Apoptosis is triggered by melatonin in an *in vivo* model of ovarian carcinoma

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Abstract

Apoptosis plays an important role in the treatment of cancer, and targeting apoptosis-related molecules in ovarian cancer (OC) is of great therapeutic value. Melatonin (Mel) is an indoleamine displaying several anti-cancer properties and has been reported to modulate apoptosis signaling in multiple tumor subtypes. We investigated OC and the role of Mel therapy on the pro-apoptotic (p53, BAX, caspase-3, and cleaved caspase-3) and anti-apoptotic (Bcl-2 and survivin) proteins in an ethanol (EtOH)-preferring rat model. To induce OC, the left ovary was injected directly with a single dose of 100 μg 7,12-dimethylbenz(a)anthracene dissolved in 10 μl of sesame oil under the bursa. Right ovaries were used as sham-surgery controls. After developing OC, half of the animals received i.p. injections of Mel (200 μg/100 g BW per day) for 60 days. Body weight gain, EtOH consumption, and energy intake were unaffected by the treatments. Interestingly, absolute and relative OC masses showed a significant reduction after Mel therapy, regardless of EtOH consumption. To accomplish OC-related apoptosis, we first observed that p53, BAX, caspase-3, and cleaved caspase-3 were downregulated in OC tissue while Bcl-2 and survivin were overexpressed. Notably, Mel therapy and EtOH intake promoted apoptosis along with the upregulation of p53, BAX, and cleaved caspase-3. Fragmentation of DNA observed by TUNEL-positive nuclei was also enhanced following Mel treatment. In addition, Bcl-2 was downregulated by the EtOH intake and lower survivin levels were observed after Mel therapy. Taken together, these results suggest that Mel induce apoptosis in OC cells of EtOH-preferring animals.

Key Words

- ovarian cancer
- melatonin
- pro-apoptotic protein
- anti-apoptotic protein

Introduction

Ovarian cancer (OC) is one of the most lethal gynecological malignancies and presents a poor prognosis at the moment of diagnosis (<50% with a 5-year survival rate; Cannistra 2004). Early-stage OC shows no apparent symptoms, and a lack of a reliable screening tool is yet an unsolved issue (Fallows et al. 2001). OC is presumed
Melatonin modulates apoptosis in ovarian cancer

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Materials and methods

Animals and experimental design

Eighty female adult UChB rats, a model of EtOH-preferring rats (50 days old, weighing 200 g), were obtained from the Department of Anatomy, Bioscience Institute/Campus of Botucatu, UNESP – Universidade Estadual Paulista. The rats were individually housed in polypropylene cages containing laboratory-grade pine shavings as bedding and maintained under constant room temperature (RT; 23 ± 1 °C) and lighting conditions (12 h light:12 h darkness cycle, with the lights switched on at 0600 h). Filtered tap water and standard rodent chow (3074 SIF, Purina Ltda., Campinas, SP, Brazil) were provided ad libitum. All animals were divided into two groups (n = 40): EtOH group, in which the rats had access to a 10% (v/v) EtOH solution ad libitum (free choice of water or EtOH), and a control group, which was composed of EtOH-naïve rats without access to EtOH. At 65 days of age, they were given a choice between two bottles containing either water or a 10% (v/v) EtOH solution ad libitum over a period of 15 days. Animals displaying EtOH consumption higher than 2.0 g of EtOH/kg per day (from 4 to 5 g of EtOH/kg per day) were selected to this study according to Chuffa et al. (2011a, 2013b).

After OC development (260 days old), the animals designated to receive Mel (M-5250, Sigma–Aldrich) were administered i.p. doses of 200 μg/100 g BW dissolved in 0.04 ml of 95% EtOH and then diluted in 0.3 ml of 0.9% NaCl (vehicle) at a final concentration of 0.3 mg/ml. The daily injections were administered at night (between 1830 and 1900 h) for 60 consecutive days (Chuffa et al. 2011b). This protocol relies on previous works in which long-term Mel therapy was able to reduce tumor size and tumor cell survival.

