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Advanced pancreatic ductal adenocarcinoma - Complexities of treatment and emerging therapeutic options

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**Abstract**

Pancreatic ductal adenocarcinoma is a devastating disease with a poor prognosis regardless of stage. To date the mainstay of therapy for advanced disease has been chemotherapy with little incremental im­provements in outcome. Despite extensive research investigating new treatment options the current practices continue to utilise fluorouracil or gemcitabine containing combinations. The need for novel the­rapeutic approaches is mandated by the ongoing poor survival rates associated with this disease. One such approach may include manipulation of ribosome biogenesis and the nucleolar stress response, which has recently been applied to haematological malignancies such as lymphoma and prostate cancer with promising results. This review will focus on the current therapeutic options for pancreatic ductal adenocarcinoma and the complexities associated with developing novel treatments, with a particular emphasis on the role of the nucleolus as a treatment strategy.

**Key words:** ribosome biogenesis; nucleolar stress; RNA polymerase I; Pancreatic ductal adenocarcinoma; chemotherapy

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**Core tip:** This manuscript is a review of the complexities involved in the treatment of advanced pancreatic ductal adenocarcinoma. It details the current approaches to therapy and the disease factors which have impacted on progress thus far. This review identifies the possible role of nucleolar stress as a treatment modality based on recent data from studies of haematological ma­lignancies and some other solid organ cancers and explains the basic science involved in this process.

**INTRODUCTION**

Advanced pancreatic ductal adenocarcinoma (PDAC) remains a significant cause of mortality accounting for up to 4% of all cancer related deaths world-wide[1]. Of concern, despite treatment, the mortality rates remain high and essentially unchanged over the last two decades[2]. Furthermore, even in the setting of adjuvant therapy, 5 year survival rates remain poor at only 25%[2], while for metastatic disease it is 1%[3]***.*** Such poor statistics are no doubt compounded by the fact that current backbones to treatment, fluorouracil or gemcitabine, have also remained unchanged over the last 10-20 years and continue to form the basis for new combination approaches[4]. As a consequence, there is an ongoing need for the development of im­proved diagnostic techniques, treatment options and surveillance markers to improve the outcomes for this devastating disease.

In recent years, various studies have focused on trialling new chemotherapeutic drug combinations for the treatment of PDAC. The PRODIGY 4/ACCORD 11 trial[5,6] demonstrated that FOLFIRINOX, a combination of fluorouracil, oxaliplatin and irinotecan, provided superior progression free survival (PFS), overall survival (OS) and response compared with gemcitabine alone for patients with metastatic disease. Unfortunately, however, the drawback was that many patients dis­played significant toxicity associated with this multidrug approach including an higher incidence of grade 3 or 4 neutropenia and thrombocytopenia[5].

Two years later, the MPACT trial compared the combination of gemcitabine and nab-paclitaxel against gemcitabine alone in a total of 842 patients and demonstrated a statistically significant improvement in median OS (8.7 mo *vs* 6.6 mo)[7] and was particularly interesting given that previous investigations comparing gemcitabine combinations against gemcitabine alone for PDAC had been negative. Importantly, this trial did not exclude more elderly patients unlike the ACCORD-11 study which only incorporated patients with a mean age of 61 years[5]. In keeping with PDAC demographics, MPACT included patients with a mean age for men of 71 and 75 for women[8]. Toxicities related to gemcitabine and nab-paclitaxel were overall very similar to those seen with FOLFIRINOX though there is a perception that FOLFIRINOX is best reserved for “fit” patients despite the two regimens never formally being compared.

More recently, the NAPOLI-1 study compared the effect of nanoliposomal irinotecan with or without fluorouracil and folinic acid as well as the combination of fluorouracil and folinic acid alone in patients with metastatic disease who progressed following a gemcitabine based approach[9]. In its nanoliposomal form, irinotecan has improved drug stability allowing its active metabolite, SN-38, to remain in circulation longer than free irinotecan, ultimately prolonging intra-tumoral levels of the active drug[10,11]. The median OS was significantly improved for those receiving nanoliposomal irinotecan, fluorouracil plus folinic acid (6.1 mo *vs* 4.2 mo) compared with patients who did not receive nanoliposomal irinotecan. Similarly, the PFS noted for those receiving combination therapy with nanoliposomal irinotecan was significantly improved compared to fluorouracil and folinic acid alone at 3.1 mo *vs* 1.5 mo. Differences between nanoliposomal irinotecan monotherapy and fluorouracil with folinic acid alone were not statistically significant[9]. Based on this study and others the FDA has approved the use of nanoliposomal irinotecan with fluorouracil and folinic acid for treatment of metastatic PDAC refractory to gemcitabine based therapy. Interestingly the toxicities seen with this newer regimen were relatively similar to those seen with FOLFIRINOX and gemcitabine with nab-paclitaxel including grade 3 or 4 toxicities in such domains as neutropenia, diarrhoea, vomiting and fatigue.

While new chemotherapy combinations have shown promise in the clinical setting, PDAC remains a difficult disease to manage due to multiple factors including patient age, medical co-morbidities and cancer related symptoms. As is often the case in advanced/metastatic malignancies, the art of medicine relies on achieving a balance between therapeutic gain, drug induced side effects and quality of life. To this end, novel approaches that are non, or at least minimally genotoxic are of particular interest. The potential advantages observed with targeted therapies in reducing systemic side effects and improving outcomes for other malignancies have gained interest in the realm of PDAC and will be briefly discussed in this review with a particular focus on the emerging role of nucleolar stress pathway and ribosome biogenesis as a potential treatment option given its promising results in the management of haematological malignancies and prostate cancer[12-14].

**DISEASE COMPLEXITIES**

When evaluating the role of novel therapies, it is important to understand potential interactions with the disease in question, specifically focusing on disease factors that could impact on drug delivery, tolerability or dosing for example. PDAC is associated with multiple complexities which have precluded the development of novel approaches and therefore require due con­sideration.

***Patient factors***

PDAC is a disease of the elderly, presenting on average at 71 years of age[3]. Up to 9% of patients present with localised disease however, the majority are diagnosed with either locally advanced or metastatic disease at their first consultation[15]. Due its anatomical location the symptoms associated with PDAC tend to occur insidiously therefore contributing to the often delayed time to investigation and subsequent diagnosis[16]. Such patient factors have important implications on treatment decisions which may in part explain the limited use of FOLFIRINOX in many patients despite its improved median OS, PFS and objective response[5].

Given the advanced age and stage at diagnosis the traditional focus on clinical outcomes and survival for interventional studies of PDAC management have shifted with increasingly more importance being placed on patient reported outcomes such as pain manage­ment and appetite. This is of particular importance for advanced PDAC given its poor prognosis[17]. Without acknowledging the impact the treatment may have on patient reported outcomes, therapeutic advancements may be futile, thus developing drugs that are minimally toxic would be beneficial, yet to date, therapeutic advancements have continued to cause very similar side effect profiles to one another.

