 Activation of pancreatic stellate cells attenuates intracellular Ca\(^{2+}\) signals due to downregulation of TRPA1 and protects against cell death induced by alcohol metabolites

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Alcohol abuse, an increasing problem in developed societies, is one of the leading causes of acute and chronic pancreatitis. Alcohol-related AP is initiated by toxic cytosolic Ca\(^{2+}\) overload, disrupting mitochondrial functions, and inducing cell death. However, activated PSCs acquire remarkable resistance against ethanol metabolites via enhanced Ca\(^{2+}\)-handling capacity, predominantly due to the downregulation of the TRPA1 channel. Inhibition or knockdown of TRPA1 reduces EtOH/POA-induced cytosolic Ca\(^{2+}\) overload and protects quiescent PSCs from cell death, similarly to the activated phenotype. Our results lead us to review current dogmas on alcoholic pancreatitis. While acinar cells and quiescent PSCs are prone to cell death caused by ethanol metabolites, activated PSCs can withstand noxious signals and, despite ongoing inflammation, deposit extracellular matrix components. Modulation of Ca\(^{2+}\) signals in PSCs by TRPA1 agonists/antagonists could become a strategy to shift the balance of tissue PSCs towards quiescent cells, thus limiting pancreatic fibrosis.

**INTRODUCTION**

Alcohol abuse is one of the major problems of modern societies, contributing to the development of many debilitating diseases—including acute pancreatitis (AP) and chronic pancreatitis (CP) [1, 2]. Although the incidence of pancreatic disorders is increasing globally, thus far, there is no authorised treatment available [3]. It is generally accepted that AP is initiated by toxic cytosolic Ca\(^{2+}\) overload in the pancreatic acinar cells (PACs), causing autodigestion due to premature activation of intracellular proteases and failure of mitochondrial ATP production. This results in necrosis of the enzyme-producing PACs [4, 5]. Alcohol-related AP is initiated by the powerful Ca\(^{2+}\) releasing effect of fatty acid ethyl esters (FAEEs), generated inside PACs by the non-oxidative combination of ethanol (EtOH) and fatty acids (FAs). The release of Ca\(^{2+}\) from intracellular stores, followed by Ca\(^{2+}\) release-activated Ca\(^{2+}\) entry from the extracellular fluid, causes the toxic global and sustained increase in the cytosolic Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]\_i), triggering alcohol-induced AP [4, 6, 7].

Pancreatic fibrosis, mediated by pancreatic stellate cells (PSCs) [1, 8], is a particularly frequent complication in alcoholic pancreatitis, but is much less common following other aetiologies [9–11]. In health, PSCs exist mainly in their quiescent phenotype, and are known to generate Ca\(^{2+}\) signals in response to physiological levels of bradykinin [4, 12, 13]. They become activated by mechanical stress or cytokines, e.g. transforming growth factor-beta (TGF-β) [12, 14, 15]. While activated PSCs play a role in tissue healing, prolonged inflammation can promote the persistent activation of these cells [12, 16, 17]. In such a scenario,
PSCs repopulate tissue and deposit excessive amounts of extracellular matrix (ECM) components, resulting in organ dysfunction, diabetes, malnutrition and may contribute to an increased risk of pancreatic cancer [18, 19].

The role of Ca\(^{2+}\) in PACs and its relationship to the pathogenesis of AP is well established [4], but the signals that trigger pancreatic fibrosis are less clear. Activated cells are a dominant fraction of PSCs present during inflammation [14, 19], and it has been suggested that alcohol may contribute to PSC activation [20]. However, so far, it is unknown whether EtOH also affects Ca\(^{2+}\) homeostasis in PSCs. Importantly, the progression from AP to fibrosis-associated CP is likely determined by the physiological consequences of PSC activation.

Here, we provide evidence that inducers of alcoholic pancreatitis act directly on PSCs, eliciting large and sustained Ca\(^{2+}\) responses in quiescent PSCs that cause cell death. However, as a result of significant alterations in Ca\(^{2+}\) homeostasis, activated PSCs develop remarkable resistance to alcohol metabolites, which prevents them from dying. We also explored the mechanisms and candidates responsible for this phenomenon and discovered that the loss of the transient receptor potential ankyrin 1 channel (TRPA1) in activated PSCs is the main contributing factor. Our results prompt us to review the current dogmas on alcoholic pancreatitis to include the TRPA1 channel and TRPA1-dependent Ca\(^{2+}\) signals in PSCs as important players in the pathogenesis of this disease.

RESULTS

EtOH and fatty acids induce pathophysiological Ca\(^{2+}\) responses in hPSCs

Since pathophysiological Ca\(^{2+}\) signalling in PACs directly triggers pancreatic pathology, we sought to investigate whether non-oxidative metabolites of alcohol and FAs can also affect Ca\(^{2+}\) homeostasis in hPSCs. To address this, we monitored the real-time changes in the concentration of cytosolic Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_{i}\)) of Fluo-4-loaded hPSCs (Fig. 1A), treated with EtOH and palmitoleic acid (POA) or palmitoleic acid ethyl ester (POAEE). Our cells responded to bradykinin, as was previously reported for mouse and human PSCs (Fig. 1B) [21–23]. Our new data revealed that EtOH (200 mM) induced only a modest increase in [Ca\(^{2+}\)]\(_{i}\), of hPSCs (Fig. 1C, F). In contrast to EtOH alone, simultaneous application of EtOH with POA (concentrations of EtOH/POA were: 10 mM/10 µM, 25 mM/25 µM, 50 mM/50 µM and 100 mM/100 µM) caused a large and sustained elevation of [Ca\(^{2+}\)]\(_{i}\) in hPSCs (Fig. 1D). This increase was dose-dependent and most prominent for EtOH/POA 50 mM/50 µM (Fig. 1G). EtOH/POA 100 mM/100 µM induced such a dramatic [Ca\(^{2+}\)]\(_{i}\) overload that it killed a substantial fraction of hPSCs during the experiment, leading to the seemingly decreased response areas (Fig. 1G). Similarly, treatment with EtOH/POAEE 100 mM/100 µM and 200 mM/200 µM resulted in a global and prolonged elevation of [Ca\(^{2+}\)]\(_{i}\) in hPSCs (Fig. 1E) and the higher concentration had a more prominent effect (Fig. 1H).

