PATHOLOGICAL STUDIES ON BCG AND HYPERICIN MEDIATED PHOTODYNAMIC THERAPY (PDT) IN TUMOR TREATMENT IN MICE

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Abstract

The purpose of the current study is to evaluate the effectiveness of Hypericin Photodynamic therapy (PDT) alone or associated with BCG immunization as an oncotherapy in Swiss mice. For this purpose, 90 mature male swiss mice were chosen and equally divided into 5 comparable groups. The first, was a negative control group, the second was hypericin control group and the third was tumor control group. The forth and fifth groups were the treated groups with Hypericin – PDT and BCG/ Hypericin – PDT, respectively. They were subjected to the treating regimen as follows: a dose of 5 mg/kg body weight of an aqueous ethanolic Hypericin solution was injected by both intra-peritoneal (i.p.) and intratumoral (i.t.) injections, followed 2 hours later by exposure to non-coherent light (590 nm) for 5 minutes in each session. Mice received 5 sessions / week for 6 weeks. Both treatments resulted in pronounced degeneration and necrosis of the neoplastic cells with replacement of tumor masses by fibrous connective tissue. The curative rate was enhanced when Hypericin-PDT was accompanied by BCG as an immuno-stimulating agent. From the pathological pictures, it can be concluded that Hypericin-PDT is an effective treatment for superficial neoplasms. BCG administration, as a non-specific immuno-stimulant, enhances the therapeutic effect of Hypericin -PDT.

INTRODUCTION

Cancer is still a disease that causes a major threatening for human and animal life, in addition to the cost effectiveness in its treatment (Epstein, 1990).

There are various therapeutical approaches for treatment of tumors, the selection of certain approach depends mainly on the tumor location.

Photodynamic therapy (PDT) might be the treatment of choice for superficial tumors, and in other situations, it can be used as an adjuvant or palliative treatment (Lilge et. al., 1997).

Photodynamic therapy involves the relatively selective uptake of a photosensitizig drug and a subsequent irradiation with light of a suitable wave length corresponding to the peaks absorption of the drug (Henderson and Dougherty, 1992).
The useful photosensitizer should have a narrow absorption band, selective accumulation in tumor tissues, fast clearance from the normal tissues and low toxicity (Bonnett, 1994). Hypericin, is considered as a second generation of photosensitizers; it showed an increased photodynamic efficiency in the treatment of tumors and reduced phototoxicity (Ochsner, 1997).

The aim of the present study is to evaluate the effectiveness of hypericin photodynamic therapy (PDT) alone or associated with BCG immunization as an oncotherapy in mice.

**MATERIALS AND METHODS**

**Experimental animals**

This study was conducted on 90 mature male swiss mice with an average weight of 25-30 g. They were left two weeks for acclimatization, then, they were equally divided into 5 comparable groups. Throughout the experimental period, they were kept under constant environmental conditions and offered food and water *ad libitum*.

**Tumor transplantation**

Ehrlichia tumor line obtained from National Cancer Institute, Cairo, Egypt with approximately \(5 \times 10^5\) tumor cells in 0.2 ml serum free medium were inoculated subcutaneously into the dorsal aspect of the upper back of 54/90 mice. Tumors were used for experimentation when they reached a surface diameter of 3-7mm measured by vernier caliper. Tumor volume was calculated by the formula (Chung *et. al.*, 2000).

\[ V = \frac{p}{6} (L \times W \times H) \]

whereas:

**Preparation of Hypericin (Hyp)**

Hypericum perforatum extract in a powder form was purchased from Seekum, Pharmaceutical Co., Al-Sharkeya Gov. Each gram powder of Hypericum perforatum extract contained 275 mg Hypericin. The extract was dissolved in 70 % aqueous ethanol to prepare a stock solution containing 192 mg hypericin /ml ethanol, and stored at -20 °C in dark place. Hypericin dose was equivalent to 5mg/Kg body weight (or 125ug hypericin / mice) (VanderWerf *et. al.*, 1996).

**Bacille Camellete - Guerin ( BCG ) vaccine**

Lyophilized BCG vaccine in vials each containing 0.5 mg was purchased from Public Organization of Medical Products, Ministry of Health, Agoza, Giza, Egypt. Each BCG vial was diluted in BCG vaccine solvent (according to the labeled information and enclosed pamphlet).
Light source and dosimetry (Non-coherent light)

Irradiation was performed using non-coherent light source using tungsten lamp with central wavelength of 590nm and a delivery system of fiberoptic cable made by Photon Co., Egypt.

