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Parental obesity programs pancreatic cancer development in offspring

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Abstract

Epidemiological studies suggest that timing of obesity onset – and underlying metabolic dysfunction – is important in determining pancreatic cancer rates: early and young adult abdominal overweight/obesity is more strongly associated with this cancer than obesity that develops later in life. Parental obesity and overweight are associated with metabolic dysfunction and obesity in their children. Here, we evaluated the impact of parental overweight on offspring’s susceptibility of pancreatic cancer using the P48Cre+/KrasG12D/+ mouse model. Male mice were fed an obesity-inducing diet (OID) before conception and mated with females raised on a control diet (CO) to generate the offspring. In a separate experiment, pregnant dams were fed CO or OID throughout gestation. The resulting OID offspring from the maternal (OID-m) or paternal lineage (OID-p) were used to study body weight, metabolic parameters and pancreatic cancer development and for molecular analysis. Parental obesity increased offspring’s body weight at birth, weaning and in adulthood compared to CO, with gender- and genotype-specific differences. OID-p and OID-m offspring showed metabolic disorder and accelerated development of high-grade PanIN/PDAC. OID offspring also had higher rates of acinar-to-ductal reprogramming assessed by CPA1+/SOX9+-positive pancreatic cells. Levels of Tenascin C (TNC), an ECM glycoprotein shown to suppress apoptosis, were elevated in OID offspring, particularly females. In line with that, OID offspring displayed increased collagen content and decreased apoptosis in pancreatic lesions compared to CO. An ancestral history of obesity through either the paternal or maternal lineages increases offspring’s susceptibility to pancreatic cancer development.

Introduction

Pancreatic ductal adenocarcinoma (PDAC) has an overall 5-year survival rate of about 9% (American Cancer Society 2018). This dismal prognosis is due to lack of early detection methods, effective therapies and our poor understanding of risk factors for this disease.

While presenting a ductal phenotype, PDAC has been experimentally shown to arise from the acinar compartment through a process of acinar-to-ductal reprogramming (Kopp et al. 2012). Kras mutations are present in early human PanIN lesions as well as invasive PDAC, and this activating mutation is generally accepted as the initiating event in PDAC progression (Kanda et al. 2013). However, PDAC development in mice carrying the KrasG120V mutation is slow (Hingorani et al. 2003), suggesting that other genetic or environmental factors are needed for tumor promotion (Dawson et al. 2013).
There are few established risk factors for PDAC with family history, chronic pancreatitis, smoking, metabolic syndrome and obesity among them (Pandol et al. 2012). Epidemiologic studies suggest that obesity, overweight and underlying metabolic dysfunction account for up to 50% of all PDACs (Bracci 2012), and findings from animal studies support this association (Dawson et al. 2013, Chang et al. 2017). Population studies also suggest that timing of overweight/obesity onset is important in determining pancreatic cancer rates: childhood and young adult abdominal overweight/obesity is more strongly associated with this cancer than overweight/obesity that develops later in life (Li et al. 2009, Genkinger et al. 2015, Nogueira et al. 2017). However, the mechanisms behind this association are not well understood.

Parental environmental and lifestyle exposures have been shown to affect phenotypic characteristics in the next generation (Jirtle & Skinner 2007). Most of the evidence comes from maternal exposures during pregnancy, given the intimate relationship between mother and the fetus (de Assis et al. 2006, Loche et al. 2018). However, recent studies demonstrate that pre-conception paternal factors can also modulate the risk of disease in their progeny (Ng et al. 2010, Rodgers et al. 2015). Children of obese parents are more likely to develop overweight, insulin resistance and diabetes in early life (Dorner & Plagemann 1994, Linabery et al. 2013, Santos et al. 2018). These findings have been recapitulated in animal studies with overweight and metabolic dysfunction being observed in offspring for both maternal and paternal obesity (Jimenez-Chillaron et al. 2009, McPherson et al. 2015, Huypens et al. 2016).

