Retrospective Study

Non-invasive splenic parameters of portal hypertension: Assessment and utility

Ayesha Karim Ahmad, Sebastiana Atzori, James Maurice, Simon D Taylor-Robinson, Adrian KP Lim

ORCID number: Ayesha Karim Ahmad 0000-0003-3763-1212; Sebastiana Atzori 0000-0002-4444-2095; James Maurice 0000-0003-1530-5328; Simon D Taylor-Robinson 0000-0002-8811-1834; Adrian KP Lim 0000-0003-3904-2876.

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Abstract

BACKGROUND
Portal hypertension is a major complication of cirrhosis that is associated with significant morbidity and mortality. The present gold-standard method to risk stratify and observe cirrhosis patients with portal hypertension is hepatic venous pressure gradient measurement or esophagogastroduodenoscopy. However, these methods are invasive, carry a risk of complications and are associated with significant patient discomfort. Therefore, non-invasive splenic parameters are of clinical interest as potential useful markers in determining the presence of portal hypertension. However, diagnostic accuracy and reproducibility remains unvalidated.

AIM
To assess the diagnostic accuracy of spleen stiffness, area and diameter in predicting the presence of portal hypertension.

METHODS
Of 50 patients with varying liver disease pathologies were prospectively recruited from the St. Mary’s Hospital Liver Unit in London; 25 with evidence of portal hypertension and 25 with no evidence of portal hypertension. Liver stiffness, spleen stiffness, spleen diameter and spleen area were measured using the Philips Affiniti 70 elastography point quantification point shear wave elastography system. The aspartate aminotransferase-to-platelet-ratio-index (APRI) score was also calculated. Performance measures, univariate and multivariate logistic
INTRODUCTION

Portal hypertension (PH) is a major complication of cirrhosis\(^1\). Esophageal varices (EV) are present in 40% of compensated advanced chronic liver disease (cACLD) patients and in 70% of decompensated cirrhosis patients. Strategies to identify individuals with clinically significant portal hypertension (CSPH) is vital to reduce morbidity and mortality\(^2\).

The hepatic venous pressure gradient measurement (HVPG) and esophageal-gastrooduodenoscopy (EGD) form the backbone of diagnosis and surveillance of EV\(^3\). However, both methods are invasive and carry a risk of complications\(^4\). Therefore, non-invasive and safe methods of diagnosis and surveillance of PH are of great clinical interest.

Current guidelines propose that non-invasive methods of assessment of liver fibrosis can predict the incidence of cirrhosis-induced PH manifestations\(^6\). The Baveno VI guidelines suggest cirrhosis patients with a liver stiffness measurement < 20 kPa and a platelet count > 150000/μL can avoid screening endoscopy\(^7\). Nevertheless, while 20% of EGDs are spared, new algorithms are still required, as up to 40% of EGDs continue unnecessarily\(^8\).

Ultrasound elastography techniques are based on the principle that tissue elastic...
properties can be distorted using shear waves to measure stiffness. Spleen stiffness measurements using ultrasound elastography have shown an association with CSPH as the spleen undergoes parenchymal remodelling and fibrogenesis, due to blood pooling in PH. Interestingly, evidence on patients with chronic hepatitis C infection also suggests that spleen stiffness is dependent on inflammation present in the liver that directly contributes to the pathogenic mechanisms underlying PH.

Transient elastography (TE) is the most validated ultrasound elastography technique and shows a sensitivity ≥ 90% in detecting patients with CSPH. However, limitations exist due to its lack of 2D imaging guidance and attenuation of wave propagation in obesity and ascites.

Point shear wave elastography (p-SWE), often referred to as acoustic radiation force impulse, overcomes these issues by providing integrated 2D-ultrasound imaging which can be used in patients who are obese or have ascites. Despite several meta-analyses on spleen stiffness measurements, it remains unclear whether TE or p-SWE has greater diagnostic accuracy. Furthermore, although it is well established that spleen stiffness and combination variables such as liver stiffness-spleen diameter to platelet ratio (LSPS) score are superior to liver stiffness for detection of EV, little is known whether a combination of splenic parameters can improve diagnostic accuracy.

