Pancreatic stellate cells contribute to regeneration early after acute necrotising pancreatitis in humans

A Zimmermann, B Gloor, A Kappeler, W Uhl, H Friess, M W Büchler

Background and aim: The aim of this study was to systematically analyse the pattern of regeneration in human acute pancreatitis by testing whether pancreatic stellate cells, their myofibroblastic offspring, and pancreatic ductules are involved in the regenerative process.

Patients and methods: Between January 1994 and November 2000, 24 necrosectomy specimens containing vital tissue were obtained for pathological examination. Formalin fixed tissue samples were routinely processed and immunostained for cytokeratins 7 and 19, smooth muscle actin, desmin, Ki-67, and CD68. Pancreatic tissue from organ donors served as normal controls.

Results: Necrosectomy specimens were obtained between 11 and 41 days after the onset of symptoms. In vital areas of necrosectomy samples, spherical hypercellular spheres consisting of loose vascular connective tissue occurred, in part showing duct-like profiles which sprouted from remnant exocrine tissue almost perpendicular to the periphery of the spheres. In normal tissue, only a few stellate cells and myofibroblasts were present around ducts and ductules. In contrast, numerous stellate cells and myofibroblasts were detected in the hypercellular regenerative spheres after acute pancreatitis, both being situated within the loose tissue and forming compact peri ductular sheaths. Stellate cells/myofibroblasts and ductule cells exhibited increased proliferative activity.

Conclusions: Pancreatic stellate cells and their activated myofibroblastic offspring may participate in regeneration after acute necrotising pancreatitis in humans. Time course studies are needed to further strengthen this regeneration concept.

Acute pancreatitis encompasses a whole spectrum of inflammatory lesions in the pancreas. In accordance with a recent proposition, these lesions are being described using the terms, severe acute pancreatitis, mild acute pancreatitis, acute fluid collections, pancreatic necrosis, and acute pseudocysts. The aetiology and pathogenesis of necrosis and haemorrhage as hallmarks of severe acute pancreatitis have been studied in detail, and several models of pathogenic pathways have recently been developed. The end result of an acute attack of acute pancreatitis is a well characterised type of pancreatic and peripancreatic tissue breakdown ranging from interstitial oedema and low grade multifocal necrosis of the pancreas (mild acute pancreatitis) to massive haemorrhagic necrosis. In patients who survive, at least some of these lesions are generally considered to be reversible, in contrast with the overall progressive character of chronic pancreatitis. If necrotic tissue is present it may become infected or forms fluid collections containing debris. These acute fluid collections may disappear spontaneously or develop into pseudocysts. What has not been specifically addressed in the question as to whether, during the time period between the initiation of necrosis and necrosectomy, repair and/or regeneration of the exocrine apparatus occurs. Theoretically, regeneration of pancreatic exocrine tissue destroyed by acute pancreatitis requires replacement of both epithelial cell populations and cells forming the matrix, allowing an ordered regrowth of lost epithelium.

Recently, stellate cells representing the homologue of the respective cells occurring in the liver have been identified in the pancreas. Pancreatic stellate cells have been shown to play a significant pathogenic role in fibrogenesis and in particular in mechanisms involved in fibrosis occurring in chronic pancreatitis. In contrast, it has not been established whether pancreatic stellate cells are involved in remodelling and regenerative mechanisms ensuing after acute necrotising pancreatitis in humans.
a pressure cooker. Primary antibodies were directed against cytokeratin (CK)-7 (clone OV-TL 12/30; Dako Diagnostics AG, Zug, Switzerland; working concentration 2 \( \mu \)g/ml; trypsin pretreatment), CK-19 (clone RCK108; Dako; 0.8 \( \mu \)g/ml; trypsin), desmin (clone D33; Dako; 5 \( \mu \)g/ml; microwave), α smooth muscle actin (SMA, clone 1A4; Sigma, St Louis, Missouri, USA; dilution 1:600; no pretreatment), CD68 (clone PG-M1; Dako; 2.5 \( \mu \)g/ml; microwave), and Ki-67 (clone MIB1; Dako; 1 \( \mu \)g/ml; pressure cooker). After the primary antibody a biotinylated goat-antimouse Ig antibody (Dako) was applied, followed by streptavidin-biotin complex/alkaline phosphatase (Dako). Sections were developed in new fuchsin-naphtol AS-BI (Sigma), counterstained with haematoxylin, and mounted. For double immunohistochemistry, stainings for the respective antibodies employed an avidin-biotin complex/horseradish peroxidase system (Vector, Burlingame, California, USA) and 3,3-diaminobenzidine as chromogen, and a streptavidin-biotin complex/alkaline phosphatase system as outlined above, respectively. For immunohistochemistry, positive control sections were processed simultaneously. For estimation of the proliferation index (in per cent; labelled nuclei/all nuclei counted \( \times \)100) of stellate cells and ductule cells, in acute pancreatitis and in normal controls, Ki-67/SMA and Ki-67/CK-7 double immunostains were used. In these preparation, five areas were randomly chosen, and in each area 300 nucleated cells of interest were counted and analysed for the presence of labelled nuclei. 

