FTIR spectroscopy of whole cells for the monitoring of yeast apoptosis mediated by p53 over-expression and its suppression by *Nigella sativa* extracts

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Abstract

p53 over expression in yeast results in cell death with typical markers of apoptosis such as DNA fragmentation and phosphatidylserine externalization. We aimed to substitute-supplement classical fluorescent techniques (TUNEL, Annexin V, ROS detection) usually used to detect biochemical changes occurring during yeast apoptosis mediated by p53 over-expression and the effect of anti-apoptotic purified molecules from Nigel (*Nigella sativa*) extracts on these same yeasts by the label free technique of FTIR spectroscopy. The comparison of the entire IR spectra highlighted clear modifications between apoptotic p53-expressing yeasts and normal ones. More precisely, DNA damage was detected by the decrease of band intensities at 1079 and 1048 cm⁻¹. While phosphatidylserine exposure was followed by the increase of νₛCH₂ and νₐsCH₂ bands of unsaturated fatty acids that were exhibited at 2855 and 2926 cm⁻¹, and the appearance of the C = O ester functional group band at 1740 cm⁻¹. In a second step, this FTIR approach was used to estimate the effect of a purified fraction of the Nigel extract. The modulation of band intensities specific to DNA and membrane status was in agreement with apoptosis suppression in presence of the Nigel extracts. FTIR spectroscopy is thus proven to be a very reliable technique to monitor the apoptotic cell death in yeast and to be used as a means of evaluating the biomolecules effect on yeast survival.

Introduction

Fourier-transform infrared (FTIR) spectroscopy is a vibrational spectroscopic technique used to solve the chemical composition information of a sample and to provide its “molecular fingerprint” [1]. This technique has been previously explored for different measurement modes in many biological systems [2,3] to become a powerful tool for the cell components analysis, such as nucleic acids [4], proteins [5] and membranes [6]. This technique has also been useful for complex biological systems, such as tissues [7] and microorganisms [3,8,9] because of its
FTIR spectroscopy for the monitoring of yeast apoptosis mediated by p53 over-expression

Yeast minimal medium supplemented with 2% galactose; MMGLU, Yeast minimal medium supplemented with 2% glucose; PBS, Phosphate buffered saline; PI, propidium iodide; RID, refractive index detector; ROS, reactive oxygen species; SD, standard deviation; TUNEL, terminal-dUTP nick end labelling.

ability to analyze both qualitative and quantitative, molecular and structural indications for samples components. FTIR spectroscopy is a non-destructive technique [10,11], that allows a rapid analysis of biochemical and structural changes (without the need for reagents) such as DNA, protein and cell membrane [10,12–14]. The analysis of spectral data provides qualitative and quantitative information of a cell component on the basis of peak’s shifts, bandwidths and band intensities. Under stress, data can simultaneously determine changes in proteins, lipids, carbohydrates and nucleic acids at the level of functional groups [10, 15]. FTIR spectroscopy was used by lot of researchers to study metabolic alterations in stressed microorganisms such as the bacterial response to antibiotics [16], the effect of starvation [17,18], and of different environmental stresses [19–24], as well as yeast alterations by chemical compounds such as ethanol [25,26]. Bellisola and Sorio [27] consider that FTIR spectroscopy in medical biology is an interesting emerging opportunity.

Apoptotic cells exhibit morphological changes including membrane blebbing, cell shrinkage and biochemical changes including chromatin condensation and DNA fragmentation [28], changes in protein synthesis [29] and translocation of phosphatidyl-serine (PS) from the cytoplasmic side of the plasma membrane to its outer side, and finally exposing PS to the medium. FTIR spectroscopy has already been used to study apoptosis [30–35], necrosis [36], cell cycle, differentiation and proliferation in different cell lines and tissues [37, 38], but never in yeast, and this is where the novelty of this research lies.

