Abstract
Significant advances in imaging technology have changed the management of pancreatic cancer. In computed tomography (CT), this has included development of multidetector row, rapid, thin-section imaging that has also facilitated the advent of advanced reconstructions, which in turn has offered new perspectives from which to evaluate this disease. In magnetic resonance imaging, advances including higher field strengths, thin-section volumetric acquisitions, diffusion weighted imaging, and liver specific contrast agents have also resulted in new tools for diagnosis and staging. Endoscopic ultrasound has resulted in the ability to provide high-resolution imaging rivaling intraoperative ultrasound, along with the ability to biopsy via real time imaging suspected pancreatic lesions. Positron emission tomography with CT, while still evolving in its role, provides whole body staging as well as the unique imaging characteristic of metabolic activity to aid disease management. This article will review these modalities in the diagnosis and staging of pancreatic cancer.

INTRODUCTION
Pancreatic cancer remains one of the most deadly cancers, ranked as the fourth most common cause of cancer related death in the United States per the American Cancer Society for 2009[1]. One of the reasons is because the pancreas is located deep in the abdomen, in close proximity to vital vascular structures that are early involved in the course of this disease. The median time between onset of usually nonspecific symptoms and presentation is 6 mo[2]. Only a small fraction of patients at presentation have resectable disease because of either already present metastases or locally advanced disease involving neighboring vasculature, and associated perineural invasion, to such an extent that surgical excision is no longer possible[3-4]. For all of these reasons, five year survival is only 5%[5].

Imaging plays a central role in the management of this disease. Imaging facilitates establishing a diagnosis, determining staging information, monitoring treatment response, and detecting tumor recurrence following surgery. Multiple modalities are involved, including computed tomography (CT), magnetic resonance imaging (MRI), positron emission tomography with computed tomography (PET/CT), and endoscopic ultrasound (EUS). These modalities have all evolved over the past
many years and so have their roles with regard to pancreatic cancer.

This article will review the use of multiple imaging modalities in many of these facets, and will show illustrative examples from a broad range of modalities. It will attempt to emphasize the unique contributions of each modality and where their strengths can best be used.

**DIAGNOSIS**

Usually patients with pancreatic cancer present because of symptoms of abdominal pain, accompanied by jaundice and/or unexplained weight loss. For this reason, patients will often undergo transabdominal ultrasound as their initial examination in order to evaluate for possible cholecystitis and/or obstructing calculi within the cystic duct, common hepatic duct or common bile duct. In our experience, while ultrasound is excellent for evaluation of gallbladder pathology, it is still limited for evaluation of the pancreas despite multiple recent advances. This is likely because of limitations of obscuring bowel gas, and dependence on the experience of the sonographer, among other factors. In pancreatic cancer, if the common bile duct is obstructed, ultrasound will likely demonstrate that ductal obstruction is within the region of the pancreatic head. In many cases in the past, patients have then undergone endoscopic retrograde cholangiopancreatography (ERCP) to diagnose and treat biliary obstruction. Unfortunately, post-ERCP changes, as well as imaging artifacts secondary to biliary stents, significantly degrade the imaging results of subsequent cross-sectional imaging, whether by CT, MRI or EUS. As will be discussed subsequently, cross-sectional imaging has assumed a central role in the diagnosis and management of pancreatic cancer. For this reason, an important change at our institution has been to do cross-sectional imaging, typically by CT or MRI, before any intervention (ERCP, stent placement, biopsy, etc.). At our institution, EUS can be done at the same setting as ERCP, and is done first to avoid the problems of imaging after stent placement.

