Cellular injury evidenced by impedance technology and infrared microspectroscopy

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**Key words:**

FTIR, RT-CES, cytostatic, cytotoxic, cancer

**Highlights:**

- FTIR detects different types of cell injury in a manner comparable to RT-CES.
- Spectral differences distinguish cytostasis from cytotoxicity.
- FTIR microspectroscopy detects the reversibility of cytostasis.

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Abstract

Fourier Transform Infrared (FTIR) spectroscopy is finding increasing biological application, for example in the analysis of diseased tissues and cells, cell cycle studies and investigating the mechanisms of action of anticancer drugs. Cancer treatment studies routinely define the types of cell-drug responses as either total cell destruction by the drug (all cells die), moderate damage (cell deterioration where some cells survive) or reversible cell cycle arrest (cytostasis). In this study the loss of viability and related chemical stress experienced by cells treated with the medicinal plant, *Plectranthus ciliatus*, was investigated using real time cell electronic sensing (RT-CES) technology and FTIR microspectroscopy. The use of plants as medicines is well established and ethnobotany has proven that crude extracts can serve as treatments against various ailments. The aim of this study was to determine whether FTIR microspectroscopy would successfully distinguish between different types of cellular injury induced by a potentially anticancerous plant extract. Cervical adenocarcinoma (HeLa) cells were treated with a crude extract of *Plectranthus ciliatus* and cells monitored using RT-CES to characterize the type of cellular responses induced. Cell populations were then investigated using FTIR microspectroscopy and statistically analysed using One-way Analysis of Variance (ANOVA) and Principal Component Analysis (PCA). The plant extract and a cancer drug control (actinomycin D) induced concentration dependent cellular responses ranging from nontoxic, cytostatic or cytotoxic. Thirteen spectral peaks (915 cm\(^{-1}\), 933 cm\(^{-1}\), 989 cm\(^{-1}\), 1192 cm\(^{-1}\), 1369 cm\(^{-1}\), 1437 cm\(^{-1}\), 1450 cm\(^{-1}\), 1546 cm\(^{-1}\), 1634 cm\(^{-1}\), 1679 cm\(^{-1}\) 1772 cm\(^{-1}\), 2874 cm\(^{-1}\) and 2962 cm\(^{-1}\)) associated with cytotoxicity were significantly (p value < 0.05, one way ANOVA, Tukey test, Bonferroni) altered, while two of the bands were also indicative of early stress related responses. In PCA, poor separation between nontoxic and cytostatic responses was evident while clear separation was linked to cytotoxicity. RT-CES detected morphological changes as indicators of cell injury and could distinguish between viable, cytostatic and cytotoxic responses. FTIR microspectroscopy confirmed that cytostatic cells were viable and could still recover while also describing early cellular stress related responses on a molecular level.
Graphical abstract
1. Introduction

Fourier Transform Infrared spectroscopy is a label free, non-destructive, high throughput method which is growing in popularity as the applications increase. The applications of the technique include structure elucidation of compounds [1] and peptides [2,3], identification of different bacterial strains [4,5] and antimicrobial compounds affecting bacterial viability [6], the discrimination of diseased and healthy tissues or cells [7–9], cell cycle analysis [10], the effects of known toxins on cells [11] and the identification of molecular targets of natural and synthetic anticancer drugs [12–15].

Anticancer drugs under development are generally tested in vitro to determine whether they can be classified as non-toxic (no or limited cell death caused), cytostatic (arrest of cell proliferation but the cells are essentially still alive) or cytotoxic (considerable cell death induced by the treatment). Cytostatic responses are usually employed to stop uncontrolled tumour growth while cytotoxic responses lead to cell death caused by apoptosis, necrosis or cell lysis [16]. Cytostatic and cytotoxic behaviour plays an important role in the progression of a drug into preclinical and clinical studies. Cytostatic drugs have been found to be metastases preventative and increase the survival rates of patients by delaying the time to disease progression [17]. Cytotoxic drugs, on the other hand, are less selective and the induced cell death could be more detrimental to the patient in terms of side effects. Rixe and Fojo (2007) found that a drug elicited either cytostatic or cytotoxic cellular responses depending on its active concentration [17].

Classifications of cytostasis / cytotoxicity are usually decided following spectrophotometric assessment of cell viability in the presence of the drug and a labelling agent, but the use of spectroscopy in the absence of labels is gaining momentum. Conventionally, biochemical labels such as trypan blue, tetrazolium salts and fluorochromes are used to measure a single parameter such as; membrane integrity, enzymatic reactions and intra- or extracellular markers. The major disadvantage of labels is that the experiment becomes an end point analysis measuring only one or a few parameters at a particular time point after which the cells are destroyed. Biochemical labels are also very expensive and can sometimes crossreact with other molecules. Accordingly, new label free methodologies that are less time-
consuming and more comprehensive in evaluating multiple cellular components should be developed.

