Pancreatic cancer (PC) is a form of malignancy of increasing incidence and poor prognosis, with an average of less than 10% of patients surviving 5 years after being diagnosed. The main reason for this unfavorable situation is the long asymptomatic course of the disease, and the absence of a simple screening method, typically leading to the late discovery of the disease. The development of the malignancy from the initial carcinogenesis into invasive pancreatic carcinoma takes approximately 10 years. However, the progression of pancreatic cancer from early into advanced stages can be, according to the latest studies, incredibly fast, just a few months. Early stages of pancreatic malignancy can be detected only by expensive, and sometimes invasive, diagnostic methods (CT, MR/MRCP, or EUS). Due to the current absence of a reliable non-invasive screening method, it is necessary to define a group of patients who have the highest risk of PC development, five to ten times higher risk compared to the regular population at a minimum. Risk factors combine in their effect; therefore, relative risks of PC development need to be summarized to obtain a total relative risk for each person. The main and non-influenceable risk factor in the development of PC is the increasing age. The other non-influenceable risk factor of PC is a genetic predisposition - family incidence of the disease can be detected in 4-16% of patients. Some specific genes and mutations which can play a role in PC development have already been identified (for example mutation of the PRSS-1 gene). Among the influenceable risk factors of PC is primarily smoking; obesity can play a part in PC development as well. A higher risk of PC is observed in patients with chronic pancreatitis. Nowadays, the relationship between PC and diabetes mellitus (DM) is hotly discussed. In the case of long-standing DM, the risk of pancreatic cancer is two times higher compared to the healthy population. However, new-onset DM can be the first sign of still asymptomatic PC. These patients, with paraneoplastic DM caused by pancreatic cancer cells, represent approximately 1% of recently diagnosed patients. However, this group of patients is still too large for screening. Because of that, it is necessary to find specific criteria to distinguish classic DM from the paraneoplastic form. The application of these criteria can help with the better stratification of risk in patients with new-onset diabetes and hence, it can help to discover PC in its early stages.

**Key words:** pancreatic cancer; pancreatic ductal adenocarcinoma; risk factors; chronic pancreatitis; PanIN lesions; mucinous cystic neoplasm; diabetes mellitus; smoking
Pancreatic ductal adenocarcinoma (PDAC), accounting for approximately 90% of pancreatic malignancies, is a therapy-resistant tumor, with a gradually increasing incidence and still poor prognosis with a median five-year survival rate still below 10% [1]. According to the data of the National Cancer Registry of the Czech Republic, there was a gradual significant increase in both the incidence of pancreatic cancer (PC) and its mortality from 1994 to 2016 [https://www.svod.cz/analyse.php?modul=incmor#]. An unfavorable prognosis is associated with the long asymptomatic course of the disease, as a result of which diagnosis is mostly late and most cases of PC are detected in the locally advanced stage, or even with already present distant metastases [1, 2].

PC oncogenesis is a multi-stage and relatively long process, occurring either by gradual progression of the severity of changes in relatively rare mucinous cystic lesions (IPMN - intraductal papillary mucinous neoplasia and MCN - mucinous cystic neoplasia) or more often by progression of the normal duct epithelium through the spectrum of pancreatic intraepithelial neoplasia (PanIN) into invasive PC [3, 4]. See Figure 1 for a summary of these pathways.

Progression from normal ductal epithelium through mild and severe dysplasia is characterized by a sequence of genetic alterations involving primarily a mutation in the \textit{KRAS} oncogene, occurring in the early stages of PC oncogenesis. In the later stages of oncogenesis, mutations in the tumor suppressor genes \textit{TP53}, \textit{CDKN2A} and \textit{SMAD4} (which are common in aggressive high-grade dysplasia) are typical [5]. See Figure 2 for details.

Overall, a development of about 10 years to the stage of invasive cancer is assumed, which on a theoretical level represents a sufficient time for early diagnosis [4, 6, 7]. However, there are alternative theories describing the possibility of significantly faster progress [8]. These accelerated genetic and epigenetic changes may explain the clinically known phenomenon of very rapid progression from localized PC to the advanced stages in months (Figure 3 and Figure 4) [9]. According to these theories, development may not be linear as expected according to the gradualism model, but due to the presence of factors such as aneuploidy, chromosomal instability and chromotripsis (detectable in up to 60% of PCs), very rapid progression can occur, as predicted by punctuated equilibrium theory (Figure 5) [10].

