



HU000025187T2

(19) **HU****MAGYARORSZÁG**
Szellemi Tulajdon Nemzeti Hivatala(11) Lajstromszám: **E 025 187**(13) **T2****EURÓPAI SZABADALOM**
SZÖVEGÉNEK FORDÍTÁSA(21) Magyar ügyszám: **E 11 749393**(22) A bejelentés napja: **2011. 08. 16.**

(96) Az európai bejelentés bejelentési száma:

EP 20110749393

(97) Az európai bejelentés közzétételi adatai:

EP 2605764 A1 **2012. 02. 23.**

(97) Az európai szabadalom megadásának meghirdetési adatai:

EP 2605764 B1 **2015. 03. 18.**(51) Int. Cl.: **A61K 314/37** (2006.01)**A61K 3147/45** (2006.01)**A61K 315/17** (2006.01)**A61K 393/95** (2006.01)**A61K 45/06** (2006.01)**A61P 35/00** (2006.01)

(86) A nemzetközi (PCT) bejelentési szám:

PCT/EP 11/064050

(87) A nemzetközi közzétételi szám:

WO 12022724

(30) Elsőbbségi adatok:

374296 P**2010. 08. 17.****US**

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Kompozíciók rák kezelésére

Az európai szabadalom ellen, megadásának az Európai Szabadalmi Közlönyben való meghirdetésétől számított kilenc hónapon belül, felszólalást lehet benyújtani az Európai Szabadalmi Hivatalnál. (Európai Szabadalmi Egyezmény 99. cikk(1))

A fordítást a szabadalmas az 1995. évi XXXIII. törvény 84/H. §-a szerint nyújtotta be. A fordítás tartalmi helyességét a Szellemi Tulajdon Nemzeti Hivatala nem vizsgálta.



(11)

EP 2 605 764 B1

(12) **EUROPEAN PATENT SPECIFICATION**

(45) Date of publication and mention
of the grant of the patent:
18.03.2015 Bulletin 2015/12

(51) Int Cl.:
A61K 31/437 ^(2006.01) **A61K 31/4745** ^(2006.01)
A61K 31/517 ^(2006.01) **A61K 39/395** ^(2006.01)
A61K 45/06 ^(2006.01) **A61P 35/00** ^(2006.01)

(21) Application number: **11749393.2**

(86) International application number:
PCT/EP2011/064050

(22) Date of filing: **16.08.2011**

(87) International publication number:
WO 2012/022724 (23.02.2012 Gazette 2012/08)

(54) **Compositions for the treatment of cancer**

Zusammensetzungen zur Behandlung von Krebs

Les compositions pour le traitement du cancer

(84) Designated Contracting States:
**AL AT BE BG CH CY CZ DE DK EE ES FI FR GB
GR HR HU IE IS IT LI LT LU LV MC MK MT NL NO
PL PT RO RS SE SI SK SM TR**

(30) Priority: **17.08.2010 US 374296 P**

(43) Date of publication of application:
26.06.2013 Bulletin 2013/26

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Description**Field of the Invention**

5 [0001] The present invention relates to a combination therapy for treating a patient suffering from a proliferative disorder, in particular a solid tumor, for example, colorectal cancer, melanoma, and thyroid cancer, comprising administering to the patient propane-1-sulfonic acid {3-[5-(4-chlorophenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluorophenyl}-amide and a EGFR inhibitor, i.e. erlotinib or cetuximab.

Background of the Invention

10 [0002] Normally functioning b-Raf is a kinase which is involved in the relay of signals from the cell membrane to the nucleus and is active only when it is needed to relay such signals. Mutant b-Raf having the V600E mutation, however, is constantly active and thus plays a role in tumor development. Such mutant b-Raf has been implicated in various tumors, for example, colorectal cancer, melanoma, and thyroid cancer.

15 [0003] Propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide (hereafter also referred to as "Compound I") is a b-raf kinase inhibitor that specifically targets mutant b-Raf having the V600E mutation. This compound is described in WO 2007/002325. Accordingly, such an inhibitor is used in the inhibition of tumors, particularly solid tumors, for example, colorectal cancer, melanoma, and thyroid cancer, which comprise b-Raf having a V600 mutation, preferably the V600E mutation.

20 [0004] Protein tyrosine kinases (PTKs) catalyze the phosphorylation of tyrosyl residues in various proteins involved in the regulation of cell growth and differentiation (Wilks et al., Progress in Growth Factor Research 97 (1990) 2; Chan, A. C., and Shaw, A. S., Curr. Opin. Immunol. 8 (1996) 394-401). Such PTKs can be divided into receptor tyrosine kinases (e.g. EGFR/HER-1, c-erbB-2/HER-2, c-met, PDGFr, FGFr) and non-receptor tyrosine kinases (e.g. src, lck).

25 [0005] It is known that receptor tyrosine kinases of the HER-family like HER-2 and EGFR (HER-1) are frequently aberrantly expressed in common human cancers such as breast cancer, gastrointestinal cancer (colon, rectal or stomach cancer), thyroid, leukemia and ovarian, bronchial and pancreatic cancer and melanoma. High levels of these receptors correlate with poor prognosis and response to treatment (Wright, C., et al., Br. J. Cancer 65 (1992) 118-121).

30 [0006] Inhibitors of PTKs, and in particular of EGFR, have been developed. Tumors comprising b-Raf having the V600E mutation, however, have been known to be resilient to treatment with EGFR inhibitors. See Prewett et al., Clin. Cancer Res. (2002), 8:994-1003 and Ouchi et al., Cancer Chemother. Pharmacol. (2006), 57:693-702. Applicants have unexpectedly found, however, that combination therapy with Compound I and an EGFR inhibitor not only is capable of reducing such resilience but also results in improved antineoplastic effects that are significantly superior to the results obtained with each compound alone without a significant increase in toxicity.

35 [0007] In addition to EGFR inhibitors, topoisomerase inhibitors are also antiproliferative agents. Tumors containing the V600E mutation, however, have also been known to be resilient to treatment with topoisomerase inhibitors. See Prewett et al., Clin. Cancer Res. (2002), 8:994-1003 and Abal et al., Oncogene (2004), 23:1737-44. Applicants have unexpectedly found, however, that the combination of Compound I with an EGFR inhibitor and a topoisomerase inhibitor not only is capable of reducing such resilience but also results in improved antineoplastic effects that are significantly superior to the results obtained with each compound alone or with the aforementioned combination therapy of Compound I and EGFR inhibitor without a significant increase in toxicity.

Summary of the Invention

45 [0008] The scope of the present application is defined by appended claims.

