Elastography of focal testicular lesions: current concepts and utility

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As a relatively new sonographic technique, tissue elastography has emerged as a qualitative and potentially quantitative adjunctive tool to provide additional information on tissue stiffness, aiming to further improve diagnostic confidence in discriminating benign from malignant focal testicular lesions. The purpose of this review is to provide an overview of the elastography techniques used to assess focal testicular lesions and their typical appearance on tissue elastography.

Keywords: Ultrasonography; Testicular neoplasms; Elasticity imaging techniques; Leydig cells tumor; Epidermal cyst

Introduction

Traditionally, assessment of the testis with B-mode and color Doppler ultrasonography (US) has been the main workhorse for identifying and characterising focal testicular lesions owing to its superb spatial resolution of anatomical details and the superficial location of the testis. B-mode US accurately identifies the size, shape, location (intra- or extra-testicular), and pattern of echogenicity of lesions. Colour Doppler US assesses the presence and pattern of vascularity in the lesion to increase diagnostic confidence, with disorganized vascularity suggesting a malignant lesion and avascularity or iso-vascularity compared to the background testicular tissue suggesting a benign lesion. Relatively new sonographic techniques such as tissue elastography have emerged as a qualitative and potentially quantitative adjunctive tool to provide additional information on tissue stiffness, aiming to further improve diagnostic confidence in discriminating benign from malignant lesions, under the assumption that malignant lesions are “harder” than benign lesions. The purpose of this review is to provide an overview of the elastography techniques used to assess focal testicular lesions and their typical appearance on tissue elastography.

Elastography Techniques

Strain Elastography

There are two types of elastography techniques, commonly referred to as strain and shear wave elastography. Strain elastography measures the tissue strain response when a compression force is applied to the tissue. The strain elastography technique was first used in clinical practice in the form of a proprietary technique known as real-time elastography (RTE, Hitachi Medical Corp., Tokyo, Japan). RTE evaluates the relative strain response of the tissue, including the region of interest and...
the adjacent tissue, to mechanical stress by analyzing echo signals before and after light compression is applied by the transducer. The strain response of the tissue is converted into an elastic modulus image known as an elastogram and displayed as a color-coded map, superimposed on to the conventional B-mode image. The color spectrum ranges from red to blue, representing soft to hard tissue stiffness. The visual elasticity score (VES), according to which the elastographic appearance is graded on a 5- or 6-point scale, adapted from breast elastography [1], has been used to categorize the stiffness of testicular abnormalities (Table 1) [2–4]. A VES cut-off of 3 may be used to classify testicular lesion stiffness as "hard" (VES>3) or "soft" (VES≤3). In addition to the visual map, the strain ratio (SR) between the abnormal testicular tissue and adjacent normal testicular parenchyma can be calculated as a semi-quantitative numerical index. It is only a semi-quantitative method, as it is used to assess the relative tissue elasticity of the region of interest and background tissue, rather than the absolute tissue elasticity.

**Shear Wave Elastography**

Shear wave elastography differs from strain elastography, as it is a quantitative method of assessing tissue elasticity by measuring the speed of acoustic radiation force impulse–induced shear waves traveling in the tissue of interest [5]. Shear waves are a type of mechanical wave whose propagation speed within the soft tissue can be measured. Shear wave velocities are measured by the scanner in m/sec, and they can be converted to the Young modulus of elasticity (kPa) using the simple equation \( E = 3 \rho c^2 \) (\( E \) = the Young modulus, \( \rho \) = tissue density, \( c \) = shear wave speed). Today, almost all commercial scanners provide values both in m/sec and kPa. However, it is preferable to report values in m/sec because it is the quantity directly measured by the machine and the above simple conversion to the Young modulus (\( E \)) is accurate only if several assumptions are true [6].

**Normal Testes**

The color map of the strain elastogram from a normal testicle demonstrates a central parenchymal green color with blue edges, which is surrounded by red bands (Fig. 1). This is known as a 3-ring structure [7]. Mean shear wave velocities of 0.76 m/sec (95% confidence interval, 0.75 to 0.78 m/sec) from 130 normal testicles and 1.92 m/sec in 19 patients with testicular cancer have been reported by Pedersen et al. [8] and Marcon et al. [9], respectively, both using virtual touch tissue imaging quantification (VTIQ; Siemens Medical Solutions, Mountain View, CA, USA). Furthermore,
Trottmann et al. [11] reported that the center of the testis showed significantly lower shear wave velocity than the inferior and superior portion of the testis, through results obtained from 60 healthy testes using SuperSonic Imaging (Aixplorer, Paris, France). Therefore, clinicians may consider using different normal values as references when comparing the results of pathological testicular tissue or focal lesions in different areas. In addition to intra-testicular shear wave velocity variability, a previous study demonstrated significant differences in shear wave velocities measured on different machines [12]. Trottmann et al. [13] showed that values obtained using SuperSonic Imaging were significantly higher than those measured using VTIQ in 58 healthy testes which suggests that shear wave velocity values are not interchangeable across different commercial systems; a similar limitation has been found for the shear wave assessment of liver fibrosis.