One of the popular label free assays currently in use is RT-CES. The principle of the instrument is based on the growth of adherent cells on gold electrodes. Impedance changes associated with the cells behaviour is measured continuously. When cells initially attach to the electrodes the impedance (electrical resistance) increases, causing an increase in the unit less Cell Index (CI). The CI will increase as the resistance increases, which are due to the cells proliferating or spreading out within the culture well. When a toxin is added to the wells, the cellular response can be measured continuously. Thus, any change in cell size and shape is monitored by changes within the CI. Generally a decrease in CI values could be due to cell volume decreasing, cells rounding up during the activation of cell death or the detachment of cells which leads to a decrease in resistance measured by the electrodes [18]. This technique successfully characterized the type of cellular responses of importance in our research group [19,20] as well as others [16,21–23].

In a previous study, we found that Raman micro-spectroscopy could be used to pre-screen a plant extract for its anticancer activity against the proliferation of leukemic U-937 cells [24]. Since FTIR spectroscopy is complementary to Raman spectroscopy and Raman spectroscopy only interrogates one cell at a time, we set out to determine whether FTIR could distinguish viable from cytostatic and cytotoxic responses in a given cell population following exposure to a cell death inducer. Many publications focus on the cytotoxic drugs and plant extracts or compounds affecting FTIR spectra of treated cells [11,13,14,25,26] while little research had been done on cytostatic responses [15].

In the evaluation of FTIR spectroscopy as a tool for the screening of anticancer drug responses, a cell death inducer is required. In this case a South African traditional medicinal plant was selected. Medicinal plants have long been studied for their effectiveness in treating cancer. Crude extracts of medicinal plants are more potent cytotoxins because of the synergy of the many phytochemicals found in the mixture [24]. The cell death inducer in this investigation was a leaves extract of Plectranthus ciliatus. In traditional medicine, plants from this genus are used to treat ailments of the digestive tract, respiratory system, nervous system, inflammation, infections, pain and different types of cancers [27]. The plant’s
anticancer activity was previously demonstrated by our group [28]. The aim of this study was to determine whether FTIR microspectroscopy could successfully distinguish between different types of cellular injury induced by a potentially anticancerous plant extract.

2. Methods

2.1 Plant material and extraction

*Plectranthus ciliatus* leaves were harvested in the Manie van der Schijff botanical garden, Pretoria, South Africa. The leaves were authenticated and a voucher specimen deposited in the Schweickerdt Herbarium, Pretoria, South Africa. The crude extract of the leaves was prepared and kindly donated by Dr. T. P. Kapewangolo, Department of Biochemistry, University of Pretoria. The plant extraction procedure was done as previously described [20]. The crude extract was stored at 4°C until use. Stock solutions were freshly prepared before each experiment by dissolving the plant extract in cell culture tested dimethyl sulfoxide (DMSO) (Sigma, Germany).

2.2 Cell culture

Human cervix adenocarcinoma (HeLa) cells were purchased from Highveld Biological (Pty) Ltd. (Johannesburg, South Africa) and grown in standard culture flasks in a humidified incubator at 37°C and 5% CO₂. The growth media was Minimum Essential Medium (MEM), supplemented with 5% heat-inactivated foetal bovine serum (FBS) (Hyclone, Separations, Johannesburg, South Africa) and antimicrobial cocktail (100 U/mL penicillin, 100 μg/mL streptomycin and 250 μg/L fungizone) (Hyclone, Separations, Johannesburg, South Africa). The cells were maintained by sub-culturing the cells when 80-90% confluence was reached.

