Mitotic Arrest and Apoptosis in Breast Cancer Cells Induced by *Origanum majorana* Extract: Upregulation of TNF-α and Downregulation of Survivin and Mutant p53

Yusra Al Dhaheri¹, Ali Eid¹, Synan AbuQamar¹, Samir Attoub², Mohammad Khasawneh³, Ghenima Aiche⁴, Soleiman Hisaindee³, Rabah Iratni¹

¹Department of Biology, College of Science, United Arab Emirates University, Al-Ain, United Arab Emirates, ²Department of Pharmacology & Therapeutics, Faculty of Medicine & Health Sciences, United Arab Emirates University, Al-Ain, United Arab Emirates, ³Department of Chemistry, College of Science, UAE University, United Arab Emirates University, Al-Ain, United Arab Emirates, ⁴Département de Biochimie et Microbiologie, Faculté des sciences biologiques et agronomiques, Université Mouloud Mammeri, Tizi-Ouzou, Algeria

Abstract

**Background:** In the present study, we investigated the effect of *Origanum majorana* ethanolic extract on the survival of the highly proliferative and invasive triple-negative p53 mutant breast cancer cell line MDA-MB-231.

**Results:** We found that *O. majorana* extract (OME) was able to inhibit the viability of the MDA-MB-231 cells in a time- and concentration-dependent manner. The effect of OME on cellular viability was further confirmed by the inhibition of colony growth. We showed, depending on the concentration used, that OME elicited different effects on the MDA-MB-231 cells. Concentrations of 150 and 300 μg/mL induced an accumulation of apoptotic-resistant population of cells arrested in mitosis and overexpressing the cyclin-dependent kinase inhibitor, p21 and the inhibitor of apoptosis, survivin. On the other hand, higher concentrations of OME (450 and 600 μg/mL) triggered a massive apoptosis through the extrinsic pathway, including the activation of tumor necrosis factor-α (TNF-α), caspase 8, caspase 3, and cleavage of PARP, downregulation of survivin as well as depletion of the mutant p53 in MDA-MB-231 cells. Furthermore, OME induced an upregulation of γ-H2AX, a marker of double strand DNA breaks and an overall histone H3 and H4 hyperacetylation.

**Conclusion:** Our findings provide strong evidence that *O. majorana* may be a promising chemopreventive and therapeutic candidate against cancer especially for highly invasive triple-negative p53 mutant breast cancer; thus validating its complementary and alternative medicinal use.

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* E-mail: R_iratni@uaeu.ac.ae

Introduction

Breast cancer is the most frequently diagnosed cancer among women and ranks second as a cause of cancer death in women after lung cancer. An estimated 226,870 new cases of invasive breast cancer are expected to occur among women in the US during 2012 [1]. Plants have been shown to be an excellent source of new drugs, including anticancer agents. Identification and development of new chemotherapeutic agents from plants have gained significant recognition in the field of cancer therapy and become a major area of experimental cancer research. The majority of the chemotherapeutic drugs used in cancer treatment, is either from plant origin or chemically-altered plant products and phytochemicals [2]. In fact, plant-derived anticancer drugs are much more effective and do not have large side-effect consequences compared to synthetic drugs. Examples of anticancer drugs derived from plants and currently in clinical use include the vinca alkaloids vinblastine and vincristine were isolated from *Catharanthus roseus*, the terpene paclitaxel from *Taxus brevifolia* Nutt., and the DNA topoisomerase I inhibitor camptothecin from *Camptotheca acuminata* [3].

Phytochemicals exert their chemoprevention effect of carcinogenesis through several mechanisms. These include inhibition of genotoxic effects, increased antioxidants and anti-inflammatory activity, modulation of cellular signaling pathways and altering gene expression to inhibit cell proliferation and/or induce apoptosis [4].

It is well known that cancer is pathological condition that has been associated with aberrantly regulated apoptosis. It is currently accepted that certain phytochemicals and whole plant extracts can affect the overall process of carcinogenesis by multiple mechanisms. Since apoptosis provides a physiologic protective mechanism for eliminating genetically damaged cells, initiated cells or
cells progressed to malignancy; phytochemicals affecting apoptosis can have an important effect on carcinogenesis [5].

Increasing number of studies have shown evidence of chemoprevention and chemotherapy by stimulating apoptosis in pre-cancerous and cancerous cells in vitro or in vivo suggesting that apoptosis is likely to be a crucial mechanism to suppress carcinogenesis. Epigallocatechin gallate (EGCG) (from green tea), curcumin (Curcuma longa), quercetin (Vegetable and fruits) are some examples of chemopreventive agents, from natural origin, that induce apoptosis in carcinogenesis models or in human chemoprevention trials [6,7]. Also, a number of plant extracts, such as blueberry [8], mushroom [9], gingerol [10], to name just a few, were shown to have anticancer effects against breast cancer cells.

*Origanum majorana* belongs to the family Lamiaceae, It is commonly known as marjoram. It is a perennial herb and widespread worldwide. A large number of known species of the genus *Origanum* are utilized worldwide as spices and flavoring agents and has a long history of both culinary and medicinal use. *O. majorana* is used as a home remedy for chest infection, cough, sore throat, rheumatic pain, nervous disorders, stomach disorders, cardiovascular diseases, and skin care [11,12]. Many of such traditional uses of *marjoram* species were confirmed in several studies utilizing both in vitro and in vivo approaches.

