A transvaginal endoscopy-based technique for performing ovarian examinations in sows

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Abstract. For examining pig ovaries, which have complex structures, laparoscopy is a useful technique, but requires general anesthesia; therefore, it cannot be performed repeatedly within a short period of time. We report a transvaginal endoscopy-based technique for conducting ovarian examinations without general anesthesia. Sows were sedated in pig stalls. Using a colonoscope, the vaginal wall was punctured with a trocar. To avoid the trocar being caught in the broad ligament of the uterus or the connective tissue around the vagina, the trocar was inserted close to the external uterine os and between the 2:00 and 3:00 or the 9:00 and 10:00 positions (in a clockwise direction). Then, a urethroscope was inserted into the abdomen, and an examination was carried out after the ovaries had been moved towards the urethroscope camera via rectal palpation. This less invasive procedure may allow repeated examinations and will increase our understanding of ovarian dynamics in pigs.

Key words: Ovarian observation, Sows, Transvaginal endoscopy

Evaluations of ovarian status are important for facilitating effective reproductive management in domestic animals. In pigs, high reproductive potential is characterized by a short open period and a large litter size. These characteristics are related in the structure of the pig ovary complex; for example, each ovary contains high numbers of follicles and/or corpus lutea (CL), which change in size rapidly. It is difficult to assess their numbers and states, and there are few tools for examining complex ovarian structures. Such technical limitations have adversely affected our knowledge of pig reproductive physiology.

As in other domestic animals, rectal palpation can be used to estimate the presence and numbers of follicles and CL; however, rectal palpation in pigs requires specific skills because of the difficulties caused by the long uterus horn and small pelvic cavity in pigs. Moreover, it is difficult to count the exact numbers of ovarian structures in pigs only by palpation. Therefore, ultrasonography (USG) is the first-choice method for examining the morphology of the ovary and can be used to monitor ovarian dynamics in a non-invasive manner [1, 2].

Ultrasonography can be used for measuring the sizes of follicles and CL and to facilitate ovum pick-up (OPU), as well as for measuring blood flow by using Doppler echography. However, USG cannot provide live color images, which can help scientists deal with more leading-edge technologies. For example, laparoscopy, which can provide live images, has also been used for studies of ovarian physiology [3, 4], diagnosis of reproductive disorders [5], and development of new reproductive technology (embryo transfer [6], oviductal insemination [7], OPU [8] and embryo collection from the reproductive tract [9]) in pigs.

In pigs, laparoscopy is performed by placing the animal on its back under general anesthesia; therefore, appropriate equipment, such as surgical beds and inhalation anesthesia apparatus, is required. Moreover, laparoscopy cannot be repeated within a short period since general anesthesia can cause severe stress and might result in alterations in the pig’s physical functions. Thus, the value of laparoscopy would increase markedly if laparoscopic examinations could be performed in the standing position and without general anesthesia.

In cows, OPU, which is a technique used to retrieve oocytes from live cows, has been performed using transvaginal ultrasound scanning [10]. This transvaginal approach is usually conducted in the standing position under caudal epidural anesthesia or mild sedation, which makes repeated examinations of ovarian structures possible. Ferguson et al. reported that this approach can be performed in pigs for OPU [11]. Less invasive and repeatable laparoscopy
can contribute to revealing ovarian dynamics more precisely such as neoangiogenesis or ovulation. Here, we report a transvaginal endoscopy-based technique for performing ovarian examinations in the standing position.

The present study was conducted according to the institutional guidelines for animal experiments of Rakuno Gakuen University (approval No. VH25C10). We used a total of 8 sows (body weight [mean ± SD]: 221.7 ± 14.4 kg) in three studies. The first study aimed to develop instruments and protocols for use under inhalation anesthesia. The second study was designed to confirm that the developed procedure could be performed under sedation in the standing position. The third study examined the practicability of performing repeated examinations using the developed protocol.

