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1	Expression of indian hedgehog signaling in murine oviductal infundibulum and its
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#### 28 Abstract

Homeostasis of the oviductal infundibulum epithelium is continuously regulated by signaling 29pathways under physiological and pathological conditions. Herein, we investigated the expression 30 of hedgehog (Hh) signaling-related components in the murine oviductal infundibulum, which is 31 known to maintain homeostasis in the adult epithelium. Additionally, using autoimmune disease-32prone MRL/MpJ-Fas<sup>lpr/lpr</sup> (MRL/lpr) mice showing abnormal morphofunction of the ciliated 33 epithelium of the infundibulum related to the oviductal inflammation, we examined the 34relationship between Hh signaling and pathology of the infundibulum. The expression and 35 36 localization of Pax8, a marker for progenitor cells in the oviductal epithelium, and Foxj1, a marker for ciliogenesis, were examined in the infundibulum. The results showed that Pax8 was 3738downregulated and Foxil was upregulated with aging, suggesting that homeostasis of the infundibulum epithelium of MRL/lpr mice was disturbed at 6 months of age. In all mice, the motile 39 cilia of ciliated epithelial cells in the infundibulum harbored Hh signaling pathway-related 40 molecules: patched (Ptch), smoothened (Smo), and epithelial cells harbor Gli. In contrast, Ptch, 41 Smo, and Gli2 were significantly downregulated in the infundibulum of MRL/lpr mice at 6 months 42of age. The expression levels of Pax8 and Foxj1 were significantly positively correlated with those 43of Ptch1, Smo, and Gli2. Hh signaling is thought to be involved in homeostasis of the ciliated 44 epithelium in the infundibulum. In MRL/lpr mice, which show exacerbated severe systemic 45

46	autoimmune abnormalities, molecular alterations in Hh signaling-related components are
47	considered to interact with local inflammation in the infundibulum, leading to disturbances in
48	epithelial homeostasis and reproductive function.
49	

50 Keywords: autoimmune abnormality, ciliated epithelium, hedgehog signaling pathway,
51 homeostasis, oviduct

#### 52 Introduction

Mammalian oviducts are divided into three parts, the infundibulum, ampulla, and isthmus, in 53order from the distal to proximal parts. The different composition ratios of ciliated epithelial cells 54and secretory cells in the epithelium of each part reflect the unique reproductive function in each 55section (Li et al. 2017; Koyama et al. 2019). Particularly, the infundibulum epithelium is composed 5657mostly of ciliated epithelial cells, so that ciliary beating effectively moves oocytes produced in the ovaries into the oviductal lumen; this process is known as oocyte pick-up. Oocyte pick-up and 58transportation are disturbed by abnormal ciliary morphofunction in the infundibulum caused by 5960 pathological conditions such as smoking (Talbot and Riveles 2005), hormonal dysregulation (Raidt et al. 2015), and inflammation (Hosotani et al. 2020). Physiologically, the proportion of ciliated 6162epithelial cells and secretory cells in the infundibulum epithelium changes under hormonal dynamics through the estrous cycle (Ito et al. 2016). In addition, the infundibulum is constantly 63 exposed to follicular fluid containing inflammatory molecules during each ovulation, which 64 damages the ciliary epithelium (Palma-Vera et al. 2017). Therefore, the infundibulum epithelium 65continuously undergoes epithelial turnover to maintain its healthy histology and reproductive 66 function under both physiological and pathological conditions. 67

Adult epithelial homeostasis is maintained by the proliferation and differentiation of epithelial
 stem cells, which are regulated by activation of signaling pathways such as the Wnt/β-catenin,

70	Notch, and Hedgehog (Hh) signaling pathways (Sancho et al. 2004; Carlier et al. 2020). In
71	oviductal epithelial homeostasis, secretory cells act as progenitors by self-renewing and/or
72	differentiating into ciliated epithelial cells (Ghosh et al. 2017). The molecular mechanism
73	underlying these effects is not fully understood; however, several studies have focused on the
74	involvement of Wnt/β-catenin signaling (Ghosh et al. 2017) and Notch signaling (Zhu et al. 2019)
75	in homeostasis of the oviductal epithelium. In contrast, although homeostasis of the adult tracheal
76	ciliated epithelium is maintained by activation of the Hh signaling pathway (Peng et al. 2015), the
77	involvement of this pathway in maintaining the oviductal epithelium has not been explored.
78	In a murine model of systemic autoimmune disease, MRL/MpJ-Fas <sup>lpr/lpr</sup> (MRL/lpr) mice
79	develop severe inflammation of the lamina propria in the oviductal infundibulum. Chronic
80	abnormal immune conditions result in abnormal morphofunction in the ciliated epithelium, such
81	as decreased numbers of ciliated epithelial cells, elongation of the cilia, and disorientation of ciliary
82	beating (Hosotani et al. 2018, 2020). These pathological conditions are closely related to
83	disturbances in epithelial homeostasis but the underlying molecular mechanism is unclear.
84	In this study, we investigated expression of the Hh signaling pathway in the murine
85	infundibulum epithelium and its relationship with homeostasis of the infundibulum epithelium. In
86	addition, we examined the expression of Hh signaling pathway-related molecules in MRL/lpr mice
87	as a destruction model of the infundibulum epithelium and compared the results with those

- obtained the healthy infundibulum of C57BL/6N mice (B6) as a general strain and MRL/MpJ mice
- 89 (MRL/+) as wild-type MRL/lpr mice.

