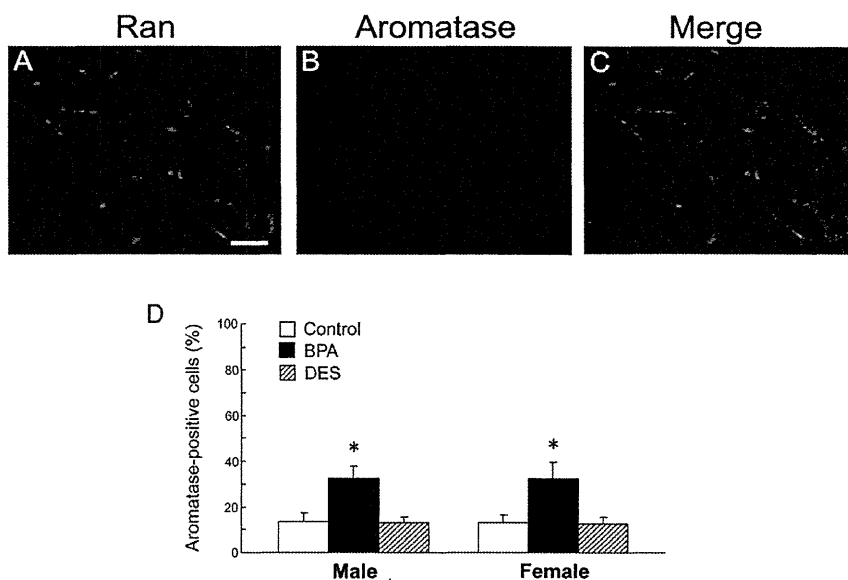


FIG. 4. BPA-specific increases of aromatase-expressing cells in primary cultured UGM. A–C) Fluorescence signals were detected for the CYP19A1 (aromatase) protein in primary cultured UGM. The nuclei were identified by Ran staining. Bar = 100 μ m, magnification \times 400. D) The number of aromatase-positive cells was counted in primary cultured UGM of the untreated control (open bar), BPA-treated UGS (closed bar), and DES-treated UGS (slashed bar), and the percentage of aromatase-positive cells was calculated from at least 10 areas. * P < 0.01 vs. control.



observed in the cytoplasm of cultured UGM. The rate of positivity (i.e., the percentage of cells that expressed CYP19A1 [aromatase] protein), was approximately 10% in the untreated and the DES-treated groups, whereas it was as high as approximately 30% in the BPA-treated group (Fig. 4D). No difference in the rate of positivity of CYP19A1 (aromatase) was found when comparing the untreated male UGM to that of the female.

Restricted BPA-Specific Up-Regulation of *Esrrg* mRNA in UGE and UGM

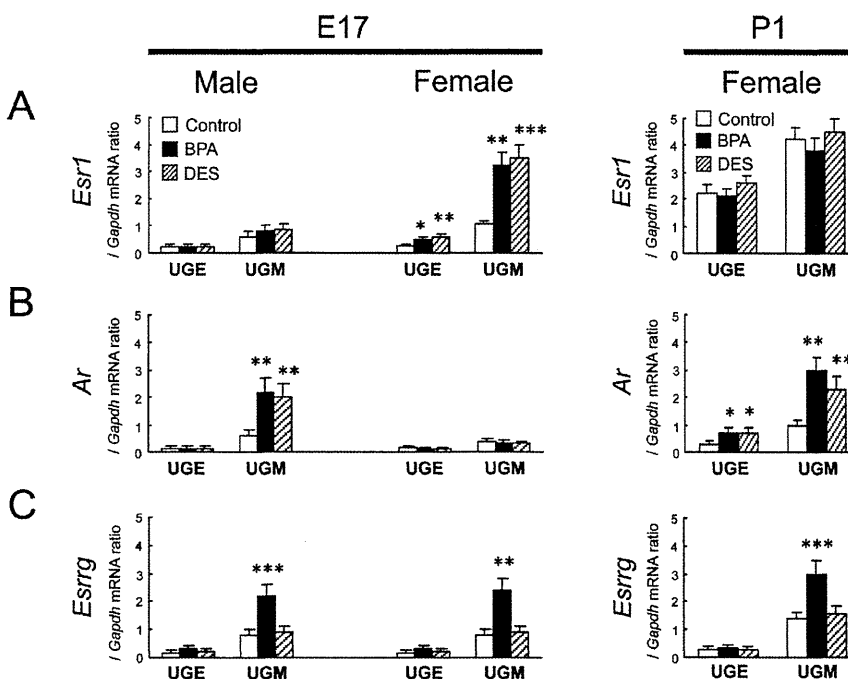
In E17 female UGM, the mRNA expression of *Esr1* was up-regulated by both BPA and DES treatment (Fig. 5A). At E17, however, the mRNA expression of *Ar* was up-regulated by both BPA and DES treatment in the male UGS (Fig. 5B). At

P1, mRNA expression of *Ar* was up-regulated by both BPA and DES treatment in the female UGS (Fig. 5B). In both the male and female, the up-regulation of *Esrrg* mRNA was observed at E17 and restricted in UGM, but not in UGE, of the BPA-treated group (Fig. 5C). In both the male and female UGE, the expression of *Esrrg* mRNA was quite low and not up-regulated, even in the BPA-treated group. At E17, no difference in mRNA expression levels was found when comparing the untreated male UGS with that of the female.

BPA-Specific Increases of *ESRRG*-Expressing Cells in Primary Cultured UGM

In both the male and female, E17 UGM was primary cultured in vitro. Representative pictures of *ESRRG*-positive

FIG. 5. Restricted BPA-specific up-regulation of *Esrrg* mRNA in UGE and UGM. The relative mRNA expressions of *Esr1* (A), *Ar* (B), and *Esrrg* (C) were determined in UGE and UGM of the untreated control (open bar), BPA-treated UGS (closed bar), and DES-treated UGS (slashed bar) at E17 and P1. * P < 0.05, ** P < 0.01, *** P < 0.001 vs. control.



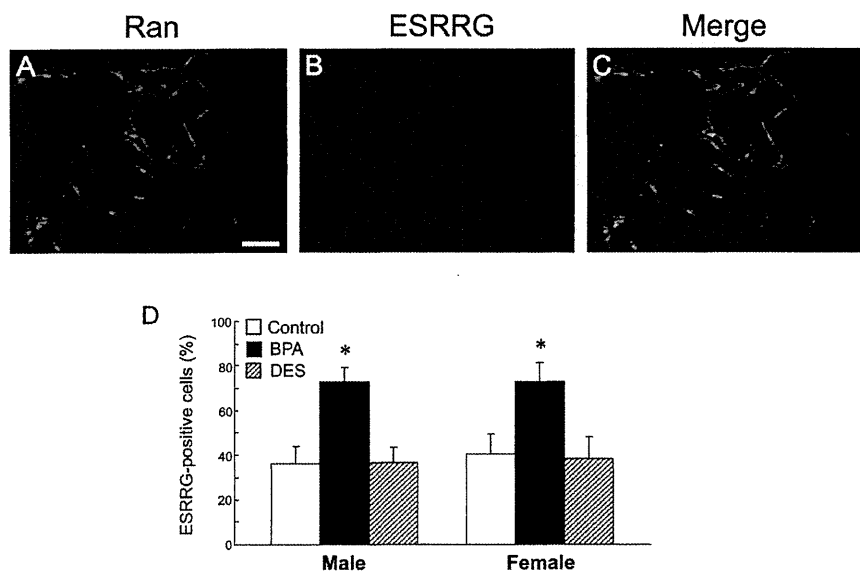


FIG. 6. BPA-specific increases of ESRRG-expressing cells in primary cultured UGM. A–C) Fluorescence signals were detected for the ESRRG protein in primary cultured UGM. The nuclei were identified by Ran staining. Bar = 100 μ m, magnification $\times 400$. D) The number of ESRRG-positive cells was counted in primary cultured UGM of the untreated control (open bar), BPA-treated UGS (closed bar), and DES-treated UGS (slashed bar), and the percentage of ESRRG-positive cells was calculated from at least 10 areas. * $P < 0.01$ vs. control.

cells are shown in Figure 6, A–C. The ESRRG-positive staining was observed in both the nucleus and the cytoplasm of cultured UGM. The number of ESRRG-positive UGM was significantly increased only in the BPA-treated group and showed a 2.2-fold increase in males and a 1.6-fold increase in females (Fig. 6D). No difference was found in the rate of positivity of ESRRG when comparing the untreated male UGM with that of the female.

BPA-Specific Up-Regulation of *Esrrg* and Steroidogenic Enzyme mRNA in Sex Hormone-Related Organs

To investigate the BPA-specific up-regulation of in situ steroidogenesis in other organs, we first examined the changes in *Esrrg* mRNA expression in sex hormone-related organs, such as the cerebellum, heart, kidney, ovary, and testis. At P1, the mRNA expression of *Esrrg* in the cerebellum, heart, kidney, and ovary, but not in the testis, was up-regulated by both BPA and DES treatment (Fig. 7A). However, no significant difference in *Ar* mRNA expression was observed in all organs examined (Fig. 7B). In the untreated group, the mRNA expression of *Esrrg* was not detected in the testis at E17 and P1 (Fig. 7C). The up-regulation of *Esrrg* mRNA was observed at E17 and restricted to the cerebellum, heart, kidney, and ovary (Fig. 7C). The BPA-specific up-regulation of *Cyp19a1*, *Cyp11a1*, and *Nr5a1* mRNA was observed only at P1 in the cerebellum, heart, kidney, and ovary, but not in the testis (Fig. 8).

