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(54) Title: METHODS TO POTENTIATE INTRAVENTOUS ESTRAMUSTINE PHOSPHATE

(57) Abstract

Estramustine phosphate is an anti-mitotic chemotherapeutic drug with proven efficacy against cancer. The invention describes methods which potentiate the therapeutic benefit of intravenous estramustine phosphate. The invention provides for intravenous estramustine phosphate to be administered at a high dosage exceeding 1300 mg as a single dose. Efficacious enhancement of estramustine phosphate pharmacokinetics is thereby achieved. Further provided, estramustine phosphate may be intravenously administered for use in combinational regimens with other chemotherapeutic agent. The therapeutic advantages achieved using the intravenous estramustine phosphate formulation are applicable to treatment of a variety of cancers including prostate cancer, breast cancer, lung cancer, colorectal cancer, pancreatic cancer, ovarian cancer, melanoma, and other cancers.
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TITLE OF THE INVENTION

METHODS TO POTENTIATE INTRAVENOUS ESTRAMUSTINE PHOSPHATE

This application claims priority to U. S. Provisional Application serial No. 60/079,542, which was filed on March 27, 1998.

BACKGROUND OF THE INVENTION

Field of the Invention:

The present invention relates to the use of estramustine phosphate, a non-nitrogen mustard carbamate derivative of estradiol-17b-phosphate, as a high dose infusion. The present invention further relates to methods to potentiate intravenously administered estramustine phosphate and to methods for treating cancer by intravenously administering estramustine phosphate.

Discussion of the Background:

Cytotoxic effects have been shown to be due to the intact estramustine molecule (Hartley-Asp. 1982). Tissue culture studies have shown that estramustine (EM) is an anti-mitotic agent, causing a dose-dependent blocking of tumor cell division in the metaphase (Hartley-Asp. 1984). Metaphase arrest is known to be caused by an interference of drugs with the microtubule structure that forms the mitotic spindle. It has been shown, with the help of immunohistochemistry, that dose-dependent disturbances of interphase microtubules occur in cultured human prostatic cells (Mareel 1988, Dahllof 1993). Treatment with EM in vitro inhibited the assembly of microtubules composed of only tubulin demonstrating a direct interaction with tubulin (Dahllof 1993). In addition, an interaction with microtubule associated
proteins (MAPs) has been demonstrated (Stearns 1988). MAPs are high molecular weight proteins that are believed to be important in stabilizing microtubules. That EM exhibits the mechanism of action of an anti-mitotic agent has been confirmed in vivo (Eklöv. 1992).

Estramustine phosphate is thus an anti-mitotic agent currently used in the treatment of advanced adenocarcinoma of the prostate. As a single agent, its activity in hormone-refractory prostate cancer is comparable to that of several other cytotoxic agents that have been studied in a series of multi-institutional, randomized trials by the National Prostatic Cancer Project (Murphy, 1983). While the drug is usually administered orally at a dose of 10-15 mg/kg/day, it is approved for intravenous administration in several countries. However, estramustine phosphate when administered intravenously has been used at dosages and according to a schedule paralleling the oral administration for the drug, i.e. at recommended dosages of 300-600 mg daily given intravenously and usually repetitively over for several consecutive days. This is then followed by orally administered drug.

In the published material, details from about 500 patients who have been treated with the intravenous formulation initially which was then followed by oral treatment can be found. Induction schedules employing 300-600 mg intravenous daily for 7-21 days, followed by daily oral doses, were typical in these studies. The drug was administered as a slow intravenous injection or as a bolus at 300 mg/day, and thrombophlebitis and local irritation at the peripheral intravenous injection sites were considered major limitations of drug administration requiring the establishment of central line administration in many patients or discontinuation of treatment. At 450 mg/day, Nagel and Kölln (1977) stated that this led to severe gastrointestinal
problems that 300 mg/day was taken as the maximum intravenous daily dose.” In a
compilation, by Andersson et al. of 245 patients receiving 300-600 mg /day for 21 days
followed by the same dose once or twice weekly for 2 months. 20% of the patients
exhibited thrombophlebitis. 17% exhibited gastrointestinal problems and 9%
exhibited liver disturbances. Toxocities resulting from such repetitive dosing
schedules often require drug discontinuation (Lundgren, 1995). Maier (1990),
administered daily intravenous doses of 900 mg/day for 7-10 days, followed by oral
therapy, without reporting phlebitis but severe liver problems did occur in 11 of 18
patients (61%) with one death due to toxic liver failure.

