New Advances in the Treatment of Metastatic Pancreatic Cancer

Marvin Schober\textsuperscript{a}  Muhammad A. Javed\textsuperscript{d}  G. Beyer\textsuperscript{b}  Nha Le\textsuperscript{e}  Alessio Vinci\textsuperscript{g}  
Malin Sund\textsuperscript{f}  Albrecht Neesse\textsuperscript{c}  Sebastian Krug\textsuperscript{a}

\textsuperscript{a}University of Halle, Department of Gastroenterology and Hepatology, Halle, \textsuperscript{b}Department of Medicine A, University Medicine, Greifswald University, Greifswald, and \textsuperscript{c}University Medical Centre Göttingen, Department of Gastroenterology and Gastrointestinal Oncology, Göttingen, Germany; \textsuperscript{d}NIHR Liverpool Pancreas Biomedical Research Unit, Department of Molecular and Clinical Cancer Medicine, Royal Liverpool University Hospital, Liverpool, UK; \textsuperscript{e}Semmelweis University, Second Internal Medicine Department, Gastroenterology Division, Budapest, Hungary; \textsuperscript{f}University of Umea, Department of Surgical and Perioperative Sciences, Umea, Sweden; \textsuperscript{g}University of Pavia, Department of Surgery, IRCCS S. Matteo University Hospital Foundation, Pavia, Italy

formulated albumin-bound paclitaxel (nab-paclitaxel) in combination with gemcitabine also resulted in a significant survival extension compared to gemcitabine monotherapy. However, both intensified therapy regimens show a broad spectrum of side effects and patients need to be carefully selected for the most appropriate protocol.

Key Message:
In this study, recent advances in the chemotherapeutic options available to treat metastatic PDAC and their implications for today’s treatment choices are reviewed.

Abstract
Background: Pancreatic ductal adenocarcinoma (PDAC) is characterised by an extremely poor overall survival (OS) compared to other solid tumours. As the incidence of the disease is rising and the treatment options are limited, PDAC is projected to be the 2nd leading cause of cancer-related deaths in the United States by 2030. A majority of patients are not eligible for curative resection at the time of diagnosis, and those that are resected will often relapse within the first few years after surgery. Summary: Until recently, the nucleoside analogue gemcitabine has been the standard of care for patients with non-resectable PDAC with only marginal effects on OS. In 2011, the gemcitabine-free FOLFIRINOX regimen (folinic acid, fluorouracil, irinotecan and oxaliplatin) showed a significant survival advantage for patients with metastatic PDAC in a phase III trial. In 2013, the Metastatic Pancreatic Adenocarcinoma Trial phase III trial with nano-

Introduction
Pancreatic ductal adenocarcinoma (PDAC) is considered to be one of the most lethal solid organ tumour. The extraordinarily poor prognosis can be attributed to the lack of early clinical symptoms and good biomarkers, early metastatic dissemination and an unusually high resistance to targeted and cytotoxic agents. The incidence of pancreatic cancer ranges worldwide from 1 to 10 cases per 100,000, out of which 85% are adenocarcinomas [1]. The
Incidence is slightly higher in developed countries, among men, and a number of risk factors as well as genetic syndromes/mutations that are associated with pancreatic cancer have been identified [1, 2]. The genetic hallmark of PDAC is an activating KRAS mutation, followed by the deletion or the inactivation of tumour-suppressor genes such as p16, DPC4 and p53 [3–5]. Extensive experimental studies in mice and men have identified and characterised a molecular and histological progression model ranging from early to advanced precursor lesions (so called pancreatic intraepithelial neoplasia lesions, PanIN I–III), to frank adenocarcinoma. Moreover, PDAC is typically accompanied by an extensive desmoplastic stroma reaction contributing to a hypovascular and hypoxic microenvironment, which possibly affects the delivery and efficacy of chemotherapeutic agents, reprogramming of cellular metabolism and evasion of tumour immunity [6–8].