Finally, the rats were divided into four subgroups (n = 20): OC, composed of DMBA-induced animals that
did not consume EtOH; OC + EtOH, composed of DMBA-induced animals that consumed 10% (v/v) EtOH during ovarian tumor development (OTD); OC + Mel, composed of DMBA-induced animals that received Mel treatment; and OC + EtOH + Mel, composed of DMBA-induced animals that consumed 10% (v/v) EtOH during OTD and received Mel treatment. After all procedures, the females were anesthetized and euthanized by decapitation (during early morning at 0400 h) for further sample collection. The present experimental protocol was accepted by the Ethical Committee of the Institute of Bioscience/UNESP (CEEA – permit number: 382).

Ovarian tumor induction

After selection, all the animals (n=80) were anesthetized using 10% ketamine (60 mg/kg, i.p.) and 2% xylazine (5 mg/kg, i.p.) during the estrous phase, a 2 cm incision through the skin and abdominal muscles was performed, and the left ovaries were accessed after grasping the fat pad surrounding the organ. The left ovary was injected under the bursa with a dose of 100 μg of DMBA (Sigma–Aldrich) dissolved in 10 μl of sesame oil (vehicle) (Hoyer et al. 2009), and was returned intact to the body cavity. Muscle and skin layers were closed using a 3-0 silk suture (Ethicon, Inc., Juarez, Mexico). Control surgery was conducted on the right ovary by administering only the vehicle. Prophylactic antibiotic (10³ units of benzylpenicillin potassium) was administered i.p. for 2 weeks. Over the next 180 days, tumor development was observed by ultrasonography (size and volume).

Mel levels

Blood was collected each 15 days from the caudal vein and Mel was extracted from the plasma (n = 20 samples/group) using HPLC-grade methanol and separated on columns (Sep-Pak Vac C-18, reverse phase, 12.5 nm; Water Corporation, Milford, MA, USA). Thereafter, 50 μl of reconstituted samples were assayed using a Coat-a-count Mel ELISA Kit and immediately read at 405 nm. Intra-assay coefficient of variation was 4%, and all the samples were assayed at the same time in duplicate. The microtiter plates and reagents were provided by IBL International (Hamburg, Germany), and the concentrations of Mel are presented as pg/ml.

Immunohistochemistry

Sections of papillary OC tissue (n = 10/group) were deparaffinized in xylene and were microwaved (700–800 W) while immersed in 0.01 M sodium citrate buffer, pH 6.0, for antigen retrieval. After blocking endogenous peroxidase activity, the tissues were incubated with 3% BSA for 1 h to avoid non-specific binding. Then, OC sections were incubated in a humidified chamber overnight at 4°C with primary antibodies (Abcam, Cambridge, UK): rabbit monoclonal anti-p53 (1:100), rabbit polyclonal anti-BAX, rabbit monoclonal anti-caspase-3 (1:50), rabbit polyclonal anti-Bcl-2 (1:100), and rabbit polyclonal anti-survivin (1:100). After immunoreactions, the slides were washed in TBS-T buffer and incubated with secondary antibody (polymer anti-mouse IgG or anti-rabbit IgG; Dako Cytomation, Carpinteria, CA, USA) at RT for 1 h. Then, the slides were reacted with diaminobenzidine (Sigma–Aldrich) for 5 min. Finally, OC sections were counterstained with hematoxylin. Negative controls were generated by omitting the primary antibody. Immunohistochemistry (IHC) results were analyzed under a Zeiss Axiopt II microscope (Carl Zeiss, Oberkochen, Germany), considering the levels of staining intensity which were scored as absent (0), weak (+), moderate (+ +), or strong (+ + +) immunoreactivity. All of the OC-positive cells for p53, BAX, caspase-3, TUNEL, Bcl-2, and survivin were counted on ten random fields (×400) considering total tumor cells (data expressed as %).

Determination of DNA fragmentation by TUNEL assay

OC tissues were washed with PBS (sodium chloride, potassium chloride, dihydrogen phosphate, and disodium hydrogen phosphate), fixed in 4% paraformaldehyde for 10 min, and permeabilized with PBS at RT. Cryostat sections (15 μm) were microwaved in 10 mM citrate buffer, pH 6, for 5 min, and in the same buffer with 0.1% Triton X-100 for 5 min at RT. The DNA strand breaks were labeled with -dUTP using TdT (0.5 U/μl) according to the manufacturer’s protocol. The incorporation of nucleotides into 3’-OH end was detected with anti-digoxigenin–fluorescein antibody (Sigma–Aldrich; dilution 1:10). Control tissues were incubated without TdT. Nuclei were stained with 6-diamidino-2-phenylindole (DAPI, 5 min) at RT. Immunopositive cells were analyzed using a fluorescence microscope (Zeiss Axiopt II, Oberkochen, Germany) at 40× magnification (excitation 590 nm and emission filter 650 nm). For DAPI staining was used excitation 365 nm; emission filter 435 nm. The quantification of fluorescence in merged figures was performed using the ImageJ Software (NIH, Bethesda, MA, USA).
Cell proliferation