Here, we evaluated the impact of diet-induced parental obesity in the susceptibility of pancreatic cancer in the next generation using the P48O/e+/KrasG12D/+ (hereafter referred to as KC) mouse model of pancreatic cancer. Our study shows that a history of obesity from either the paternal or maternal lineage is associated with accelerated development of high-grade PanIN and invasive PDAC in offspring carrying the KrasG12D/+ mutation. This phenotype was associated with increased rates of acinar-to-ductal reprogramming as well as increased desmoplasia and other alterations in the extracellular matrix (ECM) in a gender-specific manner.

**Experimental procedures**

**Animals and dietary exposures**

**Parental exposures**

*LSL-KrasG12D/+* or *P48O/e/+* male mice were fed AIN93G-based diets containing either 17.2% (Control, CO, Envigo-Teklad #TD.160018) or 57.1% energy from fat (Lard-based, Obesity-Inducing-Diet, OID, Envigo-Teklad #TD.160019) (Diet details in Supplementary Table 1, see section on supplementary data given at the end of this article) starting after weaning (3 weeks of age). Males’ body weight was recorded weekly. At 10 weeks of age, OID-fed and control-fed male mice were mated to female mice (*LSL-KrasG12D/+* or *P48O/e+*) reared on the CO diet to generate the offspring. Males were kept in female cages for 3 days. Female mice were kept on the control diet during the breeding, for the extent of pregnancy and after giving birth. The weight and number of pups per litter was determined 2 days after birth. To avoid litter effect, pups were cross-fostered 2 days after dams gave birth. Pups from 2 to 3 dams were pooled and housed in a litter of 8–10 pups per nursing dam. All pups were weaned on postnatal day 21 and fed a standard chow diet throughout the experiment. Pups’ body weight was recorded weekly.

**Maternal exposures**

*LSL-KrasG12D/+* or *P48O/e/+* females were fed the CO or OID starting 5 days before mating. OID-fed and CO-fed females were then housed together with male mice (*LSL-KrasG12D/+* or *P48O/e/+* genotype) raised on the CO diet to mate for 3 days. Pregnancy onset was assessed by the presence of a vaginal plug. Maternal body weight was recorded weekly. OID-fed dams were switched to the CO diet, starting 2 days after delivery. Pups were cross-fostered and weaned as described earlier.

The offspring of CO or OID mothers and fathers were used to study body weight, metabolic parameters, molecular analyses and pancreatic tumorigenesis as described in the following sections.

All animal procedures were approved by the Georgetown University Animal Care and Use Committee, and the experiments were performed following the National Institutes of Health guidelines for the proper and humane use of animals in biomedical research. Animals were killed if symptoms of advanced tumor development (palpable mass, jaundice, sudden weight loss) occurred.

**Genotyping**

The genotype of each animal was determined using genomic DNA extracted from tail clips and PCR using a commercially available service (Transnetyx, Inc., Cordova, TN, USA).

**Pancreas histology**

Pancreatic tissue sections were fixed in neutral buffered 10% formalin, paraffin-embedded, sectioned (5 µm) and stained with hematoxylin and eosin
Pancreatic tissue collection  Pancreatic tissue from control and OID offspring were collected at 2 months of age and again at 6–7 months (>6 months, hereafter) of age and used for pancreatic histopathology, immunohistochemistry analyses, RNA and protein extraction.

RNA sequencing  RNA extraction was performed with the miRNeasy Mini Kit (Qiagen), according to the manufacturer's protocol, and its concentration, quality and integrity were confirmed using NanoDrop ND-1000 (Thermo Scientific). The quality of the samples was assessed using a 2100 Bioanalyzer (Agilent Technologies) for RNA integrity number (>6.0) and concentration (minimum 60ng/µL). The RNA-seq analysis (n=6/group) was performed by GENEWIZ (South Plainfield, NJ, USA).

