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- (73) Patenthaver: BeyondSpring, Inc., Sertus Chambers, PO Box 2547, Cassia Court, Camana Bay, Grand Cayman, Caymanøerne
- (72) Opfinder: HUANG, Lan, No. 9 East Liaohe Road, Dalian Development Zone, Dalian, Liaoning 116001, Kina
- (74) Fuldmægtig i Danmark: Patrade A/S, Ceresbyen 75, 8000 Århus C, Danmark
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# WO-A1-2005/077940

A. C. MITA ET AL: "Phase II study of docetaxel with or without plinabulin (NPI-2358) in patients with non-small cell lung cancer (NSCLC).", JOURNAL OF CLINICAL ONCOLOGY, vol. 28, no. 15\_suppl, 20 May 2010 (2010-05-20), pages 7592-7592, XP055394409, US ISSN: 0732-183X, DOI: 10.1200/jco.2010.28.15\_suppl.7592 MICHAEL MILLWARD ET AL.: 'phase 1 study of the novel vascular disrupting agent plinabulin (NPI-2358) and docetaxel' THE JOURNAL OF NEW ANTICANCER AGENTS vol. 3, no. 30, 16 February 2011, pages 1065 - 1073, XP035052851 DOI: 10.1007/S10637-011-9642-4

MILLWARD M. ET AL: "Phase 1 study of the novel vascular disrupting agent plinabulin (NPI-2358) and docetaxel", INVESTIGATIONAL NEW DRUGS; THE JOURNAL OF NEW ANTICANCER AGENTS, KLUWER ACADEMIC PUBLISHERS, BO, vol. 30, no. 3, 16 February 2011 (2011-02-16), pages 1065-1073, XP035052851, MILLWARD M. ET AL.: "Phase I trial of NPI-2358 (a novel vascular disrupting agent) plus docetaxel.", J. CLIN. ONCOL., vol. 27, no. 15S, May 2009 (2009-05), page 3571,

SATO I. ET AL.: "Prediction of docetaxel monotherapy-induced neutropenia on the monocyte parentage.", ONCOLOGY LETTERS, vol. 3, 2012, pages 860-864,

AAVV: "ICH Harmonised Tripartite Guideline: Nonclinical Exaluation for Anticancer Pharmaceuticals", 2009, International Conference on Harmonsation of Technical Requirements for Registration of Pharmaceuticals for Human Use vol. S9

AAVV: "Febrile Neutropenia", JAMA ONCOLOGY, vol. 3, no. 12, 2017, page 251,

CRAWFORD J. ET AL.: "Myeloid Growth Factors, version 2.2017", J. NATL. COMPR. CANC. NETW., vol. 15, no. 12, 2017, pages 1520-1541,

# **DK/EP 3076972 T3**

AAVV: "Common Terminology Crteria for Adverse Events (CTCAE) v5.0", 27 November 2017 (2017-11-27), MedDRA

# **DESCRIPTION**

#### **TECHNICAL FIELD**

[0001] The present invention relates to medical field and in particular to Plinabulin for use in reducing the neutropenia rate of grade 3 or 4 neutropenia.

#### **BACKGROUND ART**

[0002] Cancer has overtaken cardiovascular diseases as the leading cause of disease incurred death in the world. In the US over 1 million people get cancer each year, while in China, over 3 million people get cancer each year. 45% of new cancer cases in the world occur in China. The most widely used treatment for cancer is chemotherapy agents, such as taxanes. However, due to taxane's unbearable side effects, including high neutropenia rates (grade 3-4 at 30-40%), its dose has to be decreased during the use or even its use has to be terminated, and thus lowering down its efficacy and treatment duration in cancer patients.

[0003] Lung cancer is the leading cause of cancer-related mortality in the United States, China, and the world. Non-small cell lung cancer (NSCLC) accounts for approximately 80% of all cases of lung cancer. By the time most patients are diagnosed with NSCLC, the disease is already advanced. Standard approved therapies for advanced NSCLC generally include successive lines of chemotherapy agents including platins, taxanes, vinca alkyloids, pemetrexed, and/or epidermal growth factor receptor (EGFR) inhibitors.

**[0004]** Docetaxel is a taxane compound approved as second line treatment of NSCLC in the US, European Union, China and multiple other countries. Docetaxel functions by disrupting the microtubule network in cells. It is generally administered as a 1-hour intravenous (IV) infusion once every 3 weeks at a dose of 75 mg/m<sup>2</sup> with dexamethasone premedication to minimize the probability of hypersensitivity reactions and fluid retention. In 2 randomized trials with docetaxel in patients with NSCLC previously treated with a platinum-based chemotherapy regimen, the median overall survival (OS) ranged from 5.7 to 7.5 months.

[0005] The most common adverse reactions with docetaxel include infections, neutropenia, anemia, febrile neutropenia, hypersensitivity, thrombocytopenia, and etc. Several additional chemotherapeutic agents have been approved as second line therapies for Stage IIIb/IV NSCLC (pemetrexed, erlotinib, and gefitinib), but have clinically equivalent OS outcomes. In wide type NSCLC patients (stage IIIb/IV, second line), OS for docetaxel cohort (75 mg/m²) was 8.2 M, much longer than OS of some other drugs. Thus docetaxel is still the treatment of choice in second line NSCLC treatment.

**[0006]** Mita AC et al 2010, Journal of Clinical Oncology 15Suppl (A7592) discloses the use of plinabulin in combination with docetaxel for treating cancer. Vainas (2012); British J. Cancer 107:814-822 teaches that the neutropenia, in particular severe one (grade 4) can be avoided by administering G-CSF.

[0007] Since the treatment of cancers is still unsatisfied, there is a clear unmet medical need for additional anticancer agents in cancer patients such as those with advanced NSCLC.

#### **SUMMARY OF THE INVENTION**

**[0008]** The invention is defined in the claims and provides plinabulin for use in reducing the neutropenia rate of a grade 3 or 4 neutropenia in a subject being administered with 75 mg/m² docetaxel, wherein the plinabulin is administered intravenously at about 20 mg/m² to about 30 mg/m² wherein the plinabulin is administered within 1 h and 24 h after administering the docetaxel

#### SUMMARY OF TECHNICAL TEACHINGS

**[0009]** The technical information set out below may in some respects go beyond the scope of the invention, which is defined exclusively by the appended claims. The additional technical information is provided to place the actual invention in a broader technical context and to illustrate possible related technical developments. Such additional technical information which does not fall within the scope of the appended claims, is not part of the invention.

**[0010]** Also described is a pharmaceutical combination comprising an active ingredient (a) of a taxane compound; and an active ingredient (b) of plinabulin.

[0011] The taxane compound comprises paclitaxel, docetaxel, and abraxane.

[0012] The combination can consist of docetaxel and plinabulin.

[0013] Also described is a use of the pharmaceutical combination for preparing drugs for treating and/or preventing cancer.

**[0014]** The cancer can be selected from the group consisting of lung cancer, colon cancer, liver cancer, breast cancer, prostate cancer, and multiple myeloma.

**[0015]** Also described is a use of a composition, a kit or a mixture comprising an active ingredient (a) of taxane and an active ingredient (b) of plinabulin for treating and/or preventing cancer.

[0016] The cancer has a tumor size of > 3cm, > 5cm, or >7cm.

[0017] Also described is a kit which contains

- 1. (i) a first container containing a first medicament which comprises a taxane compound as an active ingredient (a) and an optional pharmaceutically acceptable carrier; and
- 2. (ii) a second container containing a second medicament which comprises plinabulin as an active ingredient (b) and an optional pharmaceutically acceptable carrier;
- 3. (iii) an optional instruction recording an administration of the active ingredient (a) in combination with the active ingredient (b) for treating and/or preventing cancer.

[0018] Preferably, the instruction indicates that plinabulin has to be injected 1-24 hours after administrating taxane.

[0019] Preferably, there are 8 first medicament at 20 mg per vial and 2 second medicament at 80 mg per vial in a single kit.

[0020] Also described is a pharmaceutical composition, wherein it comprises:

an active ingredient (a) of taxane;

an active ingredient (b) of plinabulin; and

(c) a pharmaceutical acceptable carrier.

[0021] The ratio (mg: mg) of active ingredient (a) to active ingredient (b) can be 1: 100 to 50: 1; preferably, 1.5:1 to 4:1.

[0022] The total amount of the active ingredient (a) and the active ingredient (b) is 1-99wt%; and more preferably, 5-

90wt% of the composition.

[0023] Also described is a use of plinabulin for preparation of a medicine used to reduce side effect of taxane.

[0024] The side effect comprises neutropenia, anemia, febrile neutropenia, thrombocytopenia.

**[0025]** Also described is a method for reducing side effect of taxane, wherein it comprises a step of administrating plinabulin to the subject in need prior to, at same time or after administrating a taxane compound.

[0026] Also described is a method for treating and/or preventing cancer, comprising the following steps: administrating to a mammal in need thereof an active ingredient (a) of taxane and an active ingredient (b) of plinabulin

[0027] Preferably, administrating the active ingredient (a) first and then administrating and the active ingredient (b).

[0028] Also described is a method of use of plinabulin and docetaxel combination for treating cancer in a subject in that plinabulin has to be injected after docetaxel for the enhanced efficacy to treat cancer in the range of 1-24 hours after.

[0029] The subject is a patient of large tumor in various cancer types.

[0030] The subject is a patient of stage IIIb/IV NSCLC having at least one lung primary tumor > 3 cm, preferably > 5 cm, and more preferably > 7 cm.

[0031] Also described is a method of use in taxanes combined with plinabulin to decrease the toxicity in taxanes, especially its grade 3 and 4 neutropenia.

[0032] Also described us a method of use of plinabulin and docetaxel in NSCLC patient treatment in decreasing of docetaxel's neutropenia rate at all grades, and decreasing G-CSF use.

#### **DESCRIPTION OF FIGURES**

#### [0033]

Fig. 1 shows chemical structure of Plinabulin.

Fig. 2 shows mean percent weight change in Example 1 (NSCLC MV522 tumor model). Plinabulin (NPI-2358) addition to docetaxel reduced the docetaxel associated animal weight loss independent of each drug usage sequence (2 hour separation).

Fig. 3 shows mean tumor volume in Example 1 (NSCLC MV522 tumor model). The addition of Plinabulin (NPI-2358) enhanced the antitumor activity of docetaxel. The synergistic effect was more prominent in the groups receiving docetaxel first followed by NPI-2358 2 hours later.

**Fig. 4A, Fig. 4B and Fig. 4C** show mean tumor volume in Example 2 (NSCLC MV522 tumor model). The anti-tumor activity enhancement effect of Plinabulin (NPI-2358) added to docetaxel were similar in groups receiving docetaxel first followed by plinabulin 1, 4, or 24 hours later.

