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Title	An investigation of the RWPE prostate derived family of cell lines using FTIR
	spectroscopy
Туре	Article
URL	https://clok.uclan.ac.uk/4391/
DOI	##doi##
Date	2010
Citation	Baker, Matthew orcid iconORCID: 0000-0003-2362-8581, Clarke, C., Démoulin, D., Nicholson, J. M., Lyng, F. M., Byrne, H. J., Hart, C. A., Brown, M. D., Clarke, N. W. et al (2010) An investigation of the RWPE prostate derived family of cell lines using FTIR spectroscopy. The Analyst, 135 (5). pp. 887- 894. ISSN 0003-2654
Creators	Baker, Matthew, Clarke, C., Démoulin, D., Nicholson, J. M., Lyng, F. M., Byrne, H. J., Hart, C. A., Brown, M. D., Clarke, N. W. and Gardner, P.

It is advisable to refer to the publisher's version if you intend to cite from the work. ##doi##

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An investigation of the RWPE prostate derived family of cell lines using FTIR spectroscopy

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Received 30th September 2009, Accepted 10th February 2010 First published as an Advance Article on the web 4th March 2010 DOI: 10.1039/b920385k

Interest in developing robust, quicker and easier diagnostic tests for cancer has lead to an increased use of Fourier transform infrared (FTIR) spectroscopy to meet that need. In this study we present the use of different experimental modes of infrared spectroscopy to investigate the RWPE human prostate epithelial cell line family which are derived from the same source but differ in their mode of transformation and their mode of invasive phenotype. Importantly, analysis of the infrared spectra obtained using different experimental modes of infrared spectroscopy produces similar results. The RWPE family of cell lines can be separated into groups based upon the method of cell transformation rather than the resulting invasiveness/aggressiveness of the cell line. The study also demonstrates the possibility of using a genetic algorithm as a possible standardised pre-processing step and raises the important question of the usefulness of cell lines to create a biochemical model of prostate cancer progression.

Introduction

Cell lines are powerful models for Fourier transform infrared (FTIR) spectroscopic studies due to the relatively greater phenotypic homogeneity than their corresponding heterogeneous tissue and primary cell specimens.¹ Prostate cancer (CaP) cell lines have been successfully discriminated based on their infrared (IR) spectra² as well as their Raman spectra.^{3–5} However, these studies have utilised cell lines from different anatomical positions so it is arguable as to whether the spectroscopic discrimination was due to the malignancy or to the different origin of the cell lines. As the cell lines have been exposed to different environments with different levels of biomolecular compositions, the environmental effect on the cellular biochemistry cannot be controlled.

Such environmental factors can be reduced by using cell models comprising of a family of cell lines derived from a single source but with differing phenotypes/characteristics. Here we present data utilising the RWPE prostate epithelial cell line family. Epithelial cells derived from the peripheral zone of a histologically normal adult prostate were transformed with a single copy of the human papillomavirus 18 (HPV-18) to establish the nontumourigenic RWPE-1 cell line.⁶ RWPE-1 cells were further transformed by Ki-ras using the Kirsten murine sarcoma virus (Ki-MuSV) to establish the tumourigenic RWPE-2 cell line.⁶ Exposing RWPE-1 cells to *N*-methyl-*N*-nitrosourea (MNU) created a family of tumourigenic cell lines (WPE1-NA22, WPE1-NB14, WPE1-NB11 and WPE1-NB26) that show increasing invasiveness. This family of cell lines (represented schematically in Fig. 1) with a common lineage represents a unique and relevant model which mimics stages in progression from localised malignancy to invasive cancer, and can be used to study carcinogenesis, progression, intervention and chemoprevention.⁷

Spectroscopy is being increasingly used in biomedical applications with high degrees of success. IR spectroscopy is a nondestructive method for the analysis of cells, tissues and fluids.⁸ IR spectroscopy coupled with advanced computational methods has been used to detect/differentiate between different diseases and stages/grades of malignancy from tissue biopsies. These include benign and malignant prostate,^{2,9-11} colon^{12,13} and cervical¹⁴ tissues, all of which have been evaluated using IR and have resulted in high classification accuracies. However, most laboratories or projects use or require different pre-processing



Fig. 1 A schematic showing the RWPE family cell line lineage.