In order to characterise the above Ca\(^{2+}\) signals, we carried out imaging experiments in which we disrupted intracellular Ca\(^{2+}\) homeostasis either by removing extracellular Ca\(^{2+}\) or by incubation of cells with cyclopiazonic acid (CPA), a reversible inhibitor of the sarcoplasmic-endoplasmic reticulum Ca\(^{2+}\)-ATPase (SERCA) (Fig. 1) [24]. While EtOH/POA (10 mM/10 µM, 25 mM/25 µM and 50 mM/50 µM) induced elevation of [Ca\(^{2+}\)]\(_{i}\) in hPSCs in the absence of extracellular Ca\(^{2+}\), these responses were mostly modest and, unlike in the presence of extracellular Ca\(^{2+}\), returned to the baseline levels during the time course of the experiment (Fig. 1J). The magnitude of Ca\(^{2+}\) responses was very similar for all tested concentrations and did not show a clear concentration dependence (Fig. 1K). Inhibition of SERCA by CPA unmasks passive Ca\(^{2+}\) leak from the endoplasmic reticulum (ER) and led to emptying of the ER Ca\(^{2+}\) stores. After the application of 20 µM CPA, both EtOH/POA 25 mM/25 µM and 50 mM/50 µM failed to induce further Ca\(^{2+}\) signals (Fig. 1L). When the treatment was applied in reverse order, EtOH/POA 25 mM/25 µM or 50 mM/50 µM triggered a temporal [Ca\(^{2+}\)]\(_{i}\) increase, whereas a subsequent addition of 20 µM CPA did not have any effect (Fig. 1M). This suggests that EtOH/POA releases Ca\(^{2+}\) primarily from the ER store.

EtOH and POA(EE) cause activation of hPSC in vivo but not in vitro

Unlike bile-induced pancreatitis, chronic alcoholic pancreatitis is often associated with fibrosis [25, 26]. Given the above, we decided to test whether EtOH/POA is capable of activating PSCs in vivo. For this purpose, we used three different mouse models: alcoholic AP induced by EtOH (1.35 g/kg) and POA (150 mg/kg and 300 mg/kg) [27]; and two models of biliary AP, induced by tauroliothocholic acid (TC-AP) or tauroliothocholic acid 3-sulfate (TLC-S-AP) [28], all recently reviewed [29]. H/E staining (Fig. 2A, B, left panels) and histological scoring of the H/E-stained pancreatic tissue (Fig. 2C–E) confirms that all models demonstrated typical clinical characteristics of AP, such as prominent oedema, infiltration of immune cells in the parenchyma, and patchy or diffused acinar cell necrosis. POA at 300 mg/kg was substantially more harmful compared to POA at 150 mg/kg (Fig. 2A, left panel). Immunofluorescence staining for alpha-smooth muscle actin (α-SMA) was used to assess the extent of PSC activation in vivo and, therefore, the potential of the tissue to develop fibrotic complications. A scattered pattern of α-SMA staining was present in the whole tissue in the alcoholic model (Fig. 2A, right panels). In the biliary models, the staining was restricted to necrotic regions of the tissue damaged by the infusion of bile salts and was completely absent in the healthy regions (Fig. 2B, right panels).

Quantification of the fluorescence signal showed a clear, statistically significant increase of α-SMA expression in the group treated with EtOH and POA 300 mg/kg (Fig. 2F). However, in vitro experiments revealed that EtOH and FAs do not cause upregulation of α-SMA in PSCs directly, as shown by immunofluorescence staining of hPSCs fixed after 24 h treatment with EtOH/POAEE (Fig. 2G, H), Western blot for α-SMA (Fig. 2I) and comparison of ACTA2 transcript levels (Fig. 2J). The above results indicate that the activation of PSCs in vivo, as evidenced by the expression pattern of α-SMA, was not due to the direct effect of EtOH/POAEE on these cells.

Activation of hPSCs has a profound effect on pathophysiological Ca\(^{2+}\) responses

Since EtOH/POA(EE) induce substantial [Ca\(^{2+}\)]\(_{i}\) responses in hPSCs in vitro, and alcoholic AP is associated with the activation of PSCs in vivo, we next examined whether the phenotype transition could affect Ca\(^{2+}\) homeostasis and contribute to the development of the disease and its complications. To achieve this, we applied an in vitro model of hPSCs activated by TGF-β (5 ng/ml) either for 48 h (ahPSC 48 h) or 7 days (ahPSC 7 days). The phenotype of hPSCs was confirmed by immunofluorescence staining for α-SMA (Fig. 3A–C). After 48 h incubation, ~60% of cells were positive for α-SMA; and after 7 days, essentially all hPSCs were activated (Fig. 3D). Ca\(^{2+}\) elevation induced by EtOH/POA 10 mM/10 µM was almost completely abolished in ahPSCs 48 h and 7 d compared to hPSCs (Fig. 3E–H). EtOH/POA 25 mM/25 µM induced substantially reduced Ca\(^{2+}\) responses in ahPSCs (Fig. 3I–L); and a similar reduction was observed in ahPSCs treated with EtOH/POAEE 50 mM/50 µM (Fig. 3M–P). Very notable attenuation of Ca\(^{2+}\) responses also occurred in ahPSC 48 h and 7 d treated with EtOH/POAEE 100 mM/100 µM (Fig. 3Q–T) and 200 mM/200 µM (Fig. 3U–X). For most of the above treatments, Ca\(^{2+}\) signals in ahPSCs were not only reduced, but also more transient, with a tendency to return to the basal levels in the time course of the experiment.
Expression of Ca\textsuperscript{2+} channels and transporters changes in ahPSCs

