcript-3.0 (FGFR1OP2/wit3.0), Facilitates Fibroblast-Driven Wound Closure

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Wounds created in the oral cavity heal rapidly and leave minimal scarring. We have examined a role of a previously isolated cDNA from oral wounds encoding wound inducible transcript-3.0 (wit3.0), also known as fibroblast growth factor receptor 1 oncogene partner 2 (FGFR1OP2). FGFR1OP2/wit3.0 was highly expressed in oral wound fibroblasts without noticeable up-regulation of α-smooth muscle actin. In silico analyses, denaturing and nondenaturing gel Western blot, and immunocytology together demonstrated that FGFR1OP2/wit3.0 were able to dimerize and oligomerize through coiled-coil structures and appeared to associate with cytoskeleton networks in oral wound fibroblasts. Overexpression of FGFR1OP2/wit3.0 increased the floating collagen gel contraction of naïve oral fibroblasts to the level of oral wound fibroblasts, which was in turn attenuated by small-interfering RNA knockdown. The FGFR1OP2/wit3.0 synthesis did not affect the expression of collagen I as well as procontractile peptides such as α-smooth muscle actin, and transforming growth factor-β1 had no effect on FGFR1OP2/wit3.0 expression. Fibroblastic cells derived from embryonic stem cells carrying FGFR1OP2/wit3.0 (+/−) mutation showed significant retardation in cell migration. Thus, we postulate that FGFR1OP2/wit3.0 may regulate cell motility and stimulate wound closure. FGFR1OP2/wit3.0 was not up-regulated during skin wound healing; however, when treated with FGFR1OP2/wit3.0-expression vector, the skin wound closure was significantly accelerated, resulting in the limited granulation tissue formation. Our data suggest that FGFR1OP2/wit3.0 may possess a therapeutic potential for wound management. (Am J Pathol 2010, 176:108−121; DOI: 10.2353/ajpath.2010.090256)

An excisional wound created in skin does not immediately close without surgical assistance and often results in scar formation.1,2 Responding to wound-induced cytokines and growth factors, a subset of skin fibroblasts undergoes transdifferentiation and acquires the expression of α-smooth muscle actin (α-SMA). These α-SMA-expressing myofibroblasts are believed to be responsible for the late stage wound contraction.3−5 On the contrary, clinical observations and animal studies consistently demonstrate that wounds in the oral cavity heal with minimal scarring. The inflammatory response in wounded oral mucosa is less intensive than in skin,6 and the expression of cytokines is also low in oral wounds during the healing period.7−11 Saliva contains a group of cytokines,12,13 which may provide a unique oral environment suitable for wound healing. However, skin grafts transposed into the oral cavity maintain the skin wound healing phenotype,14 indicating that constitutive oral cellular phenotypes, not the oral environment, may play the primary role in the oral wound healing.

It has been pointed out that there are similarities between oral and fetal skin wound healing. Uniquely, both wounds rapidly reduce the wound size through approx-

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imation of wound margins during early healing stages.\textsuperscript{11,15–18} Wounded fetal skin epithelial cells do not proliferate but exhibit the unique “purse-string” contraction, which primarily contributes to the rapid wound closure.\textsuperscript{19} Oral wound epithelium undergoes active proliferation and migration, and does not appear to initiate the “purse-string” wound closure. Thus, the rapid and spontaneous oral wound closure must be achieved by other mechanisms. A genome-wide gene expression study using cDNA microarray revealed various distinctive gene expression patterns among fetal fibroblasts as well as adult fibroblasts isolated from different sites. The supervised hierarchical clustering analyses indicated that the gene expression profile of fetal fibroblasts was closest to that of oral fibroblasts.\textsuperscript{20} Oral fibroblasts are originated from the neural crest-derived ectomesenchymal cells, and have been postulated to play a key role in the accelerated oral wound closure.

Previously, we have isolated a unique cDNA from tooth extraction wound tissue encoding wound inducible transcript-3.0 (wit3.0).\textsuperscript{21} The GenBank database search has identified that wit3.0 is fibroblast growth factor receptor-1 oncogene partner-2 (FGFR1OP2). The FGFR1OP2/wit3.0 allele is composed of seven exons and generates two isoforms of 215-amino acid (FGFR1OP2/wit3.0a) and 253-amino acid (FGFR1OP2/wit3.0b) through alternative splicing of exon 5.\textsuperscript{22} The amino-terminal sequence of FGFR1OP2/wit3.0 encoded by the first four exons has been shown to facilitate the ligand-independent dimer formation of the FGFR1 tyrosin kinase domain in the cytogenetic abnormality (12;8) (p11; p11p22) (Figure 1A), having caused acute myeloid leukemia.\textsuperscript{23}

FGFR1OP2/wit3.0 has been identified as a small cytoplasmic peptide expressed by oral wound fibroblasts,\textsuperscript{22} however, the function of native FGFR1OP2/wit3.0 has not been elucidated. We report here that FGFR1OP2/wit3.0 is a novel cytoskeleton molecule, with a regulatory role in fibroblast migration and rapid wound closure, a hallmark of oral wound healing. The transduction of FGFR1OP2/wit3.0 in mouse skin wounds appeared to induce oral wound-like healing phenotypes: rapid closure and minimal scarring. Our data suggest that FGFR1OP2/wit3.0 may play an important role in favorable oral wound healing and possess a therapeutic potential for wound management.

