A FTIR microspectroscopic study of the uptake and metabolism of isotopically labelled fatty acids by metastatic prostate cancer

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1. Introduction

In the United Kingdom, prostate cancer (CaP) is the most commonly diagnosed cancer in men and the second most common cause of cancer related death of men [1]. A key factor directly responsible for considerable morbidity and mortality in CaP patients is metastases to the bone marrow stroma (BMS). Several epidemiological and in vitro molecular studies have demonstrated that fatty acids (FAs) play a significant role in the progression and development of CaP [3,4]. Dennis et al. [2] conducted a detailed retrospective review of 29 epidemiological studies investigating this relationship and identified associations of advanced CaP with intake of total and saturated fat. This is supported by the recent World Cancer Research Fund inquiry, which also reports that total body fat is a key factor in the development of prostate cancer (CaP) progression or prevention remains unclear through patient based studies as a result of inconsistencies in study design. Thus, there is a need for controlled systematic laboratory based studies to facilitate and potentially focus epidemiological investigations.

We have previously used imaging Fourier-Transform infrared (FTIR) microspectroscopy to demonstrate that metastatic CaP cells in bone marrow biopsy tissue with close proximity to adipocytes, give rise to higher lipid hydrocarbon signals relative to those CaP cells situated further away from these adipocyte-rich locations [6]. Adipocytes are specialized for the synthesis and storage of FAs as triacylglycerides (TAGs) as well as for FA mobilisation through lipolysis [7].

This was followed by in vitro co-culture experiments of PC-3 cells (CaP cell line derived from bone metastases) with human BMS. These experiments also showed that PC-3 cells preferentially located to adipocytes in the presence of numerous haemopoietic cell types and that these tumour cells accumulated numerous cytoplasmic lipid droplets, revealed by Oil-red O staining for TAG and Nile Red real time fluorescent microscopy [6]. A quantitative measure of this attraction was achieved using Boyden transwell invasion assays. Using this method, we found that invasion of PC-3 cells towards BMS was significantly reduced.
when the experiment was repeated with BMS grown in the absence of hydrocortisone so that adipocyte formation was inhibited [6].

Together, the above findings revealed that PC-3 cells are attracted to adipocytes and that these tumour cells exhibit high lipid signals. From these observations we asked: Could the high levels of lipids measured in metastatic CaP cells in close proximity to adipocytes, be the result of adipocyte to CaP cell lipid translocation or an increased expression of lipogenic enzymes for de novo lipid biosynthesis in the tumour cell? The latter can be induced by growth factors [8] that are also released by adipocytes [9]. To investigate whether lipids stored within adipocytes could be translocated to PC-3 cells, we grew mesenchymal stem cells in medium containing deuterated palmitic acid (D\textsubscript{31}-PA) during their differentiation into adipocytes. The mature adipocytes, comprising of numerous lipid droplets containing D\textsubscript{31}-PA, were then incubated for a further 48 h in medium containing PC-3 cells and no D\textsubscript{31}-PA. Following this period, the co-culture was fixed and analysed using imaging FTIR microspectroscopy. In the mid-IR spectral window, deuterated FAs can provide a C–D vibrational signal that does not overlap with signals arising from the functional group vibrations of endogenous FAs or other biomolecules. This specific spectral marker for the labelled FA (or its metabolites), was not only observed in high intensity in the adipocytes, confirming D\textsubscript{31}-PA uptake and storage in these cells, but was also found to illuminate PC-3 cells following background subtraction. The study demonstrated for the first time, without cell isolation and lipid extraction, that tumour cells could uptake lipid stored within adipocytes. In turn, this highlighted a potential clinical significance for the management of CaP through diet.

The importance of the role of lipids in the metastatic progression of CaP was further established by our group through biomolecular profiling of metastatic CaP cells in primary bone marrow tissue sections derived from different patients, with FTIR and histochemistry. This was carried out with comparative biomolecular profiles obtained from high-grade prostate organ-confined CaP [10]. This study identified the dominant metabolic processes driving the proliferation of metastatic cells in each of the biopsies studied: Glycerophospholipid synthesis from TAG, glycolipid synthesis as well as glycolysis.