Since ahPSCs show a different pattern of Ca\textsuperscript{2+} responses compared to qhPSCs, it is likely that not only the expression of cytoskeletal proteins changes upon PSC activation, but also proteins involved in Ca\textsuperscript{2+} homeostasis become up/down-regulated in ahPSCs. To test this hypothesis, we compared the expression of selected mRNA targets encoding cytoskeletal or ECM proteins as well as proteins engaged in Ca\textsuperscript{2+} transport and release between qhPSCs and ahPSCs (48 h and 7 days). Activation of hPSCs was evidenced by upregulated expression of the ACTA2 gene, encoding α-SMA (Fig. 4A). After the initial burst of ACTA2 expression in 48 h post-activation with TGF-β, the expression of ACTA2 stabilised at a somewhat lower level after 7 days (Fig. 4A). VIM (encoding Vimentin) increased marginally both in ahPSCs 48 h and 7 days (Fig. 4B). DES (Desmin) was elevated only after 7 days of incubation with TGF-β (Fig. 4C); in contrast, FN (Fibronectin) showed an increase in ahPSCs 48 h, but not in ahPSCs 7 days (Fig. 4D). The expression of several tested Ca\textsuperscript{2+} handling proteins initially decreased at 48 h, only to return to baseline levels after a
Fig. 1 ETOH/POA and ETOH/POAEE induce Ca\(^{2+}\) responses and deplete intracellular stores in hPSCs. A hPSCs loaded with Fluo-4-AM (Ca\(^{2+}\)-probe): left—green fluorescence of Fluo-4; right—transmitted light. Scale bar: 40 µM. B Sample trace showing cytosolic Ca\(^{2+}\) responses in an hPSC to bradykinin (test for the physiological phenotype). C Cytosolic Ca\(^{2+}\) responses (average traces ± SEM) in hPSCs to 200 mM EtOH (n = 42, N = 5); control (average traces ± SEM) shows that the application of extracellular solution alone (NaHEPES with 1 mM Ca\(^{2+}\), n = 30, N = 3) does not trigger any Ca\(^{2+}\) responses in hPSCs. D Cytosolic Ca\(^{2+}\) responses (average traces ± SEM) in hPSCs to different concentrations of ETOH/POA (mM/µM): 0 (Ctrl, same as in C, n = 30, N = 3), 10 (n = 21, N = 3), 25 (n = 24, N = 3), 50 (n = 29, N = 3), 100 (n = 29, N = 3). E Cytosolic Ca\(^{2+}\) responses (average traces ± SEM) in hPSCs to ETOH/POAEE (mM/µM): 0 (Ctrl, same as in C and D, n = 30, N = 3), 100 (n = 27, N = 3), 200 (n = 25, N = 3). F–H Bar charts show average response areas (±SEM) as well as individual response areas (black dots), which demonstrate an increase of Ca\(^{2+}\) above the baseline levels calculated between 200 and 800 s for all traces averaged in C, D and F, respectively. All data were compared to the same control (perfusion with the extracellular solution alone). I Schematic illustration of selected aspects of the cellular Ca\(^{2+}\) signalling machinery: cyclopiazonic acid (CPA) inhibits SERCA (sarcoplasmic reticulum Ca\(^{2+}\)-ATPase), which blocks ER and leads to depletion of this Ca\(^{2+}\) store. J Cytosolic Ca\(^{2+}\) responses (average traces ± SEM) in hPSCs to different concentrations of extracellular Ca\(^{2+}\) by different concentrations of ETOH/POA (mM/µM): 0 (Ctrl, same as in C, n = 30, N = 3), 10 (n = 14, N = 3), 25 (n = 19, N = 3) and 50 (n = 26, N = 3). K Bar chart shows the increase of Ca\(^{2+}\) above the baseline levels presented as average response areas (±SEM) and individual response areas (black dots) calculated between 200 and 800 s for all traces averaged in J. L Cytosolic Ca\(^{2+}\) response (representative trace) to 20 µM cyclopiazonic acid (CPA), an inhibitor of SERCA (as depicted in I), in the absence of extracellular Ca\(^{2+}\). After depletion of the ER stores by CPA, a subsequent application of ETOH/POA 25 mM/25 µM (presented in grey; n = 34, N = 3) or ETOH/POA 50 mM/50 µM (presented in blue; n = 41, N = 3) fails to trigger further Ca\(^{2+}\) responses in hPSCs. M Cytosolic Ca\(^{2+}\) response (representative trace) to ETOH/POA 25 mM/25 µM (presented in grey; n = 14, N = 2) or ETOH/POA 50 mM/50 µM (presented in blue; n = 10, N = 3) in the absence of extracellular Ca\(^{2+}\); subsequent treatment with CPA does not trigger further Ca\(^{2+}\) release from the ER. Statistical significance was calculated with the Mann–Whitney test (for data presented in F) and the Kruskal–Wallis test, followed by a post hoc analysis with the Dunn test (for data presented in G, H, K).

7-day incubation: STIM1 (Fig. 4E), ORAI1 (Fig. 4F), TRPC3 (Fig. 4G), PMCA4 (Fig. 4J), IP3R3 (Fig. 4M) and RYR2 (Fig. 4O). This was inversely correlated with a particularly high expression of ACTA2 in ahPSCs 48 h (Fig. 4A). TRPC6 was only elevated after 7 days after activation (Fig. 4H). ITPR1 (Fig. 4K), IP3R2 (Fig. 4L) and RYR1 (Fig. 4N) did not change much in ahPSCs; and RYR3 (Fig. 4P) was only slightly increased after 48 h. However, only one target of the tested mRNAs was consistently decreased in both ahPSCs 48 h and 7 d: TRPA1 (Fig. 4I).

