Image response evaluation after novel neoadjuvant treatments of pancreatic cancer

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Summary

Background Emergent neoadjuvant concepts have improved the rate of margin-free (R0) resections in borderline resectable and locally advanced pancreatic ductal adenocarcinoma. However, due to specific properties of pancreatic cancer, imaging response assessment is difficult.

Methods The concepts of histologic response patterns and their imaging correlates are outlined, together with efforts in morphologic and functional imaging to overcome the challenges of response assessment.

Results Imaging evaluation after neoadjuvant therapy remains a diagnostic dilemma. To date, better understanding of the specific transformations of PDAC under neoadjuvant agents could not be transformed into validated imaging criteria for defining response and selecting surgical candidates.

Conclusion Currently, patients with non-metastatic borderline resectable and locally advanced pancreatic ductal adenocarcinoma are brought to diagnostic laparotomy in the absence of overt radiologic progression after neoadjuvant therapy. Apart from morphologic patterns, ongoing research is focused on defining robust functional imaging parameters for response assessment.

Keywords Computed Tomography · Magnetic Resonance Imaging · 18-Fluorodeoxyglucose PET · Pancreatic ductal adenocarcinoma · Neoadjuvant therapy · Resectability

Introduction

Several trials have investigated the effect of novel neoadjuvant regimens on the resectability of both borderline resectable (BR) and locally advanced (LA) pancreatic ductal adenocarcinoma (PDAC). However, specific histopathologic changes of PDAC under neoadjuvant treatment (NAT) preclude accurate depiction of viable tumor remnants with cross-sectional imaging. This overview summarizes current knowledge of PDAC response patterns on diagnostic imaging, recent approaches to estimate downstaging and optimize patient selection for surgery based on imaging.
Table 1  NCCN guidelines version 3. 2017 criteria defining borderline resectable (BR) and locally advanced (LA) pancreatic adenocarcinoma

| Borderline resectable (BR) pancreatic adenocarcinoma | Locally advanced (LA) pancreatic adenocarcinoma |
|------------------------------------------------------|--------------------------------------------------|
| **Arterial** | **Pancreatic head/uncinated process** | **Pancreatic body/tail** |
| Arterial tumor contact ≤180° | CHA: tumor contact of ≤180° without extension to CA or hepatic artery bifurcation allowing for resection and reconstruction | CA: contact of ≤180° or contact of >180° without involvement of the aorta and with intact GDA permitting a modified Appleby procedure |
| arterial SMA: tumor contact | Contact with variant artery | |
| arterial SMV: tumor contact | Tumor contact with the aorta | |
| arterial PV: tumor contact | Tumor contact with >180° | |
| arterial contact of >180° | Tumor contact with the first jejunal SMA branch | |
| arterial contact of ≤180° | Unreconstructible SMV/PV due to tumor involvement or occlusion due to tumor or bland thrombus | |
| arterial contact of ≤180° with contour irregularity or vein thrombosis, but with suitable vessel proximal and distal for reconstruction | Contact with most proximal draining jejunal branch into SMV | |
| arterial contact of >180° | Unreconstructible SMV/PV due to tumor involvement or occlusion due to tumor or bland thrombus | |
| arterial contact of >180° with the SMA or CA | Tumor contact with the CA and aorta | |
| venous Unreconstructible SMV/PV due to tumor involvement or occlusion due to tumor or bland thrombus | Tumor contact with the IVC | |
| venous SMA: tumor contact with >180° | Tumor contact with the CA and aorta | |
| venous PV: tumor contact with >180° | Tumor contact with the IVC | |
| venous contact of >180° | Tumor contact with the SMA or CA | |
| venous contact of ≤180° | Unreconstructible SMV/PV due to tumor involvement or occlusion due to tumor or bland thrombus | |
| Distant metastasis including non-regional lymph nodes | | |
| CA celiac axis, CHA common hepatic artery, SMA superior mesenteric artery, GDA Gastroduodenal artery, PV portal vein, SMV superior mesenteric vein, IVC inferior vena cava |

Non-metastatic pancreatic adenocarcinoma is categorized into three surgical stages at the time of diagnosis: resectable (R), borderline resectable (BR), and locally advanced (LA) [1]. Currently, several classification systems are used to define resectability with a required safety margin of ≥1 mm. The consensus definitions published by the NCNN/AHPBA/SSO/SSAT, as well as definitions of resectability from the MD Anderson Cancer Centre are most widely used ([2, 3]; Table 1 and Fig. 1). Since the recognition of borderline resectability in 2008 [4], neoadjuvant concepts have been implemented to downstage patients from either BR or LA-PDAC to a more favorable surgical stage in order to achieve curative surgery ([5]; Fig. 2).