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methods. The imagined end user of these methods is quite often not a spectroscopist, statistician or chemometrician, *etc.* but a clinical pathologist. For this reason, for the successful translation of biomedical spectroscopy to the clinical environment a move towards standardisation of pre-processing methods is needed.

In this study we present the use of FTIR spectroscopy, laboratory and synchrotron based, combined with multivariate analysis for the investigation of a family of cell lines derived from the same anatomical position. We also discuss the use of a machine learning genetic algorithm (GA) as a potential source of pre-processing standardisation to allow end users maximum flexibility in using spectroscopy in the clinical environment.

Materials and methods

Cell culture and sample preparation

The RWPE-1, RWPE-2, WPE1-NA22, WPE1-NB14, WPE1-NB11 and WPE1-NB26 cell lines were all obtained from the American Type Culture Collection (ATCC) and were cultured according to identical ATCC protocols. Cells were cultured onto 2 cm \times 2.5 cm MirrIR slides (Kevley Technologies, OH, USA) until 80% confluent, fixed in 4% formalin in phosphate buffered saline and air-dried before use.¹⁵ Thirty slides per cell line representing thirty different cultures per cell line were prepared.

Invasion assay

Invasion assays were conducted according to Hart *et al.*¹⁶ Basically 1×10^5 cells in 0.25 ml RPMI 1640–0.1% fatty acid free BSA were seeded into cell culture inserts (8 µm pore size) coated with phenol red free MatrigelTM diluted 1 : 25 with phenol red free RPMI 1640 medium. The inserts were placed in a 24 well plate containing 1 ml of RPMI 1640 (w/o phenol red)–0.1% fatty acid free BSA–10 mM HEPES over tissue culture plastic (TCP) or human bone marrow stroma (BMS). 18 h post-incubation at 37 °C 5% CO₂ in humidified air, the inserts were washed in PBS and non-invading cells removed by wiping with a cotton bud. Inserts were stained with 2% crystal violet–20% methanol for 10 minutes prior to washing and allowed to air dry. Invading cells were counted using a graticule according to manufacturer's instructions.

Data acquisition

Synchrotron microspectroscopy. Single-cell spectra were collected using synchrotron radiation at beamline station 11.1 of Daresbury Laboratory Synchrotron Radiation Source (SRS) on a Nicolet Continuµm XL FTIR microscope equipped with a liquid nitrogen cooled MCT detector. The aperture used was set to fit the size of the interrogated cell, typically around 20 µm \times 20 µm. The spectra represent 150 co-added scans with a resolution of 4 cm⁻¹ and a spectral range of 700 cm⁻¹ to 6000 cm⁻¹. Background spectra were taken from a cell free area as close as possible to the analysed cell.

Laboratory microspectroscopy. Spectra were collected using a Perkin Elmer Spectrum Spotlight 300 FTIR microscope coupled to a Perkin Elmer Spectrum One Spectrometer. The



Fig. 2 RWPE-1 cultured cells on a MirrIR slide with the aperture area $150 \times 150 \ \mu\text{m}^2$ shown by the red square.

microscope is equipped with a liquid nitrogen cooled MCT detector and a CCD camera to provide an optical image of the area under interrogation. An aperture size of 150 μ m × 150 μ m was used to obtain spectra from confluent monolayers. Typically 100 co-added scans were used for the RWPE-1 cell line and 150 co-added scans for RWPE-2, WPE1-NA22, WPE1-NB11 and WPE1-NB26. An example of an RWPE-1 culture on a MirrIR slide with aperture area labelled is shown in Fig. 2. A resolution of 4 cm⁻¹ and the spectral range 700 cm⁻¹ to 6000 cm⁻¹ was used. Background spectra were collected from a separate piece of blank MirrIR slide. At least 5 spectra were acquired from each sample. Spectra tainted by water vapour were discarded.