[0009] The present invention relates to a pharmaceutical product comprising (A) a first component which comprises, as an active agent, propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b]pyridine-3-carbonyl]-2,4-difluorophenyl}-amide, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor erlotinib or cetuximab; as a combined preparation for the simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation.

[0010] The present invention also relates to a kit comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor erlotinib or cetuximab.

55 [0011] The present invention further relates to a pharmaceutical product comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor erlotinib or cetuximab; and optionally (C) a third component which comprises a topoisomerase inhibitor irinotecan or a pharmaceutically acceptable salt thereof; as a combined preparation for the simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation.

[0012] In addition, the present invention relates to the combination comprising Compound I, or a pharmaceutically-acceptable salt thereof, and an EGFR inhibitor erlotinib or cetuximab for use in the treatment of cancer comprising b-Raf having a V600 mutation.

[0013] A yet further aspect of the present invention is the use of Compound I, or a pharmaceutically-acceptable salt thereof, and an EGFR inhibitor erlotinib or cetuximab for the preparation of a medicament for the treatment of cancer comprising b-Raf having a V600 mutation.

Brief Description of the Drawings

[0014]

Figure 1 illustrates the tolerability, as demonstrated by % body weight change, of Compound I monotherapy at 75 mg/kg bid, erlotinib hydrochloride monotherapy at 67 mg/kg qd, erlotinib hydrochloride monotherapy at 100 mg/kg qd, and Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd combination therapy.

Figure 2 illustrates the antitumor activity, as demonstrated by the change in mean tumor volume over time, of Compound I monotherapy at 75 mg/kg bid, erlotinib hydrochloride monotherapy at 67 mg/kg qd, erlotinib hydrochloride monotherapy at 100 mg/kg qd, and Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd combination therapy.

Figure 3 illustrates the effect on survival, as demonstrated by percentage of surviving mice over time, of Compound I monotherapy at 75 mg/kg bid, erlotinib hydrochloride monotherapy at 67 mg/kg qd, erlotinib hydrochloride monotherapy at 100 mg/kg qd, and Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd combination therapy.

Figure 4 illustrates the tolerability, as demonstrated by % body weight change, of Compound I monotherapy at 75 mg/kg bid, Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy, and Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy.

Figure 5 illustrates the antitumor activity, as demonstrated by the change in mean tumor volume over time, of Compound I monotherapy at 75 mg/kg bid, Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy, and Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy.

Figure 6 illustrates the effect on survival, as demonstrated by percentage of surviving mice over time, of Compound I monotherapy at 75 mg/kg bid, Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy, and Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy.

Figure 7 illustrates the tolerability, as demonstrated by % body weight change, of Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, irinotecan HCl monotherapy at 40 mg/kg q4dx5, Compound I at 25 mg/kg bid and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy, and Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan HCl at 40 mg/kg q4dx5 combination therapy.

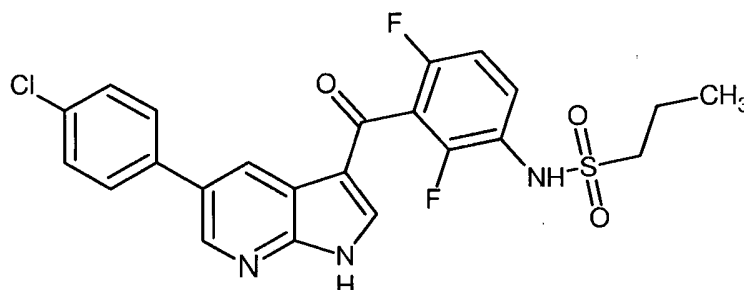
Figure 8 illustrates the antitumor activity, as demonstrated by the change in mean tumor volume over time, of Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, irinotecan HCl monotherapy at 40 mg/kg q4dx5, Compound I at 25 mg/kg bid and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy, and Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan HCl at 40 mg/kg q4dx5 combination therapy.

Figure 9 illustrates the effect on survival, as demonstrated by percentage of surviving mice over time, of Compound I monotherapy at 25 mg/kg bid, cetuximab monotherapy at 40 mg/kg 2x/wk, irinotecan HCl monotherapy at 40 mg/kg q4dx5, Compound I at 25 mg/kg bid and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 combination therapy, Compound I at 25 mg/kg bid and cetuximab

at 40 mg/kg 2x/wk combination therapy, and Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan HCl at 40 mg/kg q4dx5 combination therapy.

Detailed Description of the Invention

[0015] As stated above, "Compound I" shall herein refer to propane-1-sulfonic acid {3-[5-(4-chlorophenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide. This is a compound having the following structure.



[0016] Compound I is a b-Raf kinase inhibitor that specifically targets b-Raf having the V600E mutation.

[0017] The "V600E" mutation of b-Raf, as used herein, refers to a mutation in the b-Raf protein wherein the valine residue at residue position 600 of b-Raf is replaced by glutamic acid.

[0018] As used herein, when referring to the receptor tyrosine kinases of the HER-family like HER-2 and EGFR (HER-1), the acronym "HER" refers to human epidermal receptor and the acronym "EGFR" refers to epidermal growth factor receptor.

[0019] The term "combined preparation" as used herein means simultaneous or sequential combination. The active agents used in the present combinations, for example compound I, or a topoisomerase inhibitor such irinotecan, or an EGFR inhibitor erlotinib or a pharmaceutically acceptable salt, or cetuximab can be used in any applicable dosage form, such as for example tablets, capsules, solutions, suspensions and the like, depending on the properties of the selected active ingredient. Compound I, or a pharmaceutically acceptable salt thereof, may, for example, be administered orally. Erlotinib, or a pharmaceutically acceptable salt thereof, may, for example, be administered orally. Cetuximab may, for example, be administered intraperitoneally or intravenously. Irinotecan HCl or the pharmaceutically acceptable salt thereof may, for example, be administered intraperitoneally or intravenously. Certain combinations disclosed herein may show a therapeutic effect, which is more than additive (synergistic).

[0020] As used herein, the term "pharmaceutically acceptable carrier" indicates that the indicated carrier does not have properties that would cause a reasonably prudent medical practitioner to avoid administration thereof to a patient, taking into consideration the disease or conditions to be treated and the respective route of administration.

[0021] As used herein, the term "pharmaceutically acceptable salt" of a compound refers to any conventional salt or base addition salt that retains the biological effectiveness and properties of the compound and which is formed from a suitable non-toxic organic or inorganic acid or organic or inorganic base. As used herein, the term "therapeutically effective" means an amount of drug, or combination or composition, which is effective for producing a desired therapeutic effect upon administration to a patient, for example, to stem the growth, or result in the shrinkage, of a cancerous tumor or to increase the patient's life span.