Gemcitabine-based regimens, recently combined with nab-paclitaxel [6], and the advent of FOLFIRINOX-based schemes in 2011 [7], both with or without additional chemoradiation, hold promise for downstaging PDAC and more importantly, enhance the rate of R0 resection [8, 9].

Unfortunately, response evaluation on imaging and prediction of resectability have proven challenging due replacement of tumor by fibrosis [10, 11]. Although biomarker levels and clinical performance status are incorporated into reassessment [1], multi-phase contrast-enhanced multidetector CT (MDCT) with three-dimensional reconstructions remains the mainstay for further patient selection for surgery [12]. Meanwhile, advances in magnetic resonance imaging (MRI), notably faster acquisition techniques and enhanced image quality due to improved signal-to-noise and contrast-to-noise ratios, have brought MRI close to the spatial resolution of MDCT, while offering superior contrast resolution and the opportunities of functional imaging in terms of diffusion-weighted imaging (DWI).

Histopathologic response patterns of PDAC

It was recognized early that NAT-induced tumor cell injury in PDAC is mainly reflected by “isovolumetric” tissue replacement through fibrosis, rather than volume loss. In 1992, Evans at al. reported the histologic changes in 17 resected specimens after NAT and established a pathologic response grading sys-
main topic

Fig. 2  CT image showing a contact of ≤180° between tumor and the superior mesenteric artery (arrows), defined as borderline resectable disease according to the National Comprehensive Cancer Network (NCCN) version 2017 guidelines for PDAC [13] by describing the percentage of viable tumor cells within the specimen. In their review, Kalimuthu et al. outlined histopathologic changes after NAT in more detail [14], with emphasis on subtle residual tumor nests scattered throughout the fibrotic tissue.

A small percentage of a complete pathologic response (2% to 7%) was also reported in meta-analyses [15, 16]. Finally, patients with NAT are less likely to have positive lymph nodes and perineural invasion on histopathologic workup [17].

Imaging response assessment with RECIST1.1 (clinical stage)

The concept of tissue replacement, along with perivascular/perineural patterns of growth, makes PADC ineligible for follow-up with RECIST 1.1. It is thought to be the cause of the repeatedly documented failure to predict viable tumor nests around crucial surgical structures using MDCT or MRI.

In their meta-analysis, Dhir et al. calculated overall RECIST1.1 response rates after NAT in a subset of 61 studies [18]. Overall, partial response was seen in 20% of patients, most patients remained stable (59%), and progressive disease occurred in 16% of patients.

Pathologic response grading revealed an overall Evans grade III rate (tumor destruction of >90%) of 12%. Grade IIB (tumor destruction of 51–90%) was seen in 27%, and 37% had grade IIA (11–50% tumor destruction). Grade I (<10% tumor destruction) was found in 13% of specimens.

Thus, the majority of patients were stable or in partial response on cross-sectional imaging, and most of the resected specimens showed minor to moderate histologic response. However, as Xia [19] et al. demonstrated, radiologic and pathologic grading are not correlated on a patient basis (Fig. 3). In their work-up of 38 specimens after NAT, all pathologic response grades from minor to complete were present in both radiologic partial responders as well as in patients with stable disease. Minor pathologic response (<than 50% tumor destruction) was dominant in the group of RECIST stable disease. There were three pathologic complete responders in their group, two with RECIST stable disease, and one with partial response.

Imaging response assessment of surgical stage (resectability)

With the advent of novel NAT concepts, such as FOLFIRINOX or gemcitabine/nab-paclitaxel, several groups have investigated radiologic tumor changes after therapy in recent years, with special attention to resectability. Meta-analyses initially suggested high R0 rates in patients with BR-PDAC: when 63% of patients were selected for surgery after FOLFIRINOX, R0 margins resulted in 93% of them [20]. Patient selection becomes even more important in locally advanced PDAC: when 28% of patients were eligible for surgery after FOLFIRINOX, free margins were seen in 74% of operated patients [21]. On an intention-to-treat basis, however, overall R0 margins resulted in only 22% of treated LA-PDAC patients [22]. A single recent study, however, reported a far higher R0 rate of 62% [22].

Generally, imaging workup after NAT reveals an overestimation of residual tumor burden and the probability of R0 resection is difficult to estimate (Fig. 4). Diagnostic accuracy for predicting resectability after NAT has yielded conflicting results so far: accuracy was set at 58% by a French group [23], whereas Kim et al., in an older publication, reported a high accuracy of 83% [24].

