What is known about melatonin, chemotherapy and altered gene expression in breast cancer

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Abstract. Melatonin, synthesized in and released from the pineal gland, has been demonstrated by multiple in vivo and in vitro studies to have an oncostatic role in hormone-dependent tumors. Furthermore, several clinical trials point to melatonin as a promising adjuvant molecule to be considered for cancer treatment. In the past few years, evidence of a broader spectrum of action of melatonin as an antitumor agent has arisen; thus, melatonin appears to also have therapeutic effects in several types of hormone-independent cancer, including ovarian, leukemic, pancreatic, gastric and non-small cell lung carcinoma. In the present study, the latest findings regarding melatonin molecular actions when concomitantly administered with either radiotherapy or chemotherapy in cancer were reviewed, with a particular focus on hormone-dependent breast cancer. Finally, the present study discusses which direction should be followed in the next years to definitely clarify whether or not melatonin administration could protect against non-desirable effects (such as altered gene expression and post-translational protein modifications) caused by chemotherapy or radiotherapy treatments. As treatments move towards personalized medicine, comparative gene expression profiling with and without melatonin may be a powerful tool to better understand the antitumor effects of melatonin, the pineal gland hormone.

1. Introduction: Chemotherapy for breast cancer

According to the World Cancer Research Fund International, breast cancer is the most frequent type of tumor suffered by women in the world, with ~1.7 million newly diagnosed cases in 2012 (1). Approximately 1 in 8 women will develop a mammary tumor during her lifetime. The American Cancer Society's report for the USA in 2015 informs of 231,840 new cases of invasive breast cancer, 60,290 women with carcinoma in situ and ~40,290 mortalities (2). In addition, breast cancer is the second cause of mortality by cancer in women, only exceeded by lung tumors (3). The mortality rates for breast cancer have been declining since 1989, particularly in premenopausal women, probably as a result of earlier detection as well as improved treatments (4). One fact that may also explain this decrease is the less frequent administration of hormone replacement therapy (HRT) following the publication of a report (the Women's Health Initiative) published in 2002, which suggests that HRT may be a risk factor that could explain the increase in the incidence of breast cancer (5).

Chemotherapy consists of treatment with cancer-killing drugs administered either intravenously or orally. Chemotherapy compounds are usually applied by intravenous infusion and, through the bloodstream, they reach growing cancer cells in almost all body tissues. Chemotherapy compounds work by targeting cells with a high rate of self-renewal, which is a hallmark of cancer cells (6). Chemotherapy is recommended following surgery (adjuvant chemotherapy): Surgery is performed to dissect the tumor, and adjuvant therapy is administered to try to eliminate any cancer cells that had not been removed by the surgery and may spread out later (7). The most frequently used adjuvant treatments to be administered subsequent to surgery are radiation, chemotherapy, targeted therapy and hormone therapy (8).
Neoadjuvant therapy refers to treatments that are administered prior to surgery instead of subsequent to surgery. The benefits of neoadjuvant chemotherapy are that drugs may cause a decrease in the size of the tumor, thus facilitating tumor removal with a less extensive surgery (9). In addition, administering chemotherapy prior to the tumor being removed may aid the subsequent monitoring of the disease, since in case the first cocktail of drugs does not diminish the tumor size, other compounds may be considered (10). Finally, chemotherapy is also employed as a treatment strategy for patients whose tumor has spread outside the mammary gland and underarm area. Combinations of drugs are commonly used to treat mammary tumors detected in the early stages of carcinogenesis, while advanced cancer is more commonly treated with a unique chemotherapeutic molecule (11). Chemotherapy is usually administered in cycles, with periods of administration followed by resting periods to allow patient recovery and to minimize the side effects of treatment (12).

Since microtubules participate in the migration of chromosomes to opposite ends of mitotic cells during the anaphase, microtubule inhibitors (MIs), also known as microtubule-stabilizing agents, are molecules suitable to use in the treatment of mammary tumors. MI agents include microtubule depolymerizing compounds (Vinca alkaloids) and polymerizing agents (taxanes) (13). Vinca alkaloids derive from the periwinkle plant Catharanthus roseus (14). The first clinical trial demonstrating their efficacy in cancer was reported in 1963 (15). Nowadays, these compounds are produced synthetically and include vinblastine, vincristine, vindesine and vinorelbine (14). The main mechanism that explains their cytotoxicity is their capability to interfere with tubulin, with subsequent microtubule function disruption (particularly concerning microtubules implicated in the formation of the mitotic spindle apparatus), leading to mitosis disruption and finally resulting in metaphase arrest (16). These agents interfere with the assembly of tubulin by introducing a wedge between the contact surfaces of two tubulin molecules (17).

Taxanes are diterpenes obtained from Taxus brevifolia (18). The first reported taxane, named taxol, was initially isolated in 1971 (19). Taxanes present difficulties in formulation because they are poorly soluble in water, and for this reason, the first clinical trial including taxanes was not reported until 1987 (20). The other taxane currently in use is docetaxel, which is obtained from Taxus baccata (21). Both paclitaxel and docetaxel act as spindle poisons, stabilising the tubulin polymers against depolymerisation. In addition, they also promote microtubulin assembly. These two actions together block microtubule dynamics and consequently lead to cell cycle arrest (22). Taxanes induce changes in tubulin spatial conformation, which interferes with the depolymerisation of microtubules in a precise directional way, by binding a specific domain of tubulin located in the internal surface of the microtubule (23,24).

