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| (54) Title:                           | TUMORICIDAL ACTIVITY OF BENZOQUINONOID ANSAMYCINS AGAINST PROSTATE CANCER AND PRIMITIVE NEURAL MALIGNANCIES |

| (57) Abstract                         | The invention provides methods and compositions for treating selected human malignancies including use of certain ansamycin benzoquinones. |
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TUMORICIDAL ACTIVITY OF BENZOQUINONOID ANSAMYCINS

AGAINST PROSTATE CANCER AND PRIMITIVE NEURAL MALIGNANCIES

BACKGROUND OF THE INVENTION

Benzoquinonoid ansamycin antibiotics were isolated in the late 1970s from the culture broths of several actinomycete species. Their unusual ansa bridge structure generated considerable interest, and a number of compounds including herbimycin A (HA) and geldanamycin (GA) were screened as possible anti-retroviral and anti-tumor agents. Results against the usual test cell lines of the time such as LL210 leukemia and P-388 were disappointing. The concentrations required for anti-tumor activity were quite toxic in the whole animal and further development was abandoned.

A new wave of enthusiasm occurred in the late 1980s when it was noted that HA was able to revert the phenotype of tyrosine kinase oncogene-transformed cell lines at quite modest concentrations. Inhibition of angiogenesis and induction of differentiation in a number of model systems were also reported. HA and GA have been shown to induce differentiation in a number of in vitro model systems, reportedly due to their inhibition of src-family protein tyrosine kinases.

Pediatric and adult cancers of primitive neural derivation, whether metastatic, locally disseminated, or recurrent, are among those malignancies most refractory to cure by current multi-modality treatment regimens. New approaches to treatment are clearly needed.

SUMMARY OF THE INVENTION

The invention provides a method for treating human malignancies selected from the group comprising primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma. The method includes administering
an effective dosage of an ansamycin benzoquinone to an animal which has the malignancy. The administration is preferably parenteral, such as intravenous.

The effective dosage is usually selected from a range of about 0.1 milligram of drug per kilogram body weight of the recipient animal (mg/kg) to about 20 mg/kg. Preferably, the effective dosage is selected from a range of about 1 mg/kg to about 10 mg/kg. More preferably, the effective dosage is about 5 mg/kg.

The ansamycin benzoquinone is typically selected from a group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, and macbecin I and its derivatives including macbecin II. The general chemical formula for the ansamycin benzoquinones is displayed in Figure 1. As noted, the side groups $R_1$, $R_2$, $R_3$ and $R_4$ are typically a hydrogen, a hydroxyl, an alkyl or an alkoxy. The term "alkyl" refers to substituents that are saturated hydrocarbon radicals. The alkyl groups may be straight-chain or branched-chain, limited only by steric hindrance. Shorter alkyl groups, such as 1-4 carbon atoms are preferred. The term "alkoxy" is used to refer to an alkyl radical which also bears an oxygen substituent that is capable of covalent attachment to another hydrocarbon radical (for example, a methoxy or ethoxy group). As with alkyl groups, shorter alkoxy groups are preferred. The term "independently selected" is used to indicate that the various $R$ groups, $R_1$ to $R_4$, may be identical or different.

The invention includes a method for treating human malignancies selected from the group comprising primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma comprising parenterally administering to a human an effective dosage of an ansamycin benzoquinone selected from the group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, macbecin I and its derivatives including macbecin II. The effective dosage is an amount sufficient to ameliorate symptoms or signs of the cancer. The amount is generally a range of from about 0.1 mg/kg to about 20 mg/kg.
The geldanamycin derivatives are preferably those in the group consisting of 17-des-O-methylgeldanamycin, geldanamycin acetate, 7',8'-benzodemethoxygeldanoxazone, hydrogeldanamycin-18,21-diacetate, 7'-(and 8')-fluorodemethoxygeldanazine, 7'-bromodemethoxygeldanoxazinone, 19-piperidinogeldanamycin, geldampicin, 8'-bromodemethoxygeldanoxazinone, 7'(or 8')-fluorodemethoxygeldanazine, and 17-amino-17-demethoxygeldanamycin.

The invention further includes a pharmaceutical composition having at least one pharmaceutically acceptable excipient and an amount of an ansamycin benzoquinone effective to treat cancer in a mammal to whom at least one dose of the composition is administered. The cancer is selected from a group including primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma.