Finally, although two main p-SWE techniques exist—the elastography point quantification (ElastPQ) and Virtual Touch Quantification (VTQ)—fewer studies have looked at the performance of ElastPQ due to its novelty.

We aimed to assess whether spleen stiffness measurement, spleen area and spleen diameter can independently predict CSPH, or in combination with other biochemical or elastography parameters, and assess reproducibility of splenic area and diameter measurements.

MATERIALS AND METHODS

Study design

Patients with varying liver disease etiology were prospectively recruited as part of an ongoing comparative imaging study (REC 15/EE/0420). All subjects had evidence of chronic liver disease (CLD), were over the age of 18 and provided informed consent. Exclusion criteria included pregnancy, lack of liver disease pathology, transjugular portosystemic shunt (TIPSS) insertion or presence of hepatocellular carcinoma (HCC).

The primary analyses were conducted after all patients were recruited. The patients were divided into the following groups: Evidence of CSPH (group 1) and no evidence of CSPH (group 2). CSPH was defined either as presence of EV or portal hypertensive gastropathy (PHG) during an EGD or if patients had invasive procedures where the HVPG pressure ≥ 10 mmHg. Ultrasound elastography measurements must have been undertaken within a maximum of one year of EGD or HVPG measurements.

Ultrasound and elastography

All patients had to be fasted for up to 6h prior to scans. Participants were placed supine with arms abducted away from the ultrasound probes. The Philips Affiniti 70 (ElastPQ) (Philips Medical Systems, Seattle, WA, United States) was used to record liver stiffness measurement and spleen stiffness measurement for each patient. Ten measurements were taken from the liver and ten measurements from the spleen. Liver elastography measurements were taken from the right lobe of the liver 2.4 cm (± 1 cm) from the liver capsule. Spleen elastography measurements were taken from the middle aspect of the spleen with homogeneous elasticity with the exclusion of big vessels. The median stiffness and IQR values were recorded. Spleen area and diameter were calculated from 2D images obtained.

Clinical and biological parameters including body mass index (BMI), skin to liver capsule distance, aspartate aminotransferase (AST), alanine aminotransferase (ALT), γ-glutamyl transpeptidase (GGT), alkaline phosphatase (ALP), platelet count, prothrombin time, albumin, bilirubin and international normalized ratio (INR) were obtained for all patients at time of recruitment. APRI score was calculated as: AST (IU/L)/PLT (× 10^9/L)^10. Parameters determining presence of PH such as HVPG measurements or EGD findings were recorded. Cirrhosis was defined either by histological findings at biopsy or if decompensation had occurred.

Spleen area and spleen diameter measurements

Spleen area and diameter measurements were calculated using maximum spleen diameter and borders that included the splenic hilum in the transverse plane with the
area (Figure 1). Measurements were repeated 4 mo later by two authors independently using a random sample of 19 study patients to calculate inter-operator variabilities.

**Statistical analyses**

Descriptive statistics were carried out to compare groups 1 and 2. Ultrasound measurements, BMI and laboratory results were analysed by univariate and multivariate analyses. Correlations between variables were examined using Pearson’s correlation coefficient and P values determined using ANOVA. A multivariate logistic regression model was built using a stepwise selection to determine the association of spleen area and platelet count and spleen stiffness and platelet count with the presence of CSPH. It was ensured that the data fulfilled all necessary criteria prior to application of the logistic regression analysis. Diagnostic accuracy was assessed using receiver operating characteristic (ROC) curves. Youden’s index was used to determine the cut-off values for each parameter. P values for ROC curves were identified based on Wilcoxon’s test. As subjects were random patients and operators were fixed, a one-way random interclass correlation coefficient (ICC) model on absolute agreement to determine inter-operator variability for spleen area and diameter was carried out. P values < 0.05 were considered statistically significant. All statistical analyses were performed using SPSS version 24.0.

**Ethics**

This study was performed in accordance with the 1975 Declaration of Helsinki. Written and informed consent was obtained from all patients.