**Fig. 5** shows that side effect such as neutropenia at different grades of severity was reduced by adding plinabulin to docetaxel treatment of NSCLC patient. Both 20 mg/m<sup>2</sup> or 30 mg/m<sup>2</sup> plinabulin added to docetaxel decreased neutropenia rate of docetaxel (D: Docetaxel; DN: Plinabulin and Docetaxel)

**Fig. 6** shows overall survival Kaplan-Meier Curve in NSCLC patients (stage IIIb/IV, at least 1 prior chemotherapy). Arm DN: 30 mg/m<sup>2</sup> Plinabulin and 75 mg/m<sup>2</sup> docetaxel; Arm D: 75 mg/m<sup>2</sup> docetaxel. OS benefit increases in the Arm DN vs. Arm D as tumor size increases.

#### **DETAILED DESCRIPTION**

**[0034]** The technical information set out below may in some respects go beyond the scope of the invention, which is defined exclusively by the appended claims. The additional technical information is provided to place the actual invention in a broader technical context and to illustrate possible related technical developments. Such additional technical information which does not fall within the scope of the appended claims, is not part of the invention.

#### Context

[0035] Through comprehensive and intensive research and screening, the inventor has unexpectedly developed a method for treatment of cancer by using plinabulin and docetaxel in combination. The experiments have shown optimum efficacy benefit using NSCLC MV522 animal model. The inventor has also discovered a series of combination method of using plinabulin and docetaxel to achieve optimum efficacy and safety benefit using NSCLC MV522 animal model experiments.

[0036] In one embodiment according to the present invention, the optimum combination can dramatically decrease side effects of docetaxel in both NSCLC MV522 animal model, and NSCLC patients, and decrease G-CSF use in NSCLC patients, which are unexpected findings. No other VDA compound has been discovered with this effect in combination with docetaxel in cancer patient treatment.

[0037] Also described is that the optimum combination method achieved the most efficacy benefit in OS extension in a uniquely defined large tumor NSCLC patients, and in large tumor NSCLC MV522 animal model. No other VDA or anti-angiogenesis agent has been discovered in better efficacy in this unique defined large tumor NSCLC patient population.

#### Taxane compounds and derivatives thereof

[0038] In the combinations described herein, one important active ingredient is taxane compound or the derivatives thereof.

[0039] The term "taxane compound" or "taxane" means a member of taxane family that has anti-cancer activity similar to paclitaxel based on the same or similar mechanism of paclitaxel. The exemplary taxane compounds include but are not limited to paclitaxel, docetaxel, and abraxane and so on. This term also include the derivatives and pharmaceutically acceptable salts thereof.

**[0040]** The amount of taxane compound is preferably administrated in routine manner and in routine dosage. For example, paclitaxel is usually administrated by 50-250 or 100-175 mg/m² via intravenous injection.

#### Plinabulin

[0041] In the present invention, another important active ingredient is plinabulin.

**[0042]** As used herein, terms "plinabulin", "NPI-2358" and "t-butyl dehydrophenylahistin" are exchangeable, and each means a synthetic, low molecular weight chemical entity, with chemical name of 2, 5-piperazinedione, 3-[[5-(1,1 dimethylethyl)-1H-imidazol-4-yl]methylene]-6-(phenylmethylene)-, (3Z,6Z)] as shown in Figure 1. In the present invention, the above terms also include the pharmaceutically acceptable salts thereof.

[0043] NPI-2358 is discovered as an anti-cancer agent. WO 2004/054498 has disclosed the structure, synthesis and use of NPI-2358.

**[0044]** Plinabulin inhibits the dimerization of tubulin monomers. Its mechanism is in its effect on tumor vascularture, and thus cutting the blood supply for tumor growth, classified as vascular disrupting agent (VDA).

**[0045]** In the present invention, the amount of plinabulin is administrated in routine manner and in routine dosage. Plinabulin is administrated by 20-30 mg/m<sup>2</sup> via intravenous injection

#### Combination, pharmaceutical composition and kit

[0046] Also described is a combination, comprising an active ingredient (a) of a taxane compound; and an active ingredient (b) of plinabulin.

**[0047]** Further, a pharmaceutical composition is described, comprising an active ingredient (a) of a taxane compound; an active ingredient (b) of plinabulin; and (c) a pharmaceutically acceptable carrier.

**[0048]** The dosage forms and preparation methods for the pharmaceutical composition described herein are not particularly limited, and the composition can be made into various dosage forms such as tablets, capsules, granules, sustained-release agents, injections, and the like by conventional processes in the art. A preferred dosage form is the oral dosage form.

[0049] Also described is a kit, comprising:

- 1. (i) a first container containing a first medicament which comprises a taxane compound as an active ingredient (a) and an optional pharmaceutically acceptable carrier; and
- 2. (ii) a second container containing a second medicament which comprises plinabulin as an active ingredient (b) and a optional pharmaceutically acceptable carrier;
- 3. (iii) an instruction recording an administration of the active ingredient (a) in combination with the active ingredient (b) for treating and/or preventing cancer.

[0050] The combination, formulations and kits described herein are useful for preventing and/or treating cancer.

**[0051]** The combination described herein may be administrated together or in sequence. Preferably, the active ingredient (a) of taxane is administrated first, followed by administration of the active ingredient (b) of plinabulin within 0.5-72hr, preferably within 0.5-24hrs, more preferably within 1-24hrs, thereby significantly improving the efficacy and the patient's compliance, and significantly reducing the side effect of taxane such as neutropenia.

[0052] Of course, the effective dosage of the active ingredients can be varied according to the administration mode and the severity of the disease to be treated.

# Treatment with combination of plinabulin and taxane

[0053] Also described herein is a method to use plinabulin and docetaxel to treat cancer subjects.

**[0054]** A method for treating and preventing cancer by using two active ingredients is described, which comprises administrating to a mammal subject (such as human) an effective amount of an active ingredient (a) of a taxane compound and an active ingredient (b) of plinabulin, or administrating a first medicament comprising the active ingredient (a) and a second medicament the active ingredient (b).

**[0055]** The two active ingredients or pharmaceutical compositions described herein may be administrated by conventional routes, including (but not limited to): intramuscular, intraperitoneal, intravenous, subcutaneous, intradermal, oral, or topical administration. Preferred routes of administration include oral administration.

**[0056]** The medicaments described herein can be solid compositions for convenient administration, especially the tablets and solid filled or liquid-filled capsules. Preferably, the medicament or pharmaceutical composition is formulated as liquid formulation, or a lyophilized formulation or other suitable injection form.

**[0057]** Furthermore, the two active ingredients or drugs described herein can be used in combination with other drugs for treating cancer (such as *cis*-platin, paclitaxel, anti-tumor antibodies).

**[0058]** In the examples, the inventor evaluated the antitumor activity of plinabulin in the MV522 lung cancer xenograft model. The *in vivo* efficacy of plinabulin was determined alone and in combination with docetaxel. Tumor growth inhibition (TGI) of plinabulin (3.75 and 7.5 mg/kg), docetaxel (15 and 25 mg/kg), and the combination was determined in the MV522 tumor model. Scheduling of the drug combinations was also conducted by administering one of the agents 2 hours after the first drug was given. Significant endpoints for this experiment included mean tumor growth inhibition (TGI) or regression, animal weight loss, potential toxicity, and tumor growth delay (TGD). The primary endpoint for the TGI study was the day that the mean NPI-2358 Vehicle tumor volume reached 1.2 cm<sup>3</sup>. The endpoint for the TGD study was when each tumor reached a volume of 1.5 cm<sup>3</sup>. In addition, an amendment was conducted to compare the efficacy of docetaxel and the combination of NPI-2358 + docetaxel in large tumors (1.5 cm<sup>3</sup>).

**[0059]** Animals were implanted with cancer cells harvested from tissue culture and allowed to establish tumors in nude mice. Treatment initiated when the average tumor volume of approximately 105 mm<sup>3</sup> was reached. No significant animal weight loss occurred in any of plinabulin single agent groups demonstrating that the drug was well tolerated at the given doses and schedule. As expected, significant animal weight loss was observed following treatment with docetaxel. **Unexpectedly, the addition of NPI-2358 to docetaxel in the** 

combination groups reduced the docetaxel associated weight loss (Table 2).

**[0060]** Treatment with plinabulin as a single agent induced a slight decrease in tumor volume compared to vehicle control, while docetaxel treatment resulted in a strong decrease in tumor burden. Importantly, the addition of plinabulin enhanced the antitumor activity of docetaxel in this model.

**[0061]** This effect was most prominent in the groups receiving docetaxel first followed by plinabulin, such as 2 hours later (Table 3). Furthermore, this drug combination was more effective than single agent docetaxel at reducing tumor burden in large tumors (1.5 cm<sup>3</sup>) (Table 4).

[0062] To further explore the plinabulin and docetaxel combination schedule of administration, the inventor investigated the *in vivo* efficacy of plinabulin administered 1, 4, and 24 hours after docetaxel treatment in MV522 human lung cancer xenograph model in mice. Experimental design was similar to the animal study discussed above. The additional of plinabulin to docetaxel in the combination groups reduced the docetaxel associated animal weight loss. The weight loss trend was similar among the different groups whether plinabulin was given 1, 4, or 24 hours following docetaxel treatment (Table 6). In addition, the addition of plinabulin enhanced the antitumor activity of docetaxel.

[0063] There was not a dramatic difference in tumor burden whether plinabulin was given 1, 4 or 24 hours following docetaxel treatment (Table 7). The 1 hour plinabulin after 15 mg/kg docetaxel groups produced slightly smaller tumors than the 4 and 24-hour groups (Table 7).

[0064] Following this initial study, animals with similar tumor burden were re-treated to evaluate the effects of the plinabulin and docetaxel combination on large tumors. The plinabulin and docetaxel combination produced a more pronounced decrease in tumor burden compared to docetaxel treatment alone. Unexpectedly, the inventor

discovered that plinabulin and docetaxel combination is effective against large established tumors (Table 8).

[0065] The investor applied the optimum plinabulin and docetaxel combination treatment schedule derived from MV522 animal studies in NSCLC patients. Patients received therapy on Day 1 and Day 8 in 3-week cycles. Therapy on Day 1 consisted of 75 mg/m² docetaxel administered via intravenous infusion (IV) over 1 hour, followed 2 hours later (from the time the docetaxel infusion begins) by 30 mg/m² plinabulin administered via intravenous infusion (IV) over 30 minutes. Therapy on Day 8 consisted of 20 mg/m² or 30 mg/m² plinabulin administered via intravenous infusion (IV) over 30 minutes.