ETOH/POA-induced Ca\(^{2+}\) responses and cell death in hPSCs are regulated by TRPA1

TRPA1 caught our interest since its expression in qhPSCs could contribute to [Ca\(^{2+}\)]\(_i\) overload in these cells but not in ahPSCs. To further investigate this, we inhibited TRPA1 with HC-030031 or silenced TRPA1 expression with selective siRNA in qhPSCs (Fig. 5A). siRNA decreased levels of the TRPA1 channel, as shown by immunofluorescence staining for α-SMA and TRPA1 (Fig. 5B). This downregulation was very similar to what occurred upon activation of hPSCs. Importantly, neither pharmacological inhibition nor TRPA1 suppression caused upregulation of α-SMA on its own (Fig. 5B). However, both pharmacological inhibition and silencing of TRPA1 resulted in a very significant decrease in ETOH/POA-induced [Ca\(^{2+}\)]\(_i\) elevation, almost perfectly mirroring the extent of EtOH/POA-induced cell death in qhPSCs and ahPSCs (48 h and 7 days) are reflected in and by the condition of mitochondria, we applied tetramethylrhodamine methyl ester (TMRM), a fluorescent probe that accumulates in healthy mitochondria with intact membrane potential (Fig. 6A). The application of EtOH/POA disrupted the mitochondrial potential in qhPSCs (Fig. 6B, C). This effect was relatively modest in response to low concentrations of ETOH/POA (10 mM/10 µM, no statistical significance), but ETOH/POA 25 mM/25 µM and 50 mM/50 µM caused very substantial declines in mitochondrial potential (Fig. 6C). In contrast, the effect of ETOH/POA on mitochondria was much less pronounced in ahPSCs: while ETOH/POA 25 mM/25 µM caused only marginally lower decrease in the mitochondrial potential of ahPSCs 48 h, in ahPSCs the loss of mitochondrial potential was markedly inhibited compared to qhPSCs (Fig. 6D, E). Given that both Ca\(^{2+}\) signalling and the condition of mitochondria control cell fate, we compared the extent of ETOH/POA-induced cell death in qhPSCs and ahPSCs using annexin V-FITC, propidium iodide (PI) and Hoechst 33258. Incubation with ETOH/POA at different concentrations for 30 min caused a dose-dependent increase in cell death of qhPSCs: apoptosis was the most prevalent at lower concentrations (10 mM/10 µM and 25 mM/25 µM), whereas necrosis was dominant in the group treated with ETOH/POA 50 mM/50 µM (Fig. 6F, G). However, in ahPSCs 48 h, there was a substantial reduction in total cell death (apoptosis and necrosis) in favour of living cells for all tested concentrations of ETOH/POA (Fig. 6F, G). This effect was even more pronounced in ahPSCs 7 days and was no longer dose-dependent in the range of ETOH/POA concentrations used (Fig. 6F, G). Importantly, pharmacological inhibition of TRPA1 protected qhPSCs from cell death induced by ETOH/POA 50 mM/50 µM, reducing apoptosis and necrosis to the levels seen in ahPSCs 48 h (Fig. 6F, G).

DISCUSSION

ETOH-induced damage to the pancreas occurs via products of non-oxidative alcohol metabolism: FAEEs generated from ETOH and FAs, such as POA [4, 7]. FAEEs are present in human plasma after ETOH ingestion and were found in particularly high amounts in the pancreata of subjects intoxicated with ETOH at the time of death [31, 32]. Recently, FAEEs have been suggested as candidate biomarkers of pancreatitis [33]. FAEEs are formed by esterification of FAs with ETOH catalysed by FAEE synthases, or by transesterification of ETOH to acyl-coenzyme A by acyl-coenzyme A:ethanol O-acyltransferase [34]. Since the pancreatic tissue is characterised by high FAEE synthase activity and FAEE hydrolase activity,
comparable to that of the liver [34, 35], treatment with EtOH and POA will generate POAEE intracellularly, and vice versa. FAEEs can accumulate at the inner mitochondrial membrane [36], where they become broken down to FAs and ethanol by carboxylesterase [34]. The toxicity of alcohol metabolites is generally attributed to the generation of excessive cytosolic Ca\(^{2+}\) signals in PACs. This causes premature intracellular activation of digestive enzymes and failure of mitochondrial ATP production, leading to necrosis, which in
turn leads to a generation of Ca\(^{2+}\) signals in PSCs and pancreatic immune cells [4, 7]. Although this clarifies the core mechanism of pancreatic autodigestion, inflammation, and tissue necrosis, it does not explain why PSC-mediated fibrosis is a particularly common complication of alcoholic pancreatitis. Since Ca\(^{2+}\) signals play a major role in the mechanism of EtOH-induced toxicity in PACs, in this study, we have focused on the Ca\(^{2+}\) effects of alcohol metabolites on PSCs. EtOH alone applied at high concentration only elicited a modest elevation of baseline [Ca\(^{2+}\)] in qhPSCs (Fig. 1C), which is similar to previous observations made in PACs [4, 37]. Importantly, both EtOH/POA and EtOH/POAE consistently induced global and sustained elevations of [Ca\(^{2+}\)] in qhPSCs, in a dose-dependent manner (Fig. 1D, E, G, H), suggesting that alcohol metabolites are detrimental, not only to PACs but also to PSCs. While the influx of extracellular Ca\(^{2+}\) is a significant part of EtOH/POA(EE)-induced responses, it is the release of Ca\(^{2+}\) from intracellular stores that is the initiating signal. This is illustrated by the fact that although the responses to EtOH/POA in qhPSCs...
were substantially diminished in the absence of extracellular Ca2+; they were not completely abolished (Fig. 1J, K). In PACs, the ER is the primary source of toxic [Ca2+], elevation induced by alcohol metabolites and sustained incubation of PAGCs with POAEE or POA in the absence of external Ca2+ depletes the ER of Ca2+ [38]. However, Ca2+ entry from the extracellular space through Ca2+ release-activated Ca2+ channels is an absolute requirement for acinar necrosis to occur [4]. In qhPSCs, emptying the ER with CPA prevented EtOH/POA-driven Ca2+ release from intracellular stores (Fig. 1L) and CPA failed to induce any Ca2+ responses when it was applied after EtOH/POA, suggesting that the ER stores had already been depleted (Fig. 1M). While FAEEs in PACs cause toxic elevation of [Ca2+]i predominantly through Ca2+ release from the ER via IP3Rs [4], FAs act on mitochondria and disrupt ATP production, which results in Ca2+ pump failure (SERCA, PMCA) [38]. Our results show that EtOH/POA induces diminished and transient Ca2+ signals in qhPSCs in the absence of extracellular Ca2+ (Fig. 1J), resembling the elevation of [Ca2+]i, triggered by the inhibition of SERCA by CPA (Fig. 1M). This similarity is consistent with the failure of Ca2+ ATPases due to the lack of ATP contributing to Ca2+ overload in qhPSCs exposed to EtOH/POA or POAEE in the presence of extracellular Ca2+. This notion is strengthened by the fact that EtOH/POA induces a loss of mitochondrial potential (Fig. 6B, C), which is irrevocably associated with disrupted ATP levels. Indeed, FAs were previously shown to disrupt mitochondrial metabolism and inhibit respiration by uncoupling oxidative phosphorylation and opening the mitochondrial permeability transition pore [38, 39].