**RESULTS**

Fifty four of 155 patients recruited had an EGD/HVPG measurement taken within one year of ElastPQ measurements. Four patients were excluded as summarized in Figure 2. A total of 50 patients (mean age 57.86, 62.0% male) were included in final analysis: 25 with evidence of CSPH (group 1) (mean age 60.44, 60.0% male) and 25 with no evidence of CSPH (group 2) (mean age 55.28, 64.0% male). The median time difference between ultrasound elastography measurements and EGD/HVPG was 4 mo.

Baseline clinical and biochemical characteristics for all patients included in statistical analysis are summarized in Table 1. Patients with diagnosis of cirrhosis were found in both groups n = 25 in group 1; n = 11 in group 2) with majority classified as Child-Pugh A (n = 18, 36.0%). The most common primary etiology in group 1 was alcoholic liver disease n = 7, 28.0%), while non-alcoholic fatty liver disease (n = 8, 32.0%) was more common in group 2.

**Univariate analysis**

We hypothesized that clinical parameters associated with CLD may predict the presence of CSPH. The clinical parameters tested were BMI, ALT, AST, GGT, ALP, bilirubin, platelet count, albumin, prothrombin time, APRI score, liver stiffness, spleen stiffness, spleen area and spleen diameter. The univariate analysis showed that bilirubin, platelet count, albumin, prothrombin time, APRI score, liver stiffness, spleen area and diameter correlated with the presence of CSPH (Table 2). The best individual predictor of CSPH was platelet count (AUROC 0.846, P value < 0.001), followed by spleen area (AUROC 0.828, P value = 0.002) and APRI score (AUROC 0.827, P value < 0.001). No statistically significant discrimination was found between liver stiffness measured by the ElastPQ and CSPH (AUROC 0.657, P value = 0.061).

**Multivariate analysis**

A multiple logistic regression model showed that two combinations independently predict CSPH (Table 3). The combination with the greatest diagnostic accuracy revealed that patients with a combination of spleen area > 57.9 cm² and platelet count < 126 × 10⁹ produced an estimated area under receiver operating characteristic (AUROC) curve of 0.876 (P value < 0.001), with sensitivity of 63.2%, specificity of 100%, positive predictive value (PPV) of 100% and a negative predictive value (NPV) of 61.1%. An alternative combination of spleen stiffness > 29.99 kPa and platelet count < 126 × 10⁹ displayed a similar diagnostic accuracy with an estimated AUROC of 0.855 (P value < 0.001) and sensitivity of 88%, specificity of 75%, PPV of 78.6% and NPV of 85.7%. AUROC curves are displayed in Figure 3.
Table 1 Baseline characteristics of the patient population

| Parameter                          | Groups 1 and 2 (n = 50) | Group 1 (n = 25) | Group 2 (n = 25) |
|------------------------------------|-------------------------|------------------|------------------|
| Age                                | 57.86 (12.22)           | 60.44 (9.89)     | 55.28 (13.90)    |
| Sex (M:F)                          | 31:19                   | 15:10            | 16:9             |
| BMI                                | 29.42 (7.61)            | 28.59 (5.76)     | 30.31 (9.26)     |
| Skin to liver capsule Distance (cm)| 2.43 (0.99)             | 2.46 (1.01)      | 2.40 (0.99)      |
| METAVIR score                      | 3.13 (1.276)            | 3.74 (0.752)     | 2.52 (1.410)     |
| ALD                                | 12                      | 7                | 5                |
| NAFLD                              | 12                      | 4                | 8                |
| ALD and NAFLD                      | 4                       | 3                | 1                |
| HCV                                | 2                       | 2                | 0                |
| AIH                                | 3                       | 0                | 3                |
| Miscellaneous                      | 17                      | 9                | 9                |
| Total bilirubin (μmol/L)           | 28.00 (44.728)          | 39.92 (60.139)   | 16.08 (13.108)   |
| ALP (IU/L)                         | 127.33 (66.902)         | 144.56 (75.900)  | 109.38 (51.679)  |
| GGT (IU/L)                         | 147.68 (192.683)        | 160.96 (166.671) | 134.96 (217.570) |
| ALT (IU/L)                         | 60.84 (12.219)          | 48.52 (29.427)   | 73.16 (83.490)   |
| AST (IU/L)                         | 62.50 (49.043)          | 64.88 (32.720)   | 60.12 (61.872)   |
| Albumin (g/L)                      | 34.10 (6.129)           | 30.88 (5.761)    | 37.32 (4.679)    |
| Platelet count (× 10^9)            | 151.42 (73.016)         | 112.16 (60.023)  | 190.68 (63.804)  |
| Prothrombin time (sec)             | 12.572 (1.9886)         | 13.470 (2.2077)  | 11.540 (1.0007)  |
| INR                                | 1.184 (0.1742)          | 1.260 (0.1871)   | 1.108 (0.1222)   |
| EGD (PH present)                   | 25                      | 25               | 0                |
| HVPG (PH present)                  | 5                       | 5                | 0                |