## 90 Material and methods

### 91 Animals

Animal experiments were approved by the School of Veterinary Medicine, Rakuno Gakuen 92University (approval no. VH19A6). The animals were handled in accordance with the Guide for 93the Care and Use of Laboratory Animals, Rakuno Gakuen University, Japan. Female B6, MRL/+, 9495and MRL/lpr mice at 3 and 6 months of age were obtained from Japan SLC, Inc. (Hamamatsu, Shizuoka, Japan). Previous studies reported that autoimmune disease is severely exacerbated in 96 female MRL/lpr mice at 6 months of age compared to at 3 months of age (Hosotani et al. 2018, 97 2020). The mice were housed in groups within plastic cages at 18–26°C under a 12-h light/dark 98 cycle and had free access to a commercial diet and water. The estrous cycle of each mouse under 99 100 the natural estrous cycle was confirmed by monitoring vaginal smears (Byers et al. 2012). All mice were euthanized by either severing the carotid artery or cervical dislocation under deep anesthesia 101 using a mixture of medetomidine (0.3 mg/kg), midazolam (4 mg/kg), and butorphanol (5 mg/kg). 102The spleen was collected from the mice and weighed as a marker of autoimmune disease. 103

104

#### 105 *Immunostaining*

Mouse female reproductive organs were collected and fixed with 4% paraformaldehyde at 4°C
overnight, embedded in paraffin, and cut into 3-µm-thick sections, which were then used for

108	immunohistochemistry (IHC) and immunofluorescence (IF). Detailed information on the
109	antibodies, antigen retrieval, and serum blocking is listed in Table 1. The sections were incubated
110	in 20 mM Tris-HCl (pH 9.0) for 15 min at 110°C, 10 mM citrate buffer (pH 6.0) for 15 min at
111	110°C, or 0.1% pepsin for 5 min at 37°C. The sections for IHC were soaked in methanol containing
112	0.3% hydrogen peroxide. Sections incubated with blocking serum for 60 min at room temperature
113	were incubated overnight at 4°C with primary antibodies. Negative controls were performed with
114	normal mouse IgG (sc-2025, Santa Cruz Biotechnology Inc., Santa Cruz, CA, USA), normal rat
115	IgG (sc-2026, Santa Cruz Biotechnology) and normal rabbit IgG (sc-2027, Santa Cruz
116	Biotechnology). After three washes in 0.01 M PBS, the sections were incubated with secondary
117	antibodies for 30 min and washed. The sections for IHC were incubated for 30 min at room
118	temperature, using a streptavidin-biotin complex (SABPRO Kit, Nichirei, Tokyo, Japan), and then
119	incubated with 3,3'-diaminobenzidine tetrahydrochloride-hydrogen peroxide solution, and lightly
120	stained with hematoxylin. The stained sections of the outer infundibulum were examined using a
121	BZ-X710 microscope (Keyence, Osaka, Japan).
122	Three semi-serial sections stained with IHC for detection of Foxj1 with 50- $\mu$ m intervals were
123	used, and the percentage of expelled non-ciliated epithelial cells and the percentage of Foxj1
124	positive expelled non-ciliated epithelial cells were calculated as follows, respectively: percentage

125 of expelled non-ciliated epithelial cells (%) =  $100 \times$  number of expelled non-ciliated epithelial

130	Reverse Transcription and Quantitative Real-time Polymerase Chain Reaction
129	
128	cells / number of total 10-20 expelled non-ciliated epithelial cells in the field.
127	non-ciliated epithelial cells (%) = $100 \times$ number of Foxj1 positive expelled non-ciliated epithelial
126	cells / number of total 80-100 epithelial cells in the field. Percentage of Foxj1 positive expelled

131The oviducts were manually separated into the proximal (including the isthmus and ampulla) and distal (including infundibulum) parts and homogenized using a BioMasher (Nippi Inc., Tokyo, 132Japan). Total RNA was purified using the NucleoSpin® RNA Plus kit (Macherey-Nagel, Düren, 133134Germany) according to the manufacturer's instructions. The purified total RNA was used as a template to synthesize cDNA using ReverTra Ace qPCR RT Master Mix (Toyobo Co., Ltd., Osaka, 135136Japan). Quantitative real-time polymerase chain reaction (qPCR) analysis of the cDNA was performed using THUNDERBIRD SYBR qPCR Mix (Toyobo Co., Ltd.) and gene-specific 137primers (Table 2, Sigma-Aldrich, St. Louis, MO, USA). The qPCR cycling conditions were 95°C 138for 1 min, followed by 40 cycles of 95°C for 15 s and 52°C or 60°C for 45 s. Data were normalized 139against the expression level of actin, beta (*Actb*) and analyzed using the  $\Delta$ Ct method to compare 140 141 the expression of genes encoding hedgehog ligands; the  $\Delta\Delta$ Ct method was used to compare the expression of other genes. 142