DISCUSSION

Concern about the effects of EDCs such as BPA on human health has been increasing [24]. Although the majority of EDCs have the potential to alter functioning of the reproductive and endocrine system, the actual mechanism responsible for such alterations has not been identified thoroughly. BPA is of concern because its chemical structure resembles that of DES. Several studies have reported that BPA can mimic estrogen action, such as induction of vaginal cornification, uterine vascular permeability, growth and differentiation of the mammary gland, and synaptic plasticity in the hippocampus [25–28]. In the prostate, alterations in normal development can

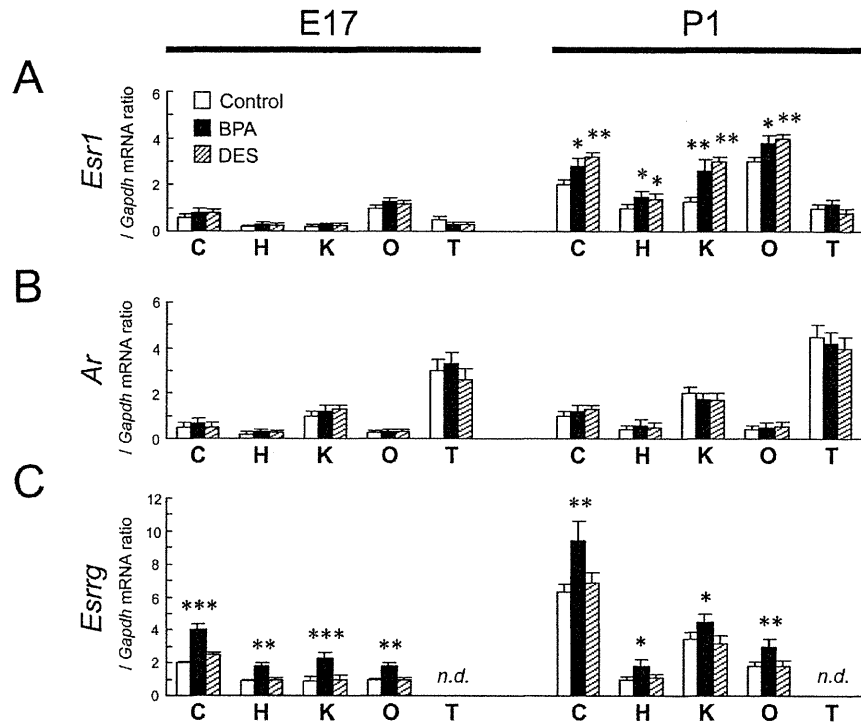
produce permanent changes that persist throughout adulthood and may increase the risk of disease in later life [9]. Thus, our objective was to investigate the biological effects of low-dose BPA on the initial development of primary ducts in the fetal prostate.

During prostatic development, alteration of sex steroid hormone synthesis may be responsible for prostatic anomalies associated with fetal exposure to EDCs. In the present study, fetal exposure to low-dose BPA increased E_2 levels in P1 UGS of both the male and female, whereas DES-induced changes were not detected. This alteration was also correlated with increased activity of CYP19A1 (aromatase) in UGS at P1, suggesting the unique action of BPA for in situ steroidogenesis in UGS. The BPA-specific increase of E_2 levels in UGS at P1 was correlated with the following: mRNA up-regulation of steroidogenic enzymes, such as *Cyp19a1* and *Cyp11a1*, and an increased number of aromatase-expressing UGM. The enzyme CYP19A1 (aromatase) is responsible for in situ E_2 production and the crucial testosterone/ E_2 balance necessary for normal embryonic and fetal development, even in males. The data presented here shows that the up-regulation of *Cyp19a1* mRNA in BPA-treated UGM was comparable to changes in both in situ E_2 production and CYP19A1 (aromatase) activity.

In the present study, we demonstrated that the BPA-specific increase in steroidogenic enzyme mRNA and aromatase-expressing cell number were observed in both the male and female UGM. During embryonic development, the mesenchymal component is involved in the induction and organogenesis of various organs, including the prostate, mammary gland, lung, kidney, and pancreas. It has been well established that subpopulations of the mesenchymal component are a source of potent molecules that regulate epithelial growth and differentiation [29]. In the prostate, androgen-responsive signals derived from UGM permissively and instructively induce UGE to form primary ducts of the prostate [30].

Comparison between the neonatal male and female UGS shows a similarity in the condensed mesenchyme of the ventral areas—that is, the ventral prostate mesenchyme (VPM) in the male and the ventral mesenchymal pad (VMP) in the female [31]. In the male, a defined VPM is specifically associated with ductal branching morphogenesis and cytodifferentiation of the ventral prostate. Females do not usually form a prostate. In a

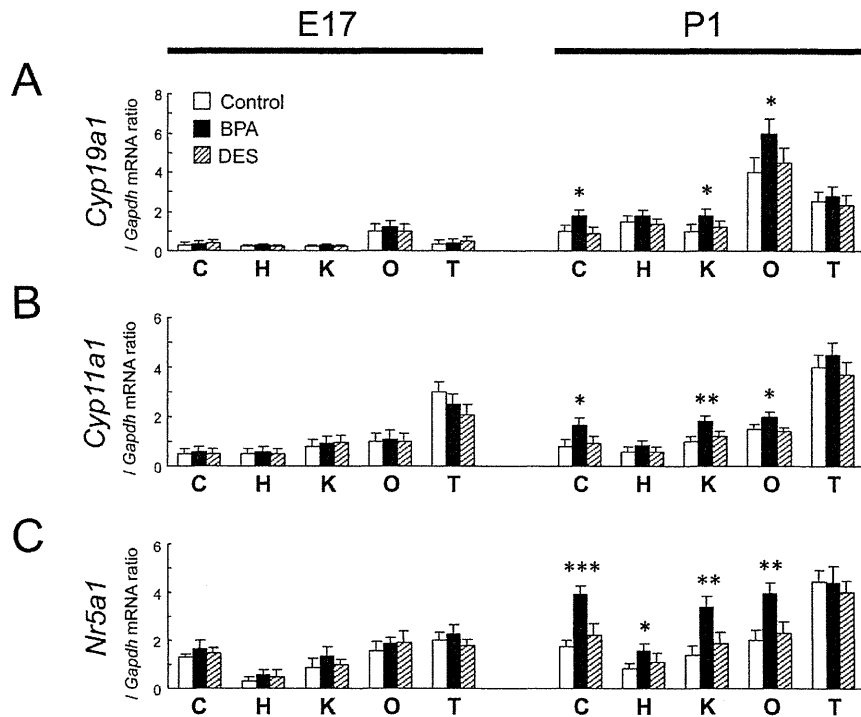
FIG. 7. BPA-specific up-regulation of *Esr1* mRNA in sex steroid hormone-related organs. The relative mRNA expressions of *Esr1* (A), *Ar* (B), and *Esr1* (C) were determined in sex steroid hormone-related organs of the untreated control (open bar), BPA-treated UGS (closed bar), and DES-treated UGS (slashed bar) at E17 and P1. C, cerebellum; H, heart; K, kidney; O, ovary; T, testis; *n.d.*, not detected. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ vs. control.



tissue recombination model, the female VMP induces prostate development in response to androgens [32], suggesting that cells within the female VMP have prostatic-inductive activity. Moreover, an earlier tissue recombination study showed that the ability of the female UGS to respond to androgens in forming prostate was gradually lost between P1 and P5 [33]. These results suggest strongly that androgen-responsive regulatory

molecules are expressed constitutively even in the female VMP. Although the female VMP forms in the absence of androgens, androgen receptor (AR) expression was observed in the neonatal female VMP in a pattern similar to that observed in the male VPM [34]. Therefore, the BPA-specific increase in E_2 levels might interact with the intracellular AR signaling in both the male VPM and the female VMP. However, to our knowledge,

FIG. 8. BPA-specific up-regulation of steroidogenic enzyme and sex-determining gene mRNA in sex steroid hormone-related organs. The relative mRNA expressions of *Cyp19a1* (A), *Cyp11a1* (B), and *Nr5a1* (C) were determined in sex steroid hormone-related organs of the untreated control (open bar), BPA-treated UGS (closed bar), and DES-treated UGS (slashed bar) at E17 and P1. C, cerebellum; H, heart; K, kidney; O, ovary; T, testis. * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$ vs. control.



the morphological changes in neonatal female UGS have not yet been investigated.

Our results suggest that BPA has a stimulatory effect on *in situ* steroidogenesis in P1 UGS of both the male and female at low-dose exposure levels. Recently, ESRRG has been reported to bind strongly with BPA [35]. Susens et al. [36] have reported that expression of ESRRG in the mouse is organ-specific: ESRRG is expressed in the brain, heart, kidney, and skeletal muscle but not in the lung, spleen, and testis. In the present study, the up-regulation of *Cyp19a1* and *Cyp11a1* mRNA by BPA treatment was detected only in organs expressing *Esrrg* mRNA. These data suggest that the possibility of a stimulatory effect on *in situ* steroidogenesis by fetal exposure to low-dose BPA may be a concern not only in UGS but also in organs expressing ESRRG, such as the brain, heart, kidney, and ovary. It is important to note that Takeda et al. [23] have recently reported that ESRRG was detected in the human testis, suggesting that the distribution of ESRRG differs slightly between mice and humans.

In the present study, the BPA-specific up-regulation of steroidogenic enzyme mRNA in UGS, cerebellum, heart, kidney, and ovary was observed only during the neonatal period (i.e., P0 and P1) and not during the prenatal period (i.e., E17 and E18). During pregnancy in rodents, large amounts of estrogens produced in the maternal ovaries are continuously delivered to the fetus through the placenta. After birth, however, the fetus may be released from the maternal, high-estrogen environment. Thus, one possibility is that the maternal, high-estrogen environment in pregnancy may protect the fetus from the effect of BPA on *in situ* steroidogenesis during the prenatal period. However, we did not investigate the effects of neonatal BPA treatment on *in situ* steroidogenesis.

The EDC-induced alterations of the *in situ* estrogen environment depend on each compound. In addition to atrazine and dioxin, the organotin compound tributyltin also increases E_2 production in human placental choriocarcinoma cells [37]. Tributyltin has been demonstrated to induce the superimposition of male sex organs, such as a penis and/or a vas deferens, over female sex organs, which is a phenomenon known as imposex [38]. These studies suggest strongly that EDCs might affect fetal development not only by mimicking the actions of sex steroid hormones but also by alteration of *in situ* steroidogenesis.

In the prostate, AR expressed in mesenchyme is required for directing growth and branching morphogenesis of epithelia, presumably by induction of growth factors [39]. In the present study, fetal exposure to BPA or DES increased *Ar* mRNA expression in E17 UGM of the male, whereas *Esrl* mRNA expression was up-regulated in E17 UGM of the female. Recently, Richter et al. [40] have reported that *in vitro* BPA treatment stimulates *Ar* and *Esrl* mRNA expression in mesenchymal cells isolated from fetal mouse prostate. Thus, our results support the idea that BPA-induced cell proliferation of the primary prostatic ducts may be caused by inducing *Ar* mRNA expression in the male UGM. In contrast, the induction of *Esrl* mRNA expression by BPA or DES may create a positive-feedback loop in the female UGM. Further investigation and morphological analysis will be necessary to confirm the effects of up-regulated ESR1 in the female UGS.