RNA library construction  The RNA samples were quantified using Qubit 2.0 Fluorometer (Life Technologies) and RNA integrity was checked with Agilent TapeStation (Agilent Technologies). RNA library preparation, sequencing reaction and initial bioinformatics analysis were conducted at GENEWIZ, LLC. RNA sequencing library preparation used the NEBNext Ultra RNA Library Prep Kit for Illumina by following manufacturer's recommendations (NEB, Ipswich, MA, USA). Briefly, mRNAs were first enriched with oligod(T) beads. Enriched mRNAs were fragmented for 15 min at 94°C. First-strand and second-strand cDNA were subsequently synthesized. cDNA fragments were end repaired and adenylated at 3'ends, and universal adapter was ligated to cDNA fragments, followed by index addition and library enrichment with limited cycle PCR. The sequencing library was validated on the Agilent TapeStation (Agilent Technologies) and quantified using Qubit 2.0 Fluorometer (Invitrogen) as well as by quantitative PCR (Applied Biosystems). The sequencing libraries were clustered on two lanes of a flowcell. After clustering, the flowcell was loaded on the Illumina HiSeq instrument according to manufacturer's instructions. The samples were sequenced using a 2 × 150 paired-end (PE) configuration. Image analysis and base calling were conducted by the HiSeq Control Software (HCS). Raw sequence data (.bcl files) generated from Illumina HiSeq was converted into fastq files and de-multiplexed using Illumina’s bcl2fastq 2.17 software. One mismatch was allowed for index sequence identification.

Data analysis  All raw data passed the FastQC quality check. We used Rsem (Li & Dewey 2011) with GRCm38 mouse reference sequence (10mm) to assess gene expression level. We compared KC gene expression profile of control group to OID group. We also performed this analysis by gender (OID females vs control females; OID males vs control males). For all sets of comparisons, we performed differential expression analysis using limma package (Ritchie et al. 2015) in R and applied fdr < 0.1 as cut off point. A heatmap was generated for each set of filtered genes, which was then uploaded to IPA (Qiagen Inc., https://www.qiagenbioinformatics.com/products/ingenuity-pathway-analysis) for network and functional analysis.

Validation of RNA-seq data by quantitative real-time polymerase chain reaction  Differentially expressed genes identified (Lama1, Tff3 and Tnc) in our RNA-seq were validated through q-PCR. Total RNA from KC offspring pancreatic tissue was extracted (n=9–13/group) using the RNeasy Lipid Tissue Mini Kit per the manufacturer’s protocol. The concentration, purity and quality of RNA samples were assessed as described earlier. Two micrograms of RNA per sample were used for the synthesis of cDNA using the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystem) according to manufacturer’s instructions. The expression levels of target genes were evaluated using a Step One Plus Real-Time PCR system (Applied Biosystems). The cDNA (2µg) was mixed with 5µL TaqMan Fast Advanced Master Mix (Applied Biosystem), 0.5µL TaqMan Assay and 3.5µL
ultrapure water (UltraPure DNase/RNase-Free Distilled Water) in 96-well plates. The amplification reactions were performed with a thermal cycling method, consisting of two initial cycles of 2 min at 50°C and 20 s at 95°C, 40 cycles of 1 s at 95°C and 20 s at 60°C. Expression of target genes were normalized to the housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (Gapdh). The variation in the expression of target genes among experimental groups was analyzed using delta Ct.

Immunohistochemistry and immunofluorescence staining

Briefly, tissues were fixed in 10% buffered formalin, embedded in paraffin and sectioned (5 μm). Sections were deparaffinized with xylene and rehydrated through a graded alcohol series. For immunohistochemistry staining, the antigen retrieval was performed by immersing the tissue sections at 98°C for 40 minutes in 1x Diva Decloaker (Biocare, Pacheco, CA, USA). Tissue sections were treated with 3% hydrogen peroxide and 10% normal goat serum for 10 min and were incubated with the primary antibody (Supplementary Table 2), overnight at 4°C. After several washes, sections were treated with the appropriate HRP-labeled polymer for 30 min and DAB chromogen (Dako) for 5 min. Slides were counterstained with hematoxylin (Fisher, Harris Modified Hematoxylin), blued using 1% ammonium hydroxide, dehydrated and mounted with Acrymount. The sections were photographed using an Olympus IX-71 Inverted Epifluorescence microscope at 20× magnification. Images were evaluated with ImageJ software (National Institutes of Health, Bethesda, MD, USA) to quantify total pancreatic area, as well as the percentage of positive cells for CPA1, SOX9 and laminin. The proliferation index (ki67 staining) was determined by immunoRatio, a plugin for the ImageJ software to quantify hematoxylin and DAB-stained tissue.