The state of the art thus typically utilized intravenous estramustine phosphate
formulations as a single-agent method for initiating a long term oral estramustine
therapy. Furthermore, intravenous administration of estramustine phosphate at higher
dosages is generally considered prohibitive due to toxicity. It is neither known nor
obvious to the art that single dose, high-dosage administration of estramustine
phosphate is feasible intravenously. While dosing up to 1200 mg/m² have been given
orally (Keren-Rosenberg, 1997), differences in drug metabolism and bioavailability do
not permit extrapolation to the high dose intravenous formulation, with relative
bioavailability of estromustine after oral administration found to be only 44%.
(Gunnarsson, 1984), with the phosphate moiety dephosphorylated in the oral
formulation in contrast to the intravenous formulation. Furthermore, it is not known
in the art that intravenous estramustine phosphate can be used in combinational
chemotherapy regimens, including the use of higher dose intravenous estramustine
phosphate. Furthermore, it is not known to the art that intravenous estramustine
phosphate has clinical utility for cancers other than for the prostate cancer indication.
In previous work, Dr. Beryl Hartley-Asp, a co-inventor of this invention, was the first to recognize the synergistic potential of estramustine phosphate with other cytotoxic agents. (Mareel 1988). In several experiments, it was demonstrated that prolonged exposure to estramustine was necessary to achieve potentiation.

Consequently, daily dosing was deemed necessary leading to the use of the ORAL preparation as previous data from the intravenous (IV) preparation suggested that achievement of constant high levels would not be clinically achievable with IV dosing.

Additive and possibly synergistic antimicrotubule effects in cells in vitro have been shown for estramustine and many other cytotoxic agents, (Mareel 1988, Speicher 1992, Pienta 1993, Batra, 1996). Thus, the combination of estramustine phosphate with other drugs in humans has been carried out using ORAL administration of estramustine phosphate. Phase II trials (Seidman, 1992, Hudes, 1992, Pienta, 1994, Hudes, 1996) with Estramustine phosphate combined with vinblastine, have been carried out in hormone refractory prostate cancer. In these trials a 50-75% decrease in prostate specific antigen was demonstrated among 88 patients. The most frequent toxicity was mild to moderate nausea. Of particular note is the 10.5% (4/37) incidence of significant cardiovascular toxicity including one deep venous thrombosis (DVT), one myocardial infarction, one episode of congestive heart failure, and one reversible neurologic event which required stopping therapy in these patients and which can be attributed to estramustine phosphate. In another Phase II trial carried out by Pienta et al. (1994), estramustine phosphate (oral) was combined with Etoposide. Fifty two patients were evaluable; including 20 patients with soft tissue disease, in which 3 complete responses (CR) (15%) and 6 partial responses (PR) (30%) were.
observed. In 32 patients with bone metastases 8 patients improved (25%), and 12 patients were stable (38%). Overall 13 men (25%) had a 75% decrease in prostate specific antigen, and 28 men (54%) had a 50% decrease. A Phase I-II study of Taxol (Hudes, 1992) and estramustine phosphate was carried out in seventeen patients with hormone refractory prostate cancer. Six patients had measurable disease and 3 of these obtained a PR of 2+, 6, and 8 months. Prostate specific antigen (PSA) decreased by ≥ 50% in 58.8%. Median duration of response was 7 months. Grade 3-4 granulocytopenia and mucositis occurred in 2 patients, nausea grade 1-2 (70.5%) and grade 3 in one patient. Edema was seen in 8 patients (47%) and transient hepatic enzyme elevation of grade 1-3 in 6 patients (35.2%).

In a recent study, Petrylak et al., (1997), using escalating doses of docetaxel with estramustine phosphate given orally demonstrated an overall prostate specific antigen response rate of 62%. In patients with bidimensionally measurable disease, 3 (43%) achieved a partial response in lymph nodes, and 1 achieved a minor response in ischial mass. This demonstrates that combination treatment with ORAL estramustine phosphate is efficacious. However, combinations of intravenous estramustine phosphate with these cytotoxic agents are not known to the art. Differences in the metabolism, particularly regarding the phosphate moiety, in the oral versus intravenous estramustine phosphate formulations make combinational therapies with the intravenous formulation non-obvious.

In contrast to other anti-mitotic agents, the effect of estramustine phosphate appears to be dependent on the presence of the estramustine binding protein (EMBP) (Eklöv, 1996). This is found under normal conditions only in the prostate (Forsgren, 1979, Flucher, 1989). However, a similar protein has also been identified in many
cancerous tissues, as well as prostate tumors. such as lung, breast glioma, colon, pancreas (Björk 1991, Bergh 1988, Eklöv 1996, Edgren 1996, Von Schoultz, 1994, Bergenheim, 1993). This protein binds estra- and estro-mustine (EaM and EoM) with very high affinity and is thought to be responsible for the selective retention of EoM in the prostate tumor. where a ratio of 1:6 to 1:11 plasma/tumor has been found in prostate cancer patients treated with estramustine phosphate oral and intravenously, respectively (Norlen 1988, Walz 1988). Recently, we have demonstrated a correlation between the levels of EMBP and the levels of EaM and EoM in human prostate tumors after a single intravenous estramustine phosphate dose to patients before radical prostatectomy. indicating that EMBP could be the cause of drug retention (Walz, 1996).