Anthracyclines such as epirubicin and doxorubicin are also commonly used in chemotherapy combined with other chemotherapeutic drugs to treat breast cancer in patients who have had surgery to remove the tumor (25). Their mechanism of action is based on their ability of insertion between two DNA strands, resulting in a DNA-anthracycline complex that inhibits both DNA and RNA synthesis (26). This mechanism also targets DNA for cleavage by topoisomerase II, which leads to a cellular cascade that eventually results in cell death (27). Epirubicin is frequently selected over doxorubicin in numerous chemotherapy protocols, since it appears to have fewer side effects (28). Eribulin is a new anthracycline approved by the Food and Drug Administration of USA in 2010 to treat women with metastatic tumors who had been previously treated with at least two chemotherapeutic compounds indicated for the treatment of metastatic breast cancer (29). This drug exerts its anticancer effects by triggering a mitotic blockade, leading cancer cells to enter apoptosis (30).

Other current approaches in the treatment of breast cancer include compounds developed against specific identified targets (molecular-targeted therapies) that contribute to tumor growth (31). As an example, trastuzumab is a monoclonal antibody used to treat patients with metastatic breast cancer. Trastuzumab is indicated against tumors overexpressing the oncogene human epidermal growth factor receptor 2 (HER2/neu, since it targets the membrane HER2/neu receptor, which normally promotes normal cell growth and is also overexpressed in certain cancerous breast tissues (32). Despite several randomized clinical trials with promising results (33), other reports indicated that both de novo and acquired resistance to trastuzumab could be developed (34). Therefore, other recently described agents have been included in the list of chemicals available to treat HER2-overexpressing mammary tumors. One of them is lapatinib, a reversible inhibitor of both epidermal growth factor receptor (EGFR) and HER2/neu tyrosine protein-kinases (35), which was approved in 2007 for women undergoing metastatic breast cancer with acquired resistance to trastuzumab (36). Lapatinib was well tolerated and displayed encouraging clinical results when used as a first-line therapy agent in ErbB2-amplified tumors, either advanced local tumors or metastatic breast cancer (37). Another molecule to be considered is HKI-272, a (Her2)/neu receptor tyrosine kinase inhibitor with an irreversible mechanism of inactivation and a demonstrated clinical activity, which is well tolerated among both high-dose trastuzumab pre-treated and non pre-treated patients with advanced ErbB2-positive mammary tumors (38). HKI-272 targets a cysteine residue located in the adenosine triphosphate-binding pocket of the ErbB2 receptor, resulting in the inhibition of the downstream signal transduction cascade triggered, and consequently altering the cell cycle regulation (39).

Among the battery of promising new chemicals, drugs that target heat shock protein (Hsp) 90 must be mentioned. Hsp90 belongs to the family of chaperones and establishes associations with a set of different proteins that are known as ‘Hsp90 client proteins’ (40). Multiple Hsp90 client proteins are implicated in breast tumor progression and resistance to chemotherapy treatments, including the receptor protein-tyrosine kinases of the ErbB2 family, estrogen receptor (ER), Akt and mutated versions of p53 (41). The efficacy of Hsp90 inhibitors has been well documented in several preclinical cancer models. One of these inhibitors is 17-AAG, which has completed phase I testing (42,43).

Angiogenesis is a physiological process that consists in new vessel formation. When a tumorous mass of cells is growing, angiogenesis is crucial to maintain both tumor growth and progression (44). Therefore, numerous drugs have been tested during the past decades in the hope of identifying specific inhibitors of the different pathways necessary for angiogenesis (45). One of the key molecules in the formation
of new vessels is vascular endothelial growth factor (VEGF), probably the most studied angiogenic factor (46). VEGF is implicated in the progression of breast cancer and is also a potential prognosis biomarker (47). Bevacizumab (Avastin®) is a recombinant humanized monoclonal antibody that recognises all the known variants of VEGF-A (48). To date, bevacizumab is the unique anti-angiogenic chemical with clearly demonstrated benefits in metastatic breast cancer treatment clinical trials (49,50). Pazopanib is another inhibitor of VEGF receptor (VEGFR) (51). Pazopanib also inhibits the signaling pathways downstream of the platelet-derived growth factor receptor and the mast/stem cell growth factor receptor c-KIT (52). Pazopanib treatment provides disease stability in patients with advanced breast cancer (53).

Estrogens are implicated in the development of the mammary gland, and are also known to be key stimulators of both the genesis and growth of mammary tumors (54). Therefore, one of the main strategies to fight breast cancer is neutralizing the stimulating actions of estrogens on the mammary gland (55). Several chemicals have been tested and commercialized, since they are selective inhibitors of the effects of estradiol on the breast. These include selective estrogen receptor modulators (SERMs), which are chemicals that directly bind to ER, thus inhibiting its actions by interfering with the binding of endogenous estrogens (56). Of these, tamoxifen (57,58) and a number of its derivatives (56) are the best known examples. Fulvestrant is an ER antagonist that has no agonist effect described and downregulates the protein levels of ERα (59). Fulvestrant is being currently administered to postmenopausal women with advanced breast cancer whose tumors are ER positive and have progressed upon receiving first-line endocrine therapy (60). Other compounds developed against estrogens are chemicals that prevent the production of steroids by downregulating the enzymes necessary for the conversion into estradiol from androgenic precursors. These drugs belong to the class known as selective estrogen enzyme modulators (SEEMs), which include both steroidal (such as formestane or exemestane) and non-steroidal (such as anastrozole and letrozole) compounds (61).

