General advice in ultrasound based elastography of pediatric patients

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Abstract

Ultrasound elastography including transient elastography (TE), point shear wave elastography, (pSWE) and two (three)-dimensional shear wave elastography (2D-SWE) have been introduced mainly for the evaluation of the liver. All the techniques are also feasible for the examination of spleen, whereas pSWE and 2D-SWE can be used for the assessment of the pancreas, kidney, gastrointestinal tract and other organs. Strain elastography also plays a role for non-liver applications. The aim of the current report is to highlight unique features and techniques for the elastographic examinations in children and to report initial results in non-liver applications.

Keywords: ultrasound; elastography; pediatric; shear wave elastography (SWE)

Introduction

Ultrasound (US) elastography was introduced into clinical routine more than twenty years ago. Guidelines were published several years later reflecting the updated knowledge of evidence based medicine [1-7]. The most recent update of Guidelines and Recommendations on the Clinical Use of Elastography of the European Federation of Societies for Ultrasound in Medicine and Biology (EFSUMB) and World Federation for Ultrasound in Medicine and Biology (WFUMB) focused on the assessment of diffuse liver disease in adults [6-10]. Compared to adults, fewer publications are available regarding the technical aspects and indications for the use of shear wave elastography in children [11]. The guidelines on adults may not be immediately transferrable to children. In this review we discuss the current evidence on the use of US elastography examination techniques in pediatric patients.
What is different in liver elastography in children compared to adults?

An invasive procedure such as liver biopsy is less accepted in children due to the need for general anesthesia, risk of complications and the physical and emotional impact on the child. Non-invasive methods are more suitable, particularly when there is a need for repeated examinations, as in the follow up of patients with chronic liver disease. Elastography is one such method that is feasible and accurate for the assessment of liver fibrosis in adults. However, children are different from adults also in terms of the disease spectrum. In addition, several factors such as age, breath-holding, and probe choice should be taken into consideration regarding the examination technique in children, since they can influence the measurement reproducibility and the diagnostic accuracy in the pediatric population.

Physiological and technical factors

Age is an important factor as there are physiological differences across pediatric age groups. For instance, in infants the liver is situated lower in the abdomen, with the inferior margin 3-3.5 cm below the costal rim; therefore, the location for measuring liver stiffness may be different. Additionally, younger children are less likely to cooperate compared to an adult in terms of keeping still and holding the breath during the measurement of liver stiffness. They may also not have the patience for the required number of measurements. In a study that measured liver stiffness by transient elastography (TE) in 240 healthy children, Engelmann et al reported 27% of invalid liver stiffness measurements in children below 6 year-old compared to 9% in older children [12]. Although breath-holding may affect the technical success rate in children, free breathing has been shown not to affect the variability and agreement of the results [13]. Deep sedation may overcome these technical challenges; however, it can be associated with other adverse effects and it is less accepted by parents. The use of sedation cancels also one of the advantages of US over CT or MR, i.e. the need for sedation with these latter techniques. There are conflicting results in terms of whether normal liver stiffness values differ across different age groups. Normal values of liver stiffness by TE are significantly different among different age groups (0-5, 6-11, 12-18 years) [12].

Although gender has not been shown to influence liver stiffness values in adults, it might have an impact in adolescence, during which there is a "normal" insulin resistance and a higher adiposity in girls. Regarding BMI, country based/WHO growth charts are needed to establish underweight/overweight/obesity status. Dillman et al showed that higher BMI may affect the success rate of liver elastography examination in children [14]. Increased BMI has also been shown to contribute to the likelihood of increased data variability (i.e., increased interquartile range over median) [15].

Finally, several technical points regarding the measurement acquisition should be highlighted. As reported in the adults, significant difference in shear wave velocities were observed between the right and left liver lobe in healthy children [16]. The location for measurement in children has not been established. In adults, measurements in the right liver lobe through intercostal spaces are more reproducible than those in the left liver lobe [17]. Canas et al compared the diagnostic accuracy of liver stiffness measurement by point shear wave elastography (pSWE) (Virtual Touch Quantification – VTQ) in right and left liver lobe to diagnose cystic fibrosis liver disease and found that measurements in the right lobe through the intercostal spaces were significantly more accurate (AUROC 0.746 vs 0.529) [18b]. However, measurements in the right liver may not be possible in children with left lateral liver transplant.