***Genetic basis of PDAC***

Inherited predispositions to PDAC account for 5%-10% of cases[18]. In ongoing studies through the Australasian Pancreatic Cancer Genome Initiative, Humphris *et al*[19] described the manifestations of inherited PDAC as occurring in 3 distinct settings, hereditary tumour predisposition syndromes, hereditary pancreatitis and familial pancreatic cancers which were further defined as occurring in a kindred in whom at least 2 first degree relatives have PDAC without diagnostic criteria for an inherited cancer syndrome[19,20]. Interestingly some of the genes identified in hereditary forms of PDAC affect pathways involved in DNA repair such as *BRCA1/2* and *PALB2* which have been noted in more recent detailed sequencing/mutational studies presented by Waddell *et al*[21] and Bailey *et al*[22] in two separate comprehensive analyses. While inherited forms of cancer are of interest the vast majority of genetic/metabolic pathway abnormalities are not inherited and form the basis of most PDAC cases.

Four major driver genes, *KRAS, TP53, CDKN2A* and *SMAD4* have been identified in the development of PDAC[22], each sharing common oncogenic signalling pathways. *KRAS* mutations occur in > 90% of tumours and may represent the underlying insult to numerous subsequent events contributing to disease development. This mutation is widely accepted as a requirement for “reprogramming” pancreatic cell metabolism to facilitate the acidic environment needed for extracellular matrix breakdown and tumour invasion common to PDAC[23,24].

Of interest, the complex genetic and metabolic pathways associated with PDAC have identified various interactions and sites which can be utilised for therapeutic means. Among these include certain growth factor receptors such as epidermal growth factor receptor (EGFR) and vascular endothelial growth factor receptor which have important roles in the RAS/RAF pathway as outlined in figure 1. Furthermore it is now clear that drugs such as nab-paclitaxel have therapeutic actions against some of these pathways either directly or downstream accounting for their efficacy.

Recent data published in Nature by Waddell *et al*[21] reported the complex mutational landscape of PDAC, identifying multiple point mutations and structural variations in key genes. This data confirmed that *KRAS* abnormalities were almost ubiquitous while *TP53* lesions were noted in 74% of samples, closely followed by *CDKN2A* lesions(35%) *and SMAD4* abnormalities (31%). Through their analysis, PDAC was sub classified into 4 subtypes based on the distribution of events such that samples fell into either stable, scattered, unstable or locally arranged groupings. Of interest, the same study established a relationship between mutational load and abnormalities affecting DNA maintenance genes, specifically *BRCA*[25] and *PALB2*[26]. It is not surprising that PDAC associated with either *PALB2* or *BRCA2* mutations should behave in a similar fashion given their roles in DNA damage and repair. *PALB2* binds and co-localizes with *BRCA2* in order to facilitate double stranded DNA damage. In cases where PDAC harbours *BRCA2* mutations there is good evidence for improved response to platinum containing therapies unlike other forms of PDAC[21]. For *PALB2* mutant disease there have been a number of reports noting improved outcomes in response to platinum based therapies or mitomycin C when compared with gemcitabine[27,28] and there is potential for targeted treatment with poly ADP ribose polymerase (PARP) inhibitors especially in the setting of *BRCA2* as seen in breast and ovarian cancers.

Further to the comprehensive studies of Waddell *et al*[21], Bailey *et al*[22] identified aggregates of point mutations in core molecular pathways affecting ce­llular functions including DNA damage and repair pathways, cell cycle regulation, transforming growth factor beta (TGF) signalling, chromatin regulation and axonal guidance. Based on the expression of 32 recurrently mutated genes found to aggregate into ten distinct pathways. From this analysis four subtypes were identified, comprising of, squamous, pancreatic progenitor, immunogenic and aberrantly differentiated endocrine and exocrine categories[22]. Similar to Waddell *et al*[21] the implications of these findings may potentially identify opportunities for therapeutic development. While these data have provided valuable insights into the complex molecular basis of PDAC many studies have previously attempted to manipulate some of these genes and/or pathways at various levels already and will be briefly discussed here.

Given the near ubiquitous nature of *KRAS* mutations in PDAC, targeting this gene and its associated pathways has been an area of interest however thus far this has not translated to clinically significant outcomes. In an effort to target as many known mutations/pathways as possible various studies have focused on novel therapies in combination with known chemotherapeutics in the hopes that treatment outcomes improve.

Selumetinib, an orally bioavailable selective MEK1/2 inhibitor showed promise in preclinical studies but failed to demonstrate a survival advantage in the second line setting when compared with the oral equivalent to fluorouracil, capecitabine[29]. Similarly, attempts at targeting other known PDAC genes/pathways including P13K/Akt/mTOR have been clinically disappointing[30,31].

Her2/neu amplification is well characterised in a number of malignancies which have shown response when used as druggable targets in breast and gastric malignancies for example[32-34]. In PDAC, Her2/neu is amplified in up to 45% of cases[35], particularly in the advanced setting. Unfortunately, while initial pre-clinical studies indicated a potential role for Her-2 directed therapy specifically trastuzumab as a monotherapy or in combination with gemcitabine[36] it did not prove clinically advantageous. Similarly, investigations combining trastuzumab with capecitabine were disa­ppointing[35]. In a related fashion EGFR which is known to co-express with Her2 has also been investigated with statistical gains noted for the EGFR inhibitor erlotinib in conjunction with gemcitabine, however this was only in the order of two weeks survival benefit and little mention of the related quality of life impact secondary to treatment was reported[34].

The JAK/STAT pathway has also been implicated as a regulator in the development of PDAC *via* its role in activating signalling cascades and gene transcription. Stimulated by oxidative stress, this pathway ultimately induces the production of inflammatory cytokines as well as cell proliferation, malignant transformation and inhibition of apoptosis in the pancreas[37,38]. This association with inflammation has prompted trials of the JAK-1/2 inhibitor ruxolotinib in combination with capecitabine for patients with metastatic PDAC after failure of first line therapy if patients expressed ele­vated inflammatory markers as assessed by C-reactive protein[39]. In this phase Ⅱ randomised trial, 127 patients were treated with either ruxolotinib and capecitabine or capecitabine and placebo. However, interim analyses failed to demonstrate sufficient efficacy with ruxolotinib combination therapy and further investigations of this drug has been suspended[40].

Together this information provides us with a raft of data for potential therapeutic targets, whether *via* drivers, alterations in signalling pathways or susceptibility genes. For example in the case of *PALB2* mutation-associated disease there are multiple reports suggesting improved outcomes in response to platinum based therapy or mitomycin C when compared with gemcitabine[27,28] suggesting that a more comprehensive understanding of the genetic complexity of PDAC will assist in treatment decisions. Thus understanding the similarities and differences between the “poor” and “exceptional responders” may provide biomarkers to identify patients who might benefit from these treatments and improve outcomes[41].

***Stromal microenvironment and drug delivery***

PDAC is characterised by the surrounding cells, specifically activated fibroblasts, myofibroblasts and pancreatic stellate cells which contribute to the composition of the surrounding matrix, elements such as hyaluronan, growth factors (*e.g.*, TGF-) and secreted protein acidic rich in cysteine (SPARC)[42,43]. These result in a unique stromal microenvironment which may not only promote tumour initiation and progression but also create a barrier to drug delivery thus rendering PDAC relatively chemoresistant. Consequently, much effort has focused on ways to deplete or manipulate the stromal microenvironment and improve therapeutic outcomes.