Testicular Neoplastic Lesions

Strain Elastography

Intra-testicular lesions can be subdivided into malignant neoplasms, benign neoplasms, and non-neoplastic benign lesions. Testicular germ cell tumors account for over 95% of primary malignant testicular tumors, and they can be further divided into non-seminomatous or seminoma germ cell tumors, which comprise 60% and 40% of germ cell tumors, respectively. Testicular seminomas are typically well-defined, homogeneously hypoechoic lesions with no features of local invasion of the tunica albuginea on B-mode US. On strain elastograms, seminomas display a uniformly stiff nature (Fig. 2). Non-seminomatous germ cell tumors include embryonal cell carcinoma, teratoma, choriocarcinoma, and mixed tumors containing more than one histological cell type (Fig. 3). These tumors also display hard tissue stiffness but may have a less homogeneous elastogram map due to their heterogenous cellular structures. Sertoli and Leydig cell tumors are gonadal stromal tumors, which represent approximately 5% of all testicular neoplasms. The majority of Leydig cell tumors are benign neoplasms; only 10% of them have been reported to have a malignant potential on histological analysis [14]. Gonadal stromal cell tumors are often small, well-defined focal testicular lesions with reduced reflectivity and frequently show increased peripheral vascularity on color Doppler imaging. Their tissue stiffness can vary from mildly hard to hard on strain elastography (Fig. 4) [10,15]. However, their vascularity pattern has

Fig. 2. Two examples of testicular seminoma.
A. Incidental finding of a testicular seminoma in a 55-year-old man is shown. Strain elastography and B-mode images demonstrate a small lesion with uniform low reflectivity and uniformly hard tissue stiffness as displayed by the homogeneous blue color within the lesion (arrows).
B. A testicular seminoma in a 40-year-old man who presented with left testicular pain is shown. Real-time elastography and B-mode images demonstrate a large multiloculated mass (arrows). On strain elastography, the lesion demonstrates uniformly hard tissue stiffness.

Fig. 3. A mixed non-seminomatous germ cell tumor in a 38-year-old man who presented with a 2-week history of mild testicular discomfort. The B-mode ultrasound image demonstrates a large heterogeneous mass with cystic areas (arrows). On the strain elastography image, the lesion demonstrates predominant hard tissue stiffness, as shown in blue.
Elastography of focal testicular lesions

A handful of studies have investigated the ability of strain elastography to differentiate malignant neoplasms from benign neoplasms using VES and SR. Whilst hard elastograms, defined based on a VES of >3, demonstrate a high sensitivity (81.1%–82.3%) have been reported to differ from that of germ cell tumors, as gonadal stromal cell tumors show a significantly shorter filling time than germ cell tumors [15] and persist longer than the normal testicular parenchymal enhancement [16]. Primary lymphoma of the testis is rare, with a reported incidence of 0.09 per 100,000, but lymphoma represents the most common testicular malignancy in older men (age >60 years) [17]. In a cohort of six patients with histological confirmation of primary testicular lymphoma, all lesions appeared hard; five of six showed a VES of 5, and the other had a VES of 4 (Fig. 5) [18].

Testicular epidermoid and dermoid cysts are the most common benign neoplastic lesions and demonstrate hard elastographic properties [10,19,20]. They are often uniformly hard on strain elastography, with SR values often exceeding 40. However, it is the often-typical B-mode US feature of an “onion ring” configuration of alternating hypoechoic and hyperechoic rings and avascularity on color Doppler US that enable differentiation of epidermoid cysts from malignant lesions (Fig. 6). Testicular hemangioma is a benign rare testicular neoplasm that occurs in the same age group (second to fourth decades) in which primary germ cell tumors predominate (Fig. 7). Bernardo et al. [21] reported a case of capillary hemangioma that demonstrated relatively soft tissue stiffness on strain elastography, with a VES of 3 and an SR of 2.3.