Clinical symptoms of PDAC patients are often very unspecific and hard to detect early. Patients may complain of abdominal discomfort (with no signs of peritoneal irritation), weight loss, painless jaundice and unspecific back pain [9, 10]. Onset of type II diabetes as well as unexplained deep vein thrombosis may also indicate the development of PDAC, but also occur for many other reasons in the elderly. Once the diagnosis of PDAC is suspected, a multi-modal diagnostic work-up using modalities such as sonography, endosonography, multi-slice CT as well as MRI should be performed in a multi-disciplinary team. In case of a metastatic disease, histological confirmation prior to cytotoxic treatment is obligatory. Surgical resection is the only potentially curative treatment and criteria of surgical resectability are based on the involvement of the local vessels [11] and presence or absence of metastatic disease. Until now, only about 20–25% of patients are eligible for surgical resection. Current treatment standards advocate the use of adjuvant chemotherapy following surgery. For locally advanced pancreatic adenocarcinoma (LAPC), there is growing interest in the use of neoadjuvant approaches, which are being evaluated, as in the currently recruiting Conko 007 trial (EUDRACT Nr. 2009-014476-21). This phase III trial is evaluating the role of a combined chemoradiotherapy after leading-in with chemotherapy in patients with LAPC, interestingly already making use of FOLFIRINOX as a potential treatment arm. The currently enrolling NEONAX-trial, a phase II trial conducted by the AIO (EUDRACT-Nr. 2013-005559-34), is addressing the question of whether or not a neoadjuvant combination treatment with nab-paclitaxel and gemcitabine plus adjuvant or only adjuvant gemcitabine and nab-paclitaxel treatment can downstage the tumour prior to surgery in patients with resectable PDAC (table 1). Despite numerous clinical trials in stage IV (metastatic) PDAC, there has only been a modest improvement of median survival rates for this group of patients [12–15]. In the last 2 decades, biomarker discovery and validation has been a key area of research to stratify patients, predict response to treatment and enable the development of a tailored treatment approach. The aim of this review is to provide a comprehensive overview of the development of chemotherapeutic regimens that have been used for the treatment of metastatic PDAC with a particular focus on the recently introduced gemcitabine-free FOLFIRINOX regimen (folinic acid, fluorouracil, irinotecan and oxaliplatin) and nab-paclitaxel + gemcitabine.
Current Management of Metastatic Disease

From 5-Fluorouracil to Gemcitabine

5-Fluorouracil (5-FU) was considered the only chemotherapeutic option for about 20 years until the introduction of gemcitabine [13, 16–18]. Several clinical studies could show that chemotherapy in advanced pancreatic cancer patients indeed leads to a modest survival benefit as well as improved quality of life (QoL) compared to best supportive care alone [19–21], but the combination of 5-FU with other drugs did not prove to be superior to the single anti-metabolite treatment [16, 22]. During the 1990s, several trials were designed to find new active drugs in pancreatic cancer treatment and the nucleoside analogue gemcitabine (2′-deoxy-2′,2′-difluorocytidine) was studied [12, 23]. In 1997, gemcitabine became the first line choice for PDAC patients due to fewer side effects (compared to 5-FU) and a significant survival benefit (table 2). This clinical study also implemented the concept of clinical benefit response (CBR) that was defined as a ≥50% reduction in pain intensity, daily analgesia consumption or ≥20 point improvement in Karnofsky performance status (PS) for ≥4 consecutive weeks. CBR was significantly improved in gemcitabine-treated patients (p = 0.0022) [13]. The CBR was shown to be 23.8% in gemcitabine-treated patients (1,000 mg/m² over 30 min) compared to 4.8% in 5-FU-treated patients (400 mg/m² over 30 min), and the median survival time in the gemcitabine group was 5.65 months compared to 4.41 months in the 5-FU-group, respectively (p = 0.0025). In the gemcitabine group, 18% compared to 2% for the 5-FU survived the first 12 months [13]. Several phase III trials with gemcitabine monotherapy revealed a median survival of 5–7 months with a
1-year survival of about 1–25%. Therefore, subsequent chemotherapy trials in PDAC needed to show superiority over standard of care gemcitabine monotherapy. This could be achieved by either of the 2 strategies: (1) combining gemcitabine with a new drug or (2) directly comparing a novel monotherapy or combination therapy against gemcitabine.

**Gemcitabine and Fluoropyrimidines**

While the results of gemcitabine clearly improved therapeutic efficacy and 1-year survival, its impact on median overall survival (OS) remained at the margin of significance. More intensive combination chemotherapies involving fluoropyrimidines and platinum analogues were investigated in numerous phase II and III trials. Two randomised phase III trials evaluated the combination of gemcitabine and 5-FU compared to gemcitabine alone. In short, the experimental arm did not improve the objective response or median OS [24, 25].