The focus of the present study is to follow on from the above experimental findings by investigating the metabolism of specific isotopically labelled unsaturated and saturated FA’s in PC-3 cells in a time-dependent manner using FTIR microspectroscopy. Metabonomic analysis through FTIR microspectroscopy enables a survey of several biomolecules simultaneously (such as lipids, phosphorylated biomolecules and protein secondary structure) to generate hypotheses that may refine targeted molecular studies. The use of deuterated analogues to investigate metabolic activity in tissue has been demonstrated by Wetzel et al. [11]. The authors used deuterated water as an alternative to radioisotopes to study the metabolic activities in various layers of the cerebellum in rat tissue following fixation.

In the present study, arachidonic acid (AA) was chosen as the candidate unsaturated FA since this omega-6 poly-unsaturated FA (ω-6 PUFA) has been shown to increase the risk of CaP by promoting the proliferation of malignant prostate epithelial cells. These effects are mediated by the increase in the cyclooxygenase-2 (COX-2) derived product, prostaglandin E\textsubscript{2} (PGE\textsubscript{2}) [12] and the lipoxygenase (LOX) derived products, 5 and 12(S)-HETE (hydroxyeicosatetraenoic acid). Of these eicosanoids, we have previously demonstrated PGE\textsubscript{2} to be a potent stimulator of PC-3 invasion [6].

As well as investigating the uptake and metabolic consequences of AA in metastatic CaP, we also study these parameters with a saturated FA, palmitic acid (PA; C16:0). PA is not an essential FA (unlike AA) and is one of the five most abundant FAs comprising adipose tissue or blood serum [13]. Molecular and ecological studies have shown a positive association of saturated fat intake with increased risk and mortality from CaP [13–15]. A molecular based patient study by Mamalakis et al. [16] found a significantly elevated adipose saturated FA composition in CaP patients as opposed to benign prostate hyperplasia (BPH) patients. Interestingly, the same study also found significantly decreased prostate tissue sum saturated FAs and elevated sum monounsaturated FAs in cancer as opposed to BPH patients. Mamalakis et al. [16] associated these findings to studies at the cellular level demonstrating that a decreased saturation index (SI; the ratio of saturated to mono-unsaturated FAs) is associated with increased tumour cell fluidity and malignant transformation. Mamalakis et al. [16] also reported that the most pronounced difference between cancer patients and BPH patients was a three-fold elevated level of the desaturated FA metabolite, palmitoleic acid (C16:1), in the former group, highlighting a possible role of this FA in the neoplastic processes.

2. Experimental

2.1. Cell culture

PC-3 cells were cultured in Ham’s F12, 7% FCS and 1% L-Glutamine on MirrIR slides until 70% confluent. Following this, one slide of cells were starved in serum-free RPMI media with 1% L-Glutamine for 24 h and then fixed in 4% formalin in PBS for 25 min prior to washing in deionised water for 3 s to remove the residue PBS [17]. This represented the control slide (no exposure to labelled FA) at 0 min. The remaining PC-3 cells were also starved in serum-free RPMI media with 1% L-Glutamine for 24 h. These cells were subsequently washed twice in PBS and then exposed to serum-free RPMI with 1% L-Glutamine, supplemented with 25 μM or 100 μM D\textsubscript{31}-AA (Sigma–Aldrich, UK) or 20 μM or 50 μM D\textsubscript{31}-PA (CK Gas Products Limited, UK) for 15, 30, 45, 60, 90, 120, 180 and 1440 min. Control slides of PC-3 cells cultured in media with no D\textsubscript{31}-PA or D\textsubscript{31}-AA were also prepared for each time-point. The slides were removed from culture media at each of the above time-points, washed twice with PBS, and fixed using the same protocol as described above.

