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5	EVALUATION OF LABORATORY TESTS FOR CIRRHOSIS AND
6	FOR ALCOHOL USE, IN THE CONTEXT OF ALCOHOLIC CIRRHOSIS
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55 **ABSTRACT** (288 words)

56 Laboratory tests can play an important role in assessment of alcoholic patients, including for 57 evaluation of liver damage and as markers of alcohol intake. Evidence on test performance should lead to better selection of appropriate tests and improved interpretation of results. We 58 59 compared laboratory test results from 1578 patients between cases (with alcoholic cirrhosis; 60 753 men, 243 women) and controls (with equivalent lifetime alcohol intake but no liver disease; 439 men, 143 women). Comparisons were also made between 631 cases who had 61 62 reportedly been abstinent from alcohol for over 60 days and 364 who had not. ROC curve analysis was used to estimate and compare tests' ability to distinguish patients with and 63 without cirrhosis, and abstinent and drinking cases. The best tests for presence of cirrhosis 64 were INR and bilirubin, with AUCs of 0.91 ± 0.01 and 0.88 ± 0.01 respectively. Confining 65 analysis to patients with no current or previous ascites gave AUCs of 0.88 ± 0.01 for INR and 66 67 0.85 ± 0.01 for bilirubin. GGT and AST showed discrimination between abstinence and 68 recent drinking in patients with cirrhosis, including those without ascites, when appropriate (and for GGT, sex-specific) limits were used. For AST, a cut-off limit of 85 units/l gave 90% 69 70 specificity and 37% sensitivity; for GGT cut-off limits of 288 units/l in men and 138 units/l in women gave 90% specificity for both and 40% sensitivity in men, 63% sensitivity in women. 71 72 INR and bilirubin show the best separation between patients with alcoholic cirrhosis (with or without ascites) and control patients with similar lifetime alcohol exposure. Although AST 73 74 and GGT are substantially increased by liver disease, they can give useful information on 75 recent alcohol intake in patients with alcoholic cirrhosis when appropriate cut-off limits are 76 used.

77 **KEY WORDS**

78 Alcohol; cirrhosis; abstinence; aspartate aminotransferase; gamma glutamyl transferase.

79 BACKGROUND

Laboratory tests play an important role in the diagnosis and monitoring of patients with alcoholic cirrhosis, both for assessing the degree of impairment of liver function by cirrhosis and for detecting ongoing alcohol intake. It is important to share information on test performance, to optimise test selection and diagnostic accuracy.

84 Many aspects of liver function are impaired in cirrhosis, and form the basis of diagnostic or 85 prognostic tests. These include excretory, synthetic, and metabolic functions, reflected in abnormal results for bilirubin; albumin and clotting factors; and glucose and ammonia, 86 87 respectively. Damage to liver cells, perhaps combined with increased enzyme expression, leads to increases in plasma or serum activity of 'liver enzymes' (gamma-glutamy) 88 transferase, GGT; aspartate aminotransferase, AST; alanine aminotransferase, ALT). 89 Although it is recognised that quite advanced cirrhosis may occur with normal liver function 90 test results (1-3), and there are recent papers comparing test results in drinking versus 91 92 abstinent alcoholics (4) and in 'heavy-drinking controls' versus patients with alcoholic 93 hepatitis (5) there is little published evidence on the comparative performance of widely available tests in distinguishing between the presence or absence of cirrhosis in heavy alcohol 94 95 drinkers. Such evidence would be valuable in its own right, and because novel tests or algorithms should be judged against the performance of currently available tests. 96