Capecitabine is an oral, tumour-selective fluoropyrimidine carbamate and pro-drug that is activated by a 3-step-targeted process (carboxylesterases, cytidine deaminase and thymidine phosphorylase, respectively). Capecitabine can be used in schedules that provide prolonged fluorouracil exposure at lower peak concentrations, thus mimicking continuous infusion of fluorouracil. Preclinical studies have shown no synergistic effect of capecitabine anti-tumour activity when combined with gemcitabine; however, relevant toxicity was noted in genetically engineered mice of pancreatic cancer more recently [26]. Herrmann and Cunningham published 2 randomised phase III trials of gemcitabine +/- capecitabine (650 or 830 mg/m² twice daily) in patients with metastatic PDAC. There was no statistically significant improvement of the median OS in the combination arm (8.4 vs. 7.2 m and 7.4 vs. 6.0 m; p = 0.234 and p = 0.08). However, a subgroup of patients with good PS (Karnofsky performance score 90–100) benefited (10.1 vs. 7.4 months, respectively; p = 0.014) from the combination treatment. In addition, in the Cunningham et al. [27] study, gemcitabine and capecitabine significantly improved the response rate (RR) (19.1 vs. 12.4%, p = 0.034) and progression-free survival (HR 0.78, p = 0.004). Interestingly, by pooling the data set of these 2 phase III trials and the phase II trial of Schreithauer in 935 patients, statistical significance could be achieved for median OS (HR 0.86; 95% CI 0.75–0.98, p = 0.02) [19, 20, 28]. Independently from this data, Hubner et al. [29] presented single-centre results of the gemcitabine plus capecitabine combination, leading to a median OS of 8.7 m (95% CI 6.7–10.7) and 1-year survival rate of 34%. Despite the fact that gemcitabine combined with fluoropyrimidine has reliable activity in metastatic PDAC, the QoL did not significantly differ between the 2 treatment arms [29–31]. Based on this data, the combination GEMCAP has replaced 5-FU therapy in clinical practice, especially in the United Kingdom.

**Gemcitabine in Combination with Platinum-Derivates and Topoisomerase Inhibitors**

There have been numerous phase II and III studies attempting to improve the efficacy and OS of chemotherapy in metastatic PDAC by using gemcitabine-based combination therapies. The combination of gemcitabine with platinum-derivates was evaluated in 11 trials in more than 2,000 patients [31]. One of the most promising trials was the GERCOR/GISCAD study where the combination of gemcitabine and oxaliplatin improved the PFS (5.58 vs. 3.7 months), CBR (38.2 vs. 26.9%) and RR (26.8 vs. 17.3%). However, no statistical significance was reached for OS [32] and the subsequent phase III trials did not confirm these encouraging results [33–35]. A recently published meta-analysis compared gemcitabine plus oxaliplatin with gemcitabine alone. Here, ORs of 1.33 (95% CI 1.05–1.69, p = 0.019) for OS and ORs of 1.38 (95% CI 1.08–1.76, p = 0.011) for PFS demonstrated and showed statistical significance [19, 20, 28, 31].

Similar results were obtained for gemcitabine and topoisomerase inhibitors (irinotecan and exatecan) combinations. Although preliminary data suggested a relevant clinical activity for topoisomerase inhibitors, 4 randomised clinical trials revealed no significant improvement of PFS or OS [36–39]. Finally, in the Cancer and Leukemia Group B (CALGB) study, 3 relevant combination therapies (gemcitabine plus cisplatin, gemcitabine plus docetaxel, gemcitabine plus irinotecan) vs. gemcitabine monotherapy demonstrated no relevant 6-month survival benefit for the experimental arms [36]. Therefore, the authors made a decision to refrain from further investigating gemcitabine-based combinations for PDAC.

**Gemcitabine and Erlotinib**

The epidermal growth factor receptor inhibitor erlotinib was the first drug that was shown to be beneficial in combination with gemcitabine in patients with advanced pancreatic cancer (table 2). In the PA.3 trial, 569 patients were randomised with both locally advanced and metastatic disease to receive gemcitabine or gemcitabine plus erlotinib. Though OS was significantly improved with combination therapy (median survival of 6.2 vs. 5.9
months), clinical relevance of a 2-week survival benefit had to be questioned [40]. This marginal benefit also came at the expense of toxicity with 6 treatment-related deaths reported during the study – all within the combination therapy group. Eight patients from the study (7 from the erlotinib cohort) also developed interstitial lung disease and the risk-benefit of this drug combination remains a controversial topic [41].