2.2. FTIR microspectroscopy

A sampling aperture size of 600 μm × 600 μm was used to collect IR spectra (n) from 5 locations within each slide. This resulted in FTIR spectra that were representative of the average biochemistry across approximately ~200 cells at each sampling location, thus ~1000 cells/sampling time-point. FTIR spectra were recorded in reflectance mode using 256 scans at 4 cm\textsuperscript{-1} spectral resolution. A background spectrum was collected from a clean MirrIR slide, using 256 scans at 4 cm\textsuperscript{-1} spectral resolution. FTIR spectra were baseline corrected and normalised to the intensity of the amide I peak.

High definition FTIR microspectroscopic maps were collected at 6.25 μm pixel-resolution in rapid-scan mode using a PerkinElmer Spotlight spectrometer with a 16 × 1 MCT linear array detector. Mid-IR spectra within the range 6000–700 cm\textsuperscript{-1} were obtained in reflection mode. The background scan was recorded at 8 cm\textsuperscript{-1} spectral resolution with 75 scans, whereas the sample scan was recorded at 8 cm\textsuperscript{-1} spectral resolution with 64 scans.

2.3. Statistics

Standard errors of the mean (±) were calculated for IR measurements collected from different locations on the same
3. Results and discussion

3.1. Uptake of palmitic acid by PC-3 cells

Firstly, we studied the influence of different growth medium compositions on FA uptake using D31-PA. Three different culture media were investigated: (i) Ham’s F-12 with 7% fetal bovine serum (FBS) and 1% l-Glutamine, (ii) RPMI with 7% FBS and 1% l-Glutamine and (iii) RPMI with 1% l-Glutamine. Both RPMI and Ham’s F-12 are complex mixtures consisting of a range of inorganic salts, amino acids, vitamins, nucleotides and glucose as well as small-molecule precursors. However, it is the Ham’s F-12 growth medium with FCS and l-glutamine, which is recommended by the European Collection of Cell Cultures (ECACC) for optimum growth of PC-3 cells [18]. RPMI is recommended by ECACC for culture of CaP cell lines, LNCaP (derived from lymph node metastases) and PNT2-C2 (immortalised normal prostate epithelial cell line) but contains no FAs, unlike Ham’s F-12, which contains the ω6-FA, linoleic acid (LA). Fig. 1A shows FTIR measurements demonstrating the influence of these media on D31-PA uptake by PC-3 cells following 24 h incubation with 50 μM D31-PA. This was carried out by baseline integration of the νas(CO2+) peaks in the spectral region 2250–2050 cm⁻¹ and using the resulting peak areas.

In Fig. 1A we find that FBS supplementation into the media resulted in a significant reduction in D31-PA uptake compared with cells incubated in media without FBS (p < 0.05). The main constituent of FBS is the globular protein, albumin, which may influence D31-PA uptake by reducing albumin bound D31-PA interactions with the plasma membrane, in-turn preventing passive diffusion of D31-PA through the plasma membrane and/or interactions of D31-PA with membrane-bound FA transporter proteins [19]. FBS is also rich in an array of other proteins, FAs and growth stimulatory factors [20] that may support growth and circumvent the requirement for D31-PA uptake. Therefore, to study the uptake and metabolism of D31-PA by PC-3 cells, we used the simplified medium composition of RPMI in the absence of FBS, thereby FA free media, enabling accurate measurement of D31-PA uptake and metabolism by PC-3 cells without the background contamination of undefined amounts of FA from the media.

