Optimisation of sample preparation using glass slides for spectral pathology

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**ABSTRACT**

The clinical translation of Fourier Transform Infrared (FTIR) microspectroscopy in pathology will require bringing this technique as close as possible to standard practice in pathology departments. An important step is sample preparation for both FTIR microspectroscopy and pathology. This should entail minimal disruption of standard clinical practice while achieving good quality FTIR spectral data. In fact, the recently described possibility of obtaining FTIR spectra of cells placed on glass substrates brings FTIR microspectroscopy closer to a clinical application. We have now furthered this work in order to identify two different types of lung cancer cells placed on glass coverslips. Two types of sample preparation which are widely used in pathology, cytospin and smear, have been used. Samples were fixed with either methanol, used in pathology, or formalin (4% paraformaldehyde) used widely in spectroscopy. Fixation with methanol (alcohol-based fixative) removed lipids from cells causing a decrease in intensity of the peaks at 2850 cm-1 and 2920 cm-1. Nevertheless, we show for the first time that using either type of sample preparation and fixation on thin glass coverslips allowed to differentiate between two different types of lung cancer cells using either the lipid region or the fingerprint region ranging from 1800 cm-1 to 1350 cm-1. We believe that formalin fixed cytospin samples would be preferred to study cells on thin coverslips using FTIR microspectroscopy. This work presents a clear indication for future advances in clinical assessment of samples within pathology units to gain a deeper understanding of cells/tissues under investigation.

Keywords

FTIR microspectroscopy, lung cancer, glass coverslips, cell fixation, cytospin, smear.

**INTRODUCTION**

It is widely accepted that Fourier Transform Infrared (FTIR) microspectroscopy has a potential as a diagnostic tool in histopathology. However, in spite of the work carried out so far, the clinical translation has not materialised yet. There are several aspects that have contributed to this absence of clinical translation.1 These include, amongst others, lack of multicentre standardisation in sample preparation, data collection and data analysis, type of substrates used in pathology, and lack of studies including big numbers of patients. Until recently, a major drawback is the type of substrate used in histopathology departments versus the substrates widely used in FTIR microspectroscopy. Pathologists use glass substrates of around 1 mm nominal thickness for both histopathology and cytology. The problem is that glass absorbs IR radiation thus reducing the amount of spectral information that can be obtained from tissue and cell samples. Previous studies have shown that, using the same glass substrate used by pathologists, information can still be obtained from the lipid region but not from the fingerprint region.2-4 Another issue is tissue samples embedded in paraffin. The latter has strong band absorption in the lipid region masking biochemical information from these samples. This landscape has recently changed with our work showing that, using glass coverslips of 0.12-0.17 mm thickness as substrates, spectral information not only from the lipid region but also from the amide I4 and, more importantly, from the fingerprint region down to 1350 cm-1 can now be obtained.5 This opens a whole new area of research for FTIR microspectroscopy by utilising substrates that are also used in pathology departments. In fact, by placing samples on coverslips, not only can spectral information be obtained but also, if further histopathological information is needed, these samples on coverslips can then be placed on and glued to standard histopathological glass slides, stained and covered with another coverslip for histopathological analysis.

This advancement in spectral pathology brings FTIR microspectroscopy closer to a clinical application. In fact, the closer the sample preparation is to a clinical set up, the faster the technique could be applied and used by clinicians. However, diverse cytological preparations (smears, cytospins, cell blocks and liquid-based cytology) fixed with different fixation procedures are used in different histopathology laboratories.6 The histopathology department at the University Hospitals of North Midlands (UHNM), like in many hospitals worldwide, prepare cell samples using either the cytospin technique or smear. Examples are cytology samples for lung cancer diagnosis (bronchio–alveolar lavage, pleural fluid) which are prepared as cytospins. Fine needle aspirations from thyroid nodules or head and neck lymph nodes are prepared as smears. For the diagnosis of haematological malignancies, bone marrow aspirates and blood samples are prepared as smears. These samples are then fixed with alcohol-based fixatives such as methanol. Furthermore, in some cases, samples are fixed and stained at the same time. Smears can lead to uneven distribution of cells, thick cell clusters, and inadequate cellularity.7 However, morphological preservation seems better in smears than in cytospins.8 On the other hand, cytospins provide cellularity even in cases where smears show unsatisfactory cellularity, increasing the diagnostic potential.8