In conclusion, we have shown the unique action of BPA in the mouse UGS. Specifically, we have demonstrated that the increases in E_2 levels and CYP19A1 (aromatase) activity were observed in the BPA-treated UGS but not in the DES-treated UGS. Ricke et al. [41] have recently reported that stromal hormone imbalance, a potential source of local E_2 production, may be responsible for prostatic disease, such as benign

prostatic hyperplasia and prostate cancer. The data in the present study give rise to the concept that the development and differentiation of UGS in mouse fetuses is very sensitive to fetal exposure to low-dose BPA via the mother. Further investigation of various aspects of BPA-specific action is necessary to fully understand the role of BPA as an EDC.

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CITED2 is activated in ulcerative colitis and induces p53-dependent apoptosis in response to butyric acid

Tsutomu Yoshida · Tsukasa Sekine ·
Ken-ichi Aisaki · Tetuo Mikami ·
Jun Kanno · Isao Okayasu

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Abstract

Background In ulcerative colitis (UC), *Fusobacterium varium* is significantly detected in patients' mucosa, and butyric acid (BA), abundantly produced by the bacterium, activates the p53 system and induces epithelial apoptosis, as we previously reported. However, factors active in the link between BA and p53 have yet to be clarified. Here, we identified a gene activated by BA specifically in UC-associated cancer cell lines and ascertained the mechanism of its activation of p53.

Methods cDNA microarray analysis based on the Percellome (per cell normalization) method was performed on BA-stimulated UC-associated cancers and sporadic colorectal cancer cell lines under conditions mimicking colonic epithelium UC. For validation of microarray results, molecular, biochemical, and histopathological analyses were performed.

Results We found the CBP/p300-interacting transactivator with glutamic acid/asparagine-rich carboxy-terminal domain 2 (CITED2) to be specifically upregulated in UC-associated cancer cell lines by BA treatment, at both mRNA and protein expression levels. CITED2 could be shown to induce p53 acetylation and p53-dependent apoptosis, accompanied by binding of CBP/p300. BA-dependent apoptosis was suppressed by an inhibitor of monocarboxylate transporter-1 and an siRNA for p53. In inflammatory foci of UC, histologically evident inflammatory activity and CITED2 expression were significantly correlated.

Conclusions CITED2 was identified as UC-associated protein by cDNA microarray based on the Percellome method under UC-mimicking conditions in vitro. CITED2 activation may induce mucosal apoptosis and erosion by activating p53 and thus play a critical role in linking enteric bacteria with mucosal inflammation in UC.

Keywords Ulcerative colitis · Inflammation · Apoptosis · CITED2 · Butyric acid · p53 · CBP/p300

The GeneChip data have been deposited in the Percellome project site (<http://www.nihs.go.jp/tox/TtgPublication.htm>) and are accessible with no restriction.

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T. Yoshida (✉) · T. Sekine · T. Mikami · I. Okayasu
Department of Pathology, Kitasato University School
of Medicine, 1-15-1 Kitasato, Minami-ku, Sagamihara,
Kanagawa 252-0374, Japan
e-mail: tyoshida@med.kitasato-u.ac.jp

K. Aisaki · J. Kanno
Division of Cellular and Molecular Toxicology,
National Institute of Health Sciences,
1-18-1 Kamiyoga, Setagaya-ku, Tokyo 158-8501, Japan

Abbreviations

BA	Butyric acid
CITED2	CBP/p300-interacting transactivator with glutamic acid/asparagine-rich carboxy-terminal domain 2
4CHC	4-Hydroxycinnamate
IL-10	Interleukin-10
MCT1	Monocarboxylate transporter-1
q-RT-PCR	Quantitative reverse transcription PCR
sCRC	Sporadic colorectal cancer
UC	Ulcerative colitis
UCCA	Ulcerative colitis-associated cancer

Introduction

Ulcerative colitis (UC) is an intractable inflammatory bowel disease whose etiology is not known in detail. With long-standing UC, adenocarcinomas tend to occur in the colon, which might be due to repeated episodes of active chronic inflammation and remission [1, 2]. In a murine model system, dextran sulfate sodium (DSS) without mutagenicity induces UC-like colitis [3], and long-term repeated treatment with DSS can induce colitic cancers [4, 5]. Therefore, UC can be considered as a model disease for carcinogenesis on a background of chronic inflammation, and elucidation of the etiology of UC is important for treatment and prevention purposes.

The bacterial flora seem to constitute an important factor for triggering inflammation in UC, in accordance with previous reports concerning spontaneous development of UC-like colitis in IL-10 knockout mice under conventional breeding but not germ-free conditions [6–8]. Previously, we detected *Fusobacterium varium* (*F. varium*) by direct culture of mucosal biopsy specimens of UC patients and found intraepithelial invasion of *F. varium* to result in production of inflammatory cytokines in colonic epithelial cell lines in vitro [9, 10]. *F. varium* produces a short-chain fatty acid, butyric acid (BA), and direct luminal injection of BA into the rectum was found to induce apoptosis and UC-like lesions in mice [11]. In addition, BA activates p53-dependent apoptosis and DNA repair in vitro, and activation of p53 by its phosphorylation has been observed in situ in inflammatory foci of UC patients [12]. Clinically, we have shown significant improvement of UC by antibiotic combination therapy targeting *F. varium* [13, 14]. Recently, this was confirmed by a double-blind, placebo-controlled, multicenter trial [15]. However, the molecular mechanisms of BA-mediated p53 activation are unknown and largely remain to be elucidated.

In order to clarify this issue, we compared the genetic profiles of UC-associated carcinoma (UCCA) and sporadic colorectal adenocarcinoma (sCRC) cell lines undergoing BA treatment. Both series of cell lines were established in our laboratory [16]. The former include both p53 wild and mutant cells, whereas the latter are p53 wild. BA upregulates early response genes especially in UCCA cell lines, acting upstream of p53. Using whole genome cDNA microarray chips and the “per cell” normalization method (Percellome method) [17], we here identified genes specifically upregulated in UCCA lines with BA treatment. By detailed validation and the further examination, we focused on one upregulated gene, encoding the CBP/p300-interacting transactivator with glutamic acid/asparagine-rich carboxy-terminal domain 2 (CITED2) [18]. The protein itself activates p53 and its downstream molecules and accelerates apoptosis, and histological analysis revealed

unique expression in the colonic mucosa of UC patients. Thus, we consider that CITED2 is an important and specific mediator of inflammation in UC, and might thus serve as a target of therapy.

Materials and methods

Cell lines and clinical cases

The human UCCA cell lines UCCA21, UCCA22, UCCA23, UCCA24, UCCA25 (p53 wild), and UCCA3 (p53 mutant), and the sCRC cell lines KE24, KE43w, and KE43p (p53 wild) were established in our laboratory [16], and cultured in Dulbecco’s Modified Eagle Medium (GIBCO, Carlsbad, CA, USA) supplemented with 10% fetal bovine serum containing penicillin and streptomycin (GIBCO) at 37°C in 5% CO₂.

Microarray and data analysis

GeneChip analysis was conducted according to the Percellome method [17, 19] described in the Electronic Supplementary Material in detail. The GeneChip data have been deposited in the Percellome project site (<http://www.nih.gov/tox/TtgPublication.htm>) and are accessible with no restriction. Differentially expressed genes were extracted by Fx algorithm [20], described in the Supplementary Methods online.

q-RT-PCR and western blotting

Total RNA was isolated from cells with an RNeasy kit (QIAGEN, Hilden, Germany), and 2-μg aliquots were used to synthesize cDNAs with a High Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Foster City, CA, USA). Pre-amplified DNAs generated with a TaqMan PreAmp Master Mix Kit (Applied Biosystems) were used as templates for quantitative PCR with TaqMan Gene Expression Master Mix (Applied Biosystems) and FAM-dye-labelled TaqMan probe and primer sets for β-actin, CITED2, p53, CDKN1A (p21), Bax, MCT1, p53R2 (Applied Biosystems) using the 7500 real-time PCR system (Applied Biosystems). Quantitative PCR was performed as multiplex PCR with a VIC-dye-labelled β-actin probe and the fold ratio against VIC-labeled β-actin was calculated. All the quantitative RT-PCR (q-RT-PCR) studies were performed in triplex, and representative results for three independent experiments are shown.

Aliquots of 2×10^6 UCCA24 or KE43p cells after various treatments were lysed with lysis buffer (20 mM Tris-HCl, pH 7.6, 170 mM NaCl, 1 mM EDTA, 0.5% NP-40, 1 mM dithiothreitol). After separation by 12% SDS-

polyacrylamide gel electrophoresis and protein transfer to polyvinyl membranes, blocking with 5% bovine serum albumin was performed. Antibodies for CITED2 (Novous Biologicals, Inc Littleton, CO, USA), p53 (Novocastra Laboratories, Newcastle upon Tyne, UK), acetylated-p53 (Bio Academia, Osaka, Japan), p21^{Cip1} (Santa Cruz Biotechnology, Santa Cruz, CA, USA), Bax (Transduction Laboratories, Lexington, KY, USA), and CBP/p300 (Santa Cruz) were used as first antibodies for western blotting. HRP-conjugated anti-mouse or HRP-conjugated anti-rabbit antibodies were then employed as secondary antibodies with ECL Western Blotting Detection Reagents (GE Healthcare, Buckinghamshire, UK) for visualization. Densitometry was performed by using ImageJ software (National Institutes of Health, USA)

Transfection and RNAi (siRNA) assays

Human CITED2 cDNA was obtained from Open Biosystems (Thermo Fisher Scientific, Huntsville, AL, USA, clone: LIFESEQ8724402) and subcloned into pcDNA-DEST40 (Invitrogen). Human CBP/p300 cDNA was also subcloned into the pcDNA-DEST40 vector. Transfection into UCCA24 and KE43p was achieved with Lipofectamin2000 (Invitrogen). siRNAs for CDKN1A, CITED2, HOXA1, SAT, TUBA3, and p53 (Ambion, Austin, TX, USA), and for non-sense control (Ambion) were transfected with the siPORT NeoFX Transfection Agent (Ambion).

Butyric acid treatment, detection of apoptosis, and MCT1 inhibition assays

Cultured cells were exposed to 2.5 mM BA as previously described [12]. Apoptosis was detected by the TUNEL method visualized by fluorescein isothiocyanate (FITC)-labeled digoxigenin according to the manufacturer's protocol with slight modifications using an ApopTag Plus Peroxidase In Situ Apoptosis Detection Kit (Intergen, Purchase, NY, USA). Apoptotic cells were counted and the percentages versus more than 100 DAPI (4',6-diamino-2-phenylindole) positive cells were calculated. 0.5 mM 4-hydroxycinnamate (4CHC, Sigma, Saint Louis, MO, USA) was added to the culture medium for MCT1 inhibition assays.