Interestingly, the combination of erlotinib and gemcitabine achieved astonishing response in a small subset of patients. It was noted that those patients developing a rash of grade ≥2 in severity showed a median survival of 10.5 months and 1-year survival of 43% in contrast with 5.3 months and 16% respectively in patients with no rash. Therefore, the German S3-guideline for exocrine pancreatic cancer recommends an initial therapeutic attempt with erlotinib and gemcitabine for an 8-week period. After this time, treatment should be continued only if a skin rash develops [11]. This interesting finding could be recapitulated in another study by Vaccaro et al. [42]; however, dose-escalation did not yield further therapeutic efficacy.

**Folinic Acid, Fluorouracil, Irinotecan and Oxaliplatin (FOLFIRINOX)**

Several studies were performed to investigate gemcitabine-based polychemotherapy regimens comprising several cytotoxic drugs. The PEFG protocol compared cisplatin, epirubicin, 5-FU and gemcitabine to gemcitabine alone [43]. The same group introduced another 2, 4-drug combinations based on the PEFG regimen called PEXG (cisplatin, epirubicin, capecitabine and gemcitabine) and PDGX (cisplatin, docetaxel, capecitabine and gemcitabine). Although median OS of 11 m were reached in this phase II study, the results were interpreted with caution, since only 2/3 of the patients had metastatic disease [44]. Another regimen called G-FLIP comprised CPT-11 (irinotecan), gemcitabine, 5-FU, leucovorin and cisplatin [45], and subsequently FOLFOX-6 and FOLFIRI-3 were evaluated in the first-line setting of advanced PDAC [46, 47]. Despite median OS times between 8 and 12 months and higher RRs, increased major toxicities, the inhomogeneity of the patient cohorts and the insufficient inclusion criteria gave rise to doubts whether polychemotherapy was the adequate treatment strategy.

FOLFIRINOX is a chemotherapy regimen comprising 4 drugs – folinic acid, 5-FU, irinotecan and oxaliplatin. This combination of drugs was supported by preclinical studies demonstrating either single agent efficacy against pancreatic cancer or synergistic activity in combination with a fellow agent. Based on ambitious phase I/II data in which FOLFIRINOX revealed a good safety profile and appealing RRs, Conroy et al. [48] conducted a French phase II/III trial (PRODIGE) with OS as primary objective, and PFS, QoL, RR and toxicity as secondary endpoints [48, 49]. In total, 342 patients were enrolled and FOLFIRINOX was administered with the following dose: oxaliplatin at a dose of 85 mg/m² over 2 h followed by leucovorin at a dose of 400 mg/m² given as a 2-hour intravenous infusion and irinotecan at a dose of 180 mg/m² in a 90-minute intravenous infusion [50]. This treatment was immediately followed by 5-FU at a dose of 400 mg/m², administered by intravenous bolus, followed by a continuous intravenous infusion of 2,400 mg/m² over a 46-hour period every 2 weeks. The patients on FOLFIRINOX demonstrated a median survival of 11.1 months, which was clearly superior when compared to the 6.8 months of the gemcitabine control group (table 2). The median progression-free survival was 6.4 months in FOLFIRINOX-treated patients versus 3.3 months in patients who received gemcitabine [50]. Compared to treatment with gemcitabine, treatment with FOLFIRINOX resulted in a superior tumour RR (9.4 vs. 31.6%); however, 50 patients (25 per arm) were not radiologically assessable. Yet in 2005, this regime showed promising results, when it had been administered to a heterogeneous group of patients just defined as histologically proven advanced PDAC, but containing both LAPC and metastatic PDAC. Thus, the treatment regimen was not exclusively restricted to patients suffering from metastatic disease [48]. However, in order to achieve the benefits shown, the FOLFIRINOX regimen requires a careful selection of patients, which implicates strict limitations as the potential real-life first line option. In the study by Conroy et al., only patients with nearly unrestricted ECOG-PS of 0–1, below 76 years of age, no cardiac ischemia within 1 year before entry, and total bilirubin levels below 1.5 times the upper normal limit were included. Interestingly, only 35–40% suffered from a pancreatic head adenocarcinoma and therefore, solely 15% received stenting of the biliary duct. No single cholangitis had been observed during the course of the trial. This difference may be related to the exclusion of patients with elevated bilirubin levels, because of the increased risk of irinotecan-induced toxicity [51]. As shown by Conroy et al., even de-escalation (n = 80 received second-line treatment in the FOLFIRINOX arm) might be necessary; yet it resulted in an impressive median OS of 4.4 months from the beginning of second-line chemotherapy. Particularly, this accounts as one explanation for the 1-year survival rate of 48.4% for patients.
treated with FOLFIRINOX versus 20.6% for patients receiving gemcitabine (p < 0.001). Despite the strict selection criteria for the study, patients on FOLFIRINOX suffered more frequently from adverse events or severe side effects compared to the patients on gemcitabine. Forty-five percent of the FOLFIRINOX patients developed neutropenia and 5.4% showed febrile neutropenia. In addition, thrombocytopenia (9.1%), diarrhea (12.7%) and sensory neuropathy (9.0%) occurred significantly more often in FOLFIRINOX patients, which implies a need for increased clinical routine check-up intervals for patients receiving this intensive regime. Moreover, supportive therapies such as nutritional support, G-CSF treatment or antibiotics are more often needed, and the discontinuation of the therapy may be required. However, G-CSF was not recommended in this trial to prevent neutropenia. Two years after the publication of the primary study data, QoL evaluations were presented [52]. Although FOLIRINOX had remarkable side effects, it significantly reduced QoL impairment compared with gemcitabine in patients with metastatic PDAC. Only diarrhea was significantly increased in the FOLIRINOX arm. Beyond the PS, age >65 years and low serum albumin were seen as independent prognostic factors for poor survival in this situation.

