2019

Vibrational Characterization of Granulosa Cells From Patients Affected by Unilateral Ovarian Endometriosis: New Insights From Infrared and Raman Microspectroscopy

Valentina Notarstefano
*Universita Politecnica delle Marche, Italy*

Giorgia Gioacchini
*Universita Politecnica delle Marche, Italy*

Hugh J. Byrne
*Dublin Institute of Technology, hugh.byrne@tudublin.ie*

Carlotta Zaca
*9. Baby Centre for Reproductive Health, Bologna, Italy*

Eleni Sereni
*9. Baby Center for Reproductive Health, Bologna, Italy*

*See next page for additional authors*

Follow this and additional works at: [https://arrow.dit.ie/nanolart](https://arrow.dit.ie/nanolart)

Part of the [Chemicals and Drugs Commons](https://arrow.dit.ie/nanolart) and the [Medical Sciences Commons](https://arrow.dit.ie/nanolart)

**Recommended Citation**

This Article is brought to you for free and open access by the NanoLab at ARROW@TU Dublin. It has been accepted for inclusion in Articles by an authorized administrator of ARROW@TU Dublin. For more information, please contact yvonne.desmond@dit.ie, arrow.admin@dit.ie, brian.widdis@dit.ie.

This work is licensed under a [Creative Commons Attribution](https://creativecommons.org/licenses/by/4.0/).
Vibrational characterization of Granulosa Cells from patients affected by Unilateral Ovarian Endometriosis: new insights from infrared and Raman microspectroscopy

Valentina Notarstefano\textsuperscript{a†}, Giorgia Gioacchini\textsuperscript{a†}, Hugh J. Byrne\textsuperscript{b}, Carlotta Zacà\textsuperscript{c}, Elena Sereni\textsuperscript{c}, Lisa Vaccari\textsuperscript{d}, Andrea Borini\textsuperscript{c}, Oliana Carnevali\textsuperscript{a}, Elisabetta Giorgini\textsuperscript{a*}

\textsuperscript{a}Dipartimento di Scienze della Vita e dell' Ambient, Università Politecnica delle Marche, via Breccie Bianche, 60131 Ancona, Italy
\textsuperscript{b}FOCAS Research Institute, Dublin Institute of Technology, Kevin Street, Dublin 2, Ireland
\textsuperscript{c}9. Baby Center for Reproductive Health, via Dante 15, 40125 Bologna, Italy
\textsuperscript{d}SISSI Beamline, Elettra-Sincrotrone Trieste, S.C.p.A., S.S. 14 – Km 163.5, 34149 Basovizza, Trieste, Italy

† These authors equally contributed to this work

*Corresponding author.
E-mail address: e.giorgini@univpm.it (E. Giorgini)

ABSTRACT

Endometriosis is a chronic gynaecological disease characterised by the presence of endometrial cells in extra-uterine regions. One of the main factors impacting on the fertility of women affected by endometriosis is the poor oocyte quality. Granulosa Cells (GCs) regulate oocyte development and maintain the appropriate microenvironment for the acquisition of its competence; hence, the dysregulation of these functions in GCs can lead to severe cellular damages also in oocytes. In this study, luteinized GCs samples were separately collected from both ovaries of women affected by Unilateral Ovarian Endometriosis and analyzed by infrared and Raman microspectroscopy. The spectral data were compared with those of GCs from women with diagnosis of tubal, idiopathic or male infertility (taken as control group). The coupling of these two spectroscopic techniques shed new light on the alteration induced by this pathology on GCs metabolism and biochemical composition. In fact, the study revealed in GCs from both ovaries of women affected by unilateral ovarian endometriosis similar biochemical modifications, such as the alteration of the protein pattern, the induction of oxidative stress mechanisms, and the deregulation of lipid and carbohydrate metabolisms. These evidences suggest that unilateral endometriosis impairs the overall ovarian functions, causing alterations not only in the ovary with endometriotic lesions but also in the contralateral “healthy” one.

Key words: Unilateral Ovarian Endometriosis, FTIR microspectroscopy, Raman microspectroscopy, Multivariate analysis
Introduction

Endometriosis is a chronic gynaecological disease characterised by the presence of epithelial, glandular and stromal endometrial cells in extra-uterine districts [1]. It is reported that 30% to 50% of women with a diagnosis of endometriosis are affected by infertility [2]. Several factors have been suggested as possible causes of endometriosis-associated infertility: pelvic adhesions, luteinized unruptured follicles, immunologic alterations, progesterone resistance, and impairment of folliculogenesis, ovulation, ovum transport, fertilization, and implantation [3–8]. The characteristics of this infertility are variable, and women affected by endometriosis can display a decrease of fertilization, implantation and pregnancy rates [9]. To avoid infertility problems, in women affected by unilateral ovarian endometriosis (UOE), oocytes are routinely collected in Assisted Reproductive Technology (ART) practice from the contralateral ovary, which does not show endometriotic lesions and hence can be considered healthy [10].

