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Endometrial stromal tumours revisited: an update based on the 2014 WHO classification

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ABSTRACT

Endometrial stromal tumours (EST) are rare tumours of endometrial stromal origin that account for less than 2% of all uterine tumours. Recent cytogenetic and molecular advances in this area have improved our understanding of ESTs and helped refine their classification into more meaningful categories. Accordingly, the newly released 2014 WHO classification system recognises four categories: endometrial stromal nodule (ESN), low-grade endometrial stromal sarcoma (LGESS), high-grade endometrial stromal sarcoma (HGESS) and undifferentiated uterine sarcoma (UUS). At the molecular level, these tumours may demonstrate a relatively simple karyotype with a defining chromosomal rearrangement (as in the majority of ESNs, LGESSs and *YWHAE*-rearranged HGESS) or demonstrate complex cytogenetic aberrations lacking specific rearrangements (as in UUSs). Herein we provide an update on this topic aimed at the practicing pathologist.

INTRODUCTION

Endometrial stromal tumours (EST) are uncommon uterine tumours responsible for less than 10% of uterine mesenchymal neoplasms¹ and less than 2% of all uterine neoplasms. The diagnostic criteria and nomenclature of ESTs have gone through several modifications in the past four decades since the earliest study by Norris and Taylor.² The rarity of these tumours coupled with the overlapping morphological features have contributed to the difficulty in classifying ESTs into clinically meaningful categories.

Briefly, endometrial stromal sarcomas (ESS) were originally divided into low-grade and high-grade based on mitotic count,² however, further studies have demonstrated that mitotic activity is prognostically irrelevant. Later on, the 2003 WHO classification system eliminated the 'high-grade' category and reclassified these tumours into 'ESS' (low-grade tumours with histological resemblance to proliferative endometrial stroma) and 'undifferentiated endometrial sarcoma' (pleomorphic tumours with no resemblance to endometrial stroma). Subsequently, it has become increasingly evident that the 'undifferentiated endometrial sarcoma' category is too broad, encompassing a wide range of heterogeneous tumours with different clinical behaviour and outcome, and that at least a subset of these tumours are morphologically and genetically distinct. The 2014 WHO classification scheme now incorporates recent molecular findings into the classification, dividing ESTs into endometrial stromal nodule (ESN), low-grade endometrial stromal sarcoma (LGESS), high-grade endometrial stromal sarcoma (HGESS) and undifferentiated uterine sarcoma (UUS).

ENDOMETRIAL STROMAL NODULE

Overview

ESNs are rare benign tumours of the uterus. The two largest series published to date showed no recurrences after a follow-up period of up to 16 years and 17.8 years, respectively.^{3,4} ESN occurs over a wide age range, from 31 to 86 years with a mean of 53 years in one study,⁴ usually presenting with abnormal vaginal bleeding or as an incidental finding in hysterectomy specimens performed for other reasons.

Morphological features

Grossly, ESN is a well-circumscribed tumour of variable size that may occur as an intramural mass or as a polypoid tumour protruding into the endometrial cavity. It usually has a fleshy and soft yellow to tan cut surface that may contain cystic areas. Examples with firm white cut surface resembling a leiomyoma usually show extensive stromal fibrosis histologically.

Microscopically, ESN is expansile in nature lacking myometrial invasion, yet focal finger-like projections into the adjacent myometrium (less than 3 in number and less than 3 mm in length) are accepted.³ Lymphovascular invasion must be absent. Tumours with clear-cut myometrial/lymphovascular invasion should be diagnosed as LGESS instead. ESNs are classically densely cellular, although occasional hypocellular examples with fibrous or fibromyxoid differentiation are encountered,⁴ and are characterised by whorling around spiral arteriole-like vessels. The tumour cells resemble those of proliferative endometrial stroma, exhibiting minimal cytological atypia. Mitotic activity is usually low (up to 5 per 10 high power fields) but a higher count does not exclude the diagnosis.⁴ ESNs demonstrate diverse morphological variations, which include smooth muscle, skeletal muscle and sex-cord stromal differentiation^{5–8} as well as a variety of other secondary features such as hyalinisation, collagen plaques, foamy macrophages and cholesterol clefts.

A definitive diagnosis of ESN can only be rendered after careful sampling and examination of the tumour border, which is only possible on hysterectomy specimens. On curettage specimens, the identification of aglandular stromal fragments ≥ 5 mm in size should raise suspicion of an underlying EST of some sort,⁹ with a caveat that the distinction between ESN and LGESS may be impossible in such specimens. Occasional cases, so-called 'EST with limited infiltration', show more tumour border irregularities than is usually expected in conventional ESNs, yet the infiltration is not as extensive as that of LGESS.⁴ In the



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current WHO classification, this is not recognised as a distinct diagnostic category, but the available limited clinical data suggest a benign course.