Materials and Methods

Generation of FGFR1OP2/wit3.0 Monospecific Antibody

The conserved sequence of human, mouse, and rat FGFR1OP2/wit3.0 [45]QYQEEIQELNEVARHRPRS [63] was selected for polyclonal antibody synthesis (Quality Controlled Biochemicals, Hopkinton, MA), resulting in the isolation of D1042. For the specificity evaluation, Western blot of cytoplasmic protein of NIH3T3 fibroblasts transfected with p3x-FLAG-human FGFR1OP2/wit3.0b expression plasmid was analyzed by D1042 antibody and M2 antibody recognizing 3x-FLAG epitope (Sigma-Aldrich, St. Louis, MO).

FGFR1OP2/wit3.0 Immunohistochemistry in an Oral Wound

An oral wound was created by simple tooth extraction from maxilla of adult (8 to 10 weeks old) male Sprague-Dawley rats following the previously described protocol.\textsuperscript{24} Seven days after extraction of three molars, maxilla were harvested, fixed in 10% buffered formalin and decalcified (Cal-Ex, Fischer Scientific, Pittsburgh, PA). Eight-micrometer thick paraffin sections were generated through the second molar. After the epitope retrieval by using a pressure cooker and Tris Buffer (pH9.0), sections were incubated in primary antibody (D1042, 1:100 to 1:500) and then in horseradish peroxidase conjugated secondary antibody (Dako Envision, Carpenteria, CA). Sections were treated with diaminobenzamine and stained with hematoxylin. The neighboring sections were separately incubated with monoclonal antibody against α-SMA (Clone 1A4, Sigma-Aldrich). Some sections were stained with H&E for references.

Expression of FGFR1OP2/wit3.0 in the Rat Oral and Skin Wounds

Oral mucosal tissue was harvested from a tooth extraction wound (Day 7) or from unwounded alveolus of rat maxilla. The dorsal rat skin was collected from full-thickness excisional wound (Day 7) with a biopsy punch (8 mm in diameter; Skalr Tru-Punch, Sklar Instruments, West Chester, PA) or from the unwounded area. Tissue specimens were immediately frozen in liquid nitrogen and total RNA samples were prepared. The steady state mRNA levels of FGFR1OP2/wit3.0a, FGFR1OP2/wit3.0b, and α-SMA were determined by the Taqman-based quantitative real-time RT-PCR. Student’s t test was performed against the unwounded tissue, and statistical significance was accepted for $P < 0.05$.

In Silico Search on FGFR1OP2/wit3.0 Peptide Structure

The human FGFR1OP2/wit3.0b peptide sequence was submitted to online computational software COILS: Prediction of Coiled Coil Regions in Proteins (http://www.ch.embnet.org/software/COILS_form.html, last accessed August 26, 2006) and the Human Protein Reference Database (http://www.hprd.org/summary?hprd_id=10589&isoform_id=10589_1&isoform_name=Isoform_1, last accessed August 26, 2006) for consensus functional structure searches, and the Imperial College Protein Homology/Analogy Recognition Engine (http://www.sbg.bio.ic.ac.uk/~phyre/, last accessed August 26, 2006) for three-dimensional molecular structure prediction.
Cell Cultures of Oral Fibroblasts, Oral Wound Fibroblasts, and Skin Fibroblasts

The tissue samples of rat gingiva, tooth extraction wound, and skin were minced in PBS with 1X antibiotic-antimycotics (Invitrogen, Carlsbad, CA) and treated with Collagenase A (200 unit/ml; Sigma-Aldrich) in complete fibroblast culture medium: Dulbecco’s modified Eagle’s medium, antibiotic-antimycotics solution, and 10% of fetal bovine serum (FBS). After incubation for 5 days, the epithelial cells were separated by centrifugation at 100xg for 3 minutes. The pellet was resuspended with complete fibroblast medium and cultured for an additional 2 weeks.

Cells from three to five passages were used in the following experiments.