On this basis, we carried out the present work to identify the best cell sample preparation (cytospin versus smear) using glass coverslips as substrates to study two different types of lung cancer cells using FTIR microspectroscopy. It is known that alcohols, used as fixatives, alter lipid content during sample fixation affecting the spectra of cells.9 On the other hand, formalin (4% paraformaldehyde), widely used as fixative in sample preparation for FTIR microspectroscopy, does not affect lipid content as much as methanol during fixation. Therefore, we tested formalin and methanol as fixatives for this work.

**MATERIALS AND METHODS**

Cells

The cells used in this study were CALU-1, an epidermoid lung cancer cell line from the European Collection of Cell Cultures (ECACC) and A549, a lung adenocarcinoma cell line (ECACC). Both cell lines were cultured in Dulbecco’s Modified Eagle Medium supplemented with 10% foetal bovine serum, 5% L-glutamine, 5% antibiotic/antimycotic, 5% HEPES (4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid) buffer, 5% non-essential amino acids, and 5% sodium pyruvate (MERCK, UK). For both cell lines, media was changed every 3-4 days. Cells were passaged before reaching confluence by removing culture medium and adding trypsin/EDTA (MERCK, UK) and incubating cells for 5 minutes. After this incubation period, cells were collected and centrifuged at 1200 rpm for 5 minutes. Supernatant was discarded and the pellet re-suspended in fresh medium. Cell viability for both cell lines was determined with the standard trypan blue exclusion method.

Sample Preparation

Two types of sample preparation were carried out, cytospin and smear. For cytospin, cells were collected as described above, transferred into 15 mL tubes and centrifuged at 1200 rpm for 5 minutes at room temperature. The supernatant was then removed and the pelleted cells were re-suspended in 0.9% NaCl. Cell concentration was brought to 106 cells/mL. 20 µL of this cell concentration was placed in a cytospin funnel and cells were then cytospun for 1 minute at 900 rpm on GalvOptics microscope coverslips (24 x 50 mm x 0.13–0.17 mm thickness, GalvOptics, UK) made of soda–lime glass.

Smears were set up as follows. 20 µL of a cell concentration of 106 cells/mL in 0.9% NaCl were placed in one of the edges of a GalvOptics microscope coverslip. The edge of a second GalvOptics microscope coverslip was then placed on the liquid sample and the sample spread out over the length of the first coverslip.

Two types of fixation were used, formalin and methanol. For formalin fixation, samples were fixed with 4% buffered paraformaldehyde in 0.9% NaCl for 20 minutes at room temperature. Excess formalin was removed by washing once with 0.9% NaCl and thrice rinsed with distilled water. Samples were then air dried at room temperature. For methanol fixation, samples were placed in ice cold methanol (VWR International, UK) for 2 minutes and then allowed to dry. No washings were carried out as methanol is a volatile compound and evaporated from the sample. This method of fixation followed the same protocol used at the pathology department at UHNM. For each type of sample and fixation, 2 replicates were prepared and 50 cells from each replicate studied.

FTIR microspectroscopy

Mid-infrared (mid-IR) spectra of the samples were obtained using a Thermo Nicolet iN10(MX) spectrometer. Spectra were collected at 4 cm-1 resolution, with 256 co-added scans using an aperture size of 15 x 15 µm centered on the cell nucleus based on our previous work.10 The time to obtain the spectrum of a single cell was less than 2 minutes. Background measurements were obtained under the same conditions from areas of coverslip without a biological sample. For all samples, spectra of 100 individual cells were obtained. Data pre-processing was carried out as previously described,4,5 i.e., spectra were cropped to the area to be analyzed and normalized using standard normal variate (SNV) which subtracts the mean spectrum and then divides the standard deviation for each spectrum removing the effect of different sample thickness and spectrum baseline offsets. The spectral analysis of cells on coverslips included the areas between 3100 cm-1 and 2700 cm-1 for the lipid region, and between 1800 cm-1 and 1350 cm-1 for the fingerprint region, the latter based on our previous work using GalvOptics microscope coverslips.5

Data analysis

Principal component analysis (PCA) was performed using Unscrambler X software (CAMO). A Levene’s test was carried out to assess homogeneity of variance. The statistical analysis was applied on the Principal Components on pair of data, thus, the T-test was carried out for data that was normally distributed. The non-parametric Mann-Whitney test, which is the equivalent non-parametric test of the T-test, was used for data not normally distributed. Statistical analysis was carried out in Excel. The statistical tests were applied with 0.05 level of significance.