Immunoprecipitation

Aliquots of 2×10^7 UCCA24 and KE43p cells were lysed in 1 ml of TNE buffer (10 mM Tris-HCl, pH 7.8, 1% nonidet-P 40, 0.15 M NaCl, 1 mM EDTA). After pre-precipitation with protein G-Sepharose for 1 h at 4°C, lysates were incubated with 1 µg of anti-CITED2 antibody or anti-CBP/p300 for 1 h at 4°C followed by incubation with protein G-Sepharose for 1 h at 4°C. After

centrifugation at 15,000 rpm for 5 min, the precipitants were washed with TNE buffer 5 times and denatured at 95°C. After SDS-polyacrylamide gel electrophoresis, they were subjected to western blotting with anti-CBP/p300 or anti-CITED2 antibodies.

Colorectal specimens and immunohistochemistry

A total of 102 surgically resected colorectal specimens from active and inactive UC patients and 20 from diverticulitis patients were obtained from the files of Kitasato University East Hospital and Kitasato University Hospital, and the histological activity of UC was determined according to the Matts' classification [21]: score 1, normal appearance; score 2, some infiltration of neutrophils in the mucosa; score 3, much cellular infiltration or presence of cryptitis; score 4, presence of crypt abscess and cellular infiltration; and score 5, presence of ulceration, erosion or necrosis, and cellular infiltration. Diverticulitis activity in diverticulosis was evaluated as none to mild and severe. Specimens were fixed with 10% formalin and embedded in paraffin. Semi-serial 4-µm-thick sections were cut for hematoxylin–eosin (H&E) staining and CITED2 immunohistochemistry with monoclonal anti-CITED2 antibodies (JA22, 1/1000, Novus Biologicals, Inc., Littleton, CO, USA) and an EnVision + kit (DakoCytomation, Glostrup, Denmark). After blocking endogenous peroxidase with 0.3% H₂O₂ in methanol, a 15-min microwave pretreatment in citrate buffer (pH 6.0, 0.01 mol/l) was performed for antigen retrieval, and after incubation with anti-CITED2 antibodies for 1 h at room temperature. 3,3'-Diaminobenzidine was applied as the final chromogen and faint nuclear counterstaining was finally achieved with methyl green. Scoring of immunoreactivity was performed as reported by Sinicrope et al. [22], and statistically analyzed by the Spearman's rank correlation test and Mann–Whitney's *U* test.

Results

Butyric acid upregulates the transcriptional expression of CITED2 specifically in UCCA lines

To identify genes upregulated by BA treatment specifically in UCCA lines, three UCCA and three sCRC lines were treated with 2.5 mM of BA (pH 7.4 in the culture medium). We earlier reported that the concentration of BA in the wells of Vero cell culture with *F. varium* was 6.42 mM (pH 6.90) [11], and 2.5 mM induced p53-mediated DNA repair [12]. The Percellome method enabled us to compare the mRNA expression profiles among the different cell lines per cell based on DNA content of each biosample. We applied the original algorithm (Fx) [20] and upregulated and

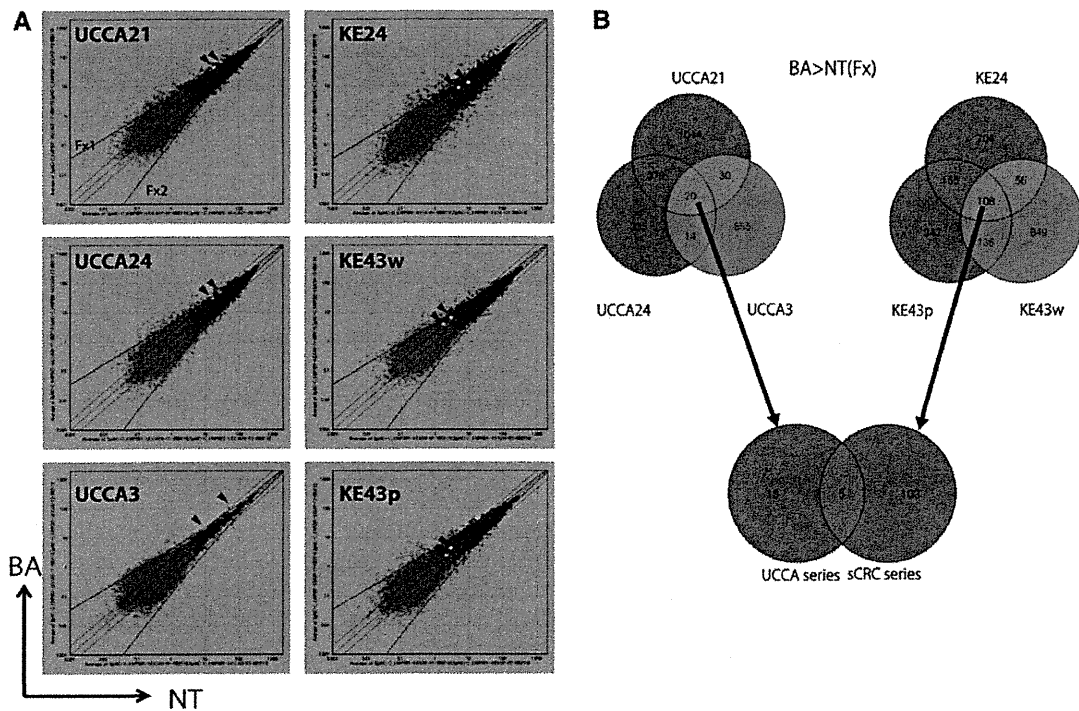


Fig. 1 a Genome-wide expression analysis after BA treatment. The analysis was performed by using the Affymetrix GeneChip Human Genome U133 plus 2.0 Array. Scatter plots are shown for BA treatment (BA) for 24 h and non-treatment (NT). The open circles show expression levels for each probe set. The colors indicate probabilities provided by the Affymetrix GeneChip Operation System: red means good and green means poor. The yellow circles show CITED2 probe sets for areas highlighted by arrowheads. The black lines show twofold, onefold, and 0.5-fold, respectively, and the blue lines indicate the empirical threshold levels (Fx1 and Fx2).

The spots above the Fx1 and below the Fx2 line were evaluated as upregulated and downregulated, respectively ($p < 0.02$). b Venn diagrams of the upregulated probe sets in butyric acid (BA) treatment compared with non-treated (NT) controls in UCCA lines (UCCA21, UCCA24, and UCCA3) and sCRC lines (KE24, KE43p, and KE43w). Numbers of upregulated probe sets are indicated. Among the 20 and 108 probe sets upregulated in the UCCA and sCRC series, respectively, 5 were commonly upregulated in both. Fx Fx algorithm for the empirical threshold calculation

downregulated genes were listed at $p < 0.02$ (Fig. 1a). As compared with non-treated controls, 20 probe sets' genes were commonly upregulated in the UCCA lines, and 108 in the sCRC lines. Five were upregulated in both UCCA and sCRC lines, and 15 and 103 only in UCCA and sCRC lines, respectively (Fig. 1b). Of the 13 known genes contained in the 15 probe sets specifically upregulated only in UCCA lines, CITED2 was upregulated with 2 independent probe sets in the same GeneChip (see Table 1 for the gene list). As we previously found that the *F. varium*-BA system induces p53-dependent cellular responses [11, 12], we focused on whether p53-signaling might be influenced. As CITED2 was identified as a CBP/p300 binding protein which regulates p53 by acetylation [18, 23, 24], we considered that CITED2 might be a candidate and subjected it to further analyses.

CITED2 acts upstream of p53

q-RT-PCR showed increased upregulation of CITED2 in UCCA cell lines as compared with sCRC cell lines on BA

Table 1 Genes upregulated by BA treatment only in the UCCA series

Probe set ID	Gene symbol	GenBank ID
237156_at	–	–
204567_s_at	ABCG1	NM_004915
210431_at	ALPPL2	J04948
200982_s_at	ANXA6	NM_001155
207980_s_at	CITED2	NM_006079
209357_at	CITED2	NM_006079
202806_at	DBN1	NM_004395
215506_s_at	DIRAS3	AK021882
211538_s_at	HSPA2	U56725
203752_s_at	JUND	NM_005354
208581_x_at	MT1X	NM_005952
201599_at	OAT	NM_000274
201364_s_at	OAZ2	AF242521
35666_at	SEMA3F	U38276
204141_at	TUBB2	NM_001069

treatment, validating the microarray results (Fig. 2a). p53 downstream molecules, p21^{Cip1} and Bax, were upregulated at the mRNA level, although p53 mRNA expression was not changed (Fig. 2b). The expression of monocarboxylate transporter-1 (MCT1), a transporter of BA, was also upregulated. When an siRNA for CITED2 was introduced into UCCA24 with or without BA treatment, it significantly suppressed CITED2 mRNA expression. Although p53 mRNA expression was upregulated by knockdown of CITED2 without BA treatment, this was abrogated by BA treatment with CITED2 knockdown, indicating an upstream location of CITED2 in the p53 signaling pathway (Fig. 2c). Suppression of CITED2 expression did not significantly affect the mRNA expression of p21, Bax, and p53R2, downstream molecules of p53, with or without BA treatment. CITED2 expression was also upregulated by BA treatment at the protein level in UCCA24 after 1–12 h, with a peak at 6 h, accompanied by p53 protein accumulation and followed by p21 and Bax expression (Fig. 3). Although Bax expression was high in the non-treatment state, Bcl-2 levels were also high and decreased with BA treatment. Upregulation of CITED2 was noted within 6 h of CITED2 transfection and p53 protein accumulation was observed in parallel. Thus, on treatment with BA, the p53 protein level was upregulated without transcriptional upregulation. As p53 downstream molecules were upregulated at the mRNA and protein levels, stabilization of p53 protein without transcriptional expression could account for this discrepancy. Therefore, CITED2 could be a candidate transducer of BA responsible for post-transcriptional upregulation of p53.