Several reports indicate that *O. majorana* is very rich in phenolic compounds. The high phenolics content in *Origanum* has a capacity to scavenge free radicals and shown to be associated with the strong antioxidant activity [13]. *O. majorana* was shown to contain phenolic terpenoids (thymol and carvacrol), flavonoids (diosmetin, luteolin, and apigenin), tannins, hydroquinone, phenolic glycosides (arbutin, methyl arbutin, vitexin, orientin, and thymoquin) and triterpenoids (ursolic acid and oleandric acid) [14].

*O. majorana* has been reported to exhibit a significant antimicrobial activity [15]. Several studies have also demonstrated that ethanolic, aqueous extracts and essential oil of *O. majorana* could protect against liver and kidney damage and genotoxicity induced by lead acetate [16–18]. *O. majorana* has also been found to inhibit platelet adhesion aggregation and secretion [19]. Furthermore, it has been shown that this plant exerts a low cytotoxicity on several hepatoma cell lines [20]. It has been shown by Al Harbi that extract of *O. majorana* reduced the side effects induced by cyclophosphamide, an established anticancer drug, without altering its cytotoxicity [12].

In the present study, we investigated the effect of *Origanum majorana* ethanolic extract (OME) on breast cancer cells. We examined the effects of OME on cell viability, cell cycle, apoptosis, and the levels of several cell cycle and apoptosis control proteins in the highly proliferative and invasive Estrogen Receptor (ER)-negative, mutant p53 breast cancer cell lines MDA-MB-231. Our results demonstrate that OME can inhibit the growth of the MDA-MB-231 cells by causing cell cycle arrest and apoptosis dependent on the downregulation of survivin and mutant p53.

**Materials and Methods**

**Preparation of the *Origanum majorana* Ethanolic Extract**

*Origanum majorana* commonly known as “marjoram” and used as a culinary herb, was obtained from a private commercial farm located in the Tyre region of Lebanon. All necessary permits were obtained for the harvesting of the leaves. The identity of the OM dried leaves used in this study was further confirmed by a plant taxonomist. 5.0 g of the dried leaves were ground to a fine powder using a porcelain mortar and pestle. The powder was suspended in 100 mL of 70% absolute ethanol and the mixture was kept in the dark for 72 hours at 4°C in a refrigerator without stirring. The mixture was then filtered through a glass sintered funnel and the filtrate was evaporated to dryness using a rota-vapor at room temperature. The green residue was kept under vacuum for 2–3 hours and its mass was recorded.

**Cell Culture and Reagents**

Human breast cancer cells MDA-MB-231 were maintained in DMEM (HyClone). The culture media was supplemented with 10% fetal bovine serum (invitrogen), 100 U/ml penicillin/ streptomycin (invitrogen). Antibodies to phosphor-H2AX (ser139) (07–164), acetyl-Histone H3 (06–599), acetyl-Histone H4 (06–866) and phosphor-Histone H3 (ser10) (05–1336), p53 (E26, 04–241) and cyclin B1 (05–373) were obtained from Millipore. Antibodies to survivin (sc-17779), β-actin (C4, sc-7334), goat anti-mouse IgG-HRP (sc-2005) and goat anti-rabbit IgG-HRP (sc-2004) were obtained from Santa Cruz Biotechnology, Inc. Antibody to TNF-α (ab9739) was obtained from abcam and antibodies AlexaFluor 488 goat anti-rabbit IgG (H+L) (A11008), AlexaFluor 594 goat anti-mouse IgG (H+L) (A11058) were obtained from invitrogen.

**Cellular Viability**

Cells were seeded in triplicate in 96-well plates at a density of 5,000 cells/well into 96-well plates. After 24 h of culture, cells were treated with increasing concentrations of OME or equal volume of vehicle [ethanol] as control and incubated for the indicated time period. Cell viability was determined using a CellTiter-Glo Luminescent Cell Viability assay (Promega Corporation, Madison, USA), based on quantification of ATP, which signals the presence of metabolically active cells. Luminescent signal was measured using Berthold LB12 Luminometer. Data were presented as proportional viability (%) by comparing the treated group with the untreated cells, the viability of which is assumed to be 100%.

**Measurement of Caspase 3/7, 8 and 9 Activities**

MDA-MB-231 cells were seeded at the density of 3,000 cells/well into 96-well plate in triplicate and treated with indicated concentrations of OME or equal volume of vehicle [ethanol] as control for 24 and 48. Caspase-3/7, 8 and 9 activities were measured using a luminescent caspase-Glo 3/7, caspase8 and caspase 9 assay kit (Promega Corporation, Madison, USA) following the manufacturer’s instructions. Briefly, caspase reagents were added to triplicate wells of a 96-well plate which was then mixed in an orbital shaker and incubated for 2.5 h at room temperature in the dark. Luminescent signal was measured as described above.

