The fetus's perineal area has not traditionally been the focus of the prenatal echoscopy, especially because adequate visualization depends on fetus's position. However, it is precisely thanks to the targeted attention to the perineal area that we sometimes manage to find not only quite curious but also clinically significant findings.

Sacrococcygeal teratoma can grow large and affect the natural birth process. When localized in the perineal region, it sometimes leads to defecation and micturition disorders. In addition, intestinal loops may be located in the tumor structure, which leads to deformation and the risk of hernia formation. In the presence of a giant sacrococcygeal teratoma, the following events may occur: heart failure leading to fetal hydrops, obstruction of the urinary tract, obstruction of the gastrointestinal tract, impaired innervation of organs due to compression, intratumoral hemorrhage, intrauterine anemia.

Hymenal pathology is quite rare in everyday clinical practice. Usually, it is one of the two most common nosologies: hymenal polyp or imperforate hymen. These two anomalies occur almost exclusively in childhood or immediately after menarche.

At a later age, the preserved or dysmorphic hymen can present difficulties in diagnosing and treating intravaginal pathological processes. In this article, we would like to bring to your attention two clinical cases with different prognosis to draw colleagues' attention to the need for a targeted assessment of the perineal area and correct prenatal counseling for a couple, because the ranking of the severity of the probable consequences is significant. Thus, some findings will require only adequate monitoring in childhood, others — may be the cause of premature birth, changes in delivery tactics or immediate surgery after birth.

**Key words:** prenatal diagnosis, sacrococcygeal teratoma, hymen polyp.
Perhaps the first question of future parents who came for prenatal ultrasound examination sounds familiar: “and who is there with us?”. But we must admit that this particular zone should become a zone of echoscopic interest, especially when visualization is not perfect due to fetal position. Thanks to the focused attention to the perineal area, we sometimes manage to find not only quite curious but clinically significant findings. We would like to present some of them to your attention in this article.

Case number 1.
A woman with a physiologic pregnancy was referred to our clinic at 32–33 weeks of gestation. Ultrasound investigation in the first and second trimesters found no pathological fetal changes. In the third trimester, a single-chamber formation with anechoic homogeneous contents, without blood flow, sized 17*15 mm was visualized in the perineal region between the anus and the labia (the genitals were developed according to the female type). This mass was not anatomically associated with either the bladder or the fetal spine (Fig. 1). Preliminary diagnosis: neoplasm in the perineal region, cystic teratoma?

The girl was born at 39–40 weeks via planned cesarean section. At three weeks old, an MRI was done; the conclusion: the formation of soft tissues of the perineum (cystic teratoma?). At the age of 3 months, two operations were performed sequentially: I – radical removal of the cystic duplication of the rectum; II – the imposition of a loop sigmoidostomy.

In the records of the «National Cancer Institute» noted: «mature cystic teratoid tumor». The child requires careful medical supervision, since small clusters of cell groups positive for NSE, S100 receptors (markers correlating with potential tumor recurrence) were determined in the tumor. At the age of 8 months, the III colostomy closure operation was performed. At the moment the girl is 2.5 years old and in general healthy.

Review of the literature
Teratoma (teratoma; Greek teras, terat [os] monster, deformity + -yma; synonyms: complex tumor, embryoma, mixed teratogenic tumor, tridermoma, monodermoma, parasitic fetus) – a tumor consisting of several types of tissues derived from one or two or three germ layers, the presence of which is not characteristic of those organs and anatomical regions of the body in which it develops [1]. According to modern concepts, teratoma belongs to the group of germ cell neoplasms. Germ cell tumors develop from a pluripotent, highly specialized germ cell epithelium of the gonads, capable of undergoing somatic trophoblastic differentiation and being a histogenetic source of tumors of various structures, for example, testicular seminoma, ovarian dysgerminoma, embryonic cancer, choriocarcinoma, polyembryomas, and also teratomas neoplasms (tumors of more than one histological type) [2].

According to the literature, teratoma accounts for approximately one third of all tumors in childhood and 7% of tumors in adults [3]. Teratoma is more common in women than in men. At the same time, in women, teratoma is usually benign, while the male sex is associated with tumor malignancy in a greater number of cases. In children aged 4 months to 5 years, teratomas are malignant in 50–60% of cases.