The effect of FOLFIRINOX on imaging resectability criteria compared to pathologic margins was presented in two publications with similar results:

Ferrone and colleagues found a shift to more advantageous resection categories on CT in a collective of 40 patients after FOLFIRINOX. While 19 patients still remained in the LA group radiographically, R0 margins were achieved in 35 patients (92%) [25].

A French multicenter study by Wagner et al. [26] noted radiologic downstaging of resectability according to NCCN criteria in only 7/36 patients after FOLFIRINOX, while the majority of patients remained stable on imaging. However, R0 resections were achieved in 31 patients (86%), among them 6 patients with persistent LA disease on CT imaging.

In this light, several studies have focused on imaging predictors for resectability and outcome, with heterogeneous approaches and results.
Fig. 3 Imaging response using RECIST (Response Evaluation Criteria In Solid Tumors) version 1.1, and Evans pathologic response grades in pancreatic specimen found by Xia et al. [19]. RECIST classification: CR complete response, PR partial response, SD stable disease, PD progressive disease. Pathologic response grades: minimal, <50% tumor destruction; moderate, 50–90% tumor destruction, marked, >90% tumor destruction; complete, no tumor cells

Regression of vessel contact was explored in two studies

Cassinotto et al. found regressive circumferential tumor contact with crucial vessels (superior mesenteric vein, portal vein, and superior mesenteric artery) being highly predictive for R0 resections (PPV: 91%). Specifically, a regression in SMA contact yielded an odds ratio of 3.82 (95% CI: 1.27–11.5) for free margins [27]. However, no correlation between persistent narrowing of SMV/PV and R0 resection was observed.

Similarly, in the publication by Wagner [26], regressive circular vascular involvement, both arterial and venous (PV and SMV), resulted in longer disease-free survival than stable circumferential contact.

Fig. 4 a Locally advanced PDAC in a 52-year-old female at baseline (arrowhead), with encasement of the superior mesenteric artery of >180° (arrow). b The same patient after neoadjuvant therapy with FOLFIRINOX. Tumor shrinkage can be seen (arrowhead); however, SMA encasement persists (arrow). The patient underwent total pancreatectomy with negative resection margins. c Local recurrence 1 year after total pancreatectomy in the same patient (arrow).

Size and resectability

Michelakos et al. confirmed tumor size after FOLFIRINOX (median 2.3 cm in resected group vs. 3 cm in non-resected) to be associated with an attempt for resection at surgical exploration (R0 in the resected group: 80%), which is consistent with other studies [27, 29]. Changes in size were not predictive for resectability [28].

Tumor enhancement

Marchegiani et al., similar to prior studies [26, 27], observed an increase in contrast uptake in pancreatic tumors after NAT [29], attributed to fibrosis. Enhancement in their study was significantly higher in R0-resected cancer. However, in the publications by Wagner [26] and Casinotto [27], neither absolute enhancement nor enhancement changes after NAT were correlated to R0 resection.

In most studies, patients proceeded to resection when they had stable disease or partial response on imaging, while non-resected patients were excluded, which potentially distorts the comparison of imaging features to histologic margins. Furthermore, all studies were retrospective in design, and apart from the French multicenter study, all were single center.

While most studies were set up with two readers in consensus or with high interobserver concordance, interestingly, a study in 2010 presented inter-reader agreement for predicting resectability as an endpoint. A multi-rater kappa value of no more than 0.28 was calculated for predicting resectability after NAT, owed mainly to a high variability of sensitivities, ranging from 14 to 86%. However, the sample size was small with only seven resectable patients after NAT.

Does cross-sectional imaging always overestimate viable tumor?

While the majority of papers suggest an overestimation of the viable tumor remnant, Amodeo et al. re-
ported an underestimation in four out of seven patients who were classified as RECIST responders but had stable disease at laparotomy [30]. Also, with regard to NCCN resectability, cross-sectional imaging can occasionally underestimate the true extent of tumor: in the study by Marchegiani, seven patients were deemed resectable after NAT, but only five of them had R0 resection.

Finally, the true extent of tumor within fibroinflammatory tissue replacement has not been systematically explored so far with radiologic-pathologic lesion-by-lesion work-up.

**Functional imaging: 18-FDG PET and diffusion-weighted MRI**

18-FDG-PET is established in the response evaluation of lymphoma [31] and seems superior to morphologic response assessment in neoadjuvant therapy of esophageal cancer [32].