The Experimental design

Negative control group

Eighteen/ninety mice did not receive any light or chemical treatment.

Hypericin control group

Eighteen/ ninety mice were injected subcutaneously (sc) into the dorsal aspect of the upper back and intraperitoneal (ip) by hypericin solution with a dose rate of 5mg/kg body weight in each route. Injections were applied 5 times / week for 6 successive weeks.

Tumor control group

Eighteen/ ninety mice were subjected to subcutaneous inoculation of Ehrlichia cell line as previously described and left without treatment along the experimental period.

Hypericin assisted photodynamic therapy group (PDT Group)

Eighteen/ ninety tumor-bearing mice were subjected to PDT protocol. As the tumor surface area reached 3-7mm in diameter, tumor sites were depilated using hair-removal cream, and Hypericin was administrated by i.p. and intratumoral (i.t.) routes, at a dose rate of 5mg/Kg by both routes. Two hours after administration, animals were anesthetized by xylazine HCl at 13 mg/kg, together with Ketamine HCl at 87 mg/kg by intramuscular route, and then, the tumors were irradiated with the light source. The light density was used at 120 J/cm² for 5 min /session (Blank et. al., 2001). Mice received 5 sessions / week for 6 weeks.

BCG/hypericin PDT-treated group

Eighteen/ ninety tumor-bearing mice received BCG at a dose rate of 25ug / mice, at the same time of tumor cell line inoculation. When the tumor surface area reached 3-7mm in diameter, animals were subjected to the previously illustrated hypericin/ PDT treated group.

Pathological examination

Three mice from each control and treated groups were examined clinically and were sacrificed weekly for 6 weeks. Careful necropsy and macroscopical examination of the sacrificed mice were done. Skin tissue specimens from control and treated groups were collected and fixed in 10% neutral formalin, dehydrated in ethyl alcohol, cleared in xylene and embedded in paraffin. Sections of 4-5 microns were obtained and stained with H&E for routine histopathological examination. Alizarin red was used...
to demonstrate calcium salts, and Masson’s Trichrome was used for fibrous connective tissue detection (Bancroft and Stevens, 1996).

**RESULTS**

Gross examination of hypericin Control mice revealed that the picture of injected area became hard in texture with loss of hair at the 4th week, while, there appeared severe hardness at the site of hypericin injection with brown colour in association with easily detached hair at the end of the 6th week.

The gross examination of tumor control group revealed slowly grown tumor masses with no clear borders till the 3rd week. Three weeks later, the tumor showed fast growth rate that gave rise to voluminous hard and ulcerate tumor masses. The most noticeable appearance was the tendency of the tumor to grow in a diffuse manner throughout the experimental period (Figs. 1 and 2).

Gross examination of PDT-group showed progressive reduction in tumor size. At the 3rd week, some tumors had cavitations on cut sections. These cavities contained liquefied necrotic debris. At the end of the 5th and 6th weeks, there was marked regression in the tumor masses which were replaced in large areas by scar tissue (Figs. 1 and 2).

Gross examination of BCG –PDT regimen exhibited progressive retardation in tumor growth with circumscribed pattern of growth and clear borders. At the end of the 6th week, there was marked improvement in the curing process than in PDT-group indicated by replacement of most tumor masses by mature scar tissue (Figs.1&2).

Microscopic examination of hypericin control group revealed appearance of heavy inflammatory cells infiltration, mainly neutrophils, lymphocytes and macrophages laden with yellow-brown pigment granules, and by the extension of the experimental period, there was marked increase in the number of macrophages. At the end of the 6th week, there was marked hyperkeratosis with thickening in the epidermal layer in some areas. The upper dermal layer had proliferated microvasculature with small areas of haemorrhage, while, the deeper dermal layer showed multiple necrotic areas and aggregations of lymphocytes and macrophages (Fig. 3).