**[0066]** The tolerability profile for plinabulin when added to docetaxel is better than docetaxel alone. Due to unbearable docetaxel side effects, the initial dose of 75 mg/m<sup>2</sup> docetaxel was decreased in 10% of patients (5 of 50) in the plinabulin 30 mg/m<sup>2</sup> plus docetaxel treatment arm whereas, the percent in the matched docetaxel alone arm was much higher at 18.2% of patients (10 of 55).

**[0067]** Of particular interest, patients in the plinabulin 30 mg/m<sup>2</sup> plus docetaxel arm had a statistically significantly lower rate of neutropenia (for all events and those events ≥ Grade 3) than patients in the matched docetaxel alone arm.

[0068] Neutropenia is the docetaxel side effect that is most severe. Neutropenia was seen in 36.4% of patients in this same docetaxel alone arm, consistent with historical data. In contrast, the incidence of neutropenia in the plinabulin 30 mg/m $^2$  plus docetaxel treated arm at 8% was significantly lower than the matched docetaxel alone arm (p<0.01).

**[0069]** Similar results to the above were observed in the plinabulin 20 mg/m<sup>2</sup> plus docetaxel study arm and its matched docetaxel treatment arm. Less G-CSF was used in the combination group compared with the docetaxel group.

[0070] The finding that plinabulin can decrease docetaxel's neutropenia rate is unexpected.

[0071] The inventor also performed extensive analyses and unexpectedly identified a subgroup which reacts to the plinabulin and docetaxel combination. Of the subgroup analyses, the one defined as "at least one tumor with a diameter size greater than 3 cm" was the subgroup in which plinabulin 30 mg/m² plus docetaxel (OS=11.5 M) had the largest significant OS benefit when compared with the matched docetaxel treated group (OS=7.8 M) (Fig. 6). The OS benefit in the combination group compared with the docetaxel group persists in other larger tumor groups (1 tumor > 5 cm, or >7 cm, Fig. 6). The inventor has discovered a uniquely defined large tumor group which benefit from the optimum plinabulin and docetaxel combination, which was never reported before in any literature. None of approved tumor vasculature targeting agent has been reported to show any favor in large tumor NSCLC populations.

**[0072]** In summary, the optimum combination scheme for plinabulin and docetaxel to achieve enhanced efficacy in uniquely defined large tumor patient population and dramatically decrease neutropenia rate of docetaxel in all patients has been described. Described is the use of Plinabulin in combination with other chemotherapy agents to treat large tumor in multiple cancer indications. Also described is the use of Plinabulin combined with other taxane compounds to decrease taxane compounds' debilitating neutropenia side effects.

[0073] The main advantages described include:

- 1. (a) It discloses the effect of taxane in combination with plinabulin in the prevention of cancer (such as lung cancer), and provides a method of treatment by using the optimum combination of taxane and plinabulin.
- 2. (b) Taxane and plinabulin combination are relatively safe.
- (c) The sequentially administration of taxane and plinabulin possesses synergistic inhibition of tumors, and has medically relevant statistical significance, thereby tolerance to taxane is increased and the side effects are reduced.

#### **EXAMPLES**

[0074] The present invention will be further illustrated below with reference to the specific examples. It should be understood that these examples are only to illustrate the invention but not to limit the scope of the invention. Invention, which is limited only by the scope of the appended claims. The experimental methods with no specific conditions described in the following examples are generally performed under the conventional conditions, or according to the manufacturer's instructions. Unless indicated otherwise, parts and percentage are calculated by weight.

#### Reference Example 1

**[0075]** *In vivo* evaluation of plinabulin (NPI-2358) as a single agent and in combination with docetaxel in the MV522 human non-small cell lung tumor xenograph model in athymic nu/nu mice

[0076] The objective of the example was to determine potential additive or synergistic effects of plinabulin in combination with docetaxel in the MV522 model by exploring the scheduling of the drug combination explored by administering one of the agents 2 hours after the first drug was given. Significant endpoints for this experiment included mean tumor growth inhibition (TGI) or regression, tumor growth delay (TGD) in large tumor, weight loss, and mortality.

#### **Experimental Design:**

#### Materials and Methods

**[0077] Model Information-** Female nude mice (Hsd: Athymic Nude-Foxn1<sup>nu</sup>) between 5 and 6 weeks of age weighing approximately 20 grams were obtained from Harlan, Inc. (Madison, WI). MV522 is a conventional human NSC metastatic lung tumor cell line (US 7,700,615 or 7,629,380). Animals were injected subcutaneously (SC) with approximately 1x10<sup>7</sup> MV522 cells harvested from tissue culture. When tumors grew to approximately 105 cubic millimeters (mm³) in size (3 days following implantation), animals were pair-matched by tumor size into treatment and control groups; each treatment group contained eight mice. Animals were ear-tagged and followed individually throughout the experiment.

[0078] Study Design and Dosing- Initial doses were administered on Day 1 following pair-matching. The experiment was carried out as a tumor growth inhibition (TGI) and tumor growth delay (TGD) study. Group 1 endpoint was when tumor volume reached 1.2 cm³ (TGI) and the treated groups endpoint was when each individual animal's tumor volume reached 1.5 cm³ (TGD). NPI-2358 in vehicle (8% Solutol® HS15, 12% PG, 80% D5W) was administered by intraperitoneal (IP) injection on a thrice weekly for two weeks - 1 day schedule (3xWKLYx2 - 1) or a twice weekly for three weeks schedule (2xWKLYx3) at the doses listed below (Table 1). To serve as negative controls, NPI-2358 vehicle and docetaxel vehicle were injected IP on a 2xWKLYx3 schedule and IV on a Q2Dx3 schedule, respectively. Docetaxel was administered by intravenous (IV) injection via tail vein at 15 mg/kg once every other day for three treatments (Q2Dx3) or at 25 mg/kg on a Q7Dx3 schedule(every 7 day for three treatment). Docetaxel and docetaxel vehicle were administered two hours prior to NPI-2358 vehicle and NPI-2358, respectively, in Groups 7-9 and 12-14. For Groups 10, 11, 15, and 16, NPI-2358 was given two hours prior to docetaxel.

Table 1: Study Design

<u>Group</u>	<u>Compound</u>	<u>Schedule</u>	Dose (mg/kg)	<u>Dosing Days</u>	<u>Route</u>
1	Untreated Control				
2	Docetaxel Vehicle	Q2Dx3		1,3,5	IV
3	NPI-2358 Vehicle	2xWKLYx3		1,4,8,11,15,18	ΙP
4	NPI-2358	3xWKLYx2-1	7.5	1,3,5,8,11	ΙP
5	NPI-2358	2xWKLYx3	3.75	1,4,8,11,15,18	ΙP
6	NPI-2358	2xWKLYx3	7.5	1,4,8,11,15,18	ΙP
7	Docetaxel NPI-2358 Vehicle	Q2Dx3 3xWKLYx2- 1	15	1,3,5 1,3,5,8,11	IV IP
8	Docetaxel NPI-2358	Q2Dx3 3xWKLYx2- 1	15 3.75	1,3,5 1,3,5,8,11	IV IP
9	Docetaxel NPI-2358	Q2Dx3 3xWKLYx2- 1	15 7.5	1,3,5 1,3,5,8,11	IV IP
10	NPI-2358 Docetaxel	3xWKLYx2-1 Q2Dx3	3.75 15	1,3,5,8,11 1,3,5	IP IV
11	NPI-2358 Docetaxel	3xWKLYx2-1 Q2Dx3	7.5 15	1,3,5,8,11 1,3,5	IP IV
12	Docetaxel NPI-2358 Vehicle	Q7Dx3 2xWKLYx3	25	1,8,15 1,4,8,11,15,18	IV IP
13	Docetaxel NPI-2358	Q7Dx3 2xWKLYx3	25 3.75	1,8,15 1,4,8,11,15,18	IV IP
14	Docetaxel NPI-2358	Q7Dx3 2xWKLYx3	25 7.5	1,8,15 1,4,8,11,15,18	IV IP
15	NPI-2358 Docetaxel	2xWKLYx3 Q7Dx3	3.75 25	1,4,8,11,15,18 1,8,15	IP IV
16	NPI-2358 Docetaxel	2xWKLYx3 Q7Dx3	7.5 25	1,4,8,11,15,18 1,8,15	IP IV

In combination groups 7-9 and 12-14, docetaxel was given 2 hours prior to NPI-2358 or NPI-2358 vehicle.

In combination groups 10, 11, 15, and 16, NPI-2358 was given 2 hours prior to docetaxel.

## **Data Collection and Statistical Analysis**

**[0079]** Animal Weights- Individual and group mean weights ±SD and percent weight change were recorded twice weekly until study completion beginning Day 1. Group weights on Day 42 and weight nadir values are reported.

[0080] Moribundity/Mortality- Animals were observed twice weekly for general moribundity and daily for mortality.

**[0081]** *Tumor Volume-* Individual and group mean tumor volumes ±SEM were recorded twice weekly (24 hours after dosing until study completion (mean control tumor volume=1.2 cm<sup>3</sup>, Day 25) beginning Day 1. Tumor measurements were converted to cubic millimeter tumor volume using the formula below:

Tumor Volume (mm $^3$ ) = Width  $^2$  (mm) x Length (mm) x 0.52

**[0082]** *Tumor Necrosis-* Degree of tumor necrosis was rated at each tumor measurement using the following arbitrary index:

N0	None	No Visible Necrosis
N1	Slight	Reddened or Inflamed; Intact Tumor
N2	Mild	<10% Tumor Necrosis
N3	Moderate	≥10 and <50% Tumor Necrosis
N4	Severe	>50% Tumor Necrosis

[0083] Notable differences in tumor necrosis between treated and control groups were reported.

**[0084]** *Tumor Growth Inhibition-* The TGI portion of the study was completed on Day 25 once the designated control group (Group 3, NPI-2358 vehicle) reached a mean tumor volume of approximately 1.2 cm<sup>3</sup> which is separate from the TGD study. Mice were weighed and caliper tumor measurements taken. Tumor growth inhibition (TGI) values were calculated for each group containing treated animals using the formula below:

1- Mean Final Tumor Volume (Treated) - Mean Initial Tumor Volume (Treated) Mean Final Tumor Volume (Control) - Mean Initial Tumor Volume (Control) X 100%

[0085] Animals experiencing complete tumor responses or animals experiencing technical or drug-related deaths were censored from final TGI calculations; however animals experiencing partial tumor responses were included in the final TGI calculations. The National Cancer Institute (NCI) criteria for compound activity is TGI>58%. TGI values for each treatment group are reported at study completion; these calculations are based on the final study day.