In study 1, which aimed to develop instruments and protocols especially for use with a rigid urethroscope (A2942A; Olympus, Tokyo, Japan), we used 2 Large White sows (body weight: 203 and 227 kg). Stainless steel outer guides and trocars were prepared for use during the transvaginal procedure (see Methods, Fig. 1), i.e., to puncture the vaginal wall from the vaginal cavity to the abdominal cavity. During standard laparoscopic examinations, pigs are placed in the supine position, and the abdominal cavity is insufflated with gas to broaden the viewing field and facilitate examination. However, this approach requires general anesthesia [3] and normally requires a special gas delivery device [12]. Instead, an inner guide was developed for our procedure to prevent the camera from coming into contact with the tissue. Pipes of two different lengths (43.5 and 44.0 cm, respectively) were prepared. The lengths were determined to be 0.5 and 1 cm longer than the effective length (43.0 cm, Fig. 1A) of the urethroscope. The sows were kept under inhalation anesthesia (described in Methods) and placed in the prone position on a surgical air bed. For the examinations, a rigid urethroscope was inserted together with an inner guide pipe into the abdominal cavity. Then, the operator grasped the ovaries in their hand via the rectum and drew them into the visual field of the scope (Fig. 2, Supplemental movie 1: online only). The procedure took about 30 min from the induction of anesthesia. The 43.5-cm inner guide provided a wider and larger viewing field than the 44-cm inner guide (Figs. 1B and C). At the end of each operation, the sows were euthanized, and the vaginal wall puncture site was examined anatomically.

In study 2, we used 2 hybrid sows (Large White × Landrace, body weight: 196 and 232 kg, parity: 5 and 7, estrus cycle: 21–22 days) to examine the practicability of using the developed protocol to perform repeated examinations in a maintaining stall (190 × 58 × 103 cm) under sedation. Two examinations were conducted in each sow with a 2- or 3-day interval between them, 8 experiments in total. In some trials, the trocar got trapped in the broad ligament of the uterus (see trials 4–6 in Table 1), and puncturing of the vaginal wall failed. During detailed examinations of the vaginal wall puncture sites performed in trials 1–6, we found that the trocar became trapped when it was inserted immediately lateral to or below the external os of the uterus (between the 3:00 and 5:00 or 7:00 and 9:00 positions in a clockwise direction from the external os). When the puncture site was located too caudal from the uterine os, the trocar sometimes became caught in the connective tissue around the vagina. Based on the results of the trials in study 3, we concluded that the vaginal wall can be punctured smoothly by inserting the trocar near to the external os of the uterus and between the 2:00 and 3:00 or 9:00 and 10:00 positions (in a clockwise direction) (Fig. 4A). It took about 5–10 min to puncture the vaginal wall. The success rate was 75% in trials 7–10 with this improved method, although it was 25% in trials 3–6. Throughout the studies including 2 trials with luteal phase sows in study 1, the ovarian status such as whether they were in the luteal or follicular phase did not affect the success of the puncture. In sow WL5, the vaginal wall could not be penetrated, even using this
Fig. 2. Transvaginal endoscopic examination procedure and images of the ovaries obtained in experiment 1. The operator grasped one of the ovaries through the rectum (A), and a ureteroscope was inserted transvaginally to view the ovaries (B, C). CL: corpus luteum, FL: follicle, CA: corpus albicans.

Fig. 3. Physical immobilization of the sow in a stall and transvaginal endoscopic examination of the ovaries. Sows under sedation were gently kept in position in a maintaining stall using a hard surgical mattress, and a lashing belt was used to sling up the sows to prevent them from lying down on the floor (A, B). Examinations were performed in an operation stall for large animals (C).
invasive and less traumatic way. Increasing our understanding of ovarian dynamics in sows in a less follicular and luteal samples. This procedure might contribute to repeated examinations to be conducted and can be used to collect ovarian sampling have been established yet, this procedure allows sedation in the standing position. Although no methods for transvaginal endoscopic examinations of the ovaries in sows under anesthesia, as described above.

**Methods**

In experiment 1, the sows were sedated via the intramuscular injection of a mixture of 40 μg/kg body weight (BW) medetomidine (Dorbene vet; Kyoritsu Seiyaku, Tokyo, Japan) and 0.2 mg/kg BW midazolam (Dormicum; Astellas Pharma, Tokyo, Japan), and anesthesia was induced with an intravenous injection of 6 mg/kg BW propofol (1% propofol injection for animals; Mylan Seiyaku, Tokyo, Japan). Then, the sows were orotracheally intubated and maintained at a surgical depth of anesthesia with 2–4% sevoflurane (SevoFlo; DS Pharma Animal Health, Osaka, Japan) in oxygen.

