Imaging features of uterine and ovarian fibromatosis in Nevoid Basal Cell Carcinoma Syndrome

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ABSTRACT

Gorlin-Goltz Syndrome also known as Nevoid Basal Cell Carcinoma Syndrome is an autosomal dominant multisystem disorder. It is characterized by basal cell carcinomas, odontogenic keratocysts, skeletal abnormalities and in a minority of female patients bilateral calcified ovarian fibromas. It is challenging to radiologically assess ovarian fibromas as they have similar imaging patterns to some malignant ovarian lesions. However, it is vitally important to differentiate between benign and malignant lesions to determine patients’ suitability for fertility-sparing surgery. This report describes a case of a 25 year-old patient with Gorlin-Goltz Syndrome and bilateral ovarian fibromas.

CASE REPORT

A 25-year-old Caucasian woman underwent gynecological examination for irregular menses. Blood tests were normal apart from a slightly raised prolactin (56 ng/ml; normal ranges 0-23 ng/ml).

Transrectal ultrasound imaging (US) had highlighted the presence of several uterine and ovarian hypoechoic masses; the largest one between uterus and bladder (Figure 1), measuring 65 x 62 x 59 mm, was adherent to anterior uterine wall, with heterogenous vascularization. Left ovary was appreciable with difficulty due to the presence of a rounded hypoechoic mass measuring 18 x 15 mm. Ovarian follicles were not depictable bilaterally. These findings required further evaluation with Magnetic Resonance Imaging (MRI) of the pelvis.

MRI detected multiple round-shaped lesions in both ovaries and uterus, the largest one in the left adnexa (67 x 73 x 55 mm); all the lesions had a similar appearance with a heterogenous signal on the T2-weighted sequences, mild diffusion restriction on diffusion weighted sequence and intense heterogenous contrast-enhancement after Gadolinium administration (Figure 2, Figure 3, Figure 4, Figure 5). The patient also had adjacent lymphadenopathy and intraperitoneal fluid. These findings were suspicious of malignant lesions and a staging computed tomography (CT) examination was performed.

CT confirmed the presence of the numerous uterine and ovarian masses, some of which contained calcific foci, but no additional lesions were found in either the thorax or abdomen. A possible differential diagnosis was multiple fibromatosis of the uterus and ovaries, even though some of ovarian masses could not be differentiated radiologically from fibrothecomas or ovarian germ-cell tumors (Figure 6).

Upon further questioning, the patient revealed a history of multiple basal cell carcinomas (BCCs), the last one had been excised on the right thigh 4 years prior to presentation, and of an odontogenic cyst, surgically removed 9 years prior to presentation. The patient underwent laparoscopic fertility-sparing surgery of the ovarian and uterine masses (Figure 7).
Histological analysis confirmed small aggregates of fibroblast cells without cytologic atypies, consistent with the diagnosis of NBCCS given the patient’s medical history (Figure 8).

**DISCUSSION**

**Etiology & Demographics:**

Gorlin-Goltz Syndrome also known as Nevoid Basal Cell Carcinoma Syndrome (NBCCS) is a rare multisystem disease. It is caused by mutations in the PTCH1 gene and is autosomal dominant with complete penetrance and variable expressivity. This syndrome has several names, but we will use NBCCS, as suggested by Professor Gorlin [1]. Patients and their relatives often use the name “Gorlin Syndrome”, as it does not contain the word “carcinoma”, even if approximately 50 % of patients will develop skin cancers [2].

The estimated prevalence of NBCCS varies from 1/57,000 to 1/256,000 and it commonly appears in adolescence. There is no difference in the prevalence between males and females; however, those affected are predominantly Caucasians [3].

**Clinical & Imaging Findings:**

NBCCS is associated with basal cell carcinomas, odontogenic keratocysts (that generally develop during first to third decades), palmar and/or plantar pits and ectopic calcifications of the falx cerebri [2].

This syndrome is a clinical and radiological diagnosis with the presence of 2 major criteria and 1 minor criterion or 1 major and 3 minor criteria, as suggested by the first classification of Evans et al in 1993 [4], recently updated by the same authors in March 2018 [5] (Table 1), who highlighted that no study has been able to assess which combination of diagnostic criteria represents the best trade-off between sensitivity and specificity. If clinical features are inconclusive, to establish the diagnosis, the identification of heterozygous germine PTCH1 is necessary.