Western blot

Protein levels were assessed by Western blot in the pancreatic tissue and tumors obtained from OID or control offspring. Total protein was extracted from pancreatic tissues and tumors using RIPA buffer with Halt Protease Inhibitor Cocktail (Thermo Fisher). Ten micrograms of protein extracts were resolved on a 4–12% denaturing polyacrylamide gel (SDS-PAGE). Proteins were transferred using the iBlot 7-Minute Blotting System (Invitrogen) and blocked with 5% nonfat dry milk for 1 hour. Membranes were incubated with the specific primary antibodies (for antibody specifications and dilutions, see Supplementary Table 2) at 4°C overnight. After several washes, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibody at room temperature for 1 hour. Membranes were developed using the Chemiluminescent HRP antibody detection reagent (Denville Scientific Inc., Metuchen, NJ, USA) and exposed to blot imaging systems (Amersham Imager 600, GE Healthcare Life Sciences). Optical density of the bands was quantified using Quantity-one software (Bio-Rad). To control for equal protein loading, expression of the proteins of interest was normalized to the β-tubulin signal or cyclophilin signal.

Collagen staining

To evaluate collagen deposition, 5 μm thick paraffin-embedded pancreatic sections were stained with picrosirius red staining. Briefly, after de-waxing and hydration, tumor sections were stained with picro-sirius red solution (0.5 g of picro-sirius red F3B (C.I. 35782) in 500 mL of

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saturated aqueous solution of picric acid) for 1 h. Sections were washed in two changes of acidified water, dehydrated in three changes of 100% ethanol, cleared in xylene and, then, mounted in a resinous medium. The sections were photographed using an Olympus IX-71 Inverted Epifluorescence microscope at 40× magnification. Staining was quantified by converting the image to gray scale and then isolating the red-stained collagen using thresholding and subsequent measure of the thresholded area with ImageJ software.

Statistical analyses

All statistical analyses were done using GraphPad Prism7 software. The results are presented as mean ± S.E. Comparisons between groups were conducted using two-tailed Student’s t test (or corresponding non-parametric test). When comparing two variables for multiple conditions, a two-way ANOVA was performed. Time-courses measurements such as body weight and glucose tolerance test were analyzed by repeated-measures ANOVA. For comparison of frequency measurements such as cell phenotyping (CPA1+, SOX9+, TNC+) and PanIN/PDAC development, a chi-square test was performed. A P value of less than 0.05 was considered significant.

Results

Parental phenotypes

Male mice fed OID gained more weight and were significantly heavier before breeding compared to males fed the CO diet (Supplementary Fig. 1A and B, P < 0.05, P = 0.02). The increase in body weight was accompanied by an increase in circulating leptin (P < 0.09), but not in insulin, levels (Supplementary Fig. 1C and D).

Pregnant dams fed OID during gestation also gained more weight than those on the CO diet (Supplementary Fig. 2A and B, P < 0.05). However, circulating leptin and insulin levels were not different between groups (Supplementary Fig. 2C and D).