143

# 144 In Situ Hybridization

145	Paraformaldehyde-fixed paraffin-embedded specimens of female reproductive organs were
146	cut into 5µm-thick sections, air-dried overnight, and baked in an oven for 60 min at 60°C. RNA in
147	situ hybridization was performed using an RNAscope® 2.5 HD Detection Regent-Brown kit
148	(Advanced Cell Diagnostics, Newark, CA, USA) according to the manufacturer's instructions. The
149	RNAscope® positive control probe-Mm-Polr2a (Cat. No. 312471, Advanced Cell Diagnostics),
150	RNAscope® negative control probe-DapB (Cat. No. 310043, Advanced Cell Diagnostics),
151	RNAscope® probe-Mm-Shh (Cat. No. 314361, Advanced Cell Diagnostics), RNAscope® probe-
152	Mm-Dhh (Cat. No. 415031, Advanced Cell Diagnostics), and RNAscope® probe-Mm-Ihh-noXHs
153	(Cat. No. 413091, Advanced Cell Diagnostics) were used.

154

## 155 Statistical Analysis

The results are expressed as the mean  $\pm$  standard error (s.e.). Data among three or more groups were compared using Tukey's test (P < 0.05). Data between two groups were compared using Student's *t*-test (P < 0.05). Correlations between two parameters were analyzed using Spearman's correlation test (P < 0.05). Statistical analysis was conducted using JMP 14.2.0 (SAS Institute, Inc., Cary, NC, USA). 161 **Result** 

162	Expression and Localization of Epithelial Homeostasis-Related Molecules in the Infundibulum
163	In the oviductal epithelium, Pax8 is a marker of secretory cells (i.e., progenitor cells), whereas
164	Foxj1 is a marker of ciliogenesis. Pax8 was localized in the nucleus of secretory cells in the
165	infundibulum of mice, except for in MRL/lpr mice at 6 months of age (Figure 1a-c and a'-c').
166	Foxj1 was localized in the nucleus of ciliated epithelial cells in the infundibulum of mice, except
167	for MRL/lpr mice at 6 months of age, in which Foxj1 expression was observed not only in the
168	nucleus of ciliated epithelial cells as well as that of non-ciliated epithelial cells expelled from the
169	epithelium (Figure 1d-f and d'-f'). There was no significant differences in the percentage of
170	expelled non-ciliated epithelial cells composing the infundibulum epithelium among the strains
171	and ages (Figure 1h), while the percentage of Foxj1 positive expelled non-ciliated epithelial cells
172	was significantly higher in MRL/lpr mice at 6 months of age than other strains and MRL/lpr mice
173	at 3 months of age (Figure 1i).
174	We compared the transcriptional expression levels of <i>Pax8</i> and <i>Foxj1</i> in the infundibulum as
175	the distal part of the oviduct with those in the ampulla and isthmus as the proximal part (Figure 2a

- and b). *Pax8* expression was lower in the distal part than in the proximal part of all strains at 3
- 177 months of age. In the entire oviduct of MRL/lpr mice at 6 months of age, *Pax8* expression showed
- an age-related decrease and was significantly lower than that in the other strains. In contrast, *Foxj1*

179	expression was higher in the distal part than in the proximal part of all mice. In the entire oviduct
180	of MRL/lpr, Foxj1 expression was higher than in other strains, particularly at the distal part which
181	showed a significant age-related increase.
182	
183	Expression and Localization of Hedgehog Signaling-Related Molecules in Infundibulum
184	The canonical Hh signaling pathway in primary cilia is activated by binding of Hh ligands such
185	as sonic, indian, and desert Hh (Shh, Ihh, and Dhh, respectively) to the transmembrane protein
186	patched (Ptch), followed by an interaction with smoothened (Smo) and the activation of GLI
187	family zinc finger 1-3 (Gli1-3) transcription factors (Briscoe and Thérond 2013). There were no
188	significant differences in the expression of Hh signaling pathway-related genes during the estrous
189	cycle (Figure 4a-e); therefore, we focused on the infundibulum of mice at estrus. In all mice, Ptch1
190	and Smo were localized in the cilia of ciliated epithelial cells in the infundibulum (Figure 3a-a""",
191	b-b"", c-c"", d-d"", e-e"", and f-f""). The fluorescence intensity of Ptch1 and Smo in the
192	infundibulum did not differ between mice of different strains and ages. In B6 and MRL/+ mice,
193	Gli2 was highly expressed in the nucleus of secretory cells and faintly in the nucleus of ciliated
194	epithelial cells in the infundibulum (Figure 3g-g"", h-h""). In MRL/lpr mice at both 3 and 6
195	months of age, Gli2 was ubiquitously observed in the nuclei of cells comprising the infundibulum
196	epithelium (Figure 3i-i"").

