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Cholesterol sulfate is a DOCK2 inhibitor that mediates tissue-specific immune evasion in the eye

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Although immune responses are essential to protect the body from infection, they can also harm tissues. Certain tissues and organs, including the eye, constitute specialized microenvironments that locally inhibit immune reactivity. Deducator of cytokinesis protein 2 (DOCK2) is a Rac-specific guanine nucleotide exchange factor (GEF) that is predominantly found in hematopoietic cells. DOCK2 plays a key role in immune surveillance because it is essential for the activation and migration of leukocytes. DOCK2 mutations cause severe immunodeficiency in humans. We found that DOCK2-mediated Rac activation and leukocyte migration were effectively inhibited by cholesterol sulfate (CS), but not by cholesterol or other sulfated steroids. CS bound to the catalytic domain of DOCK2 and suppressed its GEF activity. Mass spectrometric quantification revealed that CS was most abundantly produced in the Harderian gland, which provides the lipids that form the oily layer of the tear film. Sulfation of cholesterol is mediated by the sulfotransferases SULT2B1b and, to a lesser extent, SULT2B1a, which are produced from the same gene through alternative splicing. By genetically inactivating Sult2b1, we showed that the lack of CS in mice augmented ultraviolet- and antigen-induced ocular surface inflammation, which was suppressed by administration of eye drops containing CS. Thus, CS is a naturally occurring DOCK2 inhibitor and contributes to the generation of the immunosuppressive microenvironment in the eye.

INTRODUCTION

The immune system has evolved to recognize and interact with microorganisms to protect the body from infection. Leukocytes continually patrol the body to identify invading pathogens and elicit immune responses against them. However, immune responses generally carry a risk for damaging or impairing the function of vital tissues and therefore could threaten the survival of the host (1, 2). Certain tissues and organs, such as the brain and the pregnant uterus, constitute specialized microenvironments that locally inhibit immune reactivity (3–5). Although this phenomenon is classically known as immune privilege, various other tissues also create microenvironments that help them evade the immune system. Among these are tumors (6), in which local immunosuppressive mechanisms predominate over stimulatory immune responses. Several molecular and cellular mechanisms that suppress tissue inflammation have been reported. These include the production of cell-surface molecules, such as the Fas ligand or programmed cell death ligand 1, secretion of the anti-inflammatory cytokine interleukin-10 or transforming growth factor–β, and induction of intracellular enzymes with immunoregulatory effects, such as indoleamine 2,3-dioxygenase, arginase, or inducible nitric oxide synthase (7–10). Although it seems reasonable to expect that bioactive lipids could contribute to immune evasion, no evidence for their participation in immune evasion has been reported.

Dedicator of cytokinesis protein 2 (DOCK2) is a member of the CDM family of proteins, named for Caenorhabditis elegans cell-death abnormal (CED–)–5, mammalian DOCK1 and DOCK5, and Drosophila melanogaster Myoblast City, and is predominantly found in hematopoietic cells (11). Although DOCK2 does not contain Dbl homology (DH) and pleckstrin homology (PH) domains, which are typically found in guanine nucleotide exchange factors (GEFs), DOCK2 mediates the exchange of guanosine diphosphate (GDP) for guanosine triphosphate (GTP) on the guanosine triphosphatase Rac through its DOCK homology region–2 (DHR–2) domain (12, 13). DOCK2 is a major Rac GEF acting downstream of chemotactic receptors and antigen receptors in lymphocytes, neutrophils, and natural killer cells and plays key roles in the migration and activation of these cell types (14–17). Consistent with this, deletion of Dock2 in mice prevents cardiac allograft rejection and the development of autoimmune disease (18, 19). DOCK2 mutations also cause severe immunodeficiency in humans (20). Thus, DOCK2 is essential for immune surveillance and immune responses in both humans and mice.

Sulfonation plays an important role in the biological activity of many endogenous molecules. Cholesterol sulfate (CS) is a sulfated derivative of cholesterol and is widely distributed in various tissues and body fluids (21). In skin, CS is located predominantly in the epidermis, where it contributes to epidermal differentiation as well as to the development and maintenance of the epidermal barrier function (21–25). In addition, CS has been implicated in many biological processes including sperm capacitation, platelet adhesion, blood clotting, leukotriene biosynthesis, and T cell receptor signaling (26–30). However, its physiological functions and the underlying mechanisms are not fully understood. Here, we identified CS as a naturally occurring DOCK2 inhibitor. Under physiological conditions, CS was...
most abundantly produced in mice in the Harderian gland, which provides lipids to form the oily layer of the tear film (31). Although the lack of CS augmented ultraviolet (UV)– and antigen-induced ocular surface inflammation, administration of eye drops containing CS limited this inflammatory response. Our results thus indicate that CS mediates immune evasion in the eye by inhibiting DOCK2.