In experiments 2 and 3, the sows were sedated via the intramuscular injection of mafoprazine mesylate at a dose of 0.5 mg/kg BW (Mafopran; DS Pharma Animal Health), followed by the intramuscular injection of a mixture of 40 μg/kg BW medetomidine and 0.2 mg/kg midazolam. To avoid peristalsis and contraction of the rectum and uterus, epidural anesthesia was induced with 3 ml of 2% lidocaine (xylocaine; Astra-Zeneca, Osaka, Japan) or the intravenous injection of 1 ml of butylscopolamine bromide (20 mg/ml Buscopan; Boehringer Ingelheim International GmbH, Germany). In one sow in experiment 2, the operation time was prolonged with supplemental anesthesia, which was induced via the intramuscular injection of 2.5 mg/kg BW sodium pentobarbital (Somnopenyl; Kyoritsu Seiyaku). After each operation in experiment 3, the sows were awakened using 0.2 mg/ml atipamezole (Atipame, Kyoritsu Seiyaku). Euthanasia was achieved with an overdose of sodium pentobarbital under anesthesia, as described above.

**Transvaginal endoscopic examination procedure**

The sows were placed in pig stalls. After removing any feces, the pigs’ external genitalia were washed and cleaned. The sows were

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**Table 1. Results of the transvaginal endoscopy examinations**

| Trials | Sow ID | Ovarian status | Puncturing of the vaginal wall | Ovaries observed |
|--------|--------|----------------|------------------------------|------------------|
| Study 2 | 1 WL6   | FL (Day -1)    | Succeeded                    | Covered/Exposed  |
|        | 2 WL3297 | FL (Day -1)    | Succeeded *                  | Covered/Exposed  |
| Study 3 | 3 WL3258 | FL (Day 0)     | Succeeded                    | Covered/Exposed  |
|        | 4 WL3258 | FL (Day 1)     | Failed                       | –                |
|        | 5 WL3   | CL (Day 10)    | Failed                       | –                |
|        | 6 WL5   | CL (Day 12)    | Failed                       | –                |
|        | 7 WL4   | FL (Day -5)    | Succeeded                    | Covered/Exposed  |
|        | 8 WL3   | CL (Day 13)    | Succeeded                    | Exposed          |
|        | 9 WL5   | CL (Day 15)    | Failed                       | –                |
|        | 10 WL4  | FL (Day -3)    | Succeeded                    | Covered          |

Ovarian status: The first day of estrus was set as Day 0. CL: Luteal phase, FL: Follicular phase. * Puncturing of the vaginal wall was successful after several failures. Covered: the ovaries were covered by the mesosalpinx and fimbria ovarica. Exposed: the ovaries were observed directly.

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improved method. Necropsy showed that the vagina was torsioned near to its os.

After puncturing the vaginal wall, a transvaginal ovarian examination was successfully performed under sedation in the standing position using the procedure described in Fig. 4. The vaginal wall injuries caused by the instruments healed within 2 or 3 days. The ovaries were sometimes almost entirely covered by the mesosalpinx and fimbria ovarica (Table 1, Supplemental movie 2: online only). Of the 6 trials conducted in studies 2 and 3, the ovaries could only be directly observed in a single case (trial 8), and they were covered by the mesosalpinx and fimbria ovarica in the other 5 cases. In 4 of the 5 latter cases, the operator was able to manipulate the mesosalpinx and fimbria ovarica, then expose the ovaries by hand through the rectum. Although it is not clear when and how the mesosalpinx and fimbria ovarica covers the ovaries, it seems to often occur in the follicular phase. In such cases, instruments such as alligator forceps might be helpful for removing the mesosalpinx and fimbria ovarica. In the present study, we also attempted to obtain ovarian follicular liquid or luteal tissue using custom-made needles (22 G/62 cm for follicles, 18 G/60 cm for CL; Misawa Medical Industry, Saitama, Japan) (Supplemental movie 3: online only). In each sampling procedure, although follicles or CL were punctured at 2–4 sites, the ovarian dynamics and follicular development proceeded according to normal cycles. Three to 6 days after the final operation, all the sows were euthanized and checked anatomically. There were few puncture scars on the ovaries and no evidence of adhesion. The puncture scars in the vaginal wall were also too small to identify, and there was no evidence of inflammation or adhesion.