BA induces apoptosis through MCT1, CITED2, and p53

As we previously reported [12], BA induced apoptosis detected by the TUNEL method in the UCCA24 line (Fig. 4a, b). Apoptosis was abundantly induced by doxorubicin with wild-type p53 as previously reported [16]. It is well known that the doxorubicin-induced apoptosis is p53-dependent [25], and apoptosis was inhibited by an siRNA for p53, but not by non-sense siRNA. BA-induced apoptosis was inhibited by 4-hydroxycinnamate (4CHC), a specific inhibitor for MCT1 in both types of cell lines, and was also significantly inhibited by siRNA for p53. CITED2 overexpression also induced apoptosis in UCCA24, and this was partially and significantly reduced by p53 siRNA, but not by non-sense siRNA. As BA activates p53 by stabilization and does not induce transcriptional expression, the partial inhibition suggests reduction of the protein pool before BA treatment or CITED2 induction. The results indicate that BA induces apoptosis through MCT1, CITED2, and p53 activation. Although the BA-induced

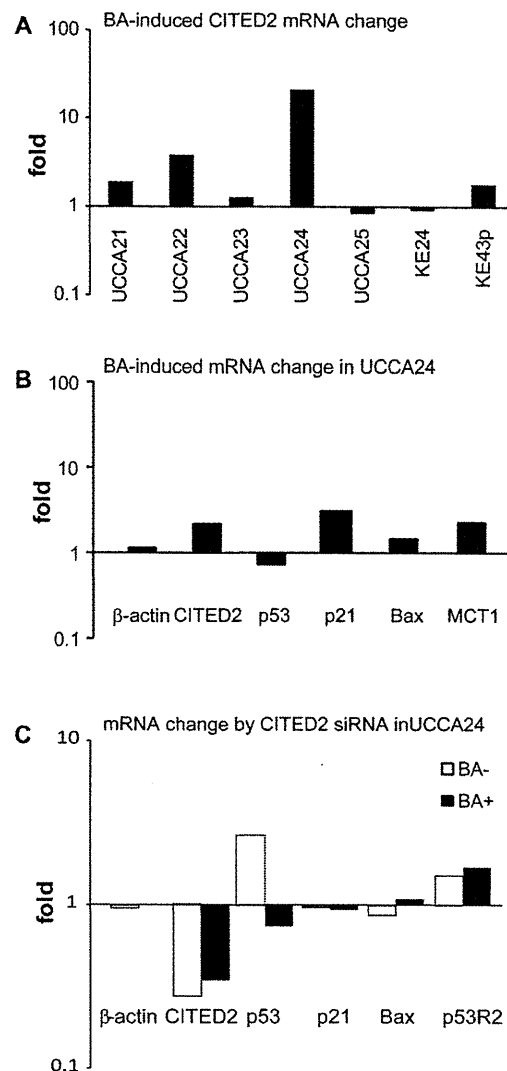


Fig. 2 q-RT-PCR for validation of CITED2 and p53-related genes, and knockdown effects of CITED2. Representative results for three independent experiments are shown. **a** CITED2 mRNA expression is induced by BA in UCCA cell lines. q-RT-PCR was performed after 2.5 mM BA was added for 24 h. UCCA cell lines, including UCCA 21 and 24 used for microarray analysis, tended to show upregulation of CITED2 mRNA expression against no BA treatment, whereas sCRC lines (KE24, KE43p) did not. **b** mRNA expression of p53-related molecules except p53 itself was increased in UCCA24 by BA treatment for 24 h, as analyzed by q-RT-PCR. Fold ratios of mRNA expression after 24 h BA treatment against no BA treatment are shown in the histogram. **c** siRNA for CITED2 was transfected 24 h before BA treatment in UCCA24. After BA treatment for 24 h, expression of β-actin, CITED2, p53, p21, Bax, and p53R2 was analyzed by q-RT-PCR. Fold expression of CITED2 siRNA transfected cells was compared with non-specific siRNA transfection. *Open* and *filled bars* show fold ratios for each gene against VIC-labeled β-actin expression with and without BA treatment

CITED2 level was almost equivalent to or greater than that in CITED2 transfected cells (Fig. 3a, b), the apoptosis level was much greater in CITED2 transfected cells (Fig. 4b).

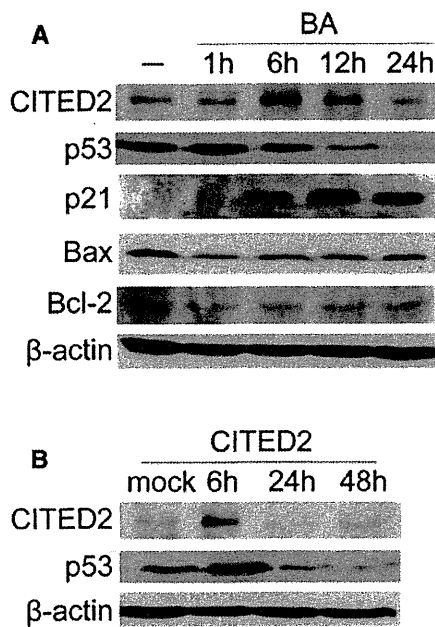


Fig. 3 CITED2, p53, and related protein expression with BA treatment and forced CITED2 expression in UCCA24. The indicated protein expression was analyzed by western blotting with BA treatment for 1, 6, 12, and 24 h (a) and CITED 2 transfection for 6, 24, and 48 h (b). CITED2 was transfected 24 h prior to BA treatment

This might be due to the direct effect of CITED2 by transfection whereas the apoptosis induced by BA could be regulated by other factors.

BA and CITED2 induce acetylation of p53 and increase protein expression

CITED2 has been identified as a co-factor for CBP/p300 [18], which induces apoptosis through acetylation of p53 [24, 26]. The amount of p53 protein was significantly increased along with the acetylated form of p53 with BA treatment (Fig. 5a) and CITED2 transfection (Fig. 5b). It is likely that CITED2 stabilizes p53 by its acetylation.

CITED2 binds to CBP/p300 in vitro

During BA treatment, we confirmed binding of CITED2 to CBP/p300 in vitro by immunoprecipitation (Fig. 5c). Binding was induced by CITED2 and CBP/p300 transfection as well as by BA treatment.

CITED2 is expressed in the surface epithelium of UC patients' colonic mucosa

Expression of CITED2 protein was evaluated in the colonic mucosa of UC patients in situ. The area (0–3) and the intensity (0–3) of CITED2 positivity were multiplied

together and the correlation of the resulting CITED2 expression score with histological evidence of inflammation was assessed according to Mats (see Fig. 6a) [21]. CITED2 expression was localized in crypt bottoms of normal colonic mucosa. In contrast, it was found to be significantly increased in the surface lining epithelium and both upper and lower halves of the crypts in UC cases, whereas the increase of expression was limited to the lower halves in diverticulitis, with non-specific inflammation. In addition to the significant differences in expression score between histological grades by the Mann–Whitney's *U* test shown in Fig. 6b, a significant correlation of the increase of CITED2 expression with histological grade by Spearman's rank correlation test was also demonstrated, especially in lining epithelium and the upper halves of crypts ($\rho = 0.641$, $p < 0.0001$, lining epithelium, $\rho = 0.691$, $p < 0.0001$, upper crypts, $\rho = 0.460$, $p = 0.0002$, lower crypts).

Discussion

There have been many studies of the mechanisms underlying UC. Concerning roles of enterobacteria in UC development, bacterial invasion into the mucosal epithelia is considered to be associated with the production of cytokines, as a first step in inflammatory processes [10]. Thus, inflammatory cytokines produced by macrophages and lymphocytes in the stroma of the colonic mucosa are considered important factors. Interleukin-10 (IL-10)-deficient mice spontaneously develop UC-like colitis under conventional conditions, but not under specific pathogen- or germ-free conditions [6, 7]. To induce colitis in IL-10-deficient mice, commensal enteric bacteria are necessary [7]. Shkoda et al. [8] clearly revealed that IL-10 inhibits inflammation-induced endoplasmic reticulum (ER)-mediated stress responses by modulating activating transcription factor (ATF)-6 nuclear recruitment to the glucose-regulated ER stress protein-78 gene promoter in colonic epithelial cells. In inflammatory foci of UC, p53 is activated in the crypt epithelium [12], which suggests protective activity against inflammatory stress. Therefore, mutations of p53, frequently found in the early stages of UC-associated tumorigenesis, including non-tumorous regenerative mucosa, in contrast to the adenoma-carcinoma sequence [27], may greatly contribute to carcinogenesis. Short-chain fatty acids including BA are important to maintain colonic functions, and BA constitutes an energy supply for colonocytes and plays a central role in homeostasis of the colonic mucosa [28]. Previously, we detected *F. varium* by direct culture of UC patients' mucosa [14] and induced experimental UC in mice by intra-rectal administration of BA secreted from *F. varium* [9, 11]. In addition, BA induces DNA repair and apoptosis with the activation of