SPARC is a glycoprotein believed to be involved in cancer development *via* its modulation of cell proliferation, progression, angiogenesis, migration, metastasis and apoptosis[44]. It’s normal role in cellular functions is thought to be multifactorial with effects on cell dispersion and chemosensitization as well as induction of apoptosis but also has antiangiogenic properties[45-48]. While the intricacies of SPARC and cancer are yet to be fully elucidated its potential to increase the invasive capacity of malignant cells and possible association with poor prognosis is recognised[7]. Of interest, SPARC methylation leading to pathogenesis correlates with both tobacco smoking and alcohol consumption, which are known associated modifiable risk factors for PDAC[49].

Interestingly gemcitabine has been reported to alter SPARC expression in a dose dependent manner in cell lines[50] however, there is also evidence to suggest that SPARC overexpression enhances PDAC cell chemosensitivity to gemcitabine[51]. In fact, SPARC may actually assist in the delivery of nab-paclitaxel to the tumour due to its affinity for albumin[52]. Nab-paclitaxel is formulated with human albumin at concentrations that closely resemble physiological albumin levels. This feature seems to enable nab-paclitaxel to penetrate the stromal environment and reach the tumour more efficiently[53]. Despite these studies however the role for SPARC within the stromal micro-environment and its implications on therapy remain controversial. For example data presented by Hidalgo *et al*[54] reported no clear association between SPARC levels and treatment efficacy with combination therapy using gemcitabine and nab-paclitaxel or gemcitabine alone in metastatic PDAC.

Growth factors such as TGF- are produced by cells within the stromal microenvironment. Interestingly TGF- levels correlate with tumour metastases and progression as well as poorer patient outcomes[55]. The specific role for TGF- in this case is not clear but may involve regulation of cell cycle arrest, apoptosis, immune response and/or wound healing. Activated TGF- signalling is mediated *via* SMADs, a known driver of PDAC, which is also reported to correlate with worse prognosis or disseminated disease[56,57].

Intriguingly TGF- has been reported as a tumour suppressor in early stages of malignancy but a pro­moter in established disease[58] further emphasising the complex nature of PDAC.

Despite the steady accumulation of knowledge from studies into genetic and molecular pathways, complex stromal characteristics and better drug delivery remains an ongoing issue further prompting investigations of novel approaches such as nucleolar stress pathways and ribosome biogenesis.

**RIBOSOME BIOGENESIS**

Ribosome biogenesis is a highly coordinated process which takes place within a dynamic compartment of the nucleus termed the nucleolus. Long before the functional role of the nucleolus was established, Pianese[59] noted that malignant cells were often characterised by enlarged, abnormal nucleoli. Since then, numerous studies have correlated morphological changes of the nucleoli with malignant disease.

It is now clear that the morphological changes affecting nucleoli in malignant cells reflects hype­ractivated transcription of ribosomal RNA (rRNA) genes by RNA polymerase I (Pol I) yielding the 47S rRNA precursor which is rapidly processed into mature 18, 5.8 and 28S rRNAs[60] while the 5S rRNA precursor is transcribed by RNA polymerase III (Pol III). These RNA’s together with ribosomal proteins (RPs) transcribed by RNA polymerase II (Pol II) are essential for ribosome assembly, central to the synthesis of cellular proteins[61,62]. Co-ordination of all three polymerases is necessary for the development of a functional mammalian ribosome (80S) composed of a small (40S) and large (60S) subunit[62,63].

During active cell division and proliferation ribosome biogenesis increases so as to maintain elevated cellular demands. It is not surprising that this process is a major consumer of cellular energy requiring vigilant regulation[64,65]. However, ribosome biogenesis is not only regulated by RNA polymerases, as depicted in figure 2, but also involves a number of tumour su­ppressors or oncoproteins[66] including c-myc, con­sidered a “master regulator” of protein synthesis[67]. Conversely p53 has an important place in inhibiting ribosome biogenesis[68] and promoting cell cycle arrest, senescence or apoptosis.

In normal cells, levels of p53 protein are typically low[69], kept in check by various mechanisms including MDM2, an E3 ubiquitin ligase[62], as shown in figure 3, allowing cell division and proliferation.

***Nucleolar stress***

Disruptions in ribosome synthesis lead to nucleolar stress which in turn results in nucleolar disruption and the release of RP’s *i.e.*, RPL5 and RPL1. These RPs are able to bind to MDM2 and interrupt its interaction with p53[60,70]. Similarly, the product of the *CDKN2A* gene, p14ARF, is also able to exert effects on MDM2[60]. Ultimately, these pathways lead to stabilisation of p53 allowing it to act on various transcriptional targets and induce apoptosis, senescence and/or cell cycle arrest depicted in figure 3. Importantly one of the transcriptional targets for p53 is the *MDM2* gene setting up an autologous feedback loop to maintain genomic and cellular homeostasis[62,71,72] keeping the process of cell death and proliferation in check.

When the above mentioned homeostatic mechanisms are impaired however, ribosome biogenesis becomes dysregulated and the balance between cell growth and arrest is compromised leading to unchecked cellular proliferation and malignant transformation. Activation of the nucleolar stress pathway may therefore provide a therapeutic strategy against aggressive malignancies.

While p53 dependent nucleolar stress is well under­stood, more recent research by Quin *et al*[73] has identified p53 independent mechanisms involving activation of ATM/ATR signalling which occurs without the induction of global DNA damage[73]. Evidently, ribosome biogenesis is a highly complex and finely balanced process involving multiple key effectors. Disruption of any of these components can mediate the nucleolar stress pathway and have detrimental effects on cell growth and proliferation. It is this feature along with the potential for reduced global DNA damage that makes targeting dysregulated ribosome biogenesis an attractive therapeutic option.

***Therapeutic targeting of dysregulated ribosome biogenesis***

Many regulators of ribosome biogenesis are also involved in malignant diseases including PDAC. For example, activation of RAS or PI3K pathways known to be involved in PDAC are critical for coordinating protein synthesising capacity (ribosome number) required for maintaining cellular growth and proliferation. Other genes and their products, such as c-myc and p53 also have significant roles in ribosome biogenesis and malignancy when overexpressed or mutated. Attempts at targeting these various genes and pathways have been disappointing in the realm of PDAC but tackling dysregulated ribosome biogenesis have shown more promise.

Pivotal studies conducted by Bywater *et al*[12], provided proof of principal for the benefits of targeting dysregulated ribosome biogenesis as demonstrated by CX5461, a novel small molecule Pol I inhibitor, investigated in the Eµ-Myc mouse model of Burkitt’s lymphoma. In this model, cells are highly proliferative and have elevated RNA levels, including rRNA, which correlates with accelerated Pol I transcription and cell growth. Pharmacological inhibition of Pol I transcription with CX5461 induced nucleolar stress leading to an increase in p53 levels resulting in apoptotic cell death of malignant cells. Most compelling however, CX5461 did not kill non-malignant B cells representing a valuable therapeutic option in the management of susceptible malignancies[12].