In this study, we have tested the capacity of different inducers of pancreatic pathology, i.e. bile acids vs ethanol metabolites, to activate PSCs in three mouse models of AP. Our new data demonstrate a marked tissue-wide elevation in α-SMA expression in the alcoholic model (Fig. 2A, B). However, this activation is clearly not due to a direct action of alcohol metabolites on PSCs, since EtOH/POA and EtOH/POAEE failed to activate qhPSCs in vitro (Fig. 2G–J). Previously, it has been suggested that rat PSCs could be activated by EtOH in vitro [20], but this does not seem to be the case for human PSCs. This issue requires further investigation as results from cultured cells may also differ from those in acutely isolated cells.

Experimental alcoholic AP shows a different pattern from experimental biliary AP, with inflammation present in the entire tissue in the former but not latter. Although the extent of this difference in human AP is not known, the perpetuation of excess ethanol consumption will continue to drive the activation of PSCs, contributing to chronicity. The presence of cytokines and inflammatory mediators likely leads to the widespread activation of PSCs manifested as increased α-SMA expression. When neighbouring PACs are damaged and undergo necrosis, they release kallikrein and trypsin [40]. Kallikrein can induce the generation of bradykinin from its precursor, which then affects Ca2+ homoeostasis in PSCs (Fig. 1B) [41]. Furthermore, Ca2+ signals in PACs may further perpetuate necrosis and ongoing inflammation in the tissue, via mechanisms that involve the production of nitric oxide [42].

Our new results for the first time show that activated PSCs develop remarkable resistance against alcohol metabolites, evidenced by significantly reduced Ca2+ responses to EtOH/POA (Fig. 3E–P) and EtOH/POAEE (Fig. 3Q–X) as well as by a markedly decreased in cell death caused by EtOH/POA in ahPSCs compared to qhPSCs (Fig. 6F, G). This is particularly important for the pathogenesis of alcoholic pancreatitis. Activated PSCs will no longer be sensitive to alcohol metabolites present in the tissue and may continue to proliferate, activate the remaining quiescent cells, and deposit ECM components. The mechanism behind the resistance of ahPSCs to alcohol metabolites lies in the enhanced Ca2+ handling capacity, compared to qhPSCs, which depends on the expression of Ca2+ channels and transporters (Fig. 4). In this study, the most important Ca2+-handling proteins were selected for analyses, including generic proteins such as STIM1, Orai1, IP3Rs and RyRs, and those highlighted in the existing literature related to PSCs and other stellate cells/tissue fibroblasts (such as TRPA1, see below). Our results show that TRPA1 expression drops to very low levels in ahPSCs 48 h and 7 d (Fig. 4I). Pharmacological inhibition as well as silencing of TRPA1 in qhPSCs results in almost identical Ca2+ responses to EtOH/POA to those recorded in ahPSCs 48 h (Fig. 5C–F). Even more notably, the TRPA1 blocker protected against EtOH/POA-induced cell death in qhPSCs (Fig. 6F, G). Since altered Ca2+ fluxes across the plasma membrane could have an impact on the intracellular Ca2+ store content [43], downregulation of TRPA1 could contribute to the apparent decrease in the resting [Ca2+]ER of ahPSCs, unmasked by inhibition of SERCA by CPA (Fig. 5G, H). TRPA1 has previously been suggested to be protective against fibrosis in intestinal myofibroblasts, both in vitro and in a mouse model of chronic colitis [44]. Very recently, downregulation of TRPA1 by TGFB was implicated in the resistance of lung myofibroblasts to cell death [45], although the detailed mechanism has not been investigated. Here, we have identified a remarkable downregulation of TRPA1 in ahPSCs and analysed its pathophysiological consequences in relation to pancreatitis. Given the results of this and previous studies, TRPA1 could play a central role.
role in the regulation of cell death induced by toxic stimuli in a large spectrum of tissue (myo)fibroblasts.