RESULTS
CS binds to the DHR-2 domain of DOCK2 and inhibits its GEF activity
Because DOCK2 deficiency suppresses allograft rejection and autoimmune disease development in mice (18, 19), DOCK2 could serve as a molecular target controlling inflammatory responses. To search for potential DOCK2 inhibitors, we screened 577 pharmacologically active compounds for their ability to inhibit DOCK2-mediated Rac activation in vitro and identified CS (Fig. 1A) as a candidate. CS inhibited the Rac GEF activity of the DHR-2 domain of murine DOCK2 in vitro in a dose-dependent manner with a half-maximal inhibitory concentration (IC50) of 2.0 μM (Fig. 1B and C). Similar results were obtained using the DHR-2 domain of human (fig. S1A). Although CS also inhibited murine DOCK1 and DOCK5 (fig. S1B), CS did not affect Rac activation mediated by the other DH-type GEFs Tiam1 and Trio at any of the tested concentrations (Fig. 1B and fig. S1C). When the sulfate moiety of CS was removed or replaced with acetate, the resulting compounds, cholesterol, or cholesterol acetate (CA), respectively, exhibited greatly reduced DOCK2 inhibitory activity (Fig. 1C). Similarly, other sulfated steroids such as pregnenolone sulfate (PREGS), dehydroepiandrosterone sulfate (DHEAS), estrone sulfate (E1S), estradiol sulfate (E2S), and estriol sulfate (E3S) failed to inhibit the Rac GEF activity of DOCK2 (Fig. 1C). These results indicate that both the sulfate moiety and the cholesterol side chain are required for the inhibitory activity of CS. Surface plasmon resonance (SPR)–based binding assays revealed that CS directly bound to the DOCK2 DHR-2 domain with a dissociation constant of 9.9 nM (Fig. 1D). This binding was specific because cholesterol derivatives other than CS did not show any binding to DOCK2 DHR-2 (Fig. 1E). Consistent with this finding, the association between DOCK2 DHR-2 and Rac1 was abrogated in the presence of CS, but not in the presence of cholesterol or CA (Fig. 1F). Thus, CS is a naturally occurring DOCK2 inhibitor that binds to the DHR-2 domain and blocks DOCK2 Rac GEF activity.

CS inhibits DOCK2-mediated Rac activation and cellular functions
Migration of lymphocytes and neutrophils in mice and humans critically depends on DOCK2 (14, 16, 20). To examine whether CS inhibits DOCK2 activity in cells, we first analyzed the effect of CS on lymphocyte migration. In Transwell chemotaxis assays, CS effectively inhibited the migratory response of murine T cells to the chemokine CCL21 in a concentration-dependent manner, similar to the knock-out of Dock2 (Fig. 2A). In addition, murine bone marrow (BM)–derived neutrophils treated with CS exhibited a migration defect toward the chemoattractant N-formyl-Met-Leu-Phe (fMLP) (Fig. 2B). Similar results were obtained when human peripheral blood T cells and neutrophils were analyzed in the same chemotaxis assays (fig. S2, A and B). This inhibitory effect was specific to CS because other cholesterol derivatives did not affect migration of murine lymphocytes (Fig. 2C). Consistent with these functional defects in chemotaxis, CS treatment markedly suppressed CCL21- and fMLP-induced Rac activation in murine T cells and neutrophils, respectively (Fig. 2D). We also found that neutrophils undergoing chemotaxis stopped migration at a defined distance when CS was added to the fMLP source (Fig. 2E and movie S1). However, such an effect was not observed by adding DHEAS to the fMLP source (Fig. 2E and movie S1). These results indicate that CS blocks leukocyte migration, likely by inhibiting the DOCK2-mediated Rac activation.

CS is most abundantly produced in the Harderian gland
Sulfation is catalyzed by members of the sulfotransferase (SULT) family of sulfate-conjugating enzymes (32). In both humans and mice, the SULT2B1a and SULT2B1b proteins are encoded by the same gene but differ in their N-terminal amino acid sequences as a result of alternative splicing (21, 33–35). Whereas human SULT2B1a and SULT2B1b have unique 8- and 23–amino acid sequences N-terminal to the conserved region, murine SULT2B1a and SULT2B1b have additional 54– and 20–amino acid sequences at the N terminus, respectively (Fig. 3A) (35). A study using human recombinant proteins indicated that SULT2B1b, but not SULT2B1a, plays a major role in cholesterol sulfation (36). Similarly, we found that the in vitro cholesterol sulfation activity of recombinant murine SULT2B1b was seven times higher than that of murine SULT2B1a, with the two enzymes exhibiting Vmax values of 11.2 and 1.6 nmol min−1 mg−1 and Michaelis constant (Km) values of 2.6 and 1.0 μM, respectively (Fig. 3B). The recombinant enzymes used in these assays were expressed and purified as GST fusions, and the GST tag was removed before the cholesterol sulfation assays. We obtained similar kinetic data using a murine SULT2B1b that was expressed as a His-SUMO–tagged fusion protein and analyzed after removing the His-SUMO moiety with SUMO protease 1 (fig. S3; Vmax of 12.3 nmol min−1 mg−1 and Km value of 2.2 μM), although the extra peptide remaining on the N terminus of the two differently expressed recombinant proteins differed (see Materials and Methods). In addition, in both assays, none of our SULT2B1b recombinant proteins showed substrate inhibition (37) below 15 μM cholesterol (Fig. 3B and fig. S3).