BRIEF DESCRIPTION OF THE FIGURES

A more complete appreciation of the invention and many of the attendant advantages thereof will be readily obtained as the same become better understood by reference to the following detailed description when considered in connection with the accompanying drawings, wherein:

Figure 1 illustrates the concentration of estramustine phosphate after a single intravenous dose of Estracyt (mean ± SEM, N = 4 + 4 + 3) given at a dosage of 1000 mg (range 980-1070 mg), 1000 mg/m², and 1500 mg/m²; and

Figure 2 illustrates the concentration of estromustine after a single intravenous dose of Estracyt (mean ± SEM, n = 4 + 4 + 3) given at a dosage of 1000 mg (range 980-1070 mg), 1000 mg/m², and 1500 mg/m².
SUMMARY OF THE INVENTION

The present invention describes methods of potentiating the therapeutic use and efficacy of intravenously administered estramustine phosphate. It provides for the intravenous administration of estramustine phosphate in dosages exceeding 1300 mg. It also provides for intravenous administration of estramustine phosphate at dosages exceeding 950 mg/m² (milligrams per square meter of body surface area). It further provides for administration of high dose intravenous estramustine phosphate as a single dose, which may further be administered on a weekly or longer schedule. The present invention enables optimization of pharmacokinetics as to maximize therapeutic advantage, and further enables use of intravenous estramustine phosphate in combination with other therapies, including other chemotherapies, providing further improved therapeutic benefit. The present invention enables use of intravenous estramustine phosphate as therapy for multiple tumor types, including prostate, breast, lung, ovarian, colorectal, melanoma, pancreatic, and brain cancers.

Thus, one application of the present invention is to provide high dose estramustine phosphate therapy intravenously, wherein the dose exceeds 950 mg/m².

Another application is to provide a schedule of intravenous administration, whereby that schedule enables optimization of pharmacokinetics of estramustine phosphate and its metabolites at minimal toxicity, and further whereby said optimization permits convenient and efficacious combination regimens of therapy.

Thus, an application of the present invention permits the use of intravenous estramustine phosphate in combination with other therapeutic regimens, including cytotoxic chemotherapy.

Another application of the present invention is to provide a method which
increases binding saturation and prolongs binding duration of estramustine phosphate or its metabolites to estramustine binding protein or estramustine binding protein-like protein (EMBP).

Thereby, the present invention provides application to treatment of cancers having EMBP, including but not limited to prostate, breast, lung, ovarian, colorectal, melanoma, pancreatic, and brain cancers, by intravenous administration.

Another application of the present invention is to provide a method of rapidly relieving symptoms secondary to cancer, inclusive of but not limited to cancer-induced pain and urinary obstruction.

Further, the present invention enables these applications for use of intravenous estramustine phosphate independent of the formulation. Thereby, the present invention provides for the infusion of estramustine phosphate as free drug, as protein-bound drug, or as drug within liposomes.

Thereby, the present invention describes a formulation of estramustine phosphate wherein the estramustine phosphate is administered intravenously in conjunction with liposomes.

Thus, the method of the present invention in which doses above 900 mg/m² (generally greater than 1300 mg per dose) can be administered safely and within an effective schedule is extremely unexpected.

The present invention teaches the advantage of intravenous estramustine in combination with other chemotherapy agents. The present invention further teaches the advantage of high dose intravenous estramustine in combination with other chemotherapeutic agents.

We teach in this invention that intravenous estramustine phosphate may be
used to treat tumors having elevated EMBP-like protein (herein referred to simply as EMBP).

The novel and non-obvious applications of the present invention can be recognized from a comparison of the pharmacokinetic data following oral administration estramustine phosphate with that following high-dose intravenous administration of estramustine phosphate. The pharmacokinetic and toxicity data regarding high-dose intravenous estramustine phosphate is not known to the art. Dephosphorylation of estramustine phosphate to estramustine (EM), followed by oxidation at the 17 position to estromustine (EoM), the estrone analogue of EM, are the major metabolic steps after administration of oral estramustine phosphate in man. EoM is the predominant metabolite found in plasma when estramustine phosphate is administered on the daily oral schedule. The relative bioavailability based on estromustine is approximately 44% (Gunnarsson, 1984). After intravenous administration estramustine phosphate is initially found in plasma but is rapidly hydrolyzed to the same metabolites as are found after oral administration, the major metabolite being estromustine. Both estramustine and estromustine are further metabolized by cleavage of the carbamic ester to yield approximately 15% estradiol and estrone, respectively (Gunnarsson, 1981, 1984). We have demonstrated an unexpected prolonged availability of the major metabolite estromustine following high-dose intravenous administration, which can lead to unexpected clinical benefits. Previous data from patients treated with a single intravenous dose of 300 mg demonstrated that the elimination of estromustine had a half lives of 10-20 hours. The main route of elimination was metabolism of estramustine phosphate to estromustine, estramustine, estradiol and estrone. The data of particular importance for the efficacy
of estramustine phosphate was the half lives of estramustine phosphate. (Figure 1), and the major cytotoxic metabolite estromustine (Figure 2). By application of the methods of this invention, we now demonstrate the novel finding that after a single high intravenous dose of estramustine phosphate at 1000mg/m² it was found that the half life of estromustine was approximately 100 hours. (Figure 2). This finding further enables therapeutic applications of high-dose intravenous estramustine phosphate.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The present invention teaches the ability to administer estramustine phosphate at doses above 950 mg/m² (i.e., greater than 1300 mg).