[0086] Tumor Growth Delay- This arm of the study was ended on Day 74 at the sponsor's request. At TGI study completion (Day 25), individual tumor volumes from control and treatment groups were reviewed and those greater than or equal to the designated TGD tumor volume endpoint (1.5 cm³) were removed from the study and each animal assigned a day of sacrifice value based on the day it reached the endpoint. Upon TGD study completion, a Median Day of Sacrifice (MDS) was calculated for the control (C) and each treatment (T) group and used to determine tumor growth delay (T-C) using the following equations:

MDS = Median 
$$[\Sigma \text{ ins}]$$
 T-C = MDS<sub>Treated</sub> - MDS<sub>Control</sub>

[0087] Where IDS (Individual Day of Sacrifice) is the day when each animal reached its tumor volume endpoint (1.5 cm<sup>3</sup>); only animals reaching this endpoint were included in TGD calculations. Gross or net log<sub>10</sub> cell kill for each treatment group was determined using the following equations:

Gross Log<sub>10</sub> Cell Kill = 
$$\frac{(T-C)}{(3.32) (Td)}$$

**[0088]** Where (T) = median day of death for the treated group, (C) = median day of death for the control group (NPI-2358 vehicle group), and (Td) is the tumor-volume doubling time estimated from a log-linear plot of log phase tumor growth (100-800 mm<sup>3</sup>) over time in the control group; (3.32) is the number of doublings required for a population to increase one  $\log_{10}$  unit. Weight and tumor data from individual animals experiencing technical or drug-related deaths was censored from final group calculations and statistical analyses. Long-term survivors (animals not reaching the tumor volume endpoint by the preset time period (LTS)) are not included in these calculations. For comparison of activity with standard agent(s), gross or net  $\log_{10}$  cell-kill values were converted to an arbitrary activity rating below:

Antitumor Activity	Gross Log <sub>10</sub> Cell-Kill
Highly Active +++++	=5.0
++++	4.0-4.9
+++	3.0-3.9
++	2.0-2.9
+	1.0-1.9
Inactive	<1.0

[0089] Net log<sub>10</sub> cell-kill is utilized for test agents administered for less than five days of total treatments while activity of agents dosed for five or more treatment days are calculated using gross log cell-kill; an active rating of (+++) to (++++) is needed to effect partial or complete responses.

**[0090]** Partial/Complete Tumor Response- Individual mice possessing tumors measuring less than on Day 1 were classified as having partial response (PR) and a percent tumor regression (%TR) value is determined using the formula below:

1- Final Tumor Volume (mm<sup>3</sup>)
Initial Tumor Volume (mm<sup>3</sup>) × 100%

**[0091]** If partial tumor responses are reported in multiple animals within one group, a mean PR value was determined. Individual mice lacking palpable tumors were classified as undergoing complete response (CR). The number of partial and complete responses and percent tumor regression is reported for each treatment group at study completion; these calculations are based on the final study day.

**[0092]** *TGI Statistics*- Statistical analyses were carried out between treated and control groups comparing final tumor volume. For two or more treatment groups, a two-tailed One-Way Analysis of Variance (ANOVA) followed by the Dunnett's multiple comparisons test was employed. An unpaired, two-tailed student t-test was used to compare one treated group to control. Weight and tumor data from individual animals experiencing technical or drug-related deaths were censored from analysis. However, weight and tumor data from animals reporting partial or complete responses were included in these calculations.

**[0093]** *TGD Statistics*- A Log-rank test was used to determine statistically significant differences in overall survival experience between each treated group compared to control and as utilized is equivalent to the Mantel-Haenszel test. If a drug is evaluated at multiple concentrations on the same route and schedule, a Log-rank test from trend is also performed. Weight and tumor data from individual animals experiencing technical or drug-related deaths was censored from analysis. However, weight and tumor data from animals reporting partial or complete regressions or long term survivors were included in these calculations. All analyses were performed using GraphPad Prism® software (version 5.0).

# Results

**[0094]** No significant animal weight loss occurred in any of plinabulin single agent groups demonstrating that the drug was well tolerated at the given doses and schedule. As expected, significant animal weight loss was observed following treatment with docetaxel.

[0095] Unexpectedly, the addition of NPI-2358 to docetaxel in the combination groups reduced the docetaxel associated weight loss (Table 2).

Table 2: Animal Weight and Drug Toxicity Results: Control, Single Agent, and Combination Groups

	GROUP	DOSE	ROUTE/SCHEDULE	FINAL WE	<b>FINAL WEIGHT DATA (DAY 25)</b>		TA (DAY 25)	WEIGHT N	<u>IADIR</u>
				MEAN (G)	±	SD	%CHANGE	%CHANG	DAY
1	Untreated Control			24	±	2	+8	0	1
2	Docetaxel Vehicle		IV/Q2Dx3	22	±	2	+6	-6	18
3	NPI-2358 Vehicle		IP/2xWKLYx3*	23	±	2	+14	0	1
4	NPI-2358	7.5	IP/ 3xWKLYx2-1§	24	±	2	+11	-1	8
5	NPI-2358	3.75	IP/2xWKLYx3*	23	±	1	+15	0	1

	GROUP	DOSE	ROUTE/SCHEDULE	FINAL WE	IGH	T DA	TA (DAY 25)	WEIGHT N	<u>ADIR</u>
ammin				MEAN (G)	±	SD	%CHANGE	%CHANG	DAY
6	NPI-2358	7.5	IP/2xWKLYx3*	25	±	2	+14	0	1
	Docetaxel	15	IV/ Q2Dx3						
7	NPI-2358 Vehicle		IP/ 3xWKLYx2-1§	24	±	2	+17	-23	11
8	Docetaxel	15	IV/ Q2Dx3	23	±	2	+19	-22	11
0	NPI-2358	3.75	IP/ 3xWKLYx2-1	23	I		שוד	-22	11
9	Docetaxel	15	IV/ Q2Dx3	24	±	2	+15	-15	11
9	NPI-2358	7.5	IP/ 3xWKLYx2-1§	2 <del>4</del>	T	_	713	-13	11
10	NPI-2358	3.75	IP/ 3xWKLYx2-1§	23	±	2	+17	-17	11
10	Docetaxel	15	IV/ Q2Dx3	23	T		717	-17	11
11	NPI-2358	7.5	IP/ 3xWKLYx2-1§	24	±	1	+20	-15	11
	Docetaxel	15	IV/ Q2Dx3	24	1	'	120	-13	11
	Docetaxel	25	IV/ Q7Dx3						
12	NPI-2358 Vehicle		IP/2xWKLYx3*	18	±	2	-10	-23	15
13	Docetaxel	25	IV/ Q7Dx3	20	±	3	-3	-11	15
13	NPI-2358	3.75	IP/2xWKLYx3*	20	_	J	-3	-11	13
14	Docetaxel	25	IV/ Q7Dx3	22	±	3	+3	-16	15
14	NPI-2358	7.5	IP/2xWKLYx3*	22	I	3	73	-10	13
15	NPI-2358	3.75	IP/2xWKLYx3*	20	±	3	-4	1.4	15
10	Docetaxel	25	IV/ Q7Dx3	<b>Z</b> U	Τ	3	-4	-14	IJ
16	NPI-2358	7.5	IP/2xWKLYx3*	20	±	2	-3	-19	18
10	Docetaxel	25	IV/ Q7Dx3	20	I		-3	-10	10

N=8/GRP ON DAY 1

\*Days of Injection: 1, 4, 8, 11, 15, 18 § Days of Injection: 1, 3, 5, 8, 11

In combination groups 7-9 and 12-14, docetaxel was given 2 hours prior to NPI-2358 or vehicle.

In combination groups 10, 11, 15, and 16, NPI-2358 was given 2 hours prior to docetaxel.

[0096] Treatment with plinabulin as a single agent induced a slight decrease in tumor volume compared to Vehicle control, while docetaxel treatment resulted in a strong decrease in tumor burden. Importantly, the addition of NPI-2358 enhanced the antitumor activity of docetaxel in this model.

**[0097]** This effect was most prominent in the groups receiving docetaxel first followed by NPI-2358 2 hours later (Table 3). Furthermore, this drug combination was more effective than single agent docetaxel at reducing tumor burden in large tumors (1.5 cm<sup>3</sup>) (Table 4).

Table 3: Tumor Volume and TGI Results: Control, Single Agent, and Combination Groups

			 				<del>.</del>		 
				FINAL TUM		<u>DLUME</u>			,
-				(DAY 25)					
accentan			ROUTE/SCHEDULE		±			#PR/CR	***************************************
7				(MM°)					3
***************************************	1	Untreated Control	 	963	±	114			 mmmmm
		Docetaxel							9

	······			FINAL TUM	OR V	<u>DLUME</u>			
				(DA	Y 25)	***************************************			***************************************
ammanna	GROUP	DOSE	ROUTE/SCHEDULE	MEAN (MM <sup>3</sup> )	±	SEM	%TGI	#PR/CR	%TR
2	Vehicle		IV/ Q2Dx3	1411	±	75		0/0	
3	NPI-2358 Vehicle		IP/2xWKLYx3*	1371	±	88		0/0	
4	NPI-2358	7.5	IP/ 3xWKLYx2-1§	1222	±	114	12	0/0	
5	NPI-2358	3.75	IP/2xWKLYx3*	1347	±	82	2	0/0	
6	NPI-2358	7.5	IP/2xWKLYx3*	1215	±	110	12	0/0	
7	Docetaxel	15	IV/ Q2Dx3	259	±	45	88	0/0	
	NPI-2358 Vehicle		IP/3xWKLYx2-1 <sup>§</sup>						
8	Docetaxel	15	IV/ Q2Dx3	93	±	33	99†	4/2	42
	NPI-2358	3.75	IP/ 3xWKLYx2-1						
9	Docetaxel	15	IV/ Q2Dx3	160	±	25	96 <sup>†</sup>	2/0	59
	NPI-2358	7.5	IP/3xWKLYx2-1 <sup>§</sup>			***************************************			*************
10	NPI-2358	3.75	IP/ 3xWKLYx2-1 <sup>§</sup>	260	±	73	88 <sup>†</sup>	2/0	57
	Docetaxel	15	IV/ Q2Dx3			***************************************			***************************************
11	NPI-2358	7.5	IP/3xWKLYx2-1 <sup>§</sup>	233	±	30	90	0/0	
	Docetaxel	15	IV/ Q2Dx3			***************************************			**********
12	Docetaxel	25	IV/ Q7Dx3	43	±	4	107 <sup>†</sup>	8/0	58
	NPI-2358 Vehicle		IP/2xWKLYx3*						
13	Docetaxel	25	IV/ Q7Dx3	36	±	6	107 <sup>†</sup>	7/1	58
	NPI-2358	3.75	IP/2xWKLYx3*						
14	Docetaxel	25	IV/ Q7Dx3	27	±	5	109 <sup>†</sup>	6/1	70
	NPI-2358	7.5	IP/2xWKLYx3*						
15	NPI-2358	3.75	IP/2xWKLYx3*	48		12	104 <sup>†</sup>	6/0	67
	Docetaxel	25	IV/ Q7Dx3						
16	NPI-2358	7.5	IP/2xWKLYx3*	66		7	103 <sup>†</sup>	6/0	46
	Docetaxel	25	IV/ Q7Dx3	***************************************					

# N=8/GRP ON DAY 1

\*Days of Injection: 1, 4, 8, 11, 15, 18 § Days of Injection: 1, 3, 5, 8, 11 In combination Groups 7-9 and 12-14, docetaxel was given 2 hours prior to NPI-2358 or NPI-2358 vehicle.