Offspring phenotypes

While both female and male (WT and KC combined) offspring of OID-fed fathers had a trend toward higher weight at birth (Fig. 1A and B, P = 0.07), only OID-p female offspring were heavier at weaning (Fig. 1C and D, P < 0.05) compared to CO. No major differences in body weight were observed after weaning in OID-p offspring (Fig. 1E, F, G and H). However, we detected a non-significant increase in body weight in adult OID-p females from the KC genotype (Fig. 1F).
Both female and male (WT and KC combined) offspring of OID-fed mothers had higher weight at birth compared to CO (Fig. 1I and J, \( P=0.01 \)), but not at weaning (Fig. 1K and L). After puberty (Fig. 1M, N, O and P), we detected an increase in body weight in both the KC females and males from the OID-m group (Fig. 1N and P), with statistically significant differences found only in OID-m males compared to CO (Fig. 1P, \( P<0.01 \) and \( P<0.05 \)).

OID-p offspring showed impaired glucose tolerance at 2 (Fig. 2A, B and C, \( P=0.02 \)) and 6 (Fig. 2D, E and F, \( P=0.04 \)) months of age, particularly in males. OID-m offspring also showed impaired glucose tolerance at 2 (Fig. 2G, H and I, \( P<0.01 \)) and at 6 (Fig. 2J, K and L, \( P<0.01 \)) months of age. Insulin intolerance was also detected in OID offspring at 6 months of age (Supplementary Fig. 3A, B and C, \( P=0.01 \)). However, no differences in circulating insulin or leptin levels were detected between the groups (Supplementary Fig. 3D, E, F, G, H and I).

**Rates of PanIN and PDAC development in OID offspring**

All KC mice in either the CO or OID offspring groups developed PanIN and PDAC as previously described (Hingorani et al. 2003). The tumorigenesis process, however, was accelerated in the OID offspring from both the paternal or maternal OID lineages (Fig. 3A, B, C, D, E, F, G, H, I, J, K, L, M, N, O, P, Q, R, S, T, U, V, W and X).

At 2 months of age, the OID-p offspring had increased ratio of high/low-grade PanIN compared to CO (Fig. 3E and F, \( P=0.02 \)). The OID-p offspring also had a non-significant increase in numbers of invasive PDAC (Fig. 3G and H, \( P=0.2 \)) at this time point, with 50% of OID-p mice

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**Figure 2**

Metabolic function in CO and OID offspring. (A, B, C, D, E and F) Glucose tolerance test (GTT) in offspring from CO- and OID-fed fathers at 2 months (A, all offspring; B, females; C, males) and at 6 months (D, all offspring; E, females; F, males) of age. (G, H, I, J, K and L) GTT in offspring from CO- and OID-fed mothers at 2 months (G, all offspring; H, females; I, males) and at 6 months (J, all offspring; K, females; L, males) of age from CO- and OID-fed mothers (\( n=5 \)/gender/age/group). The data are expressed as mean±S.E.M. Significant differences versus the CO group were determined by two-way repeated-measures ANOVA; *\( P<0.05 \).
developing PDAC, while none were observed in the CO group. At 6 months of age and older, we continued to observe accelerated development of pancreatic lesions, with increased ratio of high/low-grade PanIN in the OID-p offspring compared to CO (Fig. 3I and J, \( P=0.01 \)). Slightly higher number of invasive PDACs in the OID-p group was also observed in this age cohort; however, results did not reach statistical significance (Fig. 3K and L).

PanIN and PDAC development in OID-m offspring (Fig. 3M, N, O, P, Q, R, S, T, U, V, W and X) largely mirrored what was observed for the OID-p offspring. At 2 months of age, the OID-m offspring had increased ratio of high/low-grade PanIN (Fig. 3Q and R, \( P=0.05 \)), particularly in females (Fig. 3R, \( P<0.05 \)). A non-significant increase in numbers of invasive PDAC in OID males compared to CO was also found (Fig. 3S and T). At 6 months of age and older, we continued to observe accelerated development of lesions, with increased ratio of high/low-grade PanIN in OID-m offspring compared to CO (Fig. 3U, V and W, \( P=0.02 \)), mainly in females (\( P<0.01 \)). Although not statistically significant, higher number of invasive PDACs was also observed in the OID-m offspring at this age cohort, with all PDAC detected in males (Fig. 3V, W and X).