PERCIST1.0 criteria use a lean-body-mass-normalized standard uptake value (SUL) to compare peak values before and after therapy [33]. On the other hand, traditional parameters, such as maximum standardized uptake value (SUVmax) and body-weight-normalized SUV (SUVbw), are still in use to compare peak metabolic activities before and after NAT.

In a recent publication comprising 14 subjects and comparing both RECIST 1.1 and PERCIST1.0 criteria to pathologic residual tumor, an insignificant correlation of \( r = 0.26 \) was found between PERCIST and pathology, and none between RECIST and pathology. With known inadequacy of RECIST1.1, the limited but additional value of PERCIST was highlighted [34].

Prior studies with PET/CT similarly yielded ambiguous results. Mellon et al. [35] found a correlation between post-NAT SUVmax and pathologic response grade, but no correlation with the decrease of SUVmax. Of note, seven out of 81 patients with BR or LA-PDAC were excluded from the study, since PET failed to show any detectable baseline SUVmax.

A Japanese group confirmed a high variability of SUVmax at baseline and post NAT, as well as a high variability of regression indices [36]. For statistics, values were dichotomized along their median. Doing so, SUVmax after NAT was not associated with response, a result contrary to the work of Mellon, while baseline SUVmax and regression index were.

Diffusion-weighted MR imaging (DWI) offers functional tissue assessment by mapping the restriction of Brownian water molecule motion. Diffusion restriction is a marker for cellularity and pathologic characteristics of cellular barriers, both increased in tumors [37]. Calculating the apparent diffusion coefficient (ADC) in multiples of \( 10^{-3} \text{ mm}^2/\text{sec} \) from diffusion-weighted images allows for quantitative assessment of restricted diffusion. As such, on ADC maps, low values—depicted as dark areas—represent restricted diffusion (Fig. 5). ADC maps are widely investigated in oncologic imaging to estimate treatment response with promising results [38, 39].

Although Cuneo and colleagues first found a high correlation between ADC values and pathologic response in a small group of 7 patients [40], the variability of absolute pretreatment and posttreatment ADC values and their changes proved challenging. Okada et al. found pretreatment ADC values to be more indicative of response and surgical margins than posttreatment values and ADC changes. As another result of their study, MRI-ADC parameters outperformed the respective PET-SUVmax values [41].

In an analysis of 23 patients, however, Dalah et al. presented posttreatment ADC values and pathologic response to be moderately correlated (\( r = 0.517, p = 0.02 \)) [42]. The study is notable for using elaborate postprocessing tools such as color-coded ADC maps and histograms to demonstrate tumor heterogeneity after NAT.

Both publications pointed in the same direction by highlighting a significant difference between pre- and posttreatment mean ADC values.

Contrary to PET, which requires an interval of at least 8 weeks between NAT and restaging, in order...
to allow restitution of actinic inflammation, MRI was performed within 3–5 weeks after completing NAT.

Both functional methods may allow for estimating the overall response of a mass and predict survival [37, 43], but they lack the spatial resolution to predict surgical margins around vessels. Still, in the light of evidence that pathologic response grade might be a factor associated with survival, biomarker imaging may play a role in future decision algorithms.

**Conclusion**

The NCNN and ESMO panels currently endorse neoadjuvant therapy in borderline resectable PDAC [44, 45] and several individual studies have shown that in a minority of patients, NAT enabled R0 resection of locally advanced PDAC. However, on an intention-to-treat basis, according to meta-analyses, the overall R0 rate was only about 54% in BR-PDAC [20] and 22% in LA-PDAC [22].

This poses a considerable challenge to interdisciplinary teams seeking to identify patients with potentially resectable disease. Histologic work-up suggests an inhomogeneous response of PDAC with interspersed tumor nests within fibrotic changes.

The current consensus is that radiologic mass regression occurs in a minority, while most patients remain stable on imaging. Generally, cross-sectional imaging overestimates the amount of residual viable tumor around vessels and thus cannot predict resectability.

There is no consistency across studies with regard to predictive imaging parameters. Most studies are retrospective, single-center, observational studies, examining a range of hypotheses, such as changes in tumor size, vessel contact, or enhancement. Due to heterogeneity, results are non-comparable; furthermore, statistical power is limited by small sample sizes.

At present, guidelines recommend taking patients to surgery after NAT when there is no tumor progression on cross-sectional imaging.

Functional imaging such as 18-FDG-PET/CT and MR-DWI reveal conflicting results in the search for optimal imaging predictors of response. But while markers of low cellularity may be indicators of response, to date, functional imaging methods lack the spatial resolution to detect microscopic disease at the crucial interface between mass and vessel wall. It is thought that this limit will be overcome in the years to come.

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