To determine which tissues produce SULT2B1b, we developed a specific antibody that can distinguish SULT2B1b from SULT2B1a in mice (fig. S4, A to C). Western blot analyses revealed that SULT2B1b was present in the small intestine and skin in C57BL/6 (designated Sult2b1+/+) mice (Fig. 3C), consistent with that previously reported for Sult2b1 mRNA expression (34). However, the highest abundance of SULT2B1b was detected in the Harderian gland (Fig. 3C), an orbital gland that produces the lipids that form the oily layer of the tear film (31). This was confirmed by a different antibody that was raised against the C-terminal sequence that is conserved between murine SULT2B1a and SULT2B1b and thus recognizes both isoforms (fig. S5, A to D). SULT2B1a was not detected in any tissues tested (fig. S5D). Endogenous cholesterol sulfating activity was much higher in the Harderian gland than in the small intestine or skin, the other tissues in which we detected SULT2B1b (Fig. 3D), and its activity correlated with the amount of SULT2B1b in each tissue (Fig. 3, C and D).

To understand the physiological function of SULT2B1b in more detail, we used knockout (KO) (Sult2b1−/−) mice that had been generated by deleting exons 3 to 7 of the Sult2b1 gene (fig. S6, A to C) (30, 38). Although the Harderian gland of Sult2b1−/− mice contained more than 500 pmol of CS per milligram of tissue, the presence of CS and cholesterol sulfation activity were hardly detected in the Harderian gland of Sult2b1−/− mice (Fig. 3, E to G), indicating that CS production
CRITICALLY depends on SULT2B1b in vivo. We also found that in mice heterozygous for the mutant allele (Sult2b1+/−), the CS content in the Harderian gland was reduced to 58% of that of wild-type controls.

**CS limits ocular surface inflammation**

Acute UV exposure causes photokeratitis, which is initiated by apoptosis of corneal epithelial cells and is followed by recruitment of leukocytes into the anterior chamber, the aqueous humor–filled space between the cornea and iris, where they come into contact with and damage the corneal endothelium (39–41). To examine whether lack of CS aggravates the severity of photokeratitis, Sult2b1+/−, and Sult2b1−/− mice were UV-irradiated under anesthesia. The thickness of the corneal epithelial layer was comparably reduced in both groups of mice 24 hours after UV irradiation (Fig. 4A), indicating that the presence or absence of CS did not affect the susceptibility of corneal cells to UV-induced apoptosis. However, inflammatory infiltrates in the anterior chamber were increased in Sult2b1−/− mice compared to in Sult2b1+/− mice (Fig. 4B). The majority of the inflammatory cells infiltrating the anterior chamber in both genotypes were Gr1+ neutrophils (fig. S8). Similarly, Sult2b1 deficiency significantly augmented the infiltration of T cells into the conjunctiva in an experimental allergic conjunctivitis (EAC) model (Fig. 4, C and D), in which mice were sensitized and topically challenged with short ragweed (SRW) pollen (42).