Laboratory broadbeam spectroscopy. FTIR spectra were collected using a Varian 3100 Excalibur Series FTIR spectrometer equipped with a liquid nitrogen cooled MCT detector. Samples were placed upon a Pike Technologies 30 Spec 30 degree spectral reflectance accessory to allow spectral acquisition. Each spectrum represents 256 co-added scans collected at 4 cm⁻¹ resolution. Spectra were acquired from a large population of cells, which act to average the signal and hence allow single-cell specific characteristics (*e.g.* cell cycle stage) to be disregarded. 10 spectra were collected from each culture resulting in 300 spectra per cell line. A background spectrum was collected before starting analysis and after every 5 spectra.

Data analysis

Two different analyses were performed. The datasets acquired using synchrotron and laboratory based microspectroscopy were analysed in a typical fashion *i.e.* with the analyst choosing the pre-processing procedures and multivariate model to use, whereas the laboratory based broadbeam spectroscopic study was analysed using genetic algorithm fed support vector machines and principal component analysis. For the microspectroscopic study the cell lines used were RWPE-1, RWPE-2, WPE1-NA22, WPE1-NB26 and WPE1-NB11. The broadbeam spectroscopic study used these cell lines as well as WPE1-NB14.

Laboratory based and synchrotron based microspectroscopic study. Matlab coupled with in house written software was used for data processing. The spectra were vector normalised, corrected using the extended multiplicative signal correction (EMSC) model,¹⁷ using the average spectrum as the reference for correction and finally the Savitzky–Golay smoothed first derivative was taken using a 5-point smoothing window. (Note that the latest version of the resonant Mie scattering correction (RMieS-EMSC) was not available for this study.^{18,19})

The spectral range 900-1800 cm⁻¹ was used, resulting in 467 spectral data points for principal component analysis (PCA) and principal component-discriminant function analysis (PC-DFA). PCA is a common unsupervised multivariate method for finding patterns/structures within high dimensionality datasets. PCA was computed using the Non-linear Iterative Partial Least Squares (NIPALS) algorithm. PC-DFA utilises PCA to reduce the dimensionality of the data prior to discriminant function analysis (DFA). DFA then discriminates between groups on the basis of the resultant PCs and the a priori knowledge of the group membership that are fed into the DFA algorithm. Maximising the inter-group variance and minimising the intra-group variance achieve this. The maximum number of discriminant functions available is the number of groups minus one.²⁰ The optimum number of PCs was determined iteratively. Prior to DFA, the dataset was split into a training set and an independent test set. The spectra were randomly assigned to either set, with the constraint that 20% of the spectra collected on each cell line should belong to the independent test set. As PC-DFA is a supervised technique and the model is supplied with information about group membership, any result produced by the model needs to be tested. This testing was carried out by supplying the model with the independent test set and observing where the model places the spectra on a graphical output. Confidence ellipses or ellipsoids are added to the discriminant function plots. These are, respectively, 2D and 3D visualisation of the 95% confidence interval. This was achieved using error_ellipse.m written by A. J. Johnson and obtained from Matlab central file exchange.²¹ Covariance matrices were calculated from the discriminant function analysis score matrix for each grouping, where the centroid was defined as the mean of each discriminant function analysis score matrix for each grouping.

Laboratory based broadbeam spectroscopic study. The spectra were subjected to a quality test whose main criteria were: (1) the difference between the highest and the lowest point of the amide I peak had to be between 0.3 and 1.3 absorbance units and (2) an absence of peaks attributable to water vapour. As a result of the quality test approximately 10% of the spectra were discarded, the spectral numbers per cell line and split between training set, validation set and blind test set are shown in Table 1.