**Flow Cytometric Analysis of Cell Cycle**

MDA-MB 231 cells were seeded in 100 mm culture dishes and cultured for 24 h before addition of various concentrations of OME or equal volume of vehicle [ethanol] as control. After incubation for the indicated time, cells were harvested by trypsin release, washed twice with ice-cold PBS, resuspended in 500 μl PBS, fixed with an equal volume of 100% ethanol and incubated for at least 12 h at −20°C. Before flow cytometry analysis, cells were pelleted, washed twice with PBS, permeabilized in 0.1% Triton X-100/PBS for 15 min on ice, pelleted and then resuspended in PBS containing 40 μg/ml propidium iodide and 25 μg/ml RNase A, and incubated at 37°C for 30 min. Cell samples were analyzed on the BD FACSCanto II (Becton
PBS. After washing with PBS, sample cells were mounted in
were washed three times with PBS and placed for 1 h at RT in the
milk/PBS overnight at 4°C. Following incubation, the exponential
growth of cells was determined at the
indicated concentrations of OME or equal volume of vehicle (ethanol) as control for 24 h. Cells were then fixed in 10% formalin
solution (4% paraformaldehyde) (Sigma-Aldrich) for 5 min at RT
followed by permeabilization in PBS containing 0.1% Triton X-
100 for 5 min at RT. Cells were then washed three times with PBS,
blocked with 5% nonfat dry milk in PBS for 30 min at RT
and incubated with the primary antibody diluted, at the
concentration suggested by the manufacturer, in 1% nonfat dry
milk/PBS overnight at 4°C. Following overnight incubation,
cells were washed three times with PBS and placed for 1 h at RT in the
presence of rhodamine-conjugated or fluorescein-conjugated
primary antibody diluted at 1:200 in 1% nonfat dry milk/
PBS. After washing with PBS, sample cells were mounted in
Fluoroschield with DAPI (Sigma-Aldrich) and examined under
Nikon Ti U fluorescence microscope.

Cell Extract and Western Blotting Analysis
Cells (1.8×10⁶) were seeded in 100 mm culture dishes and
seeded for 24 h before addition of various concentrations of
OME, or equal volume of vehicle (ethanol) as control. After
incubation for the indicated times, cells were washed twice with
ice-cold PBS, released by scrapping, pelleted and lysed in RIPA
buffer supplemented with protease/phosphatase inhibitor cocktail.
Following incubation for 30 min on ice, the cell lysate was
obtained by centrifugation at 14,000 rpm for 20 min at 4°C.
Protein concentration of lysates was determined by BCA protein
assay kit (23225, Thermo Scientific) and the lysates were adjusted
with lysis buffer. Aliquots of 25 μg of total proteins were resolved
onto 10–12% SDS-PAGE. Proteins were transferred to nitrocel-
lulose membranes (88018, Thermo Scientific) and blocked for 1 h
at room temperature with 5% non-fat dry milk in TBST (TBS and
0.05% Tween 20). Incubation with specific primary antibodies was
performed in blocking buffer overnight at 4°C. Horseradish
peroxidase-conjugated anti-IgG was used as secondary antibody.
Immunoreactive bands were detected by ECL chemiluminescent
substrate (32209, Thermo Scientific). Membrane stripping by
membrane stripping by Western blot stripping buffer (21039, Thermo Scientific) according to the manufacturer’s instructions.

Colony Formation Assay in Soft Agar
Assays were performed in six-well plates. The lower (base) layer
consisted of 1 ml 2.4% Noble agar. The base layer was overlaid
with a second layer consisting of 2.9 ml growth medium, 0.3%
Noble agar, and 3x10⁶ MDA-MB-231 cells. Agar at 50°C
was mixed with medium at 37°C, plated, and left to set for 10 min.
Then 2 ml of growth medium was added on the top of the second
layer. Plates were incubated at 37°C, humidiﬁed, and 5% CO₂.
Cells were allowed to grow in absence of treatment for 14 days
until visible colonies were formed. Plates were fed twice a week
with 2 ml of regular complete growth medium (DMEM). After 2
weeks, plates were fed with medium containing either the
indicated concentrations of OME or equal volume of vehicle
(ethanol) as control and the colonies were allowed to grow for one
more week. Following treatment, the plates were washed twice
with PBS and then colonies were ﬁxed with 10% ice-cold
methanol for 10 min and washed once with PBS. The colonies
were allowed to stain for 1 h in solution containing 2% Giemsa.
The size of the colonies were measured, counted using a
microscope (10X) and the colony size was categorized as Large
(>200 μM) or small (50–200 μM). Colony sizes were expressed as
a percentage of total counted colonies and then compared to the
vehicle treated controls (ethanol). The experiment was repeated
two times.

Statistical Analysis
Results were expressed as means ± S.E.M. of the number of experiments. A Student’s t test for paired or unpaired values
was performed and a p value of <0.05 was considered statistically
significant.

Results

O. majorana Extract Inhibits the Viability of the MDA-MB-
231 Breast Cancer Cells
To examine the anticancer activity of Origanum majorana extract
(OME) on breast cancer cells, we first measured the effect of
various concentrations of the extract (0, 50, 150, 300, 450 and
600 μg/mL) on the proliferation of the MDA-MB-231 breast
cancer cell line (Figure 1A). Our results show that the exposure of
the MDA-MB-231 to OME decreased cellular viability in a
concentration- and time-dependent manner. The IC₅₀ (producing
half-maximal inhibition) was approximately 350 μg/mL at
24 h and 400 μg/mL at 48 h treatment. The observation of the
OME-treated MDA-MB231 cells under light microscopy also
revealed that the number of cells decreased when the concentra-
tion increased. Furthermore, as shown in figure 1B, light
microscopy observation of MDA-MB231 cells treated with
concentrations of 150, 300 and 450 and 600 μg/mL OME,
underwent morphological changes characterized by a loss of their
epithelial morphology visible after 24 h of treatment. Echinoid
spikes and cellular rounding, characteristics of apoptotic cells were
also observed.