[0022] The terms "cell proliferative disorder" and "proliferative disorder" refer to disorders that are associated with some degree of abnormal cell proliferation. In one embodiment, the proliferative disorder is cancer.

[0023] The terms "cancer" and "cancerous" refer to or describe the physiological condition in mammals that is typically characterized by unregulated cell growth/proliferation. Examples of cancer include, but are not limited to, colorectal cancer, melanoma, and thyroid cancer.

[0024] The term "colorectal tumor" or "colorectal cancer" refers to any tumor or cancer of the large bowel, which includes the colon (the large intestine from the cecum to the rectum) and the rectum, including, e.g., adenocarcinomas and less prevalent forms, such as lymphomas and squamous cell carcinomas.

[0025] "Inhibiting cell growth or proliferation" means decreasing a cell's growth or proliferation by at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, 95%, or 100%, and includes inducing cell death.

[0026] The phrase "substantially reduced" or "substantially different," as used herein, refers to a sufficiently high degree of difference between two numeric values (generally one associated with a molecule and the other associated with a reference/comparator molecule) such that one of skill in the art would consider the difference between the two values to be of statistical significance within the context of the biological characteristic measured by said values.

[0027] The term "tumor" refers to all neoplastic cell growth and proliferation, whether malignant or benign, and all pre-

cancerous and cancerous cells and tissues. The terms "cancer," "cancerous," "cell proliferative disorder," "proliferative disorder," and "tumor" are not mutually exclusive as referred to herein.

[0028] "Regression" of a tumor is said to occur following treatment when the volume of said tumor is reduced. If the tumor remains present (tumor volume > 0 mm³) but its volume is reduced from what it was at the initiation of treatment, "partial regression" (PR) is said to have occurred. If the tumor is palpably absent following treatment, "complete regression" (CR) is said to have occurred.

[0029] The term "small molecule" as used herein is a chemical compound with a molecular weight of less than 1000 g/mol, preferably less than 700 g/mol. Small molecules according to the present inventions can be obtained by chemical reactions known to the person of skill in the art of organic chemistry and/or medicinal chemistry. Examples of small molecules may be, without limitation, the compound I or the compound known as erlotinib, preferably erlotinib hydrochloride.

[0030] The term "large molecule" as used herein refers to chemical compounds with a molecular weight of more than 1000 g/mol. Preferably "large molecules" are compounds that can be obtained via biotechnological production processes, such as fermentation. More preferably, the term "large molecule" refers to polypeptides, such as for example antibodies, more specifically monoclonal antibodies. One example of a large molecule according to the present invention is cetuximab.

[0031] The present invention relates to a pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor erlotinib or cetuximab; the amount of said active agents being such that the combination thereof is therapeutically-effective in the treatment of cancer comprising b-Raf having a V600 mutation. The present invention further relates to the above-mentioned product as a combined preparation for the simultaneous or sequential use in the treatment of cancer comprising b-Raf carrying the V600E mutation.

[0032] In one aspect a method of treating a patient suffering from a proliferative disorder, comprising administering to the patient the combination or pharmaceutical preparations as mentioned above, is disclosed.

[0033] "Treatment of a proliferative disorder" shall be understood to include maintaining or decreasing tumor size, inducing tumor regression (either partial or complete), inhibiting tumor growth, and/or increasing the life span of a patient suffering from said disorder.

[0034] The present invention also relates to a kit or a composition comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor erlotinib or cetuximab. The kit or composition may be used, for example, in the treatment of cancer comprising b-Raf having a V600 mutation.

[0035] In an embodiment of the invention, the proliferative disorder is a solid tumor, in particular colorectal cancer, melanoma and/or thyroid cancer.

[0036] In another embodiment of the invention, the proliferative disorder is a tumor comprising b-Raf having a V600 mutation, preferably the V600E mutation.

[0037] In a further embodiment of the invention, the proliferative disorder is selected from the group consisting of colorectal cancer, melanoma, and thyroid cancer and the cancer involves a tumor comprising b-Raf having a V600 mutation, preferably the V600E mutation.

[0038] In yet a further embodiment of the invention, the proliferative disorder is a solid tumor comprising b-Raf having a V600 mutation, preferably the V600E mutation.

[0039] In yet a further embodiment of the invention, the proliferative disorder is colorectal cancer.

[0040] In yet a further embodiment of the invention, the proliferative disorder is colorectal cancer involving a tumor comprising b-Raf having a V600 mutation, preferably the V600E mutation.

[0041] In yet a further embodiment of the invention, the EGFR inhibitor is a small molecule EGFR inhibitor. In one such embodiment, the EGFR inhibitor is erlotinib, or a pharmaceutically acceptable salt thereof, for example, erlotinib hydrochloride (erlotinib HCl). Erlotinib HCl is also known under its trade name Tarceva®, and is for example sold in the U.S. by Genentech, South San Francisco, U.S.A.

[0042] In yet a further embodiment of the invention, the EGFR inhibitor is a large molecule EGFR inhibitor, for example, an antibody that targets EGFR. In one such embodiment, the EGFR inhibitor may be a monoclonal antibody that targets EGFR, for example, cetuximab. Cetuximab is also known under its trade name Erbitux® and is for example sold in the U.S. by ImClone Systems, Inc., New York, U.S.A.

[0043] In certain embodiments the present invention relates to a pharmaceutical product for the treatment of colorectal cancer involving a tumor comprising b-Raf having the V600E mutation, comprising (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, erlotinib, or a pharmaceutically acceptable salt thereof; the amount of said active agents being such that the combination thereof is therapeutically-effective in the treatment of said cancer.

[0044] In a further embodiment the present invention relates to a pharmaceutical product for the treatment of colorectal cancer involving a tumor comprising b-Raf having the V600E mutation, comprising (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which

comprises, as an active agent, cetuximab; the amount of said active agents being such that the combination thereof is therapeutically-effective in the treatment of said cancer.

[0045] The amount of each component administered by the present combinations or compositions may, but does not have to be therapeutically effective by itself. That is, this invention specifically contemplates combinations wherein the amount of Compound I, or a pharmaceutically-acceptable salt thereof, and/or the amount of EGFR inhibitor erlotinib or cetuximab, in the combination may be less than the amount that is therapeutically-effective for each active agent when said agent is administered in monotherapy.

[0046] The first component (A) and the second component (B) of the present invention are administered in any amount and for any duration that the combined amounts thereof are therapeutically effective in treating a proliferative disorder.