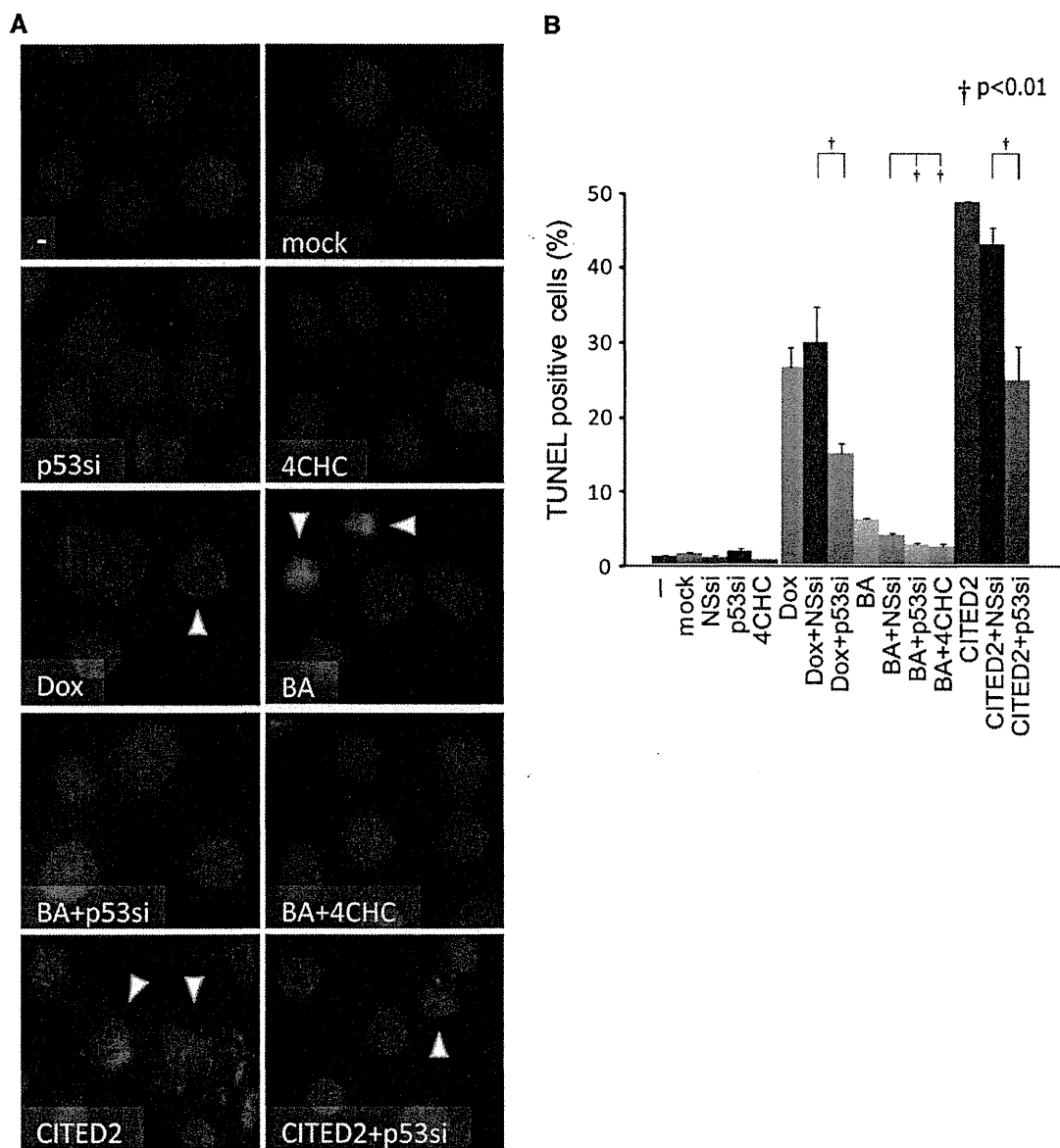


Fig. 4 Induction of p53-dependent apoptosis by BA treatment and CITED2 expression. **a** Representative images of apoptosis detected by the TUNEL method under various conditions in UCCA24 (NT no treatment, Dox treated with doxorubicin, 4CHC treated with 4CHC, BA stimulated with BA, BA + 4CHC treated with both BA and 4CHC, CITED2 CITED2 transfected).

b Average percentages of apoptotic cells in three experiments in UCCA24 in the histogram with standard deviation bars. NSsi and p53si, siRNA for non-sense and p53 transfected, respectively. †Significant differences ($p < 0.01$) by Student's *t* test for each condition are also shown

p53–p53R2 system [12]. Therefore, BA would damage colonocytes in a high concentration condition. In order to mimic the inflammatory condition of UC in vitro, we used UCCA cell lines derived from human UC-associated adenocarcinomas [16]. Although UCCA lines are tumor cell lines and may have different characteristics from non-tumorous colonic epithelium, those used here are unique in being derived from a human UC background. As BA-induced apoptosis and activated p53-dependent DNA repair

have been found in both sCRC and UCCA lines [11, 12], we here focused on their roles as a key factor for UC inflammation.

First, we established that mRNA expression of CITED2 and several other genes was upregulated by BA treatment in ulcerative colitis-associated cancer cell lines, but not in sporadic counterparts, by cDNA microarray analysis with the Percellome method [17]. Whereas two independent probe sets were listed as only upregulated in UCCA lines,

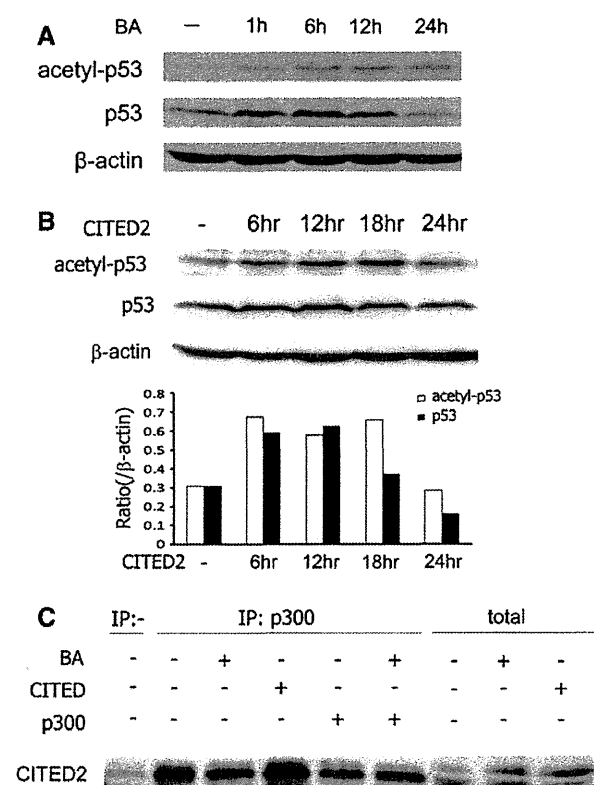


Fig. 5 Induction of p53 acetylation by BA and CITED2, accompanied by binding of CBP/p300 and CITED2 in UCCA24. **a** A 2.5 mM BA treatment induced acetylation of p53 and increased the total amount of p53 from 1 to 24 h. **b** Transfection of CITED2 24 h prior to BA treatment for 24 h also induced acetylated p53 and increased the total amount of p53. A histogram of results for relative expression of acetylated p53 and p53 is shown below. Relative expression against β -actin was calculated by densitometry of western blot data. **c** Immunoprecipitation by anti-CBP/p300 antibody and immunoblotting with anti-CITED2 antibody showed binding of CITED2 and CBP/p300. BA indicates treatment by 2.5 mM BA for 24 h. CITED2 and p300 indicate the transfection of CITED2 and CBP/p300 24 h prior to the assay, respectively. A western blot from the total lysate is also illustrated

we especially focused on CITED2 among the 13 genes, because its function seemed to be closely related to p53 via CBP/p300 [18], with an upstream action. Indeed, RNAi analysis directly indicated that CITED2 is upregulated upstream of p53. Induction of CITED2 mRNA was not evident in several UCCA lines, such as UCCA23 and UCCA25 (Fig. 2). However, we confirmed the induction of CITED2 protein in those lines (data not shown) and the upregulation of CITED2 was statistically confirmed by the immunohistochemistry from the 102 inflammatory lesions of UC (Fig. 6).

Although CITED2 has been noted as an important factor for organ or tissue development [29, 30], it was originally found to inhibit the binding to hypoxia-inducible factor 1 (HIF1) [31]. Bakker et al. [32] reported CITED2 to be

transcriptionally induced by FOXO3a, a member of the FOXO subfamily of Forkhead transcription factors, and to inhibit HIF1-induced apoptosis in hypoxia, which is proposed as a survival response of cancer cells. Although HIF1 induces p53-dependent apoptosis [33–35], it also causes p53-independent apoptosis via BNIP3 and NIX induction [34–36]. As Bakker et al. used p53 wild-type cell lines, it was uncertain whether CITED2 inhibited p53-dependent HIF1-induced apoptosis. In the present study, however, BA and CITED2 clearly induced apoptosis, which was inhibited by an siRNA for p53. Furthermore, the BA-induced apoptosis was inhibited by an inhibitor of MCT1, through which BA is transported into the cytoplasm [37, 38], and an siRNA for p53 partially but significantly inhibited the BA- and CITED2-induced apoptosis. Apoptosis induced by CITED2 transfection was more frequent than by BA treatment (Fig. 4b), although the BA-induced CITED2 level was almost equivalent to or greater than that in CITED2 transfected cells (Fig. 3a, b). This might be due to a direct effect of CITED2 by transfection or to apoptosis induced by BA being regulated by other factors. The dependence of BA-induced apoptosis on MCT1, demonstrated by 4CHC, an inhibitor of MCT1, should be verified by other approaches. Although protein stabilization of p53 was apparent on BA treatment with CITED2 overexpression, knockdown of p53 at the mRNA level could also inhibit the BA–CITED2 pathway. Therefore, CITED2 possibly plays different roles in BA-induced and HIF1-induced apoptosis, which might depend on modulation of the phosphorylation or acetylation sites of p53.

According to recent studies, p53 induces apoptosis via acetylation of lysine 120 on DNA damage or in response to stress signaling with CBP/p300 binding [24, 26]. The acetylated form of p53 is stable with regard to the Mdm2-ubiquitination system [39]. We here demonstrated acetylation of p53 by BA treatment and forced CITED2 expression accompanied by p53 protein accumulation (Fig. 5a, b). At the same time, binding of the CITED2 and CBP/p300 was also observed by immunoprecipitation (Fig. 5c). Therefore, we conclude that CITED2 is upregulated by BA, in turn activating p53-dependent apoptosis via acetylation of p53 protein (Fig. 7). Although transcription of p53 was not observed, the data support p53 protein stabilization by BA and CITED2. Further investigation of the half-life of p53 after BA treatment with or without CITED2 knockdown should clarify this point.

In UC, p53 gene alteration is observed even in non-tumorous regenerative epithelium [27]. In addition, p53 expression is upregulated in inflammatory foci of UC without gene mutation and the p53 and p53R2 DNA repair system is accelerated [12]. Therefore, the repair capacity may be overloaded in active UC. We previously found greater upregulation of Bax protein in UCCA as compared

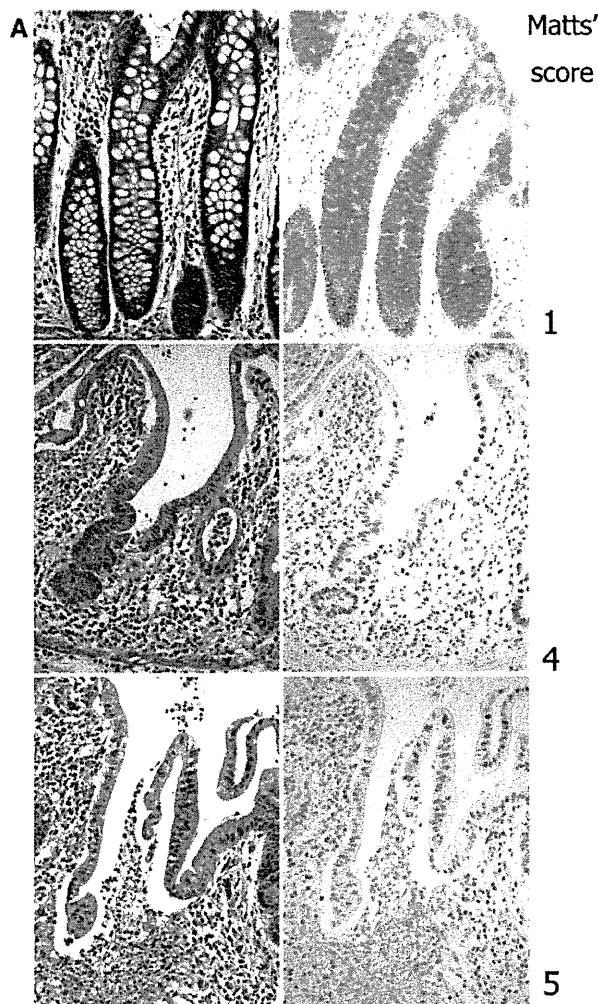


Fig. 6 CITED2 expression in inflammatory foci of ulcerative colitis. **a** Immunohistochemical staining of CITED2 is shown in the *right panels* and semi-serial H&E sections are shown in the *left panels*. Examples of inactive UC (Matts' score 1, *upper panel*) and active UC (Matts' score 4 and 5, *middle and lower panel*) are presented. Strong CITED2 expression is remarkable in epithelial nuclei in active UC. **b** CITED2 expression correlates with activity of inflammation of UC in situ. The area (0–3) and the intensity (0–3) of CITED2 expression were evaluated and the CITED2 expression score (0–9) was calculated by multiplying the two results. The inflammatory grade of the diverticulosis cases was scored from 0 to 2 (none, mild, severe). The CITED2 expression scores are shown along with histological scores. Median values are shown by the *horizontal line*, the *box* represents values between the 25th and 75th percentiles, and the *lower and upper bars* indicate the 10th and 90th percentiles, respectively. †*p* < 0.01 and ‡*p* < 0.05 by Mann–Whitney's *U* test