Early diagnosis of NBCCS is crucial due to the risk of developing malignancies such as medulloblastoma and aggressive skin cancers. Medulloblastoma is screened for in all patients with NBCCS as early diagnosis can improve outcomes [1].

Clinical history of our patient revealed the presence of 2 major criteria (basal cell carcinomas and odontogenic keratocysts) and 1 minor criteria (ovarian fibromas).

It is difficult to know the frequency of ovarian fibromas in NBCCS as they do not present unless they become multiple, large, calcified or twist on their pedicles. A population-based study, performed in the early 90s, suggests a frequency of 25%. Ovarian fibromas associated with NBCCS are most often bilateral (75%), calcified and nodular, often overlapping medially, eventually erroneously diagnosed as calcified uterine leiomyomas [6]. Ovarian fibromas not associated with this syndrome are often unilateral and calcified only in 10% of cases. Rarely, the tumor may be virilizing or renin secreting [6]. In our case, there was no raise of renin but a mild hyperprolactinemia, which is known to be associated with large uterine fibromas [7].

Sonography is generally used as the first-line imaging technique for the evaluation of ovarian pathologic abnormalities; nonetheless their sonographic features are often non-specific, therefore MRI is often needed for further differentiation [8].

On Ultrasound, fibromas most commonly manifest as solid, hypoechoic masses with sound attenuation. However, the US appearance is variable, and hyperechoic masses with increased through-transmission may be seen [9].

On CT, fibromas manifest as diffuse, slightly hypoattenuating masses; unlike most other solid masses, fibromas show poor, very slow enhancement with administration of contrast material [9].

MR is an excellent imaging modality for studying female pelvis and permits detection and excellent characterization of uterine and ovarian masses [8]. Ovarian fibromas in conventional MR imaging studies are characterized by low to intermediate signal intensity on T1-weighted images and low signal intensity on T2-weighted images caused by densely packed connective tissue.

MRI features of fibromas and fibrothecomas depend on the size of the lesion because the lesions measuring more than 60 mm can mimic malignant ovarian tumors as they present as solid adnexal masses sometimes associated with a pseudocapsule, degenerative changes, oedema, and peripheral subcapsular cystic areas. This explains the varying degree of high intensity portions on T2-weighted images and slow heterogenous enhancement [8, 10]. Only hemorrhage within the lesions would be characterized by the appearance of high signal intensity on T1-weighted images and heterogenous low and high signal mixed intensity on T2-weighted images [11].

On T2-weighted images fibromas and thecomas with a fibrotic component appear as well-circumscribed masses with low signal intensity containing scattered high-signal intensity areas representing oedema or cystic degeneration. The imaging appearance of thecomas without prominent fibrosis is similar to malignant tumors. The prominent lipid component of thecomas could be depicted with chemical-shift MR imaging [9].

The addition of diffusion-weighted imaging (DWI) and perfusion weighted imaging (PWI) to conventional MR might help to diagnose fibrothecomas [10] especially when the ADC map does not show any relevant restricted diffusion areas.

Sometimes ascites and pleural effusions may be present, with large fibromas [8]; the presence of normal ovarian tissue adjacent to an ovarian lesion is a useful morphological feature that can help exclude invasive ovarian malignancy [10]. Therefore, the association of pelvic masses with heterogenous T2-signal and pelvic intraperitoneal fluid do not necessarily imply the presence of a malignant neoplasm, as shown in literature, but it is often related to degenerative aspects in the largest fibromas and fibrothecomas.
Treatment & Prognosis:

Treatment of this syndrome must be multidisciplinary and will depend on the systems affected. Prognosis depends on the malignant progression of the lesions; however, life expectancy is no different from that of the general population [3].

Most ovarian fibromas are asymptomatic but when sufficiently large, they present with symptoms related to an abdominal mass, as gastrointestinal or genitourinary symptoms. Rarely, a patient may present with acute symptoms secondary to torsion of the fibroma, as has occurred in other studies. The decision to perform conservative surgery with preservation of ovarian function is problematic. Although ovarian malignancies have not been reported with this syndrome, Abel and Holtz have reported two ovarian fibrosarcomas in a review of 170 adolescents with ovarian tumors [12].

In the absence of gynecological symptoms, it seems wisest to preserve ovarian function in young patients, as there is no reason to sacrifice childbearing capacity. Moreover, fertility does not appear to be affected in NBCCS [13].