2. Melatonin and mammary cancer: In vitro and animal studies

Melatonin is an indollic hormone produced principally by the pineal gland. Melatonin is a ubiquitously distributed molecule with a variety of diverse functions (62). Melatonin employs a diverse set of mechanisms to regulate the physiology and molecular biology of cells (63). The majority of actions of melatonin are based on its ability to bind to melatonin membrane receptors, which are G-protein coupled receptors that trigger cellular signaling pathways (64). The pineal hormone also acts through orphan receptors or molecules such as calmodulin (65). Additionally, melatonin can detoxify free radicals and related oxygen derivatives via receptor-independent pathways (66).

Concerning tumorigenesis, numerous studies have been performed in animal models. Experimental approaches include increasing the activity of pinealocytes and administering exogenous melatonin, which cause a decrease in the number, incidence and development of chemically-induced mammary tumors (67). However, reduced levels of melatonin (for example, by removing the pineal gland) appear to stimulate breast cancer progression (68). Several reports established a lower cancer risk among totally blind women (69-71). By contrary, a moderate but significant increase in the risk of developing breast cancer among women who have been working for long periods in rotating night shifts (which implicates that they were exposed to light during the night, and consequently, the nocturnal melatonin production was inhibited) has also been documented (72).

The antiproliferative effects of melatonin on the breast cancer cell line MCF-7 have been studied for more than two decades (73). The data available suggest that the inhibitory action of melatonin on mammary cancer estrogen-positive cell lines is based on its ability to regulate either the synthesis of estrogens or estrogen signaling pathways (74). Thus, the pineal hormone is capable of downregulating both the expression and activity of the enzymes necessary for the synthesis of estrogens from androgenic precursors, therefore acting as a SEEM. In the MCF-7 cell line, melatonin at physiological concentrations exhibits anti-aromatase properties (75). The pineal hormone is able to reduce the activity of aromatase, the principal enzyme in estrogen biosynthesis. Melatonin inhibits aromatase under basal conditions or when the enzyme activity is stimulated by cyclic adenosine monophosphate (cAMP) or cortisol (76). When gene expression was evaluated and the CYP19 gene (coding for aromatase) was examined, it was observed that melatonin downregulated its expression at the transcriptional level (76). The major bloodstream circulating form of physiologically inactive estrogen is estrone sulfate, which acts as an estrogen reserve (77). The enzyme steroid sulfatase (STS) converts inactive estrogen sulfa tes into estrone and estradiol. Estrone can be further transformed into physiologically active estrogen by the action of 17β-hydroxysteroid dehydrogenase type 1 (17β-HSD1). Finally, the enzyme estrogen sulfotransferase (EST) sulfonates estrogens to form biologically inactive estrogen sulfates. Both enzymes, STS and EST, serve a role in the modulation of the in situ levels of estradiol in hormone-dependent tumors (78). Melatonin modulates the expression and activity of aromatase, STS, 17β-HSD1 and EST not only in tumor cells, but also in surrounding cells such as fibroblasts and endothelial cells (74,79-82).

Melatonin can also counteract the different actions of estrogens, thus functioning as a naturally occurring SERM (83). The mechanisms implicated in the antiestrogenic effects of melatonin are yet being elucidated. Unlike other antiestrogens molecules such as tamoxifen, melatonin does not directly bind to ER (84). In estrogen-positive breast cancer cells, melatonin decrease the expression of ERs (85) and impairs the estrogen-mediated transcriptional activation of genes through destabilization of the estradiol-ER complex, preventing its binding to DNA in both estrogen response element (ERE)- and activator protein 1-containing promoters (86). This effect appears to be mediated by calmodulin, since melatonin behaves as a calmodulin antagonist. The pineal hormone promotes structural changes in the calmodulin-ERα protein complex, thus facilitating its binding to an ERE (87,88). Remarkably, melatonin does not alter the recruitment of co-activators triggered by ERα, suggesting that melatonin mechanisms of action diverge from those of other anti-estrogen chemicals used in breast cancer treatment (88).

It is important to mention that only ERα, but not ERβ, binds to calmodulin (84). The substitution for glycine of two lysine
residues located at positions 302 and 303 of the hinge domain of ERα generated a mutant version of ERα that was incapable of binding to calmodulin, and that therefore turned into a melatonin non-regulated receptor (88). The effects of melatonin may also be explained in terms of binding to its specific membrane receptors, such as melatonin receptor type 1 (MT1), resulting in an interplay with the ER signaling pathway (89). MT1 receptors are present in normal human breast tissues and in tumor tissues (90). Melatonin and estradiol signaling pathways converge and they have opposite effects over cAMP intracellular concentrations. In breast cancer cells, estrogens trigger adenylate cyclase activation, which results in increased cAMP cytoplasmic levels in a classical short-time second-messenger mechanism that is independent of transcription (91). The increase in cAMP levels cooperates with long-time genomic effects of estradiol, thus enhancing ER-mediated transcriptional activation (91). By contrary, melatonin, through its specific binding to its membrane receptor MT1, inactivates adenylate cyclase, resulting in decreased cAMP levels (92).

