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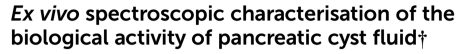
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1 Introduction

Due to its poor prognosis, pancreatic cancer (PC) is one of the world's most aggressive cancers, with an incidence rate close to its rate of mortality.¹ In 2024, PC had the lowest 5-year survival rate of any malignancy, at just 13% according to the American Cancer Society.² Indeed, PC is expected to overtake colorectal cancer as the 2nd cause of cancer-related death in next two decades.³ However, its poor survival rate is mainly due to a typically late-stage diagnosis as well as the lack of specific symp-



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Pancreatic cystic lesions (PCLs) are fluid-filled sacs often identified incidentally during abdominal imaging for unrelated pancreatic indications. While most PCLs are non-cancerous (benign) with no potential progression to pancreatic cancer (PC), some PCLs may undergo malignant transformation and are therefore more likely to progress into invasive cancer. The main challenge lies in separating PCLs that are entirely benign from those that are premalignant, as imaging tools and clinical guidelines remain inadeguate. Understanding how these PCLs arise and develop is crucial for the proper management of patients with these lesions. Well-established PDAC cell lines and pancreatic cyst fluid (PCF), the fluid contained within PCLs, represent an invaluable tool for investigating the biological processes driving PCL formation and their eventual malignant transformation. Such information could greatly improve patient risk stratification, as well as avoid unnecessary follow-up and treatment for those who do not need it. Novel approaches based on vibrational spectroscopy may represent a useful adjunct to clinical tests of this sample type, offering a mode for objective assessment as well as providing potential in vivo applications. In the present ex vivo investigation, the first of its kind in this field, a focus was brought to developing a cell line-based model with vibrational spectroscopy for discriminating the response of PDAC cell lines to exposure to PCF. We demonstrate that this approach provides a robust assay which may have potential, with further development, to provide models which are predictive of the trajectory of disease progression in precision medicine.

> toms. The two main forms of PC are pancreatic ductal adenocarcinoma (PDAC) (accounting for 85–90% of disease incidence), and pancreatic neuroendocrine tumor (PNET), which is less common (having an incidence rate less than 5%).^{4,5}

> Pancreatic cystic lesions (PCLs) are fluid-filled cavities that can be found either on or within the pancreas, and have several subtypes, including pseudocysts (non-neoplastic cysts), intraductal papillary mucinous neoplasms, mucinous cystic neoplasms, serous cystic neoplasms, and other rare cystic lesions.⁶ Since these PCLs can be either benign or precancerous, they possess variable risk of progressing to PC in any given patient.6,7 The increased identification of PCLs by cross-sectional imaging over the past few years has made their appropriate management of greater importance. Several sets of clinical guidelines are employed globally for the surveillance and management of PCLs, indicating a lack of consensus as to the clinical management of these patients.8 These guidelines differ on several cut-offs and criteria, and have been shown to perform poorly when it comes to risk stratification of patients.8 An accurate risk stratification method is hence essential to correctly identify patients with low-risk PCLs from those at increased risk of PC development and thus guide their clinical management and specify their treatment. Moreover, the diagnostic



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approaches currently adopted to discriminate between PCLs and assess their risk of malignancy fall short, underscoring an urgent need for further advanced diagnostics modalities.^{9,10}

In the past two decades, vibrational spectroscopy (VS), including Raman and FTIR spectroscopy, when combined with appropriate machine learning and chemometrics approaches, has shown promise for the detection of a variety of cancers such as lung,¹¹ breast,¹² prostate,¹³ and colorectal cancer.14 These non-invasive, sensitive, objective and reproducible techniques allow rapid label-free molecular profiling of biological samples which may be translated into clinically useful diagnostic information.¹⁵ Previous work has demonstrated the capability of VS for diagnosis of PC.¹⁶ Recent work has coupled Raman spectroscopy (RS) with KNN (K-Nearest Neighbours) and SVM (Support Vector Machine) algorithms to differentiate between tumour repopulating cells and control cells from the PDAC cell line Mia Paca-2 with high accuracy.¹⁷ Other research using surface enhanced Raman spectroscopy (SERS) combined with principal component differential function analysis (PC-DFA) investigated a diagnostic model using exosomes derived from normal and PDAC cell lines for the detection of PC.18 Further research has revealed the link between cholesterol esterification (CE) and metastasis in PC using label-free RS,¹⁹ where results demonstrated that inhibiting CE is capable of preventing PDAC cells from growing and propagating (metastasis) using a murine model. This research emphasizes the role of RS as a powerful tool to identify biomolecular variations in PC tissues and cell lines.

Szymoński et al., recently reported in two proof-of-concept studies,^{20,21} that FTIR spectroscopy and Raman hyperspectral imaging (RHM) combined with data science and machine learning approaches including PCA (Principal Component Analysis), HCA (hierarchical cluster analysis), NMF (non-negative matrix factorization), T-SNE (T-distributed stochastic neighbour embedding), and CNN (convolutional neural network), can effectively discriminate between PC subtypes, including PDAC, intraductal papillary mucinous carcinoma, and ampulla of Vater. The main outcomes of these experiments highlighted notable variations in the β -sheet secondary structure composition of proteins, as well as the DNA methylation level among the three PC subtypes. Another study revealed the potential of FTIR hyperspectral imaging to detect distinctive biochemical alterations between disease subtypes, including PDAC and PNET, from healthy and dysplastic tissues.²² Sala et al., also used Attenuated Total Reflection (ATR)-FTIR spectroscopy for detecting PC using a blood-based biopsy.²³

The primary goal of this study was to determine whether the exposure of reporter PDAC cell lines of differential phenotypes to pancreatic cyst fluid (PCF) produces biochemical alterations which are observable spectroscopically *ex vivo*. A secondary goal was to compare the ability of FTIR and Raman spectroscopy, together with linear and non-linear dimensionality reduction approaches (respectively, PCA and UMAP (Uniform Manifold Approximation and Projection)) to discriminate the response across PDAC cell phenotypes when exposed to PCF. Our *in vitro* results provide a first demonstration of the potential of spectroscopic technologies as complementary diagnostic tools, opening up a new path for prediction of the trajectory of PC using this approach.