Yeast apoptosis was detected for the first time in a strain carrying mutation in the AAA-ATPase gene CDC48 [39]. Other works identified stimulants including acetic acid [40], sugars [41], hydrogen peroxide (H2O2) [42], aspirin [43], and essential oils [44]. In addition, the heterologous expression of pro-apoptotic proteins such as Bax [45] and caspase 1 and 3 in yeast leads to apoptosis [46]. Yeast apoptosis has been described to occur in different physiological scenarios [47]. Particularly, S. cerevisiae apoptosis shares several morphological and biochemical features with the mammalian one, despite some peculiar differences. Nuclear DNA fragmentation and phosphatidyl serine externalization are common markers of both mammalian and yeast cell death. However, the characteristic feature of mammalian apoptosis is the activation of caspases “Aspartate-specific”, and proteases that initiate and achieve the cell death cascade through of the cell components degradation while yeast contains only one caspase homolog gene, called YCA1, encoding for a metacaspase which is Arginine/Lysine-specific, [48]. Several other differences do exist between both apoptosis mechanisms, due to the absence of specific genes such as bcl2, p53, bax, MDM2 in yeasts.

In a previous study our group showed that Human p53 over expression in the yeast Saccharomyces cerevisiae resulted in cell death with characteristic markers of apoptosis: an exposure of PS and DNA strand cleavage as shown by Annexin V staining and TUNEL (terminal-dUTP nick end labeling) assays respectively, including higher ROS production [49]. This system model was exploited to purify molecules from Nigel (Nigella sativa) extracts that can suppress the negative effect of the p53 expression in yeast (unpublished results).

The aim of this study was to get further insight into the capability of the non-destructive technique FTIR spectroscopy to confirm the p53 apoptosis marker in yeast found previously by fluorescence technique and to investigate the role of purified anti-apoptotic fraction from Nigel (Nigella Sativa) extracts in the inhibition of such apoptosis.

Materials and methods

Strains and media

The used yeast strain is W303-1B (MATa ade2-1; ura3-1; his3-11,15; leu2-3,11,12; trp1-1; can1-100) [50] called W303. The yeast multicopy YepDP8-1 vector (provided by Denis Pompon, C.
G.M., CNRS-Gif Sur Yvette, France), called here pDP, was used to express human p53 (wt form) under the control of the Gal10/Cyc1 galactose-inducible yeast promoter. This vector contains the 2 μ replication origin and the ura3 marker [51]. The insertion of the wt p53 cDNA into this yeast expression vector has been previously described [52].

Yeast minimal medium (MM): contained 0.67% yeast nitrogen base (Difco), tryptophan (40 mg/L), histidine (20 mg/L), leucine (120 mg/L) and adenine (40 mg/L). MM was supplemented with either 2% glucose (for normal growth) or 2% galactose (for p53 gene induction) as carbon sources and called respectively MMGLU (repression of p53 expression by glucose) and MMGAL (induction of p53 expression by galactose). The growth kinetics were performed on liquid and solid media.

**Cell growth kinetics**

Yeast culture of W303/pDP and W303/p53 was grown on glucose-containing MM (MMGLU) until stationary phase (O.D = 4 units). 400 μL from MMGLU cultures of W303/pDP and W303/p53 were inoculated in MMGAL media to induce P53 expression (W303/p53) and in MMGAL added with the active fraction from Nigel extract. Yeasts were cultured overnight under continuous agitation (150 rpm) at 30˚C.

**Apoptosis assessment by Annexin V-FITC/PI staining**

Exposure of phosphatidylserine and internal PI staining were detected using the Annexin V Staining kit (life science), essentially as described by Madeo et al. (1997) [53]. Yeast cells were harvested by 4000 rpm, washed three times with 1.2 M Sorbitol, 100 mM KH₂PO₄, 50 mM citric acid. Thereafter, the spheroplasts were obtained by incubation in lysis buffer (1.2 M Sorbitol, 100 mM KH₂PO₄, 50 mM citric acid, 0.2% β-mercaptoethanol, and 0.2 mg/mL (zymolyase) for 1 h at 37˚C. Spheroplasts were washed with 1.2 M Sorbitol, 100 mM KH₂PO₄, 50 mM citric acid, and suspended in the binding buffer (1.2 sorbitol, 10 mM HEPES pH 7.4, 140 mM NaCl, 2.5 mM CaCl₂). Spheroplastes were stained with 10 μL of Annexin V 15 min in darkness at room temperature, washed with binding buffer, then stained with 10 μL of PI., samples were immediately analyzed in the cytometer.