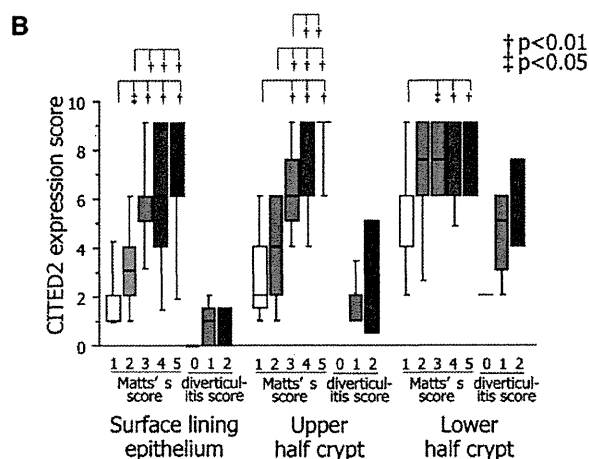


Fig. 7 Scheme for BA-induced apoptosis. BA produced by commensal bacteria, such as *F. varium*, is incorporated by MCT1 and induces CITED2 expression. The CITED2-CBP/p300 complex then acetylates p53 which stabilizes the protein, resulting in upregulated transcription of proapoptotic genes including Bax, and induction of apoptosis. Apoptosis of colonic epithelial cells leads to erosion of the mucosa, so that bacteria can more easily colonize. *Ac* acetylation; *BA* butyric acid; *MCT1* monocarboxylate transporter-1

with sCRC [40], which might have a similar effect. Matts' histological scores based on the infiltration of neutrophils and the presence of cryptitis, crypt abscess and erosion, reflect the inflammation activity of UC [21]. The fact that

CITED2 expression in UC is also increased in the upper halves of the crypts and positively correlated with the histological activity of UC, evaluated by Matts' histological scores, is clear evidence of an association with UC inflammation. In contrast, although the CITED2 expression was increased in the inflammatory foci of diverticulitis, the expression score was almost at the same level as that of the inactive UC (Matts' score 1). This might indicate that the BA-CITED2 system is specifically involved in the UC inflammation, and not under conditions of non-specific colonic inflammation such as diverticulitis.

Previously, we showed that specific enterobacteria including *F. varium* are frequently detectable in UC patients' mucosa. Here we showed BA-induced apoptosis via CITED2 induction and p53 activation, and that CITED2 expression clearly correlates with histological UC

activity. Although the reported role of CITED2 in hypoxic stress does not coincide with our present results [32], our present histopathological study supports a scenario in human UC involving another CITED2-activating pathway. Thus, highly concentrated BA produced by enteric bacteria would induce CITED2-p53-dependent apoptosis in the colonic epithelium and cause erosion, which is typical of active UC. With mucosal erosion, enteric bacteria can easily infiltrate into the colonic stroma and stimulate macrophages [41], which would enhance inflammation by producing inflammatory cytokines. Apoptosis induced by BA was here inhibited by an siRNA for p53, as well as by 4CHC, an inhibitor of MCT1 (Fig. 4b), which suggests the possibility of anti-inflammation targeted therapy. However, mechanisms of UC generation still need further analysis, particular with regard to enteric bacteria. In conclusion, we propose that p53 activation via CITED2 upregulation by BA and consequent p53 acetylation is a possible mechanism underlying UC development.

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Conflicts of interest The authors disclose no conflicts with this work.

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Original Article

Development of humanized steroid and xenobiotic receptor mouse by homologous knock-in of the human steroid and xenobiotic receptor ligand binding domain sequence

Katsuhide Igarashi¹, Satoshi Kitajima¹, Ken-ichi Aisaki¹, Kentaro Tanemura¹,
Yuhji Taquahashi¹, Noriko Moriyama¹, Eriko Ikeno¹, Nae Matsuda¹, Yumiko Saga^{2,3},
Bruce Blumberg⁴ and Jun Kanno¹

¹Division of Cellular and Molecular Toxicology, Biological Safety Research Center,
National Institute of Health Sciences, 1-18-1 Kamiyoga, Setagaya-ku, Tokyo, 158-8501, Japan

²Division of Mammalian Development, National Institute of Genetics, Yata 1111, Mishima 411-8540, Japan

³The Graduate University for Advanced Studies (Sokendai), Yata 1111, Mishima 411-8540, Japan

⁴Department of Developmental and Cell Biology, 2011 Biological Sciences 3, University of California,
Irvine, CA 92697-2300, USA

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ABSTRACT — The human steroid and xenobiotic receptor (SXR), (also known as pregnane X receptor PXR, and NR1I2) is a low affinity sensor that responds to a variety of endobiotic, nutritional and xenobiotic ligands. SXR activates transcription of Cytochrome P450, family 3, subfamily A (CYP3A) and other important metabolic enzymes to up-regulate catabolic pathways mediating xenobiotic elimination. One key feature that demarcates SXR from other nuclear receptors is that the human and rodent orthologues exhibit different ligand preference for a subset of toxicologically important chemicals. This difference leads to a profound problem for rodent studies to predict toxicity in humans. The objective of this study is to generate a new humanized mouse line, which responds systemically to human-specific ligands in order to better predict systemic toxicity in humans. For this purpose, the ligand binding domain (LBD) of the human SXR was homologously knocked-in to the murine gene replacing the endogenous LBD. The LBD-humanized chimeric gene was expressed in all ten organs examined, including liver, small intestine, stomach, kidney and lung in a pattern similar to the endogenous gene expressed in the wild-type (WT) mouse. Quantitative reverse transcription-polymerase chain reaction (RT-PCR) analysis showed that the human-selective ligand, rifampicin induced Cyp3a11 and Carboxylesterase 6 (Ces6) mRNA expression in liver and intestine, whereas the murine-selective ligand, pregnenolone-16-carbonitrile did not. This new humanized mouse line should provide a useful tool for assessing whole body toxicity, whether acute, chronic or developmental, induced by human selective ligands themselves and subsequently generated metabolites that can trigger further toxic responses mediated secondarily by other receptors distributed body-wide.

Key words: Steroid and xenobiotic receptor, Pregnane X receptor, Humanized mouse,
Ligand binding domain, Knock-in mouse

INTRODUCTION

Most orally administered xenobiotics are metabolized first by the intestine and then by the liver after portal transport. The expression levels of enzymes involved in xenobiotic metabolism are regulated at the transcriptional level by key xenobiotic sensors including the ster-

oid and xenobiotic receptor (SXR), also known as the pregnane X receptor (PXR), pregnane activated receptor (PAR) and NR1I2 (Bertilsson *et al.*, 1998; Lehmann *et al.*, 1998; Blumberg *et al.*, 1998). SXR is important in the field of toxicology for at least two reasons. Firstly, this receptor system induces the expression of CYP3A and CYP2B enzymes, the major metabolizers of pharmaceu-

Correspondence: Jun Kanno (E-mail: kanno@nihs.go.jp)

ticals and xenobiotics. Therefore, SXR is a key mediator of drug- and chemical-induced toxicity as well as drug-drug and drug-nutrient interactions (Zhou *et al.*, 2004). Secondly, the orthologous rodent and human receptors exhibit differential sensitivity for a subset of chemical ligands important in the field of toxicology. For example, rifampicin (RIF) is a specific and selective activator of human SXR, whereas pregnenolone 16 α -carbonitrile (PCN) is selective for the rodent orthologue.

Rodent-human differences in CYP3A and CYP2B-mediated responses to xenobiotics can be a profound problem in toxicologic studies where rodents are used to predict the toxicity of a compound in humans (Ma *et al.*, 2007). Therefore, development of a murine model that reconstructs the SXR-mediated systemic response of humans is of a great significance in toxicology.

Human and rodent SXRs share ~95% amino acid sequence identity in the DNA-binding domain (DBD) but only about 77% identity in the LBD. Tirona *et al.* (2004) analyzed the ligand selectivity of a human-rat chimeric protein and showed that the species differences are primarily defined by sequence differences in the LBD. Watkins and colleagues showed that the key residues responsible for the majority of the ligand selectivity were Leu 308 (human) and Phe305 (rat and mouse). Crystallographic analysis located these amino acids within or neighboring the flexible loop that forms a part of the pore to the ligand-binding cavity. Swapping the rodent and human-specific residues was shown to modulate the activation by the human-selective activator RIF *in vitro* (Watkins *et al.*, 2001). According to those findings, a simple replacement of the mouse LBD with the human sequence should be sufficient to "humanize" the ligand binding properties as well as activation of the downstream target genes.

Three kinds of humanized mice have already been generated. One is the SXR-null/Alb-SXR mouse (Alb-SXR mouse) made by crossing the SXR knockout mice with a transgenic mouse line that expresses human SXR in liver under the control of the albumin promoter (Xie *et al.*, 2000). Gonzalez and colleagues generated a transgenic mouse expressing a human BAC containing the entire hSXR gene in a SXR null background, thus controlled under human SXR promoter (SXR BAC mouse) (Ma *et al.*, 2007). Another mouse is the human SXR genome knock-in mice (hSXR genome mouse) (Scheer *et al.*, 2008). The human SXR genomic region from exon 2 to exon 9 was knocked-in to mouse SXR exon 2. This mouse expresses the human full length SXR mRNA under the control of mouse SXR promoter regulation. Although useful for toxicology studies, these mice

have disadvantages in that the human SXR is expressed only in the liver (Alb-SXR mouse), hSXR mRNA is not expressed in all of the tissues where SXR is known to be expressed (SXR BAC mouse), and there might be potential differences in the binding affinities of hSXR DNA-binding domain (DBD) to *cis*-acting elements in mouse SXR target genes (hSXR genome mouse).