At many institutions, multidetector computed tomography (MDCT) has become the primary modality for evaluating suspected pancreatic cancer. This technology has been rapidly evolving with the development of ever faster and thinner section imaging with the evolution of MDCT. As advocated by the National Comprehensive Cancer Network guidelines, this is typically through multiphase imaging technique, with pre-contrast imaging and imaging of the abdomen obtained during the peak pancreatic parenchymal phase and peak liver portal venous enhancement phase. The earlier phase best depicts the primary tumor (Figure 1) while the latter phase best demonstrates tumor involvement of venous structures and liver metastases. Use of saline flush in addition to power injection may additionally improve enhancement of the pancreatic parenchyma and adjacent vasculature. One of the newest developments, dual energy or multispectral CT, may be useful in the detection of subtle differences in enhancement with iodinated contrast but currently no studies have been published regarding its utility in pancreatic cancer.

Thin section imaging is important for not only showing the primary tumor but also for showing its relationship to vasculature not only in the original axial plane of acquisition but also in detailed multiplanar reconstructions (Figure 2). At our institution, dual phase studies are reconstructed into two data sets. Thicker 2.5 mm are used for primary interpretation, and thinner 0.625-1.25 mm axial sections are used for problem solving and for multiplanar reconstructions.

Pancreatic cancer usually manifests on CT as an ill-defined, solid mass, slightly hypodense to normal pancreatic parenchyma. Overall sensitivity of MDCT for pancreatic cancer is 86%-97% for tumors of any size, but sensitivity is probably near 77% for small (< 2 cm) lesions. It is important to pay close attention to secondary signs when the primary tumor is difficult to visualize, which is particularly the case for small lesions. These signs include focal pancreatic enlargement, extension of tumor beyond the pancreas, “upstream” pancreatic atrophy (secondary to ductal obstruction) and, most importantly, dilatation and/or cut off of the pancreatic...
duct and/or common bile duct. Gangi et al[17] reported in a study of 28 patients with pancreatic cancer who had CT imaging prior to histologic diagnosis that findings suspicious for pancreatic cancer were present in 50% of cases obtained 2-6 mo and 6-18 mo prior to the diagnosis of pancreatic cancer.

Such signs are also of particular utility in the 5.4% of pancreatic cancers that are truly isoattenuating on both phases of a dual phase pancreatic protocol[18]. In such circumstances, other modalities such as EUS can be considered. A study of isoattenuating tumors showed that MRI and PET/CT had sensitivities of 79.2% and 73.7% in such cases and can also be considered for followup when CT is equivocal[18].

Another recent development has been the use of a variety of types of reformations to enhance the conspicuity of tumor and its relationship to local structures. Prokesch et al[19] showed that use of curved multiplanar reconstructions drawn along the common bile duct, pancreatic duct, and/or mesenteric vessels may help improve sensitivity for tumor detection and the speed of interpretation over axial images alone. Salles et al[20] noted that minimum intensity projection images, besides showing ductal structures, may help improve the conspicuity of the primary tumor within the pancreas (Figure 3).

There have been multiple recent developments in MRI including parallel imaging, imaging at field strengths greater than 1.5 T, liver specific contrast agents (for detection of liver metastases), and broader implementation of diffusion weighted imaging. At our institution, a typical imaging protocol includes multiple axial acquisitions including fat suppressed T2 weighted imaging of the abdomen, thin-section T2 weighted imaging of the pancreatic region, in-and-out of phase T1 weighted imaging, fat suppressed dynamically obtained 3-D T1 weighted imaging (20 s, 60 s, 120 s, and 180 s after the start of contrast injection), fat suppressed thin-section fast imaging employing steady-state acquisition (GE Medical Systems, Milwaukee, WI), and axial diffusion weighted imaging.

Pancreatic cancer typically has the appearance on MRI of an ill defined mass, as on CT. MRI offers the advantage of multiple sequences with inherently greater soft tissue contrast without enhancement than unenhanced CT. Tumor typically appears hypointense on fat suppressed T1, and pancreatic parenchymal phase dynamically enhanced fat suppressed T1-weighted sequences. Pancreatic cancer has a variable appearance on T2 weighted images. Pancreatic cancer also has a variable appearance on diffusion weighted images (Figure 4). In a recent study of 80 patients, 38 appeared hyperintense, 12 isointense, and four hypointense[21]. Early reports from the literature are beginning to investigate the utility of diffusion weighted imaging for diagnosing pancreatic cancer, and for differentiating chronic pancreatitis from tumor[22-25].