Given the overlap in PanIN/PDAC phenotype between offspring produced from maternal and paternal OID exposures, all subsequent analyses were combined, with the OID-m and OID-p offspring collectively referred to as OID from this point on.

**Figure 3**
Pancreatic ductal adenocarcinoma development (PDAC) in CO and OID offspring. (A, B, C and D) Representative pancreas histopathology sections from offspring of CO- and OID-fed fathers at 2 months of age (A and B, H&E staining; C and D, Alcian blue staining); (E, F, G, H, I, J, K and L) Ratio of high/low-grade PanIN and invasive PDAC incidence in offspring of CO- and OID-fed fathers at 2 (E, F, G and H) and \( \geq 6 \) months (I, J, K and L) of age. (M, N, O and P) Representative pancreas histopathology sections from offspring of CO- and OID-fed mothers at 2 months of age (M and N, H&E staining; O and P, Alcian blue staining); (Q, R, S, T, U, V, W and X) Ratio of high/low-grade PanIN and invasive PDAC incidence in offspring of CO- and OID-fed mothers at 2 (Q, R, S and T) and \( \geq 6 \) months (U, V, W and X) of age. The data are expressed as mean \( \pm \) s.e.m. Significant differences versus the CO group were determined by two-way ANOVA, Student’s t test or chi-square test; **\( P<0.01 \) and *\( P<0.05 \). Inset, PanIN lesions; arrows, invasive PDAC lesion. Scale bars: 1.4 \( \mu m \). A full colour version of this figure is available at https://doi.org/10.1530/ERC-19-0016.
Rates of proliferation and apoptosis in OID offspring
PanIN and PDAC

The KRAS signaling pathway (MAPK activation) (Supplementary Fig. 4A and B) is increased in OID offspring pancreas compared to CO. Thus, we evaluated pancreatic cell proliferation and apoptosis rates in KC OID offspring (Fig. 4A, B, C, D, E, F, G, H, I, J, K, L and M). While we did not find differences in cell proliferation levels (ki67 staining) between the groups (Fig. 4E, F and G), we detected a reduction in apoptosis levels (Fig. 4H, I and J) in OID males at 2 (P<0.001) and in both OID genders at 6 months of age (P<0.01) compared to CO. We also found a concomitant increase in the pancreatic proliferation-to-apoptosis (Fig. 4K, L and M) ratio for the OID offspring at 2 and 6 months of age, but results did not reach statistical significance.

Increased rates of acinar-to-ductal reprogramming in OID offspring

It has been shown that acinar-to-ductal reprogramming is the main mechanism by which pancreatic ductal carcinoma arises (Kopp et al. 2012). Given the higher rates of PanIN/PDAC progression in OID offspring, we wondered whether pancreatic acinar-to-ductal reprogramming was also accelerated. Using co-staining for CPA1 (acinar cell marker) and SOX9 (inducer of ductal cell identity), we observed significantly higher rates of acinar-to-ductal reprogramming (CPA1+, SOX9+ cells) in pancreata of KC OID offspring compared to CO both at 2 and 6 months of age (Fig. 5A, B, C, D, E, F, G and H and Supplementary Fig. 5, P<0.001, P<0.001).

ECM alterations in OID offspring

Given the accelerated PDAC progression and acinar-to-ductal reprogramming in KC OID mice, we investigated whether there were other differences in pancreatic gene expression patterns between the groups (6 months of age) that could explain our findings. Surprisingly, we found that a comparison between the CO and OID offspring yielded little differences (three annotated genes, data not shown) when male and females in each group were analyzed together. When the analysis was
performed by gender, however, several genes showed differential expression in KC OID offspring compared to CO, particularly in females. Results are presented in heat-maps in Supplementary Fig. 6. A pathway analyses (Supplementary Fig. 7) revealed ‘organismal injury’ and ‘connective tissue function/disorder’ as some of the main bio-functions altered in OID offspring. Genes associated with those functions and upregulated in OID offspring included Tnc, Lama1 and Tff3.