2 Materials and methods

2.1 Cell culture and preparation to PCF treatment

Pancreatic cell lines HPNE, BxPC-3 and PANC-1 were purchased from ATCC, while H6c7 cell line was purchased from Kerafast Inc. (Boston, MA, USA). PDAC cell lines (BxPC-3, PANC-1) were maintained in an RPMI-1640 medium (Biosciences, San Jose, CA, USA) supplemented with 10% (v/v) foetal bovine serum (Biosciences) and 1% (v/v) penicillin/streptomycin (Lonza Group, Switzerland). H6c7-normal cells were maintained in keratinocyte serum free medium with L-glutamine, 5 ng mL⁻¹ human recombinant EGF and 50 mg mL^{-1} bovine pituitary extract (kit) (Bioscience, Dublin, Ireland), supplemented with 50 U mL^{-1} streptomycin. The base medium for HPNE-intermediate cells is 75% Dulbecco's modified Eagle's medium (DMEM) without glucose with additional 2 mM L-glutamine (ThermoFisher, UK), 1,5 g L^{-1} sodium bicarbonate, and 25% medium M3 base (Incell Corp., San Antonio, TX, USA). The growth of this cell line was optimized in an alternative growth medium: DMEM - high glucose supplemented with 10% (v/v) FBS, 10 ng mL⁻¹ human recombinant EGF (Bioscience, Ireland) and 50 U mL⁻¹ streptomycin (Lonza Group, Switzerland). All cells were maintained in humidified incubators at 37 °C, with 5% CO₂. With no apparent contamination, regular mycoplasma testing was carried out using the MycoAlert mycoplasma detection kit (Biosciences).

All cells were seeded in 100 μ L of serum-protein free media, in a flat-bottom 96-well plate at a density of 45000 cells per well. Cells were either treated with 5% (v/v) PCF or 5% (v/v) fresh serum-protein free media for 24 h. For H6c7-normal and HPNE-intermediate cell lines, PCF exposure appeared to be significantly cytotoxic, and cell concentration was reduced to such an extent that it was not possible to acquire spectra from a population of these cells. The remainder of the experiment focused therefore on analysis of the two PDAC cell lines. General biological and molecular details of the two PDAC cell lines properties are summarized in Table 1.

Table 1 Key biological and molecular characteristics of BxPC-3 and PANC-1 cell ${\rm lines}^{\rm 24}$

Cell morphologyEpithelialMesenchymalMetastasisNon-metastaticMetastaticDifferentiationModerate to poorPoor			
Cell morphologyEpithelialMesenchymalMetastasisNon-metastaticMetastaticDifferentiationModerate to poorPoor	Cell line	BxPC-3	PANC-1
KRAS statue Wild type mutated	Cell morphology Metastasis	Epithelial Non-metastatic	

KRAS: v-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog.

After the PCF-exposure period BxPC-3 and PANC-1 cells were then trypsinized, fixed in 4% paraformaldehyde, and washed with dH₂0. Fixed cells were maintained at 4 °C and suspended within 200 μ L of dH₂0. To enable their spectroscopic analysis, 100 μ L of each sample was deposited on CaF₂ slides using a Shandon Cytospin® 4 Cytocentrifuge (MM France) at 700 rpm for 7 min, which permits absorption of residual liquid by the sample chamber filter and then left to dry at room temperature. Details of the experimental workflow are illustrated in Fig. S1 in the ESI.[†]

2.2 Patient sample collection

A total of 20 samples per cell line were prepared in this experiment, including 5 controls (untreated cells) and 15 PCFtreated samples (treated cells), with no technical replicated were included, only biological replicates. PCF samples were collected prospectively by EUS-FNA (endoscopic ultrasoundguided fine-needle aspiration) from 15 patients with PCLs at one of the three institutions in Dublin, Ireland: Tallaght University Hospital, St Vincent's University Hospital, and St James's Hospital. This work was performed in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans. Patients provided informed consent for sample and data acquisition, and the study received full ethical approval from Tallaght University Hospital Joint Research Ethics Committee Review Board (ID: 0319-264). The Research Ethics Committee of TU Dublin also provided approval for conduct of the study (REC-20-64).

2.3 FTIR microspectroscopy

After visual inspection, cell aggregates were selected, and their infrared spectral images were acquired in transmission mode in the region between 4000–800 cm⁻¹ using a Spotlight 400 FTIR imaging system (PerkinElmer, France), which was purged with CO₂-free dry air and equipped with a liquid nitrogencooled MCT (mercury-cadmium-telluride) detector. FTIR images were acquired at a spatial resolution of 6.25 μ m² and a spectral resolution of 4 cm⁻¹, with each spectrum acquired using 128 scans.

A background spectrum corresponding to the CaF_2 window, and the atmospheric environment was recorded with 240 accumulations and automatically subtracted from each spectrum within the PerkinElmer SpectrumIMAGE software.

2.4 Raman microspectroscopy

Raman images were captured *via* an XPlora microspectrometer (HORIBA, France) over the 600–1800 cm⁻¹ fingerprint region using an exposure time of 15 s and a pixel size of 2 μ m². The spectrometer was equipped with a 532 nm laser diode excitation source, delivering an output power of 64 mW at the sample. Spectra were recorded using a 25% laser filter, with laser light focused on the sample using an Olympus MS Plan 100 lens with a numerical aperture of 0.95 and a focal length of 180 mm. The grating system was set at

1200 lines per mm, with a confocal hole of 300 μ m, and a slit size of 200 μ m.