 Table 1
 Number of spectra per cell line and per spectral set^a

Cell line	Training set	Validation set	Test set	Total		
RWPE-1	150 (15)	30 (3)	120 (12)	300 (30)		
RWPE-2	150 (15)	30 (3)	100 (10)	280 (28)		
WPE1-NA22	150 (15)	30 (3)	70 (7)	250 (25)		
WPE1-NB11	150 (15)	30 (3)	110 (11)	290 (29)		
WPE1-NB14	150 (15)	30 (3)	70 (7)	250 (25)		
WPE1-NB26	150 (15)	30 (3)	60 (6)	240 (24)		
^{<i>a</i>} Number of cultures shown in brackets						

 Table 2
 Range of pre-processing techniques used by the genetic algorithm

Processing	Туре	Range
Derivatisation	None	NA
	1 st Order	NA
	2 nd Order	NA
Smoothing	Savitzky–Golay 5th order	5 7 9 11 13 15 17 19 21
e	Moving average	3 5 7 9 11 13 15 17 19
Scaling	Auto-scaling	NA
e	Range-scaling	NA
EMSC	NA	NA

The blind test set was used as a double blind set as the analysis was performed at the Focas Research Institute, Dublin, Ireland and the identity of the spectra in the blind test set was kept by MJB.

The genetic algorithm (GA), principal component analysis (PCA), support vector machine (SVM) and implementation of pre-processing functions were carried out using MatlabTM. All analyses were performed using a dual quad core (Zenon) with 16 GB RAM.

Genetic algorithm (GA) implementation. A GA was used to discover the optimum pre-processing technique from a range of pre-processing techniques (Table 2). Optimisation was implemented using a modified version of the Genetic Algorithm Optimisation Toolbox for MatlabTM.²²

50 independent genetic algorithm runs were conducted retaining the highest cross-validation score, which depends upon the number of correctly classified spectra in the validation set. Using the optimum solution from each independent run, a support vector machine (SVM) was trained using the selected pre-processing regimes and selected SVM meta-parameters. Jarvis and Goodacre have successfully demonstrated the genetic algorithm optimisation approach for the selection of pre-processing methods and discriminatory spectral regions.²³

Support vector machine (SVM) implementation. Support vector machines were constructed using the LibSVM package.²⁴ Binary versions of LibSVM's symtrain and sympredict programs were controlled from MatlabTM.

Results and discussion

Invasion assay

The results of the invasion assay towards tissue culture plastic (TCP, blue) or bone marrow stroma (BMS, red) are shown in Fig. 3.

The invasion towards TCP is very low as expected, whereas when a strong chemoattractant such as BMS is introduced the invasive abilities of the cells are revealed. Bone is the most common metastatic site for prostate cancer and as such bone marrow stromal cells have been shown to enhance prostate cancer cell invasions.²⁵ The invasiveness of the cell line is compared to the invasiveness of PC-3, a cell line established from a bone metastatic site.²⁶ Previous studies have shown a range of invasiveness for these cell lines; RWPE-1 was found to be



Fig. 3 Graph showing the propensity of the different cell lines for invasion towards tissue culture plastic (TCP, blue) and bone marrow stroma (BMS, red).

non-tumourigenic/invasive whilst WPE1-NA22, WPE1-NB14, RWPE-2, WPE1-NB11 and WPE1-NB26 displayed increasing tumourigenic and invasive characteristics. The results of our invasion assay (Fig. 3), importantly, show RWPE-1 and the slow growing/tumour forming RWPE-2 to have about equal invasiveness capacity towards BMS and the WPE1 cell lines follow the general increase as reported in the literature, however, the error bars of the WPE1-NB11 and WPE1-NB26 cell lines do overlap significantly.

Laboratory based and synchrotron based microspectroscopic study

Laboratory based microspectroscopic study. The laboratory based microspectrometer was used to acquire spectra from fields of views containing tens of cells, and thus the spectra represent an average of those cells. The multiple cell diagnostic model was constructed using 672 spectra. The number of spectra per cell line for the training and independent test set is shown in Table 3.