**Screening and diagnostics of PC**

**Screening for PC-theoretical background.** For the above reasons, it is necessary to search for people with early stages of PC to improve the PC prognosis. Due to its long asymptomatic course, it is necessary to search for asymptomatic individuals-conduct screening. Despite the increase in the
incidence of PC, population screening cannot be considered justified, both for financial reasons and also for generating an unacceptably high numbers of false positive results [11, 12]. For these reasons, research is currently focusing on efforts to identify and subsequently screen high-risk groups of people using our knowledge of risk factors for PC [13]. At least a 5-10-fold increase in PC risk is considered appropriate.

The defined group of people should have an expected risk of about 4% of finding PC in the following 3 years (Figure 6) [14]. Thus, medium- and high-risk groups can be considered suitable for screening, including some genetic mutations that cause, for example, hereditary chronic pancreatitis [15, 16] and a number of other tumor syndromes (Peutz-Jeghers syndrome, hereditary non-polyposis colorectal cancer (HNPCC), familial adenomatous polyposis (FAP), familial atypical multiple mole melanoma syndrome (FAMMM syndrome) [17-20]. Risk groups also include persons with familial PC, which is defined as the occurrence of PC in at least two or more first-degree relatives [21, 22]. In some meta-analyses, the presence of *Helicobacter pylori* appears to slightly increase the risk of developing PC [23, 24]. The individual risk factors, including the degree of increase in the risk of PC development, are summarized in Table 1 (adjusted according to [11, 13, 17, 18]).

As a summary, it is possible to say that about 90% of PC are sporadic (associated with potentially modifiable risk factors), but in some individuals, PC can be attributed to familial aggregation (7%) or high-risk genetic syndromes (3%). Separate risk factors aggregate (maybe even multiply) in their effects and, therefore, it is necessary to determine the profile of risk factors for each individual and screen the ones with substantial increase. In some groups, PC screening programmes using imaging methods are already ongoing worldwide, however, the results of screening among patients from high risk groups are not unambiguous. Professor Canto's group, which has been working on this issue for a long time, published the results of a long-term follow-up (16 years, median 5.6 years) of a group of 354 patients at high risk of PC using advanced imaging methods (EUS, NMR, CT). During the follow-up period, a suspected lesion was detected in 68 patients (19%), and in 24 (7%) there was a gradual progression. 9/10 of the detected PCs were resectable with a 3-year survival of 85% [25]. In contrast, a meta-analysis of examination studies (EUS or NMR) of high-risk (≥ 5%) PC patients, including 19 studies with a total of 7,085 patients, detected 59 high-risk lesions (43 PCs). Thus, it was necessary to examine 135 high-risk patients to detect one PC, and the authors question the cost-effectiveness of this approach [26].

**Imaging methods.** Due to the current absence of reliable laboratory markers, current diagnostics, including screening, is based primarily on the use of imaging methods.
The practical and sensitive imaging methods capable of accurate imaging of a pancreatic lesion and
determining its stage are: quality CT using a pancreatic protocol, MR-MRCP, EUS with the
possibility of performing a fine-needle aspiration biopsy (FNAB) [2, 27].
Abdominal ultrasound (US) is the best initial modality for its minimal invasivity, common
availability and safety, however, due to position of the pancreas in the retroperitoneum abdominal
ultrasound is not accurate enough in imaging of the pancreas.
On the contrary, endoscopic ultrasound (EUS) is a sensitive method for identification of small
lesions of the pancreas which can be subsequently focused for fine needle aspiration biopsy
(FNAB). EUS is more sensitive than CT, especially for small lesions (≤ 2 cm) and it is also most
reliable in evaluation of infiltration of large visceral vessels and lymph node involvement [28].
However, EUS represents endoscopic = invasive procedure with relatively low availability and high
operator dependence. Therefore, quality computed tomography (CT) using pancreatic protocol
(= triphasic scan in arterial, late arterial, and venous phases) is a fairly accurate and widely available
modality with relatively high sensitivity and specificity for PC detection, which is also useful for
distinction of patients eligible for resection with curative intent and those with unresectable disease
[29]. Limitation of CT are ionizing radiation and the application of intravenous contrast which is
problematic in patients with renal failure or allergy.
In such cases, magnetic resonance imaging (MRI) with gadolinium infusion is an alternative
method that can be used to diagnose and stage PC. MRI has not been proven to perform better than
CT in PC detection while it is more expensive and less available [30]. The length and loudness of
the procedure and small gantry of the MRI scanner can be limiting for claustrophobic patients.
Conversely, because of the absence of radiation, MRI is preferably used for screening of high-risk
individuals when the need for repeated examinations can be expected.
Addition of positron emission tomography (PET) either to CT or MRI combines functional PET
imaging with anatomical images of CT/MRI. Unfortunately, it does not differentiate inflammatory
and malignant changes because both conditions manifest with increased accumulation of the tracer
and therefore, PET does not differentiate PC from chronic pancreatitis. As such, PET-CT is similar
to CT alone and does not bring much of further benefits except selected cases – detection of small
distant metastases, monitoring of cancer recurrence after chemotherapy. PET-MRI seems to be
more reliable than a PET-CT and may be useful in cystic tumours, where PET-MRI enables exact
detection of structures located inside of the lesions, such as mural nodules or intraacystic septa [31].
The overview of diagnostic work-up of suspected pancreatic mass is summarized in Figure 7 [32].
In general, a major disadvantage of imaging methods in screening is the cost (CT, EUS, MRI), potential discomfort for patient (EUS, MRI) and possibly high rate of false positive examinations. Therefore, further improvement is necessary in detection of significant precursors or early PC and also in better distinction from clinically nonsignificant lesions to avoid unnecessary surgery and psychological stress in false positive cases. Thus, most sensitive and cost-effective imaging screening protocol still needs to be defined. MRI seems more cost-effective in overall screening, with EUS more cost-effective for highest-risk individuals (relative risk > 20). But cost-effectiveness depends on MRI and EUS costs, that vary considerably among countries [33].