At most institutions, PET/CT, as currently performed without intravenous contrast, has a limited role in the diagnosis and staging of pancreatic cancer (Figure 5). A recent study of 103 patients with pancreatic cancer showed a similar detection rate between contrast-enhanced MDCT and PET/CT (89% vs 91%)[26].

EUS has developed a strong role in the detection and confirmation of pancreatic cancer. EUS has the advan-

Figure 2 Patient with locally advanced pancreatic cancer. A-C: Pancreatic parenchymal phase images show tumor (arrowheads) (A) encasing the common hepatic artery (CHA) and splenic artery (SA) (B) encasing the celiac axis (CA) and (C) abutting the superior mesenteric artery (SMA); D: Coronal reformation helps demonstrate the encasement of the celiac artery and abutment of the superior mesenteric artery. R: Right; L: Left.

Figure 3 Utility of reconstructions. A: Coronal reconstruction demonstrating craniocaudal extent of tumor (arrows); B: Coronal Minip (minimum intensity projection), same obliquity and orientation as A emphasizes low density structures, making tumor more conspicuous.
tage that tumor can be visualized in real time without the requirement of a contrast agent (Figure 6). A study from our institution showed EUS-fine needle aspiration (FNA) to have higher sensitivity (99%) than MDCT (89%-93%)\cite{16}. The technique is of particular utility for small lesions < 2 cm in size, with the sensitivity of EUS being 96%, and 70%-93% for MDCT\cite{16}. However, the technique is invasive, and is operator dependent. The presence of a biliary stent can greatly lower the ability of EUS-FNA to exclude tumor\cite{16}. For this reason at our institution, cross-sectional imaging (MDCT or MRI), is performed first, followed by EUS-FNA. As noted previously, at our institution ERCP and EUS are done at the same setting, with EUS done first.

**STAGING**

In 2002, The American Joint Committee on Cancer Staging, changed the T staging criteria to focus on tumor involvement of arteries and removed reference to venous involvement\cite{5}. This is likely secondary to advances made in the use of venous bypass grafts that allowed for

---

**Figure 4** Magnetic resonance imaging of patient with two sites of pancreatic cancer. A: T1 fat suppressed images do not show the tumor well in this patient, but are often helpful. Solid arrow indicates the pancreatic neck tumor, dashed arrow indicates the pancreatic tail tumor; B, C: However, both sites are well seen, in the neck (solid arrows) and tail (dashed arrows) on the pancreatic parenchymal phase (B) and the portal venous phase (C); D: In this case sites of tumor are not well seen on diffusion weighted imaging. Solid arrow indicates the pancreatic neck tumor, dashed arrow indicates the pancreatic tail tumor. Conspicuity of the primary lesion can be very variable on diffusion weighted imaging.

**Figure 5** Coronal fused positron emission tomography with computed tomography images of patient with pancreatic cancer and liver metastases. A: Primary tumor shows mild-moderate uptake (dashed arrow); B, C: While liver metastases in (B) and (C) show variable uptake (solid arrows).
successful en bloc resection of tumor involving veins\(^{[27]}\). Additionally, perineural invasion, a histopathologic poor prognostic factor, parallels the course of arteries, likely increasing the importance of arterial involvement\(^{[29]}\). As a result, the challenge of imaging has become to better identify the extent of tumor involvement of vasculature, particularly the celiac axis, the superior mesenteric artery and the common hepatic artery.