Group 1: Population with evidence of portal hypertension; Group 2: Population with no evidence of portal hypertension. BMI: Body mass index; ALD: Aldosterone; NAFLD: Non-alcoholic fatty liver disease; HCV: Hepatitis C virus; AIH: Autoimmune hepatitis; ALP: Alkaline phosphatase; GGT: γ-glutamyl transpeptidase; ALT: Alanine aminotransferase; AST: Aspartate aminotransferase; INR: International normalized ratio; EGD: Esophagogastroduodenoscopy; HVPG: Hepatic venous pressure gradient; PH: Portal hypertension.

Inter-observer variability for spleen area and spleen diameter measurements

An estimated single measures one-way random ICC for inter-operator variability for spleen area generated a value of 0.98 (95%CI: 0.94-0.99, P value < 0.001). Similarly, spleen diameter generated a value of 0.96 (95%CI: 0.91-0.99, P value < 0.001). Table 4 outlines the inter-operator ICC values for spleen area and diameter.

DISCUSSION

The present study aimed to assess the performance of non-invasive splenic parameters using a new generation p-SWE machine—the ElastPQ—in identifying the presence of CSPH. We demonstrated that spleen stiffness (AUROC 0.712), spleen area (AUROC 0.828) and splenic diameter (AUROC 0.804) may predict the presence of CSPH in patients with mixed underlying etiologies. Adding platelet count to either spleen area (AUROC 0.875) or spleen stiffness (AUROC 0.855) increased diagnostic accuracy. Spleenic area and diameter showed little inter-operator variability.

Our findings that spleen stiffness measured by p-SWE has a good diagnostic accuracy (cut-off > 29.99, AUROC 0.712) in identifying patients with CSPH is supported by other studies in the literature. However, these studies report varying diagnostic threshold values and performance (AUROC 0.970-0.688). Differences
Table 2 Univariate analysis showing association of clinical parameters with clinically significant portal hypertension

| Variable                  | Odds ratio | Cut-off     | P value | Estimated AUROC curve | P value of AUROC |
|---------------------------|------------|-------------|---------|------------------------|-----------------|
| BMI                       | 0.969      | > 26.42     | 0.425   | 0.506                  | 0.951           |
| ALT (IU/L)                | 0.993      | > 94.00     | 0.147   | 0.514                  | 0.869           |
| AST (IU/L)                | 1.002      | > 64.00     | 0.728   | 0.652                  | 0.065           |
| GGT (IU/L)                | 1.001      | > 85.00     | 0.639   | 0.643                  | 0.094           |
| ALP (IU/L)                | 1.001      | > 96.00     | 0.050   | 0.678                  | 0.033           |
| Bilirubin                 | 1.037      | > 11.00     | 0.013   | 0.722                  | 0.007a          |
| Platelet count (× 10^9)   | 0.979      | < 126.00    | < 0.001 | 0.846                  | < 0.001         |
| Albumin (g/L)             | 0.786      | < 33.00     | < 0.001 | 0.820                  | < 0.001         |
| Prothrombin time (sec)    | 2.721      | > 12.20     | < 0.001 | 0.800                  | < 0.001         |
| APRI score                | 2.030      | > 0.81      | 0.006   | 0.827                  | < 0.001         |
| ElastPQ median liver stiffness (kPa) | 1.048 | > 10.16     | 0.021   | 0.657                  | 0.061           |
| ElastPQ median spleen stiffness (kPa) | 1.007 | > 29.99     | 0.368   | 0.712                  | 0.010a          |
| ElastPQ spleen area (cm²)  | 1.075      | > 57.90     | < 0.001 | 0.828                  | 0.002c          |
| ElastPQ spleen diameter (cm) | 1.651 | > 13.90     | 0.002   | 0.804                  | 0.007b          |