Denature and Nondenature Gel Western Blot Analyses

Cytoplasmic proteins from oral fibroblasts and oral wound fibroblasts were extracted by using NE-PER Nuclear and Cytoplasmic Extraction Reagent (Pierce, Rockford, IL). For the denatured condition, 3.5 μg of protein samples were mixed with Laemmli loading buffer containing SDS and β-mercaptoethanol and boiled at 100°C for 10 minutes and loaded on a 12% Tris-HCl Ready Gel (Bio-Rad, Hercules, CA). For the nondenatured condition, the protein sample was mixed with Native Sample Buffer (Bio-Rad) for electrophoresis. The protein on the gel was transferred to polyvinylidene difluoride membrane (Bio-Rad) and incubated with D1045 antibody (1:100) for 1
hour at room temperature. After secondary antibody incubation, the membrane was visualized by using a chemiluminescence standard protocol (Pierce) and a CCD imaging system (LAS-3000, Fujifilm, Stamford, CT). In some experiments, the polyvinylidene difluoride membrane was stripped and reprobed with the small ubiquitin-like modifier-1 (SUMO-1) or SUMO-2/3 antibodies (Santa Cruz Biotechnology, Santa Cruz, CA).

Immunocytology of Oral Fibroblasts

Oral fibroblasts and oral wound fibroblasts were cultured in glass chamber slides (Lab Tek, Fisher Scientific) in the complete fibroblast culture medium for 24 hours. Cells were fixed with freshly prepared 3.7% formaldehyde (methanol free) followed by the treatment with 0.2% Triton X-100 and Image-IT FX signal enhancer (Invitrogen). The slides were incubated with D1042 antibody (1:100) and then with Alexa Fluor 488-conjugated goat anti-rabbit IgG antibody (1:1000) (Invitrogen). The slides were further treated with Alexa Fluor 568-conjugated phallotoxins and 4',6-diamidino-2-phenylindole (DAPI). A confocal laser scanning microscope (LSM 310, Carl Zeiss, Novi, MI) was used to scan 1-μm focal layers of each specimen.

Floating Collagen Gel Contraction Assay

Fibroblasts were seeded into the collagen gel and casted in 6-well plates (1.2 × 10^5 cells/well) following the previously established protocol. Once solidified at 37°C, fibroblast/gel was released from the well. The area of fibroblast/gel complex in digitized photographs was measured (Media Cybernetics, Bethesda, MD), and the ratio of collagen gel area against the culture well was calculated at each time point. Repeated measures analysis of variance was used to determine statistical significance for P < 0.05.

Small-Interfering RNA Knockdown of FGFR1OP2/wit3.0

Small-interfering RNAs (siRNAs) targeting the FGFR1OP2/wit3.0 were designed and generated by commercially available sources (Ambion, Austin, TX; and Dharmacon, Lafayette, CO) (Table 1). Using the optimal siRNA transfection condition for primary oral wound fibroblasts determined by the glyceraldehyde-3-phosphate dehydrogenase siRNA assay (Silencer glyceraldehyde-3-phosphate dehydrogenase siRNA, Ambion), each FGFR1OP2/wit3.0 siRNA candidate was applied to oral wound fibroblasts. The efficiency and specificity of the knock-down effect was determined by Taqman-based real time RT-PCR. The siRNA (number 4 and 5)-treated oral wound fibroblasts were applied to the floating collagen gel contraction assay as described above.

Mouse Embryonic Stem Cell-Derived Fibroblastic Cells Carrying FGFR1OP2/wit3.0 (+/−) Mutation

From the International Gene Trap Consortium (www.genetrap.org, last accessed December 21, 2008), six mouse embryonic stem (ES) cell lines were identified containing a gene trap mutagenesis in the FGFR1OP2/wit3.0 allele (Table 2). C57Bl/6J ES cell line with IST10830D12 mutation (Texas A&M Institute for Genomic Medicine, Houston, TX) and wild-type ES cells were expanded on feeder cells following a standard protocol. The trypsinized ES cells were cultured in nonadherent plates (Corning Inc, Corning, NY) to form aggregates in the fibroblast differentiation medium consisted of knockout Dulbecco’s modified Eagle's medium (Invitrogen), 20% FBS, 1 mmol/L L-glutamine, 0.1 mmol/L β-mercaptoethanol, 0.1% nonessential amino acids, and 1% antibiotic-antimycotics. The aggregates were then transferred into gelatin-coated plates and the outgrowing cells were serially passaged every 5 to 7 days. Most of

Table 1. SiRNA Target Sequences for FGFR1OP2/wit3.0

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<tr>
<td>No. 3</td>
<td>Sense</td>
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<tr>
<td></td>
<td>Antisense</td>
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<td>No. 4</td>
<td>Sense</td>
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the cells in culture appeared fibroblast-like morphology after four passages. The genotype was determined by PCR, and the mRNA levels of collagen I α2 chain (Mm00483937_m1) and FGFR1OP2/wit3.0 (Mm00470836_m1) were determined by Taqman-based real time PCR.

