Management of Incidental Pancreatic Cystic Lesions

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Keywords
Pancreatic cystic lesion · Incidental finding · Intraductal papillary mucinous neoplasia · Mucinous cystic neoplasia · Serous cystadenoma · Endoscopic ultrasound · Cross-sectional imaging

Summary
Background: Pancreatic cystic lesions (PCL) are common. They are increasingly detected as an incidental finding of transabdominal ultrasound or cross-sectional imaging. In contrast to other parenchymal organs, dysontogenetic pancreatic cysts are extremely rare. In symptomatic patients the most frequent PCL are acute and chronic pseudocysts. The majority of incidental cystic lesions, however, are neoplasias which have different risks of malignancy. Methods: PubMed was searched for studies, reviews, meta-analyses, and guidelines using the following key words: ‘pancreatic cystic lesions’ OR ‘cystic pancreatic lesions’ OR ‘intraductal papillary mucinous neoplasia’ OR ‘mucinous cystic neoplasia’ OR ‘pancreatic cyst’ OR ‘pancreatic pseudocyst’) AND (management OR treatment OR outcome OR prognosis OR diagnosis OR imaging OR ‘endoscopic ultrasound’ EUS-FNA OR EUS OR ‘endoscopic ultrasonography’ OR CT OR MRI). Retrieved papers were reviewed with regard to the diagnostic and therapeutic management of incidental PCL. Results: In addition to clinical criteria, transabdominal ultrasonography including contrast-enhanced ultrasonography, cross-sectional radiological imaging, and endoscopic ultrasound (EUS) are used for diagnostic characterization and risk assessment. EUS plays an outstanding role in differential diagnosis and prognostic characterization of incidental PCL. In a single examination it is possible to perform high-resolution morphological description, perfusion imaging, as well as fine-needle aspiration of cyst content, cyst wall, and solid components. An international consensus guideline has defined worrisome and high-risk criteria for the risk assessment of mucinous pancreatic cysts, which are mainly based on the results of EUS and cross-sectional imaging. Nevertheless, despite diagnostic progress and guideline recommendations, differential diagnosis and management decisions remain difficult. This review will discuss problems in and approaches to the diagnosis of incidental PCL. Conclusion: An evidence-based algorithm for the diagnosis of incidental PCL is proposed.