OC tissues were washed with PBS and fixed in paraformaldehyde for 10 min. Cryostat sections were microwaved in 10 mM citrate buffer, pH 6, for 5 min, and in the same buffer with 0.1% Triton X-100 for 5 min at RT.

Nonspecific binding sites were blocked with 1% BSA. Samples were incubated with anti-Ki-67 primary rabbit polyclonal antibody (dilution 1:100, overnight at 4 °C) followed by secondary polyclonal anti-rabbit IgG conjugated to FITC (1:200, sc-2012, Santa Cruz Biotechnology, Inc.) for 1 h at RT. Nuclei were stained with DAPI for 5 min. Primary and secondary antibodies were diluted in blocking buffer (1% BSA). For negative immunolabeling, no primary antibody was added. Immunopositive cells were analyzed using a fluorescence microscope (Zeiss Axioptih II) at 40× magnification. The quantification of fluorescence in merged images was performed using the ImageJ Software (NIH), and proliferation index was obtained in ten random fields considering Ki-67-positive cells by the total tumor cells (%).

Immunoblots

After treatments, ovarian tumors were rapidly removed, and 100 mg tissue samples were frozen in liquid nitrogen and stored at −80 °C. The tissues were homogenized using RIPA lysis buffer (Pierce Biotechnology, Rockford, IL, USA) supplemented with a cocktail of protease inhibitors. Also, Triton X-100 diluted 1:10 (v/v) were added to the homogenates and samples were placed on dry ice with agitation for 2 h to improve extraction. Lysates were centrifuged at 21 912 g for 20 min at 4 °C to remove insoluble material, and total protein was measured through colorimetric determination. All proteins were dissolved in 1.5× Laemmli buffer and used for SDS–PAGE (Bio-Rad Laboratories). The protein (70 μg) was loaded per well and resolved on preformed 4–12% acrylamide gradient gels (Amersham Biosciences) using a Tris–glycine running buffer electrophoresis system (60 mA, fixed for 2 h). Then, the proteins were electro-transferred (200 mA, fixed for 1.5 h) to nitrocellulose membranes in Tris-glycine–methanol buffer. Pre-stained standards (Bio-Rad) were used as molecular weight markers. After blocking the membranes with TBS-T solution containing 3% BSA at RT for 60 min, they were incubated at 4 °C overnight with the primary antibodies (1:1000 in 1% BSA): p53, BAX, caspase-3, cleaved caspase-3, Bcl-2, and survivin (Abcam). Subsequently, membranes were washed in TBS-T solution and incubated for 2 h at RT with rabbit HRP-conjugated secondary antibodies (Sigma–Aldrich; diluted 1:1000 in 1% BSA). After washing in TBS-T, signals were developed using an ECL detection kit (Thermo Fisher Scientific, Waltham, MA, USA). Immunoreactive bands were calculated from individual blots of ten rats per group using image analysis software (ImageJ Software). β-actin or α-tubulin were used as endogenous positive controls, and results were expressed as mean ± S.D. Immunoblotting concentrations (%) were represented as optical densitometry values (band intensity—pixels).

Statistical analysis

Data are presented as the mean ± S.D., and the analyses were performed using two-way ANOVA for two independent factors (Mel therapy and EtOH intake). Significant results were subjected to post-hoc analysis using Tukey’s test, and statistical significance was set at P < 0.05. SigmaPlot Version 12.5 Graphing Software was used (Systat Software Inc., San Jose, CA, USA).