97 A related issue is the value of biochemical tests as markers of alcohol use in patients with 98 liver disease, particularly alcoholic liver disease. Because the prognosis in alcoholic cirrhosis 99 is greatly improved by abstinence and treatment decisions may be affected, objective and 100 reliable measures of patients' alcohol use can be helpful. Measurement of ethanol metabolites 101 shows promise (6-8) but most either require frequent testing because of short half-lives (ethyl 102 glucuronide and ethyl sulphate in urine) or are not widely available (ethyl glucuronide or 103 fatty acid ethyl esters in hair, phosphatidylethanol in blood cell membranes). There are mixed 104 reports on whether serum disialotransferrin (carbohydrate-deficient transferrin, CDT) is affected by liver disease (9-14). A number of technical issues, depending on the method used, 105 can affect the validity of CDT results in cirrhosis (15-17). Serum GGT, which is cheap and 106 107 widely available, is a rather non-specific marker of liver damage as well as an index of alcohol intake, and it is increased in a high proportion of people with liver disease. GGT has 108 therefore been discounted for this situation, though there is little information on its potential 109 110 as an alcohol biomarker in the presence of liver disease. Nor is information readily available about the ability of other commonly available tests to distinguish abstinent from non-111 112 abstinent patients.

We have collected blood samples and clinical information, including alcohol intake history 113 and laboratory test results, from 1578 patients either with liver cirrhosis due to alcohol or 114 115 with similar alcohol intake but no history or symptoms of liver disease (18). These data allow 116 us to address the two questions outlined above. First, which tests (including biochemical liver 117 function tests, and haematology tests affected by cirrhosis) are best at distinguishing between those who do or do not have cirrhosis as a result of long-term excessive alcohol intake? 118 Second, can any of these commonly available tests assist in identifying continuing alcohol 119 use among patients with alcoholic liver disease? 120

121 METHODS

122 Information was gathered from patients recruited for the GenomALC Study (18) up to the 123 end of April 2016. Recruitment occurred in Australia, France, Germany, Switzerland, UK 124 and USA, mainly from hepatology clinics (for cases, as defined below) and from psychiatric 125 or detoxification facilities for the controls. All participants gave informed consent and the 126 study was approved by appropriate Research Ethics Committees.

127 To be eligible, participants had to have high-risk alcohol intake (greater than 80 grams per day for men, or 50 grams per day for women) for 10 years or more. Cases had alcohol-related 128 129 cirrhosis, with the diagnosis based on one or more of the following clinical, histological or FibroScan criteria as reported (18) and detailed here. Clinical cirrhosis required documented 130 evidence of one or more of the following: clinically detectable ascites (confirmed by imaging 131 or by paracentesis); spontaneous hepatic encephalopathy (grade 2 or higher); moderate or 132 large oesophageal varices on upper endoscopy. Histological cirrhosis required Metavir 133 134 fibrosis stage F4 or Ishak fibrosis stage 5 or 6. Fibroscan diagnosis required an adequately performed FibroScan with F4 stiffness; the cut off was ≥ 22 kPa (if AST <100 IU/L within 2 135 weeks of FibroScan), or ≥30 kPa (if AST between 100-200 IU/L within 2 weeks of 136 137 FibroScan). Exclusion criteria included liver transplantation for liver disease other than alcoholic cirrhosis; hepatitis B or C (by hepatitis C antibody and hepatitis B surface antigen 138 tests), known HIV, hemochromatosis (by transferrin saturation > 45% or 2+ iron on liver 139 biopsy if performed), Wilson's disease (by ceruloplasmin) or autoimmune hepatitis (by ANA 140 titre). 141

142 Control subjects had to meet the alcohol intake criteria with no history or current evidence of 143 liver disease (history of jaundice, ascites, variceal bleeding, upper gastrointestinal bleeding of 144 uncertain etiology, or blood tests which suggest impaired liver function or acute/chronic 145 alcoholic liver injury). 146 Characteristics of 1578 participants who met the eligibility criteria are summarised in Table147 1.