Typically, the mammal is a human and the amount of ansamycin benzoquinone is selected from a range of from about 0.1 mg/kg to about 20 mg/kg. Preferably, the amount of ansamycin benzoquinone is selected from a range of from about 1 mg/kg to about 10 mg/kg. More preferably, the amount of ansamycin benzoquinone is about 5 mg/kg. The composition is preferably adapted for parenteral administration such as intravenous administration.

The ansamycin benzoquinone component of the composition is usually selected from a group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, macbacin I and its derivatives including macbacin II. The geldanamycin derivative is preferably selected from the group specified above.

BRIEF DESCRIPTION OF THE FIGURES

Fig. 1 is the chemical structure of ansamycin benzoquinones.

Fig. 2 graphically shows that benzoquinonoid ansamycins inhibit cell proliferation and survival in a dose dependent fashion.
Fig. 3 is a photograph of a DNA electrophoresis assay showing that cycloheximide inhibits HA-induced DNA degradation.

Fig. 4 demonstrates lack of benzoquinonoid ansamycin toxicity to primary neurons in culture.

Fig. 5 shows HA inhibition of prostate tumor growth in nude mice.

Fig. 6 shows that systemic treatment with HA of tumor-bearing mice inhibits tumor growth.

DESCRIPTION OF THE PREFERRED EMBODIMENT

Metastatic and locally disseminated cancers of primitive neural derivation remain among the most refractory to treatment. The invention shows that benzoquinonoid ansamycins possess very potent cytotoxic activity in vitro against a select range of human tumor lines and in vivo in a mouse model of certain human tumors. The invention also found little toxicity on primary neuron cultures, several hematopoietic cell lines and a mouse fibroblast line. Cell lines of the pediatric cancers neuroblastoma, neuroepithelioma and medulloblastoma were used as model systems in which to explore the therapeutic potential of ansamycin benzoquinones.

The benzoquinonoid ansamycins are potent cytotoxins in vitro against a panel of highly malignant human tumor cell lines possessing primitive neural features. Proliferation and/or survival of fibroblasts, primary neuronal cultures and several leukemia cell lines are unaffected at concentrations resulting in greater than 99% cell loss in sensitive lines. The tumorigenicity in nude mice of sensitive cell lines can also be markedly reduced by either systemic or topical administration of these agents without apparent toxicity to the whole animal. The cytotoxic action of these ansamycins is initiated very rapidly, is irreversible and clearly distinct from the delayed inhibition of src-family kinases that has been previously reported. Due to their potency, relative selectivity and novel mechanism(s) of action, these drugs can prove clinically useful in the therapy of a number of human cancers, particularly those of neural derivation.

Preparation of derivatives involves conventional chemical techniques and is within the skill of the ordinary artisan. The ansamycin benzoquinones, such as HA and GA, are available from the National Cancer Institute (NCI) Repository for investigational use. They are commercially available from Bethesda Research Lab (BRL) in Rockville, Maryland.

When referring to treating a cancer, improved or decreased symptoms are included. That is, "treatment" is not limited to an objective regression of tumor size, but also includes the patient's report of his subjective status.

An effective amount of the compound is that amount which provides subjective relief of a symptom, decrease in the tumor burden, or decrease in an identifiable tumor marker. Administration of the compound is by any medically or pharmaceutically accepted route. Typically, the parenteral route is preferred. Examples include intravenous, intramuscular, and subcutaneous administration. Topical application is also effective. The dosing range is preferably from about 0.1 to about 20 mg/kg/day based on the patient's body weight. More specifically, the range is from about 1 to about 10 mg/kg/day.

The dosage of the benzoquinonoid ansamycin depends on many factors that are well known to those skilled in the art. The factors include, for example, the route of administration, the potency of the particular compound, the
condition being treated, and the patient's age, weight, and general state of health including cardiac, hepatic and renal function.

A preferred dosing schedule is a parenteral or topical dose usually once or twice a day for a limited period, typically about 5 to 10 days. The schedule is preferably repeated at a suitable interval, usually at about 14 to 28 days with the first day of the first cycle counted as day one. Alternatively, the doses can be given every other day or every three to four days for a total of about three to ten doses. A preferred parenteral route is intravenous (IV). Preferably, the IV dose is infused over about 30 to 60 minutes.