Results

Incidence of OC and survival rates

Although the cumulative OC incidence was increased from day 140 to 200 after the combination of DMBA with EtOH intake (Fig. 1A), no apparent difference in the OC phenotype was observed following EtOH consumption. To confirm the effect of Mel on this rat model of OC, microscopic analyses were frequently performed, and serous papillary carcinoma was the most common tumor subtype (data not shown). To evaluate the prognostic value of long-term Mel treatment associated or not to EtOH consumption, we first investigated non-treated animals bearing OC, and the cumulative survival following different treatments over the 10 months post-tumor induction. OC group was compared with the upper OC+Mel revealing a significantly shorter disease-specific survival rate (Fig. 1B). Interestingly, after developing multiple metastasis, 100% of the OC+EtOH females die at 8-month post-tumor induction, and Mel therapy promoted longer survival to these females by about 50% from 8 to 10 months (Fig. 1B).

Overall nutritional aspects and Mel concentrations

During the experiments, all of the animals were studied with the same criteria. Briefly, the final body weight, body weight gain, food and EtOH consumption, and energy intake did not vary among the treatments (Table 1). The
absolute and relative OC masses presented significant reductions after Mel treatment in animals consuming or not consuming EtOH. Furthermore, treatment with Mel was able to reduce OC volume in OC Mel group (Table 1).

To validate the treatment, plasma Mel levels were measured, and efficiently, OC Mel and OC EtOH Mel groups exhibited the highest levels of circulating Mel (Table 1).

Mel and EtOH similarly upregulated the expression of pro-apoptotic proteins in OC

We have investigated the molecular mechanism by which p53 evokes OC cell death, and if BAX is responsible for caspase-3 activation. In response to Mel treatment and EtOH consumption alone, p53 immunostaining and expression were increased (4.8- and 2.7-fold vs OC group respectively) in the epithelium of papillary carcinoma (Table 2 and Fig. 2A, B and E). However, the combination of Mel with EtOH exerted no interference on p53 expression, suggesting that this inhibitory effect may stimulate cell proliferation (Fig. 2A, B and E). Notably, BAX immunostaining and expression was also higher following Mel and EtOH alone (2.3- and 3.1-fold increased vs OC respectively), while the combination Mel + EtOH was unable to alter the protein levels of BAX (Table 2 and Fig. 2A, B and E). Although the caspase-3 immunostaining was intense in the OC + Mel, OC + EtOH, and OC + EtOH + Mel groups (Fig. 2A and Table 2), only Mel alone significantly augmented the levels of caspase-3 (4.3-fold vs OC; Fig. 2B, C and E) and cleaved caspase-3 (2.7-fold vs OC; Fig. 2C and E), which allow us to consider Mel to be important for effective regulation of downstream targets in OC apoptotic signaling. EtOH consumption alone further increased cleaved caspase-3 expressions in these cells (Fig. 2C and E).

Immunofluorescence TUNEL staining detected the level and localization of end-labeling of fragmented DNA in OC cells. Mel therapy resulted in increased TUNEL immunolabeling (fluorescence level augmented from 13 G 4.3% (OC) to 57 G 8.4% (OC Mel), Fig. 2D). Interestingly, EtOH intake significantly increased TUNEL immunolabeling (fluorescence level from 13 G 4.3% (OC) to 59 G 10.6% (OC EtOH), Fig. 2D), and the combination of Mel with EtOH was effective in promoting further high fluorescence level (66 G 12.0%). Unexpectedly, this increase possibly occurred due to another kind of cell death, since both caspase-3 and activated caspase-3 were not significantly affected by the Mel EtOH combination. To properly evaluate OC cell turnover, Ki-67 immunolabeling was carried out (Fig. 3A). Despite no significant differences were found in the proliferation rate of OC cells (Fig. 3B), there was a considerable reduction (P < 0.01) in cell proliferation/apoptosis ratio following the treatments (Fig. 3C). This low rate of cell turnover favorably argues for extending the pro-apoptotic effect of Mel and EtOH.

Mel and EtOH exerted regulatory effects on the expression of anti-apoptotic proteins in OC

To investigate the influence of Mel and EtOH intake on anti-apoptotic proteins, OC tissues displaying papillary aspect were studied for Bcl-2 and survivin expressions. The immunoreactivity for Bcl-2 was intense in OC and
To support the effectiveness of treatment, plasma Mel was measured throughout the experiment, and circulating Mel was higher in the OC+Mel and OC+EtOH+Mel groups as compared with their controls. Collectively, this study demonstrated that either Mel or EtOH intake positively regulated the expression of p53, BAX, and cleaved caspase-3, and downregulated survivin in papillary OC cells. Furthermore, despite addition of EtOH was effective to promote apoptosis, we observed an increased tumor incidence and a severe reduced survival rate in these animals. Owing to its co-carcinogenic activity, EtOH has already been shown to potentiate OC incidence (Chuffa et al. 2013a,b). However, after OC development, the deleterious effects of EtOH leading animals to experience low survival rate may occur by other systemic mechanism(s).