SULT2B1b sulfates not only cholesterol but also oxysterols such as 25-hydroxycholesterol (25HC) (35, 43–45). Although this sulfated product, 25-hydroxycholesterol-3-sulfate (25HCS) (fig. S9A), is known to induce anti-inflammatory responses (46), 25HCS did not affect...
Fig. 2. CS blocks leukocyte migration by inhibiting chemoattractant-induced Rac activation.
(A) Transwell migration assays measuring the migration of wild-type (WT) and Dock2−/− murine T cells in response to the chemoattractant CCL21 and the migration of wild-type murine T cells in response to CCL21 in the presence of the indicated concentrations of CS or vehicle only (DMSO). The percentage of T cell migration was calculated by dividing the number of T cells that migrated into the lower chamber in response to CCL21 by the number of input T cells. Data are presented as means ± SD. n = 7 experiments. *P < 0.05, **P < 0.01 (two-tailed unpaired Student’s t test).
(B) Migration of BM-derived murine neutrophils treated with CS or vehicle only (DMSO) along a 0 to 10 μM fMLP gradient over a 20-min time period. Data are representative of three independent experiments.
(C) Migration of murine T cells in response to CCL21 in the presence of the indicated steroids. Data are presented as means ± SD. n = 5 to 8 experiments per treatment. **P < 0.01 compared with control (DMSO) sample (one-way ANOVA followed by Bonferroni post hoc test).
(D) Rac activation in murine T cells and neutrophils stimulated with CCL21 and fMLP, respectively, in the presence or absence of CS. Data (means ± SD) are presented as the ratio of GTP-bound Rac to total Rac after setting the 5-s value of control (DMSO) sample to an arbitrary unit (a.u.) of 1. **P < 0.01 (two-tailed unpaired Student’s t test).
(E) Migration of BM-derived murine neutrophils toward a source of fMLP in the presence of CS, DHEAS, or vehicle only (DMSO) over a time period of 30 min. Data are representative of four independent experiments.
Fig. 3. CS is most abundantly produced in the Harderian gland by the sulfotransferase SULT2B1b. (A) Alignment of the N-terminal amino acid (a.a.) sequences of the human (h) and murine (m) SULT2B1a and SULT2B1b proteins. Conserved regions are boxed with dotted lines. (B) Cholesterol sulfation activity of recombinant murine SULT2B1b and SULT2B1a produced as GST fusion proteins. Assays were performed after the GST moiety was removed. Data (nanomole per minute per 1 mg of recombinant protein) are presented as means ± SD. n = 3 experiments. (C) Representative immunoblot showing SULT2B1b and tubulin in the indicated mouse tissues. The numbers below the blot indicate the abundance of SULT2B1b relative to that of tubulin in each tissue and normalized by setting the abundance of SULT2B1b in the Harderian gland as 1. Quantification data represent the means of three to five independent quantification measurements for each tissue type. (D) Comparison of cholesterol sulfation activity among extracts from the indicated mouse tissues. Data (picomole per minute per 1 mg of tissue lysate) are presented as means ± SD. n = 3 experiments. (E) Abundance of CS in Harderian glands from Sult2b1+/+, Sult2b1+/−, and Sult2b1−/− mice as quantified by liquid chromatography–tandem mass spectrometry (LC-MS/MS). Data (picomole per 1 mg of wet tissue) are presented as means ± SD. n = 5 glands per group. (F) Representative immunoblot showing SULT2B1b in Harderian glands from Sult2b1+/+, Sult2b1+/−, and Sult2b1−/− mice. The numbers below the blot indicate the abundance of SULT2B1b relative to actin for each tissue sample and normalized by setting the abundance of SULT2B1b in Sult2b1+/+ mice as 1. Quantification data represent the mean of three experiments for each genotype. (G) Cholesterol sulfation activity of Harderian gland extracts from Sult2b1+/+, Sult2b1+/−, and Sult2b1−/− mice. Data (picomole per minute per 1 mg of tissue lysate) are presented as means ± SD. n = 3 experiments. (H and I) The amounts of CS in various tissues (H; n = 7 to 20 tissues per group; picomole per 1 mg of wet tissue) or tears (I; n = 10 mice per group; picomole per 1 ml of tear) from Sult2b1+/+ and Sult2b1−/− mice were quantified by LC-MS/MS. Data are presented as means ± SD. **P < 0.01 (two-tailed unpaired Student’s t test). (J) Localization of CS in sections of eyeballs from Sult2b1+/+ and Sult2b1−/− mice as determined by matrix-assisted laser desorption ionization (MALDI)–imaging mass spectrometry. The color bar indicates the relative intensity of the CS signal [mass/charge ratio (m/z) 465.3]. Data are representative of three independent experiments. Scale bar, 200 μm.
**DISCUSSION**

CS has been implicated in several biological processes, yet its physiological functions are not fully understood. Here, we have identified CS as a naturally occurring DOCK2 inhibitor that binds to the catalytic DHR-2 domain and blocks its Rac GEF activity. Although CS which was suppressed by CS eye drops. In humans, CS is also a normal constituent of tears and meibum (48), an oily substance secreted by the meibomian gland. Therefore, it is likely that CS acts locally to create environments that evade immune surveillance in the eyes of both mice and humans. So far, it has been shown that DOCK2 deficiency prevents cardiac allograft rejection and autoimmune disease development in mice (18, 19). In light of these findings, CS may serve as an excellent starting place for development of DOCK2-targeting anti-inflammatory therapeutics.

**MATERIALS AND METHODS**

**Chemical compounds**

A total of 577 pharmacologically active compounds were obtained from the Drug Discovery Initiative (the University of Tokyo) and initially tested at 50 μM for their inhibitory effect on the Rac GEF activity of the DHR-2 domain of murine DOCK2. CS was purchased...
Fluorescent GEF assays were performed as described previously (49). Briefly, recombinant His-SUMO–tagged DOCK1 DHR-2 (mouse), DOCK2 DHR-2 (mouse and human), DOCK5 DHR-2 (mouse), Tiam DH-PH (mouse), and Trio DH-PH (mouse) proteins were produced in the Arctic BL21 (DE3) bacterial strain and purified by Ni-NTA DH-PH (mouse), and Trio DH-PH (mouse) proteins were produced depending on their GEF activities) in the reaction buffer [20 mM MES-NaOH, 150 mM NaCl, and 2.5 mM EDTA (pH 7.0)] for 20 min at room temperature. The samples were then mixed with GST-fusion Rac1-immobilized beads (8 μl) in a total 600 μl of the binding buffer and incubated at 4°C for 1 hour on a rotating wheel. After the beads were washed twice with the binding buffer, bound proteins were separated by SDS–polyacrylamide gel electrophoresis (PAGE) and blotted with the HisProbe HRP conjugate (1:1000 dilution).