In Vitro Wound Healing Scratch Plate Assay

The ES cell-derived fibroblastic cells with wild-type and FGFR1OP2/wit3.0 (+/+) genotypes were seeded in 12-well plates and allowed to grow in the fibroblast differentiation medium to approximately 90% confluence. In vitro wounds were created by drawing lines down the well with a 20 μl plastic pipette.27 After washing, cells were maintained in the low serum medium containing 0.5% FBS. The digital images were captured after 6 and 12 hours of incubation, and the area occupied by the cells within the scratch was measured (NIH Image J version 1.42). From the Student’s t test performed against the wild-type cells, statistical significance was accepted for \( P < 0.05 \).

3xFLAG-Human FGFR1OP2/wit3.0β Lentiviral Vector Carrying Wild-Type Sequence and Nonsynonymous Single Nucleotide Polymorphism (SNPs)

Two nonsynonymous SNPs (rs 1058701 and rs11613) were identified in the exon 5 from the National Center for Biotechnology Information SNP database, and were named SNP1 and SNP2, respectively. Using a site-directed mutagenesis (QuikChange Site-Directed Mutagenesis Kit, Agilent Technologies, La Jolla, CA), three additional cDNA constructs were generated each containing SNP1, SNP2, and both SNP1 and 2. The 3xFLAG-human FGFR1OP2/wit3.0β cDNAs with wild-type sequence as well as SNP1, SNP2, and SNP1/2 sequences were subcloned into the lentivirus-derived vector: pRLSinhCMV-IG. The construct (pRLSinhCMV-wit3.0-IG) has an internal ribosome entry site-green fluorescent protein gene downstream of 3xFLAG-human FGFR1OP2/wit3.0β fusion peptide. The construct was used to co-transfect 293T cells together with two packaging vectors, pCMVΔR8.2 and pVSVG. The lentiviral vector was concentrated to five ~20 μg p24/ml virus stock.

Table 2. Available Mouse ES Cell Lines Containing Disrupted FGFR1OP2/wit3.0 Allele (146526432-146547719) through the International Gene Trap Consortium

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<th>Identification status</th>
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<td>SIGTR</td>
<td>146526450-146526731</td>
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The Effect of FGFR1OP2/wit3.0β Wild-Type, SNP1, SNP2, and SNP1/2 on Gel Contraction

Oral fibroblasts and skin fibroblasts (2 × 10^6 cells/well) were treated with 1 μg of lentivirus (1 μg p24 = 5 × 10^7 infectious units in 293T cells) and polybrene (8 μg/ml) for 24 hours. The lentivirus transduction efficiency to oral fibroblasts and skin fibroblasts were evaluated for the internal ribosome entry site-driven green fluorescent protein expression. The effect of FGFR1OP2/wit3.0β overexpression carrying wild-type, SNP1, SNP2, and SNP1/2 sequences on floating collagen gel contraction were determined as described above.

The Effect of Transforming Growth Factor-β1 on the Expression of FGFR1OP2/wit3.0 and α-SMA

Oral fibroblasts and skin fibroblasts were seeded on a 6-well plate (1 × 10^5 cells/well). After 24 hours of starvation treatment by low serum culture medium (0.5% FBS), cells were incubated in Dulbecco’s modified Eagle’s medium containing transforming growth factor-β1 (TGF-β1; 0, 0.1, and 0.5 ng/ml). The steady state mRNA levels of FGFR1OP2/wit3.0 and α-SMA were accessed by Taqman-based real-time RT-PCR. The experiments were triplicated. From the Student’s t test performed against the untreated cells, statistical significance was accepted for \( P < 0.05 \).

Mouse Dorsal Skin Full-Thickness Excisional Wound Model

Under isofluorane inhalation anesthesia, two identical skin wounds were created on the right and left side equidistant (0.5 cm) from the dorsal mid point in male C57BL/6J mice (10 weeks old). Using an 8-mm dermal biopsy punch, full-thickness skin including panniculus carnosus layer was excised. The skin specimens containing the healing excisional wound were collected and examined for the steady state mRNA levels of FGFR1OP2/wit3.0.
Efficacy of Gene Transfer Methods to Mouse Skin Wound

The efficiency of gene transfer to mouse skin wound was compared for plasmid DNA and lentiviral vector carrying firefly luciferase reporter gene, which were mixed in ether type I collagen (Cell Prime, Cohesion, Palo Alto, CA) or cationized gelatin (CG) SM50 (generous gift from Dr. Tabata, Kyoto University, Japan). Cell Prime or CG mixed with β-galactosidase reporter gene plasmid DNA (1 µg/µl) served as the negative control. Four days after gene delivery, anesthetized mice were injected with 200 µl of filtered D-Luciferince firefly (15 mg/ml in PBS) and luciferase activity in skin wound area was measured by using the fluorescent single-view three-dimensional optical imaging system (Xenogen IVIS System, Caliper Life Science, Hopkinton, MA).