Compositions of the present invention are presented for administration to humans and animals in unit dosage forms, such as tablets, capsules, pills, powders, granules, aqueous solutions or suspensions and water-in-oil emulsions containing suitable quantities or formulations of a benzoquinonoid ansamycin.

The term "unit dosage form" refers to physically discrete units suitable as unitary dosages for human subjects and animals, each unit containing a predetermined quantity of active material calculated to produce the desired pharmaceutical effect in association with the required pharmaceutical diluent, carrier or vehicle. Some examples of suitable unit dosage forms are tablets, capsules, pills, powder packets, wafers, granules, teaspoonfuls, tablespoonfuls, droppersful, ampoules, and vials.

The benzoquinonoid ansamycin may be combined or mixed with various solutions and other compounds as is known in the art. For example, it may be administered in water, saline or buffered vehicles. The benzoquinonoid ansamycin may be administered by any conventional method including oral, topical and parenteral (e.g., intravenous or intramuscular) injection. The treatment may consist of a single dose or a plurality of doses over a period of time. The drugs can be combined with appropriate doses of compounds. A pharmaceutically effective amount of the drugs can be employed
with a pharmaceutically acceptable carrier such as an additive or diluent.

A preferred formulation for parenteral use is an emulsion including the ansamycin benzoquinone and at least one emulsifying agent. For example, the vehicle Cremophor can be used as an emulsifier. Emulsifiers which are nonionic and which contain complex fatty acids are preferred. Additionally, detergents may be useful in preparing the emulsion. Another preferred formulation includes modifying the parent compound, such as HA or GA, to a salt which would be soluble in an aqueous solution. For example, esterification of reactive groups can be carried out to produce water soluble homologues or analogues of the ansamycin benzoquinone. Because the parent compounds tend to be insoluble in water, administration as an aqueous solution without modification of the parent is a lesser preferred, although a feasible, alternative.

EXAMPLES.

EXAMPLE 1.

Cell Cultures. CHP-100 cells were obtained from Dr. A. Evans (Children's Hospital of Philadelphia). The cell lines TC-32 and NIH3T3 were obtained from Dr. M. Tsokos (NCI, Laboratory of Pathology). All other cell lines used were purchased from the American Type Culture Collection ATCC, Rockville, MD). Primary neonatal rat cortical neurons were established using standard techniques and supplied by Dr. M. Koenig (Armed Forces Research Institute, Washington, DC). All cell lines were tested using a MycotectR kit (GIBCO Laboratories, Grand Island, NY) and found free of mycoplasma contamination. All culture media were supplemented with 10% fetal bovine serum (FBS) (Whittaker Bioproducts, Walkersville, MD).

The cell lines CHP-100, TC-32, IMR-32, SKNSH, CEM and HL-60 were grown in RPMI 1640 (Biofluids, Inc., Rockville, MD). The cell lines D283 Med, D341 Med, SKNMC, SK-MEL-1, SK-MEL-2, and RPMI 7951 were cultured in Eagle's MEM with 1% non-essential amino acids and 1mM sodium pyruvate (all from
Biofluids, Inc). NIH3T3 cells were cultured in Dulbecco's MEM with 4.5 g/L glucose (Biofluids, Inc). Herbimycin A (National Service Center NSC 305978) and geldanamycin (NSC 122750) were obtained from the Drug Synthesis and Chemistry Branch, National Cancer Institute (NCI), formulated as 2mg/ml stock solutions in dimethyl sulfoxide (Sigma Chemical Co., St. Louis, MO) and stored at 4°C in the dark.

Selective Cytotoxic Activity of HA and GA Against Primitive Neural Cell Lines. Data are graphed in Fig. 2 using the following legend. Open circles refer to the cell line CHP-100. Closed boxes refer to the cell line NIH 3T3. Panel A had HA added to the culture medium and ³H-thymidine incorporation was assayed 48 hours later. Panel B had GA added to the culture medium and ³H-thymidine incorporation was assayed 48 hours later. Panel C had HA added to the culture medium and MTT reduction was assayed 5 days later. Panel D had GA added to the culture medium and MTT reduction was assayed 5 days later. Data are expressed as percent cpm or optical density relative to control wells plated simultaneously without the addition of the drug. All points represent the mean of triplicate determinations. Standard deviations are less than 10%.