In addition to the Bywater study, Devlin *et al*[13] demonstrated that combination therapy with CX5461 and AKT-mTOR inhibitors worked synergistically leading to significantly extend survival in lymphoma bearing mice[61]. Importantly, the acute lymphoblastic leukaemia cells were more sensitive to rRNA synthesis inhibition compared with normal bone marrow cells[74] further emphasising the potential reduction in global toxicity underlying Pol I inhibition. Similarly, reduced global toxicity was noted by Hannan *et al*[75] review of lymphoma models in response to Pol I inhibition.

To date, the majority of studies of Pol I inhibitors have focussed on haematological malignancies, and as such CX-5461 is the subject of a phase I clinical trial at the Peter MacCallum Cancer Centre for treatment of patients with advanced haematological malignan­cies (Australian New Zealand Clinical Trials Registry 12613001061729). However, such studies are now being extended to various solid organ malignancies including metastatic/recurrent/locally advanced/unresectable triple negative breast cancer which is currently recruiting through the Canadian Cancer Trials group (NCT02719977). This is a logical extension as there is preliminary data for efficacy of CX-5461 in solid tumours, with pre-clinical data suggesting a role in prostate cancer[14]. Similarly, studies in PDAC cell lines have also shown promise[76], as documented by Drygin *et al*[76] using MIA PaCa-2 cell lines. This study demonstrated a process of both autophagy and cell senescence in response to Pol I inhibition. In xenograft mouse models of human MIA PaCa-2 cells, treatment with CX5461 demonstrated significant tumour growth inhibition deemed at least comparable to xenograft models treated with gemcitabine and again shows promise.

**CONCLUSION**

Given the promising result for Pol I inhibition in the treatment of haematological malignancies and emer­ging evidence of efficacy in various solid organ cancers the potential for targeting dysregulated ribosome biogenesis and manipulating the nucleolar stress pathway is tantalising, particularly for advanced PDAC given the slow progress in its treatment options so far. Whether this will be in the setting of combinations with already established chemotherapeutics or in more novel ways remains to be seen but the results from other malignancies are encouraging. Of particular note is the potential for less toxicity to healthy cells which thus far has been a major limitation in improving treatment outcomes for PDAC.

**REFERENCES**

1 GLOBOCAN 2012: Estimated Cancer Incidence, Mortality and Prevalence Worldwide in 2012. 2016 Available from: URL: http://globocan.iarc.fr/Pages/fact\_sheets\_cancer.aspx

2 **Siegel R**, Ma J, Zou Z, Jemal A. Cancer statistics, 2014. *CA Cancer J Clin* 2014; **64**: 9-29 [PMID: 24399786 DOI: 10.3322/caac.21208]

3 **American Cancer Society**. 2016. Available from: URL: http://www.cancer.org/cancer/pancreaticcancer/detailedguide/pancreatic-cancer-key-statistics

4 **Lockhart AC**, Rothenberg ML, Berlin JD. Treatment for pancreatic cancer: current therapy and continued progress. *Gastroenterology* 2005; **128**: 1642-1654 [PMID: 15887156 DOI: 10.1053/j.gastro.2005.03.039]

5 **Conroy T**, Desseigne F, Ychou M, Bouché O, Guimbaud R, Bécouarn Y, Adenis A, Raoul JL, Gourgou-Bourgade S, de la Fouchardière C, Bennouna J, Bachet JB, Khemissa-Akouz F, Péré-Vergé D, Delbaldo C, Assenat E, Chauffert B, Michel P, Montoto-Grillot C, Ducreux M. FOLFIRINOX versus gemcitabine for metastatic pancreatic cancer. *N Engl J Med* 2011; **364**: 1817-1825 [PMID: 21561347 DOI: 10.1056/NEJMoa1011923]

6 **Gourgou-Bourgade S**, Bascoul-Mollevi C, Desseigne F, Ychou M, Bouché O, Guimbaud R, Bécouarn Y, Adenis A, Raoul JL, Boige V, Bérille J, Conroy T. Impact of FOLFIRINOX compared with gemcitabine on quality of life in patients with metastatic pancreatic cancer: results from the PRODIGE 4/ACCORD 11 randomized trial. *J Clin Oncol* 2013; **31**: 23-29 [PMID: 23213101 DOI: 10.1200/JCO.2012.44.4869]

7 **Von Hoff DD**, Ervin T, Arena FP, Chiorean EG, Infante J, Moore M, Seay T, Tjulandin SA, Ma WW, Saleh MN, Harris M, Reni M, Dowden S, Laheru D, Bahary N, Ramanathan RK, Tabernero J, Hidalgo M, Goldstein D, Van Cutsem E, Wei X, Iglesias J, Renschler MF. Increased survival in pancreatic cancer with nab-paclitaxel plus gemcitabine. *N Engl J Med* 2013; **369**: 1691-1703 [PMID: 24131140 DOI: 10.1056/NEJMoa1304369]

8 **Ducreux M**, Cuhna AS, Caramella C, Hollebecque A, Burtin P, Goéré D, Seufferlein T, Haustermans K, Van Laethem JL, Conroy T, Arnold D. Cancer of the pancreas: ESMO Clinical Practice Guidelines for diagnosis, treatment and follow-up. *Ann Oncol* 2015; **26** Suppl 5: v56-v68 [PMID: 26314780 DOI: 10.1093/annonc/mdv295]

9 **Wang-Gillam A**, Li CP, Bodoky G, Dean A, Shan YS, Jameson G, Macarulla T, Lee KH, Cunningham D, Blanc JF, Hubner RA, Chiu CF, Schwartsmann G, Siveke JT, Braiteh F, Moyo V, Belanger B, Dhindsa N, Bayever E, Von Hoff DD, Chen LT. Nanoliposomal irinotecan with fluorouracil and folinic acid in metastatic pancreatic cancer after previous gemcitabine-based therapy (NAPOLI-1): a global, randomised, open-label, phase 3 trial. *Lancet* 2016; **387**: 545-557 [PMID: 26615328 DOI: 10.1016/SO140-6736(15)00986-1]

10 **Kalra AV**, Kim J, Klinz SG, Paz N, Cain J, Drummond DC, Nielsen UB, Fitzgerald JB. Preclinical activity of nanoliposomal irinotecan is governed by tumor deposition and intratumor prodrug conversion. *Cancer Res* 2014; **74**: 7003-7013 [PMID: 25273092 DOI: 10.1158/0008-5472.CAN-14-0572]

11 **Roy AC**, Park SR, Cunningham D, Kang YK, Chao Y, Chen LT, Rees C, Lim HY, Tabernero J, Ramos FJ, Kujundzic M, Cardic MB, Yeh CG, de Gramont A. A randomized phase II study of PEP02 (MM-398), irinotecan or docetaxel as a second-line therapy in patients with locally advanced or metastatic gastric or gastro-oesophageal junction adenocarcinoma. *Ann Oncol* 2013; **24**: 1567-1573 [PMID: 23406728 DOI: 10.1093/annonc/mdt002]