Differential Diagnosis:

Leiomyomas: the distinction of ovarian fibromas and uterine leiomyomas may be difficult, as they both present as solid hypoechoic masses with ultrasound beam attenuation on US; on CT they are hypodense masses with heterogeneous enhancement after intravenous contrast administration; on MRI they both show low-signal either in T1-weighted and T2-weighted images. A careful detection of the pedicle extending toward the uterus, vascular signal voids between the uterus and tumor mass, as well as the evaluation of the relationship between the ipsilateral ovary, can facilitate the diagnosis [10].

Germ-cell tumors: on US germ-cell tumors present as non-specific complex mass with possible calcification; on CT and MRI calcifications, cystic components and small foci of fat are suggestive. Hemorrhage may be present. In this case these calcified solid masses showed no evidence of fat, making germ-cell tumors a less-likely diagnosis [14].

Pelvic metastases: on US they present as mixed echogenicity tumors with vascularity of solid component on Doppler. On CT they present as soft tissue density with areas of cystic necrosis and, after contrast administration, solid components demonstrate heterogeneous enhancement. On T1-weighted MRI images they are usually iso to hypointense with variable enhancement after contrast administration and on T2-weighted images they show heterogenous signal of solid component with hyperintensity of cystic component. In this case, the patient had a history of basal cell carcinoma that does not appear to be affected in NBCCS [13].

In conclusion, we presented the case of a young Caucasian woman affected by NBCCS that met three of the diagnostic criteria including BCCs, odontogenic keratocysts and ovarian fibromas. The use of several imaging modalities (ultrasound, MRI and CT) allowed us to formulate a hypothesis of benign uterine and ovarian lesions allowing the patient to have laparoscopic fertility-sparing surgery.

TEACHING POINT

The association of pelvic masses with heterogeneous T2-signal and pelvic intraperitoneal fluid do not necessarily imply the presence of a malignant neoplasia but it is often related to degenerative aspects in large fibromas and fibrothecomas. In the differential diagnosis clinical and radiological features must be taken into account globally to identify possible Nevoid Basal Cell Carcinoma Syndrome.

REFERENCES


2. Lo Muzio L: Nevoid basal cell carcinoma syndrome (Gorlin syndrome). Orphanet J Rare Dis 2008; 3:32. PMID: 19032739


Figure 1: 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome. Findings: ultrasound imaging highlights the presence of heterogeneous hypoechoic uterine round-shaped mass measuring 65 x 62 x 59 mm adherent to the anterior uterine wall (red arrow). Left ovary was appreciable with difficulty due to the presence of a rounded hypoechoic mass measuring 18 x 15 mm (yellow arrow: an ovarian mass). Ovarian follicles were not depictable bilaterally.

Technique: General Electric Voluson E6 (transrectal transducer, 4-9 MHz).

Figure 2 (left): 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: Magnetic Resonance shows multiple lesions of the uterus and both ovaries (red arrows). In left adnexal region there are at least four masses, with prevalent hypointense signal in T2-weighted sequences (a: axial T2-weighted, b: axial fat-saturated T2-weighted), with little increase in signal intensity on higher b values diffusion images (red circle) and mild heterogeneous restricted diffusion on ADC map with ADC coefficient of 0.6 mm2/s (c: Diffusion Weighted sequence with different b values: 0, 330, 660, 1000 s/mm2; d: ADC map).

Figure 3: 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: On T2-weighted sequences (a: sagittal plane; b: axial plane; c: coronal plane) the largest lesion adherent to the anterior uterine wall shows heterogenous signal (red arrows). ADC map does not show any relevant restricted diffusion areas (d).

Technique: Philips Ingenia 1.5 T Magnetic Resonance System. Spin Echo T2-weighted sequences (TR: 4050,55 - TE: 110); Diffusion-weighted sequence (TR: 1786,29 - TE: 90,06).
Figure 4: 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: On fat-saturated T1-weighted sequence before contrast administration, the lesions appear isointense compared to myometrium (a: axial plane before contrast administration); after contrast administration (b: axial plane after contrast administration; c: coronal plane after contrast administration; d: sagittal plane after contrast administration), the masses show heterogeneous intense contrast enhancement (red arrows indicate the largest lesion, yellow arrows the smallest masses).