An increase in Tnc (Tenascin C) expression was observed only in OID females in the RNA-seq analysis. Validation by q-PCR confirmed those results (Fig. 6A and B, P = 0.02). Also, in agreement with the RNA-seq results, q-PCR validation showed that Lama1 was significantly upregulated in OID female pancreas (Fig. 6C and D, P < 0.001) only. Validation of Tff3 expression by q-PCR, on the other hand, revealed a non-significant increase in both male and female OID offspring’s pancreas compared to CO (Fig. 6E and F). Next, we measured the protein levels of TNC using immunofluorescence and, in line with the RNA-seq results, found increased TNC expression (P > 0.001) in OID female, but not male, offspring pancreatic tissues at 6 months of age compared to CO (Fig. 6G and H). We also quantified protein of levels of laminin (LAMA1) using immunohistochemistry, and while we detected an increase expression in OID offspring pancreases, the results were not statistically significant (Fig. 6I and J).

PDAC is characterized by desmoplastic reaction (Whatcott et al. 2015) Given that in pancreatic tissue of OID offspring some differentially expressed genes were associated with connective tissue function/disorder, we investigated whether there were differences in desmoplasia between groups. Using picro-sirius red staining (collagen marker), we observed that pancreatic tissues of KC OID offspring displayed increased collagen content at 6 months of age (Fig. 6K and L, P < 0.01).

Discussion

Using the KC mouse model of pancreatic cancer, we demonstrated that parental obesity in pre-conception (paternal) or in pregnancy (maternal) is linked to accelerated PanIN/PDAC development in the next generation. This increase in PDAC development was associated with higher rates of acinar-to-ductal reprogramming, reduced apoptosis rates and ECM-related changes, including higher desmoplasia and TNC expression (in females) in pancreatic tissues.

There is strong evidence that parental obesity is linked to metabolic dysfunction and overweight in the
next generation in humans (Dorner & Plagemann 1994, Linabery et al. 2013, Santos et al. 2018). These findings have been recapitulated in animal studies (Jimenez-Chillaron et al. 2009, McPherson et al. 2015, Huybens et al. 2016). Our study supports those findings as we detected an increase in early life body weight as well as alterations in metabolic function for both OID-p and OID-m offspring.

In population studies, obesity/overweight and diabetes are consistently linked to PDAC (Bracci 2012, Eibl et al. 2018). Obesity and diabetes often co-exist, but can also independently increase the risk for PDAC (Jiao et al. 2010, Eibl et al. 2018). The timing of obesity has been shown to affect the association with PDAC: central obesity and higher BMI in childhood and young adulthood is associated with greater risk of pancreatic cancer and earlier disease onset compared to normal weight individuals or those who are normal weight or gain more weight later in life (Li et al. 2009, Genkinger et al. 2015, Nogueira et al. 2017). Our findings offer support to those population studies. Further, our results also suggest that metabolic dysfunction in early life accelerates PDAC development, particularly in males. However, the relationship between pancreatic cancer and diabetes is complex as diabetes can play both a causative role and also be a consequence of pancreatic cancer (Risch 2019). Given that the OID offspring with the WT genotype had metabolic alterations, it is likely that the metabolic dysfunction in our model is causal and not the end result of PDAC development. While those findings need to be further investigated in animal models, it will be important to determine whether in patients with familial pancreatic cancer, early life metabolic dysfunction by itself or underlying obesity would accelerate PDAC onset.

We have previously shown that ancestral obesity or nutritional patterns can increase other types of cancer in offspring (de Assis et al. 2012, Fontelles et al. 2016, da Cruz et al. 2018). This present study shows that parental obesity can predispose their children to PDAC as well. We also detected gender-specific effects in OID offspring. The metabolic dysfunction in OID offspring was more readily detectable in OID males compared to OID females. In line with that, most of the invasive PDACs were observed in OID males. However, OID females also had significantly higher rates of high-grade PanINs than controls females, suggesting that the oncogenesis process is accelerated within this group as well.