**Gemcitabine and Nab-Paclitaxel**

In 2013, a second intensified combination-chemotherapy regimen – gemcitabine with nab-paclitaxel – was presented for the treatment of patients suffering from metastatic PDAC. Metastatic Pancreatic Adenocarcinoma Trial (MPACT-Trial).

Nab-paclitaxel is a nanoparticle-sized, albumin-bound paclitaxel, which was originally developed to avoid toxic and immunogenic solvents of hydrophobic paclitaxel, and was first approved by the FDA in breast cancer patients in 2005 [7, 53–57].

In PDAC, Von Hoff et al. [53, 54] demonstrated in a phase II trial that patients with SPARC (secreted protein acidic and rich in cysteine) positive tumours revealed a highly significant benefit in OS when treated with gemcitabine plus nab-paclitaxel. Therefore, it was hypothesised that stromal SPARC as an albumin-binding protein would bind albumin-coated paclitaxel molecules, thereby selectively increasing drug delivery. However, preclinical data with genetically ablated SPARC mice showed comparable nab-paclitaxel concentrations within the endogenous murine pancreatic tumours compared to neoplasms overexpressing SPARC [58]. Controversial data in men and mice exist regarding the potential of nab-paclitaxel to deplete abundant tumour stroma in PDAC [55, 59, 60]. Therefore, it remains questionable whether nab-paclitaxel exerts any tumour-selective/targeted effect beyond its cytotoxicity as a mitotic inhibitor.

Nevertheless, the international multi-centre MPACT-trial revealed significant clinical benefits, achieving disease control rates of 67% in treated patients and a median OS of 8.5 months [53, 54, 61] (table 2).

MPACT trial patients (n = 431) receiving the combination treatment of gemcitabine plus nab-paclitaxel showed a 1-year survival rate of 35 vs. 22% when compared to the gemcitabine monotherapy group (n = 430), and 9% compared to 4% at the second year of follow-up. Moreover, a median progression-free survival of 5.5 months in the combination therapy cohort versus 3.7 months in the gemcitabine monotherapy group had been achieved. Severe neutropenia was significantly more frequent in the combination therapy group of the MPACT trial (gemcitabine + nab-paclitaxel: 38%, gemcitabine: 27%), while febrile neutropenia occurred in 3% of the patients. The same was true for the incidence of fatigue (17 vs. 7%) and neuropathy (17 vs. 1%) that occurred significantly more often in the nab-paclitaxel group than in the gemcitabine monotherapy group [53].