The fact that only those human mammary cancer cell lines that are ER+ are sensitive to the antimitotic actions of melatonin supports the theory that the antitumor effects of this indolamine occur through its actions on breast cancer cells' estrogen-responsive pathways (93). Melatonin is able to block, under different culture conditions, the mitogenic effects of estradiol (94). The antiproliferative effect of this indolamine could be explained through the modification of the levels of estrogen-modulated proteins, several growth factors and proto-oncogenes such as cMYC, transforming growth factor (TGF)α, Trefoil factor 1, also known as pS2, progesterone receptor (PGR), AP1 transcription factor subunit c-fos and TGFβ in human estrogen-positive breast cancer cell lines (94,95). Estradiol enhances cell proliferation and provokes cell cycle progression (96). The inhibitory effect of melatonin (as occurs for tamoxifen) on cell proliferation is cell-cycle specific, causing the presence of melatonin a delay in the G1-S transition (97). Changes in cell-cycle timing typically implicate modifications in various key proteins that regulate the process. The inhibitory action of the pineal hormone on cell cycle progression can be interpreted through its effects on the expression of certain proteins controlling the G1-S cell cycle transition. Thus, several studies have demonstrated that melatonin increases the expression of p53 and p21Waf1 in experiments performed in vitro (98,99). The upregulation of these proteins may be a crucial mechanism explaining how melatonin impedes the progression through the cell cycle at the G1-S transition. The accumulation of cells in G1 forces them to enter G2p, causing the cancer cells to undergo a higher differentiation, since G2p is characterized to be a quiescent state (100). This suggest that the anti-estrogenic, oncostatic and antiproliferative effects of melatonin on human mammary cancer estrogen-positive cell lines may be explained, at least in part, by the ability of the pineal hormone to inhibit cell proliferation at the same time that it enhances cell differentiation. Furthermore, several studies have demonstrated that melatonin inhibits human telomerase reverse transcriptase, which is the rate-limiting factor conditioning the telomerase activity in breast cancer cells (101,102).

Another effect of melatonin in MCF-7 cells is its ability to reduce their invasiveness (103,104). Melatonin treatment decreases the attachment of the cells to the basement membrane (105). The pineal hormone reduces the chemotactic response of MCF-7 cells (105). Melatonin also blocks the cell migration and invasion that occurs in response to estradiol (105). Cancer cells' motility and invasion are known for being adhesion-dependent mechanisms that require the expression of cell-surface molecules necessary for adhesion (106). In tumor progression, downregulation or loss of expression of several of these surface-adhesion proteins frequently happens, which leads to the loss of cell-cell recognition and the acquisition of an invasive phenotype by the tumor cells; these events correlate with poor cell differentiation (107). All the factors mentioned above are associated with poor prognosis in cancer progression (108).

Importantly, among the melatonin antitumor actions, it has been reported that melatonin induces the expression of both β2-integrin and E-cadherin, two main proteins essential for cell-cell and cell-matrix interactions; thus, melatonin changes estrogen-responsive tumor cells into a less invasive phenotype by inducing their differentiation (105).

Finally, melatonin exerts its modulatory effect in the tumor microenvironment by controlling the production and secretion of several cytokines. These cytokines are produced by breast cancer cells and regulate the differentiation of the fibroblasts located in close proximity to malignant epithelial cells. Additionally, it has been demonstrated that cytokines produced by malignant cells stimulate the aromatase expression and activity in these fibroblasts (109,110) and in proximal endothelial cells (111). VEGF is a growth factor that serves an essential role in angiogenesis. VEGF is produced and secreted by malignant epithelial cells and recognizes VEGFRs located in the cell surface of endothelial cells. The binding of VEGF to its receptor triggers a cascade of intracellular events that stimulate endothelial cells to undergo proliferation and migration (112,113). Therefore, pharmacological agents able to inhibit the production of this pro-angiogenic factor are of great interest, and could serve an essential role in impairing both tumor angiogenesis and tumor growth. Melatonin may be one molecule to consider, since it can regulate the paracrine mechanisms connecting tumor epithelial cells and the surrounding endothelial cells. One of the main actions of the pineal hormone is that melatonin treatment results in the downregulation of VEGF expression in estrogen-responsive breast cancer cells. As a result, the VEGF levels available for receptors expressed in endothelial cells are lower, and therefore, the number of cells producing estrogens in the proximity of the malignant cells is reduced. Reduced estrogen levels and a lower capability of formation of new vessels as a result of the presence of melatonin will diminish the tumor ability to spread and grow (114).

In summary, a unique molecule, melatonin, has anti-estrogenic properties: It selectively counterbalances the actions of estrogens in both normal and tumor breast tissues, and provides a novel strategy to reduce the local biosynthesis of estrogens from androgens (which in turn, is one of the principal objectives of antitumor pharmacological therapy) (115). These cumulative actions of the pineal hormone point to its potential application as an anticancer molecule in both the prevention and treatment of estrogen-positive tumors, since, as it has been pointed above, this molecule acts at different levels by
interfering with estradiol-dependent signaling pathways, both in tumor cells and in the surrounding endothelial cells and fibroblasts (116).