**RESULTS**

Figures 1 and 2 show the spectra of cells prepared as cytospins or smears, respectively (mean of 100 spectra). They also show the spectra of cells fixed with formalin or methanol. In Figure 1, the main difference between the two types of fixation methods is the loss of lipids when cells were fixed with methanol and, therefore, a decrease in the 2850 cm-1 and 2920 cm-1 peaks, as well as a shift of the 2920 cm-1 peak towards higher wavenumbers. The same is seen in Figure 2 where cells were prepared as smears.

It has to be indicated here that sample preparation as cytospin has the advantage over smears that yields a more uniform cell population to study cells using FTIR microspectroscopy rather than having the cells spread out over the glass substrate. Obtaining FTIR spectra from cytospin samples is a quicker process as cells are concentrated to one area. Furthermore, cytospins are fixed as soon as the sample is prepared after spinning at 900 rpm for 1 minute. Smears, in order not to lose too many cells, need to be dried first prior to fixing. This step before fixing smear samples could lead to changes in the biochemical structure and thus changes in the cells’ FTIR spectra. In fact, CALU-1 cell samples prepared as smears have changes in the shape of the amide I and II bands (Figure 2). Statistical analysis of a PCA comparing cytospin versus smear sample preparation showed statistical significant differences between these 2 types of samples regardless of type of fixation or cell line for PC1 for both the lipid region and the fingerprint region between 1800 cm-1 and 1350 cm-1 (Table I). These differences can be attributed to the differences in the drying process as described above.

The next step was to carry out further PCA in order to assess the differences between the spectra of these cells prepared as cytospins or smears, and fixed with formalin or methanol. Figures 3 and 4 show the PCA for the 2 studied cell lines prepared as cytospins for the lipid region and the fingerprint region between 1800 cm-1 and 1350 cm-1, respectively, and their corresponding loadings. Regarding the fixation method and as expected from the mean cell spectra, there was a clear separation between CALU-1 and A549 cells fixed with formalin or methanol for the lipid region (Figure 3a). Regarding the studied fingerprint region, there was still a separation between these two cell lines based on how they had been fixed (Figure 4a).

The PCA data for cells prepared as smears are shown in Figures 5 (lipid region) and 6 (fingerprint region between 1800 cm-1 and 1350 cm-1), as well as their corresponding loadings. For the lipid region, a similar pattern was seen when compared to samples prepared as cytospins, i.e., a clear separation between cells fixed with formalin or methanol was evident (Figure 5a). Regarding the studied fingerprint region, some differences could still be seen (Figure 6a). The statistical analysis of the PCAs in Figures 3 to 6 is shown in Table II. When comparing fixation with formalin or methanol, statistically significant differences can be seen between these 2 types of fixation regardless of cell line and whether samples have been prepared as cytospins or smears.

In order to assess whether the differences were not just due to sample preparation but also to biochemical differences between these 2 cell lines, statistical analysis from the principal components comparing A549 cell line (lung adenocarcinoma) and CALU-1 (lung epidermoid carcinoma) were carried out. Again, the statistical analysis in Table III is based on the PCAs in Figures 3 to 6. Statistically significant differences between these 2 types of cell lines both in the lipid region and the region between 1800 cm-1 and 1350 cm-1 within the fingerprint area for PC1 can be seen. These differences are present regardless of type of sample (cytospin or smear) and fixative used (formalin or methanol). For PC2, the differences were statistically significant for samples prepared as cytospins and fixed either with formalin or methanol (Table III). This is the first time that different types of lung cancer cell types have been separated with FTIR microspectroscopy using glass coverslips as substrates.