The subcellular preservation and localisation of D31-PA into PC-3 cells was assessed using FTIR microspectroscopic imaging. Fig. 1B shows an optical photomicrograph of PC-3 cells on MirrIR substrate, following exposure to 50 μM D31-PA for 24 h. Fig. 1C shows the distribution of the integrated intensity of the phosphate diester (νas(PO2+);1274–1181 cm⁻¹) peak area. For cells 1 and 2 in the optical image, it is clearly observed that the most intense phosphate signals localise at the nuclei due to the high concentration of nucleic acids present at this location. The integrated lipid hydrocarbon peak area (νas(C-H2+3); 3004–2837 cm⁻¹) signal localises with greater intensity at the cytoplasmic locations due to the presence of cellular organelles. Accepting that the cell thickness does not change significantly across the cell, we observe that the most intense νas(CO2+) signal co-localises with the lipid hydrocarbon signal at the cytoplasmic locations, suggesting the subcellular localisation of D31-PA or its metabolites is predominately in the cytoplasm. To demonstrate spectral quality, Fig. 1F shows raw FTIR spectra taken from the nucleus (Location 1) and cytoplasm (Location 2) of PC-3 cell 2 (See Fig. 1E).

Fig. 2 shows a graph of intracellular νas(CO2+) signal intensities as a function of time in PC-3 cells incubated with 20 μM or 50 μM D31-PA. A pairwise comparison of νas(CO2+) signal intensities at each time-point for cells incubated with 20 and 50 μM D31-PA reveals a significant difference only at 24 h (1440 min) with p = 0.00. It is possible that this difference at 1440 min may be the result of total/near complete D31-PA uptake by the PC-3 cells from the media that was initially supplemented with 20 μM D31-PA. In contrast, at the 1440 min time-point, the 50 μM D31-PA supplemented media may still contain available D31-PA for cell consumption and this may reflect the high intracellular νas(CO2+) signal intensity measured from these cells.

3.2. The temporal effects of D31-PA exposure to endogenous biomolecules in PC-3 cells

The FTIR technique enables us to probe fluctuations in several biomolecules simultaneously. Thus, in addition to monitoring D31-PA uptake directly using the νas(CO2+) signal, it is also possible to temporally determine the effects of this exposure on phosphate activity (integrated peak area 1274–1181 cm⁻¹), protein secondary structures (amide I frequency) and endogenous-non-isotopically labelled lipid biosynthesis (integrated peak area 3004–2837 cm⁻¹). The temporal fluctuations in these biomolecular domains were studied in PC-3 cells following incubation with 20 and 50 μM D31-PA in serum-free culture media (RPMI) and compared with the control (PC-3 cells incubated in identical conditions but without D31-PA) (Fig. 3A–C). Fig. 3D shows a typical raw FTIR spectrum taken from the dataset to illustrate baseline correction and integration to obtain peak area values for the lipid and phosphate signals.

Firstly, statistical tests (two-way ANOVA followed by Tukey multiple comparisons test) were used to identify whether each of the IR signals (protein amide I frequency shifts, endogenous lipid intensities, phosphate intensities) obtained over the 24 h period for control cells were significantly different from those obtained from D31-PA stimulated cells. These tests were also carried out to determine whether the measured signals were different between cells stimulated with high and low concentrations of D31-PA. Using this statistical test we found that combinations of control vs. 20 μM D31-PA stimulated cells, control vs. 50 μM D31-PA stimulated cells and 20 μM D31-PA stimulated cells vs. 50 μM D31-PA stimulated cells gave rise to protein shifts and phosphate signals that were significantly different with p ≤ 0.05. For the lipid signal, we found that all of the combinations stated above were also significantly different (p ≤ 0.05) except for the control vs. 20 μM D31-PA stimulated cells (p = 0.252), this will be discussed further below. These results demonstrate that addition of D31-PA has a significant effect on the biochemistry of the cells over the duration of the time-course experiment compared with the control.
Following this assessment, fluctuations in the different biomolecular domains between time-points for each culture condition were studied. We found that the endogenous lipid signal in the control PC-3 cells initially fell and is induced by metabolic/cytokine/growth factor imbalance resulting from the exchange of media to serum-free RPMI media at the zero minute time-point (Fig. 3A). Conversely, cells incubated with D$_{31}$-PA exhibited a rapid rise in endogenous lipids within 15 min of exposure. Since the incubation media (RPMI) contains no FAs, this increase in lipid content must be due to *de novo* biosynthesis. This correlates with previous reports demonstrating that the majority of FAs in tumour cells are derived from *de novo* FA synthesis and in contrast to normal tissue, human tumours display a higher rate of *de novo* FA synthesis [8]. Also, it has been shown that lipotoxicity from accumulation of long chain FAs is specific for saturated FAs such as PA [21]. This selectivity arises from the generation of molecules such as reactive oxidative species or the *de novo* synthesis of ceramide, among others [21]. Listenberger et al. [21]