**Flow cytometry analysis**

A cyflow®Space flow cytometer (Sysmex-Partec) equipped with a 488 argon laser was used to measure single cell fluorescence. Annexin fluorescence was measured using the green 536±25 nm band pass filter (FL1). Fluorescence of PI was measured using the orange 590±25 nm band pass filter (FL2).

Fluorescent beads of 10 μm in diameter were added in order to normalize the flow cytometer settings. The voltage for forward scatter (FSC) and side scatter SSC were fixed at 120 and 200, respectively. The flow rate was adjusted to 120 μL/ min. Data were collected for 10.000 events using logarithmic amplification. Unstained cells were used as controls and each assay was run in duplicate. Data analysis was investigated using winlist software (verity software, USA).

**FTIR spectroscopy**

Cultures were stopped after 32 h and harvested by centrifugation (7000 rpm, 10 min). The Pellets were washed twice with saline solution (0.7% NaCl) and re-suspended in 1 mL of the saline solution. Five microlitres of the concentrated yeast were deposited on a selenium multi-plate (96 wells) in 3 replicates and dried in a desiccator under 1.5 bar. FTIR measurements were
performed in transmission mode using HTS-XT Tensor 27 spectrophotometer (Bruker, Marne la vallée, France). Each experiment was replicated from 3 independent cultures, in order to optimize and control reproducibility [54].

The spectra were registered using 64 scans, at a resolution of 4 cm\(^{-1}\) baseline corrected and normalized using OPUS 6.5 (Bruker) software as previously described in [55]. An average spectrum was then calculated for each independent culture. If some spectra did not pass the “quality test” and spectrum reproducibility, 3 more independent cultures were performed to obtain 3 spectra per strain. Spectra analysis would be performed using OPUS 6.5 software (Bruker) by hierarchical cluster analysis (HCA) peaks measurements and integration. To verify the similitude between spectrums recorded from the same culture, calculation of the similitude percentage was performed in the whole spectra using search standard program: similitudes have to be more than 99% [54]

Plant extracts preparation

Nigel seeds were crushed in a blender (Rutschmuhle) to obtain powdered material. A volume of 30 mL of distilled water was added to 10 g of the powder [56]. Following continuous homogenization into a polytron apparatus (Kinematica-Gmbh), the mixture was centrifuged at 10,000 rpm for 15 min. The supernatant (of brownish orange color) was filter-sterilized on 0.45 \(\mu\)m filter. Clarified solutions were stored in sterilized tubes at 4˚C until chromatographic separation.

Preparative High Performance Liquid Chromatography (HPLC)

A preparative HPLC Knaeur system-model 1100 consisting of preparative pump wellchem K-1800 and a UV detector Eurochrom K-2501, using a PL Aquagel OH-40-10\(\mu\)m (300’’25mm) with protein separation range from 10 to 200 kDa was used. The plant extract was filtered on 0.45 \(\mu\)M and injected. The fractions were eluted at a flow rate of 5 mL/min and UV monitored at 280 nm, using an isocratic phase consisting of water. Eluted fractions were dried then dissolved in water.

Preparative Fast-Performance Liquid Chromatography (FPLC)

A preparative FPLC Biorad system made up of 2128 model using anionic unoQ12. Active G1 fraction was filtered through 0.45 \(\mu\)m, deposited on an anionic column at pH 5 (citrate buffer), with a flow rate of 3 mL/min and a UV monitoring at 280nm.