2.5 Preprocessing of FTIR and Raman spectra

FTIR data preprocessing. FTIR images were first subjected to a background removal using K-means clustering technique with a number of clusters fixed at 2, in order to extract cell spectra and eliminate spectra from CaF₂ slides.

Next, the images were corrected for atmospheric contributions, including water vapor and carbon dioxide. Following that, a resonant Mie scattering correction was employed using 10 iterations, 14 principal components of the CRMieSC (Clustered Resonant Mie Scattering Correction) algorithm,²⁵ taking Matrigel as the reference spectrum. To correct remaining distortion of the FTIR spectral baseline, the Concave Rubberband method was employed using 5 iterations. Lastly, the spectra were smoothed using the Savitzky–Golay filter with a 5th polynomial order and a window size of 15, and then vector normalized prior to spectroscopic analysis.

Due to the heterogeneity of the samples which led to diverse spectral distortions, the adjustments were performed separately for each image using the OCTAVVS software v0.1.24.²⁵ An example of the preprocessing workflow is provided in Fig. S2 in the ESI. \dagger

Raman data preprocessing. Prior to data collection, the Raman spectrometer was calibrated using silicon at its distinct peak at 520.7 cm^{-1} . The preprocessing of Raman spectra included a quality test performed in SpectraGryph 1.2-spectroscopy software v1.2.16.1, which consists of eliminating the saturated spectra (spikes removal) and those with a poor signal. The spectra were baseline corrected with the Rubberband method, then smoothed using the Savitzky–Golay filter with fifth polynomial order and a window size of 13 and then vector normalized.

The procedure of preprocessing was carried out mostly within the Orange Data mining software v3.36.1.

2.6 Multivariate statistical analysis

Unsupervised exploratory analysis. • Principal components analysis (PCA) is a linear data exploration method, also considered a dimensionality reduction technique, which is used to represent the original data in a reduced space known as the space of principal components (PCs). PCs are linear combinations of the initial variables that extract the maximum amount of chemical, biological or physical information while maximizing the total variance of the data.²⁶

• Uniform manifold approximation and projection (UMAP) unlike PCA, is a non-linear manifold learning technique for dimensionality reduction. In terms of visualization quality, the UMAP preserves a larger portion of the global structure of a dataset due to its faster runtime compared to PCA or T-SNE. Moreover, UMAP may also be used as a dimensionality reduction methodology for data pre-treatment prior to input into machine learning algorithms since it does not impose any limitations on the embedding dimension.²⁷

3 Results

3.1 Characterization of the mean Raman spectra of BxPC-3 and PANC-1 cells

Raman imaging was carried out on fixed PDAC cells, BxPC-3 and PANC-1, to interrogate the capability of Raman spectroscopy in differentiating between BxPC-3 and PANC-1 cell lines, prior to PCF exposure, as well as to highlight the effect of exposing these cells to PCF. The average normalized Raman spectra were compared and highlighted in Fig. 1. The main bands present in each cell type and their corresponding assignments reflecting their biochemical profiles are displayed in Table 1.

Visually, the spectral differences between untreated and PCF-treated cells for both cell types can be seen principally in four distinctive regions denoted by the grey shaded areas. The first region (area 1) between 600–700 cm⁻¹ is mainly attributed to DNA and RNA vibration bands and a few amino acids bands, particularly ring breathing vibrations from DNA bases (C–C stretching and twisting vibrations of phenylalanine, tyrosine and guanine) and the C–C twisting mode of phenylalanine and tyrosine linked to some protein signatures. The second region (area 2) between 1200–1360 cm⁻¹ is dominated by protein signatures, notably amide III bands, the spectral signatures originating in the CH₃/CH₂ twisting mode of lipids, and the primary and secondary structures of proteins such as α -helix, β -sheet, and random coils. The third region (area 3) between 1400–1490 cm⁻¹ is attributed to the CH₂ bending

mode of lipid and proteins. Lastly, the spectral variations of the fourth region (area 4) are marked by signatures of carotenoids principally attributed to β -carotene present in the two cell types after PCF treatment (Table 2).²⁸

3.2 Characterization of the mean FTIR spectra of BxPC-3 and PANC-1 cells

FTIR imaging was carried out on aggregate fixed cells of each PDAC cell line, BxPC-3 and PANC-1, with typical spectral images and mean spectra depicted in Fig. 2. Visual images of PDAC cells are displayed in Fig. 2A on the top left for BxPC-3 and the bottom left for PANC-1, and their corresponding FTIR images are shown on the right. In Fig. 2B, the normalized mean spectrum is depicted for each cell line before and after PCF treatment. To enhance the spectral variations between the two cell lines, the second derivative spectra was computed and is depicted in Fig. 2C. Furthermore, the attributions of the main bands are displayed in Table 3.

