BMJ Open Comparison and validation of International Consensus Diagnostic Criteria for diagnosis of autoimmune pancreatitis from pancreatic cancer in a Taiwanese cohort

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ABSTRACT

Objectives: The International Consensus Diagnostic Criteria (ICDC) designed to diagnosis autoimmune pancreatitis (AIP) has been proposed recently. The diagnostic performance of ICDC has not been previously evaluated in diffuse-type and focal-type AIP, respectively, in comparison with the revised HISORt and Asian criteria in Taiwan.

Design: Prospective, consecutive patient cohort.

Setting: Largest tertiary referred centre hospital managing pancreatic disease in Taiwan.

Participants: 188 patients with AIP and 130 with tissue proofed pancreatic adenocarcinoma were consecutively recruited.

Interventions: The ICDC, as well as revised HISORt and Asian criteria, was applied for each participant. Each diagnostic criterion of ICDC was validated with special reference to levels 1 and 2 in diffuse-type and focal-type AIP.

Outcomes: Sensitivity, specificity and accuracy.

Each diagnostic criterion of ICDC was validated with special reference to levels 1 and 2 in diffuse-type and focal-type AIP.

Results: The sensitivity, specificity and accuracy of ICDC for all AIP were the best: 89.4%, 100% and 93.7%, respectively, in comparison with the revised HISORt and Asian criteria. The area under the curve of receiver-operator characteristic of ICDC was 0.95 (95% CI 0.92 to 0.97) in all AIP and 0.93 (95% CI 0.88 to 0.97) in focal-type AIP.

Conclusions: The sensitivity, specificity and accuracy of ICDC are higher than the revised HISORt and Asian criteria. The sensitivity, specificity and accuracy of each criterion are higher in diffuse-type AIP compared with focal-type AIP. Under the same specificity, the sensitivity and accuracy of ICDC are higher than other diagnostic criteria in focal-type AIP. ICDC has better diagnostic performance compared with previously proposed diagnostic criteria in diffuse-type and focal-type AIP.

INTRODUCTION

Autoimmune pancreatitis (AIP) is a unique type of chronic pancreatitis characterised by elevated serum IgG4, swelling of the pancreas, irregular narrowing of the main pancreatic duct (MPD), histological evidence of lymphoplasmacytic inflammation and a good response to steroid therapy.1 Although some advance has been made in the diagnosis and treatment of AIP in past years,2 the diagnosis of AIP is still a great clinical challenge, especially in the differential diagnosis from pancreatic cancer (PC).3–5 Correct diagnosis of AIP could avoid delayed unnecessary resection of pancreas and vice versa, to avoid delayed treatment of PC. In 2002, the Japan Pancreas Society proposed diagnostic criteria for AIP based on imaging, serology and histology. At that time, the serological criteria included elevated γ-globulin, IgG and autoantibodies.6 In 2006, the revised Japanese criteria were modified and added IgG4 to the serological
criteria. In 2008, the Asian diagnostic criteria was established by modification of the Japanese and Korean diagnostic criteria. In western countries, the HISORt criteria were proposed from America. In 2011, the International Consensus Diagnostic Criteria (ICDC) were proposed which classified AIP into types 1 and 2. Type 1 is featured histologically by lymphoplasmacytic sclerosing pancreatitis (LPSP) and type 2 by idiopathic duct-centric pancreatitis. The ICDC included five cardinal features of AIP including parenchymal imaging, ductal imaging, serology, other organ involvement (OOI), histology of pancreas and response to steroid therapy. Each criterion was further classified into two levels (levels 1 and 2). The aim of the proposal of ICDC was intended to improve the diagnosis of AIP. AIP could be also classified into focal-type and diffuse-type AIP according to the involvement of pancreatic enlargement. In clinical settings, it is more important and also difficult to differentiate focal-type AIP from PC. To date, it still lacks a simple parameter with absolute diagnostic value. Therefore, the use of combined parameters according to different diagnostic criteria in different countries exists.