12 **Bywater MJ**, Poortinga G, Sanij E, Hein N, Peck A, Cullinane C, Wall M, Cluse L, Drygin D, Anderes K, Huser N, Proffitt C, Bliesath J, Haddach M, Schwaebe MK, Ryckman DM, Rice WG, Schmitt C, Lowe SW, Johnstone RW, Pearson RB, McArthur GA, Hannan RD. Inhibition of RNA polymerase I as a therapeutic strategy to promote cancer-specific activation of p53. *Cancer Cell* 2012; **22**: 51-65 [PMID: 22789538 DOI: 10.1016/j.ccr.2012.05.019]

13 **Devlin JR**, Hannan KM, Hein N, Cullinane C, Kusnadi E, Ng PY, George AJ, Shortt J, Bywater MJ, Poortinga G, Sanij E, Kang J, Drygin D, O’Brien S, Johnstone RW, McArthur GA, Hannan RD, Pearson RB. Combination Therapy Targeting Ribosome Biogenesis and mRNA Translation Synergistically Extends Survival in MYC-Driven Lymphoma. *Cancer Discov* 2016; **6**: 59-70 [PMID: 26490423 DOI: 10.1158/2159-8290.CD-14-0673]

14 **Rebello RJ**, Kusnadi E, Cameron DP, Pearson HB, Lesmana A, Devlin JR, Drygin D, Clark AK, Porter L, Pedersen J, Sandhu S, Risbridger GP, Pearson RB, Hannan RD, Furic L. The Dual Inhibition of RNA Pol I Transcription and PIM Kinase as a New Therapeutic Approach to Treat Advanced Prostate Cancer. *Clin Cancer Res* 2016; **22**: 5539-5552 [PMID: 27486174 DOI: 10.1158/1078-0432.CCR-16-1024]

15 **Institute NC**. Surveillance, Epidemiology and End Results Program. SEER Stat Fact Sheets: Pnacreas Cancer. cited 2016-10-24 Available from: URL: https://seer.cancer.gov/

16 **Hidalgo M**, Cascinu S, Kleeff J, Labianca R, Löhr JM, Neoptolemos J, Real FX, Van Laethem JL, Heinemann V. Addressing the challenges of pancreatic cancer: future directions for improving outcomes. *Pancreatology* 2015; **15**: 8-18 [PMID: 25547205 DOI: 10.1016/j.pan.2014.10.001]

17 **Gerritsen A**, Jacobs M, Henselmans I, van Hattum J, Efficace F, Creemers GJ, de Hingh IH, Koopman M, Molenaar IQ, Wilmink HW, Busch OR, Besselink MG, van Laarhoven HW. Developing a core set of patient-reported outcomes in pancreatic cancer: A Delphi survey. *Eur J Cancer* 2016; **57**: 68-77 [PMID: 26886181 DOI: 10.1016/j.ejca.2016.01.001]

18 **Zhen DB**, Rabe KG, Gallinger S, Syngal S, Schwartz AG, Goggins MG, Hruban RH, Cote ML, McWilliams RR, Roberts NJ, Cannon-Albright LA, Li D, Moyes K, Wenstrup RJ, Hartman AR, Seminara D, Klein AP, Petersen GM. BRCA1, BRCA2, PALB2, and CDKN2A mutations in familial pancreatic cancer: a PACGENE study. *Genet Med* 2015; **17**: 569-577 [PMID: 25356972 DOI: 10.1038/gim.2014.153]

19 **Humphris JL**, Johns AL, Simpson SH, Cowley MJ, Pajic M, Chang DK, Nagrial AM, Chin VT, Chantrill LA, Pinese M, Mead RS, Gill AJ, Samra JS, Kench JG, Musgrove EA, Tucker KM, Spigelman AD, Waddell N, Grimmond SM, Biankin AV. Clinical and pathologic features of familial pancreatic cancer. *Cancer* 2014; **120**: 3669-3675 [PMID: 25313458 DOI: 10.1002/cncr.28863]

20 **Brand RE**, Lerch MM, Rubinstein WS, Neoptolemos JP, Whitcomb DC, Hruban RH, Brentnall TA, Lynch HT, Canto MI. Advances in counselling and surveillance of patients at risk for pancreatic cancer. *Gut* 2007; **56**: 1460-1469 [PMID: 17872573 DOI: 10.1136/gut.2006.108456]

21 **Waddell N**, Pajic M, Patch AM, Chang DK, Kassahn KS, Bailey P, Johns AL, Miller D, Nones K, Quek K, Quinn MC, Robertson AJ, Fadlullah MZ, Bruxner TJ, Christ AN, Harliwong I, Idrisoglu S, Manning S, Nourse C, Nourbakhsh E, Wani S, Wilson PJ, Markham E, Cloonan N, Anderson MJ, Fink JL, Holmes O, Kazakoff SH, Leonard C, Newell F, Poudel B, Song S, Taylor D, Waddell N, Wood S, Xu Q, Wu J, Pinese M, Cowley MJ, Lee HC, Jones MD, Nagrial AM, Humphris J, Chantrill LA, Chin V, Steinmann AM, Mawson A, Humphrey ES, Colvin EK, Chou A, Scarlett CJ, Pinho AV, Giry-Laterriere M, Rooman I, Samra JS, Kench JG, Pettitt JA, Merrett ND, Toon C, Epari K, Nguyen NQ, Barbour A, Zeps N, Jamieson NB, Graham JS, Niclou SP, Bjerkvig R, Grützmann R, Aust D, Hruban RH, Maitra A, Iacobuzio-Donahue CA, Wolfgang CL, Morgan RA, Lawlor RT, Corbo V, Bassi C, Falconi M, Zamboni G, Tortora G, Tempero MA, Gill AJ, Eshleman JR, Pilarsky C, Scarpa A, Musgrove EA, Pearson JV, Biankin AV, Grimmond SM. Whole genomes redefine the mutational landscape of pancreatic cancer. *Nature* 2015; **518**: 495-501 [PMID: 25719666 DOI: 10.1038/nature14169]

22 **Bailey P**, Chang DK, Nones K, Johns AL, Patch AM, Gingras MC, Miller DK, Christ AN, Bruxner TJ, Quinn MC, Nourse C, Murtaugh LC, Harliwong I, Idrisoglu S, Manning S, Nourbakhsh E, Wani S, Fink L, Holmes O, Chin V, Anderson MJ, Kazakoff S, Leonard C, Newell F, Waddell N, Wood S, Xu Q, Wilson PJ, Cloonan N, Kassahn KS, Taylor D, Quek K, Robertson A, Pantano L, Mincarelli L, Sanchez LN, Evers L, Wu J, Pinese M, Cowley MJ, Jones MD, Colvin EK, Nagrial AM, Humphrey ES, Chantrill LA, Mawson A, Humphris J, Chou A, Pajic M, Scarlett CJ, Pinho AV, Giry-Laterriere M, Rooman I, Samra JS, Kench JG, Lovell JA, Merrett ND, Toon CW, Epari K, Nguyen NQ, Barbour A, Zeps N, Moran-Jones K, Jamieson NB, Graham JS, Duthie F, Oien K, Hair J, Grützmann R, Maitra A, Iacobuzio-Donahue CA, Wolfgang CL, Morgan RA, Lawlor RT, Corbo V, Bassi C, Rusev B, Capelli P, Salvia R, Tortora G, Mukhopadhyay D, Petersen GM, Munzy DM, Fisher WE, Karim SA, Eshleman JR, Hruban RH, Pilarsky C, Morton JP, Sansom OJ, Scarpa A, Musgrove EA, Bailey UM, Hofmann O, Sutherland RL, Wheeler DA, Gill AJ, Gibbs RA, Pearson JV, Waddell N, Biankin AV, Grimmond SM. Genomic analyses identify molecular subtypes of pancreatic cancer. *Nature* 2016; **531**: 47-52 [PMID: 26909576 DOI: 10.1038/nature16965]