In combination Groups 10, 11, 15, and 16, NPI-2358 was given 2 hours prior to docetaxel.

† Value includes tumors with PR.

Table 4: MDS and TGD Results

				MDS			3	ACTIVITY	₹ :		
mmmmm	5	 ROUTE/SCHEDULE	DAY	±	SD	T- C	CELL KILL	RATING	#LTS	#PR/C	%TR
1	Untreated Control	 	34	±	14				1	0/0	

					MDS			LOG <sub>10</sub>	ACTIVITY			
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	GROUP	DOSE	ROUTE/SCHEDULE	DAY	±	SD	T- C	CELL KILL	RATING	#LTS	#PR/C	%TR
2	Docetaxel Vehicle		IV/ Q2Dx3	27	±	3	 -			0	0/0	
3	NPI-2358 Vehicle		IP/2xWKLYx3*	29	±	4	 -			0	0/0	
4	NPI-2358	7.5	IP/ 3xWKLYx2-1 <sup>§</sup>	31	±	5	2	0.0	-	0	0/0	
5	NPI-2358	3.75	IP/2xWKLYx3*	29	±	2	0	0.0	-	0	0/0	
6	NPI-2358	7.5	IP/2xWKLYx3*	29	±	12	0	0.0	-	0	0/0	
7	Docetaxel NPI-2358 Vehicle	15	IV/Q2Dx3 IP/ 3xWKLYx2-1 <sup>§</sup>	45	±	9	16	0.4	-	0	0/0	
8	Docetaxel NPI-2358	15 3.75	IV/Q2Dx3 IP/ 3xWKLYx2-1	53	±	6	24	0.6	-	0	0/0	
9	Docetaxel NPI-2358	15 7.5	IV/Q2Dx3 IP/ 3xWKLYx2-1 <sup>§</sup>	52	±	10	23	0.6	-	0	0/0	
10	NPI-2358 Docetaxel	3.75 15	IP/ 3xWKLYx2-1 <sup>§</sup> IV/Q2Dx3	53	±	7	24	0.6	-	1	1/0	83
11	NPI-2358 Docetaxel	7.5 15	IP/ 3xWKLYx2-1 <sup>§</sup> IV/Q2Dx3	46	±	2	17	0.4	-	0	0/0	
12	Docetaxel NPI-2358 Vehicle	25	IV/ Q7Dx3 IP/2xWKLYx3*	68	±	6	39	1.0	+	2	0/0	
13	Docetaxel NPI-2358	25 3.75	IV/ Q7Dx3 IP/2xWKLYx3*	67	±	6	38	1.0	+	3	1/0	42
14	Docetaxel NPI-2358	25 7.5	IV/ Q7Dx3 IP/2xWKLYx3*	69	±	8	40	1.1	+	1	0/0	
15	NPI-2358 Docetaxel	3.75 25	IP/2xWKLYx3* IV/ Q7Dx3	67	±	3	38	1.0	+	0	0/0	
16	NPI-2358 Docetaxel	7.5 25	IP/2xWKLYx3* IV/ Q7Dx3	61	±	5	32	0.8	+	2	1/0	95

## N=8/GRP ON DAY 1

\*Days of Injection: 1, 4, 8, 11, 15, 18 § Days of Injection: 1, 3, 5, 8, 11

In combination Groups 7-9 and 12-14, docetaxel was given 2 hours prior to NPI-2358 or NPI-2358 vehicle.

In combination Groups 10, 11, 15, and 16, NPI-2358 was given 2 hours prior to docetaxel.

\*Days of Injection: 1, 4, 8, 11, 15, 18

#### Reference Example 2

[0098] In vivo evaluation of plinabulin in combination with docetaxel in the MV522 human non-small cell lung tumor xenograph model in athymic nu/nu mice: schedule of drug administration

[0099] The objective of the example was to evaluate the potential additive or synergistic effects of plinabulin (NPI-2358) in combination with docetaxel by administering NPI-2358 at various time points (1h, 4h, or 24h) after docetaxel in the MV522 model. Significant endpoints for this experiment included mean tumor growth inhibition (TGI) or regression, tumor growth delay (TGD) in large tumor, weight loss, and mortality.

#### **Experimental Design:**

#### Materials and Methods

**[0100]** *Model Information*- Female nude mice (Hsd: Athymic Nude-*Foxn1*<sup>nu</sup>) between 5 and 6 weeks of age weighing approximately 20 grams were obtained from Harlan, Inc. (Madison, WI). Animals were injected subcutaneously (SC) with approximately 1x10<sup>7</sup> MV522 cells harvested from tissue culture (1:1 matrigel:media). When tumors grew to approximately 100 cubic millimeters (mm³) in size, animals were pair-matched by tumor size into treatment and control groups. Animals were ear-tagged and followed individually throughout the experiment.

**[0101]** *Original Study Design and Dosing-* Initial doses were administered on Day 1 following pair-matching. The experiment was carried out as both a tumor growth inhibition (TGI) and tumor growth delay (TGD) study. For TGI, the endpoint was when the average tumor volume of Group 1 reached 1.2 cm<sup>3</sup>. For TGD, the endpoint was reached when each individual animal's tumor volume reached 1.0 cm<sup>3</sup>. A 10 mg/ml stock vial of docetaxel was diluted in 0.9% saline on each day of dosing and administered intravenously (IV). The schedule and doses for each agent are listed in Table 5. Docetaxel vehicle (0.9% saline; IV; Days 1, 3, 5) plus NPI-2358 vehicle (8% Solutol® HS15, 12% PG, 80% D5W; IP; Days 1, 3, 5, 8, 11) was administered to serve as negative control.

Table 5: Study Design

	#Animals	Compound	Hours after Docetaxel	Dose (mg/kg)	Route	Dosing Days
1	8	Docetaxel Vehicle	0		IV	1, 3, 5
		NPI-2358 Vehicle	1		ΙP	1, 3, 5, 8, 11
2	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358 Vehicle	1		ΙP	1, 3, 5, 8, 11
3	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358 Vehicle	4		ΙP	1, 3, 5, 8, 11
4	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358 Vehicle	24		ΙP	1, 3, 5, 8, 11
5	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	1	3.75	ΙP	1, 3, 5, 8, 11
6	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	4	3.75	ΙP	1, 3, 5, 8, 11
7	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	24	3.75	ΙP	1, 3, 5, 8, 11
8	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	1	7.5	IP	1, 3, 5, 8, 11
9	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	4	7.5	ΙP	1, 3, 5, 8, 11
10	8	Docetaxel	0	15	IV	1, 3, 5
		NPI-2358	24	7.5	ΙΡ	1, 3, 5, 8, 11
11	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358 Vehicle	1		IΡ	1, 4, 8, 11, 15, 18
12	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358 Vehicle	4		ΙP	1, 4, 8, 11, 15,

	#Animals	Compound	Hours after Docetaxel	Dose (mg/kg)	Route	Dosing Days
						18
13	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358 Vehicle	24		IΡ	1, 4, 8, 11, 15, 18
14	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	1	3.75	IΡ	1, 4, 8, 11, 15, 18
15	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	4	3.75	IΡ	1, 4, 8, 11, 15, 18
16	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	24	3.75	IΡ	1, 4, 8, 11, 15, 18
17	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	1	7.5	IΡ	1, 4, 8, 11, 15, 18
18	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	4	7.5	IΡ	1, 4, 8, 11, 15, 18
19	8	Docetaxel	0	25	IV	1, 8, 15
		NPI-2358	24	7.5	ΙP	1, 4, 8, 11, 15, 18

# **Data Collection and Statistical Analysis**

**[0102]** The study in this example was completed on Day 24 once the designated control group reached a mean tumor volume of approximately 1.2 cm<sup>3</sup>.

**[0103]** Animal Weights- Individual and group mean weights ±SD and percent weight change were recorded twice weekly until study completion beginning Day 1. Group weights through Day 24 or 53 and weight nadir values are reported.

[0104] Moribundity/Mortality-Same as in Example 1.

[0105] Tumor Volume- Same as in Example 1.

[0106] Tumor Necrosis- Same as in Example 1.

[0107] Partial/complete Tumor Response -Same as in Example 1.

[0108] Tumor Growth Inhibition- Same as in Example 1.

[0109] TGI Statistics - Same as in Example 1.

**[0110]** *Tumor Growth Delay-* Same as in Example 1. This arm of the study was ended on Day 80 or 53 at the sponsor's request. Long-term survivors (animals not reaching the tumor volume endpoint by the preset time period (LTS)) were assigned an IDS value of the last study day (Day 80) and included in these calculations.

**[0111]** For comparison of activity with standard agent(s), gross or net log<sub>10</sub> cell-kill values were converted to an arbitrary activity rating below:

Activity Rating	Treatment = < 5 Days (Net Log <sub>10</sub> Cell-Kill)	Treatment = 5-20 Days (Gross Log <sub>10</sub> Cell-Kill)	Treatment >= 20 Days (Gross Log <sub>10</sub> Cell-Kill)
Highly Active			
++++	>2.6	>2.8	>3.4
+++	1.6-2.6	2.0-2.8	2.5-3.4
++	0.9-1.5	1.3 - 1.9	1.7-2.4
+	0.5-0.8	0.7 - 1.2	1.0-1.6
Inactive			
-	<0.5	<0.7	<1.0

**[0112]** Net log<sub>10</sub> cell-kill is utilized for test agents administered for less than five days of total treatments while activity of agents dosed for five or more treatment days are calculated using gross log cell-kill; an active rating of (+++) to (++++) is needed to effect partial or complete responses.

[0113] TGD Statistics- Same as in Example 1.

#### Results

[0114] To further explore the plinabulin and docetaxel combination schedule of administration, the inventor investigated the *in vivo* efficacy of plinabulin administered 1, 4, and 24 hours after docetaxel treatment in MV522 human lung cancer xenograph model in mice. Experimental design was similar to the animal study discussed above. The additional of plinabulin to docetaxel in the combination groups reduced the docetaxel associated animal weight loss. The weight loss trend was similar among the different groups whether plinabulin was given 1, 4, or 24 hours following docetaxel treatment (Table 6).