The elution gradient consists of: a- isocratic flow with 100% A-buffer (citric buffer/tris buffer) for 3 min, b- a sample injection, c- isocratic flow with 100% A-buffer for 20 min, linear gradient with 100% B-buffer (composed of 1 M NaCl in A-buffer), d- isocratic flow with 100% B-buffer for 5 min, e- isocratic flow with 100% A-buffer for 5 min.

Analytical High-Performance Liquid Chromatography (HPLC)

An HPLC Agilent system consisting of model Agilent 1100 and equipped with a “C-18 reversed phase” column (Eurosphere 100–5 C-18” 250×8mm TC 01107) was used. Active Q1 fraction was firstly filtered through 0.45\(\mu\)m, deposited on the column and fractions were eluted at a flow rate of 0.5 mL/min and a UV monitored at 260 nm and by RID (refractive index detector).

The elution gradient, used for the reversed phase C-18, was a linear one and consisted of solvent A (1% formic acid) and solvent B (100% methanol). The solvent gradient started at 95% A and 5% B, reaching 75% A at 10 min, 65% A at 30 min, 55% A at 35 min, 55% A at 40
min, 50% A at 45 min, 45% A at 50 min, 30% A at 53 min, 25% A at 56 min, and 20% A at 60 min, followed by a post-time isocratic plateau for 10 min at 95% A before the next injection. The flow rate was 0.5 mL/min.

Northern blotting

Northern blot were performed as described by Sambrook et al. (1989) [57] using Hybond N membranes (Amersham). Formamdehyde gel was performed using 20 μg of RNA for each sample. Hybridization was performed with an [α-32P] radio-labelled 650 bp fragment of cDNA p53.

Results

p53 mediated yeast apoptosis detection by fluorescent methods

Previous studies proved that the human p53 over-expression, under the control of a tightly regulated galactose-inducible promoter, induces yeast cell death with characteristic markers of apoptosis: exposure of phosphatidyl-serine and DNA strand cleavage as shown by Annexin V staining and TUNEL (terminal dUTP nick end labeling) assays respectively [49].

Fig 1 shows the inhibition of yeast growth of W303/wt p53 and control W303/pDP cells under inducible condition on MMGAL (Fig 1a). This result was confirmed by the growth kinetic in the MMGAL liquid (Fig 1b).

Apoptosis was assessed by flow cytometry analysis of phosphatidylserine externalized at the outer phase of the cytoplasmic membrane, a phenomenon that was considered an early event of cell death and was detected by labeling cells with fluorescein coupled annexin V (green dye) PI was used to distinguish apoptosis from necrosis. Cell population stained Annexin +/PI- is considered apoptotic. The percentage of apoptotic cells increases from 34.03% in W303/pDP stain (control) to 69.62% in W303/p53 (Fig 1c).

FTIR analysis of p53 expression in S. cerevisiae

We planned here to harness the FTIR spectroscopy in the study of apoptosis in such lower eukaryote and evaluate the effect of an anti-apoptotic bio-molecule on this phenomenon. The originality of our study stems from the fact that FTIR spectroscopy is used here for the first time to study molecular changes in yeast apoptosis mediated by the human P53.

Fig 2b depicts the normalized FTIR spectra of the transformed yeast cells W303/pDP (control) and W303/p53 that are growing under inducible conditions on MMGAL (Fig 1b).

FTIR spectra of normal cells (W303/pDP) and apoptotic cells (W303/p53) were compared using Hierarchical cluster analysis (HCA) (Fig 2a). Spectral apoptosis induced by P53-overexpression markers were sought through direct observations of its IR spectrum (Fig 2b); Peak assignments are based on previous FTIR studies in cell apoptosis as described in Table 1.

The results are expressed as the mean ± standard deviation (SD). The statistical significance of the differences was evaluated by Student’s t-test; P<0.05 which was considered to indicate a statistically significant differences.