197	In transcriptional analysis, <i>Ptch1</i> expression did not significantly differ among the oviductal
198	parts of all mice (Figure 5a). Ptch1 expression showed an age-related decrease in the proximal part
199	of B6 and at both the proximal and distal parts in MRL/lpr mice. In the entire oviduct of MRL/lpr
200	mice, Ptch1 expression was significantly lower than that in B6 or/and MRL/+ mice. Smo
201	expression was significantly higher in the distal part than in the proximal part of B6 and MRL/+
202	mice at 3 months of age (Figure 5b). Smo expression showed an age-related decrease at the distal
203	part in all mice. In the entire oviduct of MRL/lpr, Smo expression was significantly lower than that
204	in the other strains. Gli1 expression was significantly lower at the distal part than at the proximal
205	part: Gli2 and Gli3 expression showed no significant differences among the oviductal parts (Figure
206	5c-e). In the entire oviduct of MRL/lpr, Gli1 expression showed an age-related decrease and was
207	significantly lower than that in the other strains (Figure 5c). The mean Ct value of Gli1 expression
208	in the distal parts was high above around 32 in all mice (data not shown), so that the localization
209	level of Gli1 protein seems to be very low in the distal part. In the entire oviduct of MRL/lpr, Gli2
210	expression was significantly higher than in other strains, particularly at the distal part at 6 months
211	of age (Figure 5d). <i>Gli3</i> expression tended to be higher in the entire oviduct of MRL/lpr mice at 3
212	months of age than in other strains, whereas that of MRL/lpr at 6 months of age showed an age-
213	related decrease and was significantly lower than that in other strains (Figure 5e).

Among the transcription factors of the hedgehog ligands (*Shh*, *Dhh*, and *Ihh*), *Shh* and *Dhh* 

215	were not expressed in the infundibulum epithelium or ovarian granulosa cells (Figure 6a-f and a'-
216	f'), whereas Ihh was highly expressed in ovarian granulosa cells and slightly in infundibulum
217	epithelial cells (Figure 6a"-f"). Ihh expression in ovarian granulosa cells was lower in MRL/lpr
218	mice than in B6 mice (Figure 6a" and e"). The Shh and Dhh expression levels were not
219	significantly different among the oviductal parts, whereas Ihh expression was lower in the distal
220	part than in the proximal part in all mice (Figure 7a-f).
221	
222	Relation between Hh Signaling Pathway and Oviductal Epithelial Homeostasis
223	Correlation analysis of transcriptional expression in the oviduct of mice at 3 months of age
224	was performed (Table 3). In the proximal part, Ptch1 and Pax8 expression was significantly
225	positively correlated with the expression of all Hh signaling pathway-related genes. Foxj1
226	expression showed the same results, except for Gli1. In contrast, at the distal part, Ptch1 expression
227	was significantly positively correlated with the expression of Smo and Gli2 and significantly
228	negatively correlated with <i>Gli3</i> expression. <i>Pax8</i> and <i>Foxj1</i> expression was significantly positively
229	correlated with the expression of <i>Ptch1</i> , <i>Smo</i> , and <i>Gli2</i> .

#### 230 **Discussion**

The molecular mechanism that maintains homeostasis of the infundibulum epithelium has been 231greatly altered in autoimmune disease model mice. This molecular alteration causes an abnormal 232morphology of the ciliated epithelium of the infundibulum of MRL/lpr mice at 6 months of age, 233such as a decreased proportion of ciliated epithelial cells, elongated cilia, and disorientation of 234235ciliary alignment (Hosotani et al. 2020). The transcription factor Pax8 governs the expression of a series of genes pivotal to tissue development in thyroid follicular cells and Müllerian ducts 236(Plachov et al. 1990; Grote et al. 2006). Pax8 in the oviductal secretory cells is a marker of the 237238self-renewal and differentiation of these cells to ciliated epithelial cells (Ghosh et al. 2017); therefore, the significant downregulation of Pax8 transcripts and proteins suggests that Pax8 239240attenuates epithelial homeostasis in the infundibulum of MRL/lpr at 6 months of age. The transcription factor Foxil is required for the late steps of ciliogenesis, including docking of 241centrioles at the apical membrane to form basal bodies and axoneme elongation (You et al. 2004). 242Foxil upregulation promotes abnormal ciliogenesis, resulting in disorganized, dense, and 243lengthened cilia in the airway ciliated cells of patients with chronic mucosal inflammation (Li et 244al. 2014). Considering that the direction of coordinated ciliary beating requires basal body 245polarization (Kunimoto et al. 2012), in the infundibulum of MRL/lpr mice at 6 months of age, 246significant upregulation of Foxi1 may promote abnormal ciliogenesis, resulting in cilia elongation 247

248	and disoriented ciliary alignment. Foxj1 localization in the expelled non-ciliated epithelial cells
249	was significantly observed in the infundibulum of MRL/lpr mice at 6 months of age, which implies
250	the increase of inadequate ciliogenesis and the promoted elimination of epithelial cells failed to
251	differentiate to ciliated epithelial cells. Otherwise, significantly high expression of Foxj1 may have
252	been a reaction to the decreased number of ciliated epithelial cells following disturbed ciliogenesis
253	and loss of $Pax8^+$ epithelial progenitor cells. The abnormal morphofunction of the ciliated
254	epithelium of the infundibulum caused by alterations in homeostasis-related molecules causes
255	dysfunction in oocyte pick-up (Hosotani et al. 2018, 2020).
256	This study revealed that the motile cilia of ciliated epithelial cells in the infundibulum harbor
257	Hh signaling pathway-related molecules: the transmembrane receptors, Ptch and transmembrane
258	protein adjacent to Ptch, and Smo (Carpenter et al. 1998). In addition, Hh signaling effectors in
259	the Gli family are expressed in the nucleus of infundibulum epithelial cells. Considering the
260	significant positive correlations between Ptch1 and Smo/Gli2 in the infundibulum, motile cilia in
261	the infundibulum epithelium contain the Hh signaling pathway, although whether this pathway is
262	canonical or non-canonical matter has not been determined. Although primary cilia (i.e., immotile
263	cilia) are generally considered as sensor cilia in which canonical/non-canonical Hh signaling is

transduced (Abou Alaiwi et al. 2009; Bangs and Anderson 2017), recent studies reported that 

motile cilia also act as sensors of the pericellular environment. The motile cilia of the tracheal 