**Vascular involvement**

The determination of the extent of vascular involvement is typically made by identifying the extent that tumor involves the cross-sectional circumference of a vessel. This can be done by describing with regard to the circular cross-section of a vessel the degrees of circumferential involvement as described by Lu et al\(^{[30]}\). Since that time, the terms “abutment” and “encasement,” have also been utilized, abutment referring to 180° of involvement or less of a vessel’s circumference and encasement referring to greater than 180° of vascular circumferential involvement (Figure 2)\(^{[29,30]}\). These terms (degrees of circumferential involvement, abutment/encasement, as well as contiguous/discontiguous involvement) are also utilized in the Radiological Society of North America’s suggested reporting template for primary pancreatic masses\(^{[31]}\). The utility of these terms includes the ability to differentiate clearly resectable tumor, from “borderline” resectable tumor, from clearly unresectable tumor\(^{[30]}\). The assessment of vascular involvement is typically made at most institutions on either CT or MRI. A prospective blinded study comparing EUS, MRI and CT showed accuracies of 62%, 60% and 74% respectively for locoregional extent of tumor\(^{[30]}\). A recent study of 116 patients comparing MRI with MR angiography against MDCT showed similar sensitivities between the modalities (approximately 75%-87%)\(^{[31]}\). Unfortunately, all of the modalities are limited in regard to the evaluation of nodal disease. For cross-sectional modalities, such as CT and MRI, the criterion for identifying adenopathy has typically been size. Unfortunately, as shown by Valls et al\(^{[41]}\) in their study of patients with pancreatic cancer undergoing preoperative assessment with CT, size is a nonspecific criterion. They showed that only 3 of 18 patients (16.7%) who had pathologically proved nodal involvement were detected when a size criterion of > 1.5 cm was utilized. Other criteria, such as poorly defined nodal boundaries, conversion of nodes to a rounded shape, and conversion to a hypodense appearance are more specific, but are less sensitive. MRI, when using fat suppressed T2 sequences, can be helpful in differentiating nodes from adjacent liver in the region of the porta hepatitis\(^{[42]}\). PET/CT has been reported to have sensitivities of 46%-71% and specificities of 63%-100% for nodal involvement\(^{[43,45]}\). EUS-FNA offers the benefit of histopathologic proof, but the number of sites that can be evaluated in this manner is limited.

**Distant metastases**

Pancreatic cancer typically metastasizes to the liver, peritoneum, and lungs, with metastases to osseous structures being less common, and in our experience, typically later in the course of the disease.

In most institutions, the assessment for liver metastases is usually made by CT or MRI. The sensitivity of intravenous contrast enhanced CT has been reported to be approximately 75%-87%\(^{[46-48]}\). Conventional gadolinium enhanced dynamic MRI has been considered to be similar or slightly better than CT. In a study directly comparing MRI and CT in 58 patients, the accuracy of MRI was 93.5% vs 87.2% for CT\(^{[47]}\). It is notable that liver metastases can have a variable appearance on MRI. Danet et al\(^{[49]}\) studied a small group of 16 patients with liver metastases with biopsy confirmed pancreatic adenocarcinoma and/or biopsy proven liver metastases, with dynamic gadolinium enhanced imaging and T2 weighted imaging. Of these patients, 75% of the patients had lesions that were 1.5 cm in size, 62% had hypovascular lesions, 38% had reportedly hypervascular lesions, and 62% showed perilesional enhancement, with 38% having this form of enhancement in a ring pattern. Newer liver specific agents such as gadobenate dimeglumine (Gd-BOPTA) can be used for dynamic imaging, allowing for arterial and portal venous phase imaging, and can also be used for delayed phase imaging because of contrast retention by hepatocytes which may
improve sensitivity for liver metastases. While, to our knowledge, specific assessment for pancreatic cancer liver metastases has not been examined, the overall mean accuracy for liver metastases using Gd-BOPTA in one study was approximately 95.5% compared to superparamagnetic iron oxide-enhanced imaging with a mean accuracy of 97.2%\(^{[90]}\). Diffusion weighted imaging has the advantage that no intravenous contrast needs to be administered (Figure 7). In a study of 31 patients, MDCT was compared against diffusion weighted imaging using \(b\) values of 0, 300 and 600 s/mm\(^2\). The sensitivities and specificities for liver metastases that had been correlated with intraoperative surgical and ultrasound findings were 53.3% and 77.8% for MDCT, and 86.7% and 97.5% for diffusion weighted imaging, which was noted to be statistically significant\(^{[10]}\).