[0047] In certain embodiments of the present invention, Compound I, or a pharmaceutically acceptable salt thereof, is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, or from about 1700 mg/day to about 2100 mg/day. In yet another embodiment, the dosage amount is about 1920 mg/day.

[0048] In another embodiment of the present invention, the foregoing amounts of Compound I, or a pharmaceutically acceptable salt thereof, may be administered as a single dose daily or divided, for example into equal doses (though this is not required), and administered twice daily (bid). For example, Compound I, or a pharmaceutically acceptable salt thereof, may be administered in a dosage amount of from about 100 mg to about 1500 mg bid, from about 500 mg to about 1250 mg bid, from about 850 mg to about 1050 mg bid, or about 960 mg bid.

[0049] In an embodiment of the present invention, the administration of Compound I, or a pharmaceutically acceptable salt thereof, occurs until disease progression or unacceptable toxicity.

[0050] In certain embodiments of the present invention, erlotinib, or a pharmaceutically acceptable salt thereof, is administered at a dosage amount of from about 20 mg/day to about 500 mg/day, from about 100 mg/day to about 400 mg/day, or from about 100 mg/day to about 200 mg/day.

[0051] In another embodiment of the present invention, the administration of erlotinib, or a pharmaceutically acceptable salt thereof, occurs until disease progression or unacceptable toxicity.

[0052] In yet another embodiment of the present invention, cetuximab is administered at a dosage amount of from about 50 mg/m²/week to about 700 mg/m²/week, from about 100 mg/m²/week to about 600 mg/m²/week, or from about 200 mg/m²/week to about 500 mg/m²/week.

[0053] In yet another embodiment, cetuximab is administered weekly with the first administration being in an amount of from about 400 mg/m² to about 500 mg/m² and each subsequent administration being in an amount of from about 200 mg/m² to about 300 mg/m².

[0054] In still another embodiment, cetuximab is administered weekly with the first administration being in an amount of about 450 mg/m² and each subsequent administration being in an amount of about 250 mg/m².

[0055] In still another embodiment of the present invention, the administration of cetuximab occurs until disease progression or unacceptable toxicity.

[0056] Therefore, in another embodiment, the present invention provides a pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, erlotinib or a pharmaceutically acceptable salt thereof, as a combined preparation for simultaneous or sequential use in the treatment of a proliferative disorder, wherein

(A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day, and

(B) is administered in an amount of from about 20 mg/day to about 500 mg/day, from about 100 mg/day to about 400 mg/day, or from about 100 mg/day to about 200 mg/day.

[0057] Within this embodiment, Compound I or a pharmaceutically acceptable salt thereof, may be administered twice daily. The proliferative disorder to be treated in this way is a solid tumor, in particular colorectal cancer, melanoma, and thyroid cancer, comprising b-Raf having the V600E mutation. More particularly, the proliferative disorder is colorectal cancer involving a tumor comprising b-Raf having the V600E mutation.

[0058] Also, within this embodiment the product of the present invention may comprises compound I or a pharmaceutically acceptable salt thereof, being administered orally in a dosage amount of from about 850 mg to about 1050 mg twice daily or about 960 mg twice daily, and erlotinib or a pharmaceutically acceptable salt thereof, being administered orally in a dosage amount of from about 100 mg/day to about 400 mg/day, or from about 100 mg/day to about 200 mg/day. Both agents may, for example, be administered until disease progression or unacceptable toxicity.

[0059] The present invention also provides a pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, cetuximab; as a combined preparation for simultaneous or sequential use in the treatment of a proliferative disorder, wherein

(A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day; and
 (B) is administered in an amount of from about 50 mg/m²/week to about 700 mg/m²/week, from about 100 mg/m²/week to about 600 mg/m²/week, or from about 200 mg/m²/week to about 500 mg/m²/week.

[0060] Within this embodiment, the proliferative disorder to be treated in this way is a solid tumor, in particular colorectal cancer, melanoma, and thyroid cancer, comprising b-Raf having the V600E mutation. More particularly, the proliferative disorder is colorectal cancer involving a tumor comprising b-Raf having the V600E mutation.

[0061] Also, within this embodiment, the product of the present invention may comprises compound I or a pharmaceutically acceptable salt thereof, being administered orally in a dosage amount of from about 850 mg to about 1050 mg twice daily or about 960 mg twice daily; and cetuximab being administered intravenously in a dosage amount of from about 200 mg/m²/week to about 500 mg/m²/week. In one embodiment, cetuximab is administered initially as a 400 mg/m² dose as a 120-minute intravenous infusion, followed after a week by 250 mg/m² doses intravenously infused over 60 minutes once weekly. Both agents may, for example, be administered until disease progression or unacceptable toxicity.

[0062] The present invention also further provides a kit or a composition comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, erlotinib, or a pharmaceutically-acceptable salt thereof.

[0063] The present invention also further provides a kit or a composition comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, cetuximab.

[0064] In another aspect of this invention, the pharmaceutical product comprising the components (A) and (B) described above is administered in conjunction with radiotherapy and/or in conjunction with the administration of another active agent.

[0065] Thus, in certain embodiments the present invention provides a pharmaceutical product comprising the components (A) and (B) as described above and a third component (C) which comprises, as an active agent, a topoisomerase inhibitor; as a combined preparation for the simultaneous or sequential use in the treatment of a proliferative disorder such as solid tumors comprising b-Raf having a V600 mutation, preferably the V600E mutation. As previously stated, the amount of each component administered with the present combination product may, but does not have to be therapeutically effective by itself and this invention specifically contemplates combinations wherein the amount of each of the active agents in the combination may be less than the amount that is therapeutically-effective for each active agent when said agent is administered in monotherapy.

[0066] In an embodiment of the present invention, the topoisomerase inhibitor is an inhibitor of type I topoisomerase. In an embodiment of the invention, the topoisomerase inhibitor is irinotecan, or a pharmaceutically acceptable salt thereof, for example, irinotecan hydrochloride (irinotecan HCl). Irinotecan HCl is sold as Camptosar® by Pfizer Inc., New York, U.S.A. Irinotecan, or the pharmaceutically acceptable salt thereof, may, for example, be administered intraperitoneally or intravenously.

[0067] In an embodiment of the present invention, irinotecan, or a pharmaceutically acceptable salt thereof, is administered in an amount of from about 1 to about 400 mg/m²/week, or from about 1 to about 250 mg/m²/week. In another embodiment, irinotecan, or a pharmaceutically acceptable salt thereof, is administered at a dosage amount of from about 50 to about 200 mg/m²/week. In yet another embodiment, irinotecan, or a pharmaceutically acceptable salt thereof, is administered at a dosage amount of about 125 mg/m²/week.