The PCA score plot is shown in Fig. 4. Utilising the first two principal components (PCs) yielded the best separation of the cell lines, PC1 accounted for 56% and PC2 21% of the variance. Explaining 8% of the variance, PC3 did not provide any better separation.

Spectra from the RWPE-1 cell line (yellow circles) formed the most discernible cluster. PC1 generally separates the nontumourigenic RWPE-1 and low invasiveness cell line WPE1-NA22 from the slow tumour forming RWPE-2 and the more

 Table 3
 Distribution of acquired multiple cell spectra per cell line for the laboratory based microspectroscopic study

Cell line	Number of samples	Total number of spectra	Spectra in training set	Spectra in test set
RWPE-1	27	133	106	27
RWPE-2	29	148	118	30
WPE1-NA22	25	125	100	25
WPE1-MB11	29	145	116	29
WPE1-NB26	24	121	97	24
Total	134	672	537	135



Fig. 4 PCA score plot of the whole dataset (PC1 *vs.* PC2). A different coloured circle as per the legend of the figure represents each spectrum of the cell lines.

invasive cell lines (WPE1-NB11 and WPE1-NB26), whereas PC2 generally separates RWPE from WPE cell lines. Observing both PC1 and PC2 together, three distinct groupings can be seen: (1) RWPE-1, (2) RWPE-2 and WPE1-NA22 and (3) WPE1-NA11 and WPE1-NB26. However, as the clusters are not wholly clear, a supervised method of multivariate analysis, such as PC-DFA, will be used to illuminate difference between the cell lines.

Fig. 5(A) shows the discriminant function plot of DF1 vs. DF2 for the multiple cell spectral model based upon the training set (coloured filled circles) and independent test set (coloured empty squares), as per the figure legend, with a 95% confidence limit drawn and Fig. 5(B) shows the discriminant function plot of DF1 vs. DF3 with the 95% confidence limit drawn. The discrimination in the plots shows different separations based upon different characteristics with Fig. 5(A) showing discrimination along DF1 based upon genetic (RWPE) versus genetic plus chemical (WPE1) transformation and DF2 has separated two different types of genetic transformation, HPV-18 for RWPE-1 compared with HPV-18 plus Ki-Ras for RWPE-2. Fig. 5(B) shows the same separation along DF1 however DF3 is separating WPE1-NA22 from WPE1-NB11 and WPE1-NB26. However, it is not clear if this separation is based upon invasiveness or the difference in amount of MNU used to achieve the chemical transformation.

As 3 discriminant functions have been used it was relevant to use a pseudo-3D discriminant function plot. Fig. 6(A) shows a 3D discriminant function plot of DF1 vs. DF2 vs. DF3 based upon the training set data (coloured filled circles) and independent test set (coloured empty squares), as per the figure legend.

To assess the quality of discrimination the measures of sensitivity and specificity are used. Sensitivity measures the ability of the model to correctly classify whereas specificity measures the ability of the model to not misdiagnose. The sensitivities and specificities for the multiple cell spectral model based upon the pseudo-3D discriminant function plot are shown in Table 4.

The sensitivities and specificities (Table 4) and the pseudo-3D discriminant function plot (Fig. 6) reveal that all the false positives for WPE1-NB11 were from WPE1-NB26 spectra and all the false



Fig. 5 Discriminant function plots showing (A) DF1 vs. DF2 and (B) DF1 vs. DF3 for the multiple cell spectral model based upon the training set (coloured filled circles) and independent test set (coloured empty squares), as per the figure legend, with a 95% confidence ellipse drawn.