The method of the present invention is performed as follows. In the preferred method, estramustine phosphate is administered at a single infusion dosage exceeding 950 mg/m². Intravenous administration is performed either through a central or peripheral intravenous route. During preparation of the intended drug, the contents of packaged estramustine phosphate intended for intravenous usage are dissolved, wherein the packaged contents may consist of but are not limited to a lyophilized powder of the meglumine salts in vials of estramustine phosphate, or similar freeze-dried estramustine phosphate which are first dissolved in sterile water such as 5 ml sterile water per 300 mg estramustine phosphate, or in 5% dextrose in water for intravenous administration. In the preferred method, 5% dextrose in water is used as the diluent. In the preferred method, during preparation of the dissolved drug the preparation should not be shaken, but should be slowly inverted to mix. The solution is then given as an intravenous infusion with the preferred duration of infusion time
being 30 minutes to 3 hours, whereby infusion over 1-2 hours is a safe and convenient method. Saline solution may result in drug precipitation and thereby its use is not preferred in the infusion.

When estramustine phosphate is administered through a peripheral intravenous route, it is preferred that a longer duration of infusion and greater total infusional volume be utilized to minimize vascular irritation. Alternatively, the estramustine phosphate solution can be mixed with various amounts but preferably 3-5% human albumin or other plasma proteins including synthetic plasma proteins to achieve protein binding of the estramustine phosphate and therefore minimize any potential vascular damage.

The invention may be further realized using other preparations or formulations of estramustine phosphate. One particularly advantageous preparation of the chemotherapeutic agent estramustine phosphate which enables infusion of estramustine phosphate through either a peripheral or central vein, both at high doses and also doses less than 1300 mg, involves the infusion of estramustine phosphate in conjunction with liposomes (herein referred to as liposome encapsulated estramustine phosphate or liposomal estramustine). In one preferred method of preparing liposomal estramustine, estramustine phosphate solution is first prepared in the manner described above and then injected into a vial containing empty liposomes available as a lyophilized powder. Following adequate hydration of the liposomes, the vials are vortexed and sonicated, followed by infusion into the patient.

When estramustine phosphate is administered through a central venous route, said administration may be performed through either a temporary or permanent venous access device, including but not limited to a triple lumen catheter. Hickman
catheter, subclavian line, jugular line, or medi-port. Said administration may be but is not necessarily performed concomitant with anticoagulant therapy or with the addition of varying amounts but preferably 3-5% human albumin or other plasma proteins or liposomal estramustine to minimize any potential vascular damage in a given patient.

While the dosage of estramustine phosphate in the present invention is greater than 1300 mg, it is preferred that the patient be treated at a dose exceeding 950 mg/m². Thereby, one preferred method is to administer a single intravenous dosage of 1000 mg/m². Another preferred method is to administer a single intravenous dosage of 1500 mg/m². Furthermore, a dosage of 2000 mg/m² may be administered. However, the invention is inclusive of other dosages above 950 mg/m² and the preferred dosages are not to imply limitation.

The most preferred schedule of estramustine phosphate administration in the invention is a single infusion given once weekly to a maximal dose of 4000 mg or 3500 mg/m². Another preferred schedule is administration of a single drug infusion once every two weeks. Another preferred schedule is administration of a single drug infusion once every three weeks. Another preferred schedule is administration of a single drug infusion once every four weeks. One schedule may be preferred over another in consideration of schedules with other concomitant therapy. These schedules may repeat in a serial or repetitive fashion.

The invention described herein enables methods to prolong blood and/or tissue levels at high elevations for estramustine phosphate metabolites, including estromustine, estramustine, estrone and estradiol. Thereby, enhanced synergistic interactions with other therapies is enabled, wherein such other therapies include but are not limited to chemotherapy, radiotherapy, monoclonal antibodies, and biologic
therapies. The present invention provides maximization of therapeutic benefit by prolongation of elevated blood and tissue levels of estramustine phosphate and its metabolites. Thereby, maximization of therapeutic benefit is achieved wherein estramustine phosphate is administered intravenously at dosages exceeding 950 mg/m², which are administered in combination with other cancer therapies, inclusive but not limited to radiotherapy, chemotherapy, monoclonal antibodies, and biologic therapies.