23 **Cid-Arregui A**, Juarez V. Perspectives in the treatment of pancreatic adenocarcinoma. *World J Gastroenterol* 2015; **21**: 9297-9316 [PMID: 26309356 DOI: 10.3748/wjg.v21.i31.9297]

24 **Chiche J**, Brahimi-Horn MC, Pouysségur J. Tumour hypoxia induces a metabolic shift causing acidosis: a common feature in cancer. *J Cell Mol Med* 2010; **14**: 771-794 [PMID: 20015196 DOI: 10.1111/j.1582-4934.2009.00994.x]

25 **Tutt A**, Gabriel A, Bertwistle D, Connor F, Paterson H, Peacock J, Ross G, Ashworth A. Absence of Brca2 causes genome instability by chromosome breakage and loss associated with centrosome amplification. *Curr Biol* 1999; **9**: 1107-1110 [PMID: 10531007]

26 **Jones S**, Hruban RH, Kamiyama M, Borges M, Zhang X, Parsons DW, Lin JC, Palmisano E, Brune K, Jaffee EM, Iacobuzio-Donahue CA, Maitra A, Parmigiani G, Kern SE, Velculescu VE, Kinzler KW, Vogelstein B, Eshleman JR, Goggins M, Klein AP. Exomic sequencing identifies PALB2 as a pancreatic cancer susceptibility gene. *Science* 2009; **324**: 217 [PMID: 19264984 DOI: 10.1126/science.1171202]

27 **Chan D**, Clarke S, Gill AJ, Chantrill L, Samra J, Li BT, Barnes T, Nahar K, Pavlakis N. Pathogenic PALB2 mutation in metastatic pancreatic adenocarcinoma and neuroendocrine tumour: A case report. *Mol Clin Oncol* 2015; **3**: 817-819 [PMID: 26171187 DOI: 10.3892/mco.2015.533]

28 **Villarroel MC**, Rajeshkumar NV, Garrido-Laguna I, De Jesus-Acosta A, Jones S, Maitra A, Hruban RH, Eshleman JR, Klein A, Laheru D, Donehower R, Hidalgo M. Personalizing cancer treatment in the age of global genomic analyses: PALB2 gene mutations and the response to DNA damaging agents in pancreatic cancer. *Mol Cancer Ther* 2011; **10**: 3-8 [PMID: 21135251 DOI: 10.1158/1535-7163.MCT-10-0893]

29 **Bodoky G**, Timcheva C, Spigel DR, La Stella PJ, Ciuleanu TE, Pover G, Tebbutt NC. A phase II open-label randomized study to assess the efficacy and safety of selumetinib (AZD6244 [ARRY-142886]) versus capecitabine in patients with advanced or metastatic pancreatic cancer who have failed first-line gemcitabine therapy. *Invest New Drugs* 2012; **30**: 1216-1223 [PMID: 21594619 DOI: 10.1007/s10637-011-9687-4]

30 **Wolpin BM**, Hezel AF, Abrams T, Blaszkowsky LS, Meyerhardt JA, Chan JA, Enzinger PC, Allen B, Clark JW, Ryan DP, Fuchs CS. Oral mTOR inhibitor everolimus in patients with gemcitabine-refractory metastatic pancreatic cancer. *J Clin Oncol* 2009; **27**: 193-198 [PMID: 19047305 DOI: 10.1200/JCO.2008.18.9514]

31 **Javle MM**, Shroff RT, Xiong H, Varadhachary GA, Fogelman D, Reddy SA, Davis D, Zhang Y, Wolff RA, Abbruzzese JL. Inhibition of the mammalian target of rapamycin (mTOR) in advanced pancreatic cancer: results of two phase II studies. *BMC Cancer* 2010; **10**: 368 [PMID: 20630061 DOI: 10.1186/1471-2407-10-368]

32 **Bang YJ**, Van Cutsem E, Feyereislova A, Chung HC, Shen L, Sawaki A, Lordick F, Ohtsu A, Omuro Y, Satoh T, Aprile G, Kulikov E, Hill J, Lehle M, Rüschoff J, Kang YK. Trastuzumab in combination with chemotherapy versus chemotherapy alone for treatment of HER2-positive advanced gastric or gastro-oesophageal junction cancer (ToGA): a phase 3, open-label, randomised controlled trial. *Lancet* 2010; **376**: 687-697 [PMID: 20728210 DOI: 10.1016/S0140-6736(10)61121-X]

33 **Slamon DJ**, Clark GM, Wong SG, Levin WJ, Ullrich A, McGuire WL. Human breast cancer: correlation of relapse and survival with amplification of the HER-2/neu oncogene. *Science* 1987; **235**: 177-182 [PMID: 3798106]

34 **Kelley RK**, Ko AH. Erlotinib in the treatment of advanced pancreatic cancer. *Biologics* 2008; **2**: 83-95 [PMID: 19707431]

35 **Harder J**, Ihorst G, Heinemann V, Hofheinz R, Moehler M, Buechler P, Kloeppel G, Röcken C, Bitzer M, Boeck S, Endlicher E, Reinacher-Schick A, Schmoor C, Geissler M. Multicentre phase II trial of trastuzumab and capecitabine in patients with HER2 overexpressing metastatic pancreatic cancer. *Br J Cancer* 2012; **106**: 1033-1038 [PMID: 22374460 DOI: 10.1038/bjc.2012.18]

36 **Kimura K**, Sawada T, Komatsu M, Inoue M, Muguruma K, Nishihara T, Yamashita Y, Yamada N, Ohira M, Hirakawa K. Antitumor effect of trastuzumab for pancreatic cancer with high HER-2 expression and enhancement of effect by combined therapy with gemcitabine. *Clin Cancer Res* 2006; **12**: 4925-4932 [PMID: 16914581 DOI: 10.1158/1078-0432.CCR-06-0544]

37 **Yu JH**, Kim H. Role of janus kinase/signal transducers and activators of transcription in the pathogenesis of pancreatitis and pancreatic cancer. *Gut Liver* 2012; **6**: 417-422 [PMID: 23170143 DOI: 10.5009/gnl.2012.6.4.417]

38 **Yu JH**, Kim H. Oxidative stress and cytokines in the pathogenesis of pancreatic cancer. *J Cancer Prev* 2014; **19**: 97-102 [PMID: 25337577 DOI: 10.15430/JCP.2014.19.2.97]

39 **Hurwitz HI**, Uppal N, Wagner SA, Bendell JC, Beck JT, Wade SM, Nemunaitis JJ, Stella PJ, Pipas JM, Wainberg ZA, Manges R, Garrett WM, Hunter DS, Clark J, Leopold L, Sandor V, Levy RS. Randomized, Double-Blind, Phase II Study of Ruxolitinib or Placebo in Combination With Capecitabine in Patients With Metastatic Pancreatic Cancer for Whom Therapy With Gemcitabine Has Failed. *J Clin Oncol* 2015; **33**: 4039-4047 [PMID: 26351344 DOI: 10.1200/JCO.2015.61.4578]