**Laboratory diagnostics.** Due to the price and possible invasiveness of the above-mentioned imaging methods, a very attractive area of research is the effort to find a laboratory marker enabling a non-invasive PC screening in order to capture the early stage of PC. Unfortunately, despite intensive research, no progress has yet been made that would allow the use in everyday clinical practice. Increased levels of the tumor marker carbohydrate antigen 19-9 (CA 19-9), which is currently the only one used in routine clinical practice, is usually associated with the advanced disease and a poor prognosis [34]. Also, in patients without a functional Lewis enzyme (7-10% of the population), levels of CA 19-9 are typically undetectable or below 1.0 U/ml. In addition, CA 19-9 positivity can be caused by a number of other (even benign) diseases including cholestasis [35]. Its role in the detection of the early stages is therefore limited and, thus, CA 19-9 is not recommended as a screening marker for PC [36].

A number of other potential markers have been and are being investigated, unfortunately their real use in clinical practice has not yet been achieved [37]. However, some markers (eg MUC1, PC-594) show better results than CA 19-9 in pilot studies [38, 39]. Immunoglobulin G4 (IgG4), which is considered the diagnostic marker of autoimmune pancreatitis (AIP) can be slightly increased also in patients with PC [40]. More than a two-fold increase in IgG4 levels should be considered specific for AIP, which in a combination with negative CA 19-9 makes a diagnosis of PC highly unlikely [41].

Promising results in PC diagnostics have been published using the methods of proteomics, genomics, metabolomics and so-called "liquid biopsy" [42-44]. Currently, the use of a combination of new markers with CA19-9 [45] and the use of panels of individual biomarkers [46] seem to be useful.

Liquid biopsy is a detection of either circulating tumor cells (CTCs) or circulating tumor DNA (ctDNA) or tumor exosomes. They are either whole cells or their particules, derived from a primary tumor, that have entered the vasculature and circulate within the blood stream. CTCs possess the
capability to seed in distant organs—to metastasize. The possible role of their detection in early diagnosis of PC is in theory very promising. The identification and isolation of CTCs, ctDNA and/or tumor exosomes in PC is quite difficult, not standartized, with limited consistency of the results, however, studies have shown their presence in the bloodstream in the early stages of PC [47]. Thus, methods of liquid biopsy represent promising path for the early detection of PC especially as a profiling of several individual markers. Findings also suggest that higher diagnostic values of liquid biopsy methods available today can be reached when analyzed in a combination with CA19-9. A standardized detection method and large-scale validation are required before wide clinical application.

In summary, while it is hard to predict future development in the field, methods of liquid biopsy (especially miRNAs), proteomics, metabolomics appear most promising. The near future probably lies in a carefully selected panel of biomarkers that would allow for earlier diagnosis of PC and easier determination of its stage and, ideally, also for tailoring of the treatment plan and indication of prognosis/outcome.