266	epithelium express sensing receptors such as the bitter taste receptor and Ptch1/Smo (Shah et al.
267	2009; Nordgren et al. 2014), which sense injury and chronic inflammation in the airway. The
268	motile cilia in oviducts harbor progesterone receptors, which regulate the ciliary beat frequency
269	(Teilmann et al. 2006; Bylander et al. 2010). Thus, motile cilia in the infundibulum epithelium
270	may play a sensing role via Hh signaling-related components. Hh ligands were not significantly
271	expressed in the infundibulum, and the present and previous studies (Russell et al. 2007) revealed
272	Ihh expression in ovarian granulosa cells, where Hh signaling may induce granulosa cell
273	proliferation. Therefore, cilia in the infundibulum may receive Ihh produced by ovarian granulosa
274	cells at the time of ovulation, rather than through the paracrine effect of Hh produced by
275	infundibulum epithelial cells.
276	In the infundibulum, Pax8 and Foxj1 expression showed a significant positive correlation with
277	the gene expression of Hh signaling pathway-related Ptch1, Smo, and Gli2. This suggests that the
278	Hh signaling pathway is closely related to the regulation of epithelial homeostasis in the

279 infundibulum. To the best of our knowledge, direct molecular interactions between Pax8 and/or

Foxj1 and the Hh signaling pathway during development and homeostasis maintenance have not

been observed previously. However, Pax8 has been predicted as a target gene activated via the Shh

signaling pathway in oncogenesis (Harter et al. 2015). During the development of central nervous

system, Foxj1 expression is regulated by the Shh signaling pathway and alters the response of cells

284	to Shh signaling (Cruz et al. 2010). Although further studies are needed to determine the molecular
285	relationship between Pax8, Foxj1, and Hh signaling-related components, the significant
286	downregulation of the Ptch1 and Smo transcripts in the infundibulum of MRL/lpr mice at 6 months
287	of age may be related to the abnormal pathology of the ciliated epithelium. The alternation of Gli2
288	localization in the infundibulum of MRL/lpr mice compared to other strains would relate to the
289	downregulation of Ptch1 and Smo transcripts and the disruption of Hh signaling
290	Aged MRL/lpr mice show autoimmune disease-prone phenotypes in systemic organs, including
291	the oviductal infundibulum with infiltration of autoreactive immune cells because of a mutation in
292	the Fas cell surface death receptor (Fas) gene (Andrews et al. 1978; Watanabe-Fukunaga et al.
293	1992; Hosotani et al. 2020). MRL/lpr at 3 months of age showed expression of Ptch1, Smo, and
294	Gli1-3 in the infundibulum, and thus specific genetic alterations in the MRL/lpr strain do not cause
295	significant downregulation of Hh signaling-related components. In contrast, alterations in Hh
296	signaling transduction were reported to trigger immunological modulation. In the skin of atopic
297	dermatitis mouse models and the central nervous system with autoimmune neuroinflammation,
298	Shh signaling suppresses immune reactions via T-regulatory cell signaling and reduces tissue
299	pathology and disease severity (Papaioannou et al. 2019; Benallegue et al. 2021). In adult intestinal
300	mesenchyme, Shh and Ihh act as anti-inflammatory epithelial modulators and modulate
301	tolerogenic versus proinflammatory signaling (Zacharias et al. 2010). Therefore, significant

302	downregulation of <i>Ptch1</i> and <i>Smo</i> transcripts may interact with severe autoimmune inflammation
303	in the oviductal lamina propria of MRL/lpr mice at 6 months of age (Hosotani et al. 2020).
304	Furthermore, the deteriorated supply of fresh Hh signaling-related components in the
305	infundibulum of MRL/lpr mice at 6 months of age may be related to abnormal homeostasis of the
306	ciliated epithelium. The damages on ciliary morphofunction caused by variations in levels of
307	inflammatory cytokines in the infundibulum of MRL/lpr mice (Hosotani et al. 2020) are thought
308	to reflect the physiological damage of ciliated epithelium of infundibulum caused by inflammatory
309	follicular fluid (Palma-Vera et al. 2017), in terms of inflammatory cytokines altering the
310	morphofunction of the ciliated epithelium of infundibulum. Therefore, the downregulation of Hh
311	signaling-related components caused by the inflammatory molecules would also relate to the
312	turnover of the infundibulum epithelium under physiological conditions.
313	We observed a difference in the molecular expression between the infundibulum and
314	ampulla/isthmus. Epithelial cells in the distal and proximal parts of the oviducts are from
315	intrinsically different lineages and are maintained separately (Ford et al. 2020). The properties and
316	populations of Pax8 <sup>+</sup> epithelial cells differ between the distal and proximal parts; the infundibulum
317	shows a low population of $Pax8^+$ secretory cells (Ford et al. 2020). Foxj1 upregulation in the
318	infundibulum compared to in the proximal part reflects greater activation of ciliogenesis and a
319	larger population of ciliated epithelial cells in the infundibulum. Hh signaling-related components

were transcribed in the proximal part, despite the sparse presence of cilia. In contrast, correlation analysis indicated that Gli1 is involved in Hh signaling in the proximal part and that Gli1 and Gli3 are involved in epithelial homeostasis. Further investigations are required to reveal the effects of the difference in molecular expression patterns among oviductal parts.