Several studies reported that the presence of endometriotic lesions can directly impair also the activity of Granulosa cells (GCs), somatic cells that surround the oocyte [11–13]. GCs are responsible for many important follicular functions, such as the production of oestradiol during follicular growth, the production of essential nutrients used as an energy source during oocyte maturation, the accumulation of oocyte secreted metabolites, and the secretion of progesterone after ovulation [14–17]. A dysregulation in these functions can lead to severe cellular damages, causing decreased rates of oocyte nuclear maturation and fertilization [18–20].

Fourier Transform InfraRed Microspectroscopy (FTIRM) and Raman MicroSpectroscopy (RMS) are powerful vibrational techniques, widely applied in life sciences for the study of the biomolecular building and composition of cells [21–23]. They present the advantage to be label-free, since each molecule has a proper IR and Raman spectrum. The analysis of IR and Raman bands in terms of position, intensity and width, makes possible to obtain a unique molecular fingerprint of the most relevant biological molecules (proteins, lipids, sugars and nucleic acids) inside the analysed samples [24–26]. This chemical information can be directly related to specific biological processes, such as cellular activity, metabolism and oxidative stress [27–29]. IR absorption and Raman spectroscopies can be considered to be mutually complementary due to the difference in physical origin of the processes. Whereas IR absorption is sensitive to polar (usually antisymmetric) group vibrations, Raman is sensitive to polarizable (usually symmetric) group vibrations. Thus, the combination of the techniques gives a more complete analysis of the complete biological specimen. In addition, in the
conventional microscopic forms, FTIR has a spatial resolution of ~10 μm, whereas with a x
100 objective, confocal Raman microscopic resolutions of <1 μm can be achieved. FTIR can
have the advantage of averaging larger areas, while Raman microspectroscopy can enable
subcellular analysis [30,31].

Recently, our team applied FTIRM for the vibrational characterization of both human
oocytes and GCs [32,33]. Furthermore, preliminary FTIRM tests were also carried out on GCs
collected from patients with diagnosis of unilateral ovarian endometriosis (UOE), and
unexpected results were obtained, shedding new light on the effects of UOE on GC
metabolism. In fact, the macromolecular profile of GCs collected from patients affected by
this disease was very similar, irrespective of the ovary they were recovered from [34,35].

Pursuing this approach further, a multidisciplinary study, which applies the analytical
spectroscopic methods of both FTIRM and RMS to the study of this gynecological disease,
was performed, aiming to provide a more fundamental understanding of the biochemical
modifications induced by unilateral ovarian endometriosis in the metabolism of GCs collected
from the contralateral “healthy” ovary with respect to those from the ovary with endometriotic
lesions.

**Experimental section**

The study, approved by the Ethics Committee of 9.Baby Center for Reproductive Health
(Bologna, Italy), was carried out in full accordance with ethical principles for experiments
involving humans, include The Code of Ethics of the World Medical Association (Declaration
of Helsinki, 2013). To participate in this investigation, patients signed an informed consent,
which included the donation of GCs. All samples were strictly anonymous, and it was
impossible to correlate them to patients.

**Luteinized GCs sample collection**

N. 20 patients (N. 10 with diagnosis of unilateral ovarian endometriosis and N. 10 with
diagnosis of tubal, idiopathic or male infertility) were enrolled in an *in vitro* fertilization
program, according to the following inclusion criteria: 35±2.8 of age; non-smokers; regular
ovulatory menstrual cycles; FSH<10 IU/I on day 3 of the menstrual cycle.