Being a selective process, apoptosis is important in both physiological and pathological conditions. Mutations of the TP53 gene are the most common molecular alterations in human cancer and in OC are drastically decreased along with the treatments (Fig. 4A); however, Bcl-2 protein levels were only significantly downregulated in the OC+EtOH group (1.38-fold reduced vs OC, Fig. 4B and C). In addition, higher levels of survivin were present in the cytoplasm of OC cells, and they were gradually decreased over the treatments (Fig. 4A). Importantly, Mel therapy and EtOH alone efficiently reduced the survivin expressions (1.33- and 1.56-fold vs OC respectively, Fig. 4B and C). Lastly, Fig. 4D illustrates a possible scheme by which Mel therapy or EtOH intake may interact positively or negatively with the apoptotic proteins in the OC cells.

**Mel and EtOH significantly alter the number of positive OC cells related to apoptosis**

Reinforcing our data, total number of immunolabeled positive cells was counted to identify a real effect of treatments and tumor tissue response. Importantly, the number of p53, caspase-3, and TUNEL-positive cells were significantly higher following the treatments. In addition, BAX-positive cells were higher in OC+Mel and OC+EtOH groups compared with OC (Fig. 5). Conversely, the number of Bcl-2-positive tumor cells was dramatically reduced by either Mel or EtOH and Mel+EtOH combination, and only Mel therapy was able to reduce the number of survivin-positive cells (Fig. 5).

**Discussion**

We have reported that Mel therapy efficiently reduced OC masses (size and volume) as well as the incidence of papillary carcinomas (Chuffa et al. 2013a). In the present study, we focused to the apoptotic mechanism of the OC cells after Mel therapy in an EtOH-prefering rat model.

### Table 1  Nutritional parameters and OC-related characteristics

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Groups</th>
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<tbody>
<tr>
<td></td>
<td>OC</td>
</tr>
<tr>
<td>Final body weight (g)</td>
<td>260 ± 9.2</td>
</tr>
<tr>
<td>Body weight gain (%)</td>
<td>30 ± 4.2</td>
</tr>
<tr>
<td>Food consumption (g/day)</td>
<td>14.31 ± 0.46</td>
</tr>
<tr>
<td>ETOH consumption (ml/100 g per day)</td>
<td>–</td>
</tr>
<tr>
<td>Energy intake (kcal/day)</td>
<td>43.66 ± 4.17</td>
</tr>
<tr>
<td>OC masses* (g)</td>
<td>0.24 ± 0.04</td>
</tr>
<tr>
<td>Relative OC masses (g/100 g BW)</td>
<td>0.10 ± 0.03</td>
</tr>
<tr>
<td>OC volume (cm³)</td>
<td>9.3 ± 2.1</td>
</tr>
<tr>
<td>Plasma Mel levels (pg/ml)</td>
<td>26.34 ± 6.7</td>
</tr>
</tbody>
</table>

Values are expressed as the mean ± s.d., n = 20/group. *P < 0.05 vs OC and OC + EtOH groups respectively.

*The left ovary that was chemically induced with DMBA. Two-way ANOVA complemented by Tukey's test.

### Table 2  Scoring of IHC of OC cells

<table>
<thead>
<tr>
<th>Proteins</th>
<th>OC</th>
<th>OC + Mel</th>
<th>OC + EtOH</th>
<th>OC + EtOH + Mel</th>
</tr>
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<tbody>
<tr>
<td>p53</td>
<td>0</td>
<td>++</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>BAX</td>
<td>0</td>
<td>++</td>
<td>++</td>
<td>0</td>
</tr>
<tr>
<td>Caspase-3</td>
<td>+</td>
<td>+</td>
<td>++</td>
<td>+</td>
</tr>
<tr>
<td>TUNEL</td>
<td>0/+</td>
<td>+/++</td>
<td>+/++</td>
<td>+/++</td>
</tr>
<tr>
<td>Ki-67</td>
<td>+++</td>
<td>++/+++</td>
<td>++/+++</td>
<td>++/++++</td>
</tr>
<tr>
<td>Bcl-2</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>0/+</td>
</tr>
<tr>
<td>Survivin</td>
<td>++</td>
<td>+/+</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

Intensity of OC staining was analyzed by pathologist visual scoring as 0 (no staining), + (weak staining), ++ (moderate staining), or +++ (strong staining). n = 10 animals/group. Five selected OC sections were analyzed per animal.
Figure 2

Immunohistochemical/fluorescence localization and western blot analysis of p53, BAX, caspase-3, and cleaved caspase-3 in serous papillary OC.