Table 6: Animal Weight and Drug Toxicity Results - All Groups

3										
	Group	Hours after	Dose	Final Weight - Day	24 AVG+/-SD (g)	NADII	R			
	0.035	Docetaxel	(mg/kg)	%Change		%change	Day			
1	Docetaxel Vehicle	0		22 +/-1	+12	0	1			
	NPI-2358 Vehicle	1								
2	Docetaxel	0	15	22 +/-1	+13	-14	10			
	NPI-2358 Vehicle	1								
3	Docetaxel	0	15	23 +/- 2	+12	-10	10			
	NPI-2358 Vehicle	4			-					
4	Docetaxel	0	15	23 +/- 2	+17	-10	10			
	NPI-2358 Vehicle	24			***************************************					
5	Docetaxel	0	15	22 +/-1	+18	-12	10			
	NPI-2358	1	3.75							
6	Docetaxel	0	15	23 +/- 2	+14	-9	10			
	NPI-2358	4	3.75							
7	Docetaxel	0	15	23 +/- 2	+15	-12	10			
4	*********	************	******	-2	,	1	3			

	Group	Hours after	Dose	Final Weight - Day	24 AVG+/-SD (g)	NADIR		
	Group	Docetaxel	(mg/kg)	%Change		%change	Day	
	NPI-2358	24	3.75					
8	Docetaxel	0	15	22 +/- 1	+15	-10	10	
	NPI-2358	1	7.5					
9	Docetaxel	0	15	23 +/-1	+19	-13	10	
	NPI-2358	4	7.5					
10	Docetaxel	0	15	22 +/- 1	+17	-11	10	
	NPI-2358	24	7.5					
11	Docetaxel	0	25	18 +/- 2	-10	-13	17	
	NPI-2358 Vehicle	1						
12	Docetaxel	0	25	17 +/- 2	-11	-14	10	
	NPI-2358 Vehicle	4						
13	Docetaxel	0	25	17 +/- 3	-12	-20	17	
	NPI-2358 Vehicle	24						
14	Docetaxel	0	25	19 +/- 2	-4	-9	21	
	NPI-2358	1	3.75					
15	Docetaxel	0	25	19 +/- 1	-4	-8	21	
	NPI-2358	4	3.75					
16	Docetaxel	0	25	18 +/- 2	-5	-9	21	
	NPI-2358	24	3.75					
17	Docetaxel	0	25	20 +/- 2	-3	-7	21	
	NPI-2358	1	7.5					
18	Docetaxel	0	25	20+/-1	+1	-3	21	
	NPI-2358	4	7.5					
19	Docetaxel	0	25	20 +/- 2	-2	-8	21	
	NPI-2358	24	7.5					

[0115] In addition, the addition of plinabulin appeared to enhance the antitumor activity of docetaxel. There was not a dramatic difference in tumor burden whether plinabulin was given 1, 4 or 24 hours following docetaxel treatment (Table 7). The 1 hour plinabulin after 15 mg/kg docetaxel groups produced slightly smaller tumors than the 4 and 24-hour groups (Table 7).

Table 7: Tumor Volume and TGI Results - All Groups

	Group	Hours after Docetaxel	Dose (mg/kg)	Final Tumor Volume (Day 24) Mean +/- SEM	%TGI	%TR (#)
1	Docetaxel Vehicle	0		1185 +/- 65	-	-
anaman	NPI-2358 Vehicle	1				
2	Docetaxel	0	15	337 +/- 29	79	-
	NPI-2358 Vehicle	1				
3	Docetaxel	0	15	313 +/- 23	81	-
	NPI-2358	4				

	Group	Hours after Docetaxel	Dose (mg/kg)	Final Tumor Volume (Day 24) Mean +/- SEM	%TGI	%TR (#)
*******	Vehicle					
4	Docetaxel	0	15	325 +/- 41	80	-
	NPI-2358 Vehicle	24				
5	Docetaxel	0	15	77 +/- 13	103	51(5)
	NPI-2358	1	3.75			
6	Docetaxel	0	15	131 +/- 35	98	67(3)
	NPI-2358	4	3.75			
7	Docetaxel	0	15	153 +/- 21	96	19(1)
	NPI-2358	24	3.75			
8	Docetaxel	0	15	92 +/- 16	101	31(3)
	NPI-2358	1	7.5	•••		
9	Docetaxel	0	15	139 +/- 29	97	45(3)
	NPI-2358	4	7.5	aa-3		
10	Docetaxel	0	15	136 +/- 18	97	39(2)
	NPI-2358	24	7.5			
11	Docetaxel	0	25	69 +/- 10	104	42(7)
	NPI-2358 Vehicle	1				
12	Docetaxel	0	25	65 +/- 9	104	45(7)
	NPI-2358 Vehicle	4				
13	Docetaxel	0	25	72 +/- 14	103	55(7)
	NPI-2358 Vehicle	24				
14	Docetaxel	0	25	37 +/- 6	106	64(8)
	NPI-2358	1	3.75	3.3.X		
15	Docetaxel	0	25	41 +/- 7	106	61(8)
	NPI-2358	4	3.75			
16	Docetaxel	0	25	33 +/- 6	107	68(8)
	NPI-2358	24	3.75	•••		
17	Docetaxel	0	25	61 +/- 10	104	49(7)
	NPI-2358	1	7.5	•••		
18	Docetaxel	0	25	44 +/- 5	106	56(8)
	NPI-2358	4	7.5			
19	Docetaxel	0	25	36 +/- 6	106	61(7)/CR(1)
	NPI-2358	24	7.5	aac		***************************************

[0116] Following this initial study, animals with similar tumor burden were re-treated to evaluate the effects of the plinabulin and docetaxel combination on large tumors. The plinabulin and docetaxel combination produced a more pronounced decrease in tumor burden compared to docetaxel treatment alone. Unexpectedly, the inventor discovered that plinabulin and docetaxel combination is effective against large established tumors (Table 8).

Table 8: MDS and TGD Results - All Groups

	Group	Hours after Docetaxel	Dose (mg/kg)	MDS Day +/- SD	T-C	LOG <sub>10</sub> Cell Kill	Activity Rating
1	Docetaxel Vehicle	0		24 +/- 3			
	NPI-2358 Vehicle	1					
2	Docetaxel	0	15	38 +/- 3	14	0.6	
	NPI-2358 Vehicle	1					
3	Docetaxel	0	15	42 +/- 6	18	0.7	+
	NPI-2358 Vehicle	4					
4	Docetaxel	0	15	40 +/- 7	16	0.6	-
	NPI-2358 Vehicle	24					
5	Docetaxel	0	15	52 +/- 4	28	1.1	+
	NPI-2358	1	3.75				
6	Docetaxel	0	15	42 +/- 6	18	0.7	+
	NPI-2358	4	3.75				
7	Docetaxel	0	15	45 +/- 4	21	0.9	+
	NPI-2358	24	3.75				
8	Docetaxel	0	15	52 +/- 11	28	1.1	+
	NPI-2358	1	7.5				
9	Docetaxel	0	15	49 +/- 7	25	1.0	+
	NPI-2358	4	7.5				
10	Docetaxel	0	15	47 +/- 6	23	0.9	+
	NPI-2358	24	7.5				
11	Docetaxel	0	25	63 +/- 10	39	1.6	++
	NPI-2358 Vehicle	1					
12	Docetaxel	0	25	56 +/- 4	32	1.3	++
	NPI-2358 Vehicle	4	<b>\}</b>				
13	Docetaxel NPI-2358 Vehicle	0	25	59 +/- 10	35	1.4	++
		24		7			
14	Docetaxel	0	25	66 +/- 9	42	1.7	++
	NPI-2358	1	3.75				
15	Docetaxel	0	25	59 +/- 6	35	1.4	++
	NPI-2358	4	3.75				
16	Docetaxel	0	25	59 +/- 12	35	1.4	++
	NPI-2358	24	3.75				
17	Docetaxel	0	25	59 +/- 9	35	1.4	++
	NPI-2358	1	7.5				
18	Docetaxel	0	25	61 +/- 9	37	1.5	++
	NPI-2358	4	7.5		9		
19	Docetaxel	24	25	63 +/- 7	39	1.6	++
	NPI-2358	24	7.5	225			-

[0117] Evaluation of the optimum plinabulin and docetaxel combination in treating advanced non-small cell lung cancer patients

#### **OBJECTIVES**

#### **Primary**

[0118] To compare the overall survival of patients with NSCLC treated with docetaxel to patients treated with docetaxel + plinabulin.

#### **Secondary**

#### [0119]

- 1. 1. To compare the response rate, duration of response, 6-month survival, and progression free survival in patients with NSCLC treated with docetaxel to patients with docetaxel + plinabulin;
- 2. 2. To compare the safety and adverse events profile of docetaxel to docetaxel + plinabulin.

#### STUDY DESIGN

**[0120]** This was an open-label study in patients with advanced NSCLC that had progressed after treatment with at least 1 chemotherapy regimen. Patients were randomized to receive either docetaxel plus plinabulin (DN) or docetaxel alone (75 mg/m²) (D). 2 dosing cohorts were investigated:

- 1. 1) 30 mg/m<sup>2</sup> dosing cohort: Approximately 110 patients were to be randomized (1:1) to receive either docetaxel plus plinabulin at 30 mg/m<sup>2</sup> (DN 30 mg/m<sup>2</sup> arm) or docetaxel alone (D arm);
- 2. 2) 20 mg/m<sup>2</sup> dosing cohort: Approximately 57 patients were to be randomized (2:1) to receive either docetaxel plus plinabulin at 20 mg/m<sup>2</sup> (DN 20 mg/m<sup>2</sup> arm) or docetaxel (D arm) alone.

#### Dosing Regimen

[0121] Patients received therapy on Day 1 and Day 8 in 3-week cycles.

**[0122]** Therapy on Day 1 consisted of 75 mg/m² docetaxel administered via intravenous infusion (IV) over 1 hour, followed 2 hours later (from the time the docetaxel infusion began) by placebo (Arm D) or 30 mg/m² or 20 mg/m² plinabulin (Arm DN) DN administered via intravenous infusion (IV) over 30 minutes. Oral dexamethasone (16 mg) was given the day prior to, the day of and the day following docetaxel infusion (Day 1). Therapy on Day 8 consisted of placebo (Arm D) or 30 mg/m² or 20 mg/m² plinabulin (Arm DN) administered via intravenous infusion (IV) over 30 minutes.