**DISCUSSION**

Several studies have already shown the advantages and difficulties of cell and/or tissue fixation for FTIR spectroscopy.11-16 However, none of these studies used glass as a substrate on which to hold the sample for analysis. It could be argued that the type of substrate would not make much difference to the final spectral data. However, if this technique is to make it into standard clinical practice, it is important to confirm that fixing cells on glass coverslips with methanol or formalin as examples, still gives robust spectral data which can then be shown to clinicians wishing to apply this technique in pathology departments for disease diagnosis and monitoring.

When a new methodology is being introduced in clinical practice, it is important that there are no major disruptions on the standard procedures used by clinicians. In the case of the introduction of FTIR microspectroscopy in histopathology, it would be crucial in a first instance that sample preparation is as close as possible to what it is considered standard clinical practice. However, it is also crucial that sample preparation does not modify cells in a way that FTIR spectra could be greatly affected. Thus, it is not always easy to find the right balance between what FTIR microspectroscopy can offer and what clinicians might expect from this technique.

Two common sample preparations in pathology are cytospins and smears. The latter is faster and requires less equipment thus reducing costs when compared to cytospin preparation. This is paramount for the already economically stretched health services worldwide. Thus, smear preparation could be the way forward when taking FTIR microspectroscopy to clinical practice. However, the fact that smear samples need drying before being fixed poses a problem when studying these samples with FTIR microspectroscopy. It is well known that drying cells could affect their FTIR spectra16 mainly in the fingerprint region17,18 as air drying could cause delocalisation of biomolecules as a result of large surface tension forces associated with the water air interface13 with protein vibration bands undergoing changes in both intensity and line shape upon drying.18 In fact, we have seen changes in the amide I when fixing cells with methanol probably due to the drying step (Figure 2). In view of the issues with air drying, preparing samples as cytospin would be in our opinion the way forward. However, it is not clear yet whether the data presented here using cytospin on glass slides together with previous reported data is robust enough to create a change of practice in pathology departments for samples that are presently prepared as smear. Further work might be needed including, amongst other, more cancer types and bigger sample populations.

Regarding cell fixation, several protocols have been described.19 The way histology samples are fixed in pathology departments is also an important point when bringing FTIR microspectroscopy to clinical practice. Fixation of cytology slides in methanol is an easy, straightforward and faster procedure when compared to formalin fixation. However, methanol is an alcohol that washes out most of the membrane lipids and thus affecting the spectra of cells.9 This can be seen in Figures 1 and 2 where the bands at 2850 cm-1 and 2920 cm-1 corresponding mainly to the CH2 stretching modes of methylene chains in membrane lipids20-22 have a lower intensity in cells fixed with methanol. Furthermore, methanol fixation also causes changes in the position of the 2920 cm-1 band (Figures 1 and 2). Conversely, formalin causes the crosslinking of the primary and secondary amine groups of proteins preserving also the lipids.13 This cross-linking locks the secondary structure of proteins11 although small spectral changes in the fingerprint region following formalin fixation have also been described.12, 15, 23 However, these changes seem to be smaller than biochemical changes caused by cancer.14 Nevertheless, we have shown for the first time that placing lung cancer cells on glass coverslips as smear or cytospin, it is possible to differentiate two different types of lung cancer cells using FTIR microspectroscopy (Table III). Further work is needed to confirm this in other cell lines and diseases.

While some of the data presented here would have been expected based on previous published data,9 our work using a glass substrate brings forward FTIR microspectroscopy towards a clinical application in pathology. The methodology we would advocate would be preparing cytology samples as cytospins using glass coverslips as substrates and fixing them with formalin. An automated system would study these samples using FTIR microspectroscopy identifying, in a first instance, “abnormal” samples and separating them from those samples deemed to be non-pathological. The setting up of these samples on glass coverslips and fixed with formalin would then allow the pathologists to further study them using staining and/or immunochemistry techniques as examples. However, if formalin fixation causes problems with immunohistochemistry,24 then, further cytology samples could be prepared from the original patient’s sample. Nevertheless, to have an automated screening tool to identify “*abnormal*“ samples would reduce working hours in pathology departments as pathologists would need to concentrate only on those samples considered “*abnormal*”. In summary, the data presented here together with our previous FTIR data obtained using glass coverslips could be used by spectroscopists to start or further the interest of pathologists in the application of FTIR microspectroscopy in cancer diagnosis.