*p < 0.05.

bP < 0.01.

cP < 0.005. AUROC: Area under the receiver operating characteristic; BMI: Body mass index; ALT: Alanine aminotransferase; AST: Aspartate aminotransferase; GGT: γ-glutamyl transpeptidase; ALP: Alkaline phosphatase; APRI: AST-to-platelet-ratio-index; ElastPQ: Elastography point quantification.

Table 3 Diagnostic performance of combination variables as predictors of clinically significant portal hypertension

| Variables                                                                 | AUROC curve | Sensitivity (%) | Specificity (%) | Positive predictive value (%) | Negative predictive value (%) |
|---------------------------------------------------------------------------|-------------|-----------------|-----------------|-----------------------------|-----------------------------|
| ElastPQ median spleen area (> 57.90 cm²) and Platelet count (< 126 × 10^9) | 0.875       | 63.2            | 100.0           | 100.0                      | 61.1                        |
| ElastPQ median spleen stiffness (> 29.99 kPa) and platelet count (< 126 × 10^9) | 0.855       | 88.0            | 75.00           | 78.6                       | 85.7                        |

AUROC: Area under the receiver operating characteristic; ElastPQ: Elastography point quantification.

Table 4 One-way random intraclass coefficient values for inter-operator variability

|                   | Spleen area | Spleen diameter |
|-------------------|-------------|-----------------|
| Number of operators | 2           | 2               |
| Number of measurements | 19          | 19              |
| Mean               | 0.98        | 0.96            |
| 95% confidence interval | 0.94-0.99  | 0.91-0.99      |

between studies may be explained by varying methodologies employed, as well as use of different p-SWE techniques. Nevertheless a recent study, which adopted a similar methodology to our own, identified a cut off of < 31 kPa to rule out the presence of EV of any grade which resonates with our findings\(^{19}\). Interestingly, our study did not identify liver stiffness as a predictor of CSPH, which differs from findings of recent studies\(^{16,18,20}\) and suggestions made by the Baveno VI Guidelines\(^{6}\). But, contrasting findings are not uncommon, as differences between
studies within the literature are also seen. A possible explanation for this may lie in the heterogeneity of populations in our study and between studies in the literature. Furthermore, studies comparing the ElastPQ technique to VTQ have shown significantly lower liver stiffness values, which may provide an added explanation for discrepancies seen[21]. Given the novelty of the ElastPQ, research focus has remained on its ability to detect fibrosis in comparison to other elastography techniques such as TE and VTQ[21-23]. As a result, there is limited data on the ability of ElastPQ to predict the presence of CSPH.

Splenic area and diameter demonstrated a modest ability to diagnose the presence of CSPH. Previous studies have explored spleen size by consideration of splenic diameter[22], which has shown to have acceptable reproducibility in the context of
platelet count/spleen diameter ratio. However, to our knowledge, there has only been one other study which has considered spleen area as a potential non-invasive diagnostic parameter. In this study, Giuffrè et al.\[^{19}\] reported similar findings with a median splenic area of 59.2 cm\(^2\) and diameter of 13.1 cm in its cohort of 210 patients\[^{19}\]. Given the excellent reproducibility seen in our study and confirmation of similar findings in one other study, spleen area may be a useful adjunct in predicting CSPH. Further research with an external cohort is needed to validate our findings.

Perhaps one of the most striking results from our study is the diagnostic accuracy of the APRI score. A cut-off value of > 0.81 generated an AUROC of 0.827 in predicting the presence of CSPH. This was comparable to diagnostic performance of spleen stiffness and spleen area. Nevertheless, these findings differ to those in the literature, which has described lower sensitivities in higher cut-off values\[^{24-26}\]. Only one study by Salzl et al.\[^{16}\] demonstrated a similar diagnostic performance (AUROC 0.805), but the cut-off value (1.90) remained higher than seen in our cohort\[^{16}\]. Differences between studies may be reflected by the smaller sample size and varied etiology within our study population. Of note, the study by Giuffrè et al.\[^{19}\], which had a similar study population, demonstrated APRI to be a statistically significant determinant of CSPH with a similar median of 0.70\[^{19}\]. Although highly applicable due to its non-invasive nature, the APRI score is affected by inflammatory processes such as acute hepatitis, which can generate false positive results that are not seen with p-SWE\[^{4}\]. We propose that the APRI score may be a useful tool in the follow-up of CLD patients in primary care while p-SWE may fare better in secondary practice.