2.3 Real time cell analysis of plant extract treated HeLa cells

An RT-CES device (Roche Diagnostics, Mannheim, Germany) in a humidified incubator was used to observe the proliferation of the cells in the absence and presence of the selected plant extract. Cell titrations (5 000 – 60 000 cells / well) were done to determine the optimal cell number for the experiments. The specific concentration (1 x 10⁵ cells/well) where the optimal exponential growth curve was found, was used for further studies as recommended by Fonteh *et al.* (2011) [19]. Three different concentrations (0.5 x cytotoxic concentration of
a treatment affecting 50% of the cell population tested (CC$_{50}$), CC$_{50}$ and 2 x CC$_{50}$ – determined by the tetrazolium dye XTT, Table 1) of the treatments were tested alongside the positive control (actinomycin D). The plant extract was dissolved in DMSO (ensure that all components were solubilized) to a stock solution of 20 mg/mL, while actinomycin D was completely dissolved in only sterile distilled water to a stock solution of 1 mg/mL. Further dilutions were made in complete media before the assays were conducted. Controls; a background (media only, no cells), untreated cells (cells and media) and a vehicle control (DMSO at 0.09%) were included every time. Electronic impedance increases as cells attach to the electrodes and is known as the Cell Index (CI). The cells were incubated in a humidified incubator for approximately 18 hours where the unit less CI $\approx$ 1, after which the treatments were added to the E-plates, final volume 250 μL. Cells were monitored for a total of 90 hours. Data sets were normalized at the point where cells were treated. The Cell Index was normalized using a build in function of the Roche xCelligence Software 1.2.1. The normalized CI is calculated automatically for each well as the CI at a given time point, divided by the CI at the normalized time point (Normalized CI$_{\text{time}}$ = CI$_{\text{time}}$ / CI$_{\text{normalized\_time}}$). Therefore, the Normalized Cell Index for all wells must be equal one (1) at the normalization time point. Cells exhibited varying response patterns represented by CI changes that either indicated the cells as growing (increasing CI), dying (decreasing CI) or experiencing proliferation arrest (unchanging CI). Three independent experiments were conducted.

Table 1. Concentrations of the cell death inducers used in the RT-CES experiment

<table>
<thead>
<tr>
<th>Cytotoxic concentration ratios</th>
<th>Plectranthus ciliatus</th>
<th>Actinomycin D</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5 x CC$_{50}$</td>
<td>4.5μg/mL</td>
<td>0.01μg/mL</td>
</tr>
<tr>
<td>CC$_{50}$(a)</td>
<td>9μg/mL</td>
<td>0.02μg/mL</td>
</tr>
<tr>
<td>2 x CC$_{50}$</td>
<td>18μg/mL</td>
<td>0.04μg/mL</td>
</tr>
</tbody>
</table>

(a) Determined by XTT

Cytotoxic concentrations of the treatment affecting 50% of the cell population tested (CC$_{50}$) based on RT-CES data were calculated using Roche xCelligence Software 1.2.1. The CC$_{50}$ values were expressed as the mean of three independent assays and the variability between repeats as standard error of the mean (SEM). RT-CES graphs were used to determine whether the responses to treatment were cytotoxic or cytostatic by using guidelines published by Kustermann et al. (2013) [16]. The line at CI = 1 divided the graphs into viable /
proliferating cells (CI > 1) and nonproliferating / dying cells (CI < 1) from each other (Fig. 1). Three typical graphs were described in Kustermann et al. (2013), where the response of cells in the absence or presence of treatments were described. Where the CI was more than 1, the cells were viable / proliferating indicative of cells under no stress. Where the CI < 1 the cells experienced toxicity. Lastly, when the growth curve moved from CI > 1 to a CI < 1 during the observation time the cells could be defined as experiencing cytostatic responses [16].

For the FTIR investigation, methods published by Zelig et al. (2009) and Machana et al. (2012) were used as a guideline during the optimization of the experiments [14,29]. Exponentially growing HeLa cells were seeded (1 x 10^5 cells) and incubated in 12-well plates. After 24 hours incubation, the cells were exposed to two different concentrations (CC50 and 2 x CC50) of the plant extract and actinomycin D respectively (Table 1). Vehicle (0.09% DMSO) and negative control (untreated) cells were included in the analyses. Cells were incubated for 72 hours, after which all the cells were collected from each well. Cells were centrifuged for 10 minutes, the supernatant discarded and the pellets washed twice with 1 mL phosphate buffered saline (PBS) to remove all traces of the culture media. The cells were incubated at room temperature in 1 mL 10% formalin (in PBS) for 10 minutes to fixate the cells. The cells were centrifuged for five minutes and the supernatant discarded. The cells were washed twice with 1 mL of sterile distilled water and centrifuged for three minutes to remove all traces of formalin and PBS, which could influence the spectra. The supernatant was removed and the cells suspended in 10 µL sterile distilled water. The cell suspensions (3 µL) were dried in the biosafety flow hood for one hour on sterile CaF2 discs (Crystran, UK).

The FTIR spectra were acquired using a Bruker V70x spectrometer equipped with a Hyperion 15x IR objective (Bruker, Germany). Using a liquid nitrogen cooled detector attached to the microscope, ten areas were analysed containing a lawn of cells (approximately 400 cells / area). Spectra in the range of 850-4000 cm^{-1} were recorded. Data were collected in transmission mode and spectra (4 cm^{-1} spectral resolution, 2 cm^{-1} intervals, co-added for 100 scans) recorded using Bruker OPUS software. Three independent experiments were conducted. All the spectra were pre-processed using OPUS 7 software. Atmospheric contributions (water vapour and CO2) were subtracted and the spectra vector normalized.
Baseline correction was done using the rubber band function. Second derivatives were calculated (17 smoothing points using the built in Savitzky-Golay algorithm to improve the signal-to-noise ratio) and used for interpretation and statistical analysis [29]. Second derivatives were analysed to assign accurately and visually maximize differences [14,29].