3. Melatonin and cancer: Clinical trials

As aforementioned, numerous experiments performed in vitro (breast cancer cell lines such as MCF-7) and in vivo (animal models) have well established the oncostatic properties of melatonin (117). Since melatonin plasma levels are diminished in estrogen-dependent breast cancer patients, various clinical trials have been performed to evaluate the potential beneficial effects of melatonin in human neoplasms. Following the pioneer clinical work of Lissoni et al (118), who evaluated the effect of melatonin in cancer patients bearing untreated advanced solid tumors, multiple studies have been performed and published to date (119). However, the value of melatonin as an adjuvant agent in cancer treatment is not totally clear, and consensus about positive melatonin actions appears to be difficult to achieve. A number of studies point to melatonin as a treatment with no beneficial effects. In cachectic patients with gastrointestinal or advanced lung cancer, including a dose of melatonin at night did not improve parameters such as appetite, weight or quality of life of patients, in comparison with patients who received a placebo (120). In patients with brain metastases, high doses of melatonin did not produce any beneficial effect (121). By contrast, there are a large number of reports supporting the potential benefits of melatonin if included in chemotherapy protocols. In breast, lung and gastrointestinal cancer patients, melatonin protected against thrombocytopenia, and stomatitis, asthenia and neuropathy were less recurrent in the group of melatonin-treated patients (122). It has also been reported that melatonin offers certain protection to hematopoietic progenitors from the toxic actions of anticancer chemotherapeutic chemicals; thus, melatonin has been reported to attenuate the damage to precursor blood cells caused by both radiotherapy and chemotherapy treatments (123). It has also been suggested that melatonin may protect patients against side effects such as asthenia, cardiotoxicity and neurotoxicity caused by chemotherapy (124). Additionally, the pineal hormone increases the 1-year survival and tumor regression rates in cancer patients with metastatic solid tumors with poor clinical status (125). In metastatic non-small cell lung cancer patients treated either with just chemotherapeutic agents or with chemotherapy plus melatonin, both the overall tumor regression rate and the 5-year survival rate of patients concomitantly receiving melatonin were significantly higher as compared with those receiving only chemotherapeutic agents. It appears that chemotherapy was better tolerated in patients who also received the pineal hormone (126). The study points to melatonin as an adjuvant drug capable of improving the effectiveness of chemotherapy in terms of both quality of life and survival of patients (126). A recent report concludes that melatonin combined with chemotherapy did not significantly improve survival or ameliorated various adverse side effects in patients with non-small cell lung advanced cancer, although certain improvement in the quality of life of these patients was observed (127). A systematic review comprising data from 21 clinical trials (all the patients enrolled in the studies were bearing solid tumors), in which the effect of melatonin concomitantly added in conjunction with chemotherapy or radiotherapy was evaluated, and supportive care, partial response, complete response, 1-year survival and chemotherapy-associated toxicities were assessed (128), concluded that melatonin may serve a beneficial role in cancer patients who are treated with chemotherapy. Patients who received melatonin experienced substantial improvements, particularly in terms of tumor remission and 1-year survival rates and melatonin also ameliorated the side effects of chemotherapy (127,128). Furthermore, another review summarizing the data from eight eligible randomized controlled trials (n=761) obtained similar conclusions (129).

4. Can melatonin enhance the beneficial and protect against the deleterious effects of chemotherapy?

As aforementioned, it is well documented that melatonin diminishes the incidence of chemically induced cancers and is able to slow down the growth of certain hormone-responsive cancers (67,117). The antitumor actions of this indolamine have been described in breast cancer, both in in vivo animal experiments (in 7,12-dimethylbenz[a]anthracene chemically induced mammary tumors in rodents) and in in vitro assays (in estrogen-positive human mammary cancer cell lines) (64). Furthermore, there are numerous reports endorsing the beneficial use of melatonin during chemotherapy in clinical trials (118,122-124). Therefore, the ultimate goal of the present review is to provide a compilation on the current knowledge concerning the interplay of melatonin and chemotherapy agents at the molecular level. The PubMed database (www.ncbi.nlm.nih.gov/pubmed) was interrogated for citations of 84 genes known to be commonly involved in the dysregulation of several normal processes during breast carcinogenesis, which are also present in breast cancer cell lines. The list includes signal transduction genes and other genes involved in usually altered pathways, including cellular adhesion, angiogenesis, proteolytic activities, cell cycle progression, cell cycle control and apoptosis (Table I). Research into carcinogenic mechanisms (130) has identified during the last decades numerous functional alterations due to somatic mutations, gene expression alterations and altered post-translational protein modifications (131). Therefore, the database search was performed including as keywords i) the name of each gene, ii) breast cancer, iii) melatonin, and iv) chemotherapy.

For the majority of genes reviewed, there were thousands of citations in the literature when the search included the name of each gene and breast cancer as key words. When melatonin was included in the selection criteria, the number of articles was markedly reduced; thus, 53 out of the 84 genes surveyed did not have a single report associated under these criteria. Searching for ER together with melatonin and breast cancer led to ~100 articles, whereas p53 and PGR searches produced 15 and 12 reports respectively, and the rest of the 28 genes evaluated produced 1-9 articles.

