Modulation of cancer endocrine therapy by melatonin: a phase II study of tamoxifen plus melatonin in metastatic breast cancer patients progressing under tamoxifen alone

P Lissoni¹, S Barni¹, S Merengalli¹, V Fossati¹, M Cazzaniga¹, D Esposti² and G Tancini¹

¹Divistione di Radioterapia Oncologica, San Gerardo Hospital, 20052 Monza, Milan, Italy; ²Istituto di Fisiologia Umana II, University of Milan, Milan, Italy.

Summary
Recent observations have shown that the pineal hormone melatonin (MLT) may modulate oestrogen receptor (ER) expression and inhibit breast cancer cell growth. On this basis, we have evaluated the biological and clinical effects of a concomitant MLT therapy regimen in metastatic breast cancer patients who had progressed in response to tamoxifen (TMX) alone. The study included 14 patients with metastasis who did not respond (n = 3) to therapy with TMX alone or progressed after initial stable disease (SD) (n = 11). MLT was given orally at 20 mg day⁻¹ in the evening, every day starting 7 days before TMX, which was given orally at 20 mg day⁻¹ at noon. A partial response was achieved in 4/14 (28.5%) patients (median duration 8 months). The treatment was well tolerated in all cases, and no MLT-induced enhancement of TMX toxicity was seen; on the contrary, most patients experienced a relief of anxiety. Mean serum levels of insulin-like growth factor 1 (IGF-1), which is a growth factor for breast cancer, significantly decreased on therapy, and this decline was significantly higher in responders than in patients with SD or progression. This pilot phase II study would suggest that the concomitant administration of the pineal hormone MLT may induce objective tumour regressions in metastatic breast cancer patients refractory to TMX alone.

Keywords: breast cancer; insulin-like growth factor 1; melatonin; pineal gland; tamoxifen

Several experimental studies have demonstrated that the anti-estrogenic action is only one of the great variety of mechanisms responsible for the antineoplastic properties of tamoxifen (TMX) in breast cancer. Oestrogens themselves would stimulate breast cancer growth by determining the paracrine release of growth factors, such as insulin-like growth factor 1 (IGF-1) (Furlanetto and Decarlo, 1984; Duclos et al., 1989). Recent investigations have shown that breast cancer cells may express IGF-1 receptors (Bonneteue et al., 1990), and their presence seems to have a positive prognostic significance. Moreover, it has been demonstrated that TMX therapy reduces IGF-1 blood levels in breast cancer patients, and this event may contribute to the therapeutic effect of TMX itself (Pollak et al., 1990). Another growth factor for breast cancer is pro lactin (PRL). High levels of PRL have been proven to be associated with a poor prognosis in metastatic breast cancer patients (Bhatavdekar et al., 1990), while the expression of PRL receptor on breast cancer cells constitutes a good prognostic factor (Bonneteue and Peryt, 1989). However, the role of PRL and PRL receptor in breast cancer is still controversial.

Recent advances in endocrinology have documented that the endocrine secrations are under a modulatory control exerted by the pineal gland (Regelson and Pierpaoli, 1987), mainly through the circadian release of its most investigated hormone melatonin (MLT). MLT has been proven to stimulate oestrogen receptor (ER) expression on breast cancer cells (Danforth et al., 1983) and to reverse some malignant phenotypic characteristics of cancer cells (Hill and Blask, 1988), perhaps by inhibiting oncogene expression, which would be responsible for the malignant characteristics themselves. Finally, MLT appears to inhibit the secretion of IGF-1 and PRL (Regelson and Pierpaoli, 1987), both involved in the stimulation of breast cancer cell proliferation. Other endocrine secretions are influenced by MLT, particularly growth hormone and cortisol (Regelson and Pierpaoli, 1987). Therefore, several effects exerted by MLT, consisting in stimulation of ER expression and inhibition of IGF-1 production and PRL release, would suggest that the pineal hormone may potentiate TMX therapeutic efficacy. In fact, preliminary experimental studies have demonstrated that MLT may amplify in vitro TMX-induced inhibition of breast cancer cell growth (Hill et al., 1992). In addition, MLT has been proven to have a direct cytostatic action against some breast cancer cell lines (Hill and Blask, 1988). The present phase II study was performed to investigate the biological and therapeutic effects of a concomitant administration of MLT in metastatic breast cancer patients who progressed under therapy with TMX alone.