In summary, we have developed a procedure for performing transvaginal endoscopic examinations of the ovaries in sows under sedation in the standing position. Although no methods for transvaginal ovarian sampling have been established yet, this procedure allows repeated examinations to be conducted and can be used to collect follicular and luteal samples. This procedure might contribute to increasing our understanding of ovarian dynamics in sows in a less invasive and less traumatic way.
then sedated as described above (in the Animal sedation, anesthesia, and euthanasia [experiment 3] section of the Material and methods). To prevent the sows from lying down on the floor, i.e., to keep them standing, hard mattresses were placed around their bodies (Figs. 3A and B). A colonoscope was inserted into the vagina, and the vaginal wall was wiped with sterile gauze and iodized cotton (0.5% povidone iodine). Local anesthesia was induced using 2% lidocaine jelly (2% xylocaine jelly, Astra-Zeneca, Osaka, Japan), and an outer guide was placed at the side of the external os of the uterus. The edge of the outer guide was pressed against the vaginal wall between the 2:00 and 3:00 or the 9:00 and 10:00 positions (in a clockwise direction) (shaded areas). A trocar was inserted through the outer guide, and a puncture was made through the vaginal wall and into the abdominal cavity (B). The trocar was removed, and an inner guide was inserted into the abdominal cavity along with the outer guide (C). The outer guide was removed and a urethroscope was inserted through the inner guide, and then an examination was performed (D).

![Diagram](image.png)

Fig. 4. Procedure for the transvaginal endoscopic examinations. The edge of the outer guide was placed on one side of the external os of the uterus. The puncture site (A) was located close to the external os of the uterus and between the 2:00 and 3:00 or the 9:00 and 10:00 positions (in a clockwise direction) (shaded areas). A trocar was inserted through the outer guide, and a puncture was made through the vaginal wall and into the abdominal cavity (B). The trocar was removed, and an inner guide was inserted into the abdominal cavity along with the outer guide (C). The outer guide was removed and a urethroscope was inserted through the inner guide, and then an examination was performed (D).
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References

1. Soede NM, Noordhuizen JP, Kemp B. The duration of ovulation in pigs, studied by transrectal ultrasonography, is not related to early embryonic diversity. Theriogenology 1992; 38: 653–666. [Medline] [CrossRef]
2. Schwarz T, Murawski M, Wierczko E, Bartlewski P. An ultrasonographic study of ovarian antral follicular dynamics in prepubertal gilts during the expected activation of the hypothalmo-pituitary-ovarian axis. J Reprod Dev 2013; 59: 409–414. [Medline] [CrossRef]
3. Wildt DE, Fujimoto S, Spencer JL, Dukelow WR. Direct ovarian observation in the pig by means of laparoscopy. J Reprod Fertil 1973; 35: 541–543. [Medline] [CrossRef]
4. Brüssow KP, Ratky J, Kanitz W, Becker F. Determination of the duration of ovulation in gilts by means of laparoscopy. Reprod Domest Anim 1990; 25: 184–190. [CrossRef]
5. Wildt DE, Morcom CB, Dukelow WR. Laparoscopic pregnancy diagnosis and uterine fluid recovery in swine. J Reprod Fertil 1975; 44: 301–304. [Medline] [CrossRef]
6. Hazleger W, Kemp B. Recent developments in pig embryo transfer. Theriogenology 2001; 56: 1321–1331. [Medline] [CrossRef]
7. Morcom CB, Dukelow WR. A research technique for the oviductal insemination of pigs using laparoscopy. Lab Anim Sci 1980; 30: 1030–1031. [Medline]
8. Brüssow KP, Ratky J. Repeated laparoscopical follicular puncture and oocyte aspiration in swine. Reprod Domest Anim 1994; 29: 494–502. [CrossRef]
9. Brüssow KP, Ratky J. Endoscopic collection of porcine embryos. Reprod Domest Anim 1996; 31: 711–715. [CrossRef]
10. Pieterse MC, Vos PL, Kruip TA, Willemsen AH, Taverne MA. Characteristics of bovine estrous cycles during repeated transvaginal, ultrasound-guided oocyte aspiration (TUGA) in the sow. J Vet Med Sci 2013; 75: 191–194. [Medline] [CrossRef]
11. Ferguson E, Bellovs S, Lemieux F, Godke R. Development of a chute to facilitate transvaginal ultrasound-guided oocyte aspiration (TUGA) in the sow. J Vet Med Sci 2013; 75: 191–194. [Medline] [CrossRef]
12. Diermack P, Van Dorsselaer T, Torp KD, Schaeffer R, Geny B. Calibrated pneumoperitoneal venting to prevent N2O accumulation in the CO2 pneumoperitoneum during laparoscopy with inhaled anesthesia: an experimental study in pigs. Anesth Analg 2002; 94: 1014–1018. [Medline] [CrossRef]