In summary, we propose the different mechanism of Hh signaling transduction between in 324325healthy and autoimmune disease conditions (Figure 8a and b). We propose that homeostasis of ciliated epithelium in the infundibulum is regulated not only by Wnt/β-catenin and Notch signaling, 326327but also possibly by Hh signaling. In addition, alterations in Hh signal transduction related to the abnormal immune condition in the oviduct of MRL/lpr disrupts the epithelial morphology in the 328infundibulum, resulting in oocyte pick-up dysfunction. Further functional study on the change of 329330 the epithelial morphology in infundibulum under the experimental manipulation of Hh signaling would strengthen the role of Hh signaling in the epithelial homeostasis in oviducts. Our results 331improve the understanding of the physiology and pathology of mammalian female reproductive 332function. 333

# **Author contributions**

335	Conceptualization: M.H. and O.I.; Methodology: M.H., O.I., Ta.N., M.A.M, Te.N. and Y. H.;
336	Validation: M.H., O.I., and T.N.; formal analysis: M.H. and O.I.; Investigation: M.H., O.I., Ta.N.
337	and M.A.M.; Resources: M.H., O.I., and M.A.M.; Data curation: M.H.; Writing - original draft:
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339	O.I.; Supervision: M.H. and Y.K.; Project administration: O.I. and YK; Funding acquisition: M.H.
340	
341	Conflicts of interest
342	The authors declare no conflicts of interest.
343	
344	Ethical approval
345	Animal experimentation was approved by the School of Veterinary Medicine, Rakuno Gakuen
346	University (approval no. VH19A6). Animals were handled in accordance with the Guide for the
347	Care and Use of Laboratory Animals, Rakuno Gakuen University, Japan.
348	
349	Data, material, and/or code availability
350	The data that support the findings of this study are available from the corresponding author
351	upon reasonable request.

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### 451 **FIGURE LEGENDS**

Fig. 1 Localization of Pax8 and Foxj1 in the oviductal infundibulum. IHC of Pax8 (a-c and a'-c'), 452Foxi1 (d-f and d'-f'), and negative control (g) are shown. Bar = 25  $\mu$ m. Black arrows: 453immunoreactive positive epithelial cells; red arrowheads: expelled non-ciliated epithelial cells 454from the epithelium. (h) The percentage of expelled non-ciliated epithelial cells from the 455456epithelium in distal parts of oviducts. There were no significant differences between ages at the same strains (Student's t-test, P < 0.05) and among strains at the same age (Tukey's test, P < 0.05). 457(i) The percentage of Foxil positive expelled non-ciliated epithelial cells from the epithelium distal 458459parts of oviducts. \*: Significant differences between 3 and 6 months of age at the same strains and same part (Student's *t*-test, P < 0.05). B&M: Significant differences between B6 or MRL/+ mice 460 461at the same age (Tukey's test, P < 0.05). B6 = C57BL/6N, MRL/+ = MRL/MpJ, MRL/lpr = MRL/MpJ-Fas<sup>lpr/lpr</sup> 462

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**Fig. 2** Expression levels of *Pax8* (a) and *Foxj1* (b) in the oviduct. Data are the mean  $\pm$  s.e. **\***: Significant differences between 3 and 6 months of age at the same strains and same part (Student's *t*-test, *P* < 0.05). *#*: Significant differences between proximal and distal parts of oviducts from mice of the same strains and age (Student's *t*-test, *P* < 0.05). B&M: Significant differences between B6 or MRL/+ mice at the same age (Tukey's test, *P* < 0.05). *Pax8*: Paired box 8; *Foxj1*: Forkhead box 469 protein J1. B6 = C57BL/6N, MRL/+ = MRL/MpJ, MRL/lpr = MRL/MpJ-Fas<sup>lpr/lpr</sup>
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471 **Fig. 3** Localization of hedgehog signaling-related proteins in the oviductal infundibulum revealed 472 by immunofluorescence. Localization of Ptch1 (a-a"", b-b"", and c-c""), Smo (d-d"", e-e"", 473 and f-f""), Gli2 (g-g"", h-h"", and i-i""), and negative control (j-j") are shown. Bar = 25  $\mu$ m. 474 Arrows: immunoreactive positive cells in the epithelium. B6 = C57BL/6N, MRL/+ = MRL/MpJ, 475 MRL/lpr = MRL/MpJ-*Fas<sup>lpr/lpr</sup>* 

476

477Fig. 4 Expression levels during estrous cycle of hedgehog signaling-related genes in the oviduct of C57BL/6N at 3 months of age. Expression levels of Ptch1 (a), Smo (b), Gli1 (c), Gli2 (d), and 478479*Gli3* (e) are shown. Data are the mean  $\pm$  s.e. \*: Significant differences between 3 and 6 months of age at the same cycle and same part (Student's *t*-test, P < 0.05). #: Significant differences between 480proximal and distal parts of oviducts from mice of the same cycle and age (Student's t-test, P < 4810.05). There were no significant differences among estrous cycles at the same age and part 482(Tukey's test, P < 0.05). P: proximal part; D: distal part 3m: 3 months of age, 6m: 6 months of age. 483Ptch1: Patched-1, Smo: Smoothened, Gli1-3: GLI family zinc finger 1-3. 484