Controlled ovarian stimulation was induced using leuprorelin (Enantone, Takeda, Rome,
Italy) and recombinant follicle-stimulating hormone (rFSH) (Gonal-F, Serono, Rome, Italy,
or Puregon Organon, Rome, Italy). Human chorionic gonadotropin (HCG) at 10,000 IU
(Gonasi, Amsa, Rome, Italy) was administered when one or more follicles reached a diameter of ca. 23 mm [36]. At the end of the treatment, from each patient, a pool of luteinized GCs was collected from all the follicles, according to the following protocol. GCs undergo luteinisation after ovulation, at a specific moment of the ovarian cycle; since they are crucial for the development of the oocyte and for pregnancy, luteinized GCs are commonly used to study of ovarian functions [37]. Follicular fluid was centrifuged at 1100x g for 10 min and the pellet was resuspended in 2 mL of Sydney IVF Gamete Buffer Medium (Cook IVF, Brisbane, Australia), overlaid on 80%-40% discontinuous gradient of silica particles suspension (1 mL PureSperm 40 and 1 mL PureSperm 80; Nidacon, Goteborg, Sweden), and centrifuged at 1600x g for 13 min to separate GCs from red blood cells. After centrifugation, three layers were observed: a top layer containing the follicular fluid, a middle ring-like layer containing GCs and a bottom layer containing erythrocytes. GCs were recovered in the middle ring-like layer using a Pasteur pipette and placed into 1 mL of NaCl 0.9% solution (Fresenius Kabi). Isolated GCs were centrifuged at 300x g for 10 minutes. The supernatant was discarded, and the pellet re-suspended in 1 mL of NaCl 0.9% solution and washed again at 300x g for 10 minutes. After additional centrifugation for 5 min at 600x g, the cell pellet was re-suspended in 100 μl of sterile NaCl 0.9% solution.

GCs samples were divided into the following experimental groups: CTRL (N. 10 GCs samples collected from the ovaries of women with diagnosis of tubal, idiopathic or male infertility, taken as control group), ENDO (N. 10 GCs samples collected from the ovary with diagnosis of unilateral ovarian endometriosis), and CONTRAL (N. 10 GCs samples collected from the contralateral “healthy” ovary).

**FTIRM measurements and data analysis**

N. 5 aliquots of each GCs sample were deposited without any fixation process onto CaF2 optical windows (1-mm thick, 13-mm diameter) and air-dried for 30 min, in order to avoid water contributions to the IR spectra [33]. FTIRM measurements were performed, within the same day of collection, at the infrared Beamline SISSI (Synchrotron Infrared Source for Spectroscopic and Imaging), Elettra Sincrotrone Trieste (Trieste, Italy). A Hyperion 3000 Vis-IR microscope equipped with a HgCdTe (MCT_A) detector and coupled with a Vertex 70 interferometer (Bruker Optics, Ettlingen, Germany) was used.

From each aliquot of GCs sample, ~10 areas containing densely packed cell monolayers were selected by visible microscopy, from which IR spectra were collected in transmission mode in the MIR region (4000 - 800 cm⁻¹) [33]. Knife-edge apertures were set at 30×30 μm²
(512 scans, spectral resolution of 4 cm\(^{-1}\), zero-filling factor of 2 in the spectral range 4000–800 cm\(^{-1}\), scanner velocity of 40 kHz). Background spectra were collected using the same parameters on clean zones of the CaF\(_2\) optical windows. All IR spectra of each aliquot of GCs sample were averaged (Averaging routine, OPUS 7.1 software) and then corrected for the contribution of atmospheric carbon dioxide and water vapour (Atmospheric compensation routine, OPUS 7.1 software). Average spectra (50 for each experimental group), obtained by this procedure, were evaluated in terms of S/N ratio on the basis of the height of the band centred at \(\sim 1660\) cm\(^{-1}\) (Amide I band of proteins), which is always the highest peak of cell spectra; average spectra having at 1660 cm\(^{-1}\) an absorbance value lower than 0.07 a.u. (\(\sim 20\%\)) were discarded [28].

The remaining IR spectra were vector normalized, and then submitted to multivariate analyses, with no further pre-processing. The pairwise PCA analysis of CTRL/ENDO, CTRL/CONTRAL and CONTRAL/ENDO spectra was performed, by exploiting an in-house developed algorithm in R Studio (R Studio: Integrated Development for R. RStudio, Inc., Boston, MA). PCA was employed as an unsupervised multivariate approach to analyse spectral data of GCs. The order of the PCs denotes their importance to the dataset; PC1 describes the highest amount of variation [38]. The PC scores were also displayed by loading spectra, which contain peaks, both positive and negative that explain the spectral variation in the dataset; this tool is used as a method to separate spectra into groups.