The number of reports was further reduced if chemotherapy was also included in the selection criteria. Only 25 of the 84 genes assessed are included in the list (Table I), indicating that, for 59 out of 84 genes, there is not a single study published including the gene name, melatonin, breast cancer and chemotherapy as keywords.
In Table II, a search was performed including as keywords melatonin (20,724 articles) melatonin and breast cancer (524 articles), and 28 drugs commonly used as chemotherapy in breast cancer treatment (6,9). Apart from tamoxifen (23 articles fulfilled the criteria ‘melatonin, breast cancer and tamoxifen’), the results show that there is limited information at the molecular level concerning the implication of co-treatment with the pineal indolamine and chemotherapy agents in breast cancer targeted therapy, and no report was available for 13 of the 28 drugs searched. In conclusion, despite the fact that during decades numerous articles have reported experimental data from in vitro and in vivo experiments showing the oncostatic actions of melatonin, the present review demonstrated that there are limited reports studying the effects of melatonin and chemotherapy agents in combination in cancer treatments, and particularly, in breast cancer. Similarly, little is known about the role of melatonin regarding the expression and functionality of the genes reported to be altered in cancer, particularly in breast cancer. There is little information available nowadays about gene expression profiles in all types of cancer, and particularly in estrogen-responsive breast tumors. There is also limited information available on how the gene expression profile may be altered by treatment with different chemotherapeutic compounds and, remarkably, whether or not melatonin has protective effects when administered together with chemotherapeutic agents. The majority of information available about melatonin, cancer, chemotherapy, and altered gene expression and function has been published (132,133) in the last few years (Fig. 1). This indicates that nowadays there is a growing field of research about this topic. Several of the most relevant findings recently reported with regard to the modulatory role of melatonin in cancer at the molecular level include the circadian interruption of melatonin production by exposure to light at night, which results in the development of resistance to tamoxifen treatment in breast cancer patients (134). In this context, it has also been described how nocturnal disruption of melatonin plasma levels originates a complete loss of tumor response to the chemotherapeutic agent doxorubicin (135). The resistance of cancer cells to chemotherapy treatments such as doxorubicin usually implicates an upregulation of P-glycoprotein, which is responsible for drug eflux from cells (136). In this context, there is a report describing that melatonin treatment increases doxorubicin intracellular concentrations in cancer cells, suggesting that melatonin may inhibit P-glycoprotein (137). In breast cancer xenografts implanted in animal models (nude mice), treatment with the pineal hormone stopped the tumor progression by reducing tumor size and cell proliferation (Ki-67), as well as by inhibiting angiogenesis (138). Melatonin treatment results in increased expression of Bcl-2-like protein 11 (Bim) parallel to lower levels of cyclooxygenase (COX)-2, which in turns potentiate tunicamycin-induced apoptosis in mammary cancer cells (139). Regulation of the COX-2, Akt, p300 and apoptotic protease activating factor-1 signaling pathways by melatonin inhibits cell proliferation and triggers apoptosis in breast cancer cells in in vitro models (140). Melatonin also regulates mouse double minute 2 homolog (MDM2), since this indolamine strongly represses MDM2 gene expression and inhibits MDM2 translocation into the nucleus of the cells (141). This can be explained because melatonin stimulates ribosomal protein L11 and inhibits the phosphorylation of MDM2 by Akt-phosphatidylinositol-4,5-bisphosphate 3-kinase (141). Melatonin downregulates sirtuin, which is a specific inhibitor of p300, and upregulates p300 and murine double minute X (MDMX) (141). As a consequence, cells exposed to melatonin exhibit significantly increased levels of p53 and of its acetylated form (141). Finally, there is a significant increase in p21 levels in melatonin-treated tumor cells (141). It has been reported that melatonin sensitizes non-small cell lung cancer cells harboring a mutated form of EGFR to gefitinib (a tyrosine kinase inhibitor) (142). In combination with cisplatin, melatonin enhances the cytotoxic effects of this chemotherapeutic agent and promotes the entry into apoptosis of lung cancer cells (143) and cervical cancer-derived HeLa cells (144). Consistently with these findings, co-treatment with melatonin and each of the following three chemotherapy agents: Cisplatin, 5-fluorouracil and doxorubicin, resulted in an enhancement of cytotoxicity and apoptosis triggered by chemotherapy in the cell line AR42J, which is derived from rat pancreatic tumors (145). There is only one report addressing the effect of melatonin combined with the purine nucleoside antimetabolite clofarabine, describing that melatonin use as co-treatment led to an enhanced cytotoxic effect of clofarabine in leukemic cell lines, which was associated with higher levels of acetylation (146). In ER+ breast cancer rat models treated with Adriamycin®, melatonin co-treatment results in lighter tumor weights, increased tumor cell apoptosis, higher expression of E-cadherin and higher survival rate (147). In combination with the nucleoside analogue gemcitabine, recent reports demonstrate that melatonin inhibits both proliferation and invasion of pancreatic ductal adenocarcinoma cells through nuclear factor-kB inhibition (148). Melatonin supports the effects of doxorubicin by activating transient receptor potential vaniloid 1 and apoptosis, thus inducing MCF-7 cell death (149). In a model of ovarian carcinoma, melatonin therapy promotes apoptosis along with the upregulation of p53, B-cell lymphoma (Bcl)-2-associated X protein (Bax) and cleaved caspase-3, suggesting that melatonin triggers apoptosis in ovarian cancer cells (150). In a gastric cancer cell line (AGS), p38, e-Jun N-terminal kinase and extracellular signal-regulated kinase were activated by melatonin, which also significantly increased caspase-3 activity, increased the expression of the pro-apoptotic gene Bax and decreased the expression of the anti-apoptotic gene Bcl-2 (151). Additionally, melatonin is able to strengthen the antitumor effects of cisplatin with low systemic toxicity (143).

5. Conclusions

Melatonin is a notable molecule to be considered in cancer treatment. A growing amount of evidence in the last few years has suggested that melatonin behaves as an oncostatic agent in a variety of cancer types in general, an in particularly, in hormone-dependent breast cancer, as documented from numerous studies performed either in animal models in vivo or in cell lines derived from human breast cancer in vitro (64,67,81). Clinical trials suggest that melatonin can have protective effects when administered along with chemotherapy in patients suffering from advanced solid tumors (129). The mechanisms underlying the oncostatic actions of the pineal hormone in
On the contrary, the limited knowledge about the interplay of melatonin and chemotherapy on molecular aspects such as gene expression profiles and gene post-translational modifications, which must be further addressed in the future. 6. Melatonin and cancer: What next? Melatonin is a pleiotropic molecule that exerts numerous physiological functions and serves important roles in different processes, including circadian rhythm, sleep and reproduction (152). A particular characteristic of the pineal hormone is the diversity of molecular mechanisms that act to regulate the above physiological processes. Melatonin reduces cell proliferation and growth of estrogen-positive breast tumors by interfering with estrogen signaling pathways. Data obtained from experiments performed in breast cancer cell lines (93,94) and animal models (81,116) have provided evidence that melatonin diminishes the incidence of mammary tumors and limits their growth in vivo, and inhibits the proliferation of human breast cancer cell lines and interferes with their metastatic behaviour in vitro (81,97). There are several proposed theories to explain the mechanisms by which melatonin reduces the growth and