Figure 2, panel A demonstrates potent growth inhibitory activity of HA against the primitive neuroectodermal cell line CHP-100 as measured by inhibition of ³H-thymidine incorporation after two days of culture with the drug. This cell line is approximately 10-fold more sensitive in vitro to the inhibitory activity of HA than the non-tumorigenic mouse fibroblast cell line NIH3T3. Panel B demonstrates a similar pattern of cell type selectivity for the related ansamycin GA, and GA is approximately ten-fold more potent than HA.

Experiments were also carried out with these two cell lines assessing the effects of HA and GA on survival rather than inhibition of DNA synthesis. Automated analysis of MTT reduction by treated cells in a 96 well microtiter format allowed generation of dose response curves similar to panels A and B, but the endpoint was the relative number of
viable cells remaining after 5 days in culture. As panels C and D demonstrate, concentrations of HA or GA that result in complete cell loss with CHP-100 do not affect the fibroblast cell line. Relative potency and selectivity were similar by MTT analysis and thymidine incorporation analysis. The effect seen in MTT experiments was not simple inhibition of growth. Values actually declined from the time of initial plating, indicating cell loss (data not shown). Microscopic examination of treated wells prior to analysis also confirmed that only cellular debris remained.

**Cell Proliferation and Survival Studies.** Thymidine incorporation studies were performed as previously described (Whitesell L, Rosolen A, Neckers LM, "Episome-generated N-myc antisense RNA restricts the differentiation potential of proimitive neuroectodermal cell lines", *Mol. Cell. Biol.* 11: 1360-1371 (1991)) except that serial dilutions of HA or GA were added to triplicate wells in 100 µL of the appropriate growth medium for the cell line being tested.

Plates were cultured for 48 to 72 hours and then pulsed with [methyl-³H]-thymidine (Dupont, Boston, MA) for 4 hours, followed by automated cell harvest and liquid scintillation counting.

As an assay of relative viable cell number, mitochondrial reduction of 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT) was used (Alley MC, et al., "Feasibility of drug screening with panels of human tumor cell lines using a microulture tetrazolium assay", *Cancer Res.*, 48: 589-601 (1988)). Ninety-six well tissue culture plates were set-up as described by Alley, but cultured for 5 to 6 days. 20 µL of a 5 mg/ml solution of MTT (Sigma, St. Louis, MO) in phosphate buffered saline (PBS) was added to all wells and plates were incubated for 4 more hours in darkness.

Plates were centrifuged, medium removed and 150 µL of dimethyl sulfoxide (Sigma, St. Louis, MO) added to each well. After a 10 minute incubation in darkness with shaking, the optical density at 540 nm was determined using a Biokinetics plate reader (Model EL-312, Bio-Tek Instruments...
Inc., Winooski, VT). Optical density was used as a measure of the formazan concentration generated in each well.

EXAMPLE 2.

Characteristics of HA and GA cytotoxicity. Having demonstrated that HA and GA are actually cytotoxic to CHP100 cells, characteristics of this cytotoxicity was examined. Cell death can be an active or passive process depending on cell type and initiating stimulus. DNA degradation is often used as an endpoint for both processes. A requirement for ongoing protein synthesis in the transduction of cell death suggests an active type of program.

CHP-100 cells (5x10^6/10 cm tissue culture dish) were plated in the presence of HA (500 nM), 80 nM cycloheximide (a concentration previously determined to inhibit protein synthesis by 75% as measured by ^3H-leucine incorporation) or a combination of the two. Sixty-eight hours after plating, the cells were harvested by trypsinization and cell number and viability determined by trypan blue exclusion in a hemocytometer chamber. The cells were washed once in PBS and high molecular weight DNA prepared according to a previously reported method (Rodriguez-Tarduchy G, Lopez-Rivas A., "Phorbol esters inhibit apoptosis in IL-2 dependent T-lymphocytes", Biochem. Biophys. Res. Comm., 164: 1069-1075 (1989)).