As reported in Fig. 2B, the spectral profiles of the two cell lines, BxPC-3 and PANC-1, before and after exposure to PCF were nearly identical, with slight variations observed mainly in four areas over the 1800–900 cm⁻¹ fingerprint region. The first region (area 1) between 930 cm⁻¹ and 1100 cm⁻¹ is related to vibrations of DNA/RNA and carbohydrates, with the bands at 965 cm⁻¹ for the C–C and C–O stretching bands in deoxyribose of DNA, 1082 cm⁻¹ for the symmetric phosphate stretching PO₂⁻ band of DNA and RNA, and the peak at 1058 cm⁻¹ for the C–O stretching and bending of carbohydrates. The second

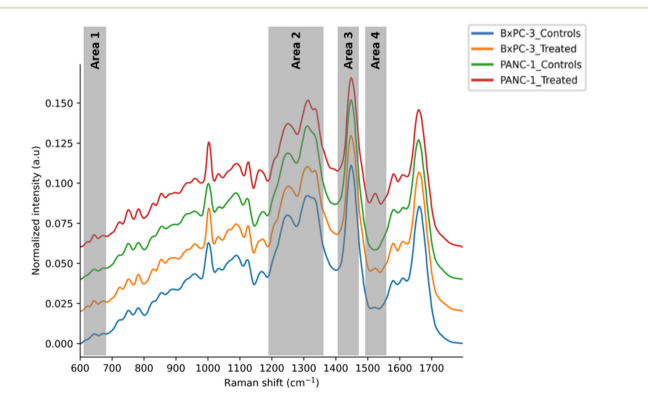


Fig. 1 Average Raman spectra of BxPC-3 and PANC-1 before and after PCF treatment.

Raman shift

Table 2 Peak assignments of Raman spectra of PDAC cell lines^{28,25}

plot shows a strong phenotypic discrimination with minimal overlap between the two cell groups, principally along (PC1, PC3) and (PC3, PC5) planes. In contrast, no clear separation is observed in the (PC1, PC2) score plot as shown in Fig. S3b.† While PC1 and PC2 capture most of the spectral variance around 70% combined, in which the two cell lines do not show any phenotypic differences due to a possible shared biochemical characteristic. The 3rd component (PC3) on the other hand with 10.54% of the explained variance (low compared to PC1 and PC2) exhibits high phenotype discrimination, as a result, the features extracted by this component are likely responsible for this separation. This loading PC3 is predominated by numerous key Raman bands, including the C-C symmetric breathing of tryptophan at 747 cm^{-1} , the C-C aromatic ring stretching phenylalanine at 1006 cm⁻¹, the amide III and I vibrations at 1311 cm⁻¹ and 1671 cm⁻¹, and the C=C in-plane bending mode of phenylalanine, tyrosine and cytosine at 1632 cm⁻¹. The band at 1585 cm⁻¹ is attributed to the ring breathing modes in the DNA bases, and the band at 1441 cm⁻¹ is linked to lipid bands (CH₂ bending mode).

UMAP analysis of Raman data from untreated cells. A further UMAP analysis was carried out on pre-processed Raman spectra of untreated PDAC cells (BxPC-3, PANC-1) using Euclidean distance as the distance metric with the number of neighbours set to 20. The intention here was to see whether UMAP as a non-linear unsupervised technique is capable of distinguishing BxPC-3 cells from PANC-1 cells before exposing them to PCF using Raman spectral data. The results are depicted in Fig. 5.

The 2D UMAP plot highlights clearly the differences between BxPC-3 and PANC-1 cell lines prior to PCF treatment using only two UMAP components (unlike PCA which required 5 PCs), with strong separation between the two groups, low intra-group variability and sufficient inter-group variability to allow group discrimination.

PCA and UMAP of FTIR data from untreated PDAC cells. The analytical pipelines used with Raman data were applied to second derivative FTIR spectra truncated to the fingerprint region between 1800-900 cm⁻¹ for both BxPC-3 and PANC-1 cell lines. The scores plot for the first five PCs are displayed in Fig. 6, while the corresponding loadings are depicted in Fig. S4b.†

In contrast with the Raman spectral data, the variance explained by FTIR data over the first five PCs (almost 60%) is relatively low (Fig. S4a†), and as depicted in Fig. 6, there remains significant overlap between the two groups along the two PCA planes (PC1, PC5) and (PC2, PC5). Although, PC5 component only account for 4.16% of explained variance, it appears to be able to distinguish between the two groups of PDAC cells.

Again, UMAP analysis (using Euclidean distance as the distance metric, and with the number of neighbours set to 20) improves slightly the discrimination between the cell phenotype as shown in Fig. 7, though with the loss of interpretive information provided by PCA loadings.

(cm^{-1})	Peak assignments	
620	C–C twisting mode of phenylalanine (proteins)	
642	C–C twisting mode of tyrosine and phenylalanine (proteins)	
670	Ring breathing vibrations of guanine and tyrosine (DNA/RNA)	
723	C–C symmetric stretch vibration A-DNA (ring breathing mode of DNA/RNA)	
751	C-C symmetric breathing of tryptophan (protein)	
782	Uracil, thymine, cytosine (ring breathing modes in the DNA/RNA)	
827	O–P–O stretching in DNA, tyrosine, proline	
852	Proline, hydroxyproline, tyrosine ring breathing modes (proteins)	
957	Carotenoids, cholesterol (symmetric stretching vibration)	
1002	C–C aromatic ring stretching phenylalanine (proteins)	
1031	C–H bending mode in phenylalanine, C–N stretching of proteins	
1061	C-C stretching in lipids, PO_2^- stretching (DNA/RNA)	
1125	C–C, C–O stretching mode in carbohydrates, β-carotene	
1173	Cytosine, guanine, tyrosine, phenylalanine (bending mode in plane of proteins)	
1206	Amide III (proteins), C- C_6H_5 stretching mode in tyrosine, phenylalanine, and tryptophan	
1247	Amide III (collagen assignment, β -sheet & random coils), guanine, cytosine (NH ₂)	
1313	CH_3CH_2 twisting mode of lipids, amide III (α -helix)	
1334	Guanine, CH ₃ CH ₂ wagging of nucleic acid	
1447	CH ₂ bending mode of proteins & lipids	
1524	Carotenoids (β-carotene)	
1578	Ring breathing modes in the DNA bases guanine, adenine	
1605	C=C in-plane bending mode of phenylalanine & tyrosine, cytosine (NH_2)	
1661	Amide I vibration mode of structural proteins	