All modalities have limited effectiveness in detecting the typically very small peritoneal implants of pancreatic cancer. Laparoscopy has been utilized to improve detection of peritoneal implants. This technique is invasive, and therefore can’t be used to monitor response to therapy. A meta-analysis in 2001 showed that laparoscopy may only alter management in 4%-15% of patients after optimized, thin section, CT\(^{[59]}\). Some studies suggest that PET/CT may offer benefits over contrast enhanced CT for the detection of distant metastases (Figure 5) as it provides a whole body scan, and when combined with IV contrast, may provide a comprehensive assessment for diagnosis and staging\(^{[83-85]}\).

**CONCLUSION**

Many advances have been made in multiple modalities in the past several years that have an impact on the diagnosis and staging of pancreatic cancer. MDCT has evolved to permit sub-millimeter imaging that allow for high quality reformation techniques that facilitate diagnosis and vascular staging. EUS has become a valuable tool for the detection and histopathologic confirmation of pancreatic tumors. MRI, with the development of thin section 3D gradient echo dynamic imaging, diffusion weighted imaging, higher field strengths, and liver specific agents combined have improved not only the identification of the primary tumor but also the evaluation for distant metastases. The role of PET/CT is continuing to evolve, and could potentially serve as comprehensive approach in the case of IV contrast enhanced PET/CT. PET/CT also offers advantages in identifying distant metastases from being a whole body assessment.

**REFERENCES**

1. American Cancer Society. Cancer Statistics Presentation 2009. Available at: URL: http://www.cancer.org/acs/groups/content/@nhs/documents/document/cancerstatis tic2009slidesrevppt.ppt
2. Conlon KC, Areu MA. Pancreas Cancer: Anatomy, Staging, Systems, and Techniques. In: Kelsen DP, Daly JM, Kern SE, Levin B, Tepper JE, Van Cutsem E, editors. Principles and practice of gastrointestinal oncology. 2nd ed. Philadelphia: Lippincott Williams and Wilkins, 2008: 1008
3. Jemal A, Siegel R, Ward E, Murray T, Xu J, Thun MJ. Cancer statistics, 2007. CA Cancer J Clin 2007; 57: 43-66 [PMID: 17237035]
4. Collins JM, Silva AC, Hayman LA. Arterial anatomy of the pancreas. Part 3: segmented computed tomography-angiography mapping of perineural invasion. J Comput Assist Tomogr 2010; 34: 961-965 [PMID: 21084917 DOI: 10.1097/RCT.0b013e3181d5fb6c]
5. Ries LAG, Melbert D, Krupcho M, Mariotto A, Miller BA, Feuer EJ, Clegg L, Horner MJ, Howlader N, Eisner MP, Reichman M, Edwards BK, editors. SEER Cancer Statistics Review, 1975-2004. Available from: URL: http://seer.cancer.gov/csr/1975_2004/
6. Modell I, Guarner L, Malagelada JR. Vagaries of clinical presentation of pancreatic and biliary tract cancer. Ann Oncol 1999; 10 Suppl 4: 82-84 [PMID: 10436792]
7. NCCN Guidelines Version 2. 2012 Panel Members Pancreatic Adenocarcinoma. 2011. Available from: URL: http://www.onko.szote.u-szeged.hu/intranet/attachments/article/87/pancreatic.pdf
8. Kim T, Murakami T, Takahashi S, Okada A, Hori M, Narumi Y, Nakamura H. Pancreatic CT imaging: effects of different injection rates and doses of contrast material. Radiology 1999; 212: 219-225 [PMID: 10405745]
9. Schueller G, Schima W, Schueller-Weidekamm C, Weber M, Stift A, Grannt M, Prokesch R. Multidetector CT of pancreas: effects of contrast material flow rate and individualized scan
delay on enhancement of pancreas and tumor contrast. Radiology 2006; 241: 441-448 [PMID: 16962815]