O. majorana Extract Leads to Mitotic Arrest and
Apoptosis in MDA-MB-231 Cells
The ability of an anticancer drug to affect cell cycle distribution
can provide information regarding its cytotoxic mechanism(s) of
action. For this reason, we investigated the effect of OME on cell
cycle distribution by flow cytometry. MDA-MB 231 cells were
treated with indicated concentrations of O. majorana for 24 h and
subjected to cell cycle analysis. At the concentration of 150 μg/
L, OME caused obvious G2/M arrest on these cells (figure 2A).
Indeed, the population of G2/M increased signiﬁcantly from 23 to
51.7% as the concentration of the OME increased to 150 μg/mL.
Figure 2. Induction of G2/M cell cycle arrest and apoptosis by *O. majorana* extract in MDA-MB-231 cells. (A) MDA-MB-231 cells (1.8×10⁶) seeded on 100 mm culture dish were exposed various concentrations of *O. majorana* extract or equal volume of vehicle (ethanol) as control for 24 h. Following treatment, cells were harvested, fixed, stained with propidium iodide, and analyzed for cell cycle distribution by flow cytometry. Data represent the mean of three independent experiments. The percentage of cells in sub-G1 (apoptosis), G1, S and G2/M appears at the upper right of each graph. (B) Expression of cell cycle regulator in OME-treated MDA-MB-231. Western blot analysis of phospho(ser10)-H3, and cyclin B1 in MDA-MB-231 cells exposed for 24 h to ethanol or indicated concentrations of OME. (C) Stimulation of caspase 3/7 activity in MDA-MB-231 cells after exposure to OME (0–600 μg/mL) for 24 h and 48 h, relative to a similar amount of viable ethanol-treated cells. The relative caspase 3/7 activity was normalized to the number of viable cells per well and is expressed as fold of induction compared to the control. (D) Concentration-dependent induction of PARP cleavage in OME-treated MDA-MB231 cells. Cells were treated with or without increasing concentrations of the extract and proteins were extracted as described in Materials and Methods. Western blot analysis was carried out using anti-PARP antibodies. (*p<0.05, **p<0.005 and ***p<0.0005).
indicating that OME-treated MDA-MB-231 cells were arrested in G2/M phase. A slight increase in the sub-G1 population (6.2%) indicates that a small population of OME-treated MDA-MB-231 cells are undergoing cell death was also observed by these concentrations. Similar results were obtained for 300 μg/mL of OME (data not shown). Interestingly, at higher concentrations of OME, flow cytometry analysis revealed a dramatic increase in the apoptotic population (sub-G1 peak) raising from 0.9% in the control to 48.4% and 56.7% in cells treated with 600 and 450 μg/mL, respectively (Figure 2A).

Figure 3. Differential regulation of survivin expression by OME in MDA-MB-231 cells. Western blot analysis showing a differential effect on survivin expression by different concentrations of OME in MDA-MB-231 cells. Whole cell protein were extracted from OME or vehicle (ethanol)-treated cells and subjected to Western blot analysis, as described in Materials and Methods, for survivin and β-actin (loading control) proteins.

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To determine whether OME induced cell cycle arrest specifically at mitosis or G2 phase, we examined the phosphorylation status of histone H3 (Ser 10). Histone H3 is phosphorylated at serine 10 during mitosis by aurora kinase and the phosphorylation status of H3 is considered a marker of mitosis [21]. We therefore, investigated the expression of p(ser10)H3 and found that treatments 150 and 300 μg/mL of O. majorana significantly increase the phosphorylation level of histone H3 (Figure 2B, upper panel). This result indicates that OME induces mitotic arrest of MDA-MB-231 cells. Next, we investigated the mechanism of OME-induced mitotic arrest. Accumulation of cyclin B1 is well known to play an important role in G2/M transition. Knock-down of cyclin B1 with siRNA produces cell cycle arrest predominantly in the G2 phase [22], while an upregulation of cyclin B1 invokes mitotic arrest [23]. We, therefore, investigated the protein level of cyclin B1 in OME-treated MDA-MB-231 cells. We found that treatment with these concentrations of OME leading to mitotic arrest, caused also an increase of cyclin B1 protein in MDA-MB-231 cells (Figure 2B, lowe panel), suggesting that cyclin B1 accumulation might play a crucial role in OME triggered mitotic arrest. Taken together, our data reveal a differential concentration effect of OME on cell cycle progression of the MDA-MB-231 cells. Whereas, lower concentrations of OME (150 and 300 μg/mL) induced a major mitotic arrest with a slight increase in the apoptotic population, higher concentrations (450 and 600 μg/mL) induced a massive cell death by apoptosis. The reduced cell viability observed in MDA-MB-231 cells treated with low concentrations of OME, is possibly due to a large extent to an inhibition of cell proliferation rather than to cell death.