Ancillary studies and differential diagnosis

ESN and LGEES share the same immunohistochemical profile (discussed below). As in LGEES, $t(7;17)$ resulting in *JAZF1-SUZ12* gene fusion is the most common chromosomal rearrangement, identified in over half of the cases, conventional as well as variant ESNs.¹⁰

The most important differential diagnoses are LGEES and cellular/highly cellular leiomyoma (table 1).¹¹ Distinction from LGEES basically relies on exclusion of myometrial and lymphovascular invasion, as the morphology is indistinguishable. Smooth muscle differentiation in ESN can be particularly confusing as one may misinterpret the interdigitating metaplastic smooth muscle as myometrial invasion. Delineating the true tumour border at gross examination is extremely helpful in this case. Highly cellular leiomyomas contain large thick-walled blood vessels, cleft-like spaces and at least focal conventional fascicular pattern.¹¹ Although smooth muscle tumours and ESTs have an overlapping immunophenotype, an immunohistochemical panel of CD10 and at least two muscle markers (such as desmin, h-caldesmon, smooth muscle heavy chain myosin and HDAC8) usually helps in this distinction.^{12–15}

LOW-GRADE ENDOMETRIAL STROMAL SARCOMA

Overview

LGEES is the second most common malignant mesenchymal tumour of the uterus,¹⁶ and it is much more common than ESN. The age range is somewhat similar to that of ESN, with the majority occurring in the perimenopausal period. It occasionally arises in young women and adolescents.^{17–20} Patients usually present with vaginal bleeding or pelvic pain. The uterine corpus is the most frequent location though it can also primarily arise in a variety of extrauterine locations such as ovary,²¹ pelvis, abdominal cavity, vulva and vagina. It often occurs in association with endometriosis.^{22–27}

LGEES is a slow-growing tumour with an indolent clinical course characterised by multiple and/or late relapses, some occurring as late as 20 years after hysterectomy.²⁸ The relapses may be local (pelvic/vaginal), distant (most common are abdominal

cavity/wall and lungs), or both (figure 1).²⁹ Relapse rates up to 56% have been reported in patients with early-stage disease possibly due to inadvertent tumour morcellation at surgery.^{30–31} The stage is the most important prognostic factor.³²

Currently, total hysterectomy and bilateral salpingo-oophorectomy, with or without adjuvant therapy, are the mainstay of treatment for stage I–II LGEES.^{33–34} Lymph node metastasis can occur,³⁵ however, lymphadenectomy is unlikely to improve survival.^{36–38} Reports on conservative fertility-sparing management in young patients are emerging though experience with this approach is still limited.^{17–20–29–39} Advanced-stage (III–IV) disease is managed with cytoreductive surgery and adjuvant hormonal therapy usually in the form of progestins or aromatase inhibitors.^{31–40} Radiation therapy may be useful for local control but is not universally recommended.^{37–41–42} Tamoxifen and oestrogen replacement therapy are contraindicated as they promote disease progression.⁴³