The objective of this study is to evaluate the diagnostic performance (sensitivity, specificity and accuracy) of ICDC of AIP from differentiating PC in a prospectively collected cohort in Taiwan, compared with the two most commonly used criteria in our country before ICDC was made (revised HISORt and Asian criteria). The diagnostic role of each cardinal feature of ICDC will be compared with the revised HISORt and Asian criteria in diffuse-type and focal-type AIP, respectively.

Methods
Study participants
Between January 1996 and December 2013, we consecutively collected 188 patients with AIP (95 men and 93 women) at the National Taiwan University Hospital, a tertiary referred centre and also the largest medical centre for management of pancreatic diseases in Taiwan. All patients with AIP fulfilled at least one of the HISORt criteria (158/188, 84%), or the Asian diagnostic criteria (162/188, 86.2%), or the ICDC criteria (168/188, 89.4%) for AIP. All patients were type 1 AIP. All patients were followed up for at least 12 months. A total of 130 consecutive patients (65 men and 65 women) with cytological or/and pathologically confirmed adenocarcinoma of the pancreas were enrolled as a control group. The patients’ mean age was 51.4 (range, 33–78 years) and 60.9 years (range 32–78 years), respectively, in patients with AIP and PC. All the patients’ medical charts were reviewed and the patients’ demographic data, including age, gender, serological studies, image studies and clinical manifestations, etc, were collected.

Findings of diagnostic criteria of ICDC
We categorised all patients with AIP and PC as level 1 findings, level 2 findings or neither for each of the five criteria (parenchymal imaging, ductal imaging, serology, other OOI, histology of pancreas and response to steroid therapy) according to ICDC. For parenchymal imaging, the frequencies of diffuse enlargement, focal enlargement and atypical imaging (pancreatic duct dilation or calcification or atrophy) were evaluated. Enlargement of the pancreas was defined as when the width of the pancreatic body or tail exceeds two-thirds of the transverse diameter of the vertebral body or if the width of the pancreatic head exceeds the full transverse diameter of the vertebral body. For ductal imaging, the MPD diameter was measured by the use of abdominal CT and/or MR cholangiopancreatography (MRCP) and/or endoscopic retrograde cholangiography. The frequencies of long stricture without marked upstream dilation, of multiple strictures without marked upstream dilation, of segmental/focal narrowing without marked upstream dilation and of marked upstream dilation of the MPD were evaluated. Pancreatic duct dilation was defined as the diameter of the MPD exceeding 5 mm. Parenchymal and ductal imaging scans were analysed by three experts (M-CC, Y-TC and P-CL).

For OOI, the frequencies of segmental/multiple proximal bile duct stricture, retroperitoneal fibrosis, symmetrically enlarged salivary/lachrymal glands and radiological evidence of renal involvement were evaluated. Bile duct stricture was evaluated by MRCP or ERC, or percutaneous transhepatic cholangiography. Retroperitoneal fibrosis was evaluated by contrast enhanced CT or MRI. Symmetrically enlarged salivary/lachrymal glands were evaluated by physical examination or/and CT.

For histology criteria of pancreas in AIP, there were 25 patients received pancreatectomy. Six patients received a biopsy of the pancreas. Pancreatic histology was evaluated by an experienced pathologist who was blinded to the other data.

Regarding steroid therapy, 126 patients (67%) received corticosteroid therapy as the initial treatment of AIP. The dose for induction therapy was started from 30 to 40 mg/ day for 2–4 weeks and then tapered 5 mg/week gradually. The treatment response of steroid was nearly 98%.

We evaluated the frequencies of level 1 findings, level 2 finding and neither of each criterion in all patients with AIP and PC. Sensitivity, specificity and accuracy were compared between ICDC, revised HISORt criteria and Asian criteria. We also evaluated the cases which did not fit the diagnosis with AIP according to each criterion.