The Effect of FGFR1OP2/wit3.0 Treatment on Excisional Skin Wound Closure In Vivo

After the full-thickness excisional skin wound was created in C57Bl/6J mice, 80 µl of CG/3x-flag-hwit3.0-WT lentivirus, CG/3x-flag-hwit3.0-SNP1 lentivirus, or CG/Luciferase-GFP lentivirus was topically applied. The standardized photographs were collected 0, 4, and 7 days after wounding. The skin wound area was measured at each time point (n = 10 in each group). One cm² of skin tissue samples framing the circular wound were collected for histological analysis (n = 6 for each time point). Six-micrometer thick paraffin sections were made through the center of the wound. The sections were stained with Goldner’s trichrome staining or Sirius Red. Collagen fiber from the sirius red-stained sections were visualized by using confocal laser scanning microscopy. Immunohistological staining by using D1042 antibody was performed as described above.

Results

Localized Expression of FGFR1OP2/wit3.0 in the Oral Wound Margin

We generated a monospecific polyclonal antibody, D1042, against FGFR1OP2/wit3.0. Western blot analysis of NIH3T3 fibroblasts expression 3XFLAG-FGFR1OP2/wit3.0 fusion peptide indicated that both M2 antibody against 3XFLAG epitope and D1042 recognized the 41 kDa peptide. Because D1042 also recognized 32 kDa peptide, we concluded that D1042 is specific to the putative epitope of 41 kDa FGFR1OP2/wit3.0 as well as 32 kDa FGFR1OP2/wit3.0 (Figure 1B).
tion of FGFR1OP2/wit3.0 and α-SMA was evaluated by immunohistology. The oral fibroblasts appear to vary in the expression of FGFR1OP2/wit3.0 and α-SMA, which did not seem to overlap each other. The fibroblasts expressing FGFR1OP2/wit3.0 showed cuboidal shapes and localized within the connective tissue immediately below the epithelial proliferation and migration front (Figure 1C). On the contrary, α-SMA was strongly expressed by vascular smooth muscle cells. The α-SMA positive myofibroblasts were abundant in the bone socket created by tooth extraction and showed elongated shape along the direction of the ligament-like structure attached to the alveolar bone (Figure 1C).

The real time RT-PCR revealed that FGFR1OP2/wit3.0 was significantly up-regulated in the oral wound, whereas wounding of skin tissues did not modulate the expression of FGFR1OP2/wit3.0 (Figure 1D). In both wounded tissues, the steady state mRNA levels of α-SMA were significantly increased albeit at different levels.

**FGFR1OP2/wit3.0 has the Characteristics as a Cytoskeleton Molecule**

The protein structure databases, Human Protein Reference Database and COIL, predicted that FGFR1OP2/wit3.0 contains two separate coiled-coil structures flanking a center domain (Figure 2A). The Imperial College Protein Homology/Analogy Recognition Engine 3D protein structure in silico analysis revealed that α-spectrin/α-actinin and apolipoprotein A-I could predict the FGFR1OP2/wit3.0 structure at the estimated precision levels of 85 to 90%. Both α-spectrin/α-actinin and apolipoprotein A-I contain short antiparallel α-helices separated by reverse turns, resulting in the dimer and polymer formation through coiled-coil structures (Figure 2B).

The denatured gel Western blot analysis showed that NIH3T3 fibroblasts expressed FGFR1OP2/wit3.0 and FGFR1OP2/wit3.0 as single molecules; however, when 3XFLAG-FGFR1OP2/wit3.0 was overexpressed, a 50 kDa product appeared. The 50 kDa FGFR1OP2/wit3.0 positive band was about 10 kDa larger than FGFR1OP2/wit3.0 and was posttranslationally modified with SUMO-1 (data not shown). FGFR1OP2/wit3.0 in oral wound fibroblasts seemed to be extensively sumoylated. In addition, D1042 positive bands were observed at 70 to 82 kDa, 100 kDa, and 150 to 160 kDa, suggesting the formation of dimers and oligomers (Figure 2C). In the non-denature gel Western blot analysis, the D1042 antibody recognized a peptide at 50 kDa that was co-recognized with SUMO-1 antibody. There were D1042-positive 100 kDa and 160 kDa bands in oral fibroblasts but not in oral wound fibroblasts. The involvement of FGFR1OP2/wit3.0 in a
large molecular moiety of over 190 kDa was also noted in both oral fibroblasts and oral wound fibroblasts; however, the size of the large molecular moiety and D1042 signal intensity increased in oral wound fibroblasts (Figure 2D).