In the preferred method, therapeutic benefit is potentiated by administering intravenous estramustine phosphate at single dosages exceeding 950 mg/m², with other cytotoxic chemotherapies. In the preferred method said combination is achieved by administering intravenous estramustine phosphate within 3 days of the other chemotherapeutic agents, preferably on the day of, or the day prior to administration of the other chemotherapeutic agents. A particularly preferred method is achieved when the other chemotherapeutic agents consist of anti-mitotic agents or anti-microtubule agents, inclusive of but not limited to taxanes, including taxol and taxotere, and agents including vinblastine, vincristine, etoposide, navelbine, doxorubicin, irinotecan (CPT-11), and liposome encapsulated chemotherapeutic agents, including liposome encapsulated taxanes such as liposome encapsulated paclitaxel. It may be further beneficial if a combination with a monoclonal therapy is utilized, that the monoclonal agent include a radionucleotide or an anti-growth factor agent.

Plasma or serum levels of estramustine are further sustained when estramustine phosphate is administered intravenously as a single infusion at a dosage exceeding 950 mg/m². The infusion may optionally be repeated in a serial or
repetitive manner to maintain elevated blood levels of the estromustine phosphate metabolites. Sustained levels of estromustine phosphate and its metabolites thereby enable sustained therapeutic benefit.

The present invention thereby provides a method to increase the binding saturation of estromustine or its metabolites to estromustine binding protein or like-protein by administering estromustine phosphate intravenously at a single infusion dose exceeding 950 mg/m². Similarly, the binding duration of estromustine phosphate or its metabolites to estromustine binding protein or estromustine binding protein-like protein (EMBP) is increased in the invention by administering the drug at intravenous dosages exceeding 950 mg/m². Thereby, all cancers having either estromustine binding protein or estromustine binding protein like-protein may be treated by intravenous estromustine phosphate. It is particularly preferred to treat prostate cancer in such manner. It is further preferred to treat breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, and cancers of the brain in such manner. It is particularly preferred that estromustine phosphate be administered intravenously wherein the single dosage exceeds 950 mg/m² when treating cancers having either estromustine binding protein or estromustine binding protein like-protein, inclusive of but not limited to the group of cancers including prostate cancer, breast cancer, ovarian cancer, pancreatic cancer, melanoma, lung cancer, and cancers of the brain.

Said cancers may further be treated using liposomal estromustine, either as a single agent or in combination with other chemotherapies. Said administrations are preferably repeated in serial or repetitive fashion at schedules of the invention, with or without combination of other therapies. Thus, said schedules may include
combinational treatment of intravenous estramustine phosphate with other chemotherapeutic therapies given on a once weekly, a once every two week, a once every three week, or a once every four week schedule, and variations therein.

It is particularly preferred that intravenously administered estramustine phosphate be administered in combination with other chemotherapeutic cytotoxic agents when used in the treatment of prostate cancer, breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian, and cancers of the brain. It is further particularly preferred that intravenously administered estramustine phosphate be administered in combination with radiation when used in the treatment of prostate cancer, breast cancer, lung cancer, pancreatic cancer, colorectal cancer, and cancers of the brain. It is further preferred that in treating cancers having estramustine binding protein or estramustine binding protein like-protein, including prostate cancer, breast cancer, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, and cancers of the brain, estramustine phosphate be administered at intravenous dosages exceeding 950 mg/m² when used in combination with other cancer therapies.

The present invention enables both objective and subjective therapeutic benefit. Benefit achieved may relate to reduction of tumor size, improved quality of life, reduction of tumor obstruction, such as urinary obstruction, reduction of cancer-induced pain, improved survival, reduction in time to cancer recurrence, or other evidence of improvement. In particular, rapid objective or subjective therapeutic benefit is achieved by administering estramustine phosphate intravenously at a dosage exceeding 950 mg/m², either as a single agent or preferably in combination with other cancer therapies. Thereby the invention enables rapid relief of cancer-induced urinary obstruction, and rapid relief of cancer-induced pain.
Other features of the invention will become apparent in the course of the following descriptions of exemplary embodiments which are given for illustration of the invention and are not intended to be limiting thereof.

EXAMPLES

The following clinical cases are provided by way of example and not limitation.

Example 1: Two patients with advanced metastatic prostate cancer were treated with estramustine phosphate intravenously given through a central line. The patients received an estramustine phosphate dosage of 2500 mg/m². Estramustine phosphate was administered as a single infusion on a weekly schedule in a repetitive fashion. Each infusional dose was administered over a 90 minute infusion. The infusions were well tolerated without serious toxicity and both patients demonstrated a response (reduction) in their prostate specific antigen (PSA).