Average spectra of the treatment groups were constructed and One-way Analysis of Variance (ANOVA) calculated. A p < 0.05 was considered significant. Bonferroni multiple comparison tests in conjunction with Tukey post-hoc testing were done in SPSS 20 to determine which variances were significant. An unsupervised method, principal component analysis (PCA) was used to identify trends in the spectra. The data sets were reduced by five principal components and mean centred. The PCA plot was constructed in OPUS 7 while Principal Component (PC) variances were calculated in SPSS 20.

### 3. Results and discussion

In the current study, the effects of a crude extract (*Plectranthus ciliatus* leaves) against HeLa cells proliferation were investigated using FTIR microspectroscopy to distinguish between cytostatic and cytotoxic cellular responses. RT-CES, a well-established technique used for making these distinctions (among others) served as a reference point for the FTIR spectroscopic data.

#### 3.1 Real Time Cell Electronic Sensing of treated HeLa cells

Distinctions between cytotoxic and cytostatic responses were determined using Kustermann and colleagues (2013) manuscript as a guideline [16] who found the area under the curve to be indicative of the type of cellular response. The analysis model created by Kustermann *et al.* [16] was constructed following on assessment of the behaviour of T3T fibroblast cells when exposed to three different compounds; Indocine (anti-inflammatory drug, nontoxic), Vandetanib (tyrosine kinase inhibitor, cytotoxic) and Dasatinib (kinase inhibitor, cytostatic affecting cell cycle). The analysis model was then tested by that same research group with 40 known compounds inducing well documented cellular responses *in vivo* and was confirmed to have good correlation with known records.
Graphs were analysed and indicated in Fig. 2. The line at CI = 1 divided the graphs into viable / proliferating cells (CI > 1) and nonproliferating / dying cells (CI < 1) from each other (Fig. 1). Three typical graphs were described in Kustermann et al. (2013), where the response of cells in the absence or presence of treatments were described. Where the CI was more than 1, the cells were viable / proliferating indicative of cells under no stress. Where the CI < 1 the cells experienced toxicity. Lastly, when the growth curve moved from CI > 1 to a CI < 1 during the observation time the cells could be defined as experiencing cytostatic responses [16].

![Graphs of cellular responses](image)

**Fig. 1.** Representative graphs of cellular responses monitored with RT-CES, where the data is normalized from compound addition until the experiment is stopped approximately 72 hours after the addition of the treatment. Data is normalized to CI = 1 and a line at 1 is indicated in each of the graphs. Untreated or vehicle treated cells where cells grow normally / as expected (CI > 1), the graph appears as shown in (a) where nontoxic responses has similar graph trends as compared to untreated and treated cells, (b) is a representation of a cytotoxic compound where the CI < 1 and finally (c) represents a cytostatic response where the cells firstly has a CI > 1 and as time progress has a CI < 1 in the observation time.
Preliminary screening of the plant extract using XTT gave a CC$_{50}$ value of 9 µg/mL while actinomycin D (positive control) produced a CC$_{50}$ value of 0.02 µg/mL (Table 1). Three concentrations based on these CC$_{50}$ values were investigated using the RT-CES system. The RT-CES system established the crude extract and actinomycin D as having CC$_{50}$ values of 15 ± 1.8 µg/mL and 0.015 ± 0.003 µg/mL respectively (Fig. 2). The difference in CC$_{50}$ determined by the two approaches is a testament to the improved sensitivity of RT-CES compared to conventional viability assays.