Apoptosis in OME-treated MDA-MB-231 cells was further examined by measuring caspase 3/7 activation in MDA-MB-231 cells treated with various concentrations (300, 450 and 600 μg/mL) of OME after 24 h of treatment. A concentration- and time-dependent activation of caspase 3/7 was detected in treated cells (Figure 2C). Interestingly, cleavage of the poly(ADP-ribose) polymerase (PARP) occurred only in cells treated with higher (450 and 600 μg/mL) but not at lower concentrations (150 and 300 μg/mL) of OME (Figure 2D). It is noteworthy to mention that at these lower concentrations of OME, only low apoptotic induction rate was observed despite the detection of caspase 3/7 activation.

Concentration-dependent Regulation of Survivin Expression by O. majorana Extract
Survivin, a member of the inhibitor of apoptosis protein (IAP) family, plays an important role in both the regulation of cell cycle and the inhibition of apoptosis. Survivin levels increase in G2/M phase conferring resistance to apoptosis to the G2/M arrested cells. However, a decrease in survivin levels sensitizes the cells to apoptosis. Several studies have reported that survivin exerts its negative effect on apoptosis by inhibiting the activity of caspase 3, 7 and 9. Therefore, we examined a possible involvement of survivin in the cell cycle arrest and apoptosis triggered by OME. Toward this, we have analyzed, by Western blotting, the expression of survivin in response to various concentrations of OME after 24 h treatment. Interestingly, we observed a differential concentration-effect of OME on survivin expression on the MDA-MB-231 cells (Figure 3). We found that low concentrations of OME led to a substantial increase in the level of survivin, while higher concentrations caused a drastic decrease of survivin (99%). Based on these results, we conclude that OME exerts a concentration-dependent effect on MDA-MB-231 cells. Low concentrations of OME induced a mitotically arrested cells accompanied by survivin upregulation which, in turn, conferred resistance to cell death to this population of cells, probably by inhibiting the activity of caspase 3/7 which was monitored by the absence of PARP cleavage at these concentrations. Treatment of MDA-MB-231 cells with higher concentrations of OME caused a dramatic decrease in survivin expression and consequently sensitized MD-MB-231 cells to apoptosis.

O. majorana Extract Activates the Extrinsic Pathway for Apoptosis via an Upregulation of TNF-α and Activation of Caspase 8
Having shown that OME induces the activation of the effector caspases 3/7, we looked at the activity of the initiator caspases of the extrinsic and intrinsic cell death pathway, namely caspase 8 and caspase 9, respectively. Surprisingly, no caspase 9 activation was detected in response to various concentrations of OME after 24 h of treatment (Figure 4A). On the other hand, caspase 8 activity increased in a concentration-dependent manner in response to OME treatment (Figure 4A). This result demonstrates that the apoptotic effect of the extract on MDA-MB-231 is dependent on caspase 8 activity, which implicates only the extrinsic cell death pathway since neither caspase 9 activation (Figure 4A) nor a change in Bax/Bcl2 ratio (data shown) were observed. After showing that the extrinsic cell death pathway is implicated in OME-dependent apoptosis, we were then interested in determining how this pathway is activated by OME. We determined the changes in the expression level of the tumor necrosis factor alpha (TNF-α) in response to OME after 24 h treatment. Western blot analysis revealed a clear increase in the level of TNFα in MDA-MB-231 cells in response to OME treatment (Figure 4B). The upregulation of TNF-α was further confirmed by immunofluorescence assay (Figure 4C). Even though we have shown that OME exerts its effect via the activation of the extrinsic pathway of cell death, we cannot rule out, at this stage,
Anti-Breast Cancer Activity of *Origanum majorana*

A. MDA-MB-231 (24 hr)

B. 24 h

C. Images of Vehicle, 150 μg/mL, and 300 μg/mL showing DAPI, TNFα, and Merge.
O. majorana Lead to Depletion of Mutant p53 in MDA-MB-231 and Upregulation of p21

Next, we tested the effect of OME on the expression of the tumor suppressor p53 in MDA-MB-231. Toward this aim, cells were treated with various concentrations of OME and the protein level of the mutant p53 determined. We found that low concentrations of 150 and 300 µg/mL of OME led to a slight increase in the protein level of mutant p53 (Figure 5, upper pannel). Most importantly, Western blotting analysis revealed apoptotic concentrations (450 and 600 µg/mL of OME) led to almost complete depletion of mutant p53 in MDA-MB-231 cells. This result is a potentially important finding because of the role of mutant p53 protein in human cancers. Because mutant p53 renders cancer cells more resistant to anticancer drugs, abolishing mutant p53 may therefore offer a promising approach for cancer prevention and therapy.

Because p21 protein has been reported to inhibit growth and apoptosis, we investigated whether the growth inhibition mediated by low concentrations (150 and 300 µg/mL of OME) was also associated with an induction of p21. Western blotting showed an upregulation of p21 protein with at least 2.5 fold increase in cells treated with low concentrations of OME, while a little or no effect on p21 expression was observed with higher concentrations of OME (Figure 5, lower pannel). Based on that, we can postulate that p21 upregulation contributes, at least partially, to the cell cycle arrest observed with lower concentrations, while it has little or no role in cell death occurring at higher concentrations of OME.

**Figure 5.** Expression levels of mutant p53 and p21 in *O. majorana*-treated MDA-MB-231 cells. Cells were treated with various concentrations of (150, 300, 450 and 600 µg/mL) of OME or equal volume of vehicle (ethanol) as control for 24 h and the expression of mutant p53, p21 and β-actin were estimated by Western blot.