**ACKNOWLEDGEMENTS**

The support from the Cancer Centre and University Hospitals of North Midlands charities is greatly acknowledged. L.M.D. is partially supported by Alba Synchrotron Light Source.

**CONFLICT OF INTEREST**

The authors declare that they have no conflict of interest.

**REFERENCES**

1. M.J. Baker, H.J. Byrne, J. Chalmers, P. Gardner, R. Goodacre, A. Henderson, S.G. Kazarian, F.L. Martin, J. Moger, N. Stone, J. Sulé-Suso. “Clinical applications of infrared and Raman spectroscopy: state of play and future challenges”. Analyst. 2018. 143(8): 1735-1757.

2. P. Bassan, J. Mellor, J. Shapiro, K.J. Williams, M.P. Lisanti, P. Gardner P. ‘‘Transmission FT-IR Chemical Imaging on Glass Substrates: Applications in Infrared Spectral Histopathology’’. Anal Chem. 2014. 86(3): 1648-1653.

3. M.J. Pilling, A. Henderson, J.H. Shanks, M.D. Brown, N.W. Clarke, P. Gardner. ‘‘Infrared Spectral Histopathology Using Haematoxylin and Eosin (H&E) Stained Glass Slides: A Major Step Forward Towards Clinical Translation’’. Analyst. 2017. 142(8): 1258-1268.

4. A.V. Rutter, J. Crees, H. Wright, D.G. van Pittius, I. Yousef and J. Sulé-Suso. “Fourier transform infrared spectra of cells on glass coverslips. A further step in spectral pathology”. Analyst. 2018. 143(23), 5711-5717.

5. A.V. Rutter, J. Crees, H. Wright, M. Raseta, D.G. van Pittius, P. Roach, J. Sulé-Suso. “Identification of a Glass Substrate to Study Cells Using Fourier Transform Infrared Spectroscopy: Are We Closer to Spectral Pathology?” Appl Spectrosc. 2019. 74(2): 178-186.

6. F. Schmitt, B. Cochand-Priollet, M. Toetsch, B. Davidson, A. Bondi, P. Vielh. “Immunocytochemistry in Europe: results of the European Federation of Cytology Societies (EFCS) inquiry”. Cytopathology. 2011. 22(4): 238-242.

7. A. Nalwa, R. Walia, V. Singh, K. Madan, S. Mathur, V. Iyer, D. Jain. “Comparison of Conventional Smear and Liquid-based Cytology Preparation in Diagnosis of Lung Cancer by Bronchial Wash and Transbronchial Needle Aspiration” J Cytol. 2018. 35(2): 94-98.

8. A.S. Kshatriya, P.M. Santwani. “Comparison of FNAC smears, cytospin smears, and cell blocks of transthoracic guided FNAC of suspected lung tumor: A study of 100 cases”. J. Cytol. 2016. 33(3): 141-144.

9. M.V. Brown, J.E. McDunn, P.R. Gunst, E.M. Smith, M.V. Milburn, D.A. Troyer, K.A. Lawton. “Cancer detection and biopsy classification using concurrent histopathological and metabolomic analysis of core biopsies”. Genome Med. 2012. 4(4): 33.

10. J. Pijanka, A. Kohler, Y. Yang, P. Dumas, S. Chio-Srichan, M. Manfait, G.D. Sockalingum, J. Sulé-Suso. ‘‘Spectroscopic Signatures of Single, Isolated Cancer Cell Nuclei Using Synchrotron Infrared Microscopy’’. Analyst. 2009. 134(6): 1176-1181.

11. J.T. Mason, T.J. O‘Leary. “Effects of formaldehyde fixation on protein secondary structure: a calorimetric and infrared spectroscopic investigation”. J. Histochem Cytochem. 1991. 39(2): 225-229.

12. E. Ó Faoláin, M.B. Hunter, J. Byrne, P. Kellehan, M. McNamara, H. Byrne, F. Lyng. “A study examining the effects of tissue processing on human tissue sections using vibrational spectroscopy”. Vibrat Spectrosc. 2005. 38(1-2): 121-127.