For a more detailed analysis, for each experimental group the average absorbance spectrum, together with its standard deviation spectra (average absorbance spectra ± standard deviation spectra) were calculated (Averaging routine, OPUS 7.1 software). Average absorbance spectra ± standard deviation spectra were then curve fitted in the following spectral regions: 3050–2800 cm\(^{-1}\), 1790–1480 cm\(^{-1}\) and 1350–900 cm\(^{-1}\). A Gaussian algorithm (GRAMS/AI 7.02, Galactic Industries, Inc., Salem, NH) was adopted on IR spectra upon straight baseline correction and vector normalization. The number of underlying bands and their centre values (expressed as wavenumbers) were precisely identified by second derivative results, and the integrated areas were calculated. The wavenumber, together with the vibrational mode, the label and the biological meaning of all the underlying bands are reported in Table 1. The mean values of the integrated areas of selected underlying bands with biological meaning were determined. In addition, the sum of the mean values of integrated areas of all the underlying bands in the 1790–1600 cm\(^{-1}\) region (corresponding to the Amide I band of proteins, named AI) and that of all the underlying bands in the 1350-900 cm\(^{-1}\) spectral range (named TOT) were also calculated. These values were used to calculate the following band area ratios:
CH/CH\(_3\), CH\(_2\)/CH\(_3\), C=O/Al, FOLDED/Al, UNFOLDED/Al, Ph1/TOT, RNA1/TOT, Ph2/TOT, GLYCO/TOT, DNA1/TOT, RNA2/TOT, DNA2/TOT, and Z-DNA/TOT. For labels and biological and vibrational meaning, see Table 1.

**Table 1**
IR vibrational modes highlighted by Peak Fitting procedure in CTRL, CONTRAL and ENDO absorbance average spectra. For each peak, the wavenumber, together with the vibrational mode, the label and the biological meaning are reported.

<table>
<thead>
<tr>
<th>Range (cm(^{-1}))</th>
<th>Wavenumber (cm(^{-1}))</th>
<th>Vibrational mode</th>
<th>Label</th>
<th>Biological meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>3000-2800</td>
<td>3012</td>
<td>Stretching vibration of =CH groups</td>
<td>CH</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2957, 2869</td>
<td>Asymmetric and symmetric stretching vibrations of CH(_3) groups</td>
<td>CH(_3)</td>
<td>Mainly lipid alkyl chains [32,56]</td>
</tr>
<tr>
<td></td>
<td>2925, 2852</td>
<td>Asymmetric and symmetric stretching vibrations of CH(_2) groups</td>
<td>CH(_2)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1790</td>
<td>Stretching vibration of C=O ester moieties</td>
<td>C=O</td>
<td>Lipid peroxidation [42]</td>
</tr>
<tr>
<td>1694</td>
<td>1694</td>
<td>Vibrational modes of β-turn secondary structures</td>
<td>FOLDED</td>
<td>Proteins secondary structure [57]</td>
</tr>
<tr>
<td></td>
<td>1680, 1627, 1613</td>
<td>Vibrational modes of β-sheet secondary structures</td>
<td>FOLDED</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1659</td>
<td>Vibrational modes of α-helix secondary structures</td>
<td>FOLDED</td>
<td></td>
</tr>
<tr>
<td>1240</td>
<td>1240</td>
<td>Vibrational modes of random coil secondary structures</td>
<td>UNFOLDED</td>
<td></td>
</tr>
<tr>
<td>1138</td>
<td>1118</td>
<td>Asymmetric stretching vibration of PO(_2)-groups</td>
<td>Ph1</td>
<td>Mainly phosphorylated proteins [42,58]</td>
</tr>
<tr>
<td></td>
<td>994</td>
<td>Stretching vibration of C-OH groups of ribose</td>
<td>RNA1</td>
<td>RNA [32,57,59]</td>
</tr>
<tr>
<td></td>
<td>1087</td>
<td>Stretching vibration of uracil ring</td>
<td>RNA2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1053</td>
<td>Symmetric stretching vibration of PO(_2)-groups</td>
<td>Ph2</td>
<td>Mainly phosphate backbone of nucleic acids [28]</td>
</tr>
<tr>
<td></td>
<td>1020</td>
<td>Stretching vibrations of C-O-C and C-OH groups of carbohydrates</td>
<td>GLYCO</td>
<td>Carbohydrates [28]</td>
</tr>
<tr>
<td></td>
<td>970</td>
<td>Stretching vibration of C=O groups in DNA</td>
<td>DNA1</td>
<td>DNA [32,57,59]</td>
</tr>
<tr>
<td></td>
<td>924</td>
<td>Vibrational modes of DNA backbone</td>
<td>DNA2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>924</td>
<td>Left-handed helix DNA vibrations</td>
<td>Z-DNA</td>
<td>Z-DNA [32,60]</td>
</tr>
</tbody>
</table>

**RMS measurements and data analysis**

RMS measurements were carried out at the FOCAS Research Institute, Dublin (Ireland). For Raman measurements, each GC samples were divided into N. 5 aliquots and fixed in a 4% paraformaldehyde (PFA) solution for 10 min, washed twice in physiological solution, and
then stored at 4°C until RMS measurements. This procedure has been demonstrated to retain
the biochemical profile of the cell as close as possible to that of live cells [39], maintaining
hydration, while preserving them from biological damage during shipping [24]. From each
GCs aliquot 10 point spectra were acquired on cells seeded on the glass slide, targeting nuclei
of the cells. No contribution of glass to the spectra was observed.