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Lifetime alcohol intake estimates were based on participants' recall of habitual daily use of

beer, wine, spirits or other alcoholic beverages (converted to grams of alcohol), and of the 149 number of years of high-risk drinking. Current abstinence was assessed by whether the 150 patient reported they had been abstinent from alcohol for 60 days or more before recruitment. 151 152 Data collection was planned before test and reference standard data were collected. Laboratory test results, as listed in Table 2, were gathered from those done for clinical 153 154 reasons at the time of recruitment or performed for this study where necessary. We also calculated AST/ALT and AST/platelet ratios. MELD scores were calculated from INR, 155 bilirubin and creatinine results (19) using the formula MELD = 3.78[Ln serum bilirubin 156 (mg/dL)] + 11.2[Ln INR] + 9.57[Ln serum creatinine (mg/dL)] + 6.43. Results for bilirubin, 157 INR and creatinine of less than 1.0 (in their respective units) were taken as 1.0, and results for 158 159 creatinine of greater than 4.0 were taken as 4.0, as recommended by the United Network for 160 Organ Sharing (UNOS) (https://www.unos.org/wpcontent/uploads/unos/MELD_PELD_Calculator_Documentation.pdf, accessed 2016-05-30). 161 For comparison of means between groups, test results showing positively skewed 162 163 distributions (bilirubin, creatinine, ALT, AST and GGT) were log₁₀-transformed. For ROC curve analysis, test results where the case mean was lower than the control mean 164 (hemoglobin; white cell count; platelet count; albumin) had the assumption of higher results 165 indicating abnormality reversed so that areas under the ROC curve (AUC) were greater than 166 0.5. Statistical analyses were performed using SPSS (IBM Corporation, Armonk, New York 167 168 10504).

169 **RESULTS**

170 The test means for abstinent and non-abstinent cases and controls are summarised in Table 2, with results for men and women shown separately in Supplementary Table 1. P-values for 171 both the effects of presence of cirrhosis and of abstinence on the means, and for case/control 172 by abstinent/non-abstinent interaction, are also shown. Most of the tests showed differences 173 between the case and control groups, but only AST and GGT showed significant effects of 174 abstinence. These two tests also showed significant case/control by abstinent/non-abstinent 175 interaction terms. Plots for AST and GGT by case-control status and by abstinence, to 176 illustrate the main effects and interaction, are shown in Figure 1; reported abstinence was 177 associated with lower AST and GGT in cases but not in controls (but very few control 178 patients had abstained from alcohol). 179

The ability of the laboratory tests to distinguish cases from controls is summarised in Table 3. 180 ROC curves (which plot test sensitivity, true positive rate, against (1-specficity), false 181 182 positive rate) are shown for the most discriminating tests (hemoglobin, platelet count, INR, 183 bilirubin and albumin) and the MELD score in Supplementary Figure 1. Because there is always a trade-off between better sensitivity and better specificity, determined by the chosen 184 185 cut-off value separating 'normal' from 'abnormal' results, comparisons of sensitivity between tests or between groups of patients should be based on the same specificity for each. For our 186 comparisons, we have chosen 90% specificity (10% false positive rate) unless otherwise 187 noted, and report the cut-off values and sensitivities associated with that specificity. 188

Most (77%) of the patients with alcoholic cirrhosis had current or prior ascites. In order to test how far this affected the test results and their diagnostic performance, we conducted further analyses on case sub-groups defined by presence or history of ascites, comparing those with and without ascites. For most tests, ascites was significantly associated with moreabnormal results (Supplementary Table 2), and exclusion of cases with reported ascites 194 decreased the case-control AUCs (Table 4). The notable exceptions were AST and GGT, where ascites was associated with lower (less abnormal) mean values and with higher AUCs. 195 196 Because only 17 of the controls reported abstinence for 60 days preceding recruitment, analysis of the ability of tests to distinguish abstinence from continued drinking was confined 197 198 to the cases (Table 3). The only tests showing AUC above 0.70 were AST and GGT, and results for these are shown in more detail in Table 5 and Figure 2. When data from men and 199 women were analysed together, the AUC for AST was 0.737 and for GGT 0.771. This 200 201 analysis was then repeated for male and female cases separately (also shown in Table 5). For AST, the AUC, test sensitivities and cut-off values were similar in men and women; but for 202 GGT the AUC was greater in women than in men and the appropriate cut-off values 203 (determined by the desired specificity) were substantially higher in men. 204