As noted above, it is known that the critical differences between human and rodent ligand-selectivity reside in the LBD. Therefore, when our project to generate a humanized SXR mouse was initiated, we reasoned that altering the LBD would be sufficient to generate a humanized ligand selectivity. We decided to retain the mouse DBD to avoid any potential differences between the binding affinities of the chimeric receptor for *cis*-acting elements in the mouse genome. To maintain the tissue-specific expression pattern of the endogenous gene, we inserted the human cDNA encoding the region carboxyl-terminal to the DBD into the mouse gene. This retains all of the 5' and 3' regulatory elements in the mouse gene, as well as introns 1 and 2, which contain important elements for regulating SXR expression (Jung *et al.*, 2006).

Here we report a new line of mouse (hSXRki mouse) in which a cDNA encoding the human LBD is homologously recombined into the mouse gene after exon 3. The tissue distribution of the resulting chimeric mouse DBD-human LBD mRNA is comparable to that of the WT mouse. The hSXRki mouse showed a fully humanized response to the human-selective activator RIF in that the Cyp3a11 mRNA was induced in liver and mucosa of small intestine in response to RIF, but not the rodent-selective compound PCN. This new mouse line should provide a useful tool for assessing the whole body toxicity induced by a human selective SXR ligand itself and its subsequently generated metabolite(s) that can trigger further toxic responses through other pathways body-wide.

MATERIALS AND METHODS

Generation of hSXRki knock-in mice

A DNA fragment of mouse SXR intron 2 to exon 3 was PCR amplified using mouse BAC DNA (BAC clone No. RP23-351P21) as a template. Primers used were BAC39486FW and mSXR462RV (for sequences of the primers see Table1). This fragment was connected to the LBD of human SXR cDNA from amino acid 105 through the carboxyl terminus amplified by the PCR primers: hSXR904FW and hSXR1887RVEcoRI (template; human SXR cDNA). The 3'UTR of bovine growth hormone (BGH) was added to 3' to the terminal codon. This concatenated fragment was introduced to a vector, which

Humanized SXR Mouse by knock-in of human SXR LBD

Table 1. List of primer pairs

Purpose	Primer name	Sequence (5' to 3')
Targeting vector construction	BAC39486FW	CCATGGGTACCACGAATAACAA
	mSXR462RV	CATGCCACTCTCCAGGCA
	hSXR904FW	AAGAAGGAGATGATCATGTCCG
	hSXR1887RVEcoRI	CCGAATTCTCATCATCAGCTACCTGTGATACCGAACA
Genotyping	NeoAL2	GGGGATGCGGTGGGCTCTATGGCTT
	SXR RC RV5	TGAGAGTGCACAAGTTCAAGCT
	WTInt5	AGTGATGGGAACCACTCCTG
	WTEx6RV	TGGTCTCAATAGGCAGGTC
	mhSXRE4	GTGAACGGACAGGGACTCAG
	mhSXRSARV	CTCTCCTGGCTCATCCTCAC
	Percellome quantitative RT-PCR	Cyp3a11 FW
Cyp3a11 RV		TCAAACAACCCCATGTTTT
Ces6 FW		GGAGCCTGAGTTCAGGACAGAC
Ces6 RV		ACCCTCACTGTTGGGGTTC
mouse SXR FW		AATCATGAAAGACAGGGTTC
mouse SXR RV		AAGAGCACAGATCTTTCCG
human SXR FW		ATCACCCGGAAGACACGAC
human SXR RV		AAGAGCACAGATCTTTCCG
mouse-human SXR FW		CCCATCAACGTAGAGGAGGA

has the neomycin resistance gene with loxP sequence at both ends, removable with Cre recombinase (Saga *et al.*, 1999). A 7kb KpnI fragment containing intron 2 was used as a long arm and 1.3kb PstI-EcoRI fragment containing from exon 8 to intron 8 was used as a short arm for homologous recombination (Fig. 1). The resulting targeting vector was linearized with SacII and introduced by electroporation to TT2 ES cell line (Yagi *et al.*, 1993) and neomycin resistant clones were selected, PCR genotyped, and confirmed by the Southern blotting. For generation of chimeric mice, these ES clones were aggregated with ICR 8-cell embryos and transferred to pseudopregnant female recipients. The chimeric mice born were bred with ICR females. Germ line transmission of the targeted allele was confirmed by PCR. A mouse was crossed with a CAG-Cre transgenic mouse (Sakai and Miyazaki, 1997) to evict the neomycin resistance gene, and back crossed to C57BL/6 CrSlc (SLC, Inc., Shizuoka, Japan) at least 6 generations and used for the analysis.

PCR Genotyping**(See Table 1 for primer sequences)**

Primers for identification of homologously recombined ES clones were NeoAL2 and SXR RC RV5. DNA purified from the tail of each mouse was used for PCR genotyping. Primers for WT detection were WTInt5 and WTEx6RV amplifying a product of 755 bp. Primers for

confirmation of removal of the neomycin resistance gene were mhSXRE4 and mhSXRSARV amplifying a product of 1,223 bp.

Southern blot analysis

To confirm homologous recombination, DNA from ES cell cultures was purified and digested with BamHI and XhoI, then electrophoresed and analyzed by Southern hybridization (Saga *et al.*, 1997). Mouse SXR exon 9 region which remains after homologous recombination was used for the probe. The restriction fragments from the WT allele and targeted allele are 2,305 bp and 1,925 bp, respectively.

Chemicals

RIF (molecular weight 822.95) and PCN (molecular weight 341.49) were purchased from Sigma-Aldrich (St. Louis, MO, USA). Corn oil was purchased from Wako Pure Chemical Industries (Osaka, Japan).

Quantitative RT-PCR (Percellome PCR)**(See Table 1 for primer sequences)**

The method for Percellome quantitative RT-PCR was described previously (Kanno *et al.*, 2006). Briefly, tissue pieces stored in RNAlater (Ambion, Austin, TX, USA) were homogenized and lysed in RLT buffer (Qiagen GmbH., Germany) and 10 µl aliquots were used

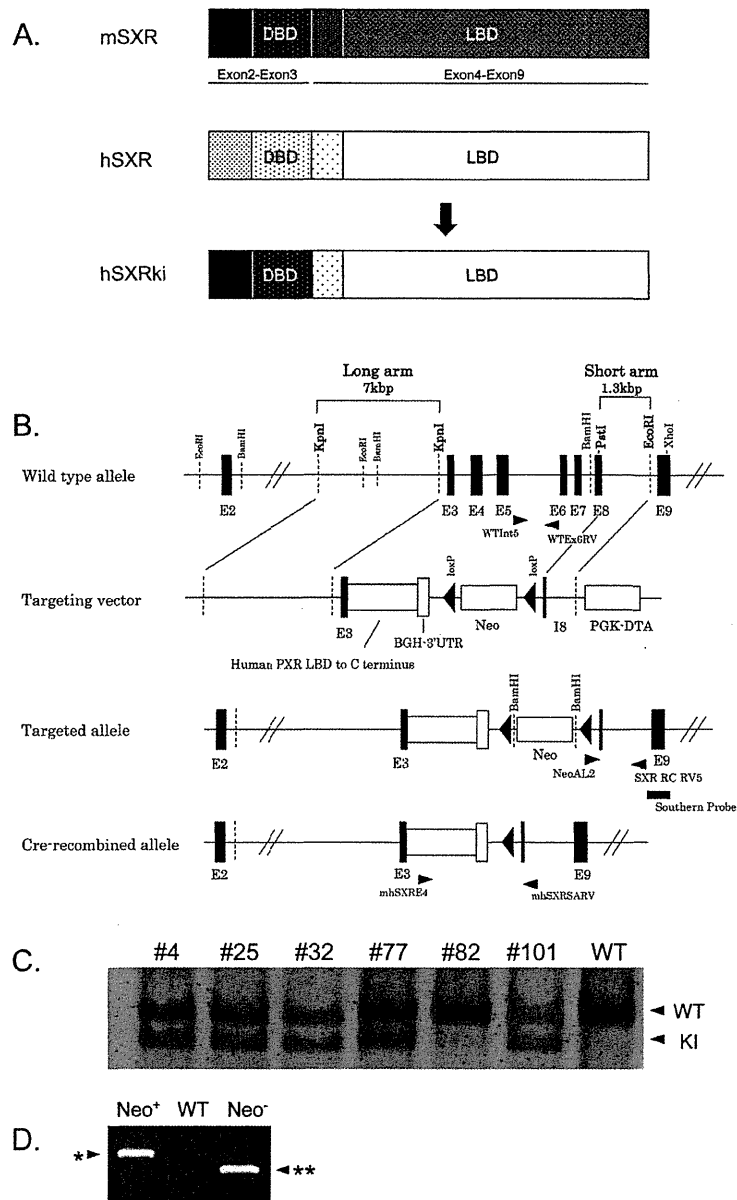


Fig. 1. Targeting strategy used to generate the hSXRki mouse. A) Diagram of hSXRki chimeric protein. Hinge region and ligand binding domain (LBD) of human SXR are knocked-in to mouse SXR, resulting in chimeric protein having murine N-terminal domain and DNA binding domain (DBD). B) Targeting strategy used to generate the hSXRki mouse. The chimeric mouse DBD and human LBD fragment, followed by the BGH 3' UTR were knocked-in to the mouse SXR gene. The genomic region spanning from exon 3 to exon 8 was substituted by the inserted fragment with the remainder of the gene remaining intact. C) Confirmation of homologous recombination by southern blot analysis. Six ES clones positive for recombination by PCR genotyping were further analyzed by southern blot (clones #4 ~ #101). Lower bands (1925 bp) indicate successful homologous recombination; upper bands (2305 bp) correspond to WT allele. Clones #4, #25, #32, #77 and #101 were confirmed as homologous recombinants; clones #4 and #25 were used for the generation of chimeric mice. D) Confirmation of Cre-mediated removal of the neomycin resistance gene. Mouse tail genome DNA was PCR amplified with the primer set, mhSXRE4 and mhSXR SARV. *: 2,858 bp (for the mice having the neomycin resistance gene), **: 1,223 bp (for the mice without the neomycin resistance gene).