The 3000–2820cm−1 region

Absorptions in this region are principally assigned to the symmetric and asymmetric stretching modes of methylene and methyl groups of lipids [58]. The intensities of bands 2855 cm−1 and 2925 cm−1 correspond respectively to CH2 symmetric stretching (νsCH2) and CH2 asymmetric stretching (νasCH2). Compared to the control strain (W303/pDP), the band intensities at 2855 and 2925 cm−1 increased from 0.270±0.009 and 0.443±0.017 to 0.364±0.020 and
0.544±0.026 in apoptotic W303/p53 cells (Fig 2c). This result was confirmed by the peak integration of 2993 to 2855 cm\(^{-1}\) area (data not shown) that increased from 14.843±0.983 in controls to 16.077±0.845 in apoptosis.

1758-1477cm\(^{-1}\) region

This region is dominated by the amide I and amide II bands, characterized by absorption at 1652 and 1547 cm\(^{-1}\), respectively. This originates from the vibrations of the amide groups (COONH) of proteins, and is often an indicator of their secondary structure.

In our study, there are no changes in amide I and amide II intensity bands. In addition amide I/amide II intensity ratio had no conformational changes of membrane proteins. However a weak band at 1740 cm\(^{-1}\) appears in apoptotic W303/p53, that arises from C = O stretching vibrations of the ester functional groups in lipids [34].

1187-945cm\(^{-1}\) region

Band changes in this region are associated with mainly macromolecules like carbohydrates and polysaccharides that are related to the amount of nucleic acids, the 1048 cm\(^{-1}\) band is attributed to the νC-O stretching [59] of nucleic acids: desoxyribose [30, 36], RNA and ribose [35]. The absorption band at 1079 cm\(^{-1}\) corresponds to the symmetric PO\(_2\) (vsPO\(_2\)−) stretching
vibrations of the phosphate groups of the DNA double stranded backbone [60,37,61], the νC–O stretching vibrations of nucleic acid polysaccharides, and the νC–O–P stretching vibration of phosphorylated lipids [35,36]. Therefore, the 1079 cm⁻¹ absorption band is reported to reflect the cell DNA content [62] and this indicates the DNA oxidative damage degree [30].

**Table 1. Spectral peak assignments.**

| Wave numbers (cm⁻¹) | Spectral peak assignment                                                   |
|---------------------|---------------------------------------------------------------------------|
| 1048                | νC–O of polysaccharides nucleic acids                                      |
| 1079                | νC–O–C of nucleic acids and phospholipids that indicates the degree of oxidative damage in DNA |
| 1406                | Carboxylates of amino acids and δSCH₂ of lipids.                         |
| 1464                | δSCH₃                                                                      |
| 1547                | Amide II: proteins, δN-H (bending) and νC–N (stretching)                 |
| 1652                | Amide I: proteins, C = O stretching                                       |
| 1740                | νC = O ester of lipids                                                   |
| 2855                | ν₇CH₂ (lipids)                                                            |
| 2926                | ν₈CH₂ (lipids)                                                            |

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In this region, the most prominent differences feature between W303/pDP and the W303/p53 apoptotic cells are the bands decreased intensities at 1048 and 1079 cm\(^{-1}\) (Fig 2d) which indicates a loss of nucleic acids content, reflecting a DNA damaged state. This finding corroborates the previous results of DNA fragmentation analyzed by flow-cytometry in p53-mediated yeast apoptosis \[49\].

The anti apoptotic effect of Nigel fraction

The current yeast genetic system was used for the screening of natural molecules having a protective effect against p53-mediated apoptosis in yeast. Nigel (\textit{Nigella sativa}) seeds were chosen as starting materials, as they are highly considered for their richness in antioxidant compounds.

Fig 3a shows that, in the presence of 4 mg dry weight of the Nigel extract that has been spread on the galactose plates, the growth of p53\(^+\) yeast cells was restored (Fig 3a). Conversely in the absence of the plant extracts, as expected, a great difference was observed between the control and the p53 expressing cells: while the growth of W303/pDP cells was normal, that of W303/p53 was strongly affected (Fig 3a).