Statistical analysis
We calculated the sensitivity, specificity and accuracy of each diagnostic criterion. The between-group demographic data were compared by the Student unpaired t test for continuous data and by the $\chi^2$ test for categorical data. Receiver-operator characteristic (ROC) curves and area under the curves (AUCs) were estimated after logistic regressions in different criteria in all AIP and focal-type AIP subgroups and presented with corresponding
RESULTS
Parenchymal imaging
Of the 188 patients, 90 (50.5%) and 93 (49.5%) with AIP were categorised as levels 1 and 2, respectively. All patients with PC were classified as level 2 (table 1). The mean serum IgG4 level was 346.6±56.2 mg/dL, which was statistically significantly higher than that in patients with PC, 119.2±23.9 mg/dL. The frequencies of serum level above 280 (level 1) and 140 mg/dL (level 2) were significantly higher in patients with AIP (p<0.001.)

Other organ involvement
Sixty-three (33.5%) of 188 patients with AIP were categorised as level 1 and none of the 130 patients with PC were categorised as level 1 or 2 (table 1). Proximal bile duct stricture was observed in 53 patients (28.2%) with AIP, and retroperitoneal fibrosis was observed in 5 patients (2.7%) with AIP, both level 1 findings. Enlarged salivary/lacrimal glands were observed in 58 patients (27.6%) with AIP and renal involvement in 9 patients (4.8%); these met level 2 criteria. Neither enlarged salivary/lacrimal glands nor renal involvement was observed in any patients with PC.

Histology of pancreas
There were 31 (16.4%) patients with AIP who were categorised as level 1 or 2 and none of the 130 patients with PC were categorised as level 1 or 2 (table 1). There were 28 patients (14.9%) with level 1 evidence and 3 patients (1.6%) with level 2 evidence in histological LPSP. No patients with PC fulfilled the level 1 or 2 criteria.

Response to steroid therapy
Of the 128 patients who received steroid treatment as the initial treatment (induction therapy), 126 (98.4%) showed steroid response with improvement clinically, serologically and morphologically. Two of the patients with diffuse pancreatic enlargement and narrowing of the MPD received steroids but there was no morphological response. Endoscopic ultrasound-guided fine-needle aspiration (EUS-FNA) was performed but no malignancy was detected. These two patients discontinued steroid use after 3-month treatment and were followed up regularly for 18 and 20 months, respectively, but no malignancy was documented, although the pancreatic enlargement did not subside.

Diagnostic on the basis of ICDC and revised HISORt and Asian criteria
Of the 188 patients with AIP, 116 patients were diagnosed as definite AIP; 35 patients were diagnosed as probable AIP and 17 were classified as not otherwise specified (NOS). Among these patients, the primary basis for diagnosis was histology in 31 patients, response to steroid therapy in 126 patients and imaging in 188 patients. There were 20 patients in this study deniable for type 1 AIP. All patients with PC were deniable for type 1 AIP on the basis of ICDC. The sensitivity, specificity and accuracy of ICDC for type 1 AIP were 89.4%,
100% and 93.7% (table 2). Using revised HISORt criteria, 158 patients (84%) were diagnosed as definitive AIP. Among these 158 patients, the primary basis of diagnosis was diffuse type in 95 patients (60.1%) and histology-based diagnosis in 31 patients (19.6%). There were 30 patients who were deniable for AIP based on the revised HISORt criteria in this study. All of the patients with PC were deniable for AIP based on the revised HISORt criteria. The sensitivity, specificity and accuracy of the revised HISORt criteria were 84%, 100% and 90.5% (table 2). Using Asian criteria, 162 patients (86.2%) were diagnosed as AIP. In total, 143 patients (88.3%) were diagnosed based on imaging plus serology; 31 patients (19.1%) were diagnosed based on histopathology and 126 patients (77.8%) were diagnosed based on steroid treatment response. There were 26 patients who were deniable for AIP based on the Asian criteria in this study. All of the patients with PC were deniable for AIP based on the Asian criteria. The sensitivity, specificity and accuracy of the revised HISORt criteria were 86.2%, 100% and 91.8% (table 2).