40 **Wire B**. Incyte Announces Decision to Discontinue JANUS Studies of Ruxolitinib plus Capecitabine in Patients with Advanced or Metastatic Pancreatic Cancer. Internet Press Release, 2016. Available from: URL: http://www.businesswire.com/news/home/20160211005321/en/Incyte-Announces-Decision-Discontinue-JANUS-Studies-Ruxolitinib

41 **Chang DK**, Grimmond SM, Evans TR, Biankin AV. Mining the genomes of exceptional responders. *Nat Rev Cancer* 2014; **14**: 291-292 [PMID: 25688402 DOI: 10.1038/nrc3723]

42 **Neesse A**, Algül H, Tuveson DA, Gress TM. Stromal biology and therapy in pancreatic cancer: a changing paradigm. *Gut* 2015; **64**: 1476-1484 [PMID: 25994217 DOI: 10.1136/gutjnl-2015-309304]

43 **Matsuoka T**, Yashiro M. Molecular targets for the treatment of pancreatic cancer: Clinical and experimental studies. *World J Gastroenterol* 2016; **22**: 776-789 [PMID: 26811624 DOI: 10.3748/wjg.v22.i2.776]

44 **Nagaraju GP**, Dontula R, El-Rayes BF, Lakka SS. Molecular mechanisms underlying the divergent roles of SPARC in human carcinogenesis. *Carcinogenesis* 2014; **35**: 967-973 [PMID: 24675529 DOI: 10.1093/carcin/bgu072]

45 **Lane TF**, Sage EH. Functional mapping of SPARC: peptides from two distinct Ca+(+)-binding sites modulate cell shape. *J Cell Biol* 1990; **111**: 3065-3076 [PMID: 2269665]

46 **Rahman M**, Chan AP, Tang M, Tai IT. A peptide of SPARC interferes with the interaction between caspase8 and Bcl2 to resensitize chemoresistant tumors and enhance their regression in vivo. *PLoS One* 2011; **6**: e26390 [PMID: 22069448 DOI: 10.1371/journal.pone.0026390]

47 **Funk SE**, Sage EH. The Ca2(+)-binding glycoprotein SPARC modulates cell cycle progression in bovine aortic endothelial cells. *Proc Natl Acad Sci USA* 1991; **88**: 2648-2652 [PMID: 2011576]

48 **Chlenski A**, Liu S, Baker LJ, Yang Q, Tian Y, Salwen HR, Cohn SL. Neuroblastoma angiogenesis is inhibited with a folded synthetic molecule corresponding to the epidermal growth factor-like module of the follistatin domain of SPARC. *Cancer Res* 2004; **64**: 7420-7425 [PMID: 15492265 DOI: 10.1158/0008-5472.CAN-04-2141]

49 **Gao J**, Song J, Huang H, Li Z, Du Y, Cao J, Li M, Lv S, Lin H, Gong Y. Methylation of the SPARC gene promoter and its clinical implication in pancreatic cancer. *J Exp Clin Cancer Res* 2010; **29**: 28 [PMID: 20338068 DOI: 10.1186/1756-9966-29-28]

50 **Toshimitsu H**, Iizuka N, Yamamoto K, Kawauchi S, Oga A, Furuya T, Oka M, Sasaki K. Molecular features linked to the growth-inhibitory effects of gemcitabine on human pancreatic cancer cells. *Oncol Rep* 2006; **16**: 1285-1291 [PMID: 17089051]

51 **Zhang J**, Jiang H, Mao Z, Wang X, Fan X, Liu Y, Wang Y. [Mechanism of SPARC-enhanced chemosensitivity of pancreatic cancer cells to gemcitabine]. *Zhonghua Zhongliu Zazhi* 2014; **36**: 335-340 [PMID: 25030587]

52 **Motamed K**. SPARC (osteonectin/BM-40). *Int J Biochem Cell Biol* 1999; **31**: 1363-1366 [PMID: 10641790]

53 **Vaz J**, Ansari D, Sasor A, Andersson R. SPARC: A Potential Prognostic and Therapeutic Target in Pancreatic Cancer. *Pancreas* 2015; **44**: 1024-1035 [PMID: 26335014 DOI: 10.1097/MPA.0000000000000409]

54 **Hidalgo M**, Plaza C, Musteanu M, Illei P, Brachmann CB, Heise C, Pierce D, Lopez-Casas PP, Menendez C, Tabernero J, Romano A, Wei X, Lopez-Rios F, Von Hoff DD. SPARC Expression Did Not Predict Efficacy of nab-Paclitaxel plus Gemcitabine or Gemcitabine Alone for Metastatic Pancreatic Cancer in an Exploratory Analysis of the Phase III MPACT Trial. *Clin Cancer Res* 2015; **21**: 4811-4818 [PMID: 26169969 DOI: 10.1158/1078-0432.CCR-14-3222]

55 **Goel G**, Sun W. Novel approaches in the management of pancreatic ductal adenocarcinoma: potential promises for the future. *J Hematol Oncol* 2015; **8**: 44 [PMID: 25935754 DOI: 10.1186/s13045-015-0141-5]

56 **Bardeesy N**, Cheng KH, Berger JH, Chu GC, Pahler J, Olson P, Hezel AF, Horner J, Lauwers GY, Hanahan D, DePinho RA. Smad4 is dispensable for normal pancreas development yet critical in progression and tumor biology of pancreas cancer. *Genes Dev* 2006; **20**: 3130-3146 [PMID: 17114584 DOI: 10.1101/gad.1478706]

57 **Iacobuzio-Donahue CA**, Fu B, Yachida S, Luo M, Abe H, Henderson CM, Vilardell F, Wang Z, Keller JW, Banerjee P, Herman JM, Cameron JL, Yeo CJ, Halushka MK, Eshleman JR, Raben M, Klein AP, Hruban RH, Hidalgo M, Laheru D. DPC4 gene status of the primary carcinoma correlates with patterns of failure in patients with pancreatic cancer. *J Clin Oncol* 2009; **27**: 1806-1813 [PMID: 19273710 DOI: 10.1200/JCO.2008.17.7188]

58 **Tian M**, Schiemann WP. The TGF-beta paradox in human cancer: an update. *Future Oncol* 2009; **5**: 259-271 [PMID: 19284383 DOI: 10.2217/14796694.5.2.259]

59 **Pianese G**. Beitrag zur Histologie und Aetiologie der Carcinoma. Histologische und experimentelle Untersuchungen. *Beitr Pathol Anat Allgem Pathol* 1896; **142**: 1-193

60 **Quin JE**, Devlin JR, Cameron D, Hannan KM, Pearson RB, Hannan RD. Targeting the nucleolus for cancer intervention. *Biochim Biophys Acta* 2014; **1842**: 802-816 [PMID: 24389329 DOI: 10.1016/j.bbadis.2013.12.009]