[0068] In yet another embodiment, dosing of irinotecan, or a pharmaceutically acceptable salt thereof, is with a six week cycle at about 75 to about 175 mg/m² weekly, for example about 125 mg/m² weekly, for the first four weeks, for example on days 1, 8, 15, and 22. In another embodiment, dosing of irinotecan is with a six week cycle at about 130 to about 230 mg/m² weekly, for example about 180 mg/m² weekly, every two weeks starting on the first week, for example on days 1, 15, and 29. In yet another embodiment, dosing of irinotecan is once every three weeks at about from 300 to about 400 mg/m², for example about 350 mg/m². In yet another embodiment, dosing of irinotecan is once every two weeks at about 130 to about 230 mg/m², for example about 180 mg/m². Dosing may be by infusion, for example, over about 90 minutes. Treatment may be until disease progression or unacceptable toxicity.

[0069] The dosage levels of each of the components (A), (B) or (C) may be modified by the physician to be lower or higher than that stated herein depending on the needs of the patient, and the reaction of the patient to the treatment. The dosages may be administered according to any dosage schedule determined by the physician in accordance with the requirements of the patient. For example, the dosages of each of the components may be administered in single or in divided doses over a period of several days, or alternating daily schedules.

[0070] The present invention also provides a pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I or a pharmaceutically-acceptable salt thereof; (B) a second component which comprises, as an active agent, cetuximab; and (C) a third component which comprises, as an active agent, irinotecan or a pharmaceutically-acceptable salt thereof.

ceutically-acceptable salt thereof; as a combined preparation for simultaneous or sequential use in the treatment of a proliferative disorder, wherein

(A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day;

(B) is administered in an amount of from about 50 mg/m²/week to about 700 mg/m²/week, from about 100 mg/m²/week to about 600 mg/m²/week, or from about 200 mg/m²/week to about 500 mg/m²/week; and

(C) is administered in an amount of from about 1 to about 250 mg/m²/week, about 50 to about 200 mg/m²/week, or about 125 mg/m²/week.

[0071] Within this embodiment, the proliferative disorder to be treated in this way is a solid tumor, in particular colorectal cancer, melanoma, and thyroid cancer, comprising b-Raf having the V600E mutation. More particularly, the proliferative disorder is colorectal cancer involving a tumor comprising b-Raf having the V600E mutation.

[0072] Also, within this embodiment the product of the present invention comprises compound I, or a pharmaceutically acceptable salt thereof, being administered orally in a dosage amount of from about 850 mg to about 1050 mg twice daily or about 960 mg twice daily; and cetuximab being administered intravenously in a dosage form and a dosage amount of from about 200 mg/m²/week to about 500 mg/m²/week; and irinotecan being administered intravenously in a dosage amount of from about 50 to about 200 mg/m²/week, or about 125 mg/m²/week. All agents may, for example, be administered until disease progression or unacceptable toxicity.

[0073] The present invention also further provides a kit or a composition comprising: (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; (B) a second component which comprises, as an active agent, cetuximab; and (C) a third component which comprises, as an active agent, irinotecan, or a pharmaceutically-acceptable salt thereof.

[0074] Compound I exists in its natural state in a crystalline form. However, the amorphous form of the compound has greater solubility in water as compared with the crystalline form and thus has an improved dissolution rate and, therefore, improved bioavailability as compared to the crystalline form. As such, the amorphous form of the compound is preferred. Accordingly, in preferred embodiments of the method and kit of the present invention, Compound I is in substantially amorphous form and, more preferably, in amorphous form. As used herein, the term "substantially amorphous" material embraces material which has no more than about 10% crystallinity; and "amorphous" material embraces material which has no more than about 2% crystallinity.

[0075] In an embodiment of the present invention, Compound I is contained in a solid molecular complex formed with hydroxypropyl methyl cellulose acetate succinate (HPMC-AS). As used herein, the term "solid molecular complex" means a composition wherein Compound I is randomly distributed ("molecularly dispersed") within a matrix formed by HPMC-AS. Preferably such composition of compound I and HPMC-AS form a one phase system, which can be characterized by X-ray powder diffraction patterns which are substantially free, or free, of crystalline signals related to crystalline form of compound I. In certain embodiments Compound I is present in the polymer in a final state of subdivision. In certain embodiments, Compound I is molecularly dispersed within the HPMC-AS matrix such that it is immobilized in its amorphous form. By "immobilized", it is meant that the molecules of Compound I interact with molecules of HPMC-AS in such a way that they are held in the aforementioned matrix and prevented from crystal nucleation due to lack of mobility. In some embodiments the polymer may prevent intramolecular hydrogen bonding or weak dispersion forces between two or more molecules of Compound I.

[0076] In some embodiments the ratio of the amount by weight of Compound I within the solid molecular complex to the amount by weight of HPMC-AS therein is from about 1:9 to about 5:5. In an embodiment, said ratio is from about 2:8 to about 4:6. In another embodiment, said ratio is about 3:7.

[0077] In certain embodiments of the method and kit of the present invention, the first component comprises the aforementioned solid molecular complex of Compound I and HPMC-AS blended with colloidal silicon dioxide. In certain embodiments, the blend is at least 0.5% by weight silicon dioxide. In an embodiment of the present invention, the blend is about 97% complex and about 3% silicon dioxide.

[0078] In another embodiment, the first component includes a composition comprising the aforementioned solid molecular complex, either blended or not blended with silicon dioxide as described above, and a pharmaceutically acceptable carrier. In certain embodiments, the aforementioned complex or blend comprising the same is suspended in the carrier. An example of a carrier is hydroxypropylcellulose (HPC). In an embodiment, the vehicle contains about 2% by weight HPC.

[0079] Each component may also contain additional agents such as preserving agents, solubilizing agents, stabilizing agents, wetting agents, emulsifying agents, sweetening agents, coloring agents, flavoring agents, salts for varying the osmotic pressure, buffers, coating agents and antioxidants.

[0080] In certain embodiments, the first component may comprise a solid molecular complex of Compound I and HPMC-AS blended with colloidal silicon dioxide, hydroxypropylcellulose, Croscopovidone (a disintegrating agent), magnesium stearate (a lubricant that may be used in tablet and capsulation operations), and/or croscarmellose sodium (a

disintegrating agent).

[0081] In an embodiment, the first component is a hard gelatin capsule comprising a solid molecular complex of Compound I and HPMC-AS blended with colloidal silicon dioxide, hydroxypropylcellulose, magnesium stearate, and croscarmellose sodium.