**Deniable cases on the basis of ICDC**

There were 20 patients who were deniable for ICDC in this study. All these 20 patients could be diagnosed by the Asian criteria. Among them, there were two patients who could be diagnosed both by the HISORt and Asian criteria. They included 6 and 14 cases with level 1 or 2 parenchymal imaging; 6 and 12 cases with level 1 or 2 ductal imaging; 9 cases with level 2 serology; 11 patients with level 1 OOI. There was one patient with a steroid treatment response. The most common factor which lead to deniable of ICDC or revised HISORt criteria was the serology criterion. There were 18 out of 20 patients who had autoantibodies which could be one of the items in the Asian serology criterion. The ICDC and HISORt only adapted IgG4 level alone as the serology criterion.

**ICDC criteria in focal-type and diffuse-type AIP**

The comparisons of frequencies of levels 1 and 2 findings in ICDC in focal-type and diffuse-type AIP are shown in table 3. The frequencies of level 1 or 2 features in ductal imaging were significantly lower in focal-type AIP (73.1% vs 97.9%, p<0.001, table 3). The frequencies of level 1 or 2 features in serology were also lower in focal-type (55.4% vs 69.3%, p=0.075). The frequencies of any level of OOI in focal-type AIP were higher than those in diffuse-type AIP (81.7% vs 53.7%, p<0.0001). The frequencies of any histological evidence of LPSP in focal-type AIP were higher than those in diffuse-type AIP (23.7% vs 9.5%, p=0.01) in our study (table 3).

**Sensitivity, specificity and accuracy of ICDC, revised HISORt criteria and Asian criteria**

The sensitivity, specificity and accuracy of ICDC were 84.9%, 100% and 93.7% (table 2). The sensitivity, specificity and accuracy of the revised HISORt criteria were 78.5%, 100% and 91.0% (table 2). The sensitivity, specificity and accuracy of the Asian criteria were 73.1%, 100% and 88.8% (table 2).

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**Table 2** Comparison of diagnostic performance of different criteria for AIP from pancreatic cancer

<table>
<thead>
<tr>
<th></th>
<th>Case number</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>All AIP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fit Asian</td>
<td>162</td>
<td>86.2</td>
<td>100.0</td>
<td>91.8</td>
</tr>
<tr>
<td>Fit revised HISORt</td>
<td>158</td>
<td>84.0</td>
<td>100.0</td>
<td>90.5</td>
</tr>
<tr>
<td>Fit ICDC</td>
<td>168</td>
<td>89.4</td>
<td>100.0</td>
<td>93.7</td>
</tr>
<tr>
<td><strong>Focal-type AIP</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fit Asian</td>
<td>68</td>
<td>73.1</td>
<td>100.0</td>
<td>99.6</td>
</tr>
<tr>
<td>Fit revised HISORt</td>
<td>73</td>
<td>78.5</td>
<td>100.0</td>
<td>96.6</td>
</tr>
<tr>
<td>Fit ICDC</td>
<td>79</td>
<td>84.9</td>
<td>100.0</td>
<td>97.3</td>
</tr>
</tbody>
</table>

AIP, autoimmune pancreatitis; ICDC, International Consensus Diagnostic Criteria.