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100%) in differentiating malignant testicular neoplasms from benign lesions, the specificity varies considerably among studies (25%–98.2%) [2,4,10,19,22]. In a cohort containing 30 testicular neoplasms, Pastore et al. [23] showed that SR provided adjunctive information about tumor characteristics along with B-mode and color Doppler imaging, aiding in the diagnosis of the neoplastic origin of testicular abnormalities. Pozza et al. [4] reported that higher SR values were found in malignant and neoplastic lesions than in benign and non-neoplastic lesions in a cohort comprising 144 testicular lesions. However, SR alone has a limited diagnostic ability to differentiate malignant from benign testicular lesions, with a reported sensitivity of 59.4% and specificity of 66.6% using a cut-off SR value of >1.41 for malignant lesions [4]. The elasticity score derived from a visual elastogram assessment was found to have a higher diagnostic accuracy (area under the curve [AUC], 0.804) than SR (AUC, 0.631) [4]. However, conflicting results were reported by Konstantatou et al. [10], who found that SR was a significantly better assessment than VES in diagnosing malignant lesions. In their cohort of 86 patients, an SR cut-off value of 3.21 was found to differentiate benign from malignant lesions with an area under the receiver operating characteristic curve of 0.722 [10]. Currently, only a single study has investigated the value of shear wave velocity in characterizing neoplastic testicular lesions [24].

Shear Wave Elastography
Unlike strain elastography, which is a qualitative assessment of tissue stiffness, shear wave elastography provides a quantitative assessment by measuring the shear wave velocity. A preliminary study involving 15 testicular germ cell tumors showed significantly higher shear wave velocities in seminomas (mean, 10.6 kPa; range, 4.5 to 15.8 kPa) than in non-seminomatous germ cell tumors (mean, 47.0 kPa; range, 29.0 to 65.5 kPa) [24]. The authors postulated that the difference in shear wave velocities may have been due to differences in histologic features between the seminomas and non-seminomas, as non-seminoma tumors consist of undifferentiated cells with a prominent stromal component, some of which may contain cartilage or bony components [24]. A cohort study by Rocher et al. described the multiparametric testicular US features of 10 burnt-out testicular tumors, for which much higher tissue stiffness was demonstrated compared to the surrounding tissue (13 kPa vs. 2 kPa) using shear wave elastography (Supersonic Imaging, Aix-en-Provence, France) [25]. Increased tissue stiffness could not distinguish burnt-out tumors from residual tumors. The diagnosis of a burnt-out tumor may be suspected in the presence no or very poor enhancement of the tissue on contrast-enhanced US. Currently, only a single study has investigated the value of shear wave velocity in characterizing neoplastic testicular lesions [24].

Testicular Non-neoplastic Lesions
Strain Elastography
A variety of non-neoplastic intratesticular abnormalities may present as focal intra-testicular abnormalities, causing a diagnostic conundrum. These indeterminate tumor-like lesions include focal

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**Fig. 6. A testicular epidermoid in a 15-year-old boy with history of microlithiasis.** On B-mode ultrasonography, the epidermoid has the typical sonographic features of an “onion ring” configuration with alternating hyperechoic and hypoechoic rings (arrows). The strain elastography color map shows that it was predominantly hard, with the blue color distributed in over 80% of the abnormality.

**Fig. 7. A testicular capillary haemangioma in a 60-year-old man.** The lesion appears relatively homogeneous with slightly increased echogenicity (arrows). The tissue stiffness color map suggests the lesion is relatively soft, with a predominantly green color mixed with some blue areas.
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abscesses, prominent retia testis, hematomas [3], segmental
testicular infarctions [26,27], venous infarctions [28], focal orchitis,
and sarcoidosis [16]. The majority of non-neoplastic testicular
lesions do not require surgical management, unlike the majority
of neoplastic lesions, for which orchidectomy or testicular-sparing
surgery is essential. Therefore, the ability to differentiate these
benign lesions from malignant lesions on imaging is of paramount
importance if orchidectomy is to be avoided.