Binding assays

SPR-based binding assays were performed with a Biacore T200 instrument (GE Healthcare Life Sciences). For this purpose, His-SUMO–tagged murine DOCK2 DHR-2 protein was immobilized onto a CM5 Sensor Chip (about 7000 RU) using an Amine Coupling Kit (GE Healthcare Life Sciences). Data were collected in Heps-buffered saline [10 mM Heps (pH 7.5) and 150 mM NaCl containing 0.005% surfactant P-20, 3% DMSO, and 0.1 mM hydroxypropyl-β-cyclodextrin (HPβCD)]. Serial concentrations of CS were injected and binding was measured.

For steroid-binding assays, CS and various steroid analogs were dissolved in methanol (50 μl of 100 μg/ml) and were immobilized onto each well of the 96-well plates (Immulon 4HBX, Thermo Fisher Scientific) by air drying under a hood for 1 hour at room temperature. After blocking with 180 μl of tris-buffered saline [20 mM tris-HCl and 150 mM NaCl (pH 7.5)] containing 5% bovine serum albumin (BSA) for 6 hours at 4°C, each sample was incubated with 0.2 μg of His-SUMO–tagged murine DOCK2 DHR-2 protein (100 μl of 2 μg/ml) for 2.5 hours at room temperature, followed by incubation with the HisProbe horseradish peroxidase (HRP) conjugate (no. 15165, Thermo Fisher Scientific; 100 μl of 1:5000 dilution) for 1.5 hours at room temperature. The bound HRP was detected by the colorimetric assay with the 3,3′, 5,5′-tetramethylbenzidine substrate solution (no. N301, Thermo Fisher Scientific) and the stop solution (no. N600, Thermo Fisher Scientific) at an absorbance of 450 nm.

To examine the effect of CS on association between DOCK2 DHR-2 and Rac1, recombinant protein encoding His-SUMO–tagged murine DOCK2 DHR-2 domain (1 μg) was incubated with CS, steroid analogs (final 50 μM), or DMSO (2%) in 200 μl of binding buffer [20 mM MES-NaOH, 150 mM NaCl, and 2.5 mM EDTA (pH 7.0)] for 20 min at room temperature. The informed consent was obtained after the nature and possible consequences of the studies were explained.

Cell preparation

Murine BM neutrophils were isolated from tibia and femur and layered onto the discontinuous Percoll (GE Healthcare) gradient. After centrifugation, cells at the 62/81% interface were recovered. Human peripheral blood samples were obtained from healthy volunteers in compliance with institutional review board protocols. Human neutrophils were prepared as described previously (50), and human CD4+ T cells were isolated from peripheral blood mononuclear cells by magnetic sorting with Dynabeads human CD4 followed by treatment with DETACHaBEAD human CD4 (Life Technologies) (51). The informed consent was obtained after the nature and possible consequences of the studies were explained.

Chemotaxis assays

Transwell chemotaxis assays were performed using CCL21 as a chemoattractant for T cells. Briefly, murine splenocytes (1 × 10^6/ml) prepared from wild-type and Dock2−/− mice (14) and human CD4+ T cells (1 × 10^6/ml) were incubated in an RPMI 1640 medium (Wako Pure Chemical Industries) containing 0.5% BSA with or without the indicated concentrations of CS, steroid analogs, or 0.2% DMSO (Transwell assay medium) at 37°C for 2 hours. Cells (1 × 10^6) for murine splenocytes and 1 × 10^6 for human CD4+ T cells) were loaded into the upper chamber of the Transwells (Costar no. 3421, 5-μm pore size), which were placed onto 24-well plates containing a Transwell assay medium supplemented with CCL21 (100 to 300 ng/ml for murine splenocytes and 600 ng/ml for human CD4+ T cells; both from R&D Systems). After incubation at 37°C for 6 hours (murine splenocytes) or 2 hours (human CD4+ T cells), cells migrating to the lower chamber were collected and stained with a phycoerythrin (PE)–conjugated antibody for murine CD90.2 (3C11, BD Pharmingen) followed by avidin-conjugated fluorescein isothiocyanate (BD Biosciences), respectively. Flow cytometric analyses were done on FACS Calibur (BD Biosciences). The percentage of migrating T cells was calculated by dividing the number of T cells in the lower chamber by the number of input T cells.

For EZ-TAXIScan chemotaxis assays, murine BM neutrophils or human peripheral blood neutrophils were incubated in an RPMI 1640 medium containing 0.1% BSA plus CS or DMSO (0.2%) for 30 min at room temperature. Cells were then allowed to migrate along an fMLP gradient (0 to 10 μM; source concentration of fMLP = 10 μM) over the 260-μm track of an EZ-TAXIScan chamber (Effector Cell
Sult2b1+/− than five generations, and age- and sex-matched backcrossed with C57BL/6 mice (purchased from Japan Clea) for more.

were probed with a monoclonal antibody for Rac1 (23A8, 1:2000 dilution; Millipore). The bound proteins and total cell lysates were separated by SDS-PAGE on a 12.5% polyacrylamide gel, and blots were probed with a monoclonal antibody for Rac1 (23A8, 1:2000 dilution; Millipore).