Patients and methods

The study included 14 consecutive women with metastatic breast cancer who did not respond to TMX therapy or progressed after initial disease stabilisation. Dominant metastasis sites were as follows: soft tissues, 3; bone, 4; visceral locations, 7 (lung, 3; pleural space, 2; liver, 2). ER estimation was made on the primary tumour by the dextran-coated charcoal method; ER was considered as positive when values were greater than 10 fmol mg⁻¹ protein. ER was positive in eight and negative in the other six cases. Patients with negative ER had been also treated with TMX since they were unable to tolerate conventional polychemotherapy because of age, low performance status (PS) and/or important medical illnesses other than cancer. The previous therapy with TMX alone resulted in stabilisation of disease in 11 patients (median duration 8 months, range 3–16), whereas the other three patients rapidly progressed on treatment. Eligibility criteria included histologically proven breast cancer, metastatic disease, measurable lesions, progression on TMX therapy alone and inability to tolerate conventional polychemotherapies because of age and/or concomitant medical illnesses. The experimental protocol was explained to each patient, and informed consent was obtained. TMX was given orally at a daily dose of 20 mg at 12.00 a.m., every day until progression. MLT, which was supplied by Medea Research (Milan, Italy), was administered orally at a daily dose of 20 mg in the evening every day of TMX therapy starting 7 days before TMX, as an induction phase. The dose of MLT was established from our previous studies (Lissoni et
al., 1989, 1991). Moreover, MLT was given during the dark period of the day because of its greater biological efficacy in this period of the day (Regelson and Pierpaoli, 1987). All patients had been off TMX for at least 1 month (median period 2 months, range 1-3) before starting MLT plus TMX therapy.

Radiological examinations were made before the onset of treatment, after each month of therapy for the first 3 months, then every 3 months. Clinical response and toxicity were evaluated according to UICC and WHO criteria respectively. All responses were confirmed by computerised tomographic (CT) scan. Complete response (CR) was defined as a complete regression of all lesions for at least 1 month; partial response (PR) was considered as a reduction of at least 50% in the sum of the products of the longest perpendicular diameters for at least 1 month; stable disease (SD) was defined as no objective cancer regression or increase greater than 25%; progressive disease (PD) was an increase of at least 25% in measurable lesions or the appearance of new lesions. Patients were considered as evaluable when they were treated for at least 2 months. PS was evaluated according to Karnofsky's score.

Routine laboratory tests were repeated at weekly intervals for the first 3 months, then every month. Moreover, serum levels of IGF-1 and PRL were also measured before treatment and at 1 month intervals for the first 3 months. IGF-1 and PRL serum levels were measured in duplicate by the radioimmunoassay (RIA) method and commercially available kits. Intra-assay and inter-assay coefficients of variation were less than 3% and 5% respectively. Normal values obtained in our laboratory (95% confidence limits) for IGF-1 and PRL were less than 2.2 U ml⁻¹ and less than 20 ng ml⁻¹ respectively. Data were statistically analysed by the chi-square test, the Student's t-test and analysis of variance as appropriate.

Results

The characteristics of patients and their clinical response are reported in Table I. All patients were evaluable for response. No CR was achieved. PR was obtained in four patients (28.5%) (median duration: 8 months, range 3–9). The first two patients had single lung nodular metastasis; the third patient showed a cytologically positive pleural effusion and pleural infiltration documented by CT scan; the last patient had multiple skin metastases. No significant difference in tumour response rate was seen between patients with positive and negative ER (2/8 vs 2/6). Eight other patients had SD, whereas the remaining two patients had progression. All patients were followed-up for at least 1 year. Survival for longer than 1 year from the onset of treatment was observed in 10/14 patients.

No toxicity was found. On the contrary, most patients experienced a relief of anxiety; moreover, a relief of depressant symptoms occurred in 3 patients. Finally, two other patients with low PS, as evaluated according to Karnofsky's score, had a clear improvement in their PS and quality of life on treatment. The improvement in the quality of life was based on specific patient report.

Changes in mean serum levels of IGF-1 observed on study are illustrated in Figure 1. Mean concentrations of IGF-1 significantly decreased on treatment with respect to the values found before therapy. Moreover, minimum values (mean ± s.e.) of IGF-1 levels observed on therapy were significantly lower in patients who responded than in those with SD or progression (0.7 ± 0.3 vs 3.1 ± 0.6 U ml⁻¹), \( P < 0.05 \), whereas no significant difference was seen before therapy (3.9 ± 0.6 vs 4.7 ± 0.9). Mean PRL levels also significantly decreased on treatment with respect to the pretreatment ones (13 ± 2 vs 25 ± 3 ng ml⁻¹, \( P < 0.05 \)), even though no difference was observed in mean PRL decrease between responding patients and those with progression or SD (14 ± 5 vs 11 ± 4 ng ml⁻¹).