In conclusion, this study sheds new light on the mechanisms underlying alcohol metabolite-induced pancreatic pathology, which involves signalling in PSCs (Fig. 7). Alcohol toxicity is mediated by EtOH, FAs, and FAEEs, which induce damage to PACs. Alcohol metabolites also act on quiescent PSCs inducing 

$[\text{Ca}^{2+}]_{i}$ overload, disrupting the mitochondrial potential, and inducing cell death. Alcoholic pancreatitis results in the activation of PSCs throughout the pancreatic tissue. Activated PSCs become resistant to alcohol metabolites owing to their enhanced $[\text{Ca}^{2+}]_{i}$ handling capacity compared to the quiescent cells; activated PSCs are protected against excessive $[\text{Ca}^{2+}]_{i}$ overload by downregulation of TRPA1. Significant resistance of activated PSCs to noxious stimuli (such as alcohol metabolites) might explain why pancreatic fibrosis is a frequent complication.
of alcoholic pancreatitis. While PACs and quiescent PSCs are prone to cell death, activated PSCs can more easily withstand the ongoing inflammation, repopulate the tissue, and deposit ECM components.

Our study also, for the first time, shows that EtOH in combination with FAs directly elicits Ca\(^{2+}\) signals in PSCs and that these signals can be sustained in the presence of extracellular Ca\(^{2+}\) (Fig. 1D, E, G, H). Recent work has highlighted the key role of PSCs in inflammation by showing that the SARS-Cov-2 spike protein primarily elicits Ca\(^{2+}\) signals in PSCs in situ in the mouse pancreas and that these Ca\(^{2+}\) signals subsequently generate Ca\(^{2+}\) signals in pancreatic macrophages, most likely via interleukins released from the PSCs [46]. The Ca\(^{2+}\) signals shown in the present work to be generated in PSCs by EtOH in combination with FAs could well turn out to be an important element in the initiation of the inflammatory response, which is such a critical part of AP.

**MATERIALS AND METHODS**

**Cell culture**

Human pancreatic stellate cells (hPSCs, cat. #3830) and dedicated stellate cell complete medium (SteCM) were purchased from ScienCell, Carlsbad, CA, USA. hPSCs were cultured in Stem in T25 flasks at 37 °C, 5% CO\(_2\) and split once a week as previously described [23]. Immediately after the first passage, multiple frozen hPSC stocks were prepared to be used later for culture revival (approximately every 5–6 weeks). The cells are regularly tested for mycoplasma contamination (by PCR). Activated hPSCs were obtained by culturing hPSCs in the presence of 5 ng/ml TGF-β (Corning, New York, NY, USA) in an "incomplete" SteCM medium i.e. deprived of foetal bovine serum (FBS) and cell growth supplements (CGS) for a set period of time (48 h or 7 days).

**Animal models**

Animal experiments were ethically reviewed and carried out in accordance with the Animals (Scientific Procedures) Act 1986 (UK); relevant project licence number: 30/2956. C57BL/6J mice (males,
Activated hPSCs are resistant to EtOH/POA-induced loss of mitochondrial potential (ΔΨ) as well as cell death. A Image shows mitochondrial localisation of TMRM in hPSCs. Scale bar: 10 μm. B The average traces (±SEM) show a decrease in TMRM fluorescence recorded in hPSCs in response to different concentrations of EtOH/POA: 0 (Ctrl, n = 24, N = 3), 10 (n = 14, N = 3), 25 (n = 12, N = 3) and 50 (n = 30, N = 3) mM/µM, respectively. CCCP (1 μM) was applied at the end of each experiment to attain the maximal decrease of ΔΨ. C The bar graph shows the average decrease (±SEM) below the baseline levels calculated between 200 and 800 s for the traces averaged in B. D Average traces (±SEM) show a decrease in TMRM fluorescence recorded in response to 25 mM EtOH and 25 µM POA in quiescent hPSCs (blue; n = 16, N = 3), TGF-β-activated hPSCs for 48 h (green; n = 17, N = 3) and TGF-β-activated hPSCs for 7 days (grey; n = 17, N = 3). CCCP (1 and 10 μM) was applied at the end of each experiment to attain the maximal decrease of ΔΨ. E The bar graph shows the average decrease (±SEM) below the baseline levels calculated between 200 and 800 s for the traces averaged in D. F Representative images of the staining of hPSCs with annexin V-FITC and propidium iodide after 30 min incubation with 50 mM EtOH and 50 µM POA: quiescent hPSCs (first column), quiescent hPSCs incubated with TRPA1 inhibitor HC-030031 (second column), TGF-β-activated hPSCs for 48 h (third column) and TGF-β-activated hPSCs for 7 days (fourth column). Scale bar: 10 μm. G The bar charts show the proportion of apoptotic (green) and necrotic (red) cells ±SEM (for all groups N = 3) calculated from the staining presented in F. Statistical significance was calculated for the combined dead cells (apoptotic + necrotic) for a given concentration of EtOH/POA between qhPSC vs qhPSC + HC-030031, ahPSC 48 h and ahPSC 7 days, using one-way ANOVA followed by a post hoc analysis with the Dunnett’s T3 test. The significance between qhPSC + HC-030031 and ahPSC 48 h was calculated using the t-test.
were excited with 488 nm laser light, and collected fluorescence was set in the range of 579–721 nm. Time series were recorded at 256×256 pixel resolution, two consecutive frames were averaged, and the interval between images was 2 s. Fluorescence signals were plotted as F/F₀, where F₀ was an averaged value calculated from the initial ten baseline images. The area under the curve was calculated for each trace. The average values obtained for each experimental group were presented as bar charts (mean ± SEM) with all individual points shown. For TRPA1 inhibition experiments, a 5-min preincubation with HC-030031 (Merck) was applied immediately before calcium recording. The inhibitor was present throughout the entire experiment. For all Ca²⁺ experiments, a pre-defined exclusion criterion was used: spontaneous activity higher than 3×(baseline values) before the application of any stimulus.