Screening—summary and implications for clinical practice. The primary goal of PC surveillance should be prevention of PC related death and prevention of PC emergence by identifying and treating its precursor lesions. The average lifetime risk of developing PC in general population is too low for population-based screening. Therefore, high risk groups of patients need to be identified by taking in account present risk factors (their total in case of more risk factors). Individuals with PC risk increased ≥5 times may be screened for PC.

Currently, patients with familial PC and hereditary pancreatic cancer syndromes caused by certain germline mutations are considered candidates for PC screening. In future, individuals with substantial increase in PC based on modifiable risk factors may be subjects to PC screening programmes as well.

Careful physical examination, family and personal history including information on smoking, dietary habits and exposure to toxins can be helpul in the identification of individuals eligible for PC screening. Furthermore, obtaining a comprehensive cancer family history from newly diagnosed PC patients can help to identify family members who may benefit from surveillance.

At present, EUS and MRI represent the most sensitive imaging methods used in PC diagnostics and they are the cornerstone of PC screening programmes. However, most sensitive and cost-effective imaging screening protocol still needs to be defined.

Unfortunately, laboratory tests available today are not capable of reliable early diagnostics of PC but, in future, we can expect rapid development of laboratory tests using the methods of proteomics,
genomics, metabolomics and so-called "liquid biopsy" [42, 44]. This may even allow population screening of PC in general population. The overview of recommended approach to initial screening and further follow-up of persons with germline mutations with increased risk of PC development is summarized in Figure 8 [20]. This approach may be in future applicable also for people with substantially increased risk of PC based on age and accumulation of environmental risk factors. Some national guidelines recommend for high risk individuals even earlier start of PC screening (at age of 35) with annual EUS and serum CA19-9 [48].

**Individual risk factors**

Risk factors of PC development can be divided into non-modifiable and modifiable risk factors. Non-modifiable risk factors are represented primarily by age and genetics of each individual, modifiable risk factors mostly by lifestyle factors such as smoking, diet and toxins. Certain risk factors, such as obesity, diabetes and CP, may be in part modifiable indirectly by life-long healthy lifestyle, but they are difficult to influence once present. Risk factors are summarized in Table 1.

**Age.** The incidence of PC is increasing with age and it is typically a disease of the elderly. Ninety percent of newly diagnosed patients are over 55 years of age, with the majority over 70 [1].

**Familial pancreatic cancer.** Familial PC is defined as a family with at least 2 first-degree relatives affected by PC without other cancers or other known genetic syndromes or familial diseases. It is estimated that up to 10% of PC may have a familial component [18]. The number of first-and second-degree relatives with PC can be used to quantify PC risk. A family history of PC in one blood relative seems not to increase PC risk significantly, however, the risk of developing PC in relatives in families with 2 affected first-degree relatives is 6 to 18-fold higher (lifetime risk 8% to 12%) and kindreds with 3 affected first-degree relatives have a 32 to 57-fold risk increase of developing PC (lifetime risk 40%) [49]. Families with member affected by PC in a younger age (< 50 years) bear a higher risk of PC development.

**Hereditary Pancreatic Cancer Syndromes.** As mentioned earlier, certain germline mutations are known to significantly increase the risk of PC. They include *BRCA2, p16, STK11/LKB1* and *PRSS1* mutations [19]. It has been shown that BRCA2 is present in 17% to 19% of families where at least 2 first-degree relatives have PC [18].
Patients with known Peutz-Jeghers syndrome represent a population with a very high risk in a wide range from 36 to 132 and a cumulative lifetime risk of 36% for the development of PC. Therefore, they may represent a group in the highest risk of PC and some sources recommend to initiate PC screening at the age of 30-35 in Peutz-Jeghers syndrome patients [19]. Hereditary pancreatitis is an autosomal dominant disease attributed mostly to the \(\text{PRSS1}\) mutation. There is a high incidence of PC 30 to 40 years after the age of onset of recurrent attacks of pancreatitis. PC risk is 50-69 times higher, with an estimated lifetime risk of PC of 40% by 70 years of age [15]. The risk is doubled in smokers, who are diagnosed with PC on average 20 years earlier compared with nonsmoking hereditary pancreatitis individuals.