Fig. 6 (A) Pseudo-3D discriminant function plot of DF1 vs. DF2 vs. DF3 based upon the training set (coloured filled circles) and independent test set (coloured empty squares) and (B) pseudo-3D discriminant function plot with 95% confidence ellipsoids.

positives for WPE1-NB26 were from the WPE1-NB11 spectra. Due to this, a new group comprising of cells from both cell lines was tested for sensitivity and specificity. Invasion assay results (Fig. 3) show that WPE1-NB26 and WPE-NB11 are very close in their invasiveness. The pseudo-3D model is able to discriminate 4 groups of cell lines RWPE-1, RWPE-2, WPE1-NA22 and WPE1-NB(11 and 26), to a high degree of accuracy, with the average sensitivity and specificity of 94% and 99.8% respectively. The specificity was exceptional in illuminating the robustness of the discrimination. Test spectra which did not fall within the confidence ellipsoid did not fall into the wrong ellipsoid.

Discriminant function 1 separated the RWPE cell lines from the WPE1 cell lines whilst discriminant functions 2 and 3 provide separation within these two groups (Fig. 5 and 6). The model is able to adequately differentiate cell lines from the RWPE and WPE families. Clusters corresponding to the chemically modified cell lines lay close to each other and the more aggressive clusters (WPE1-NB11 and WPE1-NB26) clustered together. WPE1-NA22 cells were derived from cells exposed to MNU at a concentration of 50 µg l⁻¹ whereas WPE1-NB11 and WPE1-NB26 originated from the same batch of cells exposed to MNU at 100 μ g l⁻¹ and were separated from each other only after successive steps of growth in culture and injection into immunodeficient mice.⁷ Although, cell lines are separated, there is no systematic order of separation according to level of invasiveness and thus it appears to be primarily dependent on the method of transformation rather than the difference in invasiveness which raises questions on the usefulness of cell lines in modelling cancer. Erukhimovitch et al.27 have previously questioned the use of cell lines to model non-malignant cells in their study on human and mouse cell lines, cancer cells and primary cells. This study suggests that cell lines should all be considered as premalignant cells due to the immortal character achieved by the transformation. Our study takes this further by suggesting that biochemical changes induced by different transformation methods are primarily responsible for the discrimination of the RWPE family of cell lines and it is not possible, as was the research aim, to model biochemical changes associated with

Table 4 Sensitivities and specificities for the multiple cell spectral model based upon the pseudo-3D discriminant function plot

Cell line	True positives	False negatives	Sensitivity (%)	True negatives	False positives	Specificity (%)
RWPE-1	23	4	85.2	108	0	100.0
RWPE-2	29	1	96.7	105	0	100.0
WPE1-NA22	25	0	100.0	109	1	99.1
WPE1-NB11	27	2	93.1	87	19	82.1
WPE1-NB26	22	2	91.7	86	25	77.5
WPE1(NB11 + 26)	50	3	94.3	82	0	100.0

invasiveness using FTIR spectroscopy in prostate cancer using these cell lines.

A study by Romeo *et al.*¹ on human oral mucosa cells and canine cervical cells resulted in the different cell types grouping together. This was thought to be due to the nucleus to cytoplasm ratio of the cells being more discriminatory than biochemical changes. However, a recent study²⁸ has shown that the major reason for discrimination of prostate cancer cell lines, albeit ones from different anatomical positions, by FTIR is the biochemical differences between the cell lines. Thus we can be confident that we are observing discriminatory biochemical differences between the RWPE family of cell lines but it should be stressed that these differences appear to derive from the method of transformation rather than the degree of invasiveness.

Synchrotron based microspectroscopic study. A preliminary study utilising synchrotron based FTIR microspectroscopy was performed resulting in a total of 135 spectra. Each spectrum represents the interrogation of a single cell. The breakdown for each cell line is as follows: RWPE-1 19 spectra, RWPE-2 20 spectra, WPE1-NA22 29 spectra, WPE1-NB11 33 spectra and WPE1-NB26 34 spectra. The aspect of the averaged IR spectrum, for the spectral range 900–1800 cm⁻¹, from the whole single-cell spectral dataset was very similar to that calculated from the multiple cells (Fig. 7).