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O. majorana Extract Induces Hyperacetylation of Histone H3 and H4 in the MDA-MB-231 Cells

Previously, expression of p21 and increased histone hyperacetylation have been linked to apoptosis and to growth arrest. Therefore, we examined the acetylation profile of histone H3 and H4 in MDA-MB-231 in response to treatment for 24 h to increasing concentrations of OME. As shown in figure 6A, the time course analysis showed a gradual increase in acetylated histones, H3 and H4. A marked overall increase in the acetylation status of histone H3 and H4 was also detected by immunofluorescence staining (Figure 6B). Altogether, these results showed that OME induced hyperacetylation of histone H3 and H4.

**Figure 4.** *O. majorana* induces apoptosis by activation of caspase 8 and upregulation of TNF-α. (A) *O. majorana* extract induces an activation of caspase 8 but not caspase 9 in MDA-MB-231 cells. MDA-MB-231 cells were incubated with various concentrations of the extract for 24 h. The caspase activation induced by the OME was assayed as described under Materials and methods. The relative caspase 8 and 9 activity was normalized to the number of viable cells per well and is expressed as fold of induction compared to the control. (B) Western blot analysis showing an increase in cellular TNF-α protein in the MDA-MB-231 cells treated with OME. Whole cell protein were extracted from OME-treated cells or vehicle (ethanol)-treated cells and subjected to Western blot analysis, as described in Materials and Methods, for TNF-α and β-actin (loading control) proteins. (C) Immunofluorescence staining for TNF-α in OME-treated MDA-MB-231 cells. Cells were treated with 150 and 300 µg/mL of the extract for 24 h, fixed, permeabilized, and then processed for immunofluorescence using antibodies against TNF-α protein. DAPI was used as a nuclear stain. (**p<0.05 and ***p<0.005).

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Accumulation γH2AX, a Marker of Double Strand Breaks, in *O. majorana* Extract Treated MDA-MB 231 Cells

We sought to investigate whether OME induced DNA damage in MDA-MB-231 cells. For this purpose, MDA-MB 231 cells were cultured for 6 and 24 h in complete media containing either ethanol (control) or increasing concentrations of OME (75, 150, 300, 450 and 600 µg/mL). DNA damage was determined by measuring the levels of phosphorylated H2AX (γH2AX) after 6 and 24 h of treatment of the MDA-MB-231 cells with OME. Western blotting analysis revealed a time- and a concentration-dependent increase in the levels of γH2AX in response to OME treatment (Figures 7A) indicating an accumulation of double strand breaks in these cells. The increase in DNA damage was also assessed by immunofluorescence staining of γH2AX in cells treated with 150, 300 and 450 µg/mL OME for 24 h. Figure 7C clearly shows a concentration-dependent increase of γH2AX foci in response to OME. Since the activation of γH2AX occurred as early as 6 h, a time in which no cell death (data not shown) or caspase3/7 activation were observed (Figure 7B), this rules out the possibility that the resulting DNA damage is a consequence of DNA fragmentation resulting from caspases' activities and further confirms the potential of this OME extract to induce double strand DNA breaks in a dose-dependent manner.

**Figure 6.** *O. majorana* Extract Treated MDA-MB 231 Cells

To further confirm the inhibitory potential of *O. majorana* on MDA-MB-231 cells, we sought to determine if OME could inhibit the further growth of already formed MDA-MB-231 colonies. For this purpose, MDA-MB-231 cells were first allowed to grow and form visible colonies in absence of treatment. After 14 days of growth, colonies were incubated with ethanol as control and with OME and allowed to grow for one more week. Figures 8 shows that the size of the ethanol-treated (control) colonies kept growing compared to the size of the two weeks colonies; more large colonies were obtained in the three weeks plate, while less small colonies were counted, indicating that small colonies became larger in size. Interestingly, OM-treated colonies shows regression in colony size compared to the two weeks colonies. In OM-treated plates, the number of large size colonies counted was less than what was obtained in the two weeks plate, while the number of
OME did not induce mitochondria-mediated apoptotic pathway caspase 8, caspase 3, and PARP cleavage. On the other hand, activated at 24 h and induced downstream signalling such as mediated extrinsic apoptotic pathway. TNF- through activation of caspase 9 and consequently activation of caspase 3 cleave Bax and Bak leading to the formation of the apoptosome, change in expression of the anti-apoptotic Bcl2 family members, interaction of the ligands such as TNF- surface or by the intrinsic mitochondria-mediated pathway. The ligand-receptor interaction severe depletion of the mutant p53 and survivin proteins from the body of evidence: (i) increase in the level of p(ser10)H3, a mitotic marker, and (ii) an increase of cyclin B1 protein level, whose upregulation was reported in several mitotically arrested cells. We have also shown that the cell cycle arrest correlates with an upregulation of the CDK inhibitor p21 and the anti-apoptotic protein, survivin. At high concentrations, however, OME induced a massive apoptosis demonstrated by a dramatic increase in the sub-G1 population. We have demonstrated that OME exerts its apoptotic effect by activating the cell death extrinsic pathway is mediated, at least partially, via the activation of TNF- caspase 9, and caspase 3 cleavage. We have also shown that OME-induced apoptosis is also mediated by an increase in DNA damage, revealed by an upregulation of γH2AX, severe depletion of the mutant p53 and survivin proteins from the treated cells.