Microscopic examination of tumor control group revealed highly condensed viable tumor cells varied in shape and size. Scattered necrotic foci within viable tumor cells were also seen. By advancement of the experiment, the anaplastic changes were more distinguishable in association with high mitotic activity (Fig.4). The functional blood vessels showed congestion and haemorrhages.
Microscopic picture of PDT-treated group: at the early stage, the tumor masses were composed of highly condensed viable tumor cells. At the second and third weeks, tumor cells showed swollen appearance and/or disrupted membrane. Some swollen tumor cells had autophagic vacuoles containing debris in the cytoplasm with no nuclear changes (Fig. 5). Multiple necrotic foci were detected and became more pronounced by the time. Most of the necrotic areas were separated from the surroundings by a reaction in the form of fibrous connective tissue infiltrated by inflammatory cells associated with vascular reaction manifested by newly formed vasculature with congestion and destruction (Fig. 6). At the 4th week, the tumor masses were noticeably reduced in size and composed of more loose and less viable tumor cells admixed with adipose and fine strands of fibrous connective tissue. The central areas of tumor showed necrosis and even calcification (Fig. 7). At the 5th and 6th weeks post–treatment, the viable tumor cells showed less mitotic activity with marked arrest in tumor cells proliferation. Large areas of tumor masses showed granulation tissue formation of fibrous connective tissue within which, small blood vessels were present in association with macrophages infiltration (Fig. 8).

**BCG-PDT group**: at the early stage of treatment, significant changes were reported in the form of pronounced inflammatory reaction at the periphery of the tumor masses associated with large numbers of multinucleated giant cells (Fig. 9). Moderate reduction in the number of viable tumor cells in comparison with tumor control group was seen. The picture of cell necrosis increased, especially in the area with damaged blood vessels. Tumor masses continued to regress with less density of viable tumor cells compared with PDT group (Fig. 10). The degenerated tumor cells showed the pathological criteria of necrosis, programmed cell death characterized by cell shrinkage, nuclear fragmentation and chromatin condensation, and even calcification (Fig. 11). Large number of tumor giant cells and macrophages containing phagosomes were detected infiltrating the newly formed fibrous connective tissue. At the 5th and 6th weeks, there was marked arrest of tumor cell proliferation with less significant number of viable tumor cells. Most of the tumor masses were substituted by adipose tissue and mature granulation tissue (Fig. 12).

**DISCUSSION**

Photodynamic therapy (PDT) is an experimental modality for treatment of tumor based on the presence of a photosensitizing agent (photosensitizer) which accumulates in the neoplastic tissue and could be activated by certain light source which matches the absorption spectrum of the photosensitizer. The subsequent
liberated free radicals and singlet oxygen, damage the vascular endothelium and / or the cell membrane with an end result of tumor cell necrosis (Ochsner, 1997).

In the current study, hypericin (Hyp) was used as a photosensitizing agent. The successful incorporation of hypericin into the cells was clarified microscopically in the hypericin control group by the presence of the hypericin pigment granules into the macrophages as a yellow-brown pigment.

Hypericin is known to be a lipophilic molecule (Takahashi et. al., 1989) that can be incorporated in phospholipid bilayers of the cell plasma membrane (Tang et. al., 1990) with subsequent more affinity to tumor cells (Chung et. al., 1994).

Because PDT is performed when the plasma drug level is high, it is likely that vascular damage results in the leakage of hypericin out the entire edematous area with more powerful cytotoxic effect through high yield of singlet oxygen. The i.t. route of hypericin injection was used to provide the real advantages for both of tumor localization and decreased normal tissue phototoxicity compared with conventional systemic dye administration. On the other hand companied i.p. route was used to maximum hypericin concentration in the skin (Uzdensky et. al., 2002).

Selection of the wavelength (590 nm) was depending on the aromatic ring structure of this dianthraquinone (hypericin) which resulted in strong absorption peaks at 550 and 590 nm with high power for tissue penetration at 590 nm, and subsequently, high production of ROS within the cell, which ultimately resulted in irreversible damage of vital cell components (VanderWerf et. al., 1996).

In the current study the photoactivated tumor masses showed gradual regression in the tumor size of both PDT and BCG/PDT groups, in comparison with the tumor control group.

The ability of photoactivated hypericin to induce necrosis had been related to its ability to induce degeneration of the viable tumor cells.

The most predominant type of cellular degeneration in the present study was the necrosis in the form of swollen appearance of the tumor cells with membrane disruption. Moreover, there were different stages of necrosis beginning from individual cell necrosis till massive necrosis with end step of calcium deposition. The recorded necrosis could be attributed to the secondary vascular shutdown. Tissue ischemia as a result of the decreased blood perfusion starves tumor cells till death. Inactivation of the blood vessels in the tumor center of the treated groups occurred earlier than in the tumor periphery and surrounding tissue (Hamilton et. al., 1996).