Mice
Sult2b1+/− mice, which were generated by using a targeting vector to replace exons 3 to 7 of the Sult2b1 gene with an angiotsin-converting enzyme–Cre–neomycin (ACN) cassette (52), have been described previously (30, 38) and were obtained from the Jackson Laboratory (stock no. 018773). Mice heterozygous for the mutant allele (Sult2b1+/−) were backcrossed with C57BI/6 mice (purchased from Japan Clea) for more than five generations, and age- and sex-matched Sult2b1+/− and Sult2b1−/− littermates were analyzed in functional assays. Mice were maintained under specific pathogen–free conditions in the animal facility of Kyushu University. The protocol of animal experiments was approved by the committee of Ethics on Animal Experiments of Kyushu University.

Genotyping polymerase chain reaction and Southern blot analysis
Genotyping polymerase chain reaction (PCR) was performed using the KOD FX DNA polymerase (TOYOBO) and the following primers: s1, 5′-GACAGGCAGGGCCACAC-3′; s2, 5′-CTATTCACACACACACCCAT-3′; and as1, 5′-TCCATCCCTAGTCATGGG-3′. Amplified products for wild-type allele and KO allele were 215 bp (s1 and as1) and 469 bp (s2 and as1), respectively. For Southern blot analysis, genomic DNA was isolated from the tail of each mouse, digested with Eco RI and Eco RV, separated on a 1% agarose gel, and transferred onto the Hybond-XL nylon membrane (GE Healthcare) by the alkaline transfer method. DNA fragments of 0.5 kb covering the exons 4 and 5 and 0.25 kb carrying a portion of the intron 2 were labeled with [α-32P]-dCTP (3000 Ci/mmole, PerkinElmer) using the Megaprime DNA labeling system (RPN1606, GE Healthcare) and were used as a probe for hybridization in the hybridization buffer [50% formamide, 5× SSPE, 5× Denhardt’s solution, 1% SDS, and salmon sperm DNA (100 μg/ml)]. After washing, the hybridized membrane was analyzed by BAS-2500 bio-imaging analyzer.

SULT assays
The complementary DNAs encoding murine SULT2B1a and SULT2B1b were cloned in the Eco RI–Xho I sites of the pGEX 6P-1 vector (GE Healthcare) and the pET-SUMO vector. GST-fusion SULT2B1a and SULT2B1b were produced in BL21 (DE3) bacterial strain and purified by Glutathione Sepharose 4B chromatography (GE Healthcare). The GST moiety was removed by cleavage with PreScission Protease (no. 27084301, GE Healthcare). His-SUMO–tagged SULT2B1b was produced in the Arctic BL21 (DE3) bacterial strain and purified by Ni-NTA affinity chromatography (Qiagen). The His-SUMO moiety was removed by cleavage with SUMO protease 1 (no. 4010, LifeSensors).

SULT activity was determined using radiolabeled cholesterol as described (36, 37) with slight modifications. For the kinetic analyses, 0.31 μg of SULT2B1a or 0.15 μg of SULT2B1b protein was used in the reaction mixture (50 μl) consisting of 0 to 20 μM [3H]-cholesterol (~1500 dpm/pmole), 0.1 mM 3′-phosphoadenosine 5′-phosphosulfate (PAPS; Sigma-Aldrich, A1651) in 10 mM tris-HCl (pH 7.5) containing 5 mM MgCl2, 0.2 mM HPβCD, and 5% ethanol (v/v). Reactions were initiated by adding 2 mM PAPS (2.5 μl), carried out at 37°C for 10 min, and terminated by adding 0.8 ml of ethanol. Samples were frozen at −20°C for 60 min, followed by centrifugation at 20,000g for 5 min at 4°C to remove precipitates. The supernatant was recovered in a fresh tube, evaporated using the SpeedVac (no. SPD1010, Thermo Fisher Scientific), redissolved in 8 μl of methanol containing 16 μg of CS as a carrier, and applied to a thin-layer chromatography (TLC) sheet (Chromato Sheet II, Wako, 038-24061), which was developed using a solvent system consisting of chloroform/acetone/methanol/acetic acid/water (8:4:2:2:1). After drying, the TLC sheet was developed by exposure to I2 vapor. CS spots were excised, and the radioactivity was determined by the liquid scintillation counter.