RT-CES graphs were compared to the graphs in Fig. 1 that distinguished the type of cellular responses observed (Fig. 2 and Table 2). Actinomycin D was dissolved in sterile distilled water while the plant extract was dissolved in DMSO to ensure that the mixture of components dissolved. Untreated cells were thus used as the control cells for actinomycin D treated cells while DMSO treated cells were used for plant extract treated cells. Vehicle (DMSO) treated HeLa cells had a very similar graph and proliferation profile as untreated cells and therefore nontoxic. When HeLa cells were treated with the crude extract at a concentration of 18 µg/mL (2 x CC$_{50}$ determined by XTT), there was a sudden and severe cell response within the first 10 hours after the addition of the treatment. The cell response observed included the size of the cells decreasing as well as cell detachment that led to a CI below 0.5 that remained constant (Fig. 2a). Cells treated with 18 µg/mL crude extract had a typical graph associated with cytotoxicity. Treatment of the cells with lower concentrations of the crude extract of *Plectranthus ciliatus* (4.5 µg/mL and 9 µg/mL) caused a minor decrease in CI values with similar growth trends as untreated cells (where the CI values remain similar to the CI values of treated cells, then the treatment was nontoxic). The cells treated with 9 µg/mL (the CC$_{50}$) showed a decrease of 20% in viability when comparing the CI values of DMSO treated cells. The areas were also not meaningfully different from the untreated- and DMSO cells and thus the cellular response small. The plant extract elicited nontoxic and cytotoxic cellular responses which depended upon the concentrations of the plant extract. Cytotoxicity of the extract can be traced back to the phytochemicals (including diterpenes) documented for this plant in the literature. Diterpenes isolated from the Plectranthus genus had been identified as having anti-neoplastic activity [30]. One of the speculated compounds which might have caused inhibition of cell proliferation as well as detachment of cells is rosmarinic acid which is found only in the subfamily of Nepetoideae of which *Plectranthus ciliatus* is a member [31,32].
Actinomycin D treated HeLa cells exhibited an entirely different cellular response when compared to the crude extract treated cells (Fig. 2b). When the cells were treated at higher concentrations of actinomycin D, the CI decreased more dramatically compared to the lower concentrations. The RT-CES graphs indicated that lower concentrations of actinomycin D induced minor changes and were, therefore, nontoxic. After the cells were treated with 0.04 µg/mL actinomycin D (2 x CC₅₀), the drug significantly changed the cellular response. In this case actinomycin D elicited a cytostatic cellular response. Sawicki & Godman (1971) observed that the induction of apoptosis in HeLa cells were evident at concentrations between 1 – 10 µg/mL actinomycin D, while cell deterioration was delayed when the cells were treated with lower concentrations (0.1 - 0.01 µg/mL). During the S- and G2-phase, actinomycin D is incorporated within DNA causing the cells to have cytostatic characteristics [33,34] as seen here as well.
Fig. 2. Representative graphs of the effect on proliferation of (a) *Plectranthus ciliatus* extract and (b) actinomycin D (cell death inducer) treated HeLa cells using RT-CES. Cells were incubated for approximately 18 hours after which the treatments at three different concentrations of the treatments were performed. In (a), the vehicle (DMSO) and media-only cells show minimal CI changes. The plant extract treated cells were severely affected by 18 µg/mL which led to a sudden decrease in the CI values. In (b) the affects were also more severe as compared to actinomycin D treated cells (0.04 µg/mL).
Table 2. Summary of the treatments and cellular responses after treatment.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Untreated</td>
<td>-</td>
</tr>
<tr>
<td>DMSO</td>
<td>Nontoxic</td>
</tr>
<tr>
<td><em>Plectranthus ciliatus</em> 4.5 µg/mL</td>
<td>Nontoxic</td>
</tr>
<tr>
<td><em>Plectranthus ciliatus</em> 9 µg/mL</td>
<td>Nontoxic</td>
</tr>
<tr>
<td><em>Plectranthus ciliatus</em> 18 µg/mL</td>
<td>Cytotoxic</td>
</tr>
<tr>
<td>Actinomycin D 0.01 µg/mL</td>
<td>Nontoxic</td>
</tr>
<tr>
<td>Actinomycin D 0.02 µg/mL</td>
<td>Nontoxic</td>
</tr>
<tr>
<td>Actinomycin D 0.04µg/mL</td>
<td>Cytostatic</td>
</tr>
</tbody>
</table>

3.2 FTIR spectral analysis of HeLa cells treated with actinomycin D and *Plectranthus ciliatus*

RT-CES demonstrated a cytostatic response for actinomycin D (0.04 µg/mL), while the effects of *P. ciliatus* (18 µg/mL) was cytotoxic. The sensitivity of FTIR microspectroscopy was evaluated to assess whether similar observations would be evident. Spectra were recorded for untreated and treated cells in the region of 4000-850 cm\(^{-1}\) (Fig. 3) and exhibited bands originating from lipids, proteins and carbohydrates (Table 3). Increasing signal to noise levels originated when the cells were treated with the crude extract of *P. ciliatus*. The increase in noise was concentration dependent as can be seen in Fig. 3. The noise was likely due to the crude extract containing numerous compounds causing noise in the region between 1800 and 2000 cm\(^{-1}\). The presence of the noise in the region between 1800 and 2000 cm\(^{-1}\) did not lead to any spectral shift since shifts were not apparent in the averaged spectra nor the second derivatives (Fig. S1). In Table 3, the mid infrared band assignments common to biological samples were summarized. Major vibrational bands of untreated HeLa cells in the fingerprint region were similar to proliferating HeLa cells investigated previously [10,35]. No major spectral differences were observed between vehicle control (0.09% DMSO) and untreated cells.
Fig. 3. Average spectra of FTIR spectral features in the regions between 850-4000 cm\(^{-1}\) of untreated and treated HeLa cells. The interaction between HeLa cells and the treatments *Plectranthus ciliatus* and actinomycin D at two different concentrations. Major vibrations assigned to biological molecules are indicated.