The selection of the probe should be based on anthropometric measurements rather than on age, since patients can be underweight or undeveloped for their age. There are limited and conflicting data regarding the different values obtained with the linear versus the convex probe. In one study evaluating pSWE (VTQ), Fontanilla et al found lower velocity values when using the linear (1.15±0.04 m/s) versus the convex probe (1.19±0.04 m/s), and recommended using the linear probe in children below the age of 12 months 28. For two dimensional(2D) SWE using SuperSonic Shear Imaging (SSI) values, Pienar et al found stiffness values higher with the linear probe than with the convex probe (7.8±5.1 vs 4.1±0.9 kPa) [19], whereas Franchi-Abella et al, in their pilot study on 96 children including neonates, found that liver stiffness values were significantly higher when a curvilinear 6-1 MHz transducer was used compared with a linear 15-4 MHz transducer. Amon the 51 healthy children who served as controls, average liver stiffness assessed by 2D-SWE were 5.96+/-.1.31 kPa for the SL15-4 probe and 6.94+/-.1.42 kPa for the SC6-1 probe (p=0.006) [20].

The etiologies of diseases

Although causes of chronic liver diseases in adult such as viral hepatitis (B, C, D), alcohol, NASH, and autoimmune hepatitis are also common etiologies in children, specific etiologies are more likely first detected in the pediatric population such as Cystic Fibrosis Associated Liver Disease (CFLD), metabolic liver diseases, Gaucher’s disease, Wilson’s disease, α1 antitrypsin deficiency, glycogen storage disease, Niemann-Pick disease, etc. and biliary diseases (biliary atresia, primary sclerosing...
In all these etiologies, US-based liver elastography is indicated for liver fibrosis assessment and follow-up. In addition, it can also be used to assess graft fibrosis after liver transplantation. The mechanisms leading to chronic liver disease in patients with congenital heart disease have been increasingly recognized recently. Corrective surgery, such as the Fontan procedure is associated with progressive hepatic failure and even hepatocellular carcinoma [21]. US-based elastography may be used to diagnose early fibrosis so that cardiac interventions to alleviate hepatic congestion can be attempted.

**General technical advice**

**Examination technique**

SWE (including TE, pSWE and 2D-SWE) should be performed in a supine position, with the right arm of the patient in extension to increase the width of the intercostal space. The transducer is positioned in a right intercostal space, to visualize the right liver lobe during breath hold, avoiding deep inspiration prior to the breath hold [1,5]. A transient breath hold in a neutral position is ideal [6,7]. There is little published data on children comparing liver stiffness measurement performed with breath hold and free breathing techniques. Breath hold minimizes measurement variability; however, although breath holding is feasible in older children, in children below the age of 5 years, it is almost impossible. It has been reported that 2D-SWE performed with free-breathing yields results similar to those obtained with breath-holding [20]; thus,

![Image](image1.png)

**Fig 1.** For infants, some tips are a quiet, pediatric friendly environment, positioning them in caregiver’s arms, use of appropriate age toys for settling (six-month-old boy).

Liver stiffness values can be expressed either in the unit of the Young modulus (kilopascals, kPa) when using TE, or in the units of the speed (meters/second, m/s) and kPa when using pSWE or 2D-SWE, taking into account that the shear wave speed, directly measured by the software in the equipment, is converted to the Young modulus units making some assumptions [1].

**Fasting and resting**

In adults, it has been shown that food ingestion increases liver stiffness readings for an estimated 2-3 hours after ingestion [31-33]. Exercise also increases liver stiffness [6,7,34]. Measurements should be performed in fasting conditions, ideally overnight and after at least 10 minutes of rest [6,7]. However, in babies breastfeeding should not be interrupted, because it causes distress both to the mother and the baby. In bottle fed children, the examination should be performed before the child’s next meal, if possible.
Factors influencing liver stiffness independent of liver fibrosis (confounders)