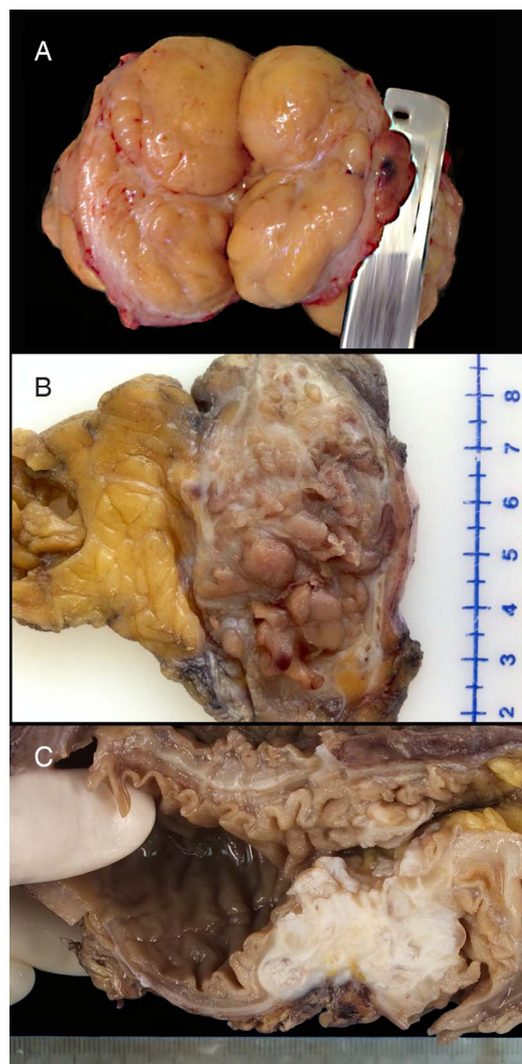


Figure 1 Gross appearance of low-grade endometrial stromal sarcoma (LGEES). (A) A 'myomectomy' specimen showing soft yellow cut surface, proved to be LGEES histologically. Courtesy of Dr Remaa Al-Safi, Department of Pathology, Maternity Hospital, Kuwait. (B) Recurrent LGEES in the soft tissues of abdominal wall following hysterectomy, showing bulging 'wormlike' masses of tissue reminiscent of that seen in the uterus. (C) Recurrent LGEES in the abdominal cavity involving intestinal wall showing white rubbery appearance corresponding to extensive hyaline plaques.

Table 1 Differential diagnoses of endometrial stromal tumours

ESN	LGEES	HGEES	UES
Cellular leiomyoma	Gland-poor adenomyosis	LGEES	Leiomyosarcoma
LGEES	Cellular leiomyoma	Leiomyosarcoma	HGEES
	Intravascular leiomyomatosis	UUS	Undifferentiated carcinoma
	Leiomyosarcoma with extensive intravascular component	GIST	Carcinosarcoma
	HGEES		Adenosarcoma with stromal overgrowth
	UTROSCT		Lymphoma
	PEComa		Melanoma
	GIST		

ESN, endometrial stromal nodule; GIST, gastrointestinal stromal tumour; HGEES, high-grade endometrial stromal tumour; LGEES, low-grade endometrial stromal tumour; PEComa, perivascular epithelioid cell tumour; UTROSCT, uterine tumour resembling ovarian sex-cord tumour; UUS, undifferentiated uterine sarcoma.

Morphological features

Grossly, LGESS may be submucosal or intramural, usually with ill-defined borders and 'wormlike' permeation within the myometrium and parametrial tissue, though some tumours might appear relatively circumscribed. Similar to ESN, the cut surface is fleshy tan to yellow (figure 1A, B), but may occasionally be firm white when there is extensive fibrous stroma (figure 1C). Areas of haemorrhage, necrosis and cystic degeneration might be present.⁴⁴ Infrequently, LGESS appears as a pure cystic mass.

The 'tongue-like' patterns of myometrial and lymphovascular invasion are classical histological features that are important in the distinction from ESN (figure 2A, E). Similar to ESN though, LGESS is composed of sheets of small uniform cells with oval to spindle nuclei, whorling around spiral arteriole-like vessels (figure 2F). The neoplastic cells resemble those of proliferative endometrial stroma and lack significant cytological atypia.^{2,45} Mitotic activity is variable. Variant histological features include smooth muscle differentiation ('star-burst' appearance), rhabdomyoblastic, rhabdoid, fibromyxoid, sex-cord, glandular, collagen plaques and others (figure 2B–D).^{46–51} As discussed above, it is not possible to separate LGESS from ESN on curettage specimens (figure 2G).

Ancillary studies and differential diagnosis

LGESS is positive for CD10 (figure 2H), oestrogen receptor (ER) and progesterone receptor (PgR). However, CD10 is not

entirely specific and may stain some smooth muscle tumours, especially leiomyosarcoma and almost 50% of highly cellular leiomyomas. Conversely, false-negative CD10 staining can be encountered with underfixation. Therefore, it is important to use a panel of immunostains that includes CD10 and at least two smooth muscle markers (eg, desmin, h-caldesmon, smooth muscle heavy chain myosin, HDAC8) as there is no single marker that is specific for ESS.⁵² Not uncommonly, LGESSs stain with smooth muscle markers, especially at foci of muscle differentiation but also in conventional areas (depending on the marker used). Adding h-Caldesmon is often helpful as it has good sensitivity and specificity for smooth muscle differentiation.⁵³ So it is important that one takes into account the relative intensity and distribution of these stains and tries to interpret them in view of the histological features, gross findings and clinical picture. Foci of sex-cord differentiation stain with calretinin, inhibin, WT1, Melan-A and CD99.

Important differential diagnoses for uterine LGESS include smooth muscle tumours (intravascular leiomyomatosis, cellular and highly cellular leiomyoma, leiomyosarcoma), adenomyosis with sparse glands, perivascular epithelioid cell tumour (PEComa) and uterine tumour resembling ovarian sex cord tumour (UTROSCT; table 1). Attention to histological features and use of appropriate immunostains are often helpful. For example, LGESS with myxoid change (figure 2C) can be