Since the introduction of liver stiffness measurements by TE, combinations of liver stiffness with spleen size have been carried out. The most common of these is the LSPS score\[^{27-29}\]. However, despite spleen stiffness being increasingly recognized as a better predictor of CSPH\[^{13}\], few studies have been carried out combining spleen stiffness measurements to other markers of CLD. Our study showed that the combination of spleen stiffness measurements and platelet counts has a high diagnostic index (AUROC 0.855). Although an exact model has not been replicated, a similar model applying spleen stiffness measurements measured by TE and the Baveno guidelines VI has shown promising results\[^{44}\]. Colecchia et al.\[^{45}\] utilised a combined model where a cut-off of ≤ 46 kPa for spleen stiffness and < 20 kPa for liver stiffness measurements by TE, and a platelet count > 150000/mm\(^3\) could effectively rule out CSPH in cACLD patients\[^{8}\]. A different study by Bota et al.\[^{31}\] used a different combination index of liver stiffness and spleen stiffness measured by VTQ, and presence of ascites which generated an AUROC of 0.721\[^{31}\]. Finally, Giuffrè et al.\[^{19}\] was perhaps the most comparable of all the studies mentioned as his team used the ElastPQ model to develop the spleen stiffness probability index\[^{19}\]. All of the findings above support the premise that a combination of non-invasive parameters may be a better diagnostic indicator than a single parameter alone. No study in the literature has considered addition of spleen area to combination variables. Further studies are needed to validate our proposed spleen area and platelet count combination and determine which set of non-invasive parameters generate the best accuracy.
**Strengths and limitations**

Both the use of a novel p-SWE machine (ElastPQ) and investigation of spleen area describe a unique approach in our study compared to others carried out in the field. To our knowledge, this is one of the first studies to assess the role of spleen stiffness, spleen area and splenic diameter measurements in predicting CSPH using the ElastPQ. As a result, our study took into consideration inter-operator variability of splenic area and diameter, which supported its potential use in clinical practice. The prospective recruitment of patients with mixed etiologies described a population representative in clinical practice, but in view of the novelty of p-SWE it would be worth determining the effect of specific etiologies on splenic measurements.

This study has some limitations, the most pertinent of which being that we only assessed for presence or absence of PH, rather than degree of PH. Furthermore, an interval gap of one year between spleen stiffness measurements and EGD/HVPG readings may represent a consistent bias within our study due to the considerable length of time between readings. However, it could be argued that the correlation between CSPH and spleen stiffness may be better if there were a shorter time interval proposed. We did not exclude patients taking pharmacological treatment for PH from the original protocol as it was suspected that non-selective beta blockers and banding of varices would be unlikely to affect splenic measurements\[^{32}\]. However, the most recent data on cirrhotic patients with high risk varices suggests that taking non-selective beta blockers can affect splenic stiffness\[^{33,34}\]. Nevertheless these studies were undertaken using Fibroscan® and VTQ (Siemens Acuson S2000TM) ultrasound systems and so, further information is still needed in order to confirm that similar findings are present with the ElastPQ.

Although IQR measurements were taken, the validity of spleen stiffness and liver stiffness could not be determined as quality criteria has not yet been established for this technique\[^{4}\]. However, Pawluś et al\[^{35}\] conducted a small study in which he measured the spleen stiffness of 59 healthy volunteers using p-SWE, which has provided a reference point of 16.6 ± 2.5 kPa as the normal range with good reproducibility of measurement results\[^{35}\]. Given the similar methodologies, this has supported our findings despite the small sample size in this study. Ultimately, further studies are needed to validate our findings, but our multivariate models suggest that findings are likely to correlate with CSPH in larger cohorts.