A Horiba Jobin-Yvon LabRAM HR800 spectrometer, equipped with a 532-nm diode laser
(\(\sim 50\) mW laser power) as source was used. All measurements were acquired by using a x100
objective (Olympus, N.A. 1). The spectrometer was calibrated to the 520.7 cm\(^{-1}\) line of silicon
prior to spectral acquisition. A 600 lines per mm grating was chosen. A 100 µm confocal
pinhole was used for all measurements. The spectra were dispersed onto a 16-bit dynamic
range Peltier cooled CCD detector. The spectral range from 1800 to 400 cm\(^{-1}\), the so-called
fingerprint region, was chosen and spectra were acquired for 3x10 seconds at each spot. For
each GCs aliquot, the average RMS spectrum was calculated (Averaging routine, OPUS 7.1
software). Average spectra were smoothed using 7 smoothing points, baseline-corrected with
the polynomial method (2 iterations) (OPUS 7.1 software), and then submitted to multivariate
analysis. The pairwise PCA analysis of CTRL/ENDO, CTRL/CONTRAL and
CONTRAL/ENDO spectra was performed, by exploiting an in-house developed algorithm
in R Studio.

The wavenumber of the most relevant peaks found in Raman spectra, together with the
vibrational mode, the label and the biological meaning are reported in Table 2. The height of
such peaks was also calculated (Integration mode K, OPUS 7.1 software).
Table 2
Vibrational modes highlighted in the 1800–400 cm⁻¹ spectral region of CTRL, CONTRAL and ENDO Raman average spectra. For each peak, the wavenumber, together with the vibrational mode, the label and the biological meaning are reported.

<table>
<thead>
<tr>
<th>Wavenumber (cm⁻¹)</th>
<th>Vibrational mode</th>
<th>Label</th>
<th>Biological meaning</th>
</tr>
</thead>
<tbody>
<tr>
<td>~1657</td>
<td>Stretching vibration of amidic C=O groups</td>
<td>PROTEINS</td>
<td>Cellular proteins [61]</td>
</tr>
<tr>
<td>~1615</td>
<td>Stretching vibrations of tyrosine and tryptophan C=C groups</td>
<td>TYR-TRP</td>
<td>Tyrosine and Tryptophan amino acid residues [62]</td>
</tr>
<tr>
<td>~1605</td>
<td>Stretching vibrations of phenylalanine and tyrosine C=C groups</td>
<td>PHE-TYR</td>
<td>Phenylalanine and Tyrosine amino acid residues [63]</td>
</tr>
<tr>
<td>~1263</td>
<td>Amidic C-N stretching and N-H bending vibrations, mainly due to helix structures</td>
<td>HELIX</td>
<td>Helical secondary structures of proteins [62]</td>
</tr>
<tr>
<td>~1003</td>
<td>Symmetric stretching breathing vibration of phenylalanine</td>
<td>PHE</td>
<td>Phenylalanine amino acid residues [64]</td>
</tr>
<tr>
<td>~980</td>
<td>C-C stretching vibration of beta-sheets structures</td>
<td>BETA</td>
<td>Beta-sheets secondary structures of proteins [65]</td>
</tr>
<tr>
<td>~855</td>
<td>C-C stretching vibration of proline sidechains</td>
<td>PRO</td>
<td>Proline amino acid residues [63]</td>
</tr>
<tr>
<td>~642</td>
<td>C-C twisting vibration of tyrosine</td>
<td>TYR</td>
<td>Tyrosine amino acid residues [38]</td>
</tr>
</tbody>
</table>

Statistical analysis

Normally distributed data deriving from RMS and FTIRM spectra were presented as mean ± S.D. Significant differences between experimental groups were determined by means of a factorial analysis of variance (one-way ANOVA), followed by Tukey's multiple comparisons test, using the statistical software package Prism6 (Graphpad Software, Inc. USA). One-way ANOVA compares the means of CTRL, CONTRAL and ENDO groups in order to make inferences about the population means. Statistical significance was set at *p*<0.05. Different letters over box charts indicate statistically significant differences among the above defined experimental groups.

Results

Luteinized GCs samples separately collected from both ovaries of women affected by UOE (the ovary with endometriotic lesions, ENDO, and the contralateral “healthy” one, CONTRAL) were analysed by FTIR and Raman microspectroscopy. The spectral data were
compared with those of GCs from women with diagnosis of tubal, idiopathic or male infertility (taken as control group, CTRL).