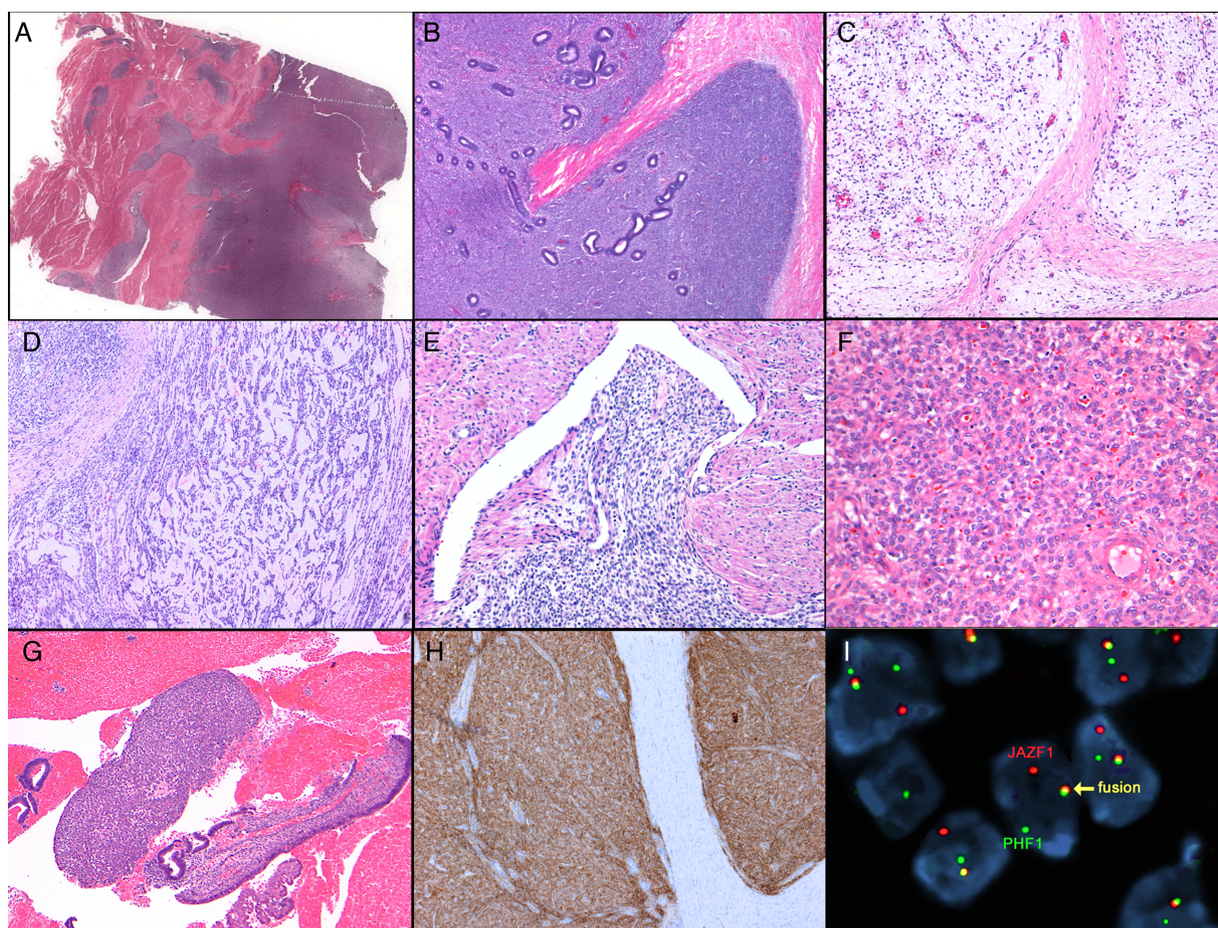


Figure 2 Microscopic appearance of low-grade endometrial stromal sarcoma (LGESS). (A) *JAZF1*-rearranged LGESS at scanning magnification with 'tongue-like' infiltration. (B) Endometrioid glandular elements are rarely seen (H&E $\times 5$). (C) *PPH1*-rearranged LGESS with striking myxoid change and preservation of arteriolar pattern (H&E $\times 10$). (D) Extensive hyaline plaques (H&E $\times 5$). (E) Lymphovascular invasion (H&E $\times 10$). (F) Spiral arteriole-like vessels and sprinkle of 'stromal granulocytes' are helpful features (H&E $\times 10$). (G) Curettage specimen showing fragments of aglandular endometrial stroma, which proved to be LGESS on hysterectomy (H&E $\times 10$). (H) CD10 immunostain positivity (CD10 $\times 5$). (I) Fluorescence in situ hybridisation (FISH) fusion assay showing *JAZF1-PPH1* fusion (yellow signal).

confused with myxoid smooth muscle lesions, but the 'tongue-like' infiltration, small arterioles, presence of classic areas of LGESS elsewhere, along with the CD10 immunostaining and muscle marker negativity, will usually resolve the diagnosis.⁵⁴

The differential diagnosis for extrauterine LGESS depends on the location of the tumour. In case of ovarian location, metastasis from a primary uterine LGESS and ovarian sex cord-stromal tumours should be excluded.²¹ In the abdominal cavity, gastrointestinal stromal tumour (GIST) comes into the differential and should be excluded with the combined use of cKit and DOG1 immunostains.