**RNA isolation, reverse transcription and real-time PCR**

RNA was isolated from cells cultured on 6-well plates using the Total RNA Mini Kit (A&A Biotechnology, Gdynia, Poland) according to the manufacturer’s protocol. RNA concentration was measured with an ND-1000 spectrophotometer (Thermo Fisher Scientific). Equal amounts of total RNA (1 µg) were taken for cDNA synthesis using the High Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific) and following the manufacturer’s protocol. For real-time PCR assays, SYBR Green PCR Mix (Thermo Fisher Scientific) was used. Specific gene primer pairs (Genomed, Warsaw, Poland) are listed in Table 1. The real-time PCR reaction was performed using a 7500 Fast Real-Time PCR thermocycler (Applied Biosystems, Waltham, MA, USA). Reaction conditions were as follows: 50 °C (20 sec), 95 °C (5 min); then 40 cycles of 95 °C (15 sec) and 60 °C (1 min) each. Relative gene expression was calculated as the cycle threshold value (Ct) normalised to glyceraldehyde-3-phosphate dehydrogenase (GAPDH) using the 2(−ΔΔCt) method [50].

**Protein isolation and Western blotting**

PSCs were lysed in RIPA buffer (Merck) supplemented with a protein inhibitor cocktail (Sigma) and the protein concentration was measured using a bicinchoninic acid assay (Sigma). 20 µg protein samples were separated by SDS-PAGE and then transferred onto polyvinylidene fluoride membranes (wet transfer, 1 h, 30 V) in an XCell SureLock® Blot Module (Thermo Fisher Scientific). To prevent non-specific binding, membranes were incubated in 3% (w/v) skimmed milk in TBST (0.03% (v/v) Tween 20 in Tris-buffered saline). The membranes were then incubated overnight at 4 °C with primary antibodies (A2547, V9131, Merck) in antibody buffer containing 1% (w/v) milk in TBST, washed with TBST three times and incubated with peroxidase-labelled secondary antibody.
secondary antibody (PI-2000, Vector Laboratories) in the same antibody
buffer. Detection was performed with the chemiluminescence HRP
substrate (Merck) in the Bio-Rad ChemiDoc imager.

Histopathology and histological scoring
Haematoxylin/eosin (H/E) staining was carried out as previously
described [28, 51]. Briefly, formalin-fixed paraffin-embedded tissue
sections (5 μm) were deparaffinised by heating in a dry oven (15 min,
65 °C) followed by two consecutive 10 min incubations in xylene.
Rehydration was achieved by subsequent washes in EtOH solutions with
increasing content of deionized H2O (dH2O): twice in 100% EtOH, then
96, 70 and 50% EtOH, and finally in dH2O. Sections were stained
in Mayer’s haematoxylin solution (Merck) for 12 min, and then washed
in running tap water (15 min). The sections were then incubated in 0.5%
eosin Y solution with phloxine (Merck) (90 s). Subsequently, the sections
were dehydrated in solutions with an increasing EtOH content; 70, 96
and 100% followed by washing in xylene. The slides were sealed with
Histofluid mounting medium (Paul Marienfeld, Lauda-Königshofen,
Germany). Microscopic evaluation of H/E samples was carried out by
assessing 10 random fields of view in a ‘blind’ manner. The severity of
each parameter—oedema, inflammatory cell infiltration, and acinar cell
necrosis—was scored using a four-level grade [28].

Immunocytofluorescence
hPSCs were seeded on Ø13 mm round glass coverslips and incubated for 24 h
in an ‘incomplete’ SteCM with ethanol (Chempur, Piekary Śląskie, Poland) and
POA or POEE. After washing with phosphate-buffered saline (PBS), cells were
fixed in methanol (Avantor Performance Materials, Gliwice, Poland) for 20 min
at −20 °C. Methanol was removed by three washes in PBS (5 min each) and
blocking of non-specific binding sites was achieved by incubation in 2% BSA
in PBS for 30 min (shaker, RT). Primary mouse antibody anti-αSMA (ab7817,
Abcam, Cambridge, UK) and/or rabbit anti-TRPA1 (NB110-40763, Novus
Biologicals, Biotechne) were applied in 1:300 dilution in 1% BSA for 1 h. The
fixed cells were then washed three times in PBS (5 min each) and
incubated with the secondary goat anti-mouse antibody Alexa Fluor 488 and/
or goat anti-rabbit antibody Alexa Fluor 635 (A11001, A31576, Thermo Fisher
Scientific) (1:500 dilution, 30 min, RT). After washing in PBS (four times, 5
min each) and rinsing in dH2O, the slides were sealed with ProLong Diamond
Antifade Mountant with DAPI (Thermo Fisher Scientific). Imaging was done
with a ZEISS LSM 880 confocal microscope.