Tublin ME, Tessler FN, Cheng SL, Peters TL, McGovern PC. Effect of injection rate of contrast medium on pancreatic and hepatic helical CT. Radiology 1999; 210: 97-101 [PMID: 9885593]

Schoellnast H, Tillich M, Deutschmann HA, Stessel U, Deutschmann MJ, Schaffler GJ, Schoellnast R, Uggowitzer MM. Improvement of parenchymal and vascular enhancement using saline flush and power injection for multi-detector-row abdominal CT. Eur Radiol 2004; 14: 659-664 [PMID: 15466425 DOI: 10.1007/s00330-003-2085-3]

A garwal B, Abu-Hamdà M, Molke KJ, Correa AM, Ho L. Endoscopic ultrasound-guided fine needle aspiration and multidetector spiral CT in the diagnosis of pancreatic cancer. Am J Gastroenterol 2004; 99: 844-850 [PMID: 15128348]

Bronstein YL, Loyer EM, Kaur H, Choi H, David C, Dubrow RA. Comparison of 2002: CT results with 110 pancreatic tumors with multiphasic helical CT. AJR Am J Roentgenol 2004; 182: 619-625 [PMID: 14975999]

Dawe P, Jowell P, Leblanc J, McHenry L, McGreavy K, Cramer H, Volmar K, Sherman S, Gress F. EUS-guided FNA of pancreatic metastases: a multicenter experience. Gastrointest Endosc 2005; 61: 689-696 [PMID: 15859793]

Fletcher JG, Wiersma MJ, Farrell MA, Bidler JL, Burgart LJ, Koyama T, Johnson CD, Stephens DH, Ward EM, Harmsen WS. Pancreatic malignancy: value of arterial, pancreatic, and hepatic phase imaging with multi-detector row CT. Radiology 2003; 229: 81-90 [PMID: 14519871]

Tamm EP, Loyer EM, Faris CA, Evans DB, Wolff RA, CharnsangAVE. Retrospective analysis of dual-phase MDCT and follow-up EUS/EUS-FNA in the diagnosis of pancreatic cancer. Abdom Imaging 2007; 32: 660-667 [PMID: 17712589]

Gangi S, Fletcher JG, Nathan MA, Christensen JA, Harmsen WS, Crownhart BS, Chari ST. Time interval between abnormalities seen on CT and the clinical diagnosis of pancreatic cancer: retrospective review of CT scans obtained before diagnosis. AJR Am J Roentgenol 2004; 182: 897-903 [PMID: 15039161]

Kim JH, Park SH, Yu ES, Kim MH, Kim J, Byun JH, Lee SS, Hwang HJ, Hwang JY, Lee SS, Lee MG. Visually isoattenuating pancreatic adenocarcinoma at dynamic-enhanced CT: frequency, clinical and pathologic characteristics, and diagnosis at imaging examinations. Radiology 2010; 257: 87-96 [PMID: 20697118]

Prokesch RW, Chow LC, Beaulieu CF, Nino-Murcia M, Mindelez RE, Bamber R, Huang J, Jeffrey RB. Local staging of pancreatic carcinoma with multi-detector row CT: use of curved planar reformations. AJR Am J Roentgenol 2004; 182: 732-740 [PMID: 12642128]

Salles A, Nino-Murcia M, Jeffrey RB. CT of pancreas: minimum intensity projections. Abdom Imaging 2008; 33: 207-213 [PMID: 17387537]

Fukukura Y, Takumi K, Kamimura K, Shindo T, Kumatage Y, Tateyama A, Nakajo M. Pancreatic adenocarcinoma: variability of diffusion-weighted MR imaging findings. Radiology 2012; 263: 732-740 [PMID: 22623694]

Fattahi R, Balci NC, Perman WH, Hsueh EC, Alkaade S, Havilgolou N, Burton FR. Pancreatic diffusion-weighted imaging (DWI): comparison between mass-forming focal pancreatic adenocystic (FP), pancreatic cancer (PC), and normal pancreas. J Magn Reson Imaging 2009; 29: 350-356 [PMID: 19161187 DOI: 10.1002/jmri.21651]