205 **DISCUSSION**

206 We have compared the performance of routine tests, and the composite MELD score, for distinguishing between patients with alcoholic cirrhosis (cases) and patients with similar 207 lifetime exposure to alcohol but no liver disease (controls). The best of these tests show good 208 discrimination, consistent with the comparison of selected groups and with clinical 209 210 experience. We have also compared results from abstinent and non-abstinent patients with alcoholic cirrhosis. The tests which perform best for making the distinction between abstinent 211 and non-abstinent cases are GGT and AST and they perform well in patients with advanced 212 liver disease as long as appropriately high cut-off limits are used. 213

It is generally accepted that conventional liver function tests have poor sensitivity in 214 detecting cirrhosis, particularly in the early stages. Although our cases have (or have had) 215 clinical symptoms, and we accept that we are comparing selected extremes of the spectrum of 216 potential patients, we find that INR, bilirubin, platelet count and albumin - in that order -217 218 give good discrimination between cases and controls (Table 3). The best single test, INR, had 219 an AUC of 0.914 and test sensitivity of 78% at a specificity of 90%. Even in less advanced disease, i.e. after restricting the analyses to patients without ascites, INR and bilirubin 220 221 continued to show good separation between the case and control groups.

The calculated AST/ALT ratio showed better discrimination than either of its components in the case/control comparison (see Table 3). The AST/platelet ratio showed no advantages, being significantly worse than platelet count for case/control discrimination or AST for drinking/abstinence (again, see Table 3); this is consistent with a previous evaluation (20). The MELD score, being based on INR, bilirubin and creatinine, gives results equivalent to (but no better than) the INR measurement alone for the case versus control comparison (although MELD may still be superior to any single test for other purposes, such asprognosis, which we did not evaluate).

To be an improvement on what is already available, any new test or test combination would 230 need to achieve either an AUC above 0.91 in patients equivalent to ours, or an equal 231 sensitivity and specificity in patients with less advanced disease. Indicators of fibrosis might 232 233 be valuable in patients with less advanced disease, and a number have been investigated. 234 Results were summarised in (3), with some markers having high reported AUCs or promising sensitivity and specificity values in comparatively small studies. A direct comparison of three 235 fibrosis markers, tissue inhibitor of metalloproteinase 1, aminoterminal propeptide of type III 236 237 collagen and hyaluronic acid, showed highly significant differences in mean values between 238 alcoholic patients with mild and advanced fibrosis but the AUCs were in the range 0.67-0.69 and sensitivity was around 33% at 90% specificity. 239

Another important clinical question is whether people with known alcoholic cirrhosis are 240 241 abstaining from alcohol. Taking the cases only, ROC curve analysis was performed to assess 242 the ability of the laboratory tests to classify people as abstinent or non-abstinent (Tables 3 and 5, Figure 2). The tests which were best at distinguishing cases from controls (INR, 243 244 bilirubin, platelet count, albumin) performed poorly in distinguishing abstinent and non-245 abstinent cases; they are detecting cirrhosis rather than drinking. It is unexpected that test results are not closer to normal in the abstinent than in the drinking cases, although the period 246 of abstinence specified (60 days or more) may be too short to have made a difference. 247

On the other hand, AST and GGT, which did not perform well for the case-control comparison, did surprisingly well in the abstinent-drinking comparison. These are tests which are primarily measuring liver cell damage and/or enzyme induction and which have not previously been considered useful in the presence of liver disease. In fact, the test performance (Table 5) for GGT is very similar to that derived from meta-analysis of data from studies on people without liver disease (21) (which estimated GGT sensitivity of 44% and AST sensitivity of 27%, each at 90% specificity). However, the GGT value giving this specificity and sensitivity in our cases (about 250 units/l) is much higher than it would be in alcoholics without known liver disease.