485

486 Fig. 5 Expression levels of hedgehog signaling-related genes in the oviduct. Expression levels of

487 *Ptch1* (a), *Smo* (b), *Gli1* (c), *Gli2* (d), and *Gli3* (e) are shown. Data are the mean  $\pm$  s.e. \*: Significant 488 differences between 3 and 6 months of age for the same strains and same part (Student's *t*-test, *P* 489 < 0.05). #: Significant differences between proximal and distal parts of oviducts from mice of the 490 same strains and age (Student's *t*-test, *P* < 0.05). B&M: Significant differences between B6 or 491 MRL/+ mice at the same age (Tukey's test, *P* < 0.05). *Ptch1*: Patched-1, *Smo*: Smoothened, *Gli1*-492 *3*: GLI family zinc finger 1-3. B6 = C57BL/6N, MRL/+ = MRL/MpJ, MRL/lpr = MRL/MpJ-493 *Fas<sup>lpr/lpr</sup>* 

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495 **Fig. 6** Expression of transcriptions of hedgehog in ovarian granulosa cells (a-a", c-c", and e-e") 496 and epithelial cells of the oviductal infundibulum (b-b", d-d", and f-f") revealed by *in situ* 497 hybridization. Negative controls are shown in (g and g'). Brown dots indicate positive reactions to 498 *in situ* hybridization. Bar = 25  $\mu$ m. Arrows: reaction-positive cells in the oviductal infundibulum. 499 B6 = C57BL/6N; MRL/lpr = MRL/MpJ-*Fas<sup>lpr/lpr</sup>* 

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Fig. 7 Expression levels of hedgehog genes in the oviduct of B6 at 3 months of age (a and d), MRL/+ at 3 months of age (b and e), and MRL/lpr at 3 months of age (c and f). Data are the mean  $\pm$  s.e. \*: Significant differences between proximal and distal parts of oviducts from mice of the same strains and age (Student's *t*-test, *P* < 0.05). 3: Significant differences between 3 and 6 months of age in mice of the same strains and from the same part (Student's *t*-test, P < 0.05). B&M: Significant differences between B6 or MRL/+ mice at the same age (Tukey's test, P < 0.05). B6 = C57BL/6N, MRL/+ = MRL/MpJ, MRL/lpr = MRL/MpJ-*Fas<sup>lpr/lpr</sup>* 

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Fig. 8 Estimated modulation mechanisms of hedgehog signaling in the oviductal infundibulum 509510between healthy (a and a') and autoimmune disease conditions (b and b'). In healthy condition, hedgehog signaling transduced by Ihh released by the ovarian granulosa cells, which is received 511by Ptch1 and Smo localized on the motile cilia of the oviductal infundibulum. The hedgehog 512513signaling mediated by Ptch1, Smo, and Gli2 in the ciliated epithelial cells would lead the constant transcription and protein production of Ptch1 and Smo and maintain the proliferation of Pax8 514positive secretory cells and differentiation of those into Foxj1 positive healthy ciliated epithelial 515cells in the oviductal infundibulum. In autoimmune disease condition, the declined levels of Ihh 516production and transcription of Ptch1 and Smo causes the modulation of hedgehog signaling 517transduction, which resulted in the disturbances in the homeostasis of ciliated epithelium of the 518oviductal infundibulum and the appearance of strongly Foxil positive ciliated epithelial cells 519which are morphologically abnormal. CEC: ciliated epithelial cell, SC: secretory cell, NCEC: non 520ciliated epithelial cell, GCs: granulosa cells, Foxj1: forkhead box J1, Pax8: paired box 8, Hh: 521Hedgehog, Ihh: indian hedgehog, Ptch1: Patched-1, Smo: Smoothened, Gli2: GLI family zinc 522

523 finger 2.

Anti gen	Cat. No	Sourc e	Ho st	Dilu tion	Antigen retrieval reagent	Blocking serum	Secondary antibody for IF (1:300)	Biotinylated secondary antibody for IHC
Gli2	NBP2 - 23602	Novu s Biolo gicals LLC., Littlet on, CO, USA	Ra bbi t	1:20 0	20 mM tris- HCl (pH 9.0)	10% goat normal serum	Goat anti-rabbit IgG H&L CF568, 20103-1 (Biotium.Inc, Hayward, CA)	-
Ptch 1	MAB 41051	R&D Syste ms Inc., Minn eapoli s, MN, USA	Ra t	1:20 0	0.1% pepsin	10% goat normal serum	Goat anti-rat IgG H&L CF568, 20096-1 (Biotium.Inc, Hayward, CA)	-
Smo	GTX 60154	Gene Tex Inc., Irvine , CA, USA	Ra bbi t	1:20 0	0.1% pepsin	10% goat normal serum	Goat anti-rabbit IgG H&L CF568, 20103-1 (Biotium.Inc, Hayward, CA)	-
Tub ulin- α	MS- 581- R7	Ther mo Scient ific, Walth am,	M ou se	Und ilute d	No need	10% goat normal serum	Goat anti-mouse IgG H&L CF488A, 20018-1 (Biotium.Inc, Hayward, CA)	_