The majority of LGESSs, including conventional and variant tumours, harbour chromosomal rearrangements,^{10 18 55–58} but at present there seems to be no correlation between the histological variant and the underlying genotype. The most common rearrangement, t(7;17)(p15;q21), results in *JAZF1-SUZ12* gene fusion.⁵⁹ Other gene fusions include *JAZF1-PHF1* (figure 2I), and the much less common *EPC1-PHF1*, *MEAF6-PHF1*,^{60 61} *ZC3H7-BCOR*⁶² and *MBTD1-CXorf67*.⁶³ Molecular testing, however, is not routinely performed but may be helpful when dealing with cases of unusual location or morphology.

HIGH-GRADE ENDOMETRIAL STROMAL SARCOMA Overview

One of the main highlights of the updated 2014 WHO classification of uterine mesenchymal tumours¹⁶ is the re-introduction of HGESS as a distinct entity. The identification of *YWHAE-NUTM2A/B* (also known as *YWHAE-FAM22A/B*) gene fusion⁶⁴ as a recurrent event in this more aggressive subset of tumours has provided objective support to the existence of a subcategory of ESS that is intermediate between LGESS and UUS. Currently, the term HGESS is reserved for those rare stromal sarcomas exhibiting round cell morphology and characteristic immunophenotype, and harbouring a t(10;17)(q22;p13) rearrangement with *YWHAE-NUTM2A/B* gene fusion. In the past, at least some of these tumours may have been lumped under 'undifferentiated endometrial sarcoma'. The paper by Kurihara *et al*⁶⁵ has been instrumental in drawing attention to the heterogeneity of the undifferentiated tumour category, stressing the importance of separating tumours within this category into those with nuclear uniformity (some may

have been *YWHAE*-rearranged) from those with nuclear pleomorphism (real undifferentiated sarcomas characterised by complex karyotypes and frequent p53 alterations). Additionally, separating HGESS from LGESS is as important since they appear to be more aggressive clinically with frequent extrauterine disease at initial presentation.^{66 67} Based on the limited data available, the prognosis of *YWHAE*-rearranged cases seems to be intermediate between that of LGESS and UUS.^{66 68} These tumours usually present as abnormal vaginal bleeding similar to LGESS. Table 2 summarises the molecularly confirmed HGESS cases reported to date.

Morphological features

On gross examination, HGESS may be polypoid and usually shows extensive permeative growth through the uterine wall akin to LGESS, frequently with haemorrhage and necrosis.⁶⁶ Microscopically, HGESS demonstrates the 'tongue-like' permeative pattern into the myometrium and angiolymphatic invasion typical of LGESS (figure 3A). The cytomorphology, however, is distinctive and does not exactly look like proliferative endometrial stromal cells. It is characterised by a monomorphic proliferation of round cells in a vaguely nested or pseudoglandular pattern⁶⁶ (figure 3B–D). About half of the tumours are biphasic, where the round-cell component is admixed with an LGESS-like component that is usually fibrous or fibromyxoid.⁶⁶ The round cells are larger than those of LGESS with slightly irregular nuclear contour, fine evenly dispersed chromatin with nuclear clearing and lack of prominent nucleoli, while the cytoplasm may be scant to moderate in amount, imparting an epithelioid appearance (figure 3D). The tumour displays delicate curvilinear vasculature that is different from the spiral arterioles seen in LGESS (figure 3C). Mitotic activity is increased and necrosis is common. While *YWHAE*-rearranged endometrial sarcomas consistently show uniform high-grade cytomorphology, the reverse is not always true (ie, not all uniform high-grade endometrial sarcomas are *YWHAE*-associated). Sciallis *et al*⁶⁹ recently identified three morphological patterns of non-pleomorphic HGESSs, and found that the *YWHAE* rearrangement was only detected in a subgroup displaying the characteristic morphological features described above along with the characteristic immunophenotype (below).