| Gene name | Gene name AND breast cancer | Gene name AND melatonin | Gene name AND breast cancer AND melatonin | Gene name AND breast cancer AND melatonin AND chemotherapy |
|-----------|-----------------------------|-------------------------|-------------------------------------------|------------------------------------------------------------|
| ER        | 68,885                      | 31,082                  | 189                                       | 96                                                         | 31                                                         | 52                                                         |
| p53       | 78,342                      | 7,698                   | 80                                        | 15                                                        | 7                                                          | 12                                                         |
| p21       | 31,647                      | 2,174                   | 35                                        | 9                                                         | 9                                                          | 5                                                          |
| VEGF      | 57,030                      | 2,737                   | 64                                        | 12                                                        | 3                                                          | 11                                                         |
| PGR       | 33,762                      | 13,582                  | 73                                        | 96                                                        | 3                                                          | 5                                                          |
| TGFβ1     | 22,043                      | 561                     | 25                                        | 4                                                         | 3                                                          | 4                                                          |
| MYC       | 29,932                      | 1,882                   | 17                                        | 5                                                         | 3                                                          | 3                                                          |
| CDH1      | 23,363                      | 2,200                   | 13                                        | 5                                                         | 3                                                          | 3                                                          |
| IL6       | 96,419                      | 1,106                   | 162                                       | 3                                                         | 2                                                          | 10                                                         |
| GSP1      | 35,897                      | 901                     | 91                                        | 2                                                         | 2                                                          | 7                                                          |
| c-JUN     | 102,575                     | 2,952                   | 106                                       | 6                                                         | 2                                                          | 6                                                          |
| RARβ      | 11,545                      | 651                     | 76                                        | 9                                                         | 2                                                          | 5                                                          |
| CCND1     | 16,332                      | 2,205                   | 16                                        | 6                                                         | 2                                                          | 2                                                          |
| AR        | 22,741                      | 1,690                   | 64                                        | 3                                                         | 1                                                          | 11                                                         |
| AKT       | 54,018                      | 3,992                   | 92                                        | 6                                                         | 1                                                          | 10                                                         |
| Ki-67     | 20,827                      | 3,129                   | 18                                        | 5                                                         | 1                                                          | 1                                                          |
| ERK1      | 26,166                      | 1,153                   | 61                                        | 4                                                         | 1                                                          | 3                                                          |
| ERBB2     | 22,268                      | 14,282                  | 72                                        | 3                                                         | 1                                                          | 3                                                          |
| EGFR      | 35,864                      | 3,550                   | 7                                        | 3                                                         | 1                                                          | 3                                                          |
| IGFBP3    | 5,003                       | 432                     | 6                                         | 2                                                         | 1                                                          | 2                                                          |
| EGF       | 29,673                      | 2,215                   | 24                                        | 6                                                         | 1                                                          | 2                                                          |
| Rb        | 19,405                      | 1,017                   | 16                                        | 1                                                         | 1                                                          | 1                                                          |
| IGFl      | 2,700                       | 160                     | 15                                        | 1                                                         | 1                                                          | 1                                                          |
| CDK2      | 6,054                       | 491                     | 5                                         | 1                                                         | 1                                                          | 1                                                          |
| p73       | 2,089                       | 113                     | 1                                         | 1                                                         | 1                                                          | 1                                                          |

* A search in PubMed database was performed for 84 genes known to be altered in breast cancer. The numbers refer to the citation found when the keywords used were: i) ‘Gene name’; ii) ‘gene name’ AND ‘breast cancer’; iii) ‘gene name’ AND ‘melatonin’; iv) ‘gene name’ AND ‘breast cancer’ AND ‘melatonin’; v) ‘gene name’ AND ‘breast cancer’ AND ‘melatonin’ AND ‘chemotherapy’; and vi) ‘gene name’ AND ‘cancer’ AND ‘melatonin’ AND ‘chemotherapy’. In the table, only the 25 genes that appear at least in one publication with the criteria ‘gene name’ AND ‘breast cancer’ AND ‘melatonin’ AND ‘chemotherapy’ are shown. The genes that have appear in any publication under these criteria of search are gelatinase A, PTGS2, Bad, Bcl-2, BIRC5, gelatinase B, CTNNB1, APC, ASC, ATM, ABCB1, ABCG2, BRCA1, TFF3, cathepsin, μ-PA, SRC, PAI-1, serpine 1, JNK1, IGF1R, CDKN2A, ADAM23, PTEN, NOTCH1, THBS1, ID1, keratin 5, GATA3, ERK2, CCNE1, XBP1, NR3C1, BRCA2, MUC1, MLH1, keratin 19, NME1, TWIST1, FOXA1, RASFF1, HIC1, SFN, MGMT, CCND2, cystatin, GRB7, keratin8, GLI1, keratin18, SFRP1, SNAI2, p57, cyclin A1, CDH13, CSF1, SLIT2, SLC39A6 and PRM2.
development of tumors: i) Through an indirect mechanism, by downregulating the synthesis of estrogens via downregulation of the hypothalamic-pituitary-reproductive axis; ii) through a direct mechanism, by interfering with the activation of estradiol receptors at the cancer cell level, thus behaving as a SERM; and iii) melatonin can regulate the enzymes necessary for the synthesis of estrogens in other tissues, therefore behaving as a SEEM (79). It has been recently demonstrated that melatonin regulates the paracrine communication that occurs between malignant epithelial cancer cells, the surrounding adipose tissue (fibroblasts and adipocytes) and endothelial cells, mainly through the downregulation of the levels of growth factors and cytokines released by breast tumor cells (116). Thus, the effects of melatonin also include anti-angiogenic actions.