Briefly, cell pellets were incubated at 37°C for at least 4 hours in 0.5mL of lysis buffer (200mM TRIS pH8.5, 100 mM EDTA, 50 µg/ml proteinase K, 1% SDS). The DNA was phenol extracted and the aqueous phase dialyzed overnight against 10 mM TRIS pH7.5, 1mM EDTA. The DNA was incubated at 37°C with 50 µg/mL RNase A. After 5 hours, 120 µg/mL proteinase K was added and the incubation continued for 5 more hours. DNA was extracted with phenol followed by chloroform and precipitated with ethanol/sodium acetate. After re-dissolving in water, 5 µg of DNA was loaded per lane on a 1.4% agarose gel and electrophoresed for 16 hours at 35 volts prior to staining with ethidium bromide and photography.
The lanes in Fig. 3 have the following meaning. CTRL indicates a control and no drugs were added. HA means that 500 nM HA were added. CHX means that 80 nM cycloheximide were added. CHX/HA means both drugs were added simultaneously. DNA was isolated and electrophoresed as described above. Degradation of high molecular weight DNA is apparent only in lane HA. Figure 3 demonstrates that protein synthesis is necessary for HA-induced cell death. In this experiment, cells were cultured in the presence of 500 nM HA with or without the non-specific protein synthesis inhibitor cycloheximide. After 68 hours, cells were harvested, assayed for viability and high molecular weight DNA was prepared. Viability of untreated cells was 82%, that of 80 nM cycloheximide-treated cells was 57%, that of HA-treated cells 26% and that of HA/cycloheximide-treated cells was 66%. Cycloheximide treatment clearly inhibited the cytotoxic activity of HA.

Analysis of high molecular weight DNA revealed substantial DNA degradation in the HA-treated cells (Figure 3, lane HA), but the nucleosomal cleavage characteristic of apoptosis was not seen. However, DNA degradation was abrogated by co-incubation of the cells with cycloheximide (lane CHX/HA). The quality of high molecular weight DNA in that lane is indistinguishable from that of either untreated (lane CTRL) or cycloheximide alone-treated cells (lane CHX). These findings confirm that HA is cytotoxic to sensitive cells and that this cytotoxicity requires active cellular participation.

To further evaluate the spectrum of activity displayed by HA and GA, a panel of primitive neural cell lines was screened using both inhibition of $^3$H-thymidine incorporation and reduction of MTT as endpoints. See Table 1 which is a summary of dose response data obtained by thymidine assay for a variety of both neural and non-neural cell lines. Non-neural cell types such as the hematopoietic lines HL-60 and CEM were quite resistant to growth inhibition by HA and GA. All primitive neuroectodermal cell lines examined, both peripheral nervous system-derived (e.g. CHP-100, SKNMC, TC-32)
and central nervous system-derived (e.g. D283 Med, D341 Med) have proven quite sensitive to HA and GA. Cells from a primary culture of the malignant pleural effusion of a patient with relapsed, heavily pre-treated Ewing's sarcoma were also sensitive to HA as determined by MTT assay (IC$_{95}$ = 400 nM).

Cells of more mature neural phenotype were relatively insensitive to the cytotoxic activity of HA. Thymidine incorporation data for the neuroblastoma cell lines IMR-32 and SKNSH are shown in Table 1. The highly differentiated rat pheochromocytoma cell line PC-12 was also relatively insensitive to HA by MTT assay (IC$_{95}$ > 500 nM).

Data are presented as bar graphs in Fig. 4. HA at 472 nM or GA at 40 nM was added to established neonatal rat cerebellar neuron cultures. Plates were incubated for a further 4 days and reduction of MTT dye reflecting viable cell number per well was then assayed as optical density at 540 nm. The height of the bars represents the mean absorbance of quadruplicate wells. Error bars indicate the standard deviation of the mean.

Figure 4 demonstrates that primary cultures of neonatal rat cerebellar neurons were not affected by culture in 472 nM HA or 40 nM GA. This experiment was also repeated several times with neonatal rat cerebral cortical neurons with the same results. Lastly, the melanoma cell lines SK-MEL-1, SK-MEL-2 and RPMI 7951 were also screened against HA by MTT assay. The one line found to be sensitive, RPMI 7951 (HA IC$_{50}$ = 125 nM) is reported to possess the most primitive, least differentiated phenotype as defined by Houghton et al., "Surface antigens of melanocytes and melanomas", J. Exp. Med., 156:1755-1766 (1982).
## TABLE 1

*Selectivity of growth inhibition by HA and GA*

Data are derived from dose response curves generated by quantification of $^3$H-thymidine incorporation by indicated cell lines 48 hours after plating in various concentrations of the indicated drug. Cell lines are grouped in decreasing order of sensitivity.