(A) The immunoreaction of p53 and BAX was intense in the surface epithelium of the OC + Mel (II and VI) and OC + EtOH (III and VII) groups (arrows), but not in the stroma. Absence of p53 and BAX reactions was observed in the OC (I and V) group (arrow), while a weak p53 reaction appeared in the epithelium of the OC + EtOH + Mel (IV) group (arrow). A weak caspase-3 reaction was notable in the epithelium of the OC (IX), in contrast to a high immunostaining in the OC + Mel (X), OC + EtOH (XI), and OC + EtOH + Mel (XII) groups (arrow). Bar = 20 μm. Negative controls were used. (B and C) Representative profiles of extracts (70 μg protein) pooled from ten samples per group. Extracts obtained from individual animals were used for densitometric analysis of the p53, BAX, caspase-3, and cleaved caspase-3 levels following normalization to the β-actin or α-tubulin. (D) Merged images of the TUNEL immunofluorescence and DAPI nuclear staining in OC (I), OC + Mel (II), OC + EtOH (III), and OC + EtOH + Mel (IV); details of the DNA fragmentation (Alexa Fluor 488, bar = 10 μm). (E) Analysis of relative optical density of the blots. All results are expressed as the mean ± s.d. (n = 10). aP < 0.05 vs OC; bP < 0.05 vs OC + Mel; and cP < 0.05 vs OC + EtOH. A full colour version of this figure is available at http://dx.doi.org/10.1530/ERC-15-0463.
associated to an early event in high-grade serous carcinoma with ubiquitous occurrence (Salani et al. 2008, Ahmed et al. 2010). Recent data have indicated that complete absence of p53 immunolabeling is associated with poor-prognosis-related mutation in OC (Yemelyanova et al. 2011). Notably, we found a significant overexpression of p53 and BAX in papillary epithelium following either Mel therapy or EtOH intake, but not after their combination. In fact, the cytotoxic effect of Mel in tumor cells is counter-intuitive, since high levels of Mel are related to protection of normal cells and inhibition of cell proliferation (Martín et al. 2006, García-Navarro et al. 2007). The increase in the levels and activity of p53 was accompanied by activation of other apoptosis-related signaling pathways in liver cancer cells and LNCaP androgen-dependent prostate cancer cells treated with 1 mM Mel (Martín-Renedo et al. 2008, Joo & Yoo 2009). Nuclear accumulation of the p53 leads to altered transcription of death-susceptible p53-responsive genes (e.g. Bax), while concomitantly suppress the expression of the Bcl-2 survival gene (Krajewski et al. 1994). In our study, the effect of Mel combined with EtOH inhibited the levels of pro-apoptotic p53 and BAX levels.

According to Ferreira et al. (2014), Mel is able to modulate tumor development and metastasis, thus attenuating the signaling pathway related to OC cell survival. Similarly, our findings demonstrated that Mel promoted a reduction in OC volume and masses displaying pro-apoptotic and anti-proliferative actions on the OC development. In breast cancer cell line, Mel significantly increased the expression of p21WAF1 mediated by p53, resulting in the imbalance of proliferation/apoptosis ratio (Mediavilla et al. 1999). Mel is an undisputed agent capable of modulating the clock genes expression (Blask et al. 2011). Alternatively, differential expression of clock genes (e.g. PER1 and PER2) may exert influences on cancer cell proliferation and growth, as evidenced in breast cancer (Yang et al. 2009) and other malignancies (Wood et al. 2009).