**Table 3** Frequencies of levels 1 and 2 findings in ICDC for focal-type and diffuse-type AIP

<table>
<thead>
<tr>
<th>Features</th>
<th>Focal-type AIP (n=93)</th>
<th>Diffuse-type AIP (n=95)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ductal imaging</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Level 1</td>
<td>16/74 (21.6%)</td>
<td>31/88 (35.2%)</td>
</tr>
<tr>
<td>Level 2</td>
<td>25/74 (33.8%)</td>
<td>30/88 (34.1%)</td>
</tr>
<tr>
<td>Level 1+2</td>
<td>41/74 (55.4%)</td>
<td>61/88 (69.3%)</td>
</tr>
<tr>
<td>Non-level 1, 2</td>
<td>33/74 (44.6%)</td>
<td>27/88 (30.7%)</td>
</tr>
<tr>
<td><strong>Serology</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Level 1</td>
<td>31 (33.3%)</td>
<td>32 (33.7%)</td>
</tr>
<tr>
<td>Level 2</td>
<td>45 (48.4%)</td>
<td>19 (20.0%)</td>
</tr>
<tr>
<td>Level 1+2</td>
<td>76 (81.7%)</td>
<td>51 (53.7%)</td>
</tr>
<tr>
<td>Non-level 1, 2</td>
<td>17 (11.3%)</td>
<td>44 (46.3%)</td>
</tr>
<tr>
<td><strong>OOI</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Level 1</td>
<td>20 (21.5%)</td>
<td>8 (8.4%)</td>
</tr>
<tr>
<td>Level 2</td>
<td>2 (2.2%)</td>
<td>1 (1.1%)</td>
</tr>
<tr>
<td>Level 1+2</td>
<td>22 (23.7%)</td>
<td>9 (9.5%)</td>
</tr>
<tr>
<td>Non-level 1, 2</td>
<td>71 (76.3%)</td>
<td>86 (90.5%)</td>
</tr>
</tbody>
</table>

AIP, autoimmune pancreatitis; ICDC, International Consensus Diagnostic Criteria; OOI, other organ involvement.
The ROC was calculated in AIP (figure 1) and focal-type AIP (figure 2). The AUC was 0.95 (95% CI 0.92 to 0.97) of ICDC (figure 1A), 0.91 (95% CI 0.92 to 0.97) of the revised HISORT criteria (figure 1B) and 0.93 (95% CI 0.92 to 0.97) of the Asian criteria (figure 1C). For focal-type AIP, the AUC was 0.93 (95% CI 0.88 to 0.97) of ICDC (figure 2A), 0.89 (95% CI 0.84 to 0.94) of the revised HISORT criteria (figure 2B) and 0.87 (95% CI 0.81 to 0.92) of the Asian criteria (figure 2C).

DISCUSSION
The sensitivity, specificity and accuracy of ICDC for all AIP were 89.4%, 100% and 93.7%. The sensitivity, specificity and accuracy of ICDC for focal-type AIP were 84.9%, 100% and 93.8%. Among the three criteria, the sensitivity and accuracy of ICDC were the best compared with the revised HISORT and Asian criteria with the same specificity (table 2). There were 12 patients with deniable revised HISORT criteria who were diagnosed as AIP in ICDC (10 definite, 1 probable and 1 NOS of ICDC). There were 26 patients with deniable Asian criteria who were diagnosed as AIP in ICDC (11 definite, 14 probable and 1 NOS in ICDC). The ICDC showed higher sensitivity than the revised HISORT and Asian criteria. The reason why patients deniable for revised HISORT or Asian criteria but fit ICDC diagnosis was partially caused by the different definition of parenchymal imaging between these criteria. ICDC includes atypical imaging and classifies atypical imaging as S2 level in the parenchymal imaging criterion. That is one reason why ICDC had higher sensitivities in diagnosis of AIP in general. The wider range of imaging criteria improved the diagnostic sensitivity but did not decrease either the specificity or the accuracy of ICDC. All the three criteria showed high specificity (table 2). The ICDC showed higher accuracy than the Asian and revised HISORT criteria in our population. Recently, studies from the
Japanese population also demonstrated better accuracy of ICDC compared with other criteria, including the Japanese pancreatic society criteria.\(^{15-18}\) Although ICDC is considered to be superior to various other criteria, it seems to be too complicated to handle for clinicians. The Japanese have therefore proposed revised diagnostic criteria by the Japanese pancreatic society very recently.\(^{19}\) It would be interested to whether the performance could even be better or easily to use in clinical practice compared to ICDC criteria. At this time moment, we did not have suitable simple amendment of ICDC of type I AIP which composed of a heterogenous population clinically in our country. With the advance of better understanding of the pathogenesis of the disease, to simplify the diagnostic criteria might be feasible and needed for clinicians.