Kartalis L, Lindholm TL, Aspelin P, Perment J, Albinni N. Diffusion-weighted magnetic resonance imaging of pancreatic tumours. Eur Radiol 2009; 19: 1981-1990 [PMID: 19308414 DOI: 10.1007/s00330-009-1384-8]

Lee SS, Byun JH, Park BJ, Park SH, Kim N, Park B, Kim JK, Lee MG. Quantitative analysis of diffusion-weighted magnetic resonance imaging of the pancreas: usefulness in characterizing solid pancreatic masses. J Magn Reson Imaging 2008; 28: 928-936 [PMID: 18821618]

Lemke A, Laun FB, Klauss M, Re TJ, Simon D, Delorme S, Schad LR, Stieljes B. Differentiation of pancreatic carcinoma from healthy pancreatic tissue using multiple b-values: comparison of apparent diffusion coefficient and intravoxel incoherent motion derived parameters. Invest Radiol 2009; 44: 769-775 [PMID: 19838121 DOI: 10.1097/RLI.0b013e3181b62271]

Iizuki K, Yamamoto Y, Sano T, Takebayashi R, Masaki T, Suzuki Y. Impact of 18-fluorodeoxyglucose positron emission tomography on the management of pancreatic cancer. J Gastroenterol Surg 2010; 14: 1151-1158 [PMID: 20443874 DOI: 10.1007/s11605-010-1207-x]

Fuhrman GM, Leach SD, Staley CA, Cusack JC, Charnsangavej C, Cleary KR, El-Naggar AK, Fenoglio CJ, Lee JE, Evans DB. Rationale for en bloc vein resection in the treatment of pancreatic adenocarcinoma adherent to the superior mesenteric-portal vein confluence. Pancreatic Tumor Study Group. Ann Surg 1996; 223: 154-162 [PMID: 859709]

Lu DS, Reber HA, Krasny RM, Kadell BM, Sayre JL. Local staging of pancreatic cancer: criteria for unresectability of major vessels as revealed by pancreatic-Phase, thin-section helical CT. AJR Am J Roentgenol 1997; 168: 1439-1443 [PMID: 9168704]

Evans DB, Erickson BA, Ritch P. Borderline resectable pancreatic cancer: definitions and the importance of multimodality therapy. Ann Surg Oncol 2010; 17: 2803-2805 [PMID: 20735218 DOI: 10.1245/s10434-012-1265-9]

Varadhachary GR, Tamm EP, Abbruzzese JL, Xiong HQ, Crane CH, Wang H, Lee JE, Pisters PW, Evans DB, Wolff RA. Borderline resectable pancreatic cancer: definitions, management, and role of preoperative therapy. Ann Surg Oncol 2006; 13: 1035-1046 [PMID: 16865597]

CT Onco Primary Pancreas Mass. Available from: URL: http://reportingwiki.rsna.org/index.php?title=Templates

Soriano A, Castells A, Ayuso C, Ayuso JR, de Caralt MT, Gines MA, Real MI, Gilabert R, Quinto L, Trilla A, Feliu F, Montanyà X, Fernández-Cruz L, Navarro S. Preoperative staging and tumor resectability assessment of pancreatic cancer: prospective study comparing endoscopic ultrasonography, helical computed tomography, magnetic resonance imaging, and angiography. Am J Gastroenterol 2004; 99: 492-501 [PMID: 15056091]

Lee JK, Kim AY, Kim PN, Lee MG, Ha HK. Prediction of vascular involvement and resectability by multidetector-row CT versus MR imaging with MR angiography in patients who underwent surgery for resection of pancreatic ductal adenocarcinoma. Eur J Radiol 2010; 73: 310-316 [PMID: 19070981 DOI: 10.1016/j.ejrad.2008.10.028]

Horton KM, Fishman EK. Volume-rendered 3D CT of the mesenteric vasculature: normal anatomy, anatomic variants, and pathologic conditions. Radiographics 2002; 22: 161-172 [PMID: 11796095]