Oral fibroblasts and oral wound fibroblasts were subjected for immunocytochemistry. The majority of oral fibroblasts exhibited elongated cellular morphology and F-actin stress fibers that were localized in the long axis of cell periphery. FGFR1OP2/wit3.0 was primarily localized in the perinuclear cytoplasm and did not overlap with the F-actin. On the contrary, oral wound fibroblasts were more spread to form cuboidal shapes and developed a fine web of cytoskeleton fiber networks (white arrows in Figure 2E). FGFR1OP2/wit3.0 was found to localize in the wider area of cytoplasm. Both F-actin and FGFR1OP2/wit3.0 appeared to associate with the cytoskeleton fiber network; however, they do not seem to co-localize but are rather seen in a somewhat mutually exclusive pattern (Figure 2E).

siRNA FGFR1OP2/wit3.0 Knockdown Attenuated the Floating Collagen Gel Contraction

Compared with skin fibroblasts, oral fibroblasts exhibited the higher rates of gel contraction activity at 12 hours and sustained its activity until 24 hours. Oral wound fibroblasts showed the greatest gel contraction rates (repeated measure analysis of variance; *P < 0.05*) (Figure 3A).

The role of FGFR1OP2/wit3.0 in fibroblast-driven wound contraction was examined in siRNA knockdown experiments. We designed five siRNA target sequences (Table 1), out of which number 5 siRNA showed the 80 to 90% knockdown efficiency followed by number 4 siRNA, which showed 20 to 30% knockdown efficiency (Figure 3B). The treatment of number 4 and number 5 siRNAs did not affect other contractile and procontractile molecules such as α-SMA, ezrin, radixin, and moesin (Figure 3B).

Considering optimal balance factor (transfection efficiency versus cell survival) and the silencing efficiency among the three transfection reagents tested, the optimal transfection efficiency of siRNA to primary oral wound fibroblasts was established: 0.25 μl Lipofectamine 2000 with 6000 cells/well cell density, generating a transfection effect of 92% (Figure 3C).

In collagen gel contraction assay, number 5 siRNA significantly decreased gel contraction rate of oral wound fibroblasts (repeated measure analysis of variance; *P < 0.05*), which became equivalent to that of unwounded oral fibroblasts. The negative control siRNA and number 4 siRNA showed no effect on the floating collagen gel contraction rates (Figure 3D).

FGFR1OP2/wit3.0 (+/−) Mutation Reduced Fibroblastic Cell Migration

To elucidate the mechanism of reduced FGFR1OP2/wit3.0 expression, migration capability of fibroblasts were evaluated. We applied the available mouse ES cell line carrying the gene trap-mediated FGFR1OP2/wit3.0 knockout mutation. The mouse ES cell line IST10830D12 was generated under the International Gene Trap Consortium and a β-geo cassette was inserted in the first intron of an FGFR1OP2/wit3.0 allele (Figure 4A). Because the translational start site is located in the second exon, this gene trap mutation would create the knockout mutation. Fibroblastic cells carrying FGFR1OP2/wit3.0 (+/−) mutation (Figure 4B) did not affect the collagen I expression but reduced the FGFR1OP2/wit3.0 mRNA level to approximately 50% of the wild-type control (Figure 4C). The in vitro wound scratch plate assay demonstrated that cell migration of FGFR1OP2/wit3.0 (+/−) cells was significantly decreased compared with the wild-type control (Figure 4D).

The Human FGFR1OP2/wit3.0β Carrying Wild-Type Sequence as well as SNP1, SNP2, and SNP1/2 Sequences Increased Floating Collagen Gel Contraction

We generated three additional FGFR1OP2/wit3.0β cDNAs representing the nonsynonymous SNP sequences in exon 5 (Figure 5A) and subcloned to a third generation lentivirus vector containing internal ribosome entry site-green fluorescent protein (Figure 5B). The transduction efficiencies of the lentivirus to primary oral fibroblasts and primary skin fibroblasts were confirmed over 80% by Fluorescence Activated Cell Sorting (FACS) (Figure 5C). The transduction of the wild-type FGFR1OP2/wit3.0β to both oral and skin fibroblasts significantly increased the floating collagen gel contraction when compared with their untransduced control fibroblasts (repeated measure analysis of variance; *P < 0.05*) (Figure 5D). In addition, the SNP1 human FGFR1OP2/wit3.0β carrying 154E→D peptide further increased floating collagen gel contraction rates (repeated measure analysis of variance; *P < 0.01*), whereas SNP2 and SNP1/2 variants showed a significant, but a less drastic, increase (repeated measure analysis of variance; *P < 0.05*) (Figure 5E).