Statistics
All clinical data were collected prospectively and entered into a statistical package program (SPSS Statistical Software, Chicago, Illinois, USA) on a personal computer. Data were analysed using Fisher’s exact test or the Mann Whitney U test where appropriate.

RESULTS
Normal pancreatic tissue: qualitative analysis
Control tissue was structurally normal. Reactivity for CK-19 was markedly present in epithelia of large and small ducts,
Acute pancreatitis: qualitative analysis

Analysis of necrosectomy samples obtained at different time points after the onset of clinical symptoms showed that some (see below) contained spherical structures of preserved tissue. These vascularised structures, which were clearly demarcated from necrotic tissue (fig 1A), were interpreted to represent a regenerative response, and were therefore termed regenerative spheres. Some of the regenerative spheres contained labelled nuclei indicating proliferation (SMA/Ki-67 double immunostain, ×400). Regenerative spheres were observed in each of the 24 necrosectomy specimens examined in the study. Of the 24 cases of acute pancreatitis analysed histologically, 18 (75%) showed one or more feeding arteries. A characteristic zonation with formation of a gradient of ductule containing granulation tissue was present in 100% of regenerative spheres. CK-7 and CK-19 reactive ductules of varying maturity were detectable in 12/24 (50%) regenerative spheres, and in 8/12 (66.6%) spheres typical pilot ductules reaching from the centre of regenerative spheres to the periphery were observed. In 100% of regenerative spheres, increased numbers of pancreatic stellate cells/myofibroblasts and pilot ductules were in evidence, being located both in a central and peripheral position of regenerative spheres in 21/24 (88%) samples. Pilot ductules with a thick and ordered ductulocentric sheath of SMA reactive cells were present in 9/12 samples where ductules were found (75%).

For samples revealing pilot ductules, the earliest time point where such structures were in evidence was 17 days after the start of symptoms, the time periods ranging from 11 days to 41 days.

For stellate cells/myofibroblasts, the mean Ki-67 proliferation index was 22.2% (range 8–35%; normal controls 0.26%, range 0–0.6%). Increased proliferative activity was also observed in ductule cells, mainly at the periphery of regenerative spheres and the tip area of ductules (mean proliferation index 3.6%, range 1.9–5.8%; normal controls 0.13%, range 0–0.4%).

There was no correlation between aetiology, sex, C reactive protein value, APACHE II score, or extent of pancreatic stellate cell activation.

DISCUSSION

In this study we have shown that regeneration after acute necrotising pancreatitis in humans evolves in a distinct and highly ordered fashion and seems to be independent of the type of pancreatic injury causing the acute disease. The regenerative process involves pancreatic stellate cells, their differentiated myofibroblastic offspring, and pancreatic ductules originating from remnant lobules, suggesting that stellate cells/myofibroblasts and pilot ductules represent a structural and functional unit growing in parallel.

Recent evidence suggests that pancreatic stellate cells represent a key cell type for pancreatic fibrogenesis and remodelling, but their role in acute pancreatitis has not yet been clarified. Vitamin A storing cells in the pancreas were originally observed in 1982 in mice fed an excess of this vitamin but these cells were first described in the human pancreas eight years later. A potential role of this cell system in pancreatic fibrogenesis was suggested via isolation of myofibroblast-like cells from the human pancreas, and it has...
been demonstrated that vitamin A storing cells from pancreas can in fact differentiate in primary culture into myofibroblasts producing extracellular matrix proteins.\textsuperscript{15} Similar to rat periportal, but not pericentral, hepatic stellate cells,\textsuperscript{16} pancreatic stellate cells can express desmin whereas myofibroblasts derived from these cells are typically reactive for SMA.\textsuperscript{17,18} In contrast with the liver, where most but not all of the stellate cells are located within the perisinusoidal space of Disse (the so-called littoral compartment), stellate cells in the pancreas appear to be mainly situated in a periacinar location.\textsuperscript{14} Their preference for this tissue space is of interest insofar as it has been shown that hepatic stellate cells are also located close to ductules in portal tracts of the liver, thus forming an extra-littoral or extrasinusoidal compartment.\textsuperscript{19}