**FTIRM analysis**

In Fig. 1, the pairwise PCA scatter plots of GCs IR spectra from CTRL/CONTRAL, CTRL/ENDO and CONTRAL/ENDO groups and the relative loadings are reported. A partial differentiation according to PC1 was detectable in CTRL/CONTRAL and CTRL/ENDO scatter plots (respectively, 71.46% and 70.26% of explained system variance) (Figs. 1A,B); conversely, no separation was detected between CONTRAL and ENDO groups (Fig. 1C). The same discriminant spectral features were observed by the analysis of PC1 loadings of CTRL/CONTRAL and CTRL/ENDO. In particular, differences were observed in the regions at 3050-2800 cm⁻¹ (stretching modes of alkyl groups in lipids), 1790-1480 cm⁻¹ (vibrational modes of Amide I and II bands of proteins), and 1350-900 cm⁻¹ (stretching modes of phosphates and carbohydrates) (Figs. 1D,E). Conversely, no relevant discriminating spectral features were observed in the CONTRAL/ENDO loading plot (Fig. 1F).

![Fig. 1 Multivariate analysis of FTIRM data. Pair-wise PCA scatter plots calculated for CTRL/CONTRAL (A), CTRL/ENDO (B) and CONTRAL/ENDO (C) spectra. PC1 loadings of CTRL/CONTRAL (D), CTRL/ENDO (E), and CONTRAL/ENDO (F) experimental groups.](image)

In Fig. 2, IR spectra of CTRL, CONTRAL and ENDO GCs in the 3100-2700 cm⁻¹ (Fig. 2A) and 1800-800 cm⁻¹ (Fig. 2B) regions are presented. As expected from the loadings analysis, tiny changes were observed by comparing the spectral profile of GCs from the
different experimental groups, above all in the lipid region (3050-2800 cm\(^{-1}\)) as well as in the 1300 – 900 cm\(^{-1}\) range ascribable to phosphate groups and carbohydrates. Hence, for a more in-depth analysis, CTRL, CONTRAL and ENDO GCs spectra were curve fitted in the 3050–2800 cm\(^{-1}\), 1790–1480 cm\(^{-1}\) and 1350-900 cm\(^{-1}\) spectral ranges and the integrated areas of the underlying bands were used to calculate specific band area ratios (see Experimental Section).

The statistical analysis of the numerical variations of these band area ratios is showed in Figs. 3 and 4. The following results have been achieved in both CONTRAL and ENDO GCs with respect to CTRL ones: (i) the ratio CH/CH\(_3\) (indicating the unsaturation degree of lipid alkyl chains) significantly increased (Fig. 3A); (ii) the ratio CH\(_2\)/CH\(_3\) (indicating the saturation degree of lipid alkyl chains) significantly decreased (Fig. 3B); (iii) the ratio C=O/AI (indicating the amount of peroxidised lipids) significantly increased (Fig. 3C); (iv) the ratio FOLDED/AI (indicating the amount of properly folded proteins) significantly decreased (Fig. 3D); (v) the value UNFOLDED/AI (indicating the degree of unfolded structures in proteins) significantly increased (Fig. 3E); (vi) the ratio Ph1/TOT (indicating the amount of phosphate groups mainly in proteins) significantly increased (Fig. 4A); (vii) the ratios RNA1/TOT and RNA2/TOT (both indicating the amount of RNA) significantly decreased (Figs. 4B,F); (viii) the ratio Ph2/TOT (indicating the amount of phosphate groups in nucleic acids) did not significantly change (Fig. 4C); (ix) the ratio GLYCO/TOT (indicating the amount of carbohydrates) significantly decreased (Fig. 4D); (x) the ratios DNA1/TOT and DNA2/TOT (both indicating the amount of DNA) did not significantly change (Figs. 4E,G), and (xi) the ratio Z-DNA/TOT (indicating the amount of Z-DNA) significantly increased (Fig. 4H).
Fig. 3 Box charts showing the numerical variation of the following band area ratios calculated for CTRL, CONTRAL and ENDO GCs: (A) CH/CH3; (B) CH2/CH3; (C) C=O/AI; (D) FOLDED/AI; (E) UNFOLDED/AI. Centre line marks the median, edges indicate the 25th and the 75th percentile, whiskers indicate standard deviation, and the black square marks the mean. Different letters over box charts indicate statistically significant differences among groups (one-way ANOVA and Tukey's multiple comparisons test). Statistical significance was set at $p<0.05$.