In the ductal imaging criterion, 161 (92.5\%) of 188 patients with AIP and 13 (1\%) of 130 patients with PC were categorised as level 1 or 2, respectively. The specificity of ductal imaging is high. One of the reasons for this high specificity is that patients with marked upstream MPD dilation (>5 mm) were excluded from level 1 or 2. In the present study, 28 of 31 patients with PC who showed focal MPD stricture were excluded from level 2 because of marked upstream MPD dilation. On the other hand, 27 (14.4\%) of 188 patients with AIP showed marked upstream MPD dilation. The frequency of any level 1 or 2 evidence in ductal imaging is close to the recently reported study (7/62, 11.3\%) by Nishino et al.\(^ {20}\) Naitoh et al.\(^ {21}\) also reported that a maximal diameter of the upstream MPD less than 5 mm was an appropriate cut-off point to differentiate mass-forming AIP from PC. In our study, we also use the 5 mm as a cut-off point to differentiate focal-type AIP and PC. Therefore, we consider that a 5 mm diameter of the upstream MPD is appropriate to discriminate AIP from PC. In the present study, four patients with PC fulfilled the level 1 serological criterion. Marked upstream MPD dilation (5 mm) was observed in these patients. If this exclusion criterion (marked upstream MPD) did not exist, this patient would have fulfilled the level 2 criterion for ductal imaging, and we would have misdiagnosed these four patients with PC as definitive type 1 AIP under ICDC. Therefore, we consider that this exclusion criterion of ductal imaging is useful for excluding PC.

The value of serum IgG4 as a serological marker of AIP was first established in 2001.\(^ {22-25}\) Hamano et al.\(^ {23}\) reported that sensitivity and specificity for differentiating AIP from PC were 90.2\% and 97.5\%. In the present study, the sensitivity and specificity of serum IgG4 (>140 mg/dL) were 79.6\% and 92.6\%, respectively. The cause of the wide range of sensitivity in the reported series might be caused by the combined analysis of types 1 and 2 in these studies. The distribution between types 1 and 2 AIP might affect the value of IgG4. Ghazale et al.\(^ {26}\) reported that the sensitivity of elevated serum IgG4 (>140 mg/dL) for PC was 10\%, and that of twofold elevation (level 1 ICDC serology criteria) was 1\%. Our present study showed that the sensitivity and specificity of twofold elevation of serum IgG4 were 36.7\% and 95.2\% in our type 1 AIP. Serum IgG4 is the only used serology marker in ICDC. In the Asian criteria, they adapted IgG, IgG4 and the presence of an autoantibody as serological criteria. In our patients with type 1 AIP, if we add the presence of an autoantibody also as a surrogate marker in serology, all of the patients deniable for ICDC could be diagnosed by this modification.

IgG4-related disease (IgG4-RD) is a new disease entity characterised by elevated serum IgG4 concentration and/or tissue infiltration by IgG4-positive cells.\(^ {24}\) Type 1 AIP is regarded as a part (pancreatic manifestation) of IgG4-RDs. In the present study, level 1 or 2 OOI was observed in 127 (67.6\%) of the patients with AIP and in none of those with PC. Therefore, this OOI criterion has high specificity for type 1 AIP.