Horton KM. Multidetector CT and three-dimensional imaging of the pancreas: state of the art. J Gastrointest Surg 2002; 6: 126-128 [PMID: 11992794]

Horton KM, Fishman EK. Multidetector CT angiography of pancreatic carcinoma: part 2, evaluation of venous involvement. AJR Am J Roentgenol 2002; 178: 833-836 [PMID: 11906858]

Horton KM, Fishman EK. Multidetector CT angiography of pancreatic carcinoma: part I, evaluation of arterial involvement. AJR Am J Roentgenol 2002; 178: 827-831 [PMID: 11906856]

Riediger H, Keck T, Wellner U, zur Hausen A, Adam U, Hopt UT, Makowiec F. The lymph node ratio is the strongest prognostic factor after resection of pancreatic cancer. J Gastrointest Surg 2009; 13: 1337-1344 [PMID: 19418101 DOI: 10.1007/s11605-009-0919-2]
magnetic resonance imaging improves the staging of pancreatic tumors. *Ann Surg* 1997; 226: 393-405; discussion 404-407 [PMID: 9351708]

48 **Calcutti L**, Casadei R, Diacono D, Caputo M, Cavina M, Minguzzi MT, Marrano D, Gavelli G. [Role of spiral computed tomography in the staging of pancreatic carcinoma] *Radiol Med* 1998; 95: 344-348 [PMID: 9676213]

49 **Danel IM**, Semelka RC, Nagase LL, Woosely JT, Leonidou P, Armao D. Liver metastases from pancreatic adenocarcinoma: MR imaging characteristics. *J Magn Reson Imaging* 2003; 18: 181-188 [PMID: 12884330 DOI: 10.1002/jmri.10337]

50 **Kim YK**, Lee JM, Kim CS, Chung CH, Kim CY, Kim IH. Detection of liver metastases: gadobenate dimeglumine-enhanced three-dimensional dynamic phases and one-hour delayed phase MR imaging versus superparamagnetic iron oxide-enhanced MR imaging. *Eur Radiol* 2005; 15: 220-228 [PMID: 15624108 DOI: 10.1007/s00330-004-2570-3]

51 **Holzapfel K**, Reiser-Erkan C, Fingerle AA, Erkan M, Eiber MJ, Rummey EJ, Fries H, Kleeff J, Gaa J. Comparison of diffusion-weighted MR imaging and multidetector-row CT in the detection of liver metastases in patients operated for pancreatic cancer. *Abdom Imaging* 2011; 36: 179-184 [PMID: 20563868 DOI: 10.1007/s00261-010-9633-5]

52 **Pisters PW**, Lee JE, Vauthey JN, Charnsangavej C, Evans DB. Laparoscopy in the staging of pancreatic cancer. *Br J Surg* 2001; 88: 325-337 [PMID: 11260096]

53 **Farma JM**, Santillan AA, Melis M, Walters J, Belinc D, Chen DT, Eikman EA, Malafa M. PET/CT fusion scan enhances CT staging in patients with pancreatic neoplasms. *Ann Surg Oncol* 2010; 18: 2465-2471 [PMID: 18551347]

54 **Nishiyama Y**, Yamamoto Y, Yokoe K, Mondon T, Sasakiya Y, Tsutsui K, Satoh K, Ohkawa M. Contribution of whole body FDG-PET to the detection of distant metastasis in pancreatic cancer. *Ann Nucl Med* 2005; 19: 491-497 [PMID: 16248386]

55 **Strobel K**, Heinrich S, Btoure U, Soyka J, Veit-Heibach P, Pestalozzi BC, Clavien PA, Hany TF. Contrast-enhanced 18F-FDG PET/CT: 1-stop-shop imaging for assessing the resectability of pancreatic cancer. *J Nucl Med* 2008; 49: 1408-1413 [PMID: 18703604 DOI: 10.2967/jnumed.108.051466]