Discussion

This preliminary phase II study would suggest that the pineal hormone MLT may amplify the therapeutic efficacy of TMX in women with metastatic breast cancer and induce objective response.

### Table 1

| Patient no. | Age | Sites of metastases | ER | Previous response to TMX alone (+) | Clinical response | Time to progression (months) | Sites of response | Sites of progression | Survival (months) |
|-------------|-----|---------------------|----|----------------------------------|------------------|------------------------------|------------------|---------------------|------------------|
| 1           | 65  | Bone                | +  | SD (6)                           | SD               | 3                            | –                | –                   | Liver            |
| 2           | 63  | Pleura              | +  | SD (13)                         | SD               | 3                            | –                | –                   | Pleura           |
| 3           | 59  | Lung                | –  | SD (8)                          | PR               | 7                            | Lung             | Skin                | 17               |
| 4           | 67  | Bone                | –  | SD (8)                          | SD               | 4                            | –                | Bone                | 16               |
| 5           | 42  | Liver, bone         | +  | SD (3)                          | SD               | 3                            | –                | Bone                | 15               |
| 6           | 74  | Pleura              | +  | SD (9)                          | PR               | 8                            | Pleura           | Bone                | 15               |
| 7           | 80  | Lung                | +  | SD (16)                         | PR               | 6                            | Lung             | Lung                | 14               |
| 8           | 59  | Skin                | –  | SD (4)                          | PD               | –                            | –                | Skin                | 7                |
| 9           | 72  | Bone                | +  | SD (14)                         | SD               | 4                            | –                | Bone                | 14               |
| 10          | 76  | Liver, bone         | –  | SD (5)                          | SD               | 8                            | –                | Liver               | 13               |
| 11          | 38  | Lung, bone          | +  | PD                              | PD               | –                            | –                | Bone                | 13               |
| 12          | 74  | Skin                | –  | PD                              | PR               | 9                            | Bone             | Skin                | 14               |
| 13          | 58  | Skin                | –  | PD                              | PR               | 9                            | Skin             | Bone                | 13               |
| 14          | 72  | Bone                | +  | PD                              | SD               | 5                            | –                | Lung                | 11               |

ER, estrogen receptor, PR, partial response; SD, stable disease, PD, progressive disease; +, time to progression (months) under TMX alone.
tumour regressions in patients who have not responded to previous therapy with TMX alone irrespective of ER status. However, measurements of other prognostic variables, such as progesterone and MLT receptors, will have to be evaluated to better define possible predictive factors for MLT efficacy. Therefore, because of its complete lack of toxicity, the combination of TMX and MLT may constitute a new effective modality of therapy for metastatic breast cancer, particularly in patients unable to tolerate conventional chemotherapies. Moreover, the results of this study, by showing declines in blood levels of tumour growth factors IGF-1 and PRL, would suggest that MLT may amplify TMX activity by blocking the production of important growth factors for breast cancer. However, the IGF-1 decrease observed in this study may be due not only to MLT action, but also at least in part to TMX itself, since TMX has been proven to inhibit IGF-1 secretion (Pollak et al., 1990). In any case, the action of MLT on IGF-1 secretion could explain the potential efficacy of the pineal hormone in patients with negative ER states. Recently, MLT receptors have been documented on some cancer cell lines (Hill et al., 1992). Therefore, further studies, by investigating the expression of MLT receptors and by analysing their existence in relation to ER, PRL and IGF-1 receptors, will be required to predict the efficacy of this pineal hormone in breast cancer. Obviously, the small number of patients considered in this study does not allow us to draw definite conclusions about the possible use of MLT to modulate the efficacy of breast cancer endocrine therapy. However, the evidence of objective tumour regressions induced by concomitant MLT treatment in breast cancer patients who did not respond to a previous therapy with TMX alone would confirm the oncostatic properties of MLT. This study does not allow us to establish whether tumour regression is due to MLT alone or to its combination with TMX. Previous studies have shown that MLT may decrease oestrogen levels in breast cancer (Regelson and Pierpaoli, 1987), but it is generally unable to induce objective tumour regression as a single agent (Lissoni et al., 1989, 1991). In addition, the contribution of a TMX withdrawal effect cannot be excluded, even though it is generally unlikely in patients non-responsive to TMX alone. In conclusion, randomised studies with TMX alone vs MLT alone vs their combination will be required to better define the influence of the pineal hormone on TMX anti-tumour activity.

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