In the present study, all resection specimens (27 patients) fulfilled the level 1 criterion. One of four biopsied specimens guided by CT fulfilled level 1, with the rest diagnosed as level 2. The present findings suggest that obtaining histopathological evidence of type 1 AIP by a biopsied specimen is difficult. EUS-FNA was considered useful for the differentiation from PC but might be insufficient for tissue collection to diagnosis as level 1. Kanno et al.\(^ {25}\) reported that 14 and 6 patients, respectively, of 25 patients were judged to have levels 1 and 2 histological findings by trucut biopsy under endoscopic ultrasound (EUS) guidance. We did not perform EUS-trucut biopsy in this study because the needle is not available and the cost is not covered by our health insurance. Further studies for the feasibility and necessarily of EUS-FNA in diagnosing type 1 AIP in ICDC histological criteria need to be studied.

Diffuse and focal enlargement of the pancreas is a characteristic feature of AIP in parenchymal imaging. ICDC included patients with atypical parenchymal imaging as level 2. For diffuse-type AIP, the diagnostic accuracy of ICDC and Asian and HISORt criteria was over 95\%, much better than that in focal-type AIP. This observation is reasonable for us to understand because it is very rare to have PC involving the whole pancreas in clinical practice. The diagnosis sensitivity in diffuse-type AIP was also higher than those in focal-type AIP in these three diagnostic criteria (table 2). In diffuse-type AIP, the Asian criteria were most sensitive with a sensitivity of 98.9\%, followed by ICDC (93.7\%) and the HISORt criteria (89.5\%). There were four patients with diffuse-type AIP who did not have elevated was the least sensitive criterion in the diagnosis of diffuse type with sensitivity only 89.5\%. The four diffuse-type AIP patients were deniable for HISOrt but fit ICDC NOS owing to these four patients did not have any collateral evidence. These four patients could be diagnosed by Asian criteria as they have presence of autoantibodies which made they fit the serology criteria in Asian diagnosis setting but not fit the serology criteria in ICDC and HISORt criteria. These four patients had received steroid response as the initial
treatment and they all had disease relapse in their follow-up. The increase of sensitivity in the Asian criteria in diagnosing diffuse-type AIP is relevant to the wide range of definition in the serology criterion (IgG, IgG4 or/and autoantibodies), compared to the use of IgG4 alone as the serology criterion in HISORt and ICDC.

It is a greater challenge to differentiate focal-type AIP with PC, compared to differentiate diffuse-type AIP from PC. In this study, we have 93 patients with focal-type AIP. The ductal imaging, serology and OOI were different from those for diffuse-type AIP. Focal-type AIP had less frequency of level 1 or 2 presentations of ductal imaging and serology. In contrast, focal-type AIP had higher frequencies of level 1 or 2 presentations of OOI. These clinical observations remind us that the collateral evidence of AIP in focal-type AIP is mostly in OOI, but not serology. Detailed physical examination and history taking and imaging interpretation imaging other than pancreas could give some hint to increase the sensitivity of diagnosis of focal-type AIP.

In focal-type of AIP, the diagnostic sensitivity of ICDC (84.9%) was higher than those of the revised HISORt (78.5%) and Asian criteria (73.1%). There were 14 (15.1%) patients with focal-type AIP deniable for the ICDC criteria. All these 14 patients could be diagnosed by the Asian criteria. There were 12 patients with the presence of autoantibodies and 9 patients with a serum IgG4 level above 140 mg/dL. Among them, there were 12 patients with level 2 criteria and 2 patients without level 1 or 2 criteria in ductal imaging. In OOI, there were nine patients with level 2 criteria and five patients without level 1 or 2 criteria. All the patients with focal-type AIP deniable for the Asian criteria could be diagnosed by the ICDC criteria.

In conclusion, ICDC shows high sensitivity, specificity and accuracy in the diagnosis of type 1 AIP. In focal-type AIP, ICDC is still the best in sensitivity, specificity and accuracy. The diagnostic sensitivity in focal-type AIP is not as good as diffuse-type AIP in all the three adapted criteria (ICDC and Asian and revised HISORt criteria). How to improve the sensitivity of diagnosis of focal-type AIP is the issue that needs to be resolved in the future.

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