TGF-β1 Did Not Affect the Expression of FGFR1OP2/wit3.0

Both in oral fibroblasts and skin fibroblasts, when treated with TGF-β1, the α-SMA expression was dose-dependently increased. However, TGF-β1 treatment did not affect the FGFR1OP2/wit3.0 expression (Figure 5F).

FGFR1OP2/wit3.0β Treatment to Mouse Excisional Wound Accelerated Early Wound Closure In Vivo

A full-thickness excisional wound was created through the panniculus carnosus layer in the dorsal skin of C57Bl/6J mice (Figure 6A). Unlike oral wound, the expression level of FGFR1OP2/wit3.0 was unchanged during early healing stages of the skin (Figure 6B). To ad-
dress the effect of FGFR1OP2/wit3.0β overexpression in this model, we first tested the in vivo gene delivery systems suitable for mouse skin wound. Plasmid DNA with collagen carriers and CG carrier did not generate sufficient reporter gene delivery to the skin wound; however, the delivery of lentivirus vector combined with the CG SM50 carrier showed the promising result (one-way analysis of variance; \( P < 0.05 \)) (Figure 6C). This method was applied to deliver the FGFR1OP2/wit3.0 expression vector to mouse full-thickness excisional wound. Compared with control wounds received green fluorescent protein-expression vector, FGFR1OP2/wit3.0β-treated wounds exhibited accelerated closure (Mann-Whitney test; \( P < 0.05 \)) (Figure 6D). It was further observed that the SNP1 FGFR1OP2/wit3.0β-treated wound showed greater wound closure (Mann-Whitney test; \( P < 0.001 \)) (Figure 6D).

The histological observation revealed that FGFR1OP2/wit3.0β was mostly expressed by fibroblasts in granulation tissue juxtaposing dermis wound margins (Figure 7A). In the mock-treated group, while epithelialized, the dermis and panniculus carnosus wound edges remained patent. The wound area was primarily filled with highly cellular granulation tissue (Figure 7B). In the FGFR1OP2/wit3.0β treated wounds, dermis wound margins appeared to be approximated toward the center of the wound, resembling the oral wound healing. Consequently, the size of granulation tissue was found smaller in the FGFR1OP2/wit3.0β treated skin wound than the untreated wound (Figure 7C).

**Discussion**

FGFR1OP2/wit3.0β was initially identified as a differentially expressed cDNA in the rat oral wound. The present immunohistological study revealed that FGFR1OP2/wit3.0 was selectively expressed by a cluster of fibroblasts in the oral wound located immediately below the epithelial proliferation and migration front of wound margins that were actively approximating over the open wound (Figure 1C). Therefore, we have postulated that FGFR1OP2/wit3.0 may play a role in oral wound closure. Oral fibroblasts isolated from gingiva or palatal mucosa have been shown strong contraction characteristics. The present study confirmed this phenotype and further revealed that oral wound fibroblasts isolated from the tooth extraction wound exhibited significantly higher collagen gel contraction ability than naïve oral fibroblasts (Figure 2A). FGFR1OP2/wit3.0β overexpression in naïve oral fibroblasts increased the collagen gel contraction to the level of that of oral wound fibroblasts (Figure 5D), which was, in turn, attenuated by the siRNA knockdown (Figure 3D). These data suggest that FGFR1OP2/wit3.0β may contribute to the fibroblast-embedded gel contraction. Wound closure is a complex physiological phenomenon, in which the mechanistic regulation may be accounted for by (1) the mediation of proinflammatory cytokines and growth factors; (2) cell migration and cell traction force; and (3) collagen and extracellular matrix (ECM) reorganization. The present...
study further examined the mechanistic model of oral wound closure by using FGFR1OP2/wit3.0 as a clue.

Wound contraction has been shown to be stimulated by wound-associated growth factors such as TGF-β and platelet-derived growth factor. TGF-β-stimulated collagen gel contraction appears to be primarily mediated by transdifferentiation of myofibroblasts and up-regulation of α-SMA, whereas platelet-derived growth factor-stimulated contraction may be contributed by cytoskeleton rearrangement. These studies were conducted by using skin fibroblasts. We demonstrated that oral fibroblasts similarly responded to TGF-β and increased the expression of α-SMA (Figure 5F). However, TGF-β1 treatment did not modulate the steady state mRNA levels of FGFR1OP2/wit3.0 in both skin and oral fibroblasts, suggesting that FGFR1OP2/wit3.0-stimulated collagen gel contraction may not be explained by the mediation of TGF-β1. In the tooth-extraction-induced oral wound, myofibroblasts expressing α-SMA were found in the ligament-like structure connecting alveolar bone of the extraction socket (Figure 1C). After tooth extraction, periodontal ligament often remains in the socket, and oral myofibroblasts may be derived from periodontal ligament fibroblasts. FGFR1OP2/wit3.0-expressing oral wound fibroblasts did not appear to express α-SMA, and thus may not be considered as “myofibroblasts.” The origin of FGFR1OP2/wit3.0-expressing oral wound fibroblasts has not been determined in the scope of this study.