13. E. Gazi, J. Dwyer, N.P. Lockyer, J. Miyan, P. Gardner, C. Hart, M. Brown, N.W. Clarke. “Fixation protocols for subcellular imaging by synchrotron-based Fourier Transform Infrared Microspectroscopy”. Biopolymers. 2005. 77(1): 18-30.

14. A.I. Mazur, E.J. (Swain) Marcsisin, B. Bird, M. Miljkoviš, M. Diem. Evaluating different fixation protocols for spectral cytopathology, Part I. Anal Chem. 2012. 84(3): 1259-1266.

15. V. Zohdi, D.R. Whelan, B.R. Wood, J.T. Pearson, K.R. Bambery, M.J. Black. “Importance of Tissue Preparation Methods in FTIR Micro-Spectroscopical Analysis of Biological Tissues: ‘Traps for New Users’.” PLoS ONE. 2015. 10(2): e0116491.

16. J. Bobrowska, J. Pabijan, J. Wiltowska-Zuber, B.R. Jany, F. Krok, K. Awsiuk, J. Rysz, A. Budkowski, M. Lekka. Data on step-by-step atomic force microscopy monitoring of changes occurring in single melanoma cells undergoing ToF SIMS specialized sample preparation protocol. Data in Brief 2016. 8: 1322-1332.

17. T. Theophanides. “Fourier transform infrared spectra of calf thymus DNA and its reactions with the anitcancer drug cisplatin”. Appl Spectrosc. 1981. 35(5): 461-465.

18. R. Manoharan, J.J. Barag, R.P. Rava, R.R. Dasari, M. Fitzmaurice, M.S. Feld. “Biochemical analysis and mapping off atherosclerotic human artery using FTIR microspectroscopy”. Atherosclerosis. 1993. 103(2): 181-193.

19. M.J Baker, J. Trevisan, P. Bassan, R. Bhargava, H.J. Butler, K.M Dorling, P.R Fielden, S.W Fogarty, N.J Fullwood, K.A Heys, C. Hughes, P. Lasch, P.L Martin-Hirsch, B. Obinaju, G.D Sockalingum, J. Sulé-Suso, R.J Strong, M.J Walsh, B.R Wood, P. Gardner, and F.L Martin. “Using Fourier transform IR spectroscopy to analyze biological materials”. Nat Protoc. 2014. 9(8): 1771-1791.

20. T. Gao, J. Feng, Y. Ci. “Human breast carcinomal tissues display distinctive FTIR spectra: implication for the histological characterization of carcinomas”. Anal Cell Pathol. 1999. 18(2): 87-93.

21. P. Lasch, M. Boese, A. Pacifico, M. Diem. “FT-IR spectroscopic investigations of single cells on the subcellular level”. Vib Spectrosc. 2002. 28(1): 147-157.

22. J. Zhou, Z. Wang, S. Sun, M. Liu. H. Zhang. “A rapid method for detecting conformational changes during differentiation and apoptosis of HL60 cells by Fourier-transform infrared spectroscopy”. Biotechnol Appl Biochem. 2001. 33(2): 127-132.

23. A. Tfayli, O. Piot, A. Durlach, P. Bernard, M. Manfait. “Discriminating nevus and melanoma on paraffion-embedded skin biopsies using FTIR microspectroccopy”. Biochim. Biophys. Acta. 2005. 1724(3): 262-269.

24. M. Werner, A. Chott, A. Fabiano, H. Battifora H. “Effect of formalin tissue fixation and processing on immunohistochemistry”. Am J Surg Pathol. 2000. 24(7):1016-1019.

**Legends to figures**

Figure 1. Mean spectrum of 100 spectra for A549 (dotted lines) and CALU-1 (solid lines) cells obtained after preparation as cytospin and fixing them with methanol (2 bottom spectra) or formalin (2 top spectra) for the lipid region and the fingerprint region between 1350 cm-1 and 1800 cm-1.

Figure 2. Mean spectrum of 100 spectra for A549 (dotted lines) and CALU-1 (solid lines) cells obtained after preparation as smear and fixing them with methanol (2 bottom spectra) or formalin (2 top spectra) for the lipid region and the fingerprint region between 1350 cm-1 and 1800 cm-1.