**PC and chronic pancreatitis.** Chronic pancreatitis (CP) is a progressive inflammation of the gland leading to irreversible morphological changes and impairment of both the exocrine and later endocrine functions of the pancreas [50]. A higher risk of tumor growth in the context of chronic inflammation has been known for a long time and has been described in a number of neoplasms not only of the gastrointestinal tract. A similar relationship can be found in the case of CP and PC [24]. Patients with the sporadic form of CP have a risk of developing PC 10-20 times higher than people of the appropriate age without CP. The risk of developing pancreatic cancer during the life of a patient with a sporadic form of CP is 1.8% after 10 years and 4% after 20 years of disease duration [49, 51]. Not all national guidelines consider increase of PC risk in long-lasting, non-genetically related CP adequate for routine PC screening [52]. However, it is crucial to aggregate all present risk factors in each individual and also to conduct an adequate diagnostic in the event of occurrence of new worrying symptoms in patients with CP (e.g. new-onset diabetes).

An exceptional group are patients with a previously mentioned genetically determined hereditary form of chronic pancreatitis. These patients account for less than 1% of all CP cases. Hereditary CP is caused by a mutation in the cationic trypsinogen gene (\(\text{PRSS1}\)) and is an autosomal dominant disease with a penetration of up to 80%. Characteristic of this disease are repeated attacks of acute pancreatitis with a high incidence of pancreatic tumor - approximately 50-80-fold increased risk of developing PC is present [15, 53].

The presence of chronic inflammation is associated with the production of free oxygen radicals, the production of cytokines and increased production of pro-inflammatory transcription factors. Similar mediators of inflammatory response pathways have been repeatedly demonstrated in CP and PC tissues. Nuclear factor kappa B (NF-\(\kappa\)B), cyclooxygenase 2 (COX-2), 5-lipoxygenase (5-LOX), interleukin-8 (IL-8) and other factors are involved in genetic damage, promote cell proliferation, inhibit apoptosis of the pancreatic cells and represent a link between chronic inflammation and
tumor growth [51]. Pancreatic chronic inflammation appears to be an early step in the development of a malignancy, with genetic alterations occurring as a result of prolonged inflammatory processes [54]. This may be evidenced, among other proofs, by the fact that PanIN lesions (representing individual steps of oncogenesis) are more common in patients with CP than in the general population [55].

In summary, CP is an established risk factor for PC because chronic inflammation promotes premalignant cell survival, autocrine stimulation of a protumorigenic environment, and desmoplasia. Progression of CP to PC occurs over one to two decades in about 5% of CP patients, with relative risk of PC about 10. The risk is much higher among patients with a hereditary predisposition. A substantial smoking history represents significant additional risk factor of PC development.

**PC and smoking.** Smoking is an independent risk factor not only for lung cancer, but also for cancer of the stomach, colon and pancreas. According to studies, smoking increases the risk of PC up to 2-3 times, 10-15% (25% by some sources) of cases of sporadic PC is expected to be conditioned by smoking [56-59]. The risk of developing PC is higher in people who smoke regularly for a long time (5 and more years) and is directly related to the number of cigarettes smoked per day (significantly with > 10 cigarettes/day). In smokers, PC is diagnosed at a younger age and more often within 1 year after the onset of the first symptoms, which may indicate faster progression of PC in smokers [60]. Cigarette smoking is also an independent risk factor for the development of the alcoholic and idiopathic form of CP and is thought to accelerate disease progression by inducing chronic inflammation. Cigarette smoking is likely to be involved in carcinogenesis by activating the inflammatory response [61].

In those with a family history and/or genetic predispositions for PC, smoking has even a greater effect (3.7-fold increased risk of developing PC) and may present with the disease one to two decades earlier. In individuals with *PRSS1* mutation smoking increases the risk of PC by 2-fold and decreases the age of PC onset by 10-20 years, as noted above.

**PC and diabetes mellitus.** A relationship between PC and diabetes mellitus (DM) has been known for decades, but the exact etiological relationship has not been fully elucidated yet. Impaired glucose tolerance or DM is present in up to 80% of PC patients [62]. The risk of PC in DM is explained by the fact that increased insulin production by Langerhans islet beta cells leads both to the exhaustion of beta cells (diabetes itself), but also to a higher local concentration of growth and stimulant factors that contribute to the malignant transformation of the surrounding exocrine tissue. This idea is justified and there are studies that demonstrate a more frequent occurrence of PC in
patients with a long history of DM with a relative risk of approximately 2 according to meta-analyses from 2011 and 2017 [63, 64]. However, the increase in relative risk is probably lower than previously thought and even becomes statistically insignificant if we exclude patients in whom DM preceded the diagnosis of PC by a short period of time (2 to 3 years).