### Table 1. List of primer sequences used in real-time PCR experiments.

| Target gene                      | Forward primer | Reverse primer |
|----------------------------------|----------------|----------------|
| ACTA2 (actin alpha 2, smooth muscle) | ACTGCCTTGGTGTGTGACAA | CACCATCACCCCCTGATGTC |
| VIM (vimentin)                  | AAATGGCTCGTACCTTCTGT | AGAAATCTCGTCCTCCTGC |
| DES (desmin)                    | CCAACAAGAACAACGAGGCC | ATCAGGAATCTTAGTGAGC |
| FN (fibronectin)                | TGTGGTTCCTTGACAGAT | GCTGTGAGGTCAGCTCAGT |
| STIM1                            | GCAGCAGAGTTTTGCGGAAT | TGTGAGTATCCAGACTGC |
| TRPC3                            | GCCCTTGGCCCTGATCTTTA | TCTGTAACGGGGGAAAACCT |
| TRPC6                            | ATGGCCGTCAAGTCTCCTG | ATCCTCCCATCTTGACAT |
| TRPA1                            | TAATGGGAAAGGCCCACCCCT | TCCCTTCTCCACTGGTGATA |
| PMCA4                            | GGGATGCACTGACCCAGATT | CCAAGACAGCTTTCAAAGG |
| ITPR1                            | GGAGACAGCGGTCATAGGG | CTCATTGACGGCTGGTTA |
| ITPR2                            | TTCATCATGACCCATGCCC | TCAGGATTAGCTCTGACAGT |
| ITPR3                            | AGCAATTCAGGCTACAGCA | AGCTTCCCCCTTCAACAC |
| RYR1 (ryanodine receptor type 1) | ATGCCACCTAAGCTCCTCAC | ATGCCCCAGAGAAGTTTCC |
| RYR2 (ryanodine receptor type 2) | GCATAGACCGTTTGCACGTC | AATTAGAGCCCGCAGACT |
| RYR3 (ryanodine receptor type 3) | GAGGAGAGATGCCACCAAC | ATGCTGCTACATGGCTCCG |
| GAPDH (glyceraldehyde-3-phosphate dehydrogenase) | GAAGGTGAAGGTCGGAGT | GAAGATGGGTATGGGATTTC |
Immunohistochemistry

For immunohistochemistry (IHF), a previously established protocol was used with modifications [23]. Before the staining, formalin-fixed paraffin-embedded tissue sections (5 μm) were deparaffinised by heating in a dry oven (15 min, 65 °C) followed by two consecutive 10 min incubations in xylene. The sections were rehydrated by subsequent washes in ethanol solutions as described above for H&E staining. The sections were then incubated in 50 mM NH₄Cl (Merck) in dH₂O (30 min). Subsequently, heat-induced antigen retrieval was carried out in TAE buffer (Merck) (pH 8.1) in an autoclave (20 min, 120 °C) and then the sections were allowed to cool at RT for 30 min before permeabilisation in 0.4% Triton X-100 (Avantor Performance Materials) in dH₂O (10 min). Later, the sections were washed three times (5 min each) in 0.1% Tween 20 in dH₂O ('washing solution'). In order to quench autofluorescence of the pancreas, the sections were incubated in 0.2% Sudan Black B (Merck) in 70% EtOH for 20 min and washed four times (5 min each) in a washing solution. Blocking of non-specific binding sites was achieved by 1 h incubation in 1% BSA in PBS with 0.1% Tween 20. The sections were then incubated with anti-α-SMA antibody (ab2817, Abcam, Cambridge, UK) in 1:500 dilution in blocking buffer overnight at 4 °C in a humidity chamber. The following day, the sections were washed four times (5 min each) in the washing solution and incubated for 1 h at RT with the secondary goat anti-mouse antibody Alexa Fluor 488 (A11001, Thermo Fisher Scientific) (1:500 dilution in blocking buffer, 30 min, RT). After incubation, the sections were washed four times (5 min each) in the washing solution, embedded in ProLong Diamond Antifade Mountant with DAPI and imaged immediately using the ZEISS LSM 880 confocal microscope. The slides were then stored at 4 °C. QuPath software was used for quantitative analysis of the staining [32].

Cell death assay

hPSCs were plated on 832 mm glass coverslips and grown for 24 h in SteCM or activated with TGF-beta (5 ng/mL) in an incomplete medium (SteCM without FBS/CGS) at 37 °C. The medium was then replaced with NaHPEBS buffer containing EtOH and POA of a given concentration; and the hPSCs were incubated for 30 min at 37 °C. 15 min before the end of the incubation, annexin V-FITC (1 µg/mL), propidium iodide (PI, 1 µg/mL) and Hoechst-33342 (5 µg/mL) (all from Thermo Fisher Scientific) were added. Green annexin V-FITC staining was used to detect apoptotic cells, PI stained necrotic cells, and Hoechst-stained nuclei. Multiple random images (15) per treatment group were taken using the ZEISS LSM 880 confocal microscope. Apoptotic, necrotic, and live cells were counted in images of each treatment group, and the results were presented as a percentage of all cells.

RNA interference-mediated gene silencing

hPSCs were cultured up to 50% confluence on glass coverslips. The cells were then transfected with 20 nM siRNA targeted against human TRPA1 (Thermo Fisher Scientific) pre-mixed with RNAiMAX Lipofectamine (Silencer™ Select, Thermo Fisher Scientific), according to the manufacturer’s protocol. The siRNA-lipid complex was prepared in Opti-MEM (Life Technologies) medium. The siRNA (Silencer™ Select) was added.

Statistical analysis

For cell death assays, three independent experiments were carried out for each treatment group. The average values and standard errors of the mean were calculated and the results were presented as bar charts. For quantitative analysis of Ca²⁺ responses or measurements of the mitochondrial potential, areas under individual traces (over baseline) were calculated and then averaged and presented as bar charts with standard errors of the mean and individual data points. Statistical analysis was performed with GraphPad software. For cell-based experiments, no statistical tests were applied to pre-estimate sample size. In animal models, group size was estimated based on Charan and Katharia, 2013 [49]. Normality of the data was assessed with the Shapiro–Wilks test and homogeneity of variances with the Levene test. In cases when the date did not pass the normality test, a non-parametric statistical test was applied. The significance threshold was set at the p value = 0.05 and, when applicable, was adjusted for multiple comparisons. Statistical tests used for data analysis are indicated in each figure legend. Where applicable, N indicates the number of independent experimental replicates, while n indicates individual cells.

DATA AVAILABILITY

All relevant data are included in the article. Raw data from individual experiments will be made available upon request.

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