Introduction: Chance and Challenge

Pancreatic cystic lesions (PCL) are common. Two Japanese autopsic studies detected small pancreatic cysts (>1–2 mm) in 73 of 300 (24.3%) and 378 of 1,374 (27.5%) consecutive autopsy cases, respectively \cite{1, 2}. The incremental dissemination and technical development of modern imaging methods facilitates the detection of PCL by transabdominal ultrasonography (TUS), computed tomography (CT), or magnetic resonance imaging (MRI). A large retrospective Japanese study reviewed the TUS findings of 12,112 consecutive patients, among them 1,012 patients with and 11,100 patients without end-stage renal disease. The prevalence of PCL in both groups proved to be 9.3 and 1.3%, respectively, with a relatively high percentage of potentially malignant mucinous neoplasms among them (2.8 and 0.2%, respectively) \cite{3}. CT and MRI studies in large cohorts of asymptomatic persons revealed unsuspected pancreatic cysts in 2.4–13.5% of the cases. There is a strong correlation of increasing age and prevalence of PCL \cite{4–6}. A recent study compared follow-up data of 2,034 patients with PCL detected incidentally at CT or MRI with follow-up data of a matched control group (n = 6,018) without PCL, showing that the detection of a PCL is associated with a 3-fold increased risk to develop pancreatic adenocarcinoma \cite{7}. However, not all inciden-
tally detected PCL carry an elevated risk of malignant transformation. There is a broad spectrum of incidental PCL, comprising 25 different types defined by the World Health Organization, among them four types of primarily cystic neoplasias: serous cystic adenoma (SCA), mucinous cystic neoplasia (MCN), as well as main-duct (MD) and branch-duct (BD) intraductal papillary mucinous neoplasia (IPMN). Solid pseudopapillary neoplasia (SPN) and cystic variants of ductal adenocarcinoma, neuroendocrine tumor, and acinus cell cancer are the best known, though rare examples of PCL resulting from necrosis and cystic degeneration of solid tumors. Dysontogenetic cysts, which are common in the kidneys and the liver, are rare in the pancreas. Contrary to symptomatic patients, pseudocysts are a very rare diagnosis in asymptomatic patients. Other non-neoplastic PCL are very rare: lymphoepithelial cysts (LEC), dermoid cysts (DC), epidermoid cysts (ECIS), retention cysts, mucinous non-neoplastic cysts (MNC), duplication cysts of the foregut, cystic hamartoma, and cystic lymphangioma. MD-IPMN, BD-IPMN, and MCN are mucinous PCL and precursor lesions of pancreatic adenocarcinoma. Their premalignant risk varies according to the particular type of lesion, its size, and some further features like (in IPMN) their histological subtype [8–10]. IPMN are subclassified into neoplasias with intestinal, pancreatobiliary, oncocytic, or gastric cellular differentiation. Intestinal, pancreatobiliary, and oncocytic subtypes predominantly involve the main pancreatic duct within the pancreatic head and body, with the pancreatobiliary subtype being the most aggressive one and developing into tubular adenocarcinoma [11, 12]. A majority of these neoplasias is already malignant at the time of diagnosis or easily progresses to invasive adenocarcinoma. Nevertheless, prognosis of MD-IPMN in non-invasive and minimally invasive stages is more favorable compared to ductal adenocarcinoma, preferably in the intestinal subtype, which develops into colloid (mucinous) adenocarcinoma [11, 13–15]. Gastric-type IPMN occur predominantly in the branch ducts, preferentially of the pancreatic head. They are frequently multifocal and have a distinctively lower risk (at the time of diagnosis approximately 15–20%) as well as slower course of progressing to invasive adenocarcinoma. In patients with gastric-type IPMN, however, simultaneous or metachronous development of ductal adenocarcinoma may occur, thus worsening the prognosis considerably [11–13, 16–19]. Prospective data on the natural history and rate of malignant transformation of mucinous precursors of pancreatic cancer are rare, and epidemiological, surgical, and retrospective data show conflicting results [17–23].

Therefore, the incidental discovery of PCL at the same time is increasingly becoming a chance as well as a challenge for modern health care systems. On the one hand, the early detection of cystic precursor lesions in asymptomatic persons opens the window widely for the prevention of a substantial portion of pancreatic cancers. On the other hand, differential diagnosis is demanding, and natural history is not sufficiently understood. In older patients with significant comorbidity and slowly developing mucinous PCL, a low or moderate risk of malignancy may be outperformed by the risks of pancreatic surgery.

Methods

A systematic literature search was performed to identify studies, reviews, meta-analyses, and guidelines evaluating diagnosis, treatment, and prognosis of PCL. PubMed was searched using the following keywords: (‘pancreatic cystic lesions’ OR ‘cystic pancreatic lesions’ OR ‘intraductal papillary mucinous neoplasia’ OR ‘mucinous cystic neoplasia’ OR ‘pancreatic cyst’ OR ‘pancreatic pseudocyst’) AND (management OR treatment OR outcome OR prognosis OR diagnosis OR imaging OR ‘endoscopic ultrasound’ EUS-FNA OR EUS OR ‘endoscopic ultrasonography’ OR CT OR MRI). Retrieved papers were reviewed with regard to the diagnostic and therapeutic management of incidental PCL.

Diagnostic Tools

Asymptomatic PCL are most often initially detected on TUS or abdominal CT. The initial imaging modality gives relevant basic information about size, localization, and gross morphological appearance. For diagnostic characterization and risk assessment, however, detailed morphological information is necessary, in particular on ductal communication, cyst content, mural nodules, and septae.