61 **Mayer C**, Grummt I. Ribosome biogenesis and cell growth: mTOR coordinates transcription by all three classes of nuclear RNA polymerases. *Oncogene* 2006; **25**: 6384-6391 [PMID: 17041624 DOI: 10.1038/sj.onc.1209883]

62 **Brighenti E**, Treré D, Derenzini M. Targeted cancer therapy with ribosome biogenesis inhibitors: a real possibility? *Oncotarget* 2015; **6**: 38617-38627 [PMID: 26415219 DOI: 10.18632/oncotarget.5775]

63 **Pelava A**, Schneider C, Watkins NJ. The importance of ribosome production, and the 5S RNP-MDM2 pathway, in health and disease. *Biochem Soc Trans* 2016; **44**: 1086-1090 [PMID: 27528756 DOI: 10.1042/BST20160106]

64 **Dang CV**. MYC, metabolism, cell growth, and tumorigenesis. *Cold Spring Harb Perspect Med* 2013; **3**: pii: a014217 [PMID: 23906881 DOI: 10.1101/cshperspect.a014217]

65 **Orsolic I**, Jurada D, Pullen N, Oren M, Eliopoulos AG, Volarevic S. The relationship between the nucleolus and cancer: Current evidence and emerging paradigms. *Semin Cancer Biol* 2016; **37-38**: 36-50 [PMID: 26721423 DOI: 10.1016/j.semcancer.2015.12.004]

66 **Dai MS**, Zeng SX, Jin Y, Sun XX, David L, Lu H. Ribosomal protein L23 activates p53 by inhibiting MDM2 function in response to ribosomal perturbation but not to translation inhibition. *Mol Cell Biol* 2004; **24**: 7654-7668 [PMID: 15314173 DOI: 10.1128/MCB.24.17.7654-7668.2004]

67 **Hsieh AL**, Walton ZE, Altman BJ, Stine ZE, Dang CV. MYC and metabolism on the path to cancer. *Semin Cell Dev Biol* 2015; **43**: 11-21 [PMID: 26277543 DOI: 10.1016/j.semcdb.2015.08.003]

68 **Pestov DG**, Strezoska Z, Lau LF. Evidence of p53-dependent cross-talk between ribosome biogenesis and the cell cycle: effects of nucleolar protein Bop1 on G(1)/S transition. *Mol Cell Biol* 2001; **21**: 4246-4255 [PMID: 11390653 DOI: 10.1128/MCB.21.13.4246-4255.2001]

69 **Golomb L**, Volarevic S, Oren M. p53 and ribosome biogenesis stress: the essentials. *FEBS Lett* 2014; **588**: 2571-2579 [PMID: 24747423 DOI: 10.1016/j.febslet.2014.04.014]

70 **James A**, Wang Y, Raje H, Rosby R, DiMario P. Nucleolar stress with and without p53. *Nucleus* 2014; **5**: 402-426 [PMID: 25482194 DOI: 10.4161/nucl.32235]

71 **Toledo F**, Wahl GM. Regulating the p53 pathway: in vitro hypotheses, in vivo veritas. *Nat Rev Cancer* 2006; **6**: 909-923 [PMID: 17128209 DOI: 10.1038/nrc2012]

72 **Liu Y**, Deisenroth C, Zhang Y. RP-MDM2-p53 Pathway: Linking ribosomal biogenesis and tumor surveillance. *Trends Cancer* 2016; **2**: 191-204

73 **Quin J**, Chan KT, Devlin JR, Cameron DP, Diesch J, Cullinane C, Ahern J, Khot A, Hein N, George AJ, Hannan KM, Poortinga G, Sheppard KE, Khanna KK, Johnstone RW, Drygin D, McArthur GA, Pearson RB, Sanij E, Hannan RD. Inhibition of RNA polymerase I transcription initiation by CX-5461 activates non-canonical ATM/ATR signaling. *Oncotarget* 2016; **7**: 49800-49818 [PMID: 27391441 DOI: 10.18632/oncotarget.10452]

74 **Negi SS**, Brown P. rRNA synthesis inhibitor, CX-5461, activates ATM/ATR pathway in acute lymphoblastic leukemia, arrests cells in G2 phase and induces apoptosis. *Oncotarget* 2015; **6**: 18094-18104 [PMID: 26061708 DOI: 10.18632/oncotarget.4093]

75 **Hannan KM**, Sanij E, Rothblum LI, Hannan RD, Pearson RB. Dysregulation of RNA polymerase I transcription during disease. *Biochim Biophys Acta* 2013; **1829**: 342-360 [PMID: 23153826 DOI: 10.1016/j.bbagrm.2012.10.014]

76 **Drygin D**, Lin A, Bliesath J, Ho CB, O’Brien SE, Proffitt C, Omori M, Haddach M, Schwaebe MK, Siddiqui-Jain A, Streiner N, Quin JE, Sanij E, Bywater MJ, Hannan RD, Ryckman D, Anderes K, Rice WG. Targeting RNA polymerase I with an oral small molecule CX-5461 inhibits ribosomal RNA synthesis and solid tumor growth. *Cancer Res* 2011; **71**: 1418-1430 [PMID: 21159662 DOI: 10.1158/0008-5472.CAN-10-1728]

Figure Legends

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**Figure 1 Schematic of major pathways associated with pancreatic ductal adenocarcinoma and site of action of current treatments.** Multiple pathways and receptors are associated with the development of pancreatic ductal adenocarcinoma (PDAC) including epidermal growth factor receptors (EGFR), human epidermal growth factor receptor 2 (Her2), and vascular endothelial growth factor receptor (VEGFR). All of these have important roles in the RAS/RAF/MEK/ERK and AKT/PI3K/mTOR pathways involved in cell growth. EGFR also has a role in the JAK/STAT pathway necessary for activation of signalling cascades and gene transcription. Transforming growth factor (TGF-) is a multifunctional cytokine involved in various processes some of which are mediated by SMAD 4, a known mutation associated with development of PDAC. Current therapeutics target these processes at various sites.

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**Figure 2 Pathways involved in ribosome biogenesis.** The first rate limiting step in ribosome biogenesis is the transcription of the rRNA genes by Pol I which forms a multiprotein transcription complex at the rDNA promotor[62]. Green arrows indicate up stream regulators which exert positive effects on the Pol I transcription complex including multiple pathways such as PI3K/AKT/mTOR, RAS/RAF/MEK and Myc which functions as the “master regulator” for cell growth[64]. The mature rRNAs together with ribosomal proteins assemble into the 40 and 60S ribosomal subunits, which then form the functional 80S ribosome[62]. Transcription of the rDNA repeat is negatively influenced by p53 (as shown by the red line) to ensure that cell growth/proliferation is tightly regulated.

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**Figure 3 Effects on ribosome biogenesis under normal conditions and in response to nucleolar stress.** Under normal conditions, p53 levels are kept at a minimum by MDM2. This allows normal cell growth and proliferation. With nucleolar stress, RPs L5 and L11 are able to bind to MDM2 as does p14ARF. This blocks the ability for MDM2 to inhibit p53. Similarly, p53 independent mechanisms also block ribosome biogenesis leading to cell cycle arrest, senescence or apoptosis.

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