For determination of tissue SULT activity, tissues were isolated, suspended in 300 μl of ice-cold phosphate-buffered saline (PBS)/water (1:1) supplemented with 1 mM dithiothreitol and protease inhibitor cocktail (cOmplete, Roche), and homogenized using the POLYTRON homogenizer (PT 1200E, KINEMATICA AG), followed by centrifugation at 20,000 g for 30 min at 4°C. The supernatant was recovered and clarified using the Millex-LH 0.45-μm filter (Millipore) to obtain tissue lysates. The reaction mixture (50 μl) consisted of the tissue lysates (2 to 10 μg of protein), 10 μM [3H]-cholesterol (~2800 dpm/pmole), 0.1 mM PAPS in 10 mM tris-HCl (pH 7.5) containing 5 mM MgCl2, 0.2 mM HPβCD, and 3.5% ethanol (v/v). Reactions were initiated by adding 2 mM PAPS (2.5 μl), carried out at 37°C for 30 min, and terminated by adding 0.8 ml of ethanol. Samples were processed as described above, and the radioactivity was determined by the liquid scintillation counter.

Western blot analyses for SULT2B1b abundance
To examine the abundance of SULT2B1b in each tissue, polyclonal antibody that specifically recognizes murine SULT2B1b was produced by immunizing rabbits with keyhole limpet hemocyanin (KLH)–coupled synthetic peptide corresponding to the N-terminal sequence of SULT2B1b with additional cysteine for coupling (MDGPQQRALWSSSEKNSVSEMWWC). A polyclonal antibody that recognizes both murine SULT2B1a and SULT2B1b was produced by immunizing rabbits with a KLH-coupled synthetic peptide corresponding to the
C-terminal conserved sequence with additional cysteine for coupling (CSPSPASDDPNPG). In some experiments, human embryonic kidney 293T cells were transfected by polyethylamine with PMX–internal ribosomal entry site–green fluorescent protein vectors encoding murine SULT2B1b, its mutant lacking N-terminal 23–amino acid residues (AN) or SULT2B1a with or without C-terminal hemagglutinin (HA) tag, and their lysates were used as controls. Freshly isolated tissues were homogenized in a 15-mL polypropylene tube containing 1 mL of lysis buffer [20 mM tris–HCl (pH 7.5), 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1% Triton X-100, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, and 1 mM sodium vanadate] supplemented with 5× protease inhibitor cocktail (comple, Roche) using the homogenizer (Ultra-Turrax T8, IKA-Werke) set at dial 4 for 1 min on ice. The homogenates were transferred to 1.5-mL tubes and centrifuged at 20,000g at 4°C for 5 min. After centrifugation, the supernatants were mixed with an equal volume of 2× sample buffer [125 mM tris–HCl (pH 6.8), 4% SDS, 20% glycerol, 0.01% bromophenol blue, and 5% 2-mercaptoethanol] and boiled for 5 min. Total protein concentration was measured by the DC Protein Assay Reagent (Bio-Rad). Tissue or cell extracts (20 μg per lane) were separated by SDS-PAGE and immunoblotted with a rabbit antibody for SULT2B1b (1:1000 dilution), rabbit antibody for SULT2B1a/1b (1:1000 dilution), goat antibody for actin (I-19, 1:1000 dilution; Santa Cruz Biotechnology), rabbit monoclonal antibody for β-tubulin (9F3, 1:1000 dilution; Cell Signaling Technology), or rat monoclonal antibody for HA (3F10, 1:2000 dilution; Roche), followed by incubation with HRP-conjugated secondary antibodies.

**Quantification of CS by mass spectrometry**

For sample preparation, tissues were freshly isolated from mice, quickly frozen with liquid nitrogen, and stored at −80°C until analyses. Frozen samples were mixed with an internal standard (IS) compound (deuterium-labeled CS; d7-CS), and homogenized in ice-cold methanol (500 μL) using a homogenizer (Finger Masher AM79330; Sarstedt). The supernatant was filtered using the ultrafiltration devices (UltrafreeMC-PLHCC; Human Metabolome Technologies), and the filtrate was directly analyzed by LC-MS/MS for CS content. Tear fluid was collected by using a homogenizer (Ultra-Turrax T8, IKA-Werke) set at dial 4 for 1 min on ice. Sections were embedded in optimal cutting temperature compound (Sakura) and fixed in 4% glutaraldehyde in PBS for 1 hour at room temperature. The spectral irradiance for the UV lamps was 280 to 400 nm, and the UVB output was monitored by a Digital UV meter (280 to 320 nm, Solarmeter). After 24 hours, the eyes were harvested for histological examinations. For induction of EAC, mice were immunized with 50 μg of SRW pollen (Greer Laboratories) in 5 mg of Imject Alum Adjuvant (77161, Thermo Fisher Scientific). Samples collected at less than 0.2 μL were omitted from the analyses. The concentrates were dissolved in 15 μL of methanol and analyzed by LC-MS/MS.