Another point to notice is that AST and GGT performed slightly better, both in the casecontrol and abstinent-drinking comparisons, in patients with cirrhosis but no ascites (Table 4). This is in contrast to the other tests, and is probably due to decreased liver cell mass in the patients with more advanced disease who have or have had ascites. As these enzymes originate from the liver, very low functioning liver cell mass will lead to less enzyme release into the circulation.

263 For the evaluation of abstinence in individuals with cirrhosis, we found relevant differences in test performance between men and women. The performance of GGT was better in women 264 265 than in men (Table 5, Figure 2) and the appropriate cut-off values for various levels of 266 specificity were higher in men. For example, a cut-off value of around 290 units/l would give 40% sensitivity and 90% specificity in men but a cut-off value of 140 units/l would give 63% 267 268 sensitivity and 90% specificity in women. (The cut-off value for equivalent specificity in the 269 absence of liver disease would be around 40-50 units/l.) On the other hand, AST (which performs about as well as GGT as an alcohol marker in the alcoholic cirrhosis context) 270 271 showed similar test performance and cut-off limits in men and women (Table 5), with a cutoff of around 85 units/l (still substantially above the appropriate value for people without 272 liver disease) giving sensitivity of about 35% and 90% specificity. 273

As mentioned above, there is a trade-off between diagnostic sensitivity and specificity. So far we have compared test performance at 90% specificity. If prevalence of the condition is low, 276 it is appropriate to use a high cut-off value to attain high specificity because of the need to minimise false positives. However, there are clinical situations where high sensitivity is 277 needed and poor specificity can be tolerated, and detection of continued drinking in patients 278 279 with alcoholic cirrhosis may be one of these. If specificity of only 70% can be accepted, then the sensitivity of GGT for detection of continued drinking in the presence of cirrhosis 280 increases to about 65% in men (at 133 units/l) and 80% in women (at 85 units/l). However, 281 282 even though GGT and AST have some ability to distinguish currently drinking from currently abstinent patients with alcoholic cirrhosis, it would be inappropriate to place too much 283 284 reliance on them. As with patients who do not have liver disease, high GGT should be considered suggestive of excessive or continuing alcohol use and a finding which warrants 285 further investigation. 286

We acknowledge some limitations due to our study design, particularly the existence of 287 288 spectrum bias because of comparison of extremes rather than unselected patients. Participants were recruited for a case-control genetic association study, so it was important to 289 290 select cases with strong evidence of cirrhosis. This limitation should be less of a problem in 291 the comparison of abstinent and non-abstinent cirrhotic patients, if we assume that alcoholics are either abstinent or drinking heavily and cannot maintain controlled drinking. Despite 292 293 assessment for the absence of past or current symptoms, a few control subjects may have had some liver damage from alcohol, though probably insignificant given our stringent eligibility 294 criteria. If liver damage was present in some controls, this would tend to decrease the 295 difference between cases and controls, and therefore impair test performance. 296

Another limitation is that test evaluations depend on having a reliable diagnosis. Liver biopsy is often used as a 'gold standard' for cirrhosis but it is invasive, not always justifiable, and may be subject to sampling error. Clinical symptoms in the presence of high long-term alcohol intake, and exclusion of alternative causes of cirrhosis, formed the basis for diagnosis in our cases; and controls were recruited with similar alcohol intake and absence ofsymptoms or history of liver disease.

Finally, we used self-report to assess alcohol intake and abstinence, again with no gold 303 standard. This has been the method of alcohol use assessment in many studies on alcohol 304 305 consumption, both those which have focused on epidemiological associations between alcohol and health or disease, and those which have evaluated alcohol biomarkers. In general, 306 self-report is a valid approach to assessment, particularly in a setting in which there are no 307 negative consequences for a participant who reports ongoing alcohol use, but it may be 308 subject to bias (22). Accuracy of self-reported alcohol use may vary according to sex, 309 country, case/control status or other unmeasured factors. However, it is reasonable to assume 310 that patients with cirrhosis who report continued drinking are giving correct information, 311 while the group who report abstinence contains some who are drinking. If so, any bias will be 312 313 conservative in that test performance will be under-estimated.