Table 2 Molecularly confirmed high-grade endometrial stromal sarcoma (HGESS) cases in the literature

Author	Year	Number of cases	Molecular method	FIGO stage at presentation	Presence of LGESS-like component histologically*
Ali <i>et al</i> ¹⁸	2014	1	FISH breakapart	IIIc	0/1
Gremel <i>et al</i> ⁶⁸	2014	3	RT-PCR	NA	NA
Sciallis <i>et al</i> ⁶⁹	2014	4	FISH breakapart	1/4 stage IIIB 1/4 stage IIIC 2/4 stage IVB	NA
Stewart <i>et al</i> ⁵⁵	2014	4	FISH breakapart	NA	1/4
Croce <i>et al</i> ⁷⁰	2013	6	FISH breakapart and RT-PCR	3/6 stage IB 1/6 stage IVB 2/6 NA	1/6 at least †
Lee <i>et al</i> ^{66‡}	2012	13	FISH breakapart and karyotyping	10/12 stages II to III 2/12 stage I	8/13
Amant <i>et al</i> ⁷¹	2011	1	t(10;17) by karyotyping and M-FISH	IIIc	NA
Regauer <i>et al</i> ⁷²	2008	1	t(10;17) by karyotyping and M-FISH	Recurrent abdominal mass	1/1
Micci <i>et al</i> ⁷³	2003	1	t(10;17) by karyotyping and RxFISH	Metastatic to lung	NA
Leunen <i>et al</i> ⁷⁴	2003	1	t(10;17) by karyotyping	III	NA

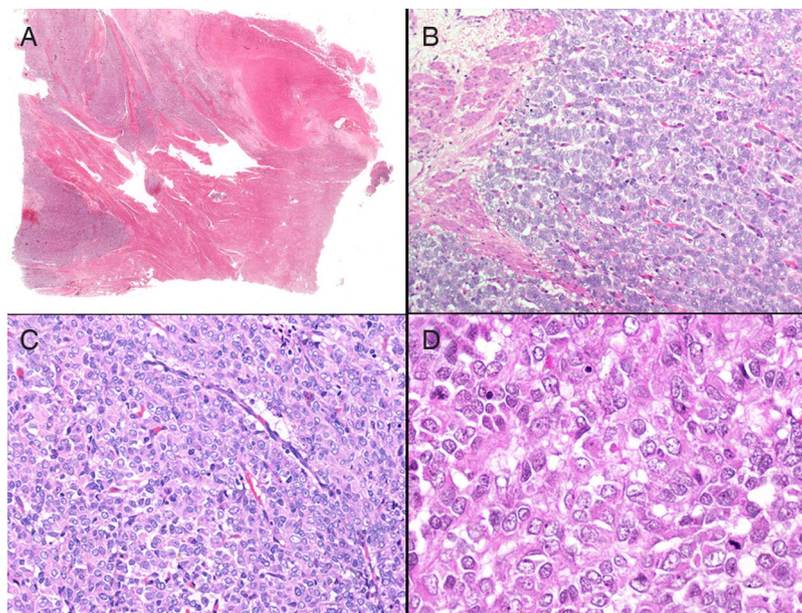
*Some *YWHAE*-rearranged HGESSs contain an LGESS-like component that is fibrous or fibromyxoid.

†Three additional cases were described as 'spindle' with no details on the grade of these spindle areas.

‡Original paper that described *YWHAE* rearrangement in HGESS.

LGESS, low-grade endometrial stromal sarcoma; RT-PCR, reverse transcriptase-PCR; NA, not available.

Figure 3 Microscopic appearance of *YWHAE*-rearranged high-grade endometrial stromal sarcoma (HGESS). (A) Scanning magnification showing 'tongue-like' infiltration similar to low-grade endometrial stromal sarcoma. (B) Nested and pseudoglandular pattern (H&E $\times 20$). (C) Delicate linear vascular pattern and lack of perivascular whorling (H&E $\times 20$). (D) Epithelioid cytomorphology with moderate amount of cytoplasm, irregular nuclear contour and nuclear clearing (H&E $\times 40$).



Ancillary studies and differential diagnosis

The high-grade round cell/epithelioid component of the tumour characteristically shows diffuse strong nuclear staining for cyclin D1,⁷⁵ lack of CD10, and weak or absent staining for ER and PR, while the low-grade component typically shows diffuse positivity for CD10, ER and PR similar to conventional LGESS⁶⁶ (figure 4A–H). The lack of hormone receptor expression may have potential treatment implications. Diffuse cytoplasmic/membranous c-Kit expression is consistently seen in the round cell areas.⁷⁶ The distinctive morphological and immunohistochemical features of HGESS are generally good surrogate

markers for the underlying rearrangement, but fluorescence in situ hybridisation and/or reverse transcriptase-PCR studies may be used in equivocal cases.^{70 77}

YWHAE-rearranged HGESS should be distinguished from LGESS because of the prognostic and treatment implications. LGESS shows smaller 'blue' cells with scanty cytoplasm and smooth nuclear contour, characteristic spiral arterioles, lack of biphasic appearance, infrequent necrosis and CD10/ER positive cyclin D1/cKit negative immunoprofile. In excision specimens, adequate sampling is necessary in order not to overlook any fibrous/fibromyxoid LGESS component that helps establish the