Figure 3. PCA for A549 (triangles) and CALU-1 (squares) cells obtained after preparation as cytospin and fixing them with methanol (open triangles and open squares) or formalin (filled triangles and filled squares) for the lipid region (a) and the corresponding loadings (b).

Figure 4. PCA for A549 (triangles) and CALU-1 (squares) cells obtained after preparation as cytospin and fixing them with methanol (open triangles and open squares) or formalin (filled triangles and filled squares) for the fingerprint region between 1350 cm-1 and 1800 cm-1 (a) and the corresponding loadings (b).

Figure 5. PCA for A549 (triangles) and CALU-1 (squares) cells obtained after preparation as smear and fixing them with methanol (open triangles and open squares) or formalin (filled triangles and filled squares) for the lipid region (a) and the corresponding loadings (b).

Figure 6. PCA for A549 (triangles) and CALU-1 (squares) cells obtained after preparation as smear and fixing them with methanol (open triangles and open squares) or formalin (filled triangles and filled squares) for the fingerprint region between 1350 cm-1 and 1800 cm-1 (a) and the corresponding loadings (b).

**Table I.** Statistical significance between different types of sample preparation (cytospin *versus* smear) based on types of fixation (formalin, methanol) and cell type (A549, CALU-1).

|  |  |  |
| --- | --- | --- |
|  | Lipid region | 1800 cm‑1 to 1350 cm-1 region |
|  | PC1 | PC2 | PC1 | PC2 |
| Formalin |  |  |  |  |
| A549 | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |
| CALU-1 | **p<0.001** | **p<0.001** | **p<0.001** | p=0.395 |
|  |  |  |  |  |
| Methanol |  |  |  |  |
| A549 | **p<0.001** | **p=0.021** | **p<0.001** | p=0.071 |
| CALU-1 | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |

**Table II.** Statistical significance between different types of fixation (formalin *versus* methanol) based on sample preparation (cytospin, smear) and cell type (A549, CALU-1).

|  |  |  |
| --- | --- | --- |
|  | Lipid region | 1800 cm‑1 to 1350 cm-1 region |
|  | PC1 | PC2 | PC1 | PC2 |
| Cytospin |  |  |  |  |
| A549 | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |
| CALU-1 | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |
|  |  |  |  |  |
| Smear |  |  |  |  |
| A549 | **p<0.001** | **p<0.001** | p=0.125 | **p<0.001** |
| CALU-1 | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |

**Table III.** Statistical significance between the different cell types (A549 *versus* CALU-1) based on sample preparation (cytospin, smear) and fixative (formalin, methanol).

|  |  |  |
| --- | --- | --- |
|  | Lipid region | 1800 cm-1 to 1350 cm-1 region |
|  | PC1 | PC2 | PC1 | PC2 |
| Cytospin |  |  |  |  |
| Formalin | **p<0.001** | **p<0.001** | **p<0.001** | **p=0.001** |
| Methanol | **p<0.001** | **p<0.001** | **p<0.001** | **p<0.001** |
|  |  |  |  |  |
| Smear |  |  |  |  |
| Formalin | **p=0.038** | p=0.162 | **p=0.008** | p=0.298 |
| Methanol | **p<0.001** | **p<0.001** | **p<0.001** | p=0.099 |

**Figure 1**

**2850 cm-1**

**2920 cm-1**

**Absorbance**

**Absorbance**

**Wavenumber (cm-1)**

**Wavenumber (cm-1)**

**Figure 2.**

**2850 cm-1**

**2920 cm-1**

**Absorbance**

**Absorbance**

**Wavenumber (cm-1)**

**Wavenumber (cm-1)**

**Figure 3a**

**PC 2 (4%)**

**PC 1 (88%)**

**Figure 3b.**

**PC2**

**PC1**

**Wavenumber (cm-1)**

**Figure 4a**

**PC 2 (14%)**

**PC 1 (25%)**

**Figure 4b**

**PC2**

**PC1**

**Wavenumber (cm-1)**

**Figure 5a**

**PC 2 (16%)**

**PC 1 (67%)**

**Figure 5b**

**Wavenumber (cm-1)**

**PC1**

**PC2**

**Figure 6a**

**PC 1 (26%)**

**PC 2 (22%)**

**Figure 6b**

**Wavenumber (cm-1)**

**PC1**

**PC2**