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*a* IC$_{50}$ = Concentration of drug resulting in 50% decrease in mean cpm of triplicate wells relative to untreated control wells.

*b* IC$_{95}$ = Concentration of drug resulting in a 95% decrement.

*c* N.D. = evaluation not done.

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### EXAMPLE 3.

**Tumorigenicity Studies.** CHP-100 cells were harvested from subconfluent monolayers and resuspended at 5x10$^7$ cells/mL in PBS. 100 µL were inoculated subcutaneously in the right and left inguinal areas of six week old male athymic nude mice (Frederick Cancer Research Facility, Frederick, MD) on day 0.

Therapy with HA or GA was initiated either on the day of tumor inoculation or 10 days post inoculation when palpable tumors were apparent. The drugs were administered either
topically by painting 5 μL of a 2 mg/mL solution of the drug in dimethyl sulfoxide on the site of tumor cell inoculation or systemically by intraperitoneal injection of drug formulated in dimethyl sulfoxide.

Control animals received identical treatments with the appropriate vehicle alone. Animals were sacrificed 21 days after cell inoculation and well encapsulated tumors were resected and weighed. All studies involving the use of mice were carried out under protocols reviewed and approved by the Animal Care and Use Sub-committee of the National Cancer Institute.

In Vivo Anti-tumor Activity. Benzoquinonoid ansamycins as potential chemotherapeutic agents in the treatment of selected human cancers were evaluated using a tumor xenograft/nude mouse model. Preliminary experiments in athymic rats have demonstrated that intravenous infusion of 4mg/kg (or 24mg/M²) of GA is well tolerated with no overt immediate or delayed toxicity.

Table 2 depicts the effects of either topical or systemic administration of drugs to animals at the time of tumor cell inoculation. Nude mice were inoculated with tumor cells subcutaneously and therapy begun on the same day. Topical therapy consisted of applying 5 μL of a 2 mg/mL solution of drug in dimethyl sulfoxide to the skin overlying the tumor inoculation site daily for 5 days. Systemic therapy consisted of intraperitoneal GA injection daily for 5 days at 4.5 mg/kg body weight, or about 90 μg. Tumor formation as defined by resection of a discrete mass of 100 mg or greater was assessed 21 days after cell inoculation. A marked reduction in subsequent tumor formation was evident following both HA and GA administration. The mean weight of tumors that formed in drug-treated animals was also significantly smaller. No overt toxicity as evidenced by weight loss, decreased motor activity or local skin reaction was noted.

On one occasion, the tumors which formed despite topical GA treatment were resected aseptically, disaggregated in tissue culture medium and grown for 9 days in vitro in the presence or absence of GA. The cell cultures recovered in
this fashion grew well and, importantly, despite previous in vivo treatment with GA they remained fully sensitive to its cytotoxic action in vitro. This finding suggests that the failure of topical GA to inhibit tumor formation in all animals was a result of inadequate exposure to active drug and not acquired tumor cell resistance.

<table>
<thead>
<tr>
<th>TABLE 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>HA and GA decrease the tumorigenicity of CHP 100 in vivo</strong></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Drug</th>
<th>Topical Tumors</th>
<th>Systemic Tumors</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean Weight (mg)</td>
<td>Mean Weight (mg)</td>
</tr>
<tr>
<td>DMSO Control 10/10(^a)</td>
<td>357 (49)(^b)</td>
<td>10/10 (436 (93))</td>
</tr>
<tr>
<td>Geldanamycin 5/10</td>
<td>302(^c) (97)</td>
<td>6/10 (252(^e) (53))</td>
</tr>
<tr>
<td>Herbimycin A 7/10</td>
<td>221(^d) (49)</td>
<td>N.D.(^f) (N.D.)</td>
</tr>
</tbody>
</table>

\(^a\) Number of tumors formed/number of sites inoculated.
\(^b\) Numbers in parentheses refer to the standard error of the mean value.
\(^c\) Comparison to control by Student's t test performed on log-transformed data p=0.037
\(^d\) As above, p=0.014.
\(^e\) As above, p=0.042.
\(^f\) N.D.=Experiment not performed.