Table 3. Major mid infrared spectral ranges in cells

<table>
<thead>
<tr>
<th>Region</th>
<th>Spectral range</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>3000 – 2800 cm(^{-1})</td>
<td>Stretching vibrations of CH(_2) and CH(_3) contained mainly in the lipid acyl chains and to a lesser extent proteins [12,36]</td>
</tr>
<tr>
<td>2</td>
<td>1800 – 1300 cm(^{-1})</td>
<td>Absorptions were due to proteins, the shoulder band around 1738-1745 cm(^{-1}) was assigned to C=O stretching mode of phospholipids. Protein amide C=O stretching took place around 1655 cm(^{-1}) (amide I) and the amide N-H bond (amide II) appears at 1546 cm(^{-1}). Bands around 1480-1300 cm(^{-1}) were caused by amino acids side chains and fatty acids [2,12,13]</td>
</tr>
<tr>
<td>3</td>
<td>1300-850 cm(^{-1})</td>
<td>The absorptions were from carbohydrates and phosphates associated with nucleic acids. The vibrational bands at 1238 cm(^{-1}) and 1088 cm(^{-1}) were associated with asymmetric and symmetric phosphodiester vibrations of nucleic acids. Various glycogen bands were superimposed between 1153 cm(^{-1}) and 1022 cm(^{-1}) while a small peak appearing at 966 cm(^{-1}) was assigned to C-O phosphodiester moieties [12,37]</td>
</tr>
</tbody>
</table>
In this investigation the vector normalized second derivatives were analysed to accurately assign and visually maximize differences [14,29] in the spectral regions (850 - 1800 cm\(^{-1}\) and 2800 – 3000 cm\(^{-1}\)) of untreated and treated HeLa cells (Fig. 4).

![Graphs showing spectral regions](image.png)

Fig. 4. The average second derivative spectra of untreated and treated HeLa cells in the regions of (a) 3000 - 2800 cm\(^{-1}\) (b) 2000 - 1500 cm\(^{-1}\) and (c) 1500 - 850 cm\(^{-1}\). Actinomycin D treated cells had very small variances in most spectral regions. When the cells were treated with the plant extract more severe vibrational band changes occurred as compared to lower concentrations of the plant extract.
3.2.1 Significantly altered vibrational bands associated with cellular injury

All the wavenumbers of the spectra between 850 - 1800 cm\(^{-1}\) and 2800 - 3000 cm\(^{-1}\) were evaluated because this is where the treatments under investigation appeared to have had the strongest impact.

3.2.1.1 Cytostatic and nontoxic responses

Nontoxic responses were identified when the cells were treated with DMSO (negative control) and actinomycin D (0.02 µg/mL) using RT-CES. Actinomycin D of higher concentrations (0.04 µg/mL) was defined as inducing a cytostatic response, but none of the vibrational bands was found to be significantly altered. FTIR microspectroscopy was unable to distinguish viable (untreated HeLa cells) from cells treated with DMSO and actinomycin D at 0.02 µg/mL nor could vibrational spectroscopy distinguish healthy from cytostatic cells treated with actinomycin D (0.04 µg/mL). Cytostatic drugs encourage cells not to proliferate and cause cell cycle arrest. During cytostasis, the cells remain viable with very low energy consumption and thus remain in the cell cycle stage in which the cytostatic drug inhibits further growth. Cells remain morphologically intact with limited expansion and movement. FTIR spectroscopy was able to confirm the data of RT-CES and literature that cytostatic cells on a molecular level were still viable and not committed to death since the cells can still recover. Spectra were similar to untreated / viable cells and / or vehicle treated cells.