Cross-Sectional Imaging

A prospective study proved a high accuracy and concordance of two readers of multidetector CT (MDCT) scans for the preoperative stratification of PCL into mucinous and non-mucinous types (82 and 85%) and the prediction of their biologic aggressiveness (85 and 86%). However, predictive values of MDCT were superior for lesions >30 mm and non-mucinous lesions [24]. Two retrospective studies suggested an equivalent accuracy of MDCT and MRI for characterizing small PCL as benign or malignant as well as mucinous or non-mucinous, and for suggesting a specific diagnosis. Whereas the accuracy for classification according to the risk of malignancy may be regarded as sufficient (75–86%), the accuracy for determining a specific diagnosis remained disappointing (40–84%) [25, 26]. In another retrospective study evaluating the accuracy of CT versus MRI and magnetic resonance cholangiopancreatography (MRCP) in the characterization of IPMN, pathologically proven ductal communication was detected by MRI in 73% of the cases, by CT, however, only in 18% of the cases. CT tended to overestimate an involvement of the pancreatic duct when compared with MRCP and surgical pathology. Moreover, CT was inferior to MRCP in identifying small-branch duct cysts and regarding the recognition of multifocality [27].

Ultrasoundography

The diagnostic value of TUS has been evaluated only in preliminary studies. One study showed a high correlation of near-isovoxel
ultrasound using matrix transducers and MRCP in evaluating the ductal communication of PCL [28]. Contrast-enhanced ultrasound (CEUS) has been prospectively shown to discriminate between pseudocyst and cystic neoplasia with a very high diagnostic accuracy, outperforming TUS without contrast enhancement [29, 30]. A retrospective study suggested that CEUS compares favorably with MRI in displaying anatomic features of PCL (septae, nodules), and demonstrated a close correlation between CEUS findings and results of surgical pathology [31].

**Endoscopic Ultrasound (EUS) and EUS-Guided Fine-Needle Aspiration**

Due to its unsurpassed spatial resolution, feasibility of vascularity analysis, and opportunity to perform fine-needle aspiration (FNA), EUS is regarded to be the most valuable diagnostic tool for the classification and prognostic evaluation of PCL [32–34]. While one retrospective study showed a comparable accuracy of EUS and MRI in the characterization of PCL and prediction of malignancy [35], most other studies are in favor of EUS. Comparing the performance of CT, MRI, and EUS in characterizing PCL in 145 patients, EUS more frequently identified PCL to be multifocal and, even more important, their presence in different surgical fields. Communication with the pancreatic duct was detected significantly more frequently by EUS compared with CT and MRI. For the detection of mural nodules and septations, EUS performed significantly superior compared with CT but not with MRI [36]. In another large cohort of patients with PCL, in which the final diagnosis was established by surgical pathology, EUS(-FNA) was superior to cross-sectional imaging in accurately classifying a cyst as neoplastic. The incremental increase in diagnostic yield of EUS(-FNA) over CT and MRI for the prediction of a neoplastic cyst in this study was 36 and 54%. Again, EUS detected considerably more mural nodules than cross-sectional imaging. Nonetheless, the neoplastic nature and malignancy of a PCL were still underestimated by EUS(-FNA) in 23 and 16% of the cases, respectively [37]. To increase the yield of EUS-FNA of PCL, aspiration not only of the cyst content but also of the cyst wall and of possible solid components was suggested [38, 39]. EUS-FNA of PCL has a higher complication rate (bleeding, pancreatitis, infection, surgical complications: 5–6% in prospective studies) compared to EUS-FNA of solid pancreatic lesions (2.4% in prospective studies) [40–43]. Two retrospective studies of large cohorts of patients undergoing resection of neoplastic pancreatic cysts did not observe an increased risk of peritoneal tumor seeding by performing EUS-FNA [40, 44]. To minimize the risk of cyst infection, complete aspiration of the cyst content and peri-interventional antibiotic treatment are recommended for EUS-FNA of PCL [41, 45].

**Diagnostic Criteria**

**FLAG(S)**

Following initial detection of a PCL, basic facts are known and should be considered: age and gender of the patient, symptoms, localization, and size of the lesion. These simple data (‘FLAGS – i.e. frequency, localization, age, gender, symptoms – criteria’) result in a relatively high pretest accuracy for various PCL and help to stratify the risk of malignancy (fig. 1, table 1). For example, it is virtually impossible that a PCL located within the pancreatic head of a 73-year-old man represents a MCN.