Strikingly, immunohistochemical analysis from MPACT phase III patients failed to confirm that SPARC is a prognostic biomarker [62], confirming the earlier published preclinical data [58].

In a recently published phase-III-trial (n = 861 patients) by Goldstein et al. [63], analysing the long-term survival of patients receiving the nab-paclitaxel plus gemcitabine combination-treatment, the superior OS and PFS could be confirmed. Moreover, subgroup analysis emphasised the relevance of CA 19–9 and neutrophil to lymphocyte ratio as reasonable prognostic markers. High CA 19–9 levels and neutrophil to lymphocyte ratio before treatment was associated with a worse OS. Survivors of more than 3 years were exclusively identified in the nab-paclitaxel plus gemcitabine arm (4%) [63]. By performing post-hoc analyses of the MPACT collective, Tabernero et al. [64] described PS score, liver metastases, age and number of metastatic sites as consistent and valid predictors of survival that are easily available clinical markers that help in decision making regarding treatment.

Notably, a retrospective analysis showed that the eligibility criteria itself for both the PRODIGE and MPACT trial, were associated with an improved survival [65]. The authors demonstrated that patients who simply fulfilled the inclusion criteria for either FOLFIRINOX or nab-paclitaxel/gemcitabine, without actually receiving the treat-
ments, had a better median OS compared to patients who failed to meet them. When comparing the FOLFIRINOX-eligible population with the nab-paclitaxel-eligible group, it becomes apparent that ECOG 2 patients were excluded in the FOLFIRINOX trial. These results were consistent with the fact that eligibility was no longer a significant prognostic factor after accounting for patient’s ECOG performance. Strictly following the trials inclusion criteria, the authors demonstrated that the majority of patients with metastatic PDAC were not eligible for either of the intensified polychemotherapeutic regimens. Comparing only eligibility for treatment for nab-paclitaxel plus gemcitabine versus the FOLFIRINOX protocol, nearly twice as many patients could have been candidates for nab-paclitaxel due to less restrictive ECOG performance criteria for this regimen in trials [65].

Furthermore, novel drug formulations that improve the pharmacokinetic properties of hydrophobic paclitaxel have been investigated. For instance, Löhr et al. [66] have shown in a phase II trial that paclitaxel embedded in cationic liposomes, the so-called EndoTAG-1, was generally well tolerated and achieved a median OS of 8.1–9.3 months when combined with gemcitabine.

Ongoing clinical trials are investigating nab-paclitaxel as the backbone for novel targeted therapies. Examples are ruxolitinib, a janus-kinase inhibitor in combination with gemcitabine or nab-paclitaxel plus gemcitabine (NCT01822756), INCB039110 a JAK1 inhibitor in combination with nab-paclitaxel plus gemcitabine (NCT01858883), or the indoleamine-2,3-dioxygenase inhibitor Indoximod in combination with nab-paclitaxel plus gemcitabine in advanced PDAC (NCT02077881). Those results might help to optimise the selection of adequate patient cohorts eligible for nab-paclitaxel, and moreover provide additional insights into novel targeted treatment avenues (table 1).

**Conclusion**

It is evident from the existing data that there is a significant survival benefit of intensive chemotherapy treatment regimens FOLFIRINOX and gemcitabine + nab-paclitaxel for metastatic PDAC. Although OS rates may be overestimated due to very stringent enrolment criteria for these trials, national and international guidelines recommend them as first-line therapy for metastatic PDAC. However, careful patient selection based on clinical and laboratory findings represents a major challenge for physicians and oncologists in the palliative situation (fig. 1). Currently, no biomarker can reliably predict the response to treatment to nab-paclitaxel + gemcitabine or FOLFIRINOX. Therefore, a careful clinical and biochemical follow-up is required to optimise the treatment effects while minimising the side effects. Often, dose reductions are a feasible approach to cope with the enhanced toxicity profile of these protocols. For the future, intensified chemotherapies such as nab-paclitaxel and FOLFIRINOX may also be extended to selected patients with LAPC in a neoadjuvant setting, as well as for adjuvant treatment strategies. Furthermore, intensified chemotherapy protocols are currently investigated as backbone for novel investigational PDAC therapies, such as anti-stromal approaches or immune cell checkpoint antagonists.

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34 Heinemann V, Quietsch D, Gieseler F, Gon- metastatic Pancreatic Cancer