To assess the preliminary data collected on single cells a PC– DFA analysis was performed. However, in this analysis instead of splitting the data into a training set and independent test set 10 separate analyses were performed with 7 randomly chosen spectra from each cell line in the training set and the remaining spectra in the independent test set each time. Fig. 8(A) shows a pseudo-3D discriminant function plot of DF1 vs. DF2 vs. DF3 based upon one of the ten analyses performed with the training set data (coloured filled circles) and independent test set (coloured empty squares). Fig. 8(B) shows the discriminant function plot with 95% ellipsoids drawn.

Spectra from the preliminary single-cell model did not cluster as well as the multiple cell spectra. Spectra from RWPE-1 and







Fig. 8 (A) Pseudo-3D discriminant function plot of DF1 vs. DF2 vs. DF3 based upon the training set (coloured filled circles) and independent test set (coloured empty squares) and (B) pseudo-3D discriminant function plot with 95% confidence ellipsoids.

RWPE-2 are clearly distinguishable from each other along discriminant function 2 and from the WPE1 cell lines along discriminant function 1, whereas the WPE1 cell lines are less distinguishable. Due to the increased variability in the spectra and the small size of the dataset, 95% confidence ellipsoids were large and overlapped. The average sensitivities and specificities for the single-cell model are shown in Table 5.

The overall average sensitivity and specificity are 67.3% and 79.8%, respectively, for this preliminary single-cell dataset. The model was able to adequately separate RWPE-1 from RWPE-2 and the RWPE cell lines from WPE1 cell lines.

The results from the preliminary single-cell spectral model are consistent with those from the multiple cell spectral model in that the same 3 main clusters consisting of HPV-18 transformed RWPE-1, HPV-18 and Ki-ras transformed RWPE-2 and HPV-18 and chemically transformed WPE1 cells are isolated. However, discrimination between the WPE1 cells could not be achieved. The standard deviation observed among the single-cell spectra was larger than that observed for the multiple spectra, attesting the large variability between single cells. A study by German *et al.* utilising synchrotron and laboratory based

Table 5 Sensitivities and specificities for the single-cell spectral model

Cell line	Sensitivity (%)	Specificity (%)		
RWPE-1	27.1	87.1		
RWPE-2	60.0	93.7		
WPE1-NA22	70.6	78.6		
WPE1-NB11	88.6	66.6		
WPE1-NB26	90.0	73.1		

A								
Derivatisation	EMSC	Filter type	Window	Normalis	ation S	caling SV	M penalty (C)	RBF gamma
1 st order	None	MA	9	None	A	uto 9.6	017	9.6626
В								
	IR assignment	RWPE-1	RWPE-2	WPE-NA22	WPE-NB11	WPE-NB14	WPE-NB26	Sensitivity (R)
Actual Cell Line	RWPE-1 RWPE-2 WPE-NA22 WPE-NB11 WPE-NB14 WPE-NB26 Specificity (S)	108 0 0 0 0 0 100.00	1 100 0 0 0 0 99.70	10 0 69 0 0 0 97.83	0 0 110 0 100.00	1 0 0 67 0 99.78	0 0 1 0 3 60 99.15	90.00 100.00 98.57 100.00 95.71 100.00

Table 6 (A) Optimum GA selected data pre-processing and SVM design parameters, and (B) confusion matrix from blind set testing of the optimum pre-processing and SVM design. The sensitivities (R) and specificities (S) are shown for each class

infrared radiation has shown that both techniques highlight similar spectral characteristic despite the increased intra-variability observed with synchrotron FTIR microspectroscopy.²⁹ Importantly this preliminary study on single cells has concurred with the multiple cell spectral study, which was performed on a different instrument with a different experimental protocol and on a different scale.