For this reason, we can define 2 subgroups of patients with co-occurring DM and PC. The first subgroup consists of patients in whom DM is a genetically and environmentally conditioned underlying disease and PC appears after a long history of DM (either with or without direct relationship). The second subgroup consists of patients with DM of a short duration (2-3 years) before diagnosis of PC. In most patients with PC, the development of DM precedes the diagnosis of PC by < 2 years, and the relative risk of diagnosis of PC is 5.38 during the first year after diagnosis of DM [65]. Thus, we can speak of paraneoplastic DM, which is classified as a separate type of pancreatogenic diabetes (T3cDM) [66]. In this group of patients, DM may be the first manifestation of an otherwise asymptomatic malignancy, and DM is most likely a direct consequence of PC cells [63].

According to retrospective studies, PC will develop within 3 years of the diagnosis of DM in approximately 1% of diabetics, and given the total numbers, it is not possible to screen all newly diagnosed diabetics. Therefore, research today focuses mainly on identifying criteria that distinguish common DM from PC-related DM [62]. In the absence of a suitable biomarker, a number of authors have defined various distinguishing criteria between common DM and PC-related DM, but these have been refuted by further research as insufficient, with no significant difference in the clinical picture of common DM and PC-related DM [67, 68]. In general, however, it can be assumed that targeted examination of elderly people with newly diagnosed, atypically manifesting DM may lead to improved detection of early stages of PC and thus to better therapeutic results [69].

Another important factor useful in stratifying the risk of developing PC could be the sex of patients. Our study showed a significantly higher prevalence of DM among women with PC (25% of men vs 43.9% of women, p=0.0008), while in the control group, DM was equally represented in both sexes (22.1% in men vs. 17.2% in women, p=0.487) [70].

Modeling represents another alternative way to define a high-risk group among newly diagnosed diabetics without knowledge of a specific laboratory marker. By analyzing the documentation of more than 1,500 patients with newly diagnosed DM, the Mayo Clinic authors identified weight change, blood glucose fluctuations, and age as major risk factors, and developed their own Enriching New-onset Diabetes for Pancreatic Cancer (END-PAC) score capable of reliably
identifying a group of patients at high risk of PC diagnosis over the next 3 years after the onset of DM (80% sensitivity and specificity with a score ≥ 3) [71]. The high negative predictive value at a score of 0 is significant, when the risk of PC development is comparable to the general population. Overall, using the END-PAC scoring system, the authors achieved the definition of a population with about 4% risk of PC development, which is considered the limit at which screening is effective (Figure 9).

Unfortunately, retrospective validation on a group of almost 14,000 patients did not reach conclusive results. PC occurred in only 2% of patients in the high-risk group defined by END-PAC within 3 years of the diagnosis of DM [72]. However, this is not the only similar attempt. The authors from the University of Pennsylvania evaluated the data from a cohort of more than 100,000 patients over the age of 35 with newly diagnosed DM. Over the next 3 years, PC was detected in 0.4% of them. Based on the data obtained, they created a prediction model including age, BMI, change in BMI, smoking, PPI, anti-diabetic medication, and HbA1C values, cholesterol, haemoglobin, creatinine, and ALP levels. With necessity to examine 6.19% of newly diagnosed DM patients, they were able to identify PCs with 44.7% sensitivity and 94.0% specificity [73]. Further research and validation on large patient populations is needed to evaluate the applicability of these scoring systems.

In summary, long lasting DM represents factor increasing mildly the risk of PC while new-onset DM can be a paraneoplastic symptom. No simple strategy how to differentiate common DM and PC-related DM has been discovered so far, however, several prediction models are being tested and research in the field of biomarkers is ongoing. At present, testing patients with newly diagnosed DM especially in presence of other PC risk factors (e.g. age, smoking, chronic pancreatitis) may lead to improved detection of early stages of PC and thus to better therapeutic results.

Conclusion

PC screening has not yet been introduced in today’s clinical practice due to the absence of simple and effective methods. Thus, detecting the early stages of PC remains very problematic. Modern imaging methods are often expensive and invasive, so they remain reserved for groups of patients at high risk of developing PC. Research on laboratory markers has not yet revealed an indicator that would be able to identify patients with an early form of PC with sufficient sensitivity and specificity. However, this area of research is promising in the future and could lead to the desired results. Risk factors for PC are known and can be used to stratify the risk of PC. The selected group of patients with the highest level of risk must then be carefully monitored or grouped into registers.
The use of risk factors for sporadic PC such as age, smoking, obesity or chronic pancreatitis for further selection of patients before imaging is problematic, as the incidence of PC in these groups is very low. However, individuals with accumulation of risk factors may be in future potential candidates for PC screening. Also, the use of knowledge about the relationship between DM and PC can be very beneficial in the future after setting fixed criteria.