Meanwhile, the necrotic effect of photoactivated hypericin is related to the release of the mitochondrial cytochrome c into the cytosol which is followed by procaspase-3 activation, and subsequently, induction of necrotic machinery.
photodynamic activation of hypericin, light is absorbed and produced an excited triplet state before transfer of energy to ground-state oxygen, creating singlet oxygen as the primary oxidant. The highly reactive singlet oxygen is toxic to the mitochondria, where it inhibits succinoxidase causing cell death (VanderWerf et al., 1996, Chung et al., 1994 and Uzdensky et al., 2002).

Concerning the BCG/PDT group, there was detectable variation in the mode of tumor growth, as tumors tend to be more circumscribed in comparison with the diffuse manner of growth which was pronounced in PDT group and tumor control group. Furthermore, BCG/PDT group exhibited a perfect response detected by the macroscopic appearance which showed marked regression in tumor masses with enhancement of the curative rate at the end of the experimental period. At the early stages of treatment, there was marked increase in the number of macrophages which helped in removal of dead and necrotic tissue. Continuous removal of necrotic tissue improved the replacement of tumor mass by fibrous connective tissue at the end of the experimental period.

From the current study, it could be concluded that, Hypericin-PDT could be considered as a treatment of choice for superficial neoplasms. Immune-stimulation with BCG enhances the response of the tumor to the PDT through removal of necrotic debris.
Figs. 1. The gross appearance of tumor masses in the tumor control group and treated groups of mice during the first three weeks.

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<thead>
<tr>
<th>Tumor Control Group</th>
<th>PDT Group</th>
<th>BCG / PDT Group</th>
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<tbody>
<tr>
<td><strong>The 1st Week</strong></td>
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<td><img src="image2.png" alt="Image" /></td>
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<tr>
<td><strong>The 2nd Week</strong></td>
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<td><img src="image4.png" alt="Image" /></td>
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<tr>
<td><strong>The 3rd Week</strong></td>
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Fig. 2. The gross appearance of tumor masses in the tumor control group and treated groups of mice during the last three weeks.

<table>
<thead>
<tr>
<th>Tumor Control Group</th>
<th>PDT Group</th>
<th>BCG / PDT Group</th>
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<tbody>
<tr>
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<tr>
<td><strong>The 5th Week</strong></td>
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<td><img src="image10.png" alt="Image" /></td>
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<tr>
<td><strong>The 6th Week</strong></td>
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Fig. 3. Hypericin control group of mice at the 6th week showing congestion of the subepidermal vasculature (asterisks) with inflammatory cells infiltration. The macrophages were laden by yellowish-brown pigment. H&E X 200.

Fig. 4. Tumor control group of mice showing highly condensed viable tumor cells under an intact epithelial surface with multiple focal areas of necrosis. H&E X 100.

Fig. 5. PDT-group of mice at the 2nd week revealing evidences of cytoplasmic swollen accompanied by loss of cell membrane integrity with multiple cytoplasmic vacuolization. H&E X 400.

Fig. 6. PDT-group of mice at the 3rd week illustrating proliferation of FCT with neutrophils infiltration associated with congestion and destruction of the newly formed vasculature. H & E 200.

Fig. 7. PDT-group of mice at the 4th week showing reduction of the tumor mass which showed central calcification and surrounded by fibrous connective tissues. Alizarin red X100.

Fig. 8. PDT-group of mice at the 6th week showing substitution of large area of tumor mass by granulation tissue formation. H&E X 200.
Fig. 9. BCG/PDT group of mice at the 1st week showing marked skin reaction between the healthy tissue and the necrotic tumor mass in the form of inflammatory cells infiltration in addition to multinucleated giant cells. H&E X 200.

Fig. 10. BCG/PDT group of mice at the 3rd week showing less density of viable tumor cells with less mitotic activity. H&E X 400.

Fig. 11. BCG/PDT-group of mice at the 4th week showing calcium salts deposition in the center of necrosed tumor mass. H & E X 200.