3.2.1.2 Early stress related responses and cytotoxicity

*P. ciliatus* (9 µg/mL) was defined as being non-toxic using RT-CES while significant vibrational band alterations were identified after statistical analysis compared with viable, non-toxic and cytostatic responses. It was postulated that RT-CES could not detect early stress related responses since those responses were based on vibrations of the molecules. Two vibrational bands were identified as significantly altered after the treatment. Those bands were assigned to amide I vibrations (1634 cm\(^{-1}\) and 1679 cm\(^{-1}\)). It was found that antiparallel β-sheets and β-sheets of amide I could be used to distinguish *P. ciliatus* (9 µg/mL and 18 µg/mL) from nontoxic and cytostatic responses (Table 4). Therefore, 1634 cm\(^{-1}\) and 1679 cm\(^{-1}\) could be used as indicators of early stress related responses that remain evident during cytotoxic responses. The dynamic biochemical reaction towards cytotoxins could, therefore, be seen in the conformational changes in amide I with the increase in β-sheets and a noteworthy increase in α-helices (1655 cm\(^{-1}\)). The change in conformation of amide I could
also be due to different proteins being expressed. Zelig et al. (2009) found that this scenario to be associated with apoptotic cells [29].

Table 4: Significantly (One-way ANOVA post hoc tests: Tukey test, Bonferroni) different bands in the second derivative spectra.

<table>
<thead>
<tr>
<th>Wavenumber (cm⁻¹)</th>
<th>One-way ANOVA p-value</th>
<th>Multiple comparison outcomes</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>915</td>
<td>0.03</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Ribose – phosphate main chain vibrations associated with nucleic acids [38,39]</td>
</tr>
<tr>
<td>931-933</td>
<td>0.007</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>C-O-C ring of deoxyribose in Z-form DNA [37,40]</td>
</tr>
<tr>
<td>989</td>
<td>0.002</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>PO₃²⁻ vibrations of nucleotides [41,42]</td>
</tr>
<tr>
<td>1192</td>
<td>0.021</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Deoxyribose - phosphate backbone with high contribution from the C3’ endo sugar moiety [37]</td>
</tr>
<tr>
<td>1369</td>
<td>0.009</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Stretching C-N of cytosine and guanine [37]</td>
</tr>
<tr>
<td>1437</td>
<td>0</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>CH₂ bending of proteins (amide II) [43]</td>
</tr>
<tr>
<td>1450</td>
<td>0.012</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Hydrogen bonded Aspartic acids and Glutamic acid (CH₂COOH) [2] Methylenes of biomolecules [44]</td>
</tr>
<tr>
<td>1546</td>
<td>0.008</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Metastable phase of Amide II [45]</td>
</tr>
<tr>
<td>1634</td>
<td>0.001</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Antiparallel β-sheets of Amide I [36,46]</td>
</tr>
<tr>
<td>1679</td>
<td>0.006</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>β-sheets of Amide I [2,36]</td>
</tr>
<tr>
<td>1772</td>
<td>0.001</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>Glycerol bound fatty acids, triacylglycerides [47]</td>
</tr>
<tr>
<td>2874</td>
<td>0.037</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>C-H stretching of lipids and proteins of CH₃ [48]</td>
</tr>
<tr>
<td>2962</td>
<td>0.016</td>
<td>Nontoxic vs. Cytotoxic</td>
<td>C-H stretching of lipids and proteins [49]</td>
</tr>
</tbody>
</table>
Other vibrational bands associated with cytotoxicity was also detected in the cells treated with 18 µg/mL (Fig. 4, Table 4). Those bands were associated to nucleic acids (915 cm\(^{-1}\), 933 cm\(^{-1}\), 989 cm\(^{-1}\), 1192 cm\(^{-1}\), 1369 cm\(^{-1}\)), proteins (1437 cm\(^{-1}\) and 1546 cm\(^{-1}\)), lipids and proteins (1450 cm\(^{-1}\), 1772 cm\(^{-1}\), 2874 cm\(^{-1}\), 2962 cm\(^{-1}\)). The 933 cm\(^{-1}\) band was assigned to the vibrational mode of the C-O-C ring of deoxyribose in Z-form DNA [37,38,44]. The region of 1069-1044 cm\(^{-1}\) was assigned to C-O vibrations of furanose and is documented to be enhanced in Z-DNA [37]. After the treatment of the cells with \textit{P. ciliatus}, the peak broke up into two new peaks ~1069 and ~1043 cm\(^{-1}\). Other bands associated with Z-DNA was also apparent in the same spectra around 1438 cm\(^{-1}\) (shoulder peak not present in the other treated cells) assigned to adenine in Z-DNA, the main Z-DNA marker at 1216 cm\(^{-1}\) as well as the band at 1264 cm\(^{-1}\) assigned to guanine-cytosine helices in Z-form DNA [37]. The 915 cm\(^{-1}\) band became less intense as the treatment increased and the band significantly decreased after 18 µg/mL plant extract treatment. The band was assigned to ribose – phosphate main chain vibrations associated with nucleic acids [38,39,49–52]. The transformation of B-DNA to Z-DNA can occur under physiological conditions but also due to increased salt concentrations or dehydration of the cells [53]. In this investigation, it is unlikely that the severity of Z-DNA formation seen in the cytotoxicity response was only due to dehydration since the control cells showed less severe changes. Lampronti \textit{et al.} (2006) found that when different plant extracts were incubated with oligonucleotides isolated from cancer cells, an inhibition of transcription factors were evident [54]. Nucleosome formation and placement, as well as the organisation of chromosomes, are affected by the formation of Z-DNA [55], which could have caused the increased absorption of the FTIR peak. Increased salt concentration could be due to additional salts found in the stomata to help with the opening and closing during respiration of the plant. During the day (when the leaves were harvested) the stomata is open which requires high concentration of potassium. Eukaryotic muscle cells have a concentration of 30 mM potassium and a slight increase to 50 mM could induce detectable transformation of B-DNA to Z-DNA [56,57]. Conformational changes in the amide II band could indicate at a glance that biochemical processes involved in chemical stress and cell death induction, such as protein expression related to chemical stress had taken place.