Table II. Number of publications identified in the MEDLINE (https://www.ncbi.nlm.nih.gov/pubmed) journal citation database (accessed November 2015), when using as search term each of the 28 molecules currently used in breast cancer research and/or treatment, alone (second column), including melatonin (third column) or including melatonin plus breast cancer (fourth column) as searching criteria.

| Drug name       | Drug alone | Melatonin AND breast cancer |
|-----------------|------------|-----------------------------|
| Vincristine     | 27,607     | 10                          |
| Eribulin        | 252        | 0                           |
| Paclitaxel      | 27,688     | 8                           |
| Docetaxel       | 11,691     | 3                           |
| Epirubicin      | 6,131      | 5                           |
| Lapanib         | 1,807      | 1                           |
| Trastuzumab     | 7,382      | 2                           |
| Pazopanib       | 825        | 0                           |
| Bevacizumab     | 11,697     | 6                           |
| Fulvestrant     | 2,278      | 5                           |
| Anastrozole     | 1,761      | 1                           |
| Irosustat       | 46         | 0                           |
| Tanespinic      | 665        | 0                           |
| Cisplatin       | 59,192     | 50                          |
| Gemcitabine     | 12,035     | 6                           |
| Pivatavastin    | 658        | 3                           |
| Pravastatin     | 4,318      | 1                           |
| Vinblastine     | 1,582      | 14                          |
| Cyclophos-phamide | 64,488    | 63                          |
| Methotrexate    | 45,411     | 20                          |
| Fluorouracil    | 48,858     | 18                          |
| Adryamicin      | 60,505     | 74                          |
| Vinorelbine     | 3,472      | 2                           |
| Mitomycin       | 18,202     | 6                           |
| Capecitabine    | 4,746      | 1                           |
| Mitoxantrone    | 5,538      | 5                           |
| Carboplatin     | 13,873     | 4                           |
| Tamoxifen       | 25,107     | 44                          |

In summary, previous studies point to melatonin as a molecule that has a great potential to be useful as an anticancer chemical without producing adverse effects (128). Therefore, melatonin should be considered for both the prevention and therapy of estrogen-positive mammary tumors. There are, in our opinion, numerous noteworthy possibilities for future clinical applications of melatonin in several types of cancer, including breast cancer.

Recently, the inhibitory effects of melatonin have been described not only for estrogen-dependent breast tumors, but also for numerous different cancers, including gastric cancer, ovarian carcinoma, pancreatic ductal carcinoma, leukemic cell lines, cervical cancer and non-small lung carcinoma cells (25,137-139,142,143,153). The majority of the results are positive, and melatonin has been described as an inhibitor of tumor growth under both in vitro and in vivo experimental conditions (64,81,97). The results arisen in the past few years also suggest that melatonin, either alone or along with chemotherapy in cancer patients diagnosed with advanced solid tumors, helps to improve the outcomes of cancer regression and life expectancy of the patients (122,128). Additionally, chemotherapies are typically better tolerated by patients who are simultaneously treated with melatonin (125). Following the pioneer clinical study of melatonin potential positive effects in untreatable advanced cancer patients performed by Lissoni et al (118) several studies have been published (124-129). The main limitation is the requirement of further studies, including additional randomized double-blind controlled trials with much larger sample sizes and implicating several international hospital centres, since the data available nowadays derive from clinical trials including only a few hundred patients (128). Furthermore, it must be considered that not all the studies performed to date point to melatonin as a molecule that improves life expectancy and ameliorates the adverse effects of chemotherapy (120,121). Thus, in patients with advanced lung or gastrointestinal cancer, melatonin did not exhibit any beneficial effect, and as consequence, the value of melatonin as an adjuvant in the treatment of cancer remains unclear from these data (120).

One of the main objectives of the present review was to summarise the current knowledge regarding the interplay of melatonin and chemotherapy. Since the majority of the potentially beneficial effects of melatonin have been described in
estrogen-responsive breast cancer, the present study searched information published on 84 genes known to be dysregulated during breast carcinogenesis (corresponding to the genes included in The Human Breast Cancer RT² Profiler PCR array; Qiagen GmbH, Hilden, Germany). These genes encode proteins implicated in signal transduction, angiogenesis, proteolysis, cell cycle and apoptosis (154).

The present study also reviewed recently published articles associating melatonin with chemotherapeutic agents (135-137,142-149). The results indicate that, for 53 out of the 84 genes evaluated, there are no current data available regarding the effect of melatonin alone or in combination with chemotherapy, either in in vivo or in in vitro studies. Our findings also show that, apart from tamoxifen, there is limited information from research performed at the molecular level addressing the potential benefits of co-treatment of melatonin with chemotherapeutic agents (155).

In summary, in our opinion, melatonin is an endogenous produced hormone with a high potential of being included as an effective anticancer molecule in the prevention and treatment of, not only hormone-dependent cancers, but also, other types of cancer, since its inhibitory effects have been demonstrated in gastric, lung, pancreatic and hematopoietic cancers (145,148,151,153). However, in the next years, additional research must be conducted to clarify if melatonin administration in combination with chemotherapeutic agents may constitute a novel anticancer treatment. In particular, future research concerning the role of melatonin as a non-toxic and low-cost drug to be considered in breast and other types of tumors must be conducted, particularly at the molecular level. Systematic screenings addressing the effects of chemotherapy on genes known to be altered in different types of cancer, and on how melatonin can modulate the expression and activity of those genes, either when acting alone or in combination with chemotherapy, should be performed. Once larger clinical trials and additional molecular studies (including gene expression profiles, post-translational modifications and individual gene tests) have been conducted, it may be reasonable to recommend melatonin as a potential drug to be considered in the treatment of breast cancer.

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