region (area 2) between 1250 cm⁻¹ and 1390 cm⁻¹ is associated with signatures of protein with the bands at 1234 cm⁻¹ and 1310 cm⁻¹ associated with the amide III bond vibration. Moreover, the peak at 1390 cm⁻¹ is associated with protein, specifically the CH₃ symmetric deformation mode. The third region (area 3) between 1450 cm⁻¹ and 1650 cm⁻¹ is dominated by protein signatures from the amide I (1650 cm^{-1}) and II bands (1550 cm^{-1}) ; these bands are suggestive of changes in the secondary structure of proteins (α -helix, β -sheet). Lastly, the fourth (area 4) between 1700 and 1800 cm⁻¹ is mainly associated with vibrations of lipids, the C=O stretching and the bands associated with fatty acid esters.

BxPC-3 vs. PANC-1 prior to PCF treatment 3.3

PCA of Raman data from untreated cells. PCA was conducted on pre-processed Raman spectra of PDAC cell lines PANC-1 and BxPC-3 prior to their exposure to PCF in order to elucidate whether Raman spectral signatures discriminate the cell phenotypes.

PCA scores and loadings are displayed respectively in Fig. 3 and 4, with the first five component explaining 90% of the total variance present in Raman data (Fig. S3a⁺). PCA scores

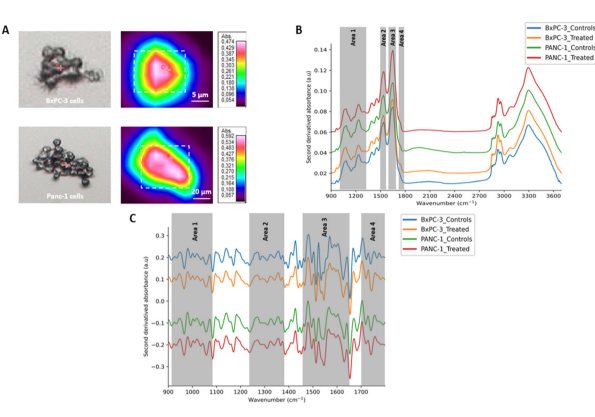


Fig. 2 FTIR imaging of PDAC cell lines BxPC-3 and PANC-1. (A) White light image of BxPC-3 (top left) and PANC-1 (bottom left) and their corresponding FTIR images on the right; (B) average FTIR spectra of BxPC-3 and PANC-1 before and after PCF treatment extracted from the marked square; (C) normalized second derivative spectra of each cell type. Spectra are shifted manually for visibility.

Table 3 Peak assignments of FTIR spectra of PDAC cell lines ³
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Wavenumber (cm ⁻¹)	Peak assignments
965	C–C and C–O stretching mode in deoxyribose of DNA
1058	C–O stretching & C–O bending mode of carbohydrates
1082	Symmetric phosphate stretching (PO ₂ ⁻) (DNA/ RNA)
1121	C–O stretching mode (proteins, carbohydrates)
1171	C–O stretching of proteins
1234	PO ₂ ⁻ asymmetric stretching (DNA/RNA), amide III
1310	Amide III
1390	CH_3 symmetric deformation (proteins)
1452	Asymmetric CH ₃ bending modes of proteins
1550	Amide II (proteins)
1650	Amide I (proteins)
1742	Fatty acid esters, C=O stretching mode of lipids
2853	CH ₂ symmetric stretching (lipids)
2872	CH ₃ symmetric stretching (lipids)
2925	CH_2 asymmetric stretching (lipids)
2958	CH ₃ asymmetric stretching (lipids, proteins)
3073	Amide B
3300	Amide A (N-H stretching in proteins)

3.4 BxPC-3, PANC-1 post exposure to PCF

PCA of Raman and FTIR data. A similar interrogation pipeline was used to interrogate the effect of PCF exposure on both BxPC-3 and PANC-1 cell lines. Firstly, the scores for PCA decomposition for both FTIR and Raman data are depicted in Fig. 8 and their corresponding loadings are provided in Fig. 9. The PCF-treated spectra are not discriminated from their controls in either PDAC cell line, and the spectral discriminants for each cell line are distinct, with scores being discriminated along PC3 for the BxPC-3 and along PC1 in the case of the PANC-1 line.

The scores along PC3 (explaining 8.41% of the variance) for BxPC-3 cells are associated with loadings to nucleic acids bands at 748 cm⁻¹ (which is attributed to the C–C symmetric breathing of tryptophan), and at 782 cm⁻¹ related to the ring breathing modes of uracil, thymine, and cytosine in the DNA/RNA bases. Additionally, the spectral bands at 1120 cm⁻¹ and 1200 cm⁻¹ are respectively associated with C–C and C–O stretching mode in the carbohydrate and protein structural bands (amide III vibration), with a negative loading at 1670 cm⁻¹ attributed to the amide I band. Lastly, lipid loadings in PC3 with the spectral features at 1377 cm⁻¹ and 1484 cm⁻¹ attributed to CH₃CH₂ twisting mode and CH₂ bending mode of lipids signatures.

Contrastingly, the loadings to PC1 captured 72.3% of the variance, with discriminating bands mainly associated to carotenoids signatures with the three specific bands of β -carotenes at 1011 cm⁻¹, 1158 cm⁻¹, and 1524 cm⁻¹ all with positive contribution to the formation of PC1.