**Localization**

Localization within the pancreas may give some clues for the diagnosis of PCL. Typically, BD-IPMNs are located within the pan-

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**Table 1. Diagnostic pretest probability: FLAG(S) criteria of PCL (modified from [33, 34])**

<table>
<thead>
<tr>
<th>Pseudocyst</th>
<th>SPN</th>
<th>SCA</th>
<th>MCN</th>
<th>BD-IPMN</th>
<th>MD-IPMN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frequency</td>
<td>very common</td>
<td>very rare</td>
<td>moderate</td>
<td>moderate</td>
<td>rare</td>
</tr>
<tr>
<td>Localization</td>
<td>unifocal (predominantly head)</td>
<td>unifocal, variable</td>
<td>unifocal (microcystic: 70% left)</td>
<td>unifocal (70% left)</td>
<td>&gt;60% multifocal, &gt;70% head, branch ducts</td>
</tr>
<tr>
<td>Median age</td>
<td>variably</td>
<td>=20 years</td>
<td>=60 years</td>
<td>=45 years</td>
<td>=65 years</td>
</tr>
<tr>
<td>Gender</td>
<td>m &gt; f</td>
<td>f &gt;&gt; m (9:1)</td>
<td>microcystic: f &gt; m (7:3); macrocystic: m &gt; f (3:2)</td>
<td>f &gt;&gt; m (20:1)</td>
<td>m = f</td>
</tr>
<tr>
<td>Symptoms</td>
<td>often, history of pancreatitis is obligatory</td>
<td>rare, only in large tumors, history of pancreatitis &lt;10% in MCN</td>
<td>in up to one-third of cases mild (recurrent) pancreatitis</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
creatic head, preferentially within the uncinate process. In up to two-thirds of the cases, however, smaller cysts can also be found in other parts of the pancreas [36, 46]. MD-IPMNs commonly develop from the pancreatic head region. Conversely, approximately 70% of microcystic SCA and most SPN are located outside the pancreatic head [33].

**Age/Gender**

Two types of cystic pancreatic neoplasias, i.e. SPN and MCN, almost exclusively occur in women. SPN is the least frequent cystic neoplasm, occurring with very rare exceptions only in girls and young women with a median age of 20 years. MCN is most commonly observed in middle-aged women.

**Symptoms**

Pseudocysts are associated with acute or chronic pancreatitis; therefore, they are only very rarely detected incidentally. Incidental PCL are smaller than symptomatic cysts and are found predominantly in older patients [47]. The majority of IPMNs are not accompanied by acute symptoms [48]. Due to transient mucinous ductal obstruction, however, IPMN of both types may cause mild and often recurrent flares of acute pancreatitis in 7–34.6% of the cases [49–52]. In addition to the cyst features, morphology of the pancreatic parenchyma as well as diameter and contour of the main pancreatic duct should be paid heed to (table 2) [33, 34].

Microcystic SCA is characterized by multiple closely agglomerated microcysts (≤20 mm, often only 1–2 mm) separated by thin but highly vascularized septae (‘honeycombing’). Sometimes a central scar (rarely with calcification) may be found. Criteria of chronic pancreatitis are lacking, and the main pancreatic duct is not involved (fig. 2).

However, differential diagnosis of the macrocystic type of SCA (multilocular cyst > 2 cm with thin wall/septae without ductal communication) to BD-IPMN and of the macrocystic type (unilocular cyst, often lobulated, thin wall) to pseudocyst or MCN may be challenging. BD-IPMN is suspected in cases with grape-like agglomerations of cysts within the pancreatic head (‘cyst by cyst’), in particular if communication between neighboring cysts as well as between cysts and side branches of the main pancreatic duct may be displayed (fig. 3). MCN are unilocular cysts with a distinct, highly vascularized wall and septae or cysts within the cyst. MD-IPMN is characterized by complete or segmental cystic dilation of the main pancreatic duct without underlying stricture. TUS or EUS may delineate hyperechoic luminal layering or mucinous plugs. In approximately 50% of MD-IPMN a dilated orifice of the papilla with mucinous secretion (patulous papilla, ‘fish-mouth papilla’) is observed (fig. 4) [55, 56]. Solid mural nodules are typical features and high-risk markers of mucinous cystic neoplasms (MCN, IPMN). However, the detection rate for small mural nod-
ules is unsatisfactory when using cross-sectional radiological imaging, and discrimination from mucin plugs is challenging [57]. Contrast-enhanced EUS was suggested to increase the diagnostic accuracy to detect neoplastic PCL, to discriminate mural nodules from mucin plugs, and to determine growth patterns of mural nodules (fig. 4, 5) [58–60].