The causes of teratomas are not fully established. The most common hypothesis is based on the phenomenon of «embryo in embryo», that is, a violation of the formation of one of the «fused» twins, as a result of which one of them becomes a «parasitic» embryonic mass. Another hypothesis is the primary impairment of embryo formation, in which the embryonic layers are displaced in the early stages of embryogenesis. The reason for this may be, among other things, chromosomal abnormalities that arise in the germ cells even before fertilization [2].

Like other germ cell tumors, teratoma can be primarily localized in the testes and ovaries, and also be located extragonadally.

The appearance of a tumor outside the gonads is explained by the delay of the germ cell epithelium on the way of its migration from the yolk sac wall to the site of the gonad initiation at 4–5 weeks of embryonic development [1, 3].

The histological structure distinguishes between mature teratoma, immature teratoma, teratoma with malignant transformation [2, 4]. A mature teratoma consists of several mature, well-differentiated tissues derived from one, two, or three germ layers; it can be solid and cystic in structure. A mature teratoma of a solid structure (benign teratoma) is a dense tumor of various sizes, with a smooth or bumpy surface. On the cut, it is heterogeneous, represented in places by a heavy, whitish-gray tissue containing foci of cartilaginous and bone density, small cysts filled with transparent liquid or mucus [5]. A mature teratoma of a cystic structure is a tumor formation, as a rule, of large size, with a smooth surface. On the cut, the tumor is formed by one or more cysts filled with a cloudy gray-yellow liquid, mucus or mushy, greasy contents, in the lumen of the cysts there may be hair, teeth, fragments of cartilage.

Microscopically mature teratomas of solid and cystic structure do not differ significantly from each other. They consist of fibrous connective tissue, in which areas of well-differentiated mature stratified squamous epithelium, intestinal and respiratory epithelium, forming organoid structures, randomly alternate. Often there are structures of peripheral nerves, apocrine glands, bones, cartilage, teeth, brain tissue, adipose tissue, smooth muscles. Less commonly, tissue of the salivary gland, pancreas, adrenal gland, kidney, lung, breast can be found in the tumor. The vast majority of mature cystic teratomas are dermoid cysts [4].

A mature teratoma is a benign tumor and, as a rule, does not give metastases, although there are isolated reports of tumor implantation in the peritoneum with a ruptured ovarian teratoma.

An immature teratoma consists of immature tissues derived from all three germ layers, which resemble the embryo’s tissues during organogenesis. The size of the tumor varies widely. It has an unevenly doughy consistency, grayish-white on the cut, with small cysts and mucous areas [5]. Microscopically, foci of proliferation of immature intestinal, respiratory, stratified squamous epithelium, immature striated muscles, cartilage are determined in the tumor. The presence of tissues of neuroectodermal origin in an immature teratoma is very characteristic. Among the immature elements of the embryonic type, there are tissue sites of a mature teratoma. It is generally accepted that an immature teratoma is a potentially malignant tumor [5, 6]. Foci of malignant germ cell tumors in mature and immature teratomas, even with their small volume, largely determine the prognosis of the disease.
Teratoma with malignant transformation is an extremely rare form of tumor. Its peculiarity lies in the appearance in a teratoma of a malignant tumor of the so-called adult type, such as squamous cell carcinoma, adenocarcinoma or melanoma. For example, cases of squamous cell carcinoma developed in a dermoid cyst have been described [1, 3].

The clinical picture is determined mainly by the localization of the teratoma. Children more often than adults have their extragonadal forms. The localization of teratoid tumors in children is diverse:
- sacrococcygeal region – 38%,
- ovaries – 31%,
- retroperitoneal space – 12%,
- testicles – 6%,
- mediastinum – 4%,
- others – 9% [2].

So, extragonadal teratomas are often observed in girls, mainly in the sacrococcygeal region.

Sacrococcygeal teratoma can grow large and affect the normal birth process. When localized in the perineal region, sacrococcygeal teratoma sometimes leads to a violation of the act of defecation and urination. In addition, intestinal loops may be located in the tumor structure, which leads to deformation and the risk of hernia formation.