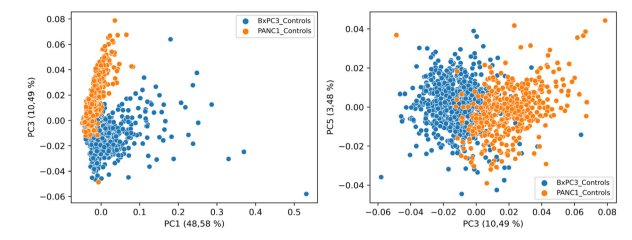
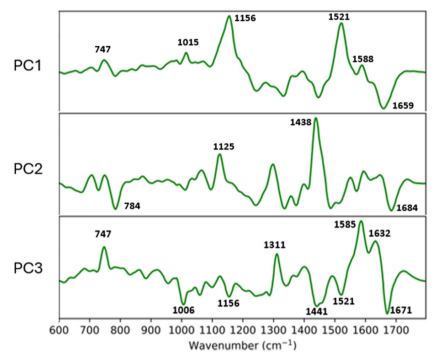
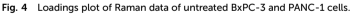


Fig. 3 Scores plot of Raman data of untreated BxPC-3 and PANC-1 cells.





As the two cell lines were exposed to the same PCF samples, the PCA scores plot of Raman data highlight different clustering across the two cell types and hence different loadings, this potentially suggests a cell type specific response to PCF treatment.

Again, for the FTIR, PCA was applied on the second derivative FTIR spectra over the 900–1800 cm⁻¹ fingerprint region for both cell lines, with the scores plot depicted in Fig. 10 and their loadings displayed in Fig. S5.† Overall, poor discrimination of the cell line response to PCF was observed due to the high intra-group variability in both PDAC cell types leading to poor explained variance by the initial PCA components. UMAP analysis of Raman and FTIR data. A further UMAP analysis was conducted on pre-processed Raman spectra and the second derivative FTIR spectra over the fingerprint region $1800-900 \text{ cm}^{-1}$ to highlight the impact of exposing BxPC-3 and PANC-1 cells to PCF (using Euclidean distance and the number of neighbours set to 20). UMAP scores plot for both Raman and FTIR data are grouped in Fig. 11.

Raman-based UMAP analysis illustrates more accurately the impact of the PCF treatment on PDAC cells (BxPC-3 and PANC-1) than the PCA does. Indeed, UMAP scores plot highlighted spectral distinction within the different PCF-treated samples for both BxPC-3 and PANC-1, where some treated samples were clearly distinguished from the untreated ones,

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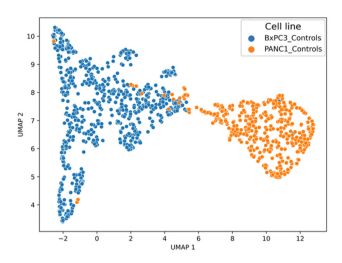


Fig. 5 UMAP plot of Raman data of untreated BxPC-3 and PANC-1 cells.

while for other treated cells no distinction with their counterpart was noticed. Moreover, PDAC cells appear to respond differently to PCF giving the results of UMAP clustering.

Unlike Raman, the impact of the PCF treatment on the cells is less noticeable in FTIR spectra, the discrimination of the cell line response to exposure to PCF appears to be more challenging. These results suggest a high intragroup variability within the treated groups for the two cell types with a low intergroup variability mainly due to baseline distortion.

3.5 BxPC-3 vs. PANC-1 pre- and post-exposure to PCF

Combined PCA and UMAP analysis of Raman data. To further investigate the ability of vibrational spectroscopy particularly Raman technique to differentiate PDAC cell phenotype pre-exposure and post-exposure induced-PCF changes on cell biology, combined PCA and UMAP analysis of the four groups was performed on pre-processed Raman data. PCA

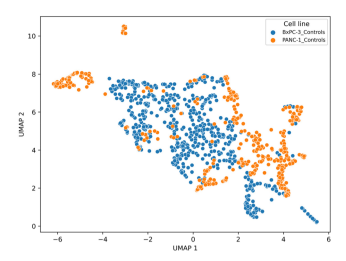


Fig. 7 UMAP plot of FTIR data of BxPC-3 and PANC-1 cells preexposure to PCF.

scores plot are highlighted in Fig. 12 and their loadings displayed in Fig. S6,† 2D and 3D UMAP scores plot are depicted in Fig. 13.

According to PCA scores plot, the first two PCA components (PC1 = 60.57%, PC2 = 14.46%) capture almost 75% of the total variance within Raman data. BxPC-3_treated group appeared distinct and separated from BxPC-3_controls along PC2 axis in both (PC1, PC2) and (PC2, PC3) planes, indicating spectral alterations induced by the treatment. In contrast, both controls and PCF-treated PANC-1 spectra show minimal spectral differences and largely overlap, suggesting a less noticeable PCF-treatment effects in this cell line. On the other hand, Ramanbased UMAP analysis highlight more effectively subtle spectral variations, that PCA unable to detect, particularly in PANC-1 groups. UMAP provides better class discrimination in complex datasets, reinforcing the capacity of Raman spectroscopy to detect delicate biochemical changes induced by PCF-exposure.

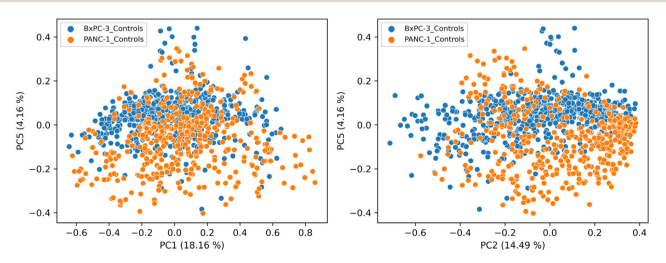


Fig. 6 Scores plot of FTIR data of untreated BxPC-3 and PANC-1 cells.