For quantification of CS with LC-MS/MS, a triple-quadrupole mass spectrometer equipped with an electrospray ionization (ESI) ion source (LCMS-8040; Shimadzu Corporation) was used in the negative-ESI and multiple reaction monitoring modes. The samples were resolved on the Mastro-C18 column (2.1 mm × 100 mm, 3-μm particle, Shimadzu GLC) by isocratic flow with mobile phase A (200 mM ammonium acetate) and mobile phase B (methanol) at ratios of 1:9 and a flow rate of 0.4 mL/min, with a column temperature of 40°C. CS and IS (d7-CS) signals were monitored by ion transitions at m/z 465.3 > 97 and 472.3 > 97, respectively. The absolute content of CS was calculated by using peak area ratios of CS against IS. For quantification of 25HCS content, the signal of ion transition at m/z 481.3 > 97 was monitored, and the peak area of 25HCS was quantified and calculated relative to that of CS.

**Disease models for ocular surface inflammation**

Photokeratitis was induced as previously described (39). Briefly, mice were irradiated with UVB at 100 mJ cm−2 s−1 for 100 s using a spot light source (LIGHTNING CURE Spot light source LC8 L9588-02, Hamamatsu Photonics) under anesthesia. The spectral irradiance for the UV lamps was 280 to 400 nm, and the UVB output was monitored by a Digital UV meter (280 to 320 nm, Solarmeter). After 24 hours, the eyes were harvested for histological examinations. For induction of EAC, mice were immunized with 50 μg of SRW pollen (Greer Laboratories) in 5 mg of Imject Alum Adjuvant (77/161, Thermo Fisher Scientific) into footpads on day 0 (42). Ten days later, 1.5 mg of SRW pollen suspended in 10 μL of PBS was topically administered into each eye once a day for 3 days. On day 13, 24 hours after the last pollen application, the eyes were harvested for histological examinations.

Histological analyses were performed as follows. The eyes were fixed in 4% glutaraldehyde in PBS for 1 hour at room temperature and then fixed in 10% paraformaldehyde in PBS overnight. After being embedded in paraffin, tissues sections (5 μm thick) were cut through the papillary–optic nerve plane and were stained with hematoxylin and eosin. The number of inflammatory cells per 0.01 mm² was counted. For immunofluorescence analyses, the eyes were embedded in optimal cutting temperature compound (Sakura Finetek) and frozen in liquid nitrogen. Cryostat sections (5 μm thick) were fixed in 4% paraformaldehyde for 30 min and blocked with 1% BSA in PBS for 1 hour at room temperature. Sections were then stained with PE-conjugated antibody for murine CD3 (17A2, 4 μg/mL; BioLegend), biotinylated antibody for murine CD45R (RA3-6B2, 5 μg/mL; BD Biosciences), or biotinylated antibody for murine Gr1 (Ly6G/Ly6C) (RB6-8C5, 5 μg/mL; BD Biosciences) followed by incubation with Alexa Fluor 546–conjugated streptavidin.
CS-containing eye drops
CS was dissolved in 40 mM HβCD before use. In UV-induced photokeratitis models, 10 μL of CS (8 μg/μL) or vehicle only (40 mM HβCD) was administered topically to the eye of Sult2b1−/− mice for a total of six times: once 5 min before irradiation and five times subsequently at 4-hour intervals. In EAC models, 10 μL of CS (8 μg/μL) or vehicle (40 mM HβCD) was administered topically to the eye of Sult2b1−/− mice three times per day at 4-hour intervals on days 10, 11, and 12.

Reverse transcription PCR
Total RNA was isolated from tissue using ISOGEN (Nippon Gene). After treatment with ribonuclease-free deoxyribonuclease I (Life Technologies), RNA samples were reverse-transcribed with oligo(dt) primers (Life Technologies) and SuperScript III reverse transcriptase (Life Technologies) for amplification by PCR. Real-time PCR was performed on a CFX Connect Real Time System (Bio-Rad) using the SYBR Green PCR Master Mix (Applied Biosystems). The following PCR primers were used: for Gapdh, 5′-TGGTCCGCTGGATCCTGA-3′ and 5′-TTGCTGGTAAGTCGAGGAG-3′; and for Ch25h, the commercial validated primers qMmuCED0001765 (Bio-Rad).

Statistical analyses
Statistical analyses were performed using GraphPad Prism. The data when two groups were compared or a one-way ANOVA when two groups were compared.

**REFERENCES AND NOTES**


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Cholesterol sulfate is a DOCK2 inhibitor that mediates tissue-specific immune evasion in the eye

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Evading the immune system with a lipid

Some tissues, such as the eye, locally suppress immune cells or limit their ability to infiltrate the tissue. Sakurai et al., found that cholesterol sulfate (CS) inhibited the guanine nucleotide exchange factor DOCK2 and thus suppressed the migration of neutrophils and T cells in vitro. In mice, CS was produced by the gland that secretes the lipids that form the outer layer of the tear film covering the eye. Mice lacking the major sulfotransferase that produces CS from cholesterol exhibited increased infiltration of immune cells into the conjunctiva and the cornea in different in vivo models of ocular surface inflammation, which was reversed by topical application of CS. Identifying a role for CS in endogenous immunosuppression suggests that it or other bioactive lipids may play a role in other tissues that evade immune surveillance, such as tumors.