There are many confounding factors that need to be taken into account [6,8,35-47]. In summary, liver stiffness increases with hepatic necro-inflammation (often, but not exclusively shown by an elevated transaminase level >5 times the upper limit of normal), obstructive cholestasis, hepatic congestion, acute hepatitis and infiltrative liver diseases. In addition, general anaesthesia significantly increases liver stiffness in healthy children [11]. Probe choice equally influenced results in paired comparisons, as did food intake and intercurrent, non-hepatological illnesses [48]. It has been reported that, in children with post-transplant liver and without fibrosis, liver stiffness values were significantly higher than those of healthy children with native liver [49]. pSWE measurements show higher reproducibility and minimal data dispersion in patients with a BMI <30 kg/m² compared to the values in patients with a higher BMI [15]. Shear wave velocities by VTQ differed between normal-weight and obese children (1.08 ± 0.14 vs 1.44 ± 0.39 m/s; p<0.001), but were not affected by gender. Multivariate linear regression demonstrated shear wave velocity to be primarily associated with age in normal-weight children (p<0.05) and with BMI in obese children (p<0.001). In the obese group, mean shear wave velocity was statistically higher in children with abnormal echogenic livers than in those with normal-appearing livers (1.53 ± 0.38 vs 1.17 ± 0.27). The difference was not significant in the normal-weight group [50].

Comparison of results between systems in pediatric patients

There are several sources of variability including technical and patient dependent factors that could affect the comparability between systems. TE is technically feasible in children of all age groups. The upper limit of normal values increases significantly with age. Median values of stiffness are significantly age dependent, while the interquartile range decreased with age [48,49]. pSWE was feasible in children at any age with an acceptable reliability. The depth of measurements in the liver had no influence on the results. There was no statistical difference between measurements taken at different ages [51,52]. Conversely, in another study pSWE speed was primarily associated with age, BMI, but not with gender [50]. Other important factors are the transducer’s frequency and the operator experience [53-57]. Proprietary elastographic technologies can give different estimates of the shear wave speed within the same liver. Thus, threshold values for fibrosis stages are needed for each specific US system. In addition, studies have also shown that liver stiffness values taken from different commercial machines produce different values. Therefore, values taken from different commercial machines are not interchangeable and follow-up studies should be performed using the same machine.

Other applications of elastography

Spleen

Conventional US is sensitive in detecting splenic abnormalities such as asplenia, focal (neoplasms, infection etc.) or diffuse (infection, splenomegaly) abnormalities. Splenomegaly can be due to extrasplenic or splenic causes, both causes can induce microscopical changes which lead to changes in spleen elasticity that can be detected with elastography. In order to achieve the most reliable results an optimal setting is essential. During the spleen elastography examination, the patient should be placed in a supine position with the left arm in maximal abduction. The transducer should then be placed on the left intercostal space to provide an adequate acoustic window. It is advisable that the patients hold their breath. This might not be possible in small children; however studies have shown that valid results can be obtained also with free breathing [58]. Convex or linear US probes can be used; the convex probes show higher variability [58,59].

The normal values for children using pSWE (VTQ) are around 2.25 m/s [60]. Also, the size of the spleen has an influence on the stiffness values [58,61]. Obesity may reduce the quality of the B-mode US image, but this has not yet been investigated in studies.

Pathological changes of spleen stiffness are seen in patients with liver diseases, systemic diseases and diseases of the bone marrow. Several studies, mostly performed in adults, have shown that increased spleen stiffness is correlated with portal hypertension. In fact, spleen stiffness is more reliable than liver stiffness in detecting variceal bleeding risk in adult patients with clinically significant portal hypertension [62]. Interestingly, spleen stiffness is increased in patients with non-cirrhotic portal hypertension (extrahepatic portal vein obstruction and idiopathic portal hypertension) in whom liver stiffness is normal or only mildly elevated [63,64]. This suggests that spleen stiffness and the ratio between liver and spleen stiffness could be used in patients with portal hypertension of unknown origin to help differentiating between cirrhotic and non-cirrhotic causes.

Spleen stiffness value correlates with the stages of liver fibrosis [65]. A study for the prediction of significant varices in children indicated that the TE cut-off value was 38 kPa [61]. Uchida et al suggested that spleen stiffness could also play a role in selecting children with biliary atresia after the Kasai procedure for liver trans-
plant [66]. In general, systemic diseases that cause splenic inflammation may increase spleen stiffness.

**Thyroid**

The adult thyroid has been extensively examined by US techniques [9,67-72] but little is known for the use of US elastography in pediatric patients [73].

**Healthy thyroid gland**

The mean SWV by pSWE (VTQ) of the normal thyroid gland was 1.22±0.20 m/s. There was no correlation between age and the mean SWV (Spearman Rho=0.049, p=0.556). There was also no correlation between the thyroid gland volume or the whole body surface area and the mean SWV. The only correlation detected was between body surface area and total thyroid gland volume (p<0.001) [74].