![Image of FTIR microspectroscopy results](image-url)
have shown that in Chinese hamster ovary (CHO) cells, endogenously generated unsaturated FAs rescue PA-induced apoptosis by promoting PA incorporation into TAC. It may be the case that this mechanism is initiated in the PC-3 cells shortly following exposure to D31-PA and is responsible for the significant up-regulation in endogenous lipids. To the author’s knowledge, this mechanism has not previously been proposed or demonstrated for CaP cells.

For PC-3 cells incubated with 50 μM D31-PA, the initial rise in endogenous lipid signal was followed by a fall, which we attribute to metabolic breakdown into adenosine triphosphate (ATP), which is a major product of lipid metabolism. This notion is supported by a phosphate spike at 30 min (Fig. 3B) accompanied by a marked shift in the amide I frequency (Fig. 3C), together indicating protein phosphorylation. Although protein phosphorylation is also observed in the control cells at a later time-point (45 min, Fig. 3C), it is not known at this juncture whether the proteins that are phosphorylated are the same as those phosphorylated when PC-3 cells are incubated with D31-PA. This requires further metabolomic and phosphopeptide-based investigations.

Earlier in this section we reported that the overall trend in endogenous lipid signal between control vs. 20 μM D31-PA stimulated cells were not significantly different. However, we can identify significant differences in endogenous lipid signals between these conditions at earlier time-points (30 and 45 min). Together, this data indicates an expected smaller perturbation to the metabolites in cells exposed to the lower concentration of D31-PA compared with exposure to higher concentrations. The endogenous lipid intensity in cells exposed to 20 μM D31-PA demonstrates a maximum at 45 min post D31-PA induction (Fig. 3A). This time-delay in reaching a lipid maximum, compared with cells incubated with 50 μM D31-PA, could suggest that if the uptake of D31-PA into the cell is greater with higher gradients, then the critical-point at which lipid biosynthesis is initiated to sequester excess D31-PA must also be delayed for culture media containing lower concentrations of D31-PA. Interestingly, the 50 μM D31-PA exposed cells show a significantly higher maximum lipid signal compared with the 20 μM D31-PA exposed cells, within this period (0–45 min). This suggests that endogenous lipid demand is greater in cells incubated with higher concentrations of D31-PA compared with cells incubated in lower concentrations of D31-PA. This supports the assumption that a greater influx of D31-PA requires for a greater number of synthesised lipids. Unfortunately, it was not possible to measure the \( v_{\text{VCD}}(\text{CD}_{2+3}) \) intensity at time-points below 45 min since the amount in the cells was below the detection limit with this experimental set-up.

The anticorrelation of lipid with phosphate intensity was not only observed in cells incubated with 50 μM D31-PA, but also for cells incubated with 20 μM D31-PA between 45 and 90 min (Fig. 3B). However, unlike the control and cells exposed to 50 μM D31-PA, the rise in phosphate signal did not induce a concomitant change in bulk protein secondary structure. In contrast, we observed a more gradual change in protein secondary structure over-time for cells incubated with 20 μM D31-PA. It may be the case that these phosphates are partitioned between other cellular processes such as phospholipid generation as well as protein phosphorylation, resulting in a more subtle change to the latter.