Example 2: Three patients with advanced metastatic prostate cancer were treated with estramustine phosphate administered intravenously through a central line at a dosage of 1000 mg/m². Estramustine phosphate was administered as a single infusion on a weekly schedule in a repetitive fashion. Each infusional dose was administered over a 30 minute infusion. The infusions were well tolerated with several patients demonstrating PSA response.

Example 3: Three patients with advanced metastatic prostate cancer were treated with estramustine phosphate administered intravenously through a central line
at a dosage of 1500 mg/m². Estramustine phosphate was administered as a single infusion on a weekly schedule in a repetitive fashion. The infusional dose was administered either over 30 minutes or over 1 hour. The infusions were well tolerated with one patient demonstrating a response in bulky tumor adenopathy.

Example 4: Three patients with advanced metastatic prostate cancer were treated with estramustine phosphate intravenously given through a central line. The patients received an estramustine phosphate dosage of 2000 mg/m². Estramustine phosphate was administered as a single infusion on a weekly schedule in a repetitive fashion. Each infusional dose was administered over a 60 minute infusion. An anti-thrombotic agent was additionally administered for venous thrombosis prophylaxis. The estramustine phosphate infusions were well tolerated without serious toxicity, and with evidence of PSA response.

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This application is based on U. S. Provisional Application serial No. 60/079,542, which was filed on March 27, 1998, which is incorporated herein by reference in its entirety.

Obviously, numerous modifications and variations of the present invention are possible in light of the above teachings. It is therefore to be understood that, within the scope of the appended claims, the invention may be practiced otherwise than as specifically described herein.
CLAIMS:

1. A method of administering estramustine phosphate as an intravenous dose, whereby the dosage of a single infusion exceeds 1300 mg.

2. A method of administering estramustine phosphate as an intravenous dose, whereby the dosage of a single infusion exceeds 950 mg/m².

3. The method of claims 1 or 2, wherein estramustine phosphate is administered as a single infusion on a once weekly schedule.

4. The method of claims 1 or 2, wherein estramustine phosphate is administered as a single infusion on a once every two week schedule.

5. The method of claims 1 or 2, wherein estramustine phosphate is administered as a single infusion on a once every three week schedule.

6. The method of claims 1 or 2, wherein estramustine phosphate is administered as a single infusion on a once every four week schedule.

7. The method of claims 1 or 2, wherein estramustine phosphate is administered in combination with other anti-cancer therapies.

8. The method of claim 7, wherein estramustine phosphate is administered intravenously in combination with other chemotherapeutic agents.
9. A method of potentiating the therapeutic benefit of a multi-drug chemotherapeutic regimen, wherein one of the drugs in the regimen comprises estramustine, by administering estramustine phosphate as an intravenous formulation.

10. The method of claim 9, wherein said intravenous formulation comprises estramustine phosphate given at high dose.

11. The method of claim 9, wherein another drug in the regimen comprises an anti-mitotic agent or anti-microtubule agent.

12. The method of claim 10, wherein the dosage of a single infusion of estramustine phosphate exceeds 1300 mg.

13. The method of claim 10, wherein the dosage of a single infusion of estramustine phosphate exceeds 950 mg/m².

14. A method to produce prolonged elevated plasma levels of estramustine to promote synergistic interaction between estramustine and a second chemotherapeutic agent, wherein:

   estramustine is administered as an intravenous formulation; and
   estramustine is administered on the day of, or within 3 days of administration of said second chemotherapeutic agent.

15. The method of claim 14, wherein said second chemotherapeutic agent
comprises an anti-mitotic agent or anti-microtubule agent.

16. The method of claim 14, wherein the intravenous formulation comprises estramustine phosphate.

17. The method of claim 16, wherein the dosage of a single infusion of estramustine phosphate exceeds 1300 mg.

18. The method of claim 16, wherein the dosage of a single infusion of estramustine phosphate exceeds 950 mg/m².

19. A method to produce elevated plasma levels of the estramustine metabolite estramustine, to promote synergistic interaction between estromustine and a second chemotherapeutic agent, wherein:

- estrastine is administered as an intravenous formulation; and
- estramustine is administered on the day of, or within 3 days of administration of said second therapeutic agent.

20. The method of claim 19, wherein said second therapeutic agent comprises an anti-mitotic agent or anti-microtubule agent.

21. The method of claim 19, wherein the intravenous formulation comprises estramustine phosphate.
22. The method of claim 21, wherein the dosage of a single infusion of estramustine phosphate exceeds 1300 mg.