314

315 Conclusions

We have documented and compared tests related to liver function in alcoholic cirrhosis, and shown the best performance for INR and bilirubin. AST and GGT are increased by liver disease but they may still give useful information on recent alcohol intake in patients with alcoholic cirrhosis if appropriately higher and sex-specific cut-off values are used.

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Figure 1. Boxplots of AST and GGT results by Case-Control and Abstinent/Non-Abstinent status. Boxes show 25th, 50th and 75th centiles, whiskers indicate 95% range. For the legend 'Abstinent 60 days' 1 = Yes (abstinent) and 2 = No (drinking). For each test, values differ significantly by both case/control and abstinent drinking status but there is also case/control by abstinent/drinking interaction (see Table 2). Abstinent/drinking status has significant effects in cases but not in controls.

Figure 2. Comparison of ROC curves for AST and GGT in men and women. Classification of Cases as Abstainer for past 60 days versus Non-Abstainer.

Table 1. Descriptive data on the 1578 GenomALC Cases and Controls included in theanalysis. High-risk drinking is defined as equal to or greater than 80 grams of alcohol per dayfor men or 50 grams/day for women, for 10 years or more.

	Cases (N	N = 997)	Controls	(N = 581)
	Male	Female	Male	Female
Number of subjects	754	243	438	143
Age (Mean ± SD, in years)	52.6 ± 8.7	50.1 ± 9.6	50.2 ± 10.0	50.3 ± 10.1
Usual alcohol intake, g/day	211 ± 148	162 ± 118	243 ± 135	197 ± 109
Years of high-risk drinking	25.3 ± 11.2	19.7 ± 9.8	22.2 ± 9.6	18.4 ± 7.5
Lifetime alcohol intake, kg	1953 ± 1754	1148 ± 1022	2002 ± 1582	1346 ± 1039
Number with ascites (ever)	573 (76%)	193 (79%)	0	0
Number with oesophageal varices (ever)	404 (54%)	126 (52%)	0	0
Number with encephalopathy (ever)	247 (33%)	89 (37%)	0	0
Number abstinent for ≥ 60 days	476 (63%)	155 (64%)	13 (3%)	4 (3%)

Table 2. Effects of alcoholic cirrhosis (case versus control) and recent drinking (reported abstinence for previous 60 days) on laboratory test results. For bilirubin, creatinine, AST, ALT and GGT the significance of differences in means and of the interaction term was assessed on log-transformed data to reduce the effects of skewed distributions. To allow for multiple testing, p-values less than 0.0038 (0.05/13) may be considered significant.

		(Controls Cases			p-values					
		Mean	SD	Ν	Mean	SD	Ν	Case-Control	Abstinence	Interaction	
Haemoglobin,	Abstinent	143.6	15.7	16	117.4	23.6	618	1.16×10^{-19}	0.944	0.840	
(g/l)	Non-Abstinent	142.9	15.7	557	117.7	25.6	362	1.10 x 10	0.944	0.072	
White cell count	Abstinent	7.907	2.549	17	6.267	2.819	616	0.063	0.036	0.029	
$(cells/l x 10^{-9})$	Non-Abstinent	7.877	2.575	556	8.006	4.413	359	01002	01020	0.02)	
			<i></i>	15	105.0	- 1 <	(10)				
Platelet count	Abstinent	251.9	64.3	17	135.8	/1.6	613	3.02 x 10 ⁻²⁷	0.742	0.480	
$(\text{cells/l x } 10^{-5})$	Non-Abstinent	248.3	81.3	555	146.0	82.8	361				
		1.000	0.040	17	1 402	0 455	505				
INR	Abstinent	1.008	0.243	17	1.402	0.455	595	3.14 x 10 ⁻¹⁷	0.817	0.495	
(ratio)	Non-Abstinent	0.986	0.154	497	1.447	0.508	326				
	A.1	41.5	4 4	17	25.4	6.0	506				
Albumin	Abstinent	41.5	4.4	17	35.4	6.9	596	4.01 x 10 ⁻¹⁸	0.864	0.134	
(g/l)	Non-Abstinent	43.0	5.6	545	34.2	7.7	333				
	A.1	10.6	0.1	17	50.0	01.6	(01				
Bilirubin	Abstinent	10.6	8.1	17	50.8	81.6	621	5.20 x 10 ⁻³¹	0.176	0.032	
(µmol/I)	Non-Abstinent	9.3	7.3	553	88.7	130.1	363				
~ · ·		75.1	10 6	17	04.5	67 1	(22)				
Creatinine	Abstinent	75.1	13.6	17	94.5	67.1	622	0.235	0.0068	0.122	
(µmol/l)	Non-Abstinent	71.9	17.8	558	75.3	39.2	360				
			110 -	15	24.5	10 7	()				
ALT	Abstinent	56.4	110.7	17	34.5	48.7	620	0.920	0.075	0.030	
(units/l)	Non-Abstinent	38.0	34.4	554	45.0	38.8	363	0.720	0.070	0.020	