Cyst Fluid Analysis

EUS-FNA allows optical inspection, biochemical analysis (amylase or lipase; carcinoembryonic antigen (CEA)), and cytological examination of cyst fluid (table 3). Typically the pseudocysts’ content is a muddy brown fluid with low CEA concentration, containing neutrophils and/or histiocytes [61]. The gross appearance of aspirates of cystic lymphangioma (milky fluid) and of LEC, DC, and ECIS (thick milky, creamy, or frothy) may afford diagnosis [62]. Viscosity of fluid may be obvious if a string of fluid can be lifted with a needle from the slide (‘string sign’) [63]. For biochemical analysis only 0.5 ml of cyst fluid is necessary. Amylase or lipase are used as surrogate markers for the communication between PCL and pancreatic duct system. In a recent study, however, cyst fluid amylase was significantly higher in pseudocysts compared with MCN and IPMN but did not differ between IPMN and MCN [64]. CEA is a valid marker for mucinous pancreatic cysts but does not correlate with malignancy [65]. A high CEA concentration of cyst fluid was measured not only in MCN and IPMN but also in LEC and MNC. An international multicenter study figured out the optimal cut-off value of 192 ng/ml for the differentiation between mucinous and non-mucinous cysts (sensitivity 73%; specificity 84%; accuracy 79%) [66]. However, recent single-center studies reported cut-off values of 50, 67, and 110 ng/ml (accuracy 85, 84, and 86%, respectively) [64, 65, 67]. A recent meta-analysis including 18 studies with 1,438 patients revealed a pooled sensitivity and specificity of cyst fluid CEA levels of 63 and 88%, respectively [68]. CEA level in cyst fluid depends on the epithelial differentiation of IPMN. Recently, it was shown to be highest in the gastric subtype, followed by the pancreatobiliary and the intestinal subtype. CEA level in IPMN of the oncocytic subtype
was found to be as comparably low as in non-mucinous cysts [69]. Preliminary data suggest that the CA 125 level may be helpful to differentiate between MCN (high) and IPMN (low) [67]. The incremental value of molecular (DNA) over biochemical analysis of cyst fluid is low. For most parameters (DNA content, mutations of kRAS and GNAS, loss of heterozygosity mutations), specificity is high, while sensitivity is reported to be only between 20 and 50%. The combination of molecular analysis and CEA or cytology results in a higher diagnostic performance for the diagnosis of neoplastic mucinous PCL than either of the individual tests [70–72]. The value of cytology is limited by the low cellularity of PCL. Specificity of diagnosis of a malignant PCL is sufficient but sensitivity is very low (in meta-analyses: 88–93% and 54–65%, respectively) [68, 73, 74]. High-grade atypia should be included in the definition of ‘positive cytology’ for a high risk of malignancy [75, 76]. Mucin expression has a limited accuracy for the diagnosis of mucinous neoplasia [77]. The mucin expression profile may be used to differentiate between the various epithelial subtypes of IPMN as well as for risk assessment (table 3). MUC5AC is expressed in MCN and in all epithelial subtypes of IPMN, and therefore may be an additional marker to distinguish MCN from non-mucinous PCL [78].

Table 3. Typical results of cyst fluid analysis of PCL (modified from [33]; a data from [69], b data from [79–81], c periodic acid–Schiff stains for detection of glycogen and mucin)