In order to check the results obtained on solid media, the growth of cells was tracked in liquid MMGAL media with or without addition of Nigel extract. As expected, in the absence of extracts, the growth of W303/p53 was inhibited (Fig 3b). When plant extracts were added to

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Fig 3. Effect of purified fraction from Nigel extract on the yeast growth of W303/p53. (a) Yeast clone W303/p53 growth on solid and (b) on liquid media MMGAL in absence and presence of purified Nigel extract. And (c) northern blot analysis of p53 mRNA level in control cell W303/p53 in inducible condition in absence (1) and in presence of Nigel extract (2). (d) Quantification of the blot.

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the medium culture, the growth of p53+ cells was almost completely restored (Fig 3b). Nigel extract can act on the p53 expression itself or on the resulting effect of p53 expression, so that a northern blot was performed to study transcriptional level of p53 in presence of Nigel extract. Fig 3c shows clearly the presence of mRNA p53 in W303/p53 grown in MMGAL in presence of Nigel extract, which mean that Nigel extract did not affect p53 expression.

Guided fractionation was applied using many chromatographic columns as described in the materials and methods part, and there was just one active fraction responsible for the anti apoptotic effect.

FTIR study of the anti apoptotic effect of Nigel fraction extract

The beneficial effect of Nigel fraction was shown either on MMGAL solid or liquid media. In this part, we investigated the contribution of the FTIR technique to study the effect of Nigel fraction on apoptosis features in W303/p53 recombinant yeast.

In order to verify that Nigel fraction has no effect on yeast growth, FTIR spectra of control cells W303/pDP grown in MMGAL with and without the active Nigel fraction were analyzed and showed no significant difference between both spectra (Fig 4a). Moreover, the spectra cluster analysis shows that there are no spectral conformational changes (Fig 4c). It is known that FTIR spectra can be divided into spectral regions where strong absorption is related to specific component. These spectral regions include 3050-2800cm-1 where fatty acids predominates, 1750-1500cm-1 that ascribed to amide bands of proteins and peptides, 1500-1250cm-1 for carboxylic acids, and 1200-900cm-1 the polysaccharide region define the cell metabolic
fingerprint. Since all these regions of the two spectra superimpose perfectly, we conclude that there would be no effect of the Nigel extract on cellular metabolism yeast cells (Fig 4).

3000–2820 cm\(^{-1}\) region

Concerning W303/p53, Fig 5 shows that in the presence of the nigel fraction, the intensities of the marker band of the membrane disorder (2855 and 2926 cm\(^{-1}\)) decreased significantly from 0.364±0.020 and 0.544±0.026 in the absence of nigel extract) to 0.273±0.008 and 0.448±0.018 (in presence of Nigel extract), respectively. Even the vsCH\(_2\) to vasCH\(_2\) ratio decreased from 0.669±0.004 to 0.610±0.007. These results show that the Nigel fraction reestablished the normal synthesis of unsaturated fatty acid.

1187–945 cm\(^{-1}\) region

DNA fragmentation in W303/p53 cells was highlighted by the intensity decrease of the band 1079 and 1048 cm\(^{-1}\) being at 0.615±0.019 and 0.595±0.020 (Fig 5), respectively. In the same p53+ cells incubated with the Nigel fraction, these intensities reached similar values for the
control W303/pDP, being at 0.715±0.023 and 0.716±0.023 respectively. Thus the Nigel fraction prevents the DNA fragmentation.

Moreover, νC = O stretching vibrations of the ester functional groups, observed at 1740 cm⁻¹ in W303/p53 cultured in MMGAL, disappeared in the presence of the Nigel fraction, which means that this fraction inhibited the PS exposure.

**Discussion**

The most typical markers of apoptosis are DNA fragmentation and phosphatidylserine externalization. Both markers are well conserved in yeast and are harnessed here to show that p53 overexpression induces apoptosis ([49]; this work). The present FTIR study of yeast apoptosis highlighted several intensity changes of bands that are characteristic of the same markers. These findings prove that the FTIR technique is a good tool to study the biochemical changes in yeast during apoptosis.