The process of apoptosis can be induced either by the extrinsic pathway which involves signalling from death receptors at the cell surface or by the intrinsic mitochondria-mediated pathway [24]. Activation of the death receptor-mediated apoptosis requires the interaction of the ligands such as TNF- and Fas with their transmembrane receptors [25]. The ligand-receptor interaction leads to the activation of the effector caspase 8, which in turn activates the effector caspase 3 directly and/or through mitochondria [26]. The mitochondria-mediated apoptosis pathway is associated with permeabilization of the mitochondria outer membrane, reduced mitochondrial membrane potential (Δψm), change in expression of the anti-apoptotic Bcl2 family members, such as Bcl2 and Bcl-XL and the pro-apoptotic members such as Bax and Bak leading to the formation of the apoposome, activation of caspase 9 and consequently activation of caspase 3 [27]. Intrinsinc- and extrinsic-pathways activated caspase 3 cleave poly(ADP-ribose) polymerase (PARP), thus resulting in apoptosis [28]. In the present study, we showed that OME induced TNF-α mediated extrinsic apoptotic pathway. TNF-α receptor was activated at 24 h and induced downstream signalling such as caspase 8, caspase 3, and PARP cleavage. On the other hand, OME did not induce mitochondria-mediated apoptotic pathway since no change in the BAX/Bcl2 ratio or activation of caspase 9 were detected. We suggest that OME induces apoptosis solely through TNF-α activated signal pathway in MDA-MB-231 cells.

Studies have reported that DNA damage is one of the molecular events associated with cell cycle arrest and apoptosis. Indeed, and many anti-cancer drugs have been shown to induce DNA damage [29,30]. Moreover, cancer cells are reported to be more susceptible than normal cells to DNA damaging agents [31], therefore there is a growing interest in dietary phytochemicals that possess DNA-damaging activity. In the present study we showed that OME elicited DNA damage measured by an increase in a concentration-dependent manner of the marker of DNA damage, γ-H2AX, after treatment with OME for 6 or 24 h. The differential response to the different concentrations of OME (G2/M arrest and/or apoptosis), may be partially mediated by the extent of DNA damage occurred within the genome. Low levels of DNA damage may trigger recruitment of DNA repair complexes, expression of anti-apoptotic and survival proteins leading to arrested cell cycle until the genotoxic lesions are repaired. In this case, survival protein such as survivin gets activated in order to maintain the viability of G2/M arrested cells. On the other hand, when genomes are overwhelmed by DNA damage, cells are eliminated by apoptosis [32,33]. In this study, we have found that high concentrations of OME triggered high level of DNA damage to the genome, causing cell to enter apoptosis. Our data suggest that Origanum majorana possesses a genotoxic effect on MDA-MB-231 cells. At this stage, the mechanism(s) by which OME induces DNA damage remain(s) unknown, and certainly deserves further studies.

Inhibitor of apoptosis proteins (IAPs), which includes survivin, represents a family of anti-apoptotic proteins that bind and inactivate active caspase 3, 7 [34,35] and caspase 9 [36] and can modulate cell division and cell cycle progression [37].

Interestingly, survivin has no effect on caspase 8 activity. Survivin has been shown to be highly expressed in most cancers, where it functions as inhibitor of apoptosis. In breast cancer, overexpressed survivin was shown to protect cells against apoptosis induced by chemotherapeutic agents, such as etoposide [34]. Based on these reports, survivin protein represents an attractive target of particular importance in cancer therapy at large and in breast cancer therapy in particular. In consideration of the recognized role for survivin as a custodian of cancer cell survival, our results suggest that OME might exert its cytotoxic anti-cancer effects at least partly via the down-regulation of survivin. In our study, we have shown that survivin expression is differentially regulated in a concentration-dependent manner by OME. Lower concentrations of OME induced an upregulation of survivin which causes cells to arrest the cell division and to resist apoptosis by inhibiting the cell death program. In fact, we showed that in these arrested cells, PARP cleavage was not detected despite the activation of caspase 3/7. This effect might be mediated by the inhibition of active 3/7 by the upregulated survivin. Survivin function could also account for the mitotic arrest induced by OME. In fact, Survivin, has also been shown to be required for mitotic arrest of Hela cells induced by the anticancer drug UCN-01 [38].

The tumor suppressor protein, p53 is found to be mutated in about 50% of human cancers [39]. Mutant p53 is reported to play a key role in cancer cells resistance to certain anticancer drugs and

Figure 6. O. majorana induces hyperacetylation of histone H3 (AcH3) and H4 (AcH4). (A) Protein levels of Ac-H3 and Ac-H4, extracted from OME-treated cells, were detected by western blot using antibodies specific for the modified histone (Ac-H3) and Ac-H4. β-actin was used as loading control. (B) Immunofluorescence staining of Ac-H3 and Ac-H4 in MDA-MB-231 cells treated with 150, 300 and 450 μg/mL of OME or equal volume of vehicle (ethanol) as control for 24 h. Cells were fixed, permeabilized, and then processed for immunofluorescence using antibodies against the indicated modified histones. DAPI was used as a nuclear stain.