[0082] In an embodiment, the first component is a tablet comprising Compound I, or a pharmaceutically acceptable salt thereof. In an embodiment, the tablet comprises a solid molecular complex of Compound I, or a pharmaceutically acceptable salt thereof, and HPMC-AS. The complex may, for example, be blended with colloidal silicon dioxide, hydroxypropylcellulose, magnesium stearate, and croscarmellose sodium. The tablet may, for example, be coated with a film coating. The film coating may, for example, comprise polyvinyl alcohol, titanium dioxide, polyethylene glycol 3350, talc, and iron oxide red.

[0083] In certain embodiments, the second component may comprise cetuximab in solution. In an embodiment, the solution is about 2 mg/ml cetuximab.

[0084] In certain embodiments, the second component may comprise a tablet comprising erlotinib, or a pharmaceutically-acceptable salt thereof, for example erlotinib hydrochloride.

[0085] In certain embodiments, the third component may comprise a solution comprising irinotecan, or a pharmaceutically acceptable salt thereof, for example irinotecan hydrochloride. In an embodiment, the solution is an about 5% dextrose solution. In an embodiment, each ml of the solution contains about 20 mg irinotecan hydrochloride, about 45 mg sorbitol, and about 0.9 mg lactic acid. In an embodiment, the solution has a pH of from about 3.0 to about 3.8, for example, about 3.5.

[0086] In addition, the present invention provides the composition comprising Compound I, or a pharmaceutically-acceptable salt thereof, and an EGFR inhibitor erlotinib or cetuximab, optionally also in combination with a topoisomerase inhibitor irinotecan, for the treatment of a proliferative disorder, in particular a solid tumor, more particularly colorectal cancer, melanoma and/or thyroid cancer, all comprising b-Raf having the V600E mutation.

[0087] The invention further provides the composition comprising Compound I, or a pharmaceutically-acceptable salt thereof, and an EGFR inhibition erlotinib or cetuximab, optionally also in combination with a topoisomerase inhibitor irinotecan, for the preparation of a medicament for the treatment of a proliferative disorder, in particular a solid tumor, more particularly colorectal cancer, melanoma and/or thyroid cancer, all comprising b-Raf having the V600E mutation.

[0088] In one aspect methods for treating a patient suffering from a proliferative disorder, in particular a solid tumor, more particularly colorectal cancer, melanoma and/or thyroid cancer, all comprising b-Raf having the V600E mutation, comprising administering to said patient any of the combinations of (A) and (B) or (A), (B) and (C) together with the dosages and dosage schemes as disclosed herein before, are also disclosed.

[0089] Applicants have conducted studies using mice containing a human colorectal cancer xenograft. Applicants found that the combination of Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd produced tumor growth inhibition (TGI) and increased life span (ILS) results that were significantly better than correlative monotherapy results at $p < 0.05$ as well as results achieved with erlotinib hydrochloride monotherapy at 100 mg/kg qd. In addition, 9 out of the 10 mice subjected to the combination therapy had partial regressions whereas no regressions (partial or complete) were observed with any of the monotherapy groups.

These studies indicate that treating patients with a combination of Compound I and erlotinib hydrochloride is superior to treatment with either agent alone. Further, they indicate that combining the two agents allows for at least reduction in the dose of erlotinib hydrochloride to obtain equivalent or better results.

[0090] Applicants also found that the combination of Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk growth inhibition (TGI) and increased life span (ILS) results that were significantly better than correlative monotherapy results at $p < 0.05$ and also better than the results achieved with monotherapy of Compound I at 75 mg/kg bid. Applicants also found that both the combination of Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk produced TGI and ILS results that were significantly better than correlative monotherapy results at $p < 0.05$ and also better than the results achieved with monotherapy of Compound I at 25 mg/kg bid. In addition, 7 out of the 9 mice subjected to the Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy had partial regressions and 10 out of 10 mice subjected to the Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy exhibited regression with 7 being partial and 3 being complete. By contrast, no regressions (partial or complete) were observed with any of the monotherapy groups.

[0091] In addition to the above, applicants found that the combination of Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan hydrochloride at 40 mg/kg q4dx5 produced tumor growth inhibition (TGI) and increased life span (ILS) results that were significantly better than correlative monotherapy results at $p < 0.05$ and also better than the results achieved with Compound I at 25 mg/kg bid and irinotecan hydrochloride at 40 mg/kg q4dx5 combination therapy and Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy. In the study, the Compound I at 25 mg/kg bid and irinotecan hydrochloride at 40 mg/kg q4dx5 combination therapy resulted in 4 out of 10 partial regressions and no complete regressions and the Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy resulted in 5 out of 10 partial regressions and no complete regressions. The cetuximab at

40 mg/kg 2x/wk and irinotecan at 40 mg/kg q4dx5 combination therapy and the correlative Compound I, cetuximab, and irinotecan hydrochloride monotherapies resulted in no regressions. By contrast, the Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan hydrochloride at 40 mg/kg q4dx5 therapy produced 10 out of 10 regressions with 9 being partial and one being complete.

[0092] These studies indicate that treating patients with a combination of Compound I and cetuximab is superior to treatment with either agent alone. Further, they indicate that combining the two agents allows for at least reduction in the dose of Compound I to obtain equivalent or better results. In addition, the studies indicate that treating patients with a combination of Compound I, cetuximab, and irinotecan hydrochloride produces even more superior results.

Examples

[0093] Abbreviations used herein are as follows:

q.s.	as much as needed
x	times
po	orally
ip	intraperitoneally
bid	twice daily
wk	week
qd	once daily
q4d x5	once every four days for a total of five doses
BWL	body weight loss

[0094] In the examples below, weight loss was graphically represented as percent change in mean group body weight, using the formula: $((W - W_0)/W_0) \times 100$, where 'W' represents mean body weight of the treated group at a particular day, and 'W₀' represents mean body weight of the same treated group at initiation of treatment. Maximum weight loss was also represented using the above formula, and indicated the maximum percent body weight loss that was observed at any time during the entire experiment for a particular group.

[0095] Efficacy data was graphically represented as the mean tumor volume $\pm$ standard error of the mean (SEM). In addition, tumor volumes of treated groups were presented as percentages of tumor volumes of the control groups (%T/C), using the formula: $100 \times ((T - T_0)/(C - C_0))$, where T represented mean tumor volume of a treated group on a specific day during the experiment, T₀ represented mean tumor volume of the same treated group on the first day of treatment; C represented mean tumor volume of a control group on the specific day during the experiment, and C₀ represented mean tumor volume of the same treated group on the first day of treatment.