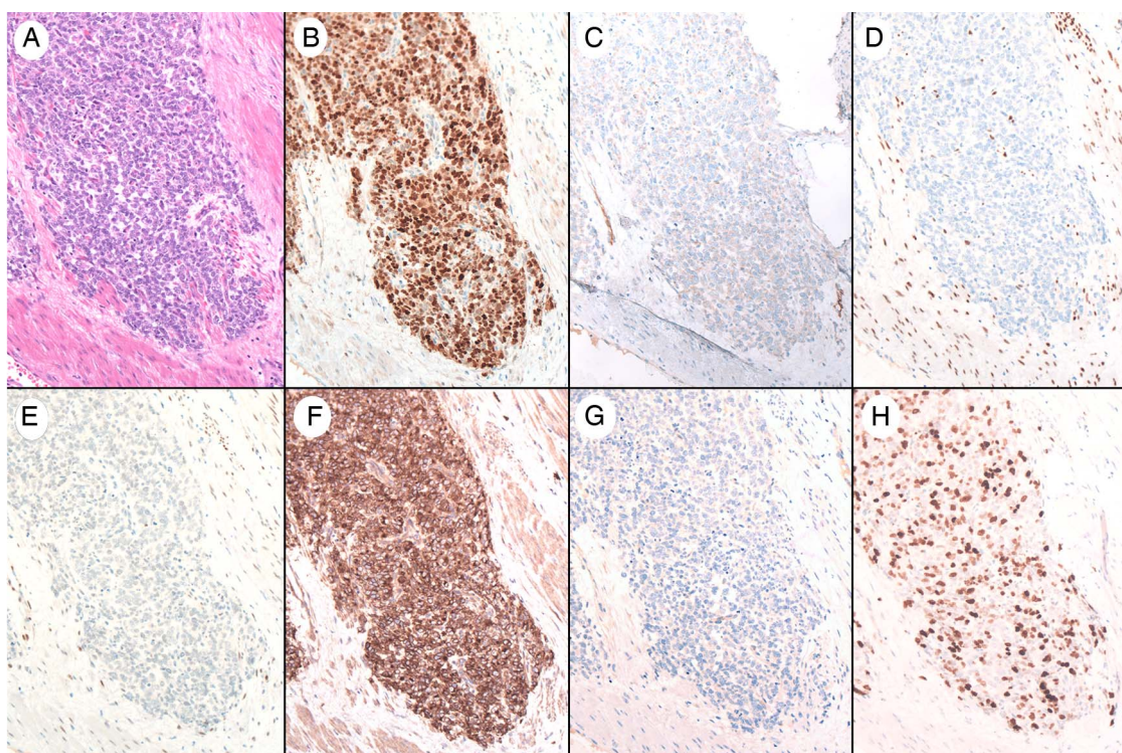


Figure 4 Immunophenotype of *YWHAE*-rearranged high-grade endometrial stromal sarcoma. (A) H&E of the high-grade round cell component of the tumour. (B) Diffuse cyclin D1 nuclear immunostaining. (C) CD10 negative. (D) Oestrogen receptor negative. (E) Progesterone receptor negative. (F) Diffuse cKit staining. (G) DOG1 negative. (H) Ki67 shows a high proliferative index (20 \times magnification).

diagnosis of HGEES. Epithelioid leiomyosarcoma should be excluded by appropriate smooth muscle immunomarkers, presence of at least focal marked atypia, \pm presence of conventional leiomyosarcoma. Perivascular epithelioid cell tumour (PEComa) of gynaecological origin may show 'finger-like' projections analogous to those of ESS but typically shows epithelioid and spindled cells with clear to eosinophilic cytoplasm, nested growth pattern, prominent vascular network and coexpression of melanocytic and muscle markers.⁷⁸ HGEES may also be potentially confused with epithelioid gastrointestinal stromal tumour particularly in intraperitoneal/pelvic locations in view of the frequent extrauterine spread at presentation and cKit positivity. In these instances radiological–pathological correlation for possible uterine origin along with lack of DOG1 staining are often helpful.⁷⁶

UNDIFFERENTIATED UTERINE SARCOMA

Overview

In order to encompass tumours of endometrial and non-endometrial derivation, the term UUS has now replaced the older term 'undifferentiated endometrial sarcoma' in the updated WHO book. This wastebasket category comprises a heterogeneous group of high-grade tumours that fail to fulfil the morphological and immunohistochemical criteria of translocation-positive endometrial sarcomas. These are highly aggressive tumours, typically presenting in older patients as postmenopausal bleeding with/without manifestations of high-stage extrauterine disease. The vast majority appear to arise through genetic pathways distinct from those of LGEES and HGEES,^{79–80} demonstrating complex karyotypes and genomic gains and losses but lacking specific translocations.^{80–81} Rare cases with a coexisting LGEES component have been reported suggesting that at least a subset may arise through 'dedifferentiation' in a low-grade sarcoma.^{65–82–83}

Morphological features

Grossly, UUSs are relatively large fleshy tumours demonstrating destructive infiltrative growth into the uterine wall, typically associated with extensive necrosis and/or haemorrhage.⁸⁴ Histologically, they show sheets or fascicles of highly atypical non-descript cells that do not resemble proliferative-phase endometrial stroma (figure 5). Lymphovascular invasion is common and mitotic activity is brisk and commonly associated with atypical mitotic figures.