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thus is considered as a potential cancer-specific target for pharmacologic interventions [40,41]. Studies have shown that inhibition of mutant p53 by RNA interference sensitizes cancer cell to cell death by chemotherapeutic agents [42]. Wang et al. 2011, showed that the naturally occurring isothiocyanates (ITCs) phenetyl isoisothiocyanate (PEITC), derived from watercress plant, and the synthetic ITC, 2,2-di phenetyl isoisothiocyanate selectively deplete mutant, but not the wild-type p53, and induce apoptosis in many cancer cells, including the MDA-MB-231 breast cancer cells [43]. Here, we showed that OME led to dramatic decrease in the mutant p53 level in MDA-MB-231 cells. As such, mutant p53 depletion may be an important target for chemoprevention and therapy by O. majorana for TNBC.

Increase in the expression of the cyclin-dependent kinase inhibitor, p21 has been shown to augment G2/M arrest via a p53-independent mechanism in human breast cancer [44]. In most cases, growth arrest was found to be associated with apoptosis. In this study, we showed that low concentrations of OME treatment led to G2/M arrest without significant increase in cell death after 24h treatment. Histone hyperacylation has been demonstrated to be directly linked to the upregulation of p21 and this activation can also occur independently of p53 [45]. Moreover, histone hyperacylation was also shown to be associated with growth suppression and apoptosis. Our data showed that OME induced histone H3 and H4 hyperacylation in MDA-MB-231 cells, suggesting that the anti-breast cancer effects of OME were at least partly mediated by histone H3 and H4 hyperacylation by regulating the expression of the genes controlling these two events. The mechanism by which OME induces histone hyperacylation might involve a histone deacetylase inhibitor (HDI) activity. Interestingly, the plant, O. majorana, contains luteolin, a dietary flavonoid with HDI activity [14].

Figure 7. O. majorana extract induces a dose-dependent activation of γH2AX, a marker of DNA double-strands breaks, in MDA-MB-231 cells. (A) Western blot analysis of phosphor-H2AX (ser 139) in MDA-MB231 cells exposed for 6 and 24h with the indicated concentrations of OME or equal volume of vehicle (ethanol) as control. (B) Immunofluorescence staining for γH2AX in OME-treated MDA-MB-231 cells. Cells were treated with vehicle or 150, 300 and 450 μg/mL extract for 24 h, fixed, permeabilized, and then processed for immunofluorescence using antibodies against p- H2AX (ser 139) protein. DAPI was used as a nuclear stain. doi:10.1371/journal.pone.0056649.g007

Figure 8. Inhibition of colony growth by O. majorana extract. Inhibition of colony growth was assessed by measuring the size of the colonies obtained in vehicle (ethanol)- and OME-treated plates. Data were compared with those obtained for the 2 weeks colonies. Two types of colonies were counted and depending on their diameter were categorized as large (≥200 μm) and small (<200–<50 μm).

|                | 2 weeks | 3 weeks (Vehicle) | 3 weeks (300 μg/mL OME) |
|----------------|---------|------------------|-------------------------|
| % Large size colonies | 38 ± 7.65 | 58 ± 0.3        | 18.2 ± 3.65             |
| % Small size colonies  | 61.3 ± 7.85 | 41.8 ± 0.1      | 81.5 ± 3.77             |

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fact, luteolin was able to decrease the viability of lung, colon, liver and breast cancer cells and induce hyperacetylation of histone H3 and H4 [46]. In light of these data, we conclude that the histone hyperacetylation induced by OME is involved with the HDI activity of luteolin. We are currently undertaking further investigations to better understand the mechanism(s) by which OME induces histone hyperacetylation.

In conclusion, our data are consistent with a model shown in figure 9 which shows the concentration-dependent differential effect of  O. majorana extract on mutant p53, triple negative MDA-MB-231 cells. At low concentrations, OME induced a mitotic arrest associated with low level of DNA damage (Figure 9, thin arrow), upregulation of the CDK inhibitor p21 and the inhibitor of apoptosis, survivin. We believe that these events along with other “yet to be identified” events contribute to the cell cycle arrest. In addition we also propose that, at these concentrations, survivin is also implicated in the blockade of the TNF-mediated apoptosis pathway, by directly inhibiting the activity of the active caspase 3. On the other hands, high concentrations, OME induce massive apoptosis via the activation of the TNF-α extrinsic pathway which is associated with high level of DNA damage (Figure 9, thick arrow) and almost complete depletion of the mutant p53 and surviving proteins from these cells. Our findings provide the first instance of a potential role for OME as an anti-breast cancer agent in vitro which certainly deserves more attention for further explorations to identify novel compounds for breast cancer.

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Author Contributions
Performed most of the cellular and biochemical studies and participated in data analysis: YD. Provided the OM plant and performed cellular viability and caspase experiments: AE. Performed some biochemical analysis, provided antibodies against acetylated histones H3 and H4 and critically revised the manuscript: SAQ. Prepared the OME extract used in this study: MK SH. Performed the colony assay experiments, participated in data analysis of cellular studies and critically revised the manuscript: SA. Carried out FACS experiments: GA. Participated in the conception, design, execution, data analysis, coordination of the study and drafted the manuscript: RI. Read and approved the final version of the manuscript: YAD AE SAQ SA MK GA SH RI.

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Figure 9. The proposed signal pathways on O. majorana-induced cell cycle arrest, at low concentrations, and apoptosis, at high concentrations, in the triple negative mutant p53 human breast cancer MDA-MB-231 cells.

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