[0096] Tumor volume (in cubic millimeters) was calculated using the ellipsoid formula: $(D \times (d^2))/2$, where "D" represents the large diameter of the tumor and "d" represents the small diameter.

[0097] Also, tumor regression and/or percent change in tumor volume was calculated using the formula: $((T - T_0)/T_0) \times 100$, where 'T' represents mean tumor volume of the treated group at a particular day, and 'T₀' represents mean tumor volume of the same treated group at initiation of treatment.

[0098] Statistical analysis was determined by the rank sum test and One Way Anova and a post-hoc Bonferroni t-test (SigmaStat, version 2.0, Jandel Scientific, San Francisco, CA, USA). Differences between groups were considered to be significant when the probability value (p) was ≤ 0.05 .

[0099] For survival assessment, the percent of increased life space (ILS) was calculated as: $100 \times [(\text{median survival day of treated group} - \text{median survival day of control group})/\text{median survival day of control group}]$. Median survival was determined utilizing Kaplan Meier survival analysis. Survival in treated groups was statistically compared with the vehicle group and survival comparisons were done between groups using the log-rank test (Graph Pad Prism, La Jolla, CA, USA). Differences between groups were considered significant when the probability value (p) was ≤ 0.05 .

Example 1

[0100] This example describes the formation of a suspension comprising Compound I.

[0101] A solid molecular complex comprising Compound I and hydroxypropyl methyl cellulose acetate succinate (HP-MC-AS) was first formed.

[0102] Compound I and HPMC-AS in a ratio of approximately 3:7, respectively, were dissolved in dimethylacetamide (DMA). The resulting solution was then added with stirring to very cold dilute hydrochloric acid resulting in the co-precipitation of Compound I and HPMC-AS as a solid molecular complex wherein Compound I was present in a nano-

particulate size range. The ratio of DMA to acid was in the range of 1:5 to 1:10.

[0103] The co-precipitate was then washed with water to remove DMA, filtered, dried to < 2% moisture content and passed through a # 30 mesh screen prior to evaluation. The resulting solid molecular complex was 30% by weight Compound I and 70% by weight HPMC.

[0104] The complex was then blended with colloidal silicon dioxide (available as Aerosil® 200 from Evonik Industries AG, Essen, Germany) such that, per 100g of the blend, 97g was the complex and 3g was colloidal silicon dioxide.

[0105] An aqueous vehicle containing 2% hydroxypropylcellulose (available as Klucel® LF from Aqualon, Wilmington, Delaware, USA) and 1N HCl at Qs to pH4 for the purpose of pH adjustment was then prepared.

[0106] 23.2 ml of the vehicle was equilibrated to room temperature and slowly transferred into 773.2 mg of the aforementioned blend. The resulting preparation was then slowly mixed until a homogenous suspension was obtained. The resulting suspension contained 9.375 mg/ml of Compound I. The suspension was stored at 2-8°C and protected from light.

Example 2

[0107] Mice were implanted with human HT-29 cell xenografts. The mice, cell line used, and implantation are described below.

[0108] Female athymic Crl:NU-Foxnlnu mice were used for efficacy testing (Charles River, Wilmington, MA, USA). Mice were 10-12 weeks of age and weighed 23-25 grams. The health of the mice was assessed daily by observation and analysis of blood samples taken from sentinel animals on shared shelf racks. All animals were allowed to acclimate and recover from shipping-related stress for one week. Autoclaved water and irradiated food (5058-ms Pico Lab mouse chow, Purina Mills, Richmond, IN, USA) were provided *ad libitum*, and the animals were kept in a 12 hour light and dark cycle. Cages, bedding and water bottles were autoclaved before use and changed weekly. All animal experiments were conducted in accordance with the Guide for the Care and Use of Laboratory Animals, local regulations, and protocols approved by the Roche Animal Care and Use Committee in our AAALAC accredited facility.

[0109] HT-29 cells (American Type Culture Collection, Rockville, MD) were grown in McCoy-5 medium supplemented with 10% Fetal Bovine Serum (FBS) and 1% of 200 nM L-glutamine, scaled up, harvested, and prepared so that each mouse received 3×10^6 cells in 0.2 ml calcium and magnesium free phosphate-buffered saline (PBS). Cells were implanted subcutaneously in the right flank of each of the mice.

Example 3

[0110] This example describes the preparation of a suspension of erlotinib hydrochloride.

[0111] One gram of Tween 80 was added to approximately 950 ml of water. While stirring at high speed, three grams of sodium carboxymethyl cellulose were added to the solution. Mixing was continued until the sodium carboxymethyl cellulose was dissolved. Water was then added q.s. until 1 liter. 12.5 grams of erlotinib hydrochloride (available from Genentech as Tarceva®) was then suspended in the solution and passed through a dissolver. The solution was then deaerated with nitrogen.

[0112] The contents of the final suspension are as follows:

Component	Amount
Erlotinib hydrochloride	12.5 g
Sodium carboxymethyl cellulose	3 g
Tween 80	1 g
Water for injection	q.s. to 1 liter

[0113] This provided a solution that was 12.5 mg/ml erlotinib hydrochloride. The solution was stored at 2 to 8°C.

Example 4

[0114] A suspension comprising Compound I was produced as described in Example 1.

[0115] The 12.5 mg/ml erlotinib hydrochloride solution was made as described in Example 3. A further 8.30 mg/ml solution was made in a manner similar to that in example 3 with the exception that 8.30 g of erlotinib hydrochloride was used instead of 12.5 g.

[0116] HT-29 xenograft-containing mice as produced in the manner described in Example 2 were randomized into groups of 10 mice each according to tumor volume so that all groups had similar starting mean tumor volumes. The approximate starting mean tumor volume for this study was 136 mm³.

[0117] Treatment of the mice began on day 12 post-cell implant and ended at day 29 post cell implant. Each group was subjected to a different therapy as follows:

- (1) mice receiving Compound I vehicle bid po and erlotinib hydrochloride vehicle qd po;
- (2) mice receiving Compound I at 75 mg/kg bid po;
- (3) mice receiving erlotinib hydrochloride at 67 mg/kg qd po;
- (4) mice receiving erlotinib hydrochloride at 100 mg/kg qd po;
- (5) mice receiving Compound I at 75 mg/kg bid po and erlotinib hydrochloride at 67 mg/kg qd po.