**Hashimoto’s thyroiditis (HT) and Graves’ disease**

The mean elastographic strain index (SI) of adult healthy individuals and patients with HT was 0.26±0.77 and 1.75±1.46, respectively (p<0.001). For the diagnosis of HT with strain elastography (SE), the sensitivity was 92.1%, and specificity was 66% when the optimal mean SI cut-off value was 0.31, (p<0.001). The use of the SI in patients with HT was highly promising for objective and countable results compared with conventional US. The sensitivity of the increase in SI was determined to be high for the diagnosis of HT. It was also found that the SI had higher sensitivity and specificity than the conventional US in HT patients with moderate to advanced tissue hardening [75]. There is not any published study on the use of elastography in Graves’ disease but personal experience shows homogenous intermediate stiffness (fig 2).

**Type 1 diabetes mellitus and thyroid gland**

Type 1 diabetes mellitus (T1D) affects the thyroid gland stiffness even in patients without autoimmune thyroiditis. The mean SWV in the thyroid gland by pSWE (VTQ) in T1D patients (1.11±0.21 m/s) was lower than that in the control group (1.29±0.23 m/s) and this difference was statistically significant. It has been suggested that pSWE may be a useful method in determining early changes in the thyroid gland in pediatric T1D and may be used as a screening tool [76].

**Thyroid gland neoplasia**

Almost all papillary carcinomas are stiff whereas follicular carcinomas may appear soft on elastography [67,68,70,71]. All other thyroid malignancies are stiffer than the surrounding tissue. Lymph node metastases are stiff as well [77,78] (fig 3).

**Myocardial stiffness**

A pilot study was conducted with the aim to systematically investigate the feasibility of using cardiac SWE in children, and to provide myocardial stiffness control data for cancer patients. The parasternal long-axis (L-A) and short-axis (S-A) views of the interventricular septum (IVS) were feasible for pediatric cardiac SWE assessment. The L-A and S-A views of the basal and mid IVS provided better success rates than that of the apical IVS. Two-dimensional SWV measurements were 1.26, 1.22, 1.71 and 1.67 m/s for L-A base, L-A mid, S-A base and S-A mid IVS, respectively, in healthy children [79].

**Muscle stiffness**

pSWE elastography for identifying structural changes, that occur in the spastic muscle after botulinum toxin A injection in children with cerebral palsy, can yield more valuable information with combined use of the modified Ashworth scale [80]. The normal abdominal layers and muscles are shown in figure 4.
Renal parenchyma

The main causes of chronic kidney disease (CKD) in the pediatric population are renal hypoplasia, dysplasia or an association of both conditions (57.5% of cases) and vesicoureteral reflux (25.8%). pSWE (VTQ) values decrease from kidneys with secondary vesicoureteral reflux (6.59±1.45), to kidneys with primary reflux (5.3±1.72), to unaffected kidneys contralateral to the reflux (4.09±0.97) and to normal kidneys (3.13±0.09) [81].

pSWE seems to have some advantages in predicting pediatric glomerular disease compared to conventional US. Comparisons of SWV measurements between left and right kidneys in the diseased and control groups all showed significant differences. In this case SWV assessment contributed to the early diagnosis of the disease [82].

Gastrointestinal tract

The gastrointestinal tract has been extensively examined by US techniques in children [77,78,83-86], including the B mode US and color flow imaging. Elastography might play a role in assessing inflammation, fibrosis and infiltration. A correct diagnosis can often only be made by colonoscopy with biopsies or surgery. Elastography would save pediatric patients from invasive procedures in uncertain cases with unclear digestive symptoms [87,88].

Considering that elastography is relatively new in this field, there are currently very few pediatric studies for reference. The latest studies in adults have shown promising ways of applying elastography in Crohn’s disease [83,84,89]. In order to have comparable values, it is unavoidable to have a standardized setup. There are different kinds of settings, where measurements can be done transdermally or in a trans-rectal endoscopic setting [53,58,90]. EUS elastography offers a high resolution of the digestive tract, close view of organs and lymph nodes [91,92]. Elastography techniques have been used in different ways in different studies; dynamic elastography shows promising results while SE has a lower cost, which makes this technique more accessible [73]. The general approach used in normal bowel ultrasonography can also be applied to elastography of the bowel. Elastography should be performed on a patient placed in supine position, with relaxed abdominal muscles. For the study of specific regions, linear probes with high frequencies (7-11 MHz) are recommended [73]. Although it is not necessary in all cases, fasting for 4-6 h decreases bowel motility. A cup of water might be helpful for the visualization of the duodenum, as well as 250-800 ml polyethylene glycol solution for the terminal ileum, which can be seen after 30 minutes [93].