Fig. 4 Box charts showing the numerical variation of the following band area ratios calculated for CTRL, CONTRAL and ENDO GCs: (A) Ph1/TOT; (B) RNA1/TOT; (C) Ph2/TOT; (D) GLYCO/TOT; (E) DNA1/TOT; (F) RNA2/TOT; (G) DNA2/TOT; (H) Z-DNA/TOT. Centre line marks the median, edges indicate the 25th and the 75th percentile, whiskers indicate standard deviation, and the black square marks the mean. Different letters over box charts indicate statistically significant difference among groups (one-way ANOVA and Tukey's multiple comparisons test). Statistical significance was set at $p<0.05$. 
RMS analysis

In Fig. 5, the pairwise PCA scatter plots of GCs Raman spectra from CTRL/CONTRAL, CTRL/ENDO and CONTRAL/ENDO groups and the relative loadings are reported. A clear segregation was observed along PC1 axis both for CTRL/CONTRAL and CTRL/ENDO spectra (61.39% and 65.12% of explained system variance, respectively) (Figs. 5A,B). Conversely, no separation was detected in CONTRAL/ENDO plot (Fig. 5C), as confirmed also by PC1 loading of CONTRAL/ENDO populations (Fig. 5F). Conversely, the PC1 loadings of CTRL/CONTRAL and CTRL/ENDO analyses showed relevant modifications in the whole examined spectral range (1800-400 cm\textsuperscript{-1}) (Figs. 5D,E).

![Multivariate analysis of RMS data. Pair-wise PCA scatter plots calculated for CTRL/CONTRAL (A), CTRL/ENDO (B) and CONTRAL/ENDO (C) spectra. PC1 loadings of CTRL/CONTRAL (D), CTRL/ENDO (E), and CONTRAL/ENDO (F) experimental groups.](image)

In Fig. 6, RMS spectra of CTRL, CONTRAL and ENDO GCs in the 1800-400 cm\textsuperscript{-1} range are presented. As already identified by the analysis of loadings, changes in the spectral profile of CONTRAL and ENDO GCs with respect to CTRL ones were observed above all in the band centered at ~1660 cm\textsuperscript{-1}, assigned to proteins, and also in the bands associated with protein secondary structures (~1263 and ~980 cm\textsuperscript{-1}, attributable respectively to helical and beta structures) and aromatic amino acids (~1615, ~1605, ~1003 and ~642 cm\textsuperscript{-1}, corresponding to phenylalanine, tyrosine, and tryptophan).
The statistical analysis of the numerical variations of heights of the peaks, reported in Table 2, is shown in Fig. 7. The following results have been achieved in both CONTRAL and ENDO GCs with respect to CTRL ones: (i) the height of PROTEINS (indicating the total amount of cellular proteins) showed a significant decrease (Fig. 7A); (ii) the height of HELIX (indicating the amount of helical secondary structures of proteins) showed a significant decrease (Fig. 7B); (iii) the height of BETA (indicating the amount of $\beta$-sheet secondary structures of proteins) showed a significant decrease (Fig. 7C); (iv) the height of TYR-TRP (indicating the amount of tyrosine and tryptophan amino acids) showed a significant decrease (Fig. 7D); (v) the height of PHE-TYR (indicating the amount of phenylalanine and tyrosine amino acids) significantly decreased (Fig. 7E); (vi) the height of PHE (indicating the amount of phenylalanine amino acid) significantly decreased (Fig. 7F); (vii) the height of PRO
(indicating the amount of proline amino acid) significantly increased (Fig. 7G); (viii) the height of TYR (indicating the amount of tyrosine amino acid) significantly decreased (Fig. 7H). All the other bands highlighted as different by the PC1 loading were investigated but did not show significant alterations among the experimental groups (data not shown).

**Discussion**

FTIRM and RMS techniques are currently applied in the biomedical field, due to the possibility to obtain, at the same time, in a label-free way and on the same sample, a unique molecular fingerprint of the most relevant biological molecules. In this study, a vibrational approach has been performed to profoundly investigate the effects of UOE on the biochemical composition and metabolism of GCs collected from the ovary with endometriotic lesions, and the contralateral “healthy” one.

**Fig. 7** Box charts showing the variation of height values of the following bands calculated for CTRL, CONTRAL and ENDO GCs: (A) PROTEINS; (B) HELIX; (C) BETA; (D) TYR-TRP; (E) PHE-TYR; (F) PHE; (G) PRO, and (H) TYR. Centre line marks the median, edges indicate the 25th and the 75th percentile, whiskers indicate standard deviation, and the black square marks the mean. Different letters above box charts indicate statistically significant difference among groups (one-way ANOVA and Tukey’s multiple comparisons test). Statistical significance was set at $p<0.05$. 
Endometriosis is known to be related to an impairment of folliculogenesis [6]. Given the importance of GCs on follicle development, a dysregulation of their functions can lead to severe damages to the oocyte [14,40,41].