There are striking similarities in the phenotypic changes as well in PDAC development in both the offspring from overweight fathers and mothers, indicating that the triggering mechanisms behind those phenotypes
are likely the same for either ancestral lineage. While we have not performed mechanistic studies, some reports suggest that a common alteration underlying paternal and maternal exposures' effects on offspring could be alterations in placenta development (Rosenfeld 2015, Watkins et al. 2017). Further, we and others have shown that paternal obesity and nutrition leads to changes in sperm small non-coding RNAs (Fontelles et al. 2016, da Cruz et al. 2018) in rodents and humans (Donkin et al. 2016). The small RNAs present in sperm have been shown to alter embryonic development after fertilization in animals (Sharma et al. 2016).

Interestingly, while both OID fathers and mothers gained more weight than controls, levels of leptin were higher in OID fathers only. There are two possible explanations for this finding. First, fathers were fed the OID for a longer period of time than mothers (7 weeks versus ~4 weeks). More likely, however, the reason why we did not observe increased leptin levels in OID mothers is that this adipokine increases with pregnancy (Domali & Messinis 2002) and would have increased in control pregnant mothers as well, masking the effects of increased body weight.

Although having a ductal phenotype, PDAC has been experimentally shown to arise from the acinar compartment through a process of acinar-to-ductal reprogramming (Kopp et al. 2012). Using cell population-specific tracing experiments, a recent study showed that this process depends on ectopic expression of the ducal gene Sox9 in acinar cells that, in the presence of mutation oncogenic Kras, progress to PanIN (Kopp et al. 2012). Our data show that OID offspring have greater numbers of CPA1+/SOX9+ cells, indicating that acinar cells transition more quickly to a ductal-like state, in the presence of oncogenic Kras. Whether this is the case in young individuals that are overweight/obese or have metabolic dysfunction still needs to be investigated. Studies also have shown that SOX9 expression is activated by the MAPK pathway (Ling et al. 2011), which is in line with the increased activation of this pathway in OID offspring.

ECM abnormality is one of the hallmarks of cancer (Ritchie et al. 2015), and it has been demonstrated that obesity increases desmoplasia in pancreatic cancer, interfering with therapeutic efficacy (Incio et al. 2016). Further, Tenascin C has been shown to suppress apoptosis and rescue pancreatic cancer cells from gemcitabine-induced apoptosis (Shi et al. 2015). In line with that, our studies suggest that having a history of ancestral obesity leads to higher collagen deposition and other ECM-related alterations and reduced apoptosis levels in pancreatic tissues.

In this study, OID offspring were fed a balanced diet throughout their lives and whether postnatal dietary intake and other lifestyle changes can further modulate PDAC development still needs to be evaluated. It will be important also to elucidate if the pancreatic tissue environment or systemic changes are driving the increased PDAC observed in OID offspring. Metabolic disorders such obesity and diabetes are characterized by low-grade inflammatory state with increased levels of adipokines and pro-inflammatory cytokines (Monteiro & Azevedo 2010, Eibl et al. 2018), creating conditions for cancer development (van Kuijlsdijk et al. 2009).

The control diet in our study has a high carbohydrate, moderate fat profile, considered a balanced dietary pattern (Odermatt 2011). This control diet is clearly more beneficial to both parents and offspring health compared to the OID. While more comprehensive studies in both in humans and animal models are needed, those dietary patterns may possibly be recommended to parents trying to conceive as means to reduced metabolic diseases and cancer in the next generation.

Our study is the first to describe that parental obesity in pre-conception (fathers) or in pregnancy (mothers) is associated with accelerated development of PDAC in the next generation. Our study gives support to epidemiologic findings suggesting that childhood and early life overweight/obesity and metabolic dysfunction are more strongly associated with pancreatic cancer than obesity that develops later in life. Interventions to reduce adiposity and metabolic dysfunction in early life may be an effective way to prevent pancreatic cancer and likely other malignancies.
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