Ancillary studies and differential diagnosis

No consistent immunophenotype is identified reflecting the heterogeneity of tumours in this category. CD10 is variably expressed⁸⁴ and can be diffuse, therefore this marker is not usually helpful in this morphological context. Hormone receptors may be negative or weakly positive.

UUS is a diagnosis of exclusion and should only be made after extensive sampling of the tumour in order to exclude other more common high-grade neoplasms with evidence of differentiation. These include leiomyosarcoma (desmin/h-caldesmon positive, CK/EMA frequently positive), carcinosarcoma (at least focal malignant epithelial component present), rhabdomyosarcoma (rhabdomyoblasts, desmin/myogenin/MyoD1 positive), sarcomatous overgrowth in a Mullerian adenocarcinoma (at least focal low-grade areas with benign epithelial elements), undifferentiated endometrial carcinoma (uniform dyshesive cells resembling lymphocytes \pm low-grade endometrioid carcinoma component, at least focal keratin/EMA)⁸⁵ and diffuse large B-cell lymphoma (resolved by appropriate immunohistochemistry).

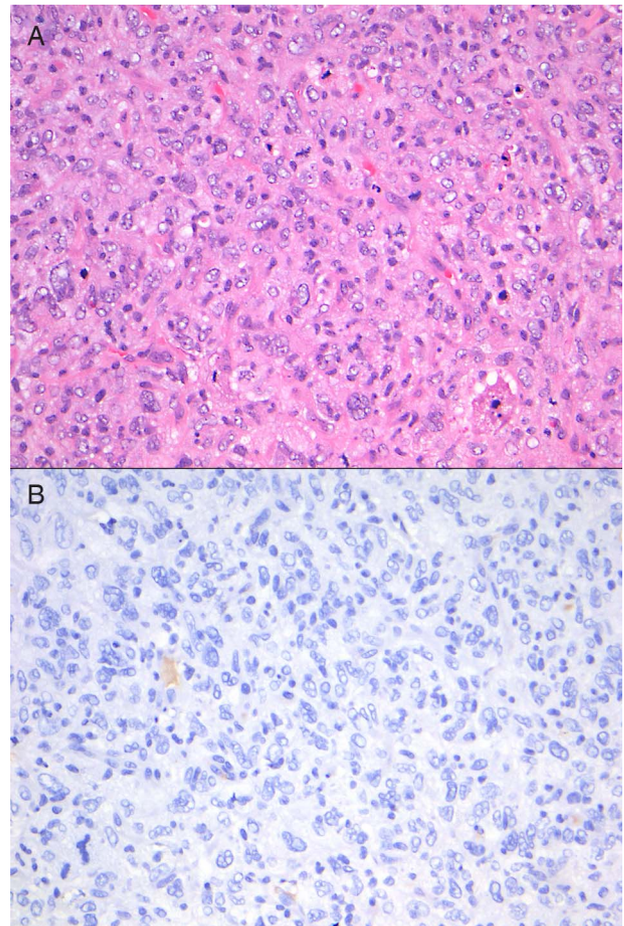


Figure 5 Undifferentiated uterine sarcoma. (A) Pleomorphic cytomorphology and brisk mitosis. (B) h-caldesmon immunostain is negative (20 \times magnification).

To sum up, our understanding of ESTs has evolved dramatically in the past couple of decades despite the rarity of these tumours and overlapping morphological features. In particular, molecular advances have played a major role in characterising ESTs more objectively. Further research is required to shed more light on the underlying molecular pathways, gain more insights into pathogenesis and optimise treatment strategies as an ultimate goal.

Take home messages

- ▶ Endometrial stromal tumours are rare uterine mesenchymal neoplasms that may mimic a variety of other uterine tumours.
- ▶ The rarity of endometrial stromal tumours has made them difficult to study and classify into meaningful clinicopathological categories.
- ▶ Recent molecular advances, which are now incorporated into the 2014 World Health Organization (WHO) classification system, have objectively assisted in the classification of these tumours.
- ▶ The WHO classification system now recognizes four tumour categories with a clear distinction between low-grade and high-grade endometrial stromal sarcomas.

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