FTIR microspectroscopy can directly probe structural modifications of D31-PA resulting from its metabolism by the PC-3 cells. This is carried out using the \( v_{\text{VCD}}(\text{CD}_{2})/v_{\text{VCD}}(\text{CD}_{3}) \) signal intensity, which can determine changes in D31-PA chain length (higher \( v_{\text{VCD}}(\text{CD}_{2})/v_{\text{VCD}}(\text{CD}_{3}) \) values indicate longer FA chains). The inset in Fig. 4 shows the method by which the \( v_{\text{VCD}}(\text{CD}_{2}) \) and \( v_{\text{VCD}}(\text{CD}_{3}) \) values are obtained, which is based on peak height. Note that for PC-3 cells incubated with 20 μM D31-PA, we found that the subtle \( v_{\text{VCD}}(\text{CD}_{3}) \) signal that exists as a shoulder on the \( v_{\text{VCD}}(\text{CD}_{2}) \) peak was not resolvable in spectra obtained at the 90 min and 120 min sampling.

Fig. 2. A graph to show mean intracellular \( v_{\text{VCD}}(\text{CD}_{2+3}) \) signal intensities (±SE) as a function of time in PC-3 cells incubated with 20 μM or 50 μM D31-PA. Standard error of the mean bars are displayed. Test for significance between \( v_{\text{VCD}}(\text{CD}_{2+3}) \) signals were obtained for 20 and 50 μM D31-PA stimulated cells at each time-point, where * is significant when \( p \leq 0.05. \)

Fig. 3. Temporal fluctuations in various biomolecular domains probed by FTIR, for PC-3 cells exposed to 20 μM or 50 μM D31-PA or no D31-PA (Control). (A) Endogenous mean lipid hydrocarbon peak area intensities (±SE); (B) mean phosphate diester peak area (\( v_{\text{PO}}(\text{PO}) \)) intensities (±SE); (C) amide I frequency shifts (±SE). * Significance test values between time-points are displayed, where * is significant when \( p \leq 0.05. \)
time-points. Fig. 4 shows that PC-3 cells incubated with 20 μM D31-PA for 3 h exhibit significantly higher ν(CD) values (p = 0.03) compared with cells exposed to 50 μM D31-PA for 90 and 180 min (p = 0.02, respectively). This data indicates lower metabolism of D31-PA in PC-3 cells exposed to 20 μM D31-PA compared with cells incubated with 50 μM D31-PA. This low rate of D31-PA metabolism may be due to the requirement for lipid derived products being satisfied by metabolism of endogenous lipids, which we find to provide lower values (≥60 min) in the cells exposed to 20 μM D31-PA for 1440 min compared with cells exposed to 50 μM D31-PA (Fig. 3A). At 1440 min post D31-PA induction, we find that there is a significant reduction in the ν(CD) values for cells stimulated with 20 μM D31-PA, generating ν(CD) values similar to those stimulated with 50 μM D31-PA for 1440 min (Fig. 4). Thus, D31-PA initially stored upon exposure may be metabolised eliciting the reduction in ν(CD) values between 90 and 1440 min time-points for the 20 μM D31-PA stimulated cells. This reduction supports the inference made above, in which we hypothesised significant depletion of D31-PA in the culture media between 90 and 1440 min. Thus, between these time-points metabolised D31-PA may not have been replenished in the cells.

Note that in the above experiments, the reproducibility of FTIR signals obtained from cells within each time-point has been measured by the standard error. Further investigations are required to address the reproducibility of temporal fluctuations in FTIR signals from cells exposed to D31-PA in independent experiments.

3.3. Uptake and metabolism of arachidonic acid by PC-3 cells

Previously, we determined the rate of AA uptake into PC-3 cells since it is known to decrease in the adipose tissue of CaP patients [6]. In that experiment, PC-3 cells were cultured in the presence of 10 μM AA and stained with Nile Red for fluorescence-assisted cell sorting (FACS) analysis. Nile Red is a fluorescent tracer that stains all neutral lipids such as TAG, cholesteryl esters (lipoproteins) and phospholipids [22]. Using this method, we observed rapid uptake of AA from the local environment, reaching a maximum level at 45 min. Following this, we observed decay in the fluorescence signal that suggested metabolism of AA into PG, 2E or HETEs.