In the presence of a giant sacrococcygeal teratoma, the following complications may occur: heart failure, leading to fetal hydrops, obstruction of the urinary tract, obstruction of the gastrointestinal tract, violation of the innervation of organs due to compression, intratumoral hemorrhage, intrauterine anemia.

The mortality rate among such fetuses is high, and averages 50%. The most common cause of death is heart failure associated with volumetric overload of the heart, preterm labor due to polyhydramnios, bleeding after teratoma rupture, hypoxemia associated with anemia. In case of obstruction urinary tract tumor, oligohydramnios may occur, which leads to lung hypoplasia [5, 6].

As a rule, mediastinal teratoma, is localized in the anterior mediastinum; as it grows, it can protrude into one or another pleural cavity or the posterior mediastinum. For a long time, both sacrococcygeal and mediastinal teratomas may not manifest clinically and are detected by chance during X-ray examination.

Retroperitoneal teratoma occurs mainly in children and often manifests itself in the same way as nephroblastoma or retroperitoneal neuroblastoma.

Treatment of mature and immature teratomas is operative. For teratomas combined with other malignant germ cell tumors, as well as for teratomas with malignant transformation, complex treatment is used. It includes surgical removal of the tumor, the use of anticancer drugs and radiation therapy [6].

The prognosis is determined by the histological structure’s variant, the primary localization of the tumor, and timely and adequate treatment. The prognosis for teratoma, combined with embryonic cancer and chorionepithelioma, is the most unfavorable.

**Case №2.**

A woman with a physiologic pregnancy was referred to our clinic at 32–33 weeks of gestation. Ultrasound investigation in the first and second trimesters found no pathological fetal changes. In the third trimester, a small echogenic mass with a diameter of 2 mm was visualized in the area of the labia of the fetus (female genital organs) (Figure 2 in 2D mode), located closer to the posterior commissure of the vulvar ring (Figure 3 in 3D mode). No other fetal malformations were found. The girl was born naturally at 39–40 weeks. A polypoid formation was visualized in the labial area (Figure 4). Presumptive diagnosis: hymenal polyp. Observation by a gynecologist was recommended.
Review of the literature

The hymen is a fold of the mucous membrane located between the vestibule and the vaginal canal; it has a connective tissue base containing muscle fibers, nerves and blood vessels.

The hymen is formed at the end of the canal process of the vaginal plate in the fifth month of intrauterine development, and separates the canalized vagina from the urogenital sinus.

Hymenal pathology is quite rare in everyday clinical practice. Usually, it is one of the two most common nosologies: hymenal polyp or imperforate hymen. These two anomalies occur almost exclusively in childhood or immediately after menarche. At a later age, the preserved or dysmorphic hymen can present difficulties in diagnosing and treating intravaginal pathological processes.

Under the influence of maternal estrogens, the hymen of a newborn girl at birth and in the first days after it is thickened and edematous, and in some cases a round or oblong polyp is found in the back of it. Hymen polyp is a benign process. Hymenal polyps are not uncommon during neonatal and early childhood. Berglan and Selander found this anomaly in 6% of the 1000 newborn girls examined. Their size is usually less than 5 mm and is probably due to estrogen stimulation during the prenatal period. Most polyps disappear by age 3 [7].

CONCLUSION

In this article, we aimed to draw the attention of colleagues to the need for a targeted assessment of the perineal area in any trimester and correct prenatal counseling for a couple as the ultrasound findings in the area are not so common but could be very important during the birth process and further in life.

REFERENCES

1. «Тератоми различной локализации у детей», Н.Ш.Загашева, Н.А.Дуйков, Ф.М.Хуррамов, А.А.Рахматулаев, Д.С.Нурматов, 2016.
2. Дурнов Л.А, Опухоли у детей, М., 1982; Патологоанатомическая диагностика опухолей человека /Под ред. Н.А.Краевского и др. – М., 1982 – С. 236.
3. Журнал: SonoAce Ultrasound №27, Рубрика: Эхография в акушерстве, Мазырко М.А.
8. Medicalplanet.su/gynecology/no- voorbrzovania_pоловix_organov_novogo_roDENNIX.html