Several cohort studies have tried to compare the tissue elasticity
of non-neoplastic lesions with that of neoplastic lesions. In
general, non-neoplastic lesions appear softer than neoplastic
lesions. Konstantatou et al. [10] showed that the median SR for
non-neoplastic lesions was 1.95, compared to 4.64 and 3.78 for
neoplastic malignant and neoplastic benign lesions. Aigner et al. [29]
showed that 100% of cases of orchitis (n=6), 75% of cysts (n=4)
and 80% of partial testicular infarctions (n=5) were soft on strain
elastography. Several case reports and case series have reported
strain elastography findings for non-neoplastic lesions, such as
segmental testicular infarctions, testicular hematomas [3], abscesses,
atypical retia testis, and cysts. Intra-testicular hematoma is often a
consequence of trauma, although upon presentation patients often
have no recollection of injury (Fig. 8). Yusuf et al. [3] evaluated a
cohort of 16 hematomas with strain elastography, 13 of which had
a VES of 3 and the reported mean SR was 1.19 (range, 0.41 to 2.36).
Intra-testicular hematomas are often soft lesions in a sub-capsular
position. Segmental testicular infarction is a rare cause of testicular
pain and discomfort. Its B-mode US and tissue elastography
features depend on the timing of sonographic examination after
the trauma. The area of the abnormality may initially show soft
characteristics due to the presence of swelling and increased water
content. As it evolves, the area may demonstrate hard features
on an elastography color map (Fig. 9). Focal orchitis may also
present as an indeterminate testicular lump. On B-mode US, it
appears as an ill-defined abnormality with reduced echogenicity.
Ancillary findings may include hydrocele or epididymitis. The area
of abnormality is often hypervascular on color Doppler. On strain
elastography, the abnormality initially appears as a hard lesion, but
as it evolves it becomes softer (Fig. 10). Testicular torsion leads to an
ischemic insult to the testes; it is a surgical emergency if it presents
acutely, and a sonographic investigation should not be attempted,
particularly if it causes delay in surgical exploration. However, US
has a role in differentiating delayed presentation of testicular torsion

Fig. 8. A testicular hematoma in a 30-year-old man who presented
with persistent right testicular pain. The lesion is oval-shaped with
low echogenicity located in the periphery of the testis (arrows). The
strain elastography color map shows a soft lesion, as indicated by
the green color in over 80% of the lesion with a scattered red color.

Fig. 9. Testicular segmental infarction in a 27-year-old man who presented with a 2-day history of testicular pain and discomfort.
A. On B-mode ultrasonography, at presentation, the lesion appears as a peripheral wedge-shaped area with slightly reduced echogenicity
(arrow). On a strain elastography color map, the lesion appears as a soft lesion, as indicated by the predominant green color (arrow). B. Two
weeks later, the lesion had evolved, becoming hard, as shown by the predominant blue color (arrows).
from other pathologies. By comparing the pathological evaluation of the surgical specimen with pre-operative testicular torsion at different time points and different degrees of torsion, Herek et al. [30] showed that testicular tissue stiffness gradually increased from 360° to 720° of torsion and became softer at 1,080°, where necrosis occurred and damage to the testes became irreversible. They also showed that the contralateral testis was also affected and presented increased tissue stiffness. Therefore, in cases of partial torsion where testicular viability and salvageability of the testis is in question, strain elastography of the testis may add additional benefit to a color Doppler US examination [30]. Non-neoplastic testicular lesions tend to be softer than their neoplastic counterparts. However, Konstantaou et al. [10] found that 16 of 38 lesions had a VES of 4 (n=4), 5 (n=7), or 6 (n=5), indicating there is significant overlap between the tissue stiffness of neoplastic and non-neoplastic lesions.

Shear Wave Elastography
A shear wave elasticity value of 1.7 kPa using the SuperSonic Imagine apparatus (Aixplorer Provence) was described in a case report of a segmental testicular infarction, compared to an elasticity value of 2.6 kPa in the adjacent normal testicular parenchyma [31]. A study of 248 patients including 130 with normal testicular tissue, 99 with microlithiasis, and 19 with testicular cancer showed that the mean shear wave velocity from three measurements taken from the centre, upper pole, and lower pole of the testis or testicular lesion was significantly higher in patients with testicular cancer (mean velocity, 1.92 m/sec) than in those with normal testicular tissue (0.76 m/sec) or testicular microlithiasis (0.79 m/sec) [8].

Conclusion
Strain or shear wave elastography allows further characterization of tissue stiffness on US. It has been shown that neoplastic lesions are harder than non-neoplastic lesions, and malignant neoplastic lesions are harder than neoplastic benign lesions. However, significant overlap exists between types of lesions. Assessment of the B-mode US appearance, lesion vasculature with color Doppler US, and contrast-enhanced US in addition to tissue elastography as part of a multiparametric ultrasound examination [32] is indispensable for increasing diagnostic accuracy and confidence when placing indeterninate lesions under surveillance or subjecting the patient to surgery [16].

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Conceptualization: Fang C, Huang DY, Sidhu PS. Data acquisition: Fang C, Huang DY, Sidhu PS. Data analysis or interpretation: Fang C, Huang DY, Sidhu PS. Drafting of the manuscript: Fang C, Huang DY, Sidhu PS. Critical revision of the manuscript: Fang C, Huang DY, Sidhu PS. Approval of the final version of the manuscript: all authors.

Conflict of Interest
No potential conflict of interest relevant to this article was reported.
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2017;36:821-831.