It should be noted that, while the tightly focused laser spot for the RMS measurements sampled only the nucleus, the lower resolution FTIRM measurements sampled typically 3-4 whole cells, including cytoplasm and nucleus. Nevertheless, individually and in combination, the analysis of FTIRM and RMS data revealed, in both the ovaries of women with a diagnosis of UOE, several layers of GCs impairment. In particular, the occurrence of lipid peroxidation in CONTRAL and ENDO GCs with respect to CTRL ones was evidenced by the significant increase of $\equiv$CH groups (CH/CH3) and C=O ester moieties (C=O/AI), together with the decrease of CH$_2$ groups (CH2/CH3) [42–45].

It is known that cellular oxidation targets not only lipids, but also proteins, leading to their misfolding and creating internal cell stressors [46]. In fact, in both CONTRAL and ENDO GCs, a significant impairment in proteins secondary structure was highlighted (decrease of FOLDED/AI and HELIX and BETA, and increase of UNFOLDED/AI), together with the decrease of tyrosine, tryptophan and phenylalanine amino acids (TYR, PHE, TYR-TRP and PHE-TYR). All these findings suggested the attack of reactive oxygen species (ROS) to proteins in general, and, in particular, to the aromatic amino acid side chains [47].

This oxidation picture is consistent with the well-known reduced follicular antioxidant ability in women with diagnosis of endometriosis [48] and with the role played by oxidative stress and ROS in the progression of the disease [49,50]. Hence, the occurrence in GCs from both ovaries, of these alterations let hypothesize that oxidative processes triggered by endometriotic lesions act not only locally, but also in a systemic way.

In addition, for the first time, the impairment in the carbohydrate metabolism of GCs caused by UOE was observed (decreased value of GLYCO/TOT in CONTRAL and ENDO GCs). Considering that glucose is fundamental for mammalian oocytes, and that it cannot be utilized unless previously transformed into pyruvate by GCs, this alteration, caused by UOE, may be considered crucial for oocytes functionality [51].

In this study, the occurrence of epigenetic effects caused by endometriosis on GCs, already reported in literature, was confirmed (increased value of Z-DNA/TOT and decreased value of RNA1/TOT and RNA2/TOT) [52,53]. Z-DNA is an elongated left-handed conformation of DNA, which can be found in segments with specialized sequences, characterized by alternations of purines and pyrimidines, especially alternating deoxycytidine and deoxyguanosine residues [54]. Z-DNA spatial conformation is thought to be able to influence
transcriptional activity by excluding transcription factors [55]. This epigenetic effect was also observed for the first time in GCs from contralateral ovary, confirming the systemic effects of UOE.

**Conclusions**

In this context, the present vibrational study sheds new light on the alterations induced by UOE on the metabolic status and biochemical composition of Granulosa Cells retrieved from the ovary with endometriotic lesions; moreover, it highlights that the same impairment also characterizes Granulosa Cells collected from the contralateral ovary, usually considered “healthy”. In fact, in GCs from both the ovaries of women affected by UOE, a similarly profound alteration in the protein pattern was found, together with the same activation of oxidative stress mechanisms, the dysregulation of carbohydrate metabolism, and the modification in DNA methylation. Hence, these results open a new scenario, suggesting that unilateral ovarian endometriosis acts not only locally, but also in a systemic way, causing changes in the metabolic and macromolecular composition of GCs of both ovaries. Due to the close relationship between Granulosa Cells and their companion oocyte, we suppose that UOE likely causes an impairment in the whole ovarian functionality.

These findings could improve the knowledge on the infertility often observed in women affected by UOE. In fact, in ART practice, oocytes are routinely collected from the contralateral ovary, which does not show endometriotic lesions and hence is be considered healthy. In contrast, our results suggest that the quality of oocytes retrieved from the contralateral “healthy” ovary may be impaired in the same manner as those retrieved from the affected ovary, suggesting the need to significantly revise the existing ART protocols.

**Conflicts of interest**

There are no conflicts to declare.

**Acknowledgements**

We thank Elettra Sincrotrone Trieste for the access to experimental facilities (accepted Proposal n. 20135178).

**References**


[58] P.T.T. Wong, R.K. Wong, T.A. Caputo, T.A. Godwin, B. Rigas, Infrared spectroscopy of