23. The method of claim 21, wherein the dosage of a single infusion of estramustine phosphate exceeds 950 mg/m².

24. A method according to claim 14 to potentiate therapeutic benefit.

25. A method according to claim 19 to potentiate therapeutic benefit.

26. A method to sustain plasma levels of estramustine and estromustine, wherein estramustine phosphate is administered intravenously as a single infusion at a dosage exceeding 1300 mg; optionally repeating the infusion in a serial manner.

27. A method to sustain plasma levels of estramustine and estromustine, wherein estramustine phosphate is administered intravenously as a single infusion at a dosage exceeding 950 mg/m², optionally repeating the infusion in a serial manner.

28. A method according to claim 26, wherein therapeutic benefit is sustained.

29. A method according to claim 27, wherein therapeutic benefit is sustained.

30. A method of increasing binding saturation of estramustine to estramustine binding protein, or like-protein wherein:
estramustine is administered as an intravenous formulation as estramustine phosphate at a single infusion dosage exceeding 1300 mg; and
binding saturation of estramustine binding protein is thereby enhanced.

31. A method of increasing binding saturation of estramustine or its metabolites to estramustine binding protein, or like-protein wherein:
estramustine is administered as an intravenous formulation as estramustine phosphate at a single infusion dosage exceeding 950 mg/m²; and
binding saturation of estramustine binding protein is thereby enhanced.

32. A method of prolonging binding duration of estramustine or its metabolites to estramustine binding protein, or like-protein wherein:
estramustine is administered as an intravenous formulation as estramustine phosphate at a single infusion dosage exceeding 1300 mg; and
binding duration of estramustine binding protein is thereby prolonged.

33. A method of prolonging binding duration of estramustine or its metabolites to estramustine binding protein, or like-protein wherein:
estramustine is administered as an intravenous formulation as estramustine phosphate at a single infusion dosage exceeding 950 mg/m²; and
binding duration of estramustine binding protein is thereby prolonged.

34. The method of claim 30, wherein the method is used to treat a cancer having estramustine binding protein, or estramustine binding protein-like protein.
35. The method of claim 34, wherein said cancer is selected from the group consisting of prostate cancer, breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, and cancers of the brain.

36. The method of claim 31, wherein the method is used to treat a cancer having estramustine binding protein, or like-protein.

37. The method of claim 36, wherein said cancer is selected from the group consisting of prostate cancer, breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, and cancers of the brain.

38. The method of claim 32, wherein the method is used to treat a cancer having estramustine binding protein, or like-protein.

39. The method of claim 38, wherein said cancer is selected from the group consisting of prostate cancer, breast cancer, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, and cancers of the brain.

40. The method of claim 33, wherein the method is used to treat a cancer having estramustine binding protein, or like-protein.

41. The method claim 40, wherein said cancer is selected from the group consisting of prostate cancer, breast cancer, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer, melanoma, and cancers of the brain.
42. A method of treatment for breast cancer in which estramustine phosphate is administered intravenously.

43. A method of treatment for lung cancer in which estramustine phosphate is administered intravenously.

44. A method of treatment for pancreatic cancer in which estramustine phosphate is administered intravenously.

45. A method of treatment for colorectal cancer in which estramustine phosphate is administered intravenously.

46. A method of treatment for ovarian cancer in which estramustine phosphate is administered intravenously.

47. A method of treatment for brain cancer in which estramustine phosphate is administered intravenously.

48. The method of claim 1, wherein the infusion is given over 30 minutes to 3 hours.

49. The method of claim 2, wherein the infusion is given over 30 minutes to 3 hours.
50. The method of claim 1, wherein relief from cancer-induced urinary
obstruction is achieved.

51. The method of claim 2, wherein relief from cancer-induced urinary
obstruction is achieved.

52. The method of claim 1, wherein rapid relief of cancer-induced pain is
achieved.

53. The method of claim 2, wherein rapid relief of cancer-induced pain is
achieved.

54. The method of claim 42, in which estramustine phosphate is administered
in combination with one or more other chemotherapeutic agents.

55. The method of claim 43, in which estramustine phosphate is administered
in combination with one or more other chemotherapeutic agents.

56. The method of claim 44, in which estramustine phosphate is administered
in combination with one or more other chemotherapeutic agents.

57. The method of claim 45, in which estramustine phosphate is administered
in combination with one or more other chemotherapeutic agents.
58. The method of claim 46, in which estramustine phosphate is administered in combination with one or more other chemotherapeutic agents.

59. The method of claim 47, in which estramustine phosphate is administered in combination with one or more other chemotherapeutic agents.

60. A method of treatment for melanoma in which estramustine phosphate is administered intravenously.

61. The method of claim 60, in which estramustine phosphate is administered in combination with one or more other chemotherapeutic agents.