The diagnosis and monitoring of Crohn’s disease represent a diagnostic challenge in which imaging plays an important role [94]. In a pilot study on 48 bowel segments of 14 pediatric patients with Crohn’s disease, Fufezan et al showed the reproducibility of the elastography technique [95]. In addition, a study on adult patients with and without appendicitis revealed the benefits of elastography (2D-SWE, SSI). Patients with appendicitis were identified with a sensitivity of 93%, specificity of 100% considering a 12.5 kPa cut-off value [96]. A statistically significant correlation was found between the intestinal wall hydrosonography changes, presence of complications, activity markers and SE score. SE, along with hydrosonography, represented a reliable investigation for the correct diagnosis and monitoring of pediatric patients with Crohn’s disease and the SE score can be used for the assessment of disease activity [95]. Another possible application of elastography is to differentiate colon adenomas from adenocarcinomas. Although adenocarcinomas are rare in the pediatric population, they represent 1% of all childhood neoplasms [97].

The most important error in SE is due to peristaltic bowel movements, which may give false results. Modern software tries to cancel these errors with correction filters, but a complete success is still not guaranteed [3,94]. At this point in time there are no normal values for these diseases in children. Therefore, further efforts should be made to provide noninvasive diagnostics to the pediatric population.

Postoperative undescended testicles

2D-SWE was used to determine the testicular volume and elasticity changes in young children with unde-
scended testes [98,99]. pSWE (VTQ) was used to assess the difference in stiffness between scrotally placed testes and postoperative undescended testicles. The SWVs of undescended testicles were 0.75-2.8 (median, 1.1) m/s, whereas they were 0.65-1 (median, 0.82) m/s in healthy controls [100].

**Neonatal brain**

Some early reports on the use of transcranial SWE of the periventricular brain parenchyma, in preterm infants and infants with hydrocephalus, suggest that SWE is possible and technically feasible [101,102] (fig 5, fig 6). Albayrak et al showed that differences between brain stiffness values in preterm and term neonates can be demonstrated by using 2D-SWE. Brain stiffness measured from both the thalamus and periventricular white matter were found to be significantly lower in preterm neonates compared with term neonates (cut-off values for determining prematurity less than 8.28 kPa for mean thalamus stiffness and less than 6.59 kPa for periventricular white matter stiffness). The authors suggested that the results might be reference points for evaluating neonatal brain stiffness in research on patients with various illnesses. 2D-SWE also seems to have the ability to depict increased intracranial pressure (ICP) in infants, with a positive linear correlation between SWE values and ICP [102]. Infants with ICP seem to have increased 2D-SWE values (mean 24.2±5.1 kPa) compared to healthy infants (mean 14.1±6.6 kPa). However, larger prospective studies are still not available. If these preliminary observations of the benefits of transcranial SWE of the neonatal brain will be confirmed by further studies, SWE might be a useful method for additional diagnostic imaging and monitoring in premature infants and children with proven or suspected increased ICP. When performing SWE of the neonatal brain, potential risks and harms of applying high energy levels by US to the neonatal brain should be considered. Recently, an experimental study on mice dealing with the potential biological effects associated with 2D-SWE on the neonatal brain was published [103]. The results indicated that 2D-SWE does not cause detectable histologic changes in the brain of neonatal mice, nor does it have long-term effects on the learning and memory abilities. However, some temporary effects were observed when the scanning lasted for more than 30 min. Thus, it is recommended to pay attention to the scanning duration when assessing neonatal brains with 2D-SWE elastography.

**Investigator training**

The examiner should acquire appropriate knowledge and training in US elastography [104,105]. The operator must distinguish a good B mode US image from suboptimal images. In fact, for pSWE and 2D-SWE experience in B-mode US is mandatory. Data acquisition should be undertaken by specialists. Operators experienced both in ultrasonography and elastography are needed to obtain reliable liver stiffness measurements in children, considering the different anatomy, especially in babies (liver situated lower in the abdomen), and the fact that cooperation from a small child is sometimes difficult. The location for measurements can be more difficult to establish in children and here the operator’s experience can play a role.

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References