<table>
<thead>
<tr>
<th>Pseudocyst</th>
<th>SPN</th>
<th>SCA</th>
<th>MCN</th>
<th>BD-IPMN</th>
<th>MD-IPMN</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Gross appearance</strong></td>
<td>non-viscous, muddy brown</td>
<td>non-viscous, old-bloody</td>
<td>non-viscous, water-clear, sometimes bloody</td>
<td>variably viscous, water-clear</td>
<td>variably viscous, water-clear</td>
</tr>
<tr>
<td><strong>Pancreatic enzymes</strong></td>
<td>high</td>
<td>no data</td>
<td>low</td>
<td>low</td>
<td>variably high</td>
</tr>
<tr>
<td><strong>CEA</strong></td>
<td>low, &lt;5 ng/ml</td>
<td>no data</td>
<td>low, &lt;5 ng/ml</td>
<td>high</td>
<td>high, depending on histological subtype (high in gastric and pancreatobiliary subtype)*, no marker of malignancy!</td>
</tr>
<tr>
<td><strong>DNA</strong></td>
<td>KRAS mutation absent</td>
<td>KRAS mutation highly specific</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Epithelium</strong></td>
<td>no, amorphic yellow material</td>
<td>yes, branching papillae with myxoid stroma</td>
<td>only in 20–25%, glycogen-rich, cuboid, non-mucinous</td>
<td>yes, mucin-containing (PAS-positive)c columnar cells, variable atypia</td>
<td></td>
</tr>
<tr>
<td><strong>Mucin phenotype</strong></td>
<td>MUC5AC+; gastric differentiation: MUC6+, MUC5AC+; intestinal differentiation: CDX2+, MUC2+, MUC5AC+; pancreatobiliary differentiation: MUC1+, MUC5AC+; oncocytic differentiation: MUC6+, MUC5AC+</td>
<td>MUC1 expression in MCN and IPMN is a marker of invasive growth,</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Blood cells</strong></td>
<td>histiocytes, leukocytes, erythrocytes</td>
<td>erythrocytes</td>
<td>hemosiderin-filled macrophages</td>
<td>rarely</td>
<td>rarely</td>
</tr>
</tbody>
</table>

Fig. 5. EUS images of a malignant BD-IPMN. a Large cystic lesions with solid parts and thick septae. b Contrast-enhanced EUS shows vascularization of solid parts and septae.
Reliability of Preoperative Diagnosis

Despite the diversity of high-resolution imaging tools and sophisticated cyst fluid markers, the diagnosis and risk assessment of PCL remains difficult. Even in tertiary referral centers with unquestionable experience, up to one-third of preoperative diagnoses of PCL proved to be incorrect [83, 84]. In one retrospective analysis of 136 patients with incidentally detected PCL which were operated on at a high-volume center due to preoperative diagnosis of mucinous PCL, 5% of resected cysts turned out not to be neoplastic. Even more worrying: when preoperative diagnosis was BD-IPMN or MCN, diagnosis failed in 40%. As many as 20% of presumed BD-IPMNs turned out to have main-duct involvement (‘mixed type’) and, therefore, carry a much higher risk of malignancy [83]. Accordingly, data from a German high-volume center showed histological main-duct extension in 67 out of 233 suspected BD-IPMNs (29%), which was not evident in preoperative imaging [85].

Interobserver agreement is disappointing in assessing morphological features, establishing a diagnosis, and estimating the risk of malignancy of PCL for MRI [86, 87] as well as for EUS [88, 89]. There is also substantial interobserver variability for the grading of cellular atypia in pancreatic cyst fluid [90, 91].

Risk Assessment

Several clinical, morphological, biochemical, and cytological criteria defining a high risk of malignancy of PCL have been evaluated (table 4). Combinations of different predictors in several studies have been shown to increase the accuracy of predicting malignancy [70, 77, 92]. However, the relative weight of predictive factors differs. The results of two meta-analyses of imaging features predicting the risk of malignancy of IPMN were not congruent [93, 94]. One meta-analysis included data from 41 studies on cyst features of both types of IPMN (MD and BD type). A cyst size > 30 mm was found to be most predictive of malignancy (odds ratio (OR) 62.4), followed by the presence of mural nodules (OR 7.3) and MD versus BD type (OR 4.7) [93]. The second meta-analysis focused on BD-IPMN and included 23 studies. Presence of mural nodules was the most important predictor of malignancy of BD-IPMN (OR 6.0), followed by dilatation of the main pancreatic duct (OR 3.4), thick septum/wall (OR 3.3), and cyst size > 30 mm (OR 2.3) [94]. For MCN, the absence of mural nodules and a cyst size < 40 mm are associated with no malignancy [95–97].