**Phospholipids changes**

Conformational changes in 3000–2820cm⁻¹ are reported in ethanol-stressed yeast cells [24] and can possibly be associated to biochemical changes resulting from the disorganization of the cell plasma membrane itself due to the increase of unsaturated fatty acids leading to an increase of the yeast membrane fluidity [63].

Increased intensities in this spectral area can be considered as an important apoptosis marker that was previously shown in many studies on mammalian cells indicating a conformational disorder in cell membrane [32,33,35,36,61]. It could also reflect the variation of lipid amounts around the nucleus, also known to increase in apoptosis [34].

The CH₂-symmetric ratio to CH₂-antisymetric stretching is another way to label the cell membrane morphology alterations [64]; in our study it increased significantly in W303/p53 cells that underwent apoptosis.

It has been reported that apoptotic cells change in morphology [65], consisting in plasma membrane pores, or leaky spots created by rigid patches of tightly packed phospholipids in the inner plasma membrane. These rigid patches are formed due to an increased amount of saturated fatty-acids [64] and increased membrane fluidity [65]. This spectral change could be correlated with the major exposure of PS in the outer leaflet of the plasma membrane [66].

**Protein changes**

This originates from the vibrations of the amide groups (COONH) of proteins, and is often an indicator of their secondary structure. The vibrational band at~1652 cm⁻¹ results from the stretching vibrations of C-O that is a characteristic of the α-helix structure [67]. However the 1547 cm⁻¹ absorption band, which is a combination of the (N–H) bending and n(C–N) stretching vibrations, is attributed to the β-sheet secondary structure of proteins [68].

The amide bands were harnessed to study all the protein structures including the membrane proteins. Indeed, several studies described the association of an increase in β-sheet structures with apoptosis [34,36,38,69]. Caspari et al 2003 [37] suggested that conformational changes of protein structures might be correlated with proteolysis exerted by apoptotic caspases [70,71]. Differences in yeast cell composition after exposure to nanosilver revealed a significant downshift (20 cm⁻¹) of the amide I peak at 1655 cm⁻¹ and a loss in α-helix structures, thereby indicating the disordered secondary structures of proteins [72].

This conformational change allows the identification of apoptosis spectral marker that could be assigned to protein oxidation [73]. The PS is a negatively-charged phospholipid that is normally predominant in membrane leaflets facing the cytosol; the loss of the membrane
phospholipid asymmetry (during apoptosis) results in the inversion of the PS exposure. Thus, the conformational disorder of ester stretching in PS, corresponding to the increase of C = O ester groups of phosphatidylserine, could be the signal of phospholipids alteration that increases the membrane permeability and fluidity [74]. In this study the phospholipids spectral changes substantiate the exposure of the PS at the outer surface of the plasma membrane induced by the p53 expression in yeast [49],

Nucleic acids changes
DNA damages were expressed by the decrease of 1048 cm\(^{-1}\) and 1079 cm\(^{-1}\) absorption bands that reflect the cell DNA content [52] and indicate the DNA oxidative damage degree [25].

Anti-apoptotic effect of *Nigella sativa* seeds
Given that the yeast has similarities with mammalian cells at the molecular and functional levels, it would contribute to understand some molecular mechanisms of higher eukaryotes such as apoptosis. In this study, p53+ yeast cells were used as a model for the screening of anti-apoptotic molecules. *Nigella sativa* extract was shown to prevent yeast apoptosis by following PS externalization and DNA fragmentation. In addition, there was no significant conformational changes between W303/pDP MMGAL and W303/pDP MMGAL+Nigel fraction. This means that *Nigella sativa* did not change the yeast metabolism nor its growth.

Conclusion
This study confirms that FTIR spectroscopy technique is a very sensitive and useful tool for the detection and monitoring of biochemical changes of apoptosis mediated by p53 over-expression in yeast *S. cerevisiae*, previously analyzed by flow-cytometry. Moreover, highlighted apoptosis marker bands were used to evaluate the protective effect of *Nigella sativa* extract on p53-mediated apoptosis. This rapid and specific method could undoubtedly be used for a large screening of other natural biomolecules on yeast p53-mediated apoptosis.

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