	Controls			Cases			p-values			
		Mean	SD	Ν	Mean	SD	Ν	Case-Control	Abstinence	Interaction
AST (units/l)	Abstinent Non-Abstinent	48.2 41.1	66.2 33.9	17 552	50.1 83.4	48.9 59.5	606 356	1.24 x 10 ⁻⁹	3.25 x 10 ⁻⁴	8.71 x 10 ⁻⁴
GGT (units/l)	Abstinent Non-Abstinent	285.5 113.6	924.2 156.8	17 553	126.4 424.0	171.4 627.2	581 348	1.35 x 10 ⁻⁹	7.39 x 10 ⁻⁷	3.54 x 10 ⁻⁴
AST/ALT ratio	Abstinent Non-Abstinent	1.136 1.184	0.386 0.435	17 547	1.743 2.113	1.029 0.952	606 355	8.18 x 10 ⁻¹³	0.051	0.124
AST/platelet ratio	Abstinent Non-Abstinent	0.190 0.201	0.225 0.247	17 547	0.524 0.841	0.595 1.017	596 353	5.24 x 10 ⁻²⁵	0.017	0.061
MELD score	Abstinent Non-Abstinent	7.24 7.05	1.78 1.36	17 490	13.57 14.52	5.95 6.94	591 324	7.42 x 10 ⁻²⁵	0.566	0.387

Table 3. Results of ROC curve analysis; for alcoholic cirrhosis (Cases versus Controls), and for abstinence among patients with alcoholic cirrhosis. To allow for multiple testing, p-values less than 0.0038 (0.05/13) may be considered significantly different from chance (i.e. from AUC = 0.500).

		Case	es versus	Controls		Abstinent Cases versus Drinking Cases				
	N Cases	N Controls	AUC	Std. Error	p-value	N Drinking	N Abstinent	AUC	Std. Error	p-value
Haemoglobin	982	573	0.802*	0.011	7.63 x 10 ⁻⁸⁸	362	618	0.501	0.019	0.960
White cell count	977	574	0.644*	0.014	2.05 x 10 ⁻²¹	359	616	0.616	0.019	1.43 x 10 ⁻⁹
Platelet count	976	573	0.852*	0.010	7.23 x 10 ⁻¹¹⁹	361	613	0.528	0.019	0.143
INR	923	515	0.914	0.008	2.80 x 10 ⁻¹⁵⁰	326	595	0.522	0.020	0.273
Bilirubin	986	571	0.875	0.009	2.50 x 10 ⁻¹³⁴	363	621	0.599	0.019	2.40 x 10 ⁻⁷
Albumin	931	563	0.821*	0.011	1.78 x 10 ⁻⁹⁶	333	596	0.543	0.020	0.031
AST	964	570	0.685	0.014	8.35 x 10 ⁻³⁴	356	606	0.737	0.017	8.85 x 10 ⁻³⁵
ALT	985	572	0.483	0.015	0.275	363	620	0.649	0.018	6.19 x 10 ⁻¹⁵
GGT	931	571	0.643	0.014	1.35 x 10 ⁻²⁰	348	581	0.771	0.016	2.03 x 10 ⁻⁴³
Creatinine	984	576	0.573	0.014	1.52 x 10 ⁻⁰⁶	360	622	0.643	0.018	6.26 x 10 ⁻¹⁴
AST/ALT ratio	963	565	0.774	0.012	2.00 x 10 ⁻⁷¹	355	606	0.627	0.019	4.61 x 10 ⁻¹¹
AST/platelet ratio	951	565	0.815	0.011	5.73 x 10 ⁻⁹⁴	353	596	0.641	0.018	3.60 x 10 ⁻¹³
MELD score	917	508	0.914	0.008	7.89 x 10 ⁻¹⁴⁸	324	591	0.527	0.020	0.173