Fig. 12. BCG/PDT-group of mice at the 6th week showing marked substitution of tumor mass by granulation tissue, macrophages, well organized FCT and even adipose tissue. Masson’s trichrome X400.
REFERENCES


دراسات باثولوجية على علاج الأورام في الجرذان بواسطة الهيبيرسين المستمحت ضونياً

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تم إجراء هذه التجربة على عدد 90 فأراً أربعة صغاراً (مستوى وزنها 20-25 جم) تم تقسيمها إلى خمس مجموعات متساوية كل منها تحتوي على 18 فأراً. المجموعة الأولى استخدمت مجاعة ضاحية سلامة. المجموعة الثانية استخدمت مجاعة ضاحية لدادة الهيبيرسين وتتم حقنها بجرعة 5 جم/كم في جرعة متساوية لكل من الحقن البروتيني والحقن تحت الجلد 5 مرات أسبوعياً لمدة 6 أسابيع. أما المجموعة الثالثة فقد استخدمت مجاعة ضاحية لسرطان حيث تم زرع خلايا إبريلخ (500,000 خلايا/فأر) تحت الجلد. المجموعة الرابعة تم فيها زرع الخلايا السرطانية (وبدع أسبوعين من نمو السرطان) تم تطبيق العلاج الضوئي كما يلي: حقن الهيبيرسين بجرعة 5 جم/كم في جرعة متساوية لكل من الحقن البروتيني والحقن داخل السرطان ثم تعرض السرطان للضوء بقوة 20 جول لمدة 5 دقائق في المرة الواحدة وتكرار ذلك 5 مرات يومياً لمدة 6 أسابيع. المجموعة الخامسة تم علاجها مثل المجموعة الثانية بالإضافة إلى تحصينها بسلقاح بي. سي. جي مزمناً مع استزراع الخلايا السرطانية.

بين أن فترات المجموعة الثانية الضخمة للصبيحة قد ظهر بها تصد ي لدى بالحنال عند مواسم الحقن والتي أظهرت أيضاً لوناً غامقاً وقد وضح هذا مجهرياً كزيادة في طبقة الكيراتين مع وجود خلايا منتهمة مماثلة صبغة بنيه متمتد للذن الأيسر.

المجموعة الثالثة أظهرت زيادة تدريجية في كتلة السرطان منذ بداية الأسبوع الأول وقد أصبح نمو السرطان نشطاً عند الأسبوعين الخامس والست.

أظهر الفحص عيني لمجموعات السرطان المعالجة ضوئياً عدم وجود تغير واضح في معدل نمو السرطان مقارنة بالمجموعات الضخمة لسرطان حتى الأسبوع الثالث ولكن المقتريات العرضية أظهرت تجويضات ملموسة بمقدار متكررة وخلايا منتشرة مع وجود عدد كبير من الخلايا الإنتهاجية والخلايا الإنتهاجية الأخرى بالإضافة إلى عدد كبير من الخلايا السرطانية العضالة مختارة لكليات السرطانية.

بالتقدم في التجربة ظهر ضمور في حجم الكتل السرطانية مع تكون أنماط ندبة والتي كانت واضحة وسريعة في المجموعة الخامسة مقارنة بالمجموعة الرابعة.

أظهرت النتائج أن استخدام العلاج الضوئي بواسطة الهيبيرسين فقط أو بالإضافة إلى التحصين بسلقاح بي. سي. جي يحدث تدميراً كبيراً في النخر الخلوى والإنتهاج الخلوى. وقد اختلقت درجة التكاثر الخلوى في البروتوكول العلاجي المختبر استناداً على مدة العلاج. حيث
أنه إتباعاً لهذا البروتوكول العلاجي كان هناك علامات إنتحار خلوى واضحة وذلك عند المراحل المبكرة من العلاج بينما كانت مظاهر السندر الخلوى وقلة الامتصاص للخلية مع قلة تعدد الأشكال ظهرت مع طول فترة العلاج حيث تم إحلال الكتلة السرطانية المرتدة بنسيج متحسوب مكون من نسيج ليفي ضام وشعيرات دموية متكررة.

أوصت الدراسة الحالية باستخدام العلاج الضوئي بواسطة الهيبيرسين مع التحسين بلقاح سي. أوصت الورام الجلد في الجلد أن هذا العلاج يحدث تدميراً كبيراً للخلايا السرطانية بالإضافة إلى إحلال الكتلة السرطانية المتكررة بنسيج ليفي ضام وشعيرات دموية متكررة (أنسجة ندبية).