Laboratory based broadbeam spectroscopic study

Genetic algorithm fed support vector machine (SVM). The genetic algorithm identified the pre-processing conditions in Table 6(A) from the available conditions supplied (Table 2), as the optimum pre-processing conditions and SVM parameters for discriminating the RWPE cell lines from the laboratory based broadbeam spectroscopic study. SVM penalty is a measure of the misclassification of the training data and RBF gamma is the use of a radial basis function to determine the area of influence the support vector has over the data space. The output from the SVM prediction of blind set classification is assessed *via* a confusion matrix (Table 6(B)).

The genetic algorithm fed SVM is able to discriminate the RWPE family of cell lines to an average overall sensitivity and specificity of 97.37% and 99.41% respectively. The main errors in the model arise from RWPE-1 cells misclassified as WPE1-NA22 and WPE1-NB14 misclassified as WPE1-NB26. Although these misclassifications are small in number they are important since they are to cell lines with very different degrees of invasiveness.

As the imagined end user of these technologies will not be a spectroscopist or chemometrician and the ultimate aim is to translate this research into the clinical environment it is necessary to generate a robust set of pre-processing functions into which the pathologist can easily input spectral data and acquire a clinically relevant output. The use of genetic algorithms (GAs) to select pre-processing conditions and/or discriminatory regions of the spectrum can allow this research community to provide a standard list of options which are acceptable to be supplied to the GA and hence allow optimum separation.



Fig. 9 PCA score plot of the dataset processed using the optimum GA chosen pre-processing methods (PC1 vs. PC2). A different coloured circle as per the legend of the figure represents each spectrum of the cell lines (ellipses drawn as a guide to the eye).

Principal component analysis. The PCA score plot for the GA chosen pre-processing method is shown in Fig. 9. PC1 accounted for 83.62% and PC2 4.53%. Using PC3 did not improve the separation.

Observing the score plot (Fig. 9) for the GA fed SVM, it can again be seen that the groups are not differentiating on invasiveness of the cell line but appear similar to the PC–DFA results obtained on the laboratory based microspectrometer study with a differentiation being made between the RWPE cell lines (genetically transformed) and the WPE1 cell lines (genetically and chemically transformed) along PC1. General clustering can be seen for all cell lines apart from WPE1-NA22.

Conclusions

Laboratory based and synchrotron based microspectroscopic study

FTIR microspectroscopy has been used to distinguish between cells derived from the same origin, same anatomical position, having a close genetic background but differing on tumourigenic behaviour and as such we have further demonstrated the use of FTIR as a sensitive tool for evaluating biological samples and processes. The discrimination has been achieved to a high degree of classification accuracy and repeated with a preliminary study on single cells. The differentiation classification accuracy is better within the laboratory based study compared to the synchrotron based study, primarily due to significantly higher variance in single-cell data and the smaller datasets available. It should be remembered, however, that the single-cell data provide information concerning cell populations and not just the average which can be a significant advantage. The model presented here. however, discriminates based upon differences between the way these closely related cell lines have been transformed and not their invasiveness, showing their unsuitability to model prostate cancer using FTIR and raising important questions on the use of cell lines as cancer models.

Laboratory based broadbeam spectroscopic study

This study has shown the use of a genetic algorithm to select optimum pre-processing methods. This allows us to determine the pre-processing methods which can be used whilst allowing the determined end user maximum flexibility in the application of the technologies and methods concerned with this research. Importantly it has also validated discrimination results observed in the other studies presented in this paper.

Overall, the study demonstrates the potential of FTIR coupled with multivariate analysis technique for pathological screening applications although further studies involving primary cells and tissue are clearly required. The use of genetic algorithms (GAs) to selecting pre-processing conditions and/or discriminatory regions of the spectrum can allow the research community to provide a standard list of options which are acceptable to be supplied to the GA and hence allow optimum separation. Once all the issues regarding spectral correction and pre-processing have been resolved there is no reason why this technology cannot be used routinely in a clinical environment to augment current practice.

Acknowledgements

The authors would like to acknowledge the EPSRC Life Science Interface Scheme (grant reference: EP/E039855/1) and MIMIT[™] (Manchester: Integrating Medicine and Innovative Technology) for financial support.

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