An international consensus guideline from 2006 (updated in 2012) recommended criteria (‘Sendai criteria’) as well as a diagnostic algorithm for the management of mucinous neoplastic cysts of the pancreas. Clinical and morphological features were defined to be ‘worrisome features’ or ‘high-risk stigmata’ (table 5) [115, 116].

<table>
<thead>
<tr>
<th>Table 4. Predictors of malignancy of mucinous neoplastic cysts (data from [60, 68, 73–75, 80, 82, 92–117])</th>
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</thead>
<tbody>
<tr>
<td><strong>Predictors of malignancy</strong></td>
</tr>
<tr>
<td>Epidemiological data</td>
</tr>
<tr>
<td>Clinical data</td>
</tr>
<tr>
<td>Laboratory findings</td>
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<tr>
<td>Morphological features</td>
</tr>
<tr>
<td>Cyst fluid markers</td>
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<td>Cytology</td>
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<table>
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<tr>
<th>Table 5. Sendai criteria of the international consensus guidelines 2012 for the management of IPMN and MCN of the pancreas: worrisome features and high-risk stigmata [116]</th>
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</thead>
<tbody>
<tr>
<td><strong>Worrisome features</strong></td>
</tr>
<tr>
<td>Clinical</td>
</tr>
<tr>
<td>Imaging</td>
</tr>
<tr>
<td><strong>High-risk stigmata</strong></td>
</tr>
<tr>
<td>Clinical</td>
</tr>
<tr>
<td>Imaging</td>
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<tr>
<td>EUS</td>
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<tr>
<td>EUS-FNA</td>
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</table>

The guideline recommends surgical treatment for all surgically fit patients with MD-IPMN and MCN. For patients with BD-IPMN, surgical treatment should be performed in the case of high-risk stigmata and should also be considered in patients without high-risk stigmata and a cyst size > 30 mm. Surveillance (cross-sectional imaging in lesions < 20 mm, EUS or MRI in lesions ≥ 20 mm) is proposed for BD-IPMN without high-risk stigmata, with the time interval depending on the size of the lesion and its growth [116]. Several cohort studies have been performed and initiated to evaluate the safety of these recommendations. Based on the guideline, surgery is indicated in less than 20% of BD-IPMNs [48]. Most
Fig. 6. Clinical algorithm for the diagnosis and treatment of incidentally detected PCL (modified from [34]).

published studies identified these recommendations to be reasonable and safe or suggested a more liberal management in the case of BD-IPMNs with small mural nodules [17, 60, 101, 109, 118–120]. One study demonstrated that absence of worrisome features, high-risk stigmata, and high-grade atypia or malignancy in EUS-FNA provided a predictive value of 99% for a safe non-surgical management [121]. Moreover, a management strategy based on a risk stratification using EUS-FNA and cyst fluid analysis proved to be most cost-effective in comparison with a conservative follow-up strategy or an aggressive surgical approach [122]. EUS proved to be the most effective method for the surveillance of BD-IPMNs and allows for an early detection of the majority of IPMN-derived or concomitantly developing pancreatic adenocarcinomas [22]. Nonetheless, weighting of the international consensus guideline criteria for the prediction of malignancy does not seem to be adequate. There is no stepwise increase in the rate of malignant or invasive IPMNs with the number of worrisome features [112]. Further research will be necessary to improve the risk stratification of the international consensus guidelines.

Conclusion and Proposal of a Diagnostic Algorithm

Despite the improvements of cross-sectional imaging and EUS-FNA, differential diagnosis, risk stratification, and clinical management of PCL remain challenging. Morphological criteria and cyst fluid analysis are not sufficient in some cases to establish a definitive diagnosis and to differentiate between benign and (pre-)malignant PCL. Interobserver variability of imaging findings and cytology is considerably high. Therefore, the standardization of criteria for diagnosis and risk stratification of incidental PCL, training of examiners, and multidisciplinary management decisions is necessary. Diagnostic management should follow a multi-step algorithm. Diagnosis results from a synthesis of the patients’ history, clinical data, as well as findings of TUS, cross-sectional imaging, EUS, and EUS-FNA. EUS plays a pivotal role in the diagnosis and surveillance of patients with incidentally detected PCL (fig. 6).

Disclosure Statement

The authors have no conflict of interest do declare.

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