[0123] In patients experiencing drug related Grade > 2 treatment emergent adverse events (except alopecia, anorexia, and fatigue) according to the CTCAE (v3.0) treatment may be delayed until the adverse event has recovered to < Grade 1. Safety laboratory tests must meet the following criteria prior to treatment with docetaxel at

the beginning of each subsequent cycle: AST  $\leq$ 2.5 x ULN, ALT  $\leq$ 2.5 x ULN ( $\leq$ 1.5 x ULN if alkaline phosphatase is >=2.5 x ULN); bilirubin < =ULN; hemoglobin >= 9 g/dL, absolute neutrophil count >= 1.5 x 10<sup>9</sup> /L and platelets >=100 x 10<sup>9</sup> /L. Dose reductions may be implemented for patients who experience recurrent or specific severe toxicities.

**[0124] TARGET POPULATION** Patients with stage IIIb/IV non-small cell lung cancer that has progressed after treatment with at least one chemotherapy regimen.

#### **INCLUSION CRITERIA**

#### [0125]

- 1. 1. Male and females >=18 years of age.
- 2. 2. ECOG performance status ≤ 1.
- 3. 3. Pathologically or histologically confirmed advanced non-small cell lung cancer (unresectable Stage IIIb or IV) that has progressed after treatment with at least one chemotherapy regimen. Measurable disease is not required for enrollment into this trial.
- 4. 4. All Adverse Events of any prior chemotherapy, surgery, or radiotherapy, must have resolved to CTCAE (v. 3.0) Grade ≤ 2.
- 5. 5. The following laboratory results, within 14 days:
  - Hemoglobin > 9 g/dL
  - Absolute neutrophil count >= 1.5 x 10<sup>9</sup>/L
  - Platelet count >= 100 x 10<sup>9</sup> /L
  - Serum bilirubin ≤ ULN
  - ∘ AST and ALT ≤ 2.5 x ULN (≤ 1.5 x ULN if alkaline phosphatase is >=2.5 x ULN).
- 6. 6. Signed informed consent.

## **EXCLUSION CRITERIA**

#### [0126]

- 1. 1. Administration of certain chemotherapy, biological, immunotherapy, radiation therapy or investigational agent (therapeutic or diagnostic) within 21 days prior to receipt of study medication. Major surgery, other than diagnostic surgery, within 6 weeks before first study drug administration.
- 2. 2. Significant cardiac history:
  - History of myocardial infarction or ischemic heart disease;
  - History of clinically significant arrhythmias; uncontrolled arrhythmia or a requirement for anti-arrhythmics;
  - History of congenital QT prolongation;
  - Left bundle branch block;
  - ECG findings consistent with ischemic heart disease;
  - New York Heart Association Class III or IV cardiac disease
  - Uncontrolled hypertension: blood pressure consistently greater than 150 mm Hg systolic and 100 mm Hg diastolic in spite of antihypertensive medication.
- 3. 3. Prior treatment with tumor vascular disrupting agents.
- 4. 4. Prior seizure disorder.
- 5. 5. Brain metastases. Patients who demonstrate signs or symptoms of brain metastases should be imaged with CT or MRI. Patients who have brain metastases that have been previously treated and reimaged after treatment and whose lesions are stable without interim development of new lesions may be enrolled.
- 6. 6. History of significant gastrointestinal disease such as ileus, bowel obstruction, hemorrhagic diarrhea, inflammatory bowel disease, active uncontrolled peptic ulcer disease. (Concomitant therapy with ranitidine or its equivalent and/or omeprazole or its equivalent is acceptable).

- 7. 7. History of peri-operative pelvic radiation therapy, whole abdomen radiation therapy, or >= Grade 2 residual gastrointestinal symptoms from radiation therapy.
- 8. 8. Active uncontrolled bacterial, viral, or fungal infection, requiring systemic therapy.
- 9. 9. Known infection with human immunodeficiency virus (HIV), or active hepatitis A, B, or C.
- 10. 10. Patients with a prior hypersensitivity reaction to any product containing polysorbate 80, taxanes, Solutol and/or propylene glycol.
- 11. 11. Pregnant or breast-feeding women. Female patients must be postmenopausal, surgically sterile or they must agree to use acceptable methods of birth control (i.e., a hormonal contraceptive with barrier method, intra-uterine device, diaphragm with spermicidal or condom with spermicide, abstinence) for the duration of the study and for one month following study completion. Female patients with childbearing potential must have a negative serum pregnancy test within 10 days before the first study drug administration. Male patients must be surgically sterile or agree to use an acceptable method of contraception.
- 12. 12. Concurrent, active second malignancy for which the patient is receiving therapy, excluding basal cell carcinoma of the skin or carcinoma *in situ* of the cervix.
- 13. 13. Any medical conditions that, in the Investigator's opinion, would impose excessive risk to the patient. Examples of such conditions include infection requiring parenteral anti-infective treatment, hydronephrosis, liver failure, any altered mental status or any psychiatric condition that would interfere with the understanding of the informed consent.
- 14. 14. Unwilling or unable to comply with procedures required in this protocol.

#### LENGTH OF STUDY

**[0127]** Stable and responding patients were treated in this investigation as long as he/she has evidence of clinical benefit (stable disease or a response) in the absence of unacceptable adverse events. Study closure was planned 12 months after the last patient is accrued.

#### INVESTIGATIONAL PRODUCTS/ DOSE/ROUTE/REGIMEN

#### Plinabulin or Placebo:

**[0128]** The initial dose of plinabulin was 30 mg/m² or 20 mg/m². Dose adjustments depended on observed adverse events. Volume of administration varied based on assigned dose and patient body surface area. The clinical formulation were supplied as a concentrated solution in 40% Solutol® HS-15/60 % propylene glycol in amber vials containing 80 mg of drug in 20 mL (4 mg/mL) and were stored at room temperature. Each vial was designated for single use. The correct volume of drug (at a concentration of 4 mg/mL in the vial) was diluted in dextrose 5% in water (D5W) at a dilution of 1:20 and administered IV peripherally or centrally. Infusion time may be increased as clinically indicated at the direction of the Sponsor. Plinabulin and placebo must be protected from light at all times including storage, dilution and administration. Plinabulin and placebo should be administered within 6 hours of dilution.

**[0129]** Premedication with anti-emetics (including significant anti-emetic regimens such as substance P inhibitor, corticosteroid and/or dopamine antagonist combinations) per institutional practice should be provided before each dose of plinabulin or placebo. 5-HT<sup>3</sup> antagonists should not be routinely administered subsequent to or between doses of plinabulin or placebo unless clearly necessary. Patients should receive a motility enhancing agent such as metaclopromide as part of their antiemetic regimen.

**[0130]** Bowel motility should be maintained per institutional practice as used for drugs such as vincristine, including use of agents such as stool softeners, bulking agents, stimulating agents and/or dopamine antagonists, as well as minimizing use of motility reducing agents such as opiates to when clearly indicated or managing opiate induced constipation with agents such as methylnaltrexone as indicated. If significant constipation develops, it should be

managed immediately and plinabulin or placebo administration should be delayed until resolution. Careful observance for signs of ileus and early diagnostic evaluation with radiographic and/or ultrasound studies is recommended.

**[0131]** If a >20% increase in systolic blood pressure is observed after administration of plinabulin or placebo, oral amlodipine 10 mg or an equivalent calcium channel blocker should be administered one hour before each subsequent dose. Increases in systolic blood pressure above 180 mmHg should be managed with nitroprusside or similar regimen per institutional practice.

#### Docetaxel:

[0132] The initial dose of docetaxel was 75 mg/m<sup>2</sup>. Dose adjustments depended on observed adverse events.

[0133] Volume of administration varied based on assigned dose and patient body surface area.

**[0134]** As a standard approved and commercially available chemotherapy agent, the investigator and study site staff should be experienced in the use of docetaxel and familiar with the formulation and docetaxel prescribing information provided by the manufacturer.

**[0135]** Docetaxel should be obtained from the institutional pharmacy and prepared per institutional protocol. Administration should be carried out with a 1 hour IV infusion per institutional protocol at the dose prescribed by this clinical trial protocol.

**[0136]** Oral dexamethasone (16 mg) was given the day prior to, the day of and the day following docetaxel infusion (Day 1). A similar corticosteroid premedication regimen may be used in accordance with local institutional practices. The dose of dexamethasone or other corticosteroid should be appropriately reduced for patients already utilizing corticosteroids.

#### **PROCEDURES**

**[0137]** Screen: (within 28 days prior to start of treatment (i.e., Day -28 to 1)) Informed consent, medical history and concomitant medications; ECG, radiographic tumor assessments and tumor markers as appropriate.

**[0138]** <u>Baseline Assessments:</u> (within 14 days of start of treatment, i.e., day -14 to 1) Physical examination, vital signs, ECOG performance status, concomitant medication usage, safety laboratory tests. Women of childbearing potential must have a negative serum pregnancy test within 10 days of start of treatment. If there was any history or findings suggestive of significant heart disease, a cardiology consultation should be obtained.

**[0139]** <u>Treatment Phase:</u> Safety assessments (including a complete physical examination) were performed prior to study drug infusion. Safety assessments (including complete physical examination) were performed prior to each subsequent cycle (2+). Additionally, the following were evaluated:

- CBC with differential/platelets and clinical chemistry were performed up to 72 hours prior to Day 1 of each cycle; an additional assessment occurred in Cycle 1/Day 15.
- Vital signs (heart rate, respiratory rate, blood pressure and temperature) were taken on the days of infusion immediately before and after each study drug infusion and at 30 and 60 minutes following last infusion on the first cycle. During subsequent cycles, vitals were taken prior to and after each infusion during the physical examination.

[0140] Assessment of response to treatment occurred during the rest period of the second cycle (and approximately every 2 cycles thereafter).

**[0141]** Treatment continued until there was evidence of progressive disease, unacceptable treatment-related adverse events, the study is closed, -or the patient is withdrawn from the study (either due to withdrawal of consent or Investigator judgment).

**[0142]** End-of Study (Off Study) visit: All patients receiving at least one dose of study drug and discontinuing treatment for any reason except death completed this assessment, within 28 days of the last study drug administration. Patients underwent physical examination, vital signs, body weight, documentation of ECOG performance status, and routine lab tests including a pregnancy test will be drawn.

**[0143]** Follow-up visits: Follow-up visits were required to monitor ongoing drug-related adverse events and survival. Patients with drug related adverse events of Grade  $\geq$  2 observed at the End of Study assessment, should be followed-up monthly until the adverse event has resolved to Grade  $\leq$  1 or the event is believed to be chronic or patient received other anti-cancer therapy. Follow-up for survival should occur at 3-month intervals.

#### **ASSESSMENTS OF:**

**[0144]** <u>EFFICACY:</u> Comparisons were made of efficacy endpoints between Arm D and Arm DN. The primary efficacy endpoint is overall survival. Secondary endpoints include response rate, duration of response, quality of life, neutropenia rate and G-CSF usage.

**[0145]** <u>SAFETY:</u> Adverse events spontaneously declared by the patients or noted during physical examination, vital signs, ECOG performance status and laboratory tests.