**Clinical implications**

p-SWE is a non-invasive, rapid tool that carries minimal complications. It is painless, better tolerated than current gold-standard techniques and is more applicable than TE. Furthermore, this technique can be implemented on regular ultrasound machines and performed during routine screening for HCC in cirrhosis, which is likely to be cost-effective and less time-consuming.

**CONCLUSION**

Combinations of spleen area and platelet count, or spleen stiffness and platelet count as measured by the ElastPQ may be safe and effective methods to diagnose CSPH. Currently, this non-invasive technique cannot replace gold-standard as further studies are needed to create validation criteria and assess the diagnostic accuracy of non-invasive parameters in patients with differing degrees of PH.

**ARTICLE HIGHLIGHTS**

**Research background**

Portal hypertension is a major complication of cirrhosis with a significant morbidity and mortality associated with it. Many of those with advanced chronic liver disease have esophageal varices and so, many patients undergo the gold-standard invasive procedures of performing an esophago-gastroduodenoscopy (EGD) or having the hepatic venous pressure gradient measurement taken through interventional radiology. However, both of these methods are invasive and carry a risk of complications.
Research motivation
Current guidelines propose that non-invasive methods can predict the incidence of clinically significant portal hypertension (CSPH). The latest guidelines suggest cirrhosis patients with a liver stiffness measurement < 20 kPa and a platelet count > 150000/μL can avoid screening endoscopy. Nevertheless, new algorithms are still required, as up to 40% of EGDs continue unnecessarily.

Research objectives
The aim of this study was to assess whether spleen stiffness measurement, spleen area and spleen diameter can independently predict CSPH, or in combination with other biochemical or elastography parameters. We also aimed to assess reproducibility of splenic area and diameter measurements.

Research methods
This was a single-centre prospective cohort study where a total of 50 patients were split into two groups and included in a retrospective analysis: 25 with evidence of CSPH (group 1) and 25 with no evidence of CSPH (group 2). The Philips EPIQ7 [elastography point quantification (ElastPQ)] (Philips Medical Systems, Seattle, United States) was used to record liver stiffness, spleen stiffness, spleen area and spleen diameter measurements for each patient. Univariate, multivariate and one-way random interclass correlation coefficient analyses were performed to assess the diagnostic accuracy of splenic parameters.

Research results
Body mass index, alanine aminotransferase, aspartate aminotransferase, γ-glutamyl transpeptidase, alkaline phosphatase, bilirubin, platelet count, albumin, prothrombin time, aspartate aminotransferase-to-platelet-ratio-index (APRI) score, liver stiffness, spleen stiffness, spleen area and spleen diameter were assessed in their ability to predict the presence of CSPH. A univariate analysis showed the best individual predictor of CSPH was platelet count [area under the receiver operating characteristic (AUROC) 0.846, P value < 0.001], followed by spleen area (AUROC 0.828, P value = 0.002) and APRI score (AUROC 0.827, P value < 0.001). A multiple logistic regression model revealed that two combinations independently predict CSPH. The combination with the greatest diagnostic accuracy included a combination of spleen area > 57.9 cm² and platelet count < 126 × 10⁹ which had 63.2% sensitivity, 100% specificity, 100% positive predictive value (PPV), 61.1% negative predictive value (NPV) (AUROC 0.876, P value < 0.001). An alternative combination of spleen stiffness > 29.99 kPa and platelet count < 126 × 10⁹ displayed a similar diagnostic accuracy with 88% sensitivity, 75% specificity, 78.6% PPV, 85.7% NPV (AUROC 0.855, P value < 0.001). Spleen area and spleen diameter demonstrated little inter-operator variability as measured by a one-way random interclass correlation coefficient (spleen area: 0.98, P value < 0.001; spleen diameter: 0.96, P value < 0.001).

Research conclusions
Combinations of spleen area and platelet count, or spleen stiffness and platelet count as measured by the ElastPQ may be safe and effective methods to diagnose CSPH. At present this cannot replace the gold standard.

Research perspectives
Performing large scale prospective studies with long-term follow-up and are needed to validate our findings.

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