**Future Prospects**

Unfortunately, laboratory tests available today are not capable of reliable early diagnostics of PC but, in future, we can expect rapid development of laboratory tests using the methods of proteomics, genomics, metabolomics and so-called „liquid biopsy”. This may even allow population screening of PC in general population.

In the field of imaging methods, improvement is also necessary to enhance the detection of significant precursors or early PC but also to allow better distinction from clinically nonsignificant lesions to avoid unnecessary surgery. Most sensitive and cost-effective imaging screening protocol still needs to be defined as well.

Prospective studies of the effect of surveillance programs on survival benefit, surgical morbidity, postoperative quality of life and psychological stress are needed.

More research (e.g. large, prospective studies) on the role of PC risk factors and their interaction is necessary for precise identification of high-risk individuals.

**References**


DAMIANO J, BORDIER L, LE BERRE JP, MARGERY J, DUPUY O et al. Should pancreas imaging be recommended in patients over 50 years when diabetes is discovered


**Figure Legends**

**Figure 1.** Three distinct pathological pathways to invasive pancreatic carcinoma (edited according to [4] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 2.** Molecular changes during the progression of normal pancreatic tissue through the spectrum of PanIN lesions into invasive PC (edited according to [4] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 3.** Time progression of PC – fast progression from PC stage T1 into T3 and T4 in 12.5 and 14.3 months respectively (edited according to [9] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 4.** Time progression of PC - fast progression from PC stage T1-2N1 into T3N1 in 9.5 months (edited according to [9] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 5.** Different patterns of tumor progression in gradualistic and punctuated equilibrium models of pancreatic cancer progression (edited according to [10] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).
**Figure 6.** Increasing risk of pancreatic cancer in presence of currently known risk factors (edited according to [14] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 7.** Diagnostic work-up of suspected pancreatic mass (according to [32] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 8.** Flowchart of screening and follow-up of individuals in increased risk of PC development (according to [18, 19] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).

**Figure 9.** Stratification of risk of PC development in patients with new-onset diabetes using END-PAC score allowing the identification of high-risk individuals eligible for PC screening (edited according to [71] - created in collaboration with Service Center for E-Learning at Masaryk University, Faculty of Informatics).
Table 1. Risk factors of PC development (according to [11, 13, 17, 18]).

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<tr>
<th>Increase in pancreatic cancer risk</th>
<th>Clinical risk factors</th>
<th>Genetic risk factors</th>
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<tr>
<td><strong>Mild (&lt; 5 times)</strong></td>
<td>BMI &gt; 30 (RR 1.2-1.5)</td>
<td>Hereditary breast and ovarian cancer (HBOC) - <em>BRCA1</em> mutation (RR 1.9-5.3)</td>
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<td>Diabetes mellitus (RR 1.4-2.2)</td>
<td>Familial adenomatous polyposis (FAP)-<em>APC</em> mutation (RR 4.46)</td>
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<td>Family history of 1 first degree relative with PC</td>
<td>Hereditary nonpolyposis colorectal cancer (HNPCC)/Lynch syndrome-<em>MSH2, MLH1</em> etc. mutation (defective DNA mismatch repair, microsatellite instability) (RR 8.6-10.7)</td>
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<td>Smoking (RR 2-3.7)</td>
<td>Hereditary breast and ovarian cancer (HBOC)-<em>PALB2</em> mutation</td>
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<td>Carcinogens exposure (polycyclic and chlorinated hydrocarbons etc.)</td>
<td>Li-Fraumeni Syndrome-<em>p53</em> mutation</td>
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<td>High alcohol intake (RR 1.5)</td>
<td>Ataxia telangiectasia-<em>ATM</em> mutation (RR 2.7)</td>
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<td><em>Helicobacter pylori</em> infection (RR 1.5)</td>
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<td><strong>Medium (5-10 times)</strong></td>
<td>Chronic pancreatitis</td>
<td>Hereditary breast and ovarian cancer (HBOC)-<em>BRCA2</em> mutation (RR 5.9)</td>
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<td>Family history of 2 first degree relatives with PC (RR 18)</td>
<td>Cystic fibrosis-<em>CFTR</em> mutation</td>
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<td><strong>High (&gt; 10 times)</strong></td>
<td>Family history of ≥ 3 relatives of any degree with PC (RR 57)</td>
<td>Familial atypical multiple mole melanoma (FAMMM) syndrome-<em>CDKN2A</em> mutation (RR 16-46.6)</td>
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<td>Hereditary pancreatitis-<em>PRSS1</em> mutation (RR 50-69)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Peutz-Jeghers syndrome-<em>STK11</em> mutation (RR 36-132)</td>
</tr>
</tbody>
</table>
Fig. 2 color  Download full resolution image