It has been demonstrated that pancreatic stellate cells, similar to their hepatic counterpart, play a pathogenic role in fibrosis.\textsuperscript{20} In chronic alcoholic pancreatitis, active synthesis of collagen by stellate cells appears to co-localise with lipid peroxidation derived aldehydes,\textsuperscript{21} and exposure to ethanol or acetaldehyde led to cell activation in cultured rat pancreatic stellate cells.\textsuperscript{22} Mechanisms involved in the activation of pancreatic stellate cells have been shown to include transforming growth factor \( \beta \), in part derived from activated macrophages,\textsuperscript{20,22} interleukin 1\( \beta \), and tumour necrosis factor \( \alpha \), inducing secretion of interleukin 8, monocyte chemotactic protein 1, and RANTES,\textsuperscript{24} and platelet derived growth factors.\textsuperscript{25}

It is of particular interest that pancreatic stellate cells and their activated offspring develop and grow in a non-random fashion within the regenerative spheres. On the one hand, the cells appear to evolve at the same rate as the growing regenerative tissue, forming highly oriented structures. On the other hand, they form a distinct compound structure or unit together with proliferating ductules, establishing a myofibroelastic periductular sheath. These epitheliomesenchymal sprouts are most probably recruited from pre-existing structures located in remnant lobules, and progressively extend into the growing regenerating sphere ductules thereby acting as pilot elements. A ductular reaction ensuing after acute pancreatitis, termed tubular complexes, has previously been observed in experimental models,\textsuperscript{26} and these complexes were derived from altered acinar cells proliferating 4–7 days after initiation of pancreatitis. In addition, it has been shown that pancreatic repair following trypsin induced necrohaemorrhagic pancreatitis involved proliferation of cells from intact acini and from tubular complexes.\textsuperscript{27}

These findings suggest that pancreatic stellate cells may not only be involved in fibrogenesis but also in tissue remodelling.

Figure 3  
(A) Loose regenerating tissue of a hypercellular regenerative sphere contains several desmin reactive cells representing elongated forms of pancreatic stellate cells (desmin immunostain, \( \times 400 \)). (B) Zones 1 and 2 of a regenerative sphere. Several \( \alpha \) smooth muscle actin (SMA) reactive myofibroblasts are seen with a burst-like orientation towards the interface with necrosis. One myofibroblast follows an immature blood vessel (SMA immunostain, \( \times 120 \)). (C) SMA reactive myofibroblasts situated in peripheral zones of regenerative spheres exhibit cytoplasmic projections, in part producing a stellate morphology (SMA immunostain, \( \times 400 \)). (D) A terminal (peripheral) segment of a pilot ductule (red) presents in the form of epithelial cell clusters. Note that myofibroblasts (brown) are in close contact with ductular cells (CK-7 and SMA double immunostain, \( \times 400 \)). (E) Pilot ductule (cytokeratin 7 (CK-7) immunostain; red). A periductular mantle-like spindle cell sheath is detectable, encircled by a vascular arcade (\( \times 200 \)). (F) Proximal and therefore more mature segments of pilot ductules are encircled by a compact sheath of myofibroblasts. The centre shows a pilot ductule with irregular arrangement of epithelial cells. The spindle cells forming the now compact periductular sheath are markedly positive for SMA, these cells representing myofibroblasts (double immunostain: CK-7, red; SMA, brown) \( \times 400 \).
Thus they mimic their hepatic analogues where it has recently been shown that myofibroblasts in the rat liver reflect the degree of hepatic remodelling rather than cirrhosis inasmuch as the myofibroblast volume fraction inversely reflects hepatocyte volume bimodality, suggesting that ductular complexes and stellate cells act as pacemakers in tissue remodelling.

The mechanisms operational in the phenomena observed in the present study are not known.

In conclusion, the results of this study suggest that pancreatic stellate cells and their activated myofibroblastic offspring may participate in regeneration after acute necroisising pancreatitis. Time course studies are needed to further strengthen this regeneration concept.

REFERENCES


Pancreatic stellate cells contribute to regeneration early after acute necrotising pancreatitis in humans
A Zimmermann, B Gloor, A Kappeler, W Uhl, H Friess and M W Büchler

Gut 2002 51: 574-578
doi: 10.1136/gut.51.4.574

Updated information and services can be found at:
http://gut.bmj.com/content/51/4/574

These include:
References
This article cites 28 articles, 4 of which you can access for free at:
http://gut.bmj.com/content/51/4/574#BIBL

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Topic Collections
Articles on similar topics can be found in the following collections
Pancreas and biliary tract (1949)
Pancreatitis (531)

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/