Technique: Philips Ingenia 1.5 T Magnetic Resonance System. Pre and post intravenous contrast administration (contrast agent used: Gadolinium-DTPA 0.2 ml/Kg). Fat-saturated Gradient Echo T1-weighted sequence before contrast administration (TR: 4; TE 1.92); fat-saturated Gradient Echo T1-weighted sequence after contrast administration (TR: 4; TE 1.92).
**Figure 5:** 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: T2-weighted sequences (a: axial plane; b: coronal plane; c: sagittal plane) show a uterine hypointense pedunculated mass measuring 30 x 24 x 20 mm.


**Figure 6** (left): Computed Tomography. 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: Computed Tomography examination on axial (a: before IV contrast administration; b: after contrast administration in venous phase), sagittal (c) and coronal planes (d) demonstrates the presence of numerous uterine and ovarian masses (red arrows), partially calcified (asterisks) on precontrast images which show heterogeneous contrast-enhancement. Anteverted uterus, with regular zonal anatomy. Small amount of fluid in Douglas pouch. Ovarian follicles were not depictable bilaterally.

Technique: Computed Tomography General Electric LightSpeed VCT 128-slice, 410 mAs, 120 kV, 2.5 mm slice thickness, 1.25 mm gap, 120 ml Iomeprol 300, DLP 612.48 mGycm.
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Figure 7: 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: laparoscopic image showing the left adnexa with multiple fibroids.

Technique: laparoscopic surgery.

Figure 8: 25-year-old woman with multiple ovarian and uterine fibromas in Nevoid Basal Cell Carcinoma Syndrome.

Findings: a) small aggregates of fibroblast cells without cytologic atipies; b) calcifications in the context of cellular aggregates and fibrous tissue.

Technique: hematoxylin and eosin 10 x.
ETIOLOGY

Mutations in the PTCH1 gene

INCIDENCE
1/57,000 - 1/256,000

GENDER RATIO
No difference

AGE PREDILECTION
Adolescence

RISK FACTORS
No known risk factors

TREATMENT
Surgical resection (if possible, fertility-sparing surgery).

PROGNOSIS
Good

FINDINGS ON IMAGING
Pelvic Fibromas
CT: mass effect, possible calcifications. MRI: low to intermediate signal in T1w and low signal in T2w. In DWI no relevant restricted diffusion areas in ADC map.

Table 2: Summary table for Gorlin-Goltz Syndrome with imaging features of associated pelvic fibromas.
Table 3: Differential diagnosis table for pelvic fibromas.

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<tr>
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<th>US</th>
<th>CT</th>
<th>MRI</th>
</tr>
</thead>
<tbody>
<tr>
<td>FIBROMAS/ LEIOMYOMAS/ FIBROTHECOMAS</td>
<td>Variable appearance; generally solid, hypoechoic masses with ultrasound beam attenuation.</td>
<td>Hypoattenuating masses. Calcifications and bilaterality are both uncommon.</td>
<td>T1w: low signal. T2w: low signal, sometimes hyperintense areas representing edema or cystic degeneration.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>After IV contrast administration: heterogenous delayed enhancement.</td>
<td>T1w after contrast administration: heterogenous enhancement.</td>
</tr>
<tr>
<td>GERM-CELL TUMORS</td>
<td>Non-specific complex mass with possible calcifications.</td>
<td>Calcifications, cystic components and small foci of fat are suggestive. Hemorrhage may be present.</td>
<td>Calcifications, cystic components and small foci of fat are suggestive. Hemorrhage may be present.</td>
</tr>
<tr>
<td>PELVIC METASTASES</td>
<td>Mixed echogenicity tumors with vascularity of solid component on Doppler.</td>
<td>Soft tissue density with areas of cystic necrosis. After contrast administration, solid components demonstrate heterogenous enhancement.</td>
<td>T1: iso to hypointense with variable enhancement after contrast administration. T2: heterogenous signal of solid component with hyperintensity of cystic component.</td>
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**ABBREVIATIONS**

BCCs = Basal Cell Carcinomas  
CT = Computed Tomography  
DWI = Diffusion Weighted Imaging  
IV = intravenous  
MR = Magnetic Resonance  
MRI = Magnetic Resonance Imaging  
NBCCS = Nevoid Basal Cell Carcinoma Syndrome  
PWI = Perfusion Weighted Imaging  
US = Ultrasound

**KEYWORDS**

Gorlin-Goltz syndrome; NBCCS; ovarian fibromas; uterine fibromas; MRI

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