To compliment the Nile Red experiments, we studied uptake and metabolism of AA using FTIR and deuterated AA (D8-AA) (Fig. 5). This was carried out because Nile Red is not specific to AA and excludes the measurement of free FAs. The latter is important for including into measurements of AA intracellular concentrations, since unesterified AA is the structural form of this FA upon initial uptake into the PC-3 cell or following release from plasma membrane stores by phospholipases A2 [23]. D8-AA uptake and or that of its metabolites were followed using the ν(C–D) intensity by integration of the ν(C–D) peak at ~2250 cm⁻¹ (Fig. 5A). This signal appears ~54 cm⁻¹ higher than the ν(CH) peaks of D31-PA. Since the deuterium atoms in D31-PA are located along the carbon chain, they can give rise to the ν(CD) and ν(ν(CD)) vibrations, as well as to the ν(ν(CD)) and ν(ν(CD)) vibrations from the terminal carbon. In contrast, the deuterium atoms in D8-AA are located only at the double bonds (Fig. 5A). The unique vibrational signatures for these lipids, highlights the potential pharmokinetic use of FTIR to simultaneously study uptake and metabolism of both saturated and unsaturated FAs in tumour cells.

Fig. 5B shows the mean intracellular ν(C–D) signal intensities as a function of time in PC-3 cells incubated with 25 μM or 100 μM D8-AA and stained with Nile Red for fluorescence-assisted cell sorting (FACS) analysis. Nile Red is a fluorescent tracer that stains all neutral lipids such as TAG, cholesteryl esters (lipoproteins) and phospholipids [22]. Using this method, we observed rapid uptake of AA from the local environment, reaching a maximum level at 45 min. Following this, we observed decay in the fluorescence signal that suggested metabolism of AA into PG, 2E or HETEs.
D$_8$-AA. We found that the $\nu(\mathrm{C-D})$ signals could be measured at time-points $\geq$ 30 min. Prior to 30 min, signal intensity of the $\nu(\mathrm{C-D})$ peak could not be detected. We observe a significant rise in $\nu(\mathrm{C-D})$ signal between 30 and 90 min for 100 $\mu$M D$_3$-AA exposed cells. This rise is also demonstrated by the mean $\nu(\mathrm{C-D})$ signal for cells incubated with 25 $\mu$M D$_3$-AA, although not significant at $p = 0.05$. Following this, there is a fall in the mean $\nu(\mathrm{C-D})$ signal between 90 and 120 min for both 25 and 100 $\mu$M D$_3$-AA exposed cells. Although this fall is not significant, there does exist a significant increase in $\nu(\mathrm{C-D})$ signal between 120 and 180 min for both 25 and 100 $\mu$M D$_3$-AA exposed cells. There is no significant difference in the $\nu(\mathrm{C-D})$ signal between 90 and 180 min or 180 min and 24 h for cells exposed to 25 and 100 $\mu$M D$_3$-AA.

The time-point of maximum AA uptake prior to metabolism was determined as 45 min by the Nile Red method [4]. However, the FTIR data demonstrates a maximum at 90 min (Fig. 5B). This difference is due to the different molecules being measured by FTIR compared with the Nile Red technique, since FTIR records signals from stored intracellular AA as well as its metabolic products that may be synthesised following the 45 min time-point, accumulatively contributing to a net increase in $\nu(\mathrm{C-D})$ signal at 90 min. The AA-derived metabolite fraction cannot be detected using the Nile Red measurement. The fall in $\nu(\mathrm{C-D})$ signal post 90 min D$_8$-AA stimulation, can be due to the efflux of D$_8$-AA-derived eicosanoids such as PGE$_2$, from the PC-3 cells into the extracellular environment, to convey their biological effects in an autocrine or paracrine manner. Supporting evidence for this is given by Hughes-Fulford [24] and Chen et al. [25], who observed a significant increase in COX-2 protein activity and PGE$_2$ synthesis within three hours of exposure to AA. In comparison with lipid uptake data obtained from PC-3 cells incubated with D$_3$-PA (Fig. 2), we do not find any reduction in $\nu_{\text{CD}}(\mathrm{CD_2+3})$ Signal over the time-points sampled for these cells. In-turn, this indicates the absence of D$_3$-PA derived metabolites secreted from the cell.