62. The method of claim 11, wherein said anti-microtubule agent is a taxane.

63. The method of claim 11, wherein said anti-microtubule agent is a liposome encapsulated taxane.

64. The method of claim 63, wherein said liposome encapsulated taxane is liposome encapsulated paclitaxel.

65. The method of claim 9, wherein another drug in said regimen is CPT-11.

66. The method of claim 9, wherein another drug in said regimen is doxorubicin.
67. The method of claim 9, wherein another drug in said regimen is etoposide.

68. The method of claim 9, wherein another drug in said regimen is navelbine.

69. The method of claim 9, wherein another drug in said regimen is vinblastine.

70. A method of potentiating the therapeutic benefit of a multi-drug chemotherapeutic regimen wherein one drug in the regimen comprises a taxane, and wherein another drug in the regimen comprises estramustine phosphate, and wherein estramustine phosphate is administered intravenously at a dosage exceeding 950 mg/m².

71. The method of claim 15, wherein said anti-microtubule agent is a taxane.

72. The method of claim 20, wherein said anti-microtubule agent is a taxane.

73. A method of administering estramustine phosphate, wherein estramustine phosphate is first encapsulated within liposomes, and then administered intravenously.

74. A formulation of estramustine phosphate, wherein estramustine is encapsulated within liposomes.

75. A method of treating cancer wherein liposome-encapsulated estramustine
phosphate is administered.

76. A chemotherapeutic agent consisting of estramustine phosphate encapsulated within liposome.

77. A method of treatment for prostate cancer, wherein liposome-encapsulated estramustine phosphate is administered.

78. A method of treatment for breast cancer, wherein liposome-encapsulated estramustine phosphate is administered.


80. A method of treatment for pancreatic cancer, wherein liposome-encapsulated estramustine phosphate is administered.

81. A method of treatment for colorectal cancer, wherein liposome-encapsulated estramustine phosphate is administered.

82. A method of treatment for ovarian cancer, wherein liposome-encapsulated estramustine phosphate is administered.

83. A method of treatment for melanoma, wherein liposome-encapsulated
84. A formulation according to claim 74 intended for intravenous administration.

85. A product comprising estramustine phosphate suitable for intravenous administration and one or more chemotherapeutic agents, as a combined preparation for simultaneous, separate or sequential use in anticancer therapy.

86. A product according to claim 85, wherein said one or more chemotherapeutic agents are selected from the group consisting of CPT-11, doxorubicin, etoposide, navelbine and a taxane derivative.

87. A product according to claim 85, wherein said estramustine phosphate suitable for intravenous administration is used as a single dosage infusion exceeding 1300 mg.

88. A product according to claim 85, wherein said estramustine phosphate suitable for intravenous administration is used as a single dosage infusion exceeding 950 mg/m².

89. A product according to anyone of claims 85 to 88 for the treatment of prostate cancer, breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer or cancers of the brain.
90. A method for the treatment of prostate cancer, breast cancer, melanoma, lung cancer, pancreatic cancer, colorectal cancer, ovarian cancer and cancers of the brain, comprising the administration of a product according to anyone of claims from 85 to 88.
Fig. 1

Conc. (μmol/l)

- 1000 mg
- 1000 mg/m2
- 1500 mg/m2
Fig. 2

Conc. (nmol/l)

- 1000 mg
- 1000 mg/m²
- 1500 mg/m²

Time (h)
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : A61K 31/56, 31/65, 31/44, 31/335, 9/127

US CL : 514/182, 152, 283, 449, 452; 424/450

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 514/182, 152, 283, 449, 452; 424/450

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Please See Extra Sheet.

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tbody>
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<td>Y</td>
<td>US, 5,077,056 A (BALLY et al) 31 December 1991, see particularly column 8 and the claims.</td>
<td>1-90</td>
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<tr>
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<td>US 5,424,073 A (RAHMAN et al) 13 June 1995, see entire document.</td>
<td>1-90</td>
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<td>Y</td>
<td>US 5,616,341 A (MAYER et al) 01 April 1997, see columns 5 and 6 and the claims.</td>
<td>1-90</td>
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</tbody>
</table>

Further documents are listed in the continuation of Box C. See patent family annex.

Date of the actual completion of the international search

18 MAY 1999

Date of mailing of the international search report

03 JUN 1999

Name and mailing address of the ISA/US Commissioner of Patents and Trademarks

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<table>
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<tr>
<th>Category</th>
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</table>
B. FIELDS SEARCHED
Electronic data bases consulted (Name of data base and where practicable terms used):

APS, STN(REG, CA, BIOSIS, MEDLINE, DRUGU, EMBASE)
search terms: estramustine, melanoma, prostate, breast, lung, pancreas, colorectal, colon, ovarian, brain, cancer, tumor, taxane, taxol, paclitaxel, cpa-11, doxorubicin, etoposide, navelbine, vinorelbine, vinblastine, liposome