* Positive status (Case) is associated with lower test result.

Table 4. Comparison of selected ROC curve results for all Cases, and for Cases with or without current or

past ascites.

	All	With ascites	No ascites
	$AUC \pm SE$	$AUC \pm SE$	$AUC \pm SE$
Case versus Control comparison			
INR	0.914 ± 0.008	0.924 ± 0.008	0.884 ± 0.014
MELD score	0.913 ± 0.008	0.928 ± 0.008	0.865 ± 0.016
Bilirubin	0.875 ± 0.009	0.881 ± 0.009	0.853 ± 0.015
Platelet count	0.852 ± 0.010	0.855 ± 0.010	0.842 ± 0.017
Albumin	0.821 ± 0.011	0.838 ± 0.011	0.762 ± 0.021
Hemoglobin	0.802 ± 0.011	0.831 ± 0.011	0.703 ± 0.022
AST	0.685 ± 0.014	0.669 ± 0.015	0.738 ± 0.020
GGT	0.643 ± 0.014	0.606 ± 0.016	0.762 ± 0.019
Cases only, Abstinent versus Drinking com	parison,		
AST	0.737 ± 0.017	0.717 ± 0.021	0.784 ± 0.031
GGT	0.771 ± 0.016	0.753 ± 0.020	0.762 ± 0.032

Table 5. Details of ROC curve analysis for AST and GGT in distinguishing between Cases with reported abstinence for 60 days and Cases reported as nonabstinent.

		AST		GGT			
	Combined	Female	Male	Combined	Female	Male	
AUC (95% CI) Standard Error	0.737 (0.705 to 0.770) 0.017	0.774 (0.713 to 0.835) 0.031	0.726 (0.688 to 0.764) 0.020	0.771 (0.739 to 0.802) 0.016	0.851 (0.798 to 0.904) 0.027	0.744 (0.706 to 0.781) 0.019	
p-value	8.85 x 10 ⁻³⁵	2.30 x 10 ⁻¹²	2.31 x 10 ⁻²⁴	$2.03 \text{ x } 10^{-43} \qquad 2.28 \text{ x } 10^{-18}$		1.92 x 10 ⁻²⁷	
70% Specificity: Sensitivity	0.67	0.70	0.66	0.69	0.82	0.66	
Cut-off (units/l)	53	53	53	122	85	133	
80% Specificity: Sensitivity	0.54	0.59	0.53	0.60	0.74	0.54	
Cut-off (units/l)	63	64	63	168	108	200	
85% Specificity: Sensitivity	0.46	0.50	0.45	0.51	0.68	0.49	
Cut-off (units/l)	72	73	72	215	126	232	
90% Specificity: Sensitivity	0.37	0.34	0.36	0.46	0.63	0.40	
Cut-off (units/l)	85	87	84	265	138	288	
95% Specificity: Sensitivity	0.23	0.22	0.24	0.35	0.54	0.28	
Cut-off (units/l)	105	108	103	363	220	422	





Figure 2.