An increase in triacylglycerides and glycerol bound free fatty acids (1772 cm\(^{-1}\)) increased during the cytotoxic response. An increase in the absorbance was likely due to increased lipid content within the cells such as the formation of lipid droplets [58]. Lipid droplets formed in T-lymphoblastoid (Jurkat) cells when apoptosis was induced by anthracyclins and other
cancer drug controls [59]. The reason for the occurrence of the lipid droplets are still not clear, although the degradation of lipids, due to inhibition of the mitochondrial fatty acid β-oxidation pathway during apoptosis could be a possible reason [58,60]. The increase of high wavenumber peaks at 2874 cm\(^{-1}\) and 2962 cm\(^{-1}\) could be indicative of plasma membrane modifications caused by the cytotoxic treatment [29,61].

### 3.2.2 Principal component analysis

The analysis is an unsupervised data reduction method used to identify trends within data based on similarities and differences within the data [62]. Untreated and treated cells at different concentrations of actinomycin D and *P. ciliatus* showed spectral variance which could be described by the first two principal components (PC). The total percentage variance per PC was indicated in brackets (Fig. 5). A total of 59% of the spectral variance were explained by the data reported. It was seen that PC 1 separated the plant extract treated cells from the untreated and actinomycin D treated cells. FTIR spectra caused clustering of *P. ciliatus* treated cells although the cellular injury was concentration dependent using RT-CES. A possible reason for the clustering could be due to significant alterations of amide I (as found with ANOVA). Clustering of actinomycin D treated cells with untreated cells and DMSO (vehicle) cells confirms that only small spectral changes occurred and it can be assumed that minimal cellular injury was caused by the addition of actinomycin D (0.02 µg/mL). These cells treated with actinomycin D or DMSO were classified as nontoxic. Cells treated with 0.04 µg/mL had a cytostatic RT-CES curve which could not be distinguished from the non-toxic responses using PCA. The cells were reversibly inhibited to proliferate, but still being able to maintain cellular functions for survival. It was postulated that RT-CES could measure the after-effects of cytotoxicity while FTIR could measure the molecular changes using vibrational differences before any morphological effects may become evident.
RT-CES classified cellular injury based on changed morphology of the cells and the ability to attach to the electrodes. Detachment is characteristic of cell death induced by cytotoxins. It was concluded that 18 µg/mL *P. ciliatus* treated HeLa cells induced a cytotoxic response while the other treatments had less severe effects on the cells. RT-CES is an established technique for distinguishing non-toxic, cytostatic and cytotoxic responses which cannot be distinguished by conventional viability assays. FTIR microspectroscopy is widely used to distinguish between different cell types; cell cycle stages as well as cytotoxic treatments. FTIR microspectroscopy confirmed the reversible nature of cytostatic cellular status by detecting that the cells were still viable. Additionally, FTIR detected changes in the molecular vibrations initiated by 9 µg/mL *P. ciliatus* treatment of cells which was associated with the early cellular stress responses.

In conclusion, RT-CES and FTIR microspectroscopy complemented each other by RT-CES detecting morphological changes as indicators of cell injury and could distinguish between viable, cytostatic and cytotoxic responses. FTIR microspectroscopy confirmed that cytostatic cells were not committed to death and could still recover while also describing early cellular stress related responses on a molecular level. Using FTIR microspectroscopy it can be determined whether a treatment were inducing cytostatic responses, early stress related responses or cytotoxic responses.
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5. Conflict of interest statement

The authors declare no conflict of interest.

6. References


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