In a similar experiment, nude mice were given a subcutaneous injection of human prostate cancer cells. These mice were treated with 5 mg/kg of HA intraperitoneally (IP). See Fig. 5. Five million human prostate cancer cells were inoculated subcutaneously. Treatment began on the day of tumor cell inoculation and was repeated every other day for a total of 3 doses. Treatment consisted of 5 mg/kg herbimycin A as an intraperitoneal injection. The animals were sacrificed and the tumors were resected and weighed two weeks after inoculation.

The ability of HA to inhibit the growth of established tumors was also examined in the nude mouse model. Initiation
of HA treatment 10 days after tumor cell inoculation at a dose of 1.5 mg/kg, given intraperitoneally every third day for 4 doses, resulted in decreased tumor mass as determined on the 21st day after tumor cell inoculation (Fig. 6). Nude mice with established subcutaneous tumors received intraperitoneal injections of either HA 1.5 mg/kg or an equivalent volume of vehicle on days 10, 13, 16 and 19 post tumor cell inoculation. Each point depicts the weight of an individual tumor mass resected on day 21. The dashed horizontal line indicates the mean weight of tumors resected from the vehicle-treated mice. Comparison of control to HA-treated tumor weights by Student's t test on log-transformed data yields p=0.057.

Because HA and GA appear to be selectively cytocidal, it is of great interest to define their precise mechanism of action from the points of view of both basic biology as well as clinical drug development. For example, understanding the mechanism of action of these drugs may allow for identification of other sensitive tumor types. See Whitesell, et al., "Benzoxquinonoid Ansamycins Possess Selective Tumoricidal Activity Unrelated to src Kinase Inhibition," Cancer Research 52:1721-1728 (April 1, 1992). Melanoma cell lines are being evaluated as an approach to defining the developmental/phenotypic specifically of these ansamycins.

In addition, a panel of GA derivatives was screened for potency and selectivity in tissue culture model systems. See Table 3 which lists drugs which were active against tumor development in culture. The structure-activity relationships generated should pinpoint the critical features required for activity and may suggest likely intracellular targets for these drugs.

Finally, the xenograft tumor results reported suggest promising in vivo anti-tumor activity for the benzoxquinonoid ansamycins. The detection of any anti-tumor activity in vivo is encouraging given the near complete lack of baseline pharmacologic information. The antibiotics discussed here are small, lipophilic molecules that should be relatively easy to produce in bulk and, following systemic administration, should readily penetrate solid tumors and even the blood-brain
Compounds such as these may well serve as models for the development of a generation of novel drugs possessing both biologic specificity and tumoricidal pharmacology.

TABLE 3

5
17-des-O-methylgeldanamycin,
geldanamycin acetate,
7',8'-benzodemethoxygeldanoxazone,
hydrogeldanamycin-18,21-diacetate,
7'-(and 8')-fluorodemethoxygeldanazine,
10
7'-bromodemethoxygeldanoxazinone,
19-piperidinogeldanamycin,
8'-bromodemethoxygeldanoxazinone,
7'(or 8')-fluorodemethoxygeldanazine, and
17-amino-17-demethoxygeldanamycin.

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The preceding description and examples are illustrative, and all references are incorporated by reference. Those skilled in the art will promptly recognize appropriate variations from the procedures as to dosing, scheduling, indications and toxicity. Thus, the invention is limited only by the scope of the appended claims.
WHAT IS CLAIMED IS:

1. A method for treating human malignancies selected from the group comprising primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma comprising administering an effective dosage of an ansamycin benzoquinone to an animal which has the malignancy.

2. The method of claim 1 wherein the administration is parenteral.

3. The method of claim 2 wherein the administration is intravenous.

4. The method of claim 1 wherein the effective dosage is selected from a range of about 0.1 mg/kg to about 20 mg/kg.

5. The method of claim 4 wherein the effective dosage is selected from a range of about 1 mg/kg to about 10 mg/kg.

6. The method of claim 5 wherein the effective dosage is about 5 mg/kg.

7. The method of claim 1 wherein the ansamycin benzoquinone is selected from a group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, and macbacin I and its derivatives.