Diagram showing the progression of events from normal pancreatic tissue to invasive carcinoma:
- Telomere shortening, KRAS
- CpG island hypermethylation
- MUC5AC
- MUC1
- P16/CDKN2
- MUC1
- TP53, SMAD4/DPC4
Fig. 3 color  Download full resolution image

T1 ≤ 2 cm

12.5 months
P = 0.03

T3

T1 ≤ 2 cm

14.3 months
P = 0.03

(average adjusted age difference)

T4
Fig. 4 color  Download full resolution image

T1N1

≤ 2 cm

T2N1

> 2 cm

9.5 months

P = 0.06

T3N1
Pancreatic cancer risk

Warrants clinical work-up → 4%

3-year Incidence (%)

Average
Low
High
Very high

0.1%
0.15%
0.85%

General population ≥ 50
Smoking* Obesity*
Long-standing DM*
1.5-2 fold
2-FDR with PDAC* New-onset DM
6-8 fold
Hereditary pancreatitis*
Chronic pancreatitis*
ENDPAC model for NOD
25-50 fold

DM - Diabetes mellitus
FDR - First-degree relative
PDAC - Pancreatic ductal adenocarcinoma
ENDPAC - Enriching new-onset diabetes for pancreatic cancer
NOD - New-onset diabetes

*Lifetime risk
Pancreatic lesion

CT in pancreatic protocol
(MRI/MRCP in specific situations - e.g. cystic lesions)
(PET in specific situations - e.g. suspicion of small distant metastases)

Resectable lesion of typical PC appearance without metastases

Locally advanced lesion

Metastatic disease

Urgent surgery (EUS confirmation of resectability in some centers)

EUS with biopsy

EUS with biopsy or metastasis biopsy
Fig. 8  Download full resolution image

Familial PC, Peutz-Jeghers syndrome (STK11 mutation)
Initial surveillance starts at 50-55 or 10 years younger than the youngest affected blood relative (some sources recommend 30-35 for Peutz-Jeghers syndrome)

The baseline screening: MRI/ MKLP, EUS, fasting serum glucose or serum HbA1C

No alarming abnormalities:
- normal pancreas
- mild chronic pancreatitis
- cyst without worrisome features

Alarming abnormalities:
- solid lesion
- cyst with worrisome features
- MPD stricture and/or dilation without clear mass

Non-functioning NET < 10mm
EUS with FNA, CT, CA 19-9

No clear suspicion of malignancy
Suspicion of malignancy

Cystic lesion with one of
- size ≥ 3cm
- MPD 5-9mm
- lymphadenopathy
- increased serum CA19-9
- growth rate ≥5mm/2 years

Solid lesion
- <5mm or uncertain significance
- MPD 5-9mm
- MPD stricture and/or dilation ≥ 6mm without a clear mass

FU 12 months  FU 6 months  FU 3 months
FOLLOW UP - MRI/ MRCP, EUS, fasting serum glucose or serum HbA1C, CA 19-9 if alarming abnormalities

Surgical resection

Germline mutations in: PRSS-1, BRCA1+2, ATM, CDKN2A, PALB2, MSH1+2, TP53
Initial surveillance starts at 40-50 or 10 years younger than the youngest affected blood relative
General Population ≥ age 50 years: 3-year Pancreatic cancer risk ~0.11%

New-onset diabetes

3-year Pancreatic cancer risk ~0.9%
(Pancreatic cancer n=10/1000)

Enriching New-onset Diabetes for Pancreatic Cancer (ENDPAC) score

>3
Very high risk: ~3.6%
(n=8/10)

1-2
Intermediate risk: ~0.5%
(n=2/10)

≤0
Low risk: <0.1%
(n=0/10)

clinical work-up