#### **STATISTICAL ANALYSES**

**[0146]** Efficacy: The distributions of overall survival and any other time-to-event endpoints were summarized using the Kaplan-Meier method. The log-rank test was used to compare the efficacy endpoints between treatment groups. All statistical tests were carried out using one-sided tests at the 5% level of significance. The primary objective of this trial is to assess the effect of the addition of plinabulin on overall survival.

**[0147]** <u>Safety:</u> All patients were evaluated for safety analysis if they received at least one dose of study drug. The safety data were presented by study arm in individual listings and summary tables, including frequency tables for adverse events and frequency and shift tables for laboratory variables.

# Results:

#### 1) Neutropenia is reduced in the combination of Plinabulin and Docetaxel group

**[0148]** The tolerability profile for plinabulin when added to docetaxel is better than docetaxel alone. Due to unbearable docetaxel side effects, the initial dose of 75 mg/m<sup>2</sup> docetaxel was decreased in 10% of patients (5 of 50) in the plinabulin 30 mg/m<sup>2</sup> plus docetaxel treatment arm whereas, the percent in the matched docetaxel alone arm was much higher at 18.2% of patients (10 of 55). In plinabulin 20 mg/m<sup>2</sup> cohort, the same result was seen, in which a lower proportion of patients required docetaxel dose reductions when treated with the combination (2.5%) than the companion D arms (22.2%).

**[0149]** There was a lower incidence of neutropenia in patients in the DN 30 mg/m<sup>2</sup> arm compared with its companion D arm (8.0% versus 36.4%, p<0.001) and the DN 20 mg/m<sup>2</sup> arm compared with its companion D arm (7.5% versus

22.2%). The DN 30 mg/m² arm (n=50) had a significantly lower incidence of all grades of neutropenia, especially  $\geq$  Grade 3 neutropenia compared with the pooled D arm (n=73) at 8.0% versus 27.4%, respectively (p=0.010). Similar results were observed for the 20 mg/m² arm (5.0% versus 27.4%, respectively; p=0.050). The neutropenia reduction effect is shown in Fig. 5. The proportion of patients who required G-CSF and the rate of docetaxel dose reduction were also lower in both DN arms compared with the D arms. The G-CSF use percentage decrease in combined DN arm vs. combined D arm is statistically significant at 0.0013%.

Table 9: Comparison of Docetaxel dose redution, neutropenia rate and G-CSF use in DN Arm and D Arm

	30 mg/m	<sup>2</sup> Cohort	20 mg/m <sup>2</sup> Cohort			
	DN arm N=50	D arm N=55	DN arm N=40	D arm N=18		
Docetaxel dose reductions (n [%])	5 (10.0%)	10 (18.2%)	1 (2.5%)	4 (22.2%)		
Neutropenia (Grade 3, 4)	8.0% (p=0.01)	27.4%	5% (p=0.05)	27.4%		
G-CSF use (n [%])	7 (14%)	16 (29.1%)	2 (5%)	6 (33.3%)		

Abbreviations: D = docetaxel; DN = docetaxel + plinabulin; G-CSF = granulocyte colony stimulating factor

Note: G-CSF included the following concomitant medications: pegfilgrastim, filgrastim, neupogen, neulasta

## 2) Large Tumor patient population overall survival (OS) benefit from plinabulin and docetaxel combination

**[0150]** In the above study, comparing plinabulin (30 mg/m², Day 1 and Day 8 of each 21-day cycle) plus docetaxel (75 mg/m², Day 1 of each cycle) with docetaxel alone in patients with locally advanced or metastatic NSCLC who had failed at least 1 prior chemotherapy regimen, in the 30 mg/m² group there was no significant difference between the plinabulin plus docetaxel treatment group (OS=8.7 M (month)) compared to the docetaxel control treatment group (OS=7.5 M) in OS (Table 11, Fig. 6), progression-free survival [PFS], and response rate. The inventor then performed extensive analyses of the data to identify a subgroup which react to the plinabulin and docetaxel combination.

**[0151]** Of the subgroup analyses, there is a clear division of OS at tumor size of 3 cm: in the group of at least 1 tumor > 3 cm, > 5cm, or > 7 cm, there was a clear OS benefit in the DN Arm compared with the D Arm, but in all tumor  $\le 3$  cm group, there was no OS difference in (6.45 M vs.6.47 M, Table 11 Fig. 6). The larger the tumor size, the more significant the OS benefit, hazard ratio, and response rate in the combination group compared with the docetaxel group alone (Table 11, Fig. 6).

**[0152]** Thus the inventor has discovered a uniquely defined large tumor group which benefit from the optimum plinabulin and docetaxel combination, which was never reported before in any literature.

Table 10: Comparison of OS and Response Rate in Arm DN (30  $mg/m^2$ ) and Arm D <u>at various size of the largest primary tumor</u>

	All Patients		≤ 3 cm		>3 cm		>5 cm		>7 cm	
Treatment Group	DN	D	DN	D	DN	D	DN	D	DN	D
OS (month)	8.7	7.5	6.5	6.5	9.0	7.5	9.0	6.7	7.32	5.03
Response Rate (%)	14	14.5	12.5	15.8	14.7	13.9	20	15	25	20
Hazard Ratio	0.9	72	0.9	34	0.9		0.7		0.5	

# REFERENCES CITED IN THE DESCRIPTION

# Cited references

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# Patent documents cited in the description

- WO2004054498A [0043]
- <u>US77006158</u> [0077]
- <u>US7629380B</u> [0077]

#### Non-patent literature cited in the description

- MITA AC et al. Journal of Clinical Oncology 15Suppl (A7592) discloses the use of plinabulin in combination with docetaxel for treating cancer Vainas, 2010, [0006]
- British J. Cancer, vol. 107, 814-822 [0006]

# **PATENTKRAV**

5

10

- 1. Plinabulin til anvendelse til reducering af neutropeni hyppigheden af en grad 3 eller 4 neutropeni hos et individ hvor der administreres med 75 mg/m² docetaxel, hvor plinabulinen administreres intravenøst ved ca. 20 mg/m² til ca. 30 mg/m², hvor plinabulinen administreres inden for 1 time og 24 timer efter administration af docetaxel.
- 2. Plinabulin til anvendelse ifølge krav 1, hvor dosis af plinabulin er ca.  $20 \text{ mg/m}^2$  eller ca.  $30 \text{ mg/m}^2$ .

3. Plinabulin til anvendelse ifølge krav 1 eller 2, hvor individet har ikke-småcellet lungecancer (NSCLC).

# **DRAWINGS**

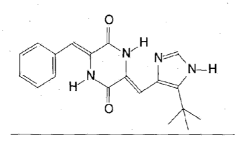
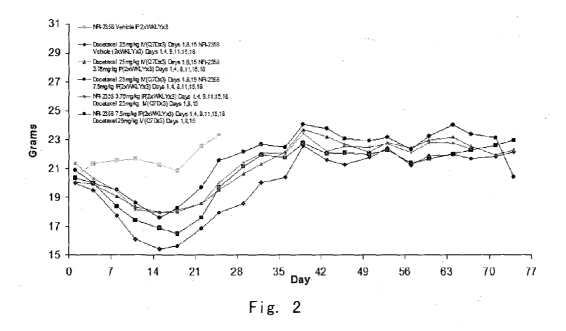
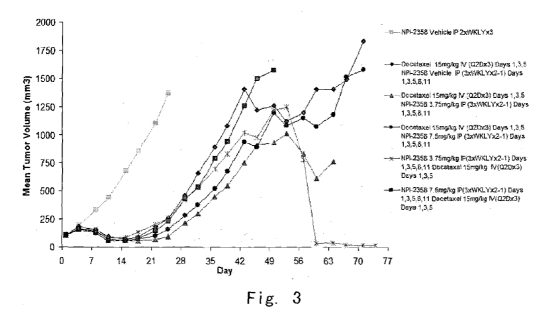
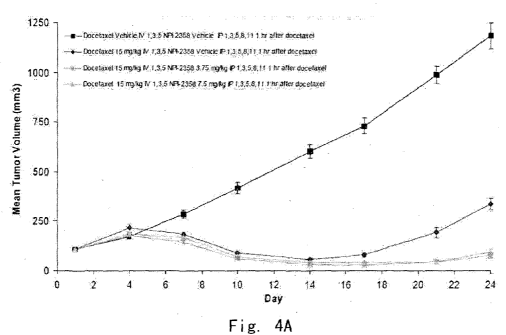
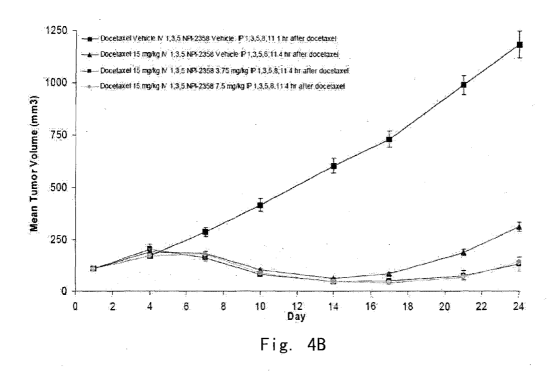


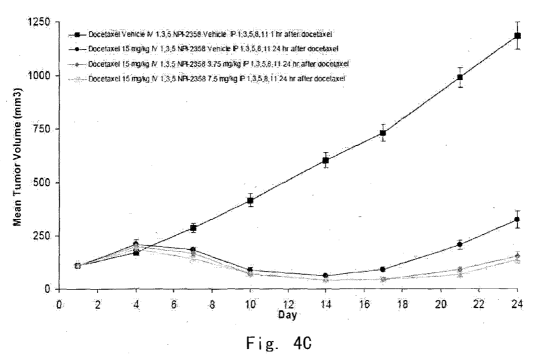
Fig. 1











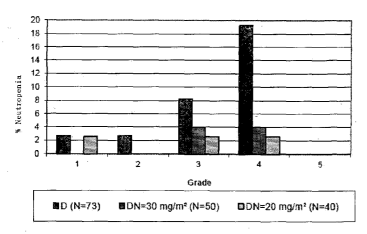
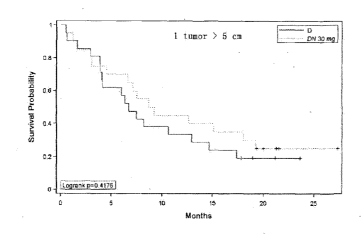


Fig. 5



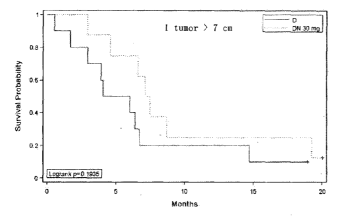


Fig. 6