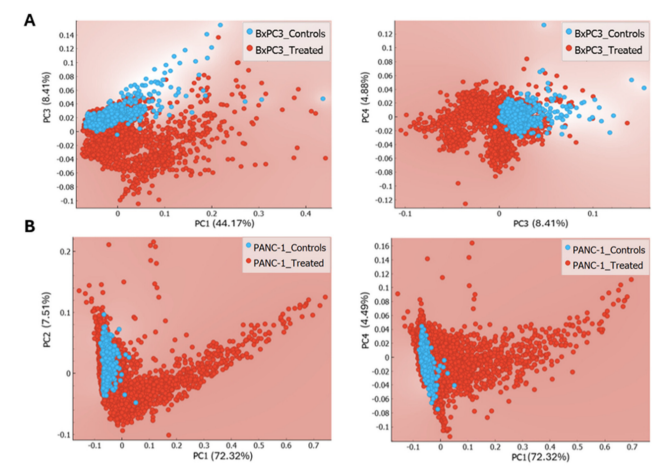


Fig. 8 Scores plot of Raman data of BxPC-3 (A) and PANC-1 cells (B) post exposure to PCF.

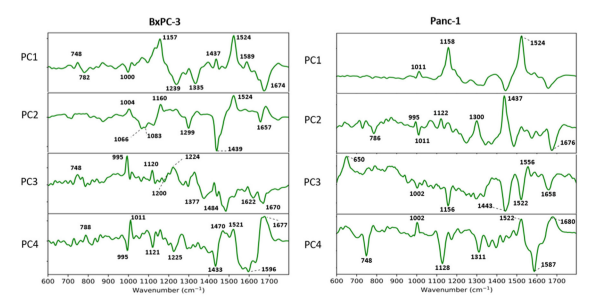


Fig. 9 Loadings plot of Raman data of BxPC-3 and PANC-1 cells after exposure to PCF.

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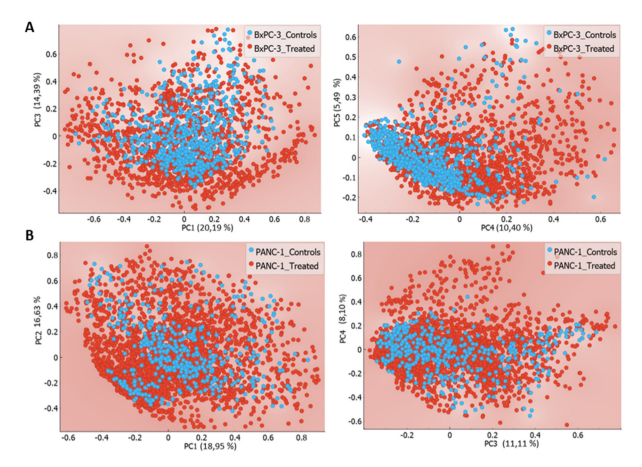


Fig. 10 Scores plot of FTIR data of PDAC cells for BxPC-3 (A) and PANC-1 (B) following their exposure to PCF.

4 Discussion

As PC is usually discovered at a late stage it remains one of the most challenging malignancies for early detection, despite recent advancement in imaging technologies and rising clinical expertise. Patients with PCLs are considered at increased risk of developing PC, as some PCLs can be pre-cancerous and may progress to invasive malignancy, finding and monitoring these PCLs are essential for early PC detection,^{31,32} However, current diagnostics tools lack sufficient sensitivity and specificity to accurately distinguish between harmless and potentially pre-malignant PCLs.^{33,34}

Although, the use of PCF for malignancy risk stratification appears promising it remains at first stages. Prior research about the application of spectroscopy for pancreatic cancer diagnosis was primarily used for analysing pancreatic tissues,^{20–22} serum/plasma,³⁵ or exosomes¹⁸ from healthy controls and PDAC patients or cell line. To our knowledge, no research has ever looked at the biological effects of PCF on the epithelial cells that surrounds PCLs. Thus, the role of PCF in the potential PCL to PC malignant transformation remains unclear and unexplored. Our approach addresses uniquely for the first time these aspects by applying FTIR and Raman spectroscopy combined with multivariate data analysis tools such PCA and UMAP techniques, to characterize these PCF-induced biomolecular changes in order to improve the risk stratification of PCL patient.

In the present work, BxPC-3 and PANC-1 PDAC cell lines were exposed ex vivo to PCF from patients with PCLs. The effect of exposure to PCF was elucidated spectrally using FTIR and Raman microspectroscopy together with dimensionality reduction methods, which demonstrated, on the one hand, the capacity of these techniques to discriminate PDAC cell phenotypes and, on the other hand, highlighted their response to PCF exposure. While our original intention was to profile the response across both normal and PDAC cell lines, we feel that the choice of PC cell lines as a reporter model here is better overall as this is a system that is primed to expand strongly to stimulants within the PCF. Importantly, Raman-based PCA and UMAP projections revealed a strong basal-level separation between BxPC-3 and PANC-1 prior to PCF-treatment. This separation is likely associated to their molecular, proteomic, and metabolic differences.^{36,37} For instance, BxPC-3 cells characterised by a wild-type KRAS and mutant SMAD4, whereas PANC-1 cell possess mutations in KRAS, CDKN2A, while retaining wild-type SMAD4. These genetic differences are also associated

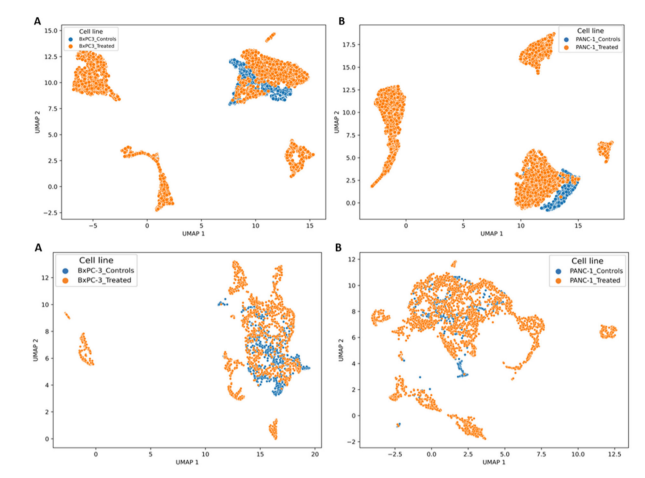


Fig. 11 UMAP plot of FTIR data of PDAC cells (A) BxPC-3 and (B) PANC-1 following exposure to PCF.