8. The method of claim 1 wherein the ansamycin benzoquinone has the formula
wherein $R_1$, $R_2$, $R_3$ and $R_4$ are independently selected from the group consisting of H, lower alkyl, lower alkoxy, and hydroxy.

9. The method of claim 8 wherein $R_1$, $R_2$, $R_3$ and $R_4$ are independently selected from the group consisting of H, methyl, methoxy, and hydroxy.

10. A method for treating human malignancies selected from the group comprising primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma comprising:

   parenterally administering to a human an effective dosage of an ansamycin benzoquinone selected from the group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, and macbacin I and its derivatives, said effective dosage being an amount sufficient to ameliorate symptoms or signs of said cancer, the amount being in a range of from about 0.1 mg/kg to about 20 mg/kg.

11. The method of claim 10 wherein the ansamycin benzoquinone is a geldanamycin derivative selected from the group consisting of:

   17-des-O-methylgeldanamycin,
   geldanamycin acetate,
   7',8'-benzodemethoxygeldanoxazone,
   hydrogeldanamycin-18,21-diacetate,
   7'-(and 8')-fluorodemethoxygeldanazine,
   7'-bromodemethoxygeldanoxazinone,
   19-piperidinogeldanamycin,
   8'-bromodemethoxygeldanoxazinone,
   7'(or 8')-fluorodemethoxygeldanazine, and
   17-amino-17-demethoxygeldanamycin.

12. A pharmaceutical composition comprising at least one pharmaceutically acceptable excipient and an amount of an ansamycin benzoquinone effective to treat cancer in a mammal to whom at least one dose of said composition is administered wherein said cancer is selected from a group comprising
primitive neuroectodermal tumors, prostate cancer, melanoma, and metastatic Ewing's sarcoma.

13. The method of claim 12 wherein the ansamycin benzoquinone has the formula

![Chemical Structure](image)

wherein R₁, R₂, R₃ and R₄ are independently selected from the group consisting of H, lower alkyl, lower alkoxy, and hydroxy.

14. The method of claim 13 wherein R₁, R₂, R₃ and R₄ are independently selected from the group consisting of H, methyl, methoxy, and hydroxy.

15. The composition of claim 12 wherein the mammal is a human.

16. The composition of claim 12 wherein the amount of ansamycin benzoquinone is selected from a range of from about 0.1 mg/kg to about 20 mg/kg.

17. The composition of claim 16 wherein the amount of ansamycin benzoquinone is selected from a range of from about 1 mg/kg to about 15 mg/kg.

18. The composition of claim 17 wherein the amount of ansamycin benzoquinone is about 5 mg/kg.

19. The composition of claim 12 wherein the composition is adapted for parenteral administration.

20. The composition of claim 19 wherein the composition is adapted for intravenous administration.
21. The composition of claim 12 wherein the ansamycin benzoquinone is selected from a group consisting of geldanamycin and its derivatives, herbimycin A and its derivatives, and macbacin I and its derivatives.

22. The composition of claim 12 wherein the ansamycin benzoquinone is a geldanamycin derivative selected from the group consisting of

- 17-des-O-methylgeldanamycin,
- geldanamycin acetate,
- 7',8'-benzodemethoxygeldanoxazine,
- hydrogeldanamycin-18,21-diacetate,
- 7'-(and 8')-fluorodemethoxygeldanazime,
- 7'-bromodemethoxygeldanoxazinone,
- 19-piperidinogeldanamycin,
- 8'-bromodemethoxygeldanoxazinone,
- 7'(or 8')-fluorodemethoxygeldanazime, and
- 17-amino-17-demethoxygeldanamycin.
FIG. 1. (PRIOR ART)

FIG. 4.
SUBSTITUTE SHEET
FIG. 2A.

FIG. 2B.

SUBSTITUTE SHEET
FIG. 2C.

FIG. 2D.
**FIG. 5.**

![Graph showing tumor mass (grams) for control and Herbimycin A groups.]

**FIG. 6.**

![Graph showing tumor weight (grams) for vehicle and Herbimycin A groups.]

**SUBSTITUTE SHEET**