TGF-β1 not only up-regulates the α-SMA expression but also significantly increases collagen synthesis causing fibrotic disorders. Inhibition of matrix metalloproteinases by ilomastat has been shown to significantly decrease collagen production, resulting in the reduction in fibroblast-embedded collagen gel contraction. The reorganizing ECM has been shown to affect the wound contraction. To further elucidate the functional role of FGFR1OP2/wit3.0, we have developed fibroblastic cells that were derived from mouse ES cells carrying
FGFR1OP2/wit3.0 (+/-) mutation. ES cell-derived fibroblastic cells with wild-type and FGFR1OP2/wit3.0 (+/-) mutation expressed collagen I at the equivalent level (Figure 4C). Our data, although limited, did not immediately support the involvement of ECM reorganization in FGFR1OP2/wit3.0-stimulated collagen gel contraction.

In the present study, fibroblastic cells derived from ES cells were further tested for the scratch plate assay, in which FGFR1OP2/wit3.0 (+/-) mutation exhibited the unexpectedly robust impairment in cell migration (Figure 4D). Fibroblast migration is an essential process during wound healing, and may, in part, result in the transmission of intracellular forces to the ECM. FGFR1OP2/wit3.0 appears to be involved in the mechanism of cell migration, which may influence the collagen gel contraction and wound closure.

The up-regulation of α-SMA dose-dependently correlated with the magnitude of cell traction force, however, the α-SMA knockout mutation resulted in the increased cell motility. Myosin IIB null mutation did not affect the generation of traction force but caused the defect in directional movement resulting in the impairment of cell migration. It has been well established that cytoskeleton molecules such as α-SMA and myosin primarily regulate cell migration and traction force generation. FGF1OP2/wit3.0 isoforms are small peptides possessing N-terminal and C-terminal coiled-coil domains, which may facilitate the self-dimerization and self-oligomerization through anti-parallel coiled-coil domains similar to α-actinin and apolipoprotein A-I. The dimer formation of FGFR1OP2-FGFR1 fusion peptide in the patient with the cytogenetic abnormality (12;8)(p11;p11p22) further support that FGFR1OP2/wit3.0 peptide can be dimerized and oligomerized through the coiled-coil domains. The non-denature gel Western blot suggested the incorpo-
The topical application of FGFR1OP2/wit3.0 in larger molecular moieties in oral wound fibroblasts (Figure 2D). Confocal laser scanning microscopy revealed the association of FGFR1OP2/wit3.0 with thin web of cytoskeleton networks of oral wound fibroblasts. F-actin was also found in the cytoskeleton network; however, F-actin and FGFR1OP2/wit3.0 appeared to be localized in a mutually exclusive fashion (Figure 2E). Taken together, FGFR1OP2/wit3.0 possesses characteristic features as cytoskeleton molecule, which may be suitable to participate in the regulation of cell migration and potentially wound closure.

The relationship between cell motility, migration, and ECM remodeling relevant to wound healing should occur in the three-dimensional ECM environment. Grinnell has proposed that the strained collagen gel may provide a more suitable experimental model to investigate the role of fibroblasts in wound healing. In the present study, we further evaluated the role of FGFR1OP2/wit3.0 in the wound healing in vivo by using the mouse dorsal skin wound model. The full-thickness excisional wound created in dorsal skin of C57Bl/6J mice underwent slow onset of wound closure during the first week followed by the progressive wound contraction during the second week. The topical application of FGFR1OP2/wit3.0β-expression vector significantly accelerated the wound closure as compared with controls (Figure 6D). The FGFR1OP2/wit3.0β-treated wound exhibited the unique approximation of dermis wound margins toward the center of excisional wound, resulting in the limited area of granulation tissue (Figure 7C). Because the blunted panniculus carnosus remained patent, we postulate that the unique approximation of dermis wound margins was likely to be contributed by FGFR1OP2/wit3.0β-induced fibroblast activities within the granulation tissue and/or dermis wound margins.

The primary goals of skin wound management are to achieve rapid wound closure and to minimize the scar formation. This study demonstrated that a previously unknown molecule, FGFR1OP2/wit3.0, possessed the characteristics consistent with a cytoskeleton molecule and played a role in the wound closure. Our data further suggest that the single molecule treatment of FGFR1OP2/wit3.0 seems to be sufficient for inducing the accelerated skin wound closure, similar to the oral wound healing phenotype. As such FGFR1OP2/wit3.0 therapy may offer a novel approach for wound management, in which the critical initial wound closure may be better achieved.
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