# **Table 1. Primary antibody information in immunostaining.**

		MA,					
		USA					
Foxj 1	14- 9965- 80	Invitr ogen, Carls bad, CA, USA	M ou se	1:50 0	10 mM citrate buffer (pH 6.0)	10% rabbit normal - serum	Rabbit anti-mouse IgG+IgA+IgM antibody, 426031, undiluted (Nichirei, Tokyo, Japan)
Pax 8	ACR 438A	Bioca re Medi cal, Pache co, CA, USA	M ou se	1:10 0	20 mM tris- HCl (pH 9.0)	10% rabbit normal - serum	Rabbit anti-mouse IgG+IgA+IgM antibody, 426031, undiluted (Nichirei, Tokyo, Japan)

525 IF: immunofluorescence, IHC: immunohistochemistry.

Genes	Accession Number	Primer Sequence (5'-3') F: Forward, R:	Product size (bp)	Annealing temp. (°C)	
		Reverse			
Actb	NR 007202 5	F: TGTTACCAACTGGGACGACA	175	(0	
	NM_007393.5	R: GGGGTGTTGAAGGTCTCAAA	165	60	
Dhh	NM 007857 5	F: TTGGCACTCTTGGCACTATCT	277	60	
Dnn	1111_007837.3	R: CTTTGCAACGCTCTGTCATC	211	00	
Foril	NM_008240.3	F: ACTATGCCACCAACCCACA	171	60	
1'03/1		R: GGATGGAATTCTGCCAGGT			
Clil	NM_010296.2	F: CGACCTGCAAACCGTAATC	280	60	
011		R: CTTGCCAACCATCATATCCA	269	00	
		F:			
Gli2	NM_001081125.1	TGGAGAAGAAAGAAGCCAAGAG	159	60	
		R: TCATGTCAATCGGCAAAGG			
Gli3	NIM 009120.2	F: CCTGCTCCAACATTTCCAAC	240		
	INIVI_008130.3	R: CTTGACTAGGGTTGTTCCTTCC	240	00	

# **Table 2. Primer list used for quantitative PCR analysis.**

F: CCTCTTGCCTACAAGCAGTTC

Ihh	NM_010544.3	R· AGATGGCCAGTGAGTTCAGAC	214	60
	NM_011040.4	F: AAGCATCGACTCACAGAGCA		
Pax8		R: GAATGAGGATCTGCCACCAC	285	60
Dtahl	NM_001328514.1	F: CCATGACAAAGCCGACTACA	293	60
		R: GGAAGACTGCGCACACTAGAA		
Shh	NM 000170 3	F: CAAGTACGGCATGCTGGCTC	254	52
Snn	NM_009170.3	R: AAGGTGAGGAAGTCGCTGTA	234	52
Smo	NIM 176006 4	F: GTCTCTGCACGCTCTTCACA	260	60
	181M_1/0990.4	R: CCAGACTACTCCAGCCATCAA	207	00

527 Dhh: desert hedgehog, Foxj1: forkhead box J1, Gli1-3: GLI family zinc finger 1–3, Ihh: Indian

hedgehog, *Pax8*: paired box 8, *Ptch1*: Patched 1, *Shh*: sonic hedgehog, *Smo*: smoothened, frizzled

529 class receptor.

Table 3. Spearman's correlation coefficient (ρ) between hedgehog signaling-related genes
expression and secretory/ciliated epithelial cells marker expression in the oviduct of mice at
3 months of age.

		Parameters								
		Hedgehog	Secretory and ciliated		Hedgehog	Secretory and ciliated				
		receptor gene	epithelial cells marker		receptor gene	epithelial cells marker				
		expression	expression		expression	expression				
		at proximal part			at distal part					
		Ptchl	Pax8	Foxjl	Ptch1	Pax8	Foxj1			
Ptchl	ρ	1	0.55***	0.38**	1	0.63***	0.62***			
	Р	-	<0.0001	0.0071	-	<0.0001	<0.0001			
Smo	ρ	0.60***	0.62***	0.42**	0.48***	0.48***	0.59***			
	Р	<0.0001	<0.0001	0.0026	0.0007	0.0007	<0.0001			
Glil	ρ	0.63***	0.49***	0.20	0.19	0.13	0.23			
	Р	<0.0001	0.0005	0.1690	0.1981	0.3862	0.1226			
Gli2	ρ	0.56***	0.72***	0.69***	0.43**	0.61***	0.69***			

	Р	<0.0001	<0.0001	<0.0001	0.0027	<0.0001	<0.0001
Gli3	ρ	0.41**	0.39**	0.35*	-0.30*	-0.09	0.07
	Р	0.0038	0.0059	0.0141	0.0431	0.5554	0.6623

534 Ptch1, Patched-1; Smo, Smoothened; Gli1-3, GLI family zinc finger 1-3, Pax8, paired box 8;

*Foxj1*, forkhead box J1.













MRL/lpr at 3 months of age

B6 at 3 months of age

Shh



Dhh

lhh

g q

Negative control



ΔC<sub>t</sub> vs Actb