Caspases represent one important player in the initiation and execution of apoptosis, and is a consensus that low levels or impairment in effector caspase function may decrease apoptosis and facilitate carcinogenesis (Wong 2011). Devarajan et al. (2002) reported that caspases-3 mRNA levels in commercially available total samples from breast and cervical tumors were undetectable or substantially decreased in ovarian tumors. We observed a reduced expression of total and cleaved-caspase-3 in OC cells, and conversely, Mel therapy over the 60 days significantly increased total caspase-3 expressions. Furthermore, nuclear active caspase-3 was upregulated by either Mel or EtOH intake. These findings are in accordance to Kim et al. (2012), which Mel treatment resulted in the sub-G1 DNA contents and TUNEL-positive cells in OC cell line (SK-OV-3), and combined treatment of Mel with cisplatin increased the cleavage of caspase-3 and poly-(ADP-ribose) polymerase. Importantly, fragmentation of DNA evidenced by the TUNEL-positive cell immunofluorescence was higher in the OC+Mel, OC+EtOH, and OC+EtOH+Mel, thereby proving that either Mel or EtOH are able to undergo apoptosis in OC cells of EtOH-prefering rats. Like a double-edged sword, defects or abnormalities in apoptotic pathways may be an interesting target of cancer therapy.
New adjuvant drugs or treatment strategies that can restore the apoptotic signaling have a potential to eliminate cancer cells. Targeting Bcl-2 and other anti-apoptotic proteins or silencing the upregulated anti-apoptotic proteins or genes are also a promising approach for the treatment of OC. In our study, only EtOH-consuming animals bearing OC had a significant reduction of Bcl-2 levels. Although our results were unable to demonstrate a real effect of Mel in the downregulation of Bcl-2, there was a profound reduction in survivin expression following either Mel therapy or EtOH intake. It has been extensively studied the role of high concentrations of Mel in determining the pro-apoptotic actions related to early oxidative stress in tumor cells (Trubiani et al. 2005, Rubio et al. 2007). For example, Trubiani et al. (2005) described activation of caspase-3 and cytochrome c, and a decrease in the levels of Bcl-2 after treatment with 2 mM Mel in lymphoma cell.
Acute myeloid leukemia HL60 cells also exhibit an increase in activated caspase-9 and caspase-3, and in the levels of BAX and cytochrome c released to the cytoplasm, as well as a reduction in Bcl-2 levels (Rubio et al. 2007). Although we did not explore any parameter of the extrinsic pathway, these data strongly support our results given that high and prolonged treatment with Mel activates the intrinsic apoptotic pathway in OC cells. More specifically, the relationship between apoptosis and reactive oxygen species formation following Mel therapy will be further investigated in other studies.

Survivin is a bifunctional protein capable of regulating cell proliferation and suppressing apoptotic cell death (O’Driscoll et al. 2003). Survivin overexpression in tumor and normal cells shows inhibition of cell death induced by many apoptotic stimuli, including Fas, BAX, caspases, and anti-cancer drugs. Notably, OC samples in our study showed higher expression of cytoplasmic survivin, and surprisingly, Mel therapy efficiently reduced their levels. Recently, Mel therapy significantly inhibited the growth of human hepatocellular carcinoma and induced apoptosis along with the downregulation of survivin and XIAP (Fan et al. 2013). However, the exact mechanism(s) by which Mel modulates survivin remain inconclusive. As a number of reports elicited survivin and other inhibitors of apoptosis to have poor prognosis in various cancer types, targeting these anti-apoptotic molecules may provide a therapeutic potential for the treatment of OC.

In summary, Mel therapy efficiently reduced OC by altering both volume and masses, while leading to apoptosis of the OC cells. Importantly, either Mel or EtOH intake promoted an increase of p53, BAX, and cleaved caspase-3 with concomitant DNA fragmentation in OC tissue, thus acting as a positive regulator of pro-apoptotic proteins. Conversely, Mel exerted a negative regulation upon anti-apoptotic proteins which suggest Mel to be an inducer of apoptosis in OC of EtOH-prefering rats. Although additional studies are needed, Mel could be used as an adjuvant agent to promote apoptosis in OC tissue.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Author contribution statement
L G A Chuffa, F E Martinez, and M S Alves: collected and analyzed the data, drafted the manuscript, and conceived the main idea of the study. M Martinez, P F F Pinheiro, L A L Júnior, R F Domeniconi, and I C C Camargo: participated in the acquisition of data and in the design and intellectual
conception of the study. All authors performed the statistical analysis and approved the final version of the manuscript.

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