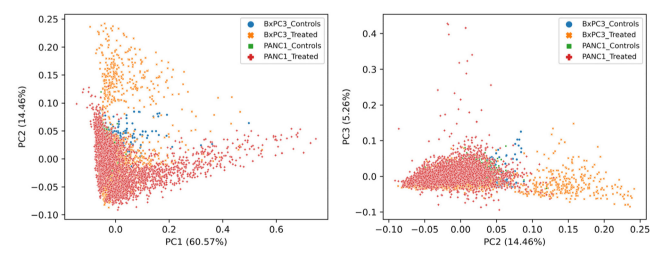


Fig. 12 Scores plot of Raman data for BxPC-3 and PANC-1 before and following their PCF-treatment.

to a differential epithelial to mesenchymal transition (EMT) profiles, with BxPC-3 exhibit more epithelial features while PANC-1 harbour mesenchymal characteristics.^{24,36}

In order to examine PCF-induced cellular responses, our work combined two complementary spectroscopic approaches

(FTIR, Raman) with dimension reduction tools such as PCA and UMAP. While PCA, and UMAP were both effective in reducing the data dimension and identifying clustering patterns, there were some discernible differences between their findings. PCA as a linear technique allow the capture of linear vari-

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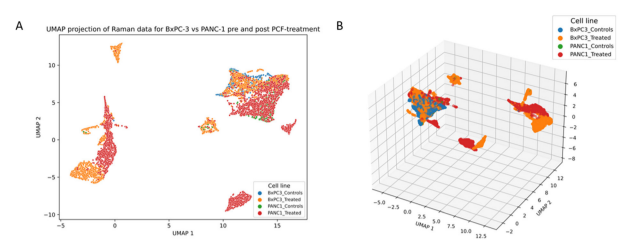


Fig. 13 UMAP plot of Raman data for BxPC-3 and PANC-1 before and following PCF-treatment. (A) 2D-UMAP projection. (B) 3D-UMAP projection.

abilities inside the data linked the cell phenotype on the one hand and the cell response to PCF on the other hand, although the discrimination between the two cell lines postexposure to PCF wasn't completely achieved especially for FTIR data with strong overlapping between the two groups. In contrast, UMAP projections was more efficient than PCA at detecting non-linear variabilities and smaller group separations particularly for Raman data as it preserve both the global and local structure of the data.²⁷ This demonstrates the increasing utility of UMAP analysis over PCA in spectroscopy data exploration. Nevertheless, the superiority of Raman over FTIR was also noticed and elucidated in this study in capturing biomolecular variations between each cell line (BxPC-3 and PANC-1) before and following the PCF treatment. Similarly, between the two complementary techniques, Raman frequently outperformed FTIR in capturing PCF-induced alterations to the cell biology. Raman high spatial resolution and capability of detecting differences at the intracellular level might explain its superior performance over FTIR. While FTIR remains beneficial and helpful for global biomolecular profiling, it showed poor class discrimination compared to Raman.

While these preliminary findings appear promising and provide insights into how PCF alters the spectral profile of PDAC cells ex vivo, further investigations are still required to overcome the current limitations and allow advancement to clinical use. One of the primary limitations is the limited sample size as only 15 PCF patient samples were used to treat the cells. Expanding the sample size using larger cohort with diverse ethnicities and various PCLs subtypes is crucial to capture full PCF variability across a variety of PCLs and enhance the statistical validity. Additionally, the current methodology involved recording spectra from remaining cancer cells after 24 h of exposure to a low PCF concentration (5% v/v). Increased concentrations of PCF with a prolonged exposure time are still unexplored and may not yield to similar findings and remain to be elucidated. Interestingly, PCF can be extremely cytotoxic to the cells even at reduced concentrations leading to a significant cell death. As such, PCF

samples included in this study should considered low cytotoxic and may not represent full spectrum of PCF effects on cancer cell biology.

Future work should investigate the biological effects of PCF on non-cancerous cell lines such as H6c7 and HPNE cell lines and its potential role during the PCL to PC malignant transformation. Multi-omics integration could greatly improve the biochemical interpretation of the detected spectral markers. Moreover, longitudinal investigations assessing the effects of PCF on normal cells may allow the examination of the ability of spectroscopy to monitor PCL progression overtime and assess its potential in the early detection of PC. By addressing the above aspects PCL risk classification and patient quality of life may improve tremendously.

5 Conclusion

Our approach demonstrates the capacity of vibrational spectroscopy particularly Raman technique in conjunction with unsupervised dimensionality reduction tools to discriminate PDAC cell phenotype and highlight PCF-induced spectral alterations on PDAC cell biology *ex vivo*. This approach, upon further validation, could have potential applications as a novel first-stage diagnostic screening method for PC and PCL risk stratification.

Conflicts of interest

The authors declare no conflicts of interest.

Data availability

Data for the control cell lines referred to in this article will be made available on Zenodo at the time of publication. Data collected from human participants, described throughout the publication are not available for confidentiality reasons.

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