At 180 min, we observe a second rise in $\nu(\mathrm{C-D})$ signal, reaching a level that is not significantly different from that measured at the 90 min time-point (Fig. 5B). This indicates regulated uptake of D$_8$-AA. Supporting this notion is the comparison of $\nu(\mathrm{C-D})$ signal intensities for PC-3 cells exposed to 25 and 100 $\mu$M D$_3$-AA over all time-points. Two-way analysis of variance was applied to the data set in Fig. 5B to test whether there exists a significant difference in $\nu(\mathrm{C-D})$ signals obtained from cells cultured in 25 and 100 $\mu$M D$_3$-AA, over identical time-points. This confirmed that combinations of culture media (with 25 $\mu$M or 100 $\mu$M D$_3$-AA) and time gave rise to no significant difference ($F = 0.097, p = 0.996$) in $\nu(\mathrm{C-D})$ signals. Thus, the amount of D$_8$-AA in PC-3 cells is tightly regulated, even in the presence of high concentrations of D$_3$-AA in the extracellular environment. Stringent control of intracellular unesterified AA has previously been reported for other mammalian cells [23].

**4. Conclusions**

In this paper we have demonstrated that FTIR microspectroscopy can be used to measure the relative uptake of a labelled FA in tumour cells under different environmental conditions, without cell isolation or lipid extraction and enrichment. We observed that incorporation of D$_3$-PA into the PC-3 cell is not as tightly regulated as was found for cells incubated with D$_8$-AA, since the level of D$_3$-PA in PC-3 cells rose as a function of availability in the culture medium. This study also confirms our previous findings [6] and others [12] that AA is rapidly taken up from the extracellular environment and metabolised within three hours. This can be correlated with the low levels of AA found in prostate cancer patients.

FTIR microspectroscopy is well-suited for surveying several biomolecular domains simultaneously for generating hypotheses to refine targeted molecular studies. This was demonstrated by temporal FTIR analysis of D$_8$-AA uptake and metabolism, where we can identify the time-point corresponding to significant eicosanoid generation, as well as differences in the response of PC-3 cells to low and high concentrations of D$_8$-AA or D$_3$-PA. We found that stimulation of PC-3 cells with the saturated FA, D$_3$-PA, resulted in significant de novo lipid biosynthesis, which may be associated with D$_3$-PA channelling into TAG pools. The time taken to develop these lipid pools is dependent on the dose of D$_3$-PA administered to the PC-3 cells. D$_3$-PA stimulation also results in significant protein phosphorylation within 3 h of exposure to PC-3 cells, which may be associated with the metabolic breakdown of these biosynthesised lipids. Thus, if the formation of TAG pools is required for PC-3 cell survival to high intracellular concentrations of D$_3$-PA or D$_8$-AA, as is the case for CHO cells [21], then lipogenic enzymes involved in the synthesis of TAG may be targets for therapeutic intervention.

**Acknowledgements**

We gratefully thank the Association of International Cancer Research (AICR) for funding (E.G) and Dr. James Nicholson (Daresbury Laboratory, UK) for use of the Spotlight instrument.

**References**


[23] F.H. Chilton, A.N. Fonteh, M.E. Surette, M. Triggiani, J.D. Winkler, Research (AICR) for funding (E.G) and Dr. James Nicholson (Daresbury Laboratory, UK) for use of the Spotlight instrument.