[0118] The Compound I suspension and its corresponding vehicle were dosed using a sterile 1cc syringe and 18-gauge gavage needle (0.2 ml/animal) twice daily. The erlotinib hydrochloride solution and its corresponding vehicle were dosed using a sterile 1cc syringe and 18-gauge needle (0.2 ml/animal) once daily starting on day 12 and ending on day 29 post-implantation. The 12.5 mg/ml solution was used for the erlotinib hydrochloride at 100 mg/kg groups and the 8.30 mg/ml solution was used for the erlotinib hydrochloride at 67 mg/kg qd groups. All dosing was based on an average mouse weight of 25 grams.

[0119] Tumor measurements were taken once or twice per week. All animals were individually followed throughout the experiment.

Toxicity

[0120] In general, no major signs of toxicity were noted in any dose group in this study described as assessed by measuring changes in body weight and gross observation of individual animals. Erlotinib hydrochloride at 100 mg/kg qd is historically not well tolerated in combination (Higgins et al., Anticancer Drugs, 15:503-12 (2004)), hence use of 67 mg/kg qd for the combination arm to ensure tolerability. Erlotinib hydrochloride at 100 mg/kg qd was included as a monotherapy arm for comparison. Compound I however is very well tolerated and was dosed at 75 mg/kg bid even in combination with erlotinib hydrochloride. EGFR inhibitor related skin rash was common in mice treated with erlotinib hydrochloride with a self limiting nature even under continuous treatment. See Table 1 and Figure 1.

Table 1

Group	Frequency	Route	% Change in Body Weight at end of Study Day 29	Max % Weight Loss	Max % Weight Gain	# animals $\geq$ 20% BWL	Mortality
Combo Vehicle	bid, qd	po	3.5	1.7	4.4	0	0
Compound I 75 mg/kg	bid	po	2.4	-0.1	3.6	0	0
Erlotinib HCl 67 mg/kg	qd	po	3.1	0.1	3.1	0	0
Erlotinib HCl 100 mg/kg	qd	po	2.9	-1.6	3.0	0	0
Compound I 75 mg/kg + Erlotinib HCl 67 mg/kg	bid, qd	po	2.5	-1.0	2.5	0	0

Tumor Growth Inhibition (TGI)

[0121] The group receiving Compound I monotherapy at 75 mg/kg bid exhibited 91% TGI whereas the group receiving erlotinib hydrochloride at 100 mg/kg qd exhibited 51 % TGI and the group receiving erlotinib hydrochloride at 67 mg/kg qd exhibited 38% TGI. No tumor regression was observed with any of the aforementioned groups. The group receiving combination therapy of Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd exhibited greater than 100% TGI with 9 out of 10 partial regressions (PRs). See Tables 2 and 3 and Figure 2.

Table 2

Group	Frequency	Route	Mean Tumor Volume (mm ³) Start Study	SEM	SD	Mean Tumor Volume (mm ³) End Study	SD	SEM
Combo Vehicle	bid, qd	po	137.99	±1.81	±5.74	1580.20	±74.00	±23.40
Compound I 75 mg/kg	bid	po	134.27	±2.11	±6.67	270.24	±68.06	±21.52
Erlotinib HCl 67 mg/kg	qd	po	136.46	±2.69	±8.49	1025.10	±142.96	±45.21

Group	Frequency	Route	Mean Tumor Volume (mm ³) Start Study	SEM	SD	Mean Tumor Volume (mm ³) End Study	SD	SEM
Erlotinib HCl 100 mg/kg	qd	po	133.82	±3.09	±9.76	838.75	±215.21	±68.06
Compound I 75 mg/kg + Erlotinib HCl 67 mg/kg	bid, qd	po	137.16	±2.08	±6.59	102.14	±21.26	±6.72

Table 3

Group	% T/C end of study Day: 29	% Inhibition end of study Day: 29	P value End of study Day: 29	Average % Regression per Group	Partial Regression	Complete Regression	Animals per Group	% Tumor Growth Inhibition
Combo Vehicle	---	---	---	---	0	0	10	---
Compound I 75 mg/kg bid	9	91	<0.001	---	0	0	10	91
Erlotinib HCl 67 mg/kg qd	62	38	<0.001	---	0	0	10	38
Erlotinib HCl 100 mg/kg qd	49	51	<0.001	---	0	0	10	51
Compound I 75 mg/kg bid + Erlotinib HCl 67 mg/kg qd	-2	regression	<0.001	26	9	0	10	>100

Assessment of Survival

[0122] The group receiving Compound I monotherapy at 75 mg/kg exhibited 100% increased life span (ILS). The group receiving erlotinib hydrochloride monotherapy at 100 mg/kg qd exhibited 38% ILS. The group receiving erlotinib hydrochloride monotherapy at 67 mg/kg qd exhibited 35% ILS. The group receiving combination therapy of Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd exhibited 142% ILS. See Table 4 and Figure 3.

Table 4

Group	ILS Calculations			
	50% Treatment Days	50% Vehicle Days	% ILS	p value
Combo Vehicle	---	---	---	---
Compound I 75 mg/kg bid	52	26	100	< 0.0001
Erlotinib HCl 67 mg/kg qd	35	26	35	< 0.0001
Erlotinib HCl 100 mg/kg qd	36	26	38	< 0.0001
Compound I 75 mg/kg bid + Erlotinib HCl 67 mg/kg qd	63	26	142	< 0.0001

Statistical Analysis

[0123] The %TGI in the Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd combination therapy was statistically superior to that of all monotherapy arms ($p < 0.05$). The %ILS in the Compound I at 75 mg/kg bid and erlotinib hydrochloride at 67 mg/kg qd combination therapy group was also statistically superior to that of all monotherapy arms tested ($p < 0.05$ for all comparisons). See Table 5.

Table 5

Treatment	versus	Treatment	TGI p value*	ILS p value**
Compound I 75 mg/kg bid		Erlotinib HCl 67 mg/kg qd	<0.05	<0.0001

(continued)

	Treatment	versus	Treatment	TGI p value*	ILS p value**
5	Compound I 75 mg/kg bid		Erlotinib HCl 100 mg/kg qd	<0.05	<0.0001
	Compound I 75 mg/kg bid		Erlotinib HCl 67 mg/kg qd + Compound I 75 mg/kg bid	<0.05	0.0003
	Erlotinib HCl 67 mg/kg qd		Erlotinib HCl 100 mg/kg qd	<0.05	0.2041
10	Erlotinib HCl 67 mg/kg qd		Erlotinib HCl 67 mg/kg qd + Compound I 75 mg/kg bid	<0.05	<0.0001
	Erlotinib HCl 100 mg/kg qd		Erlotinib HCl 67 mg/kg qd + Compound I 75 mg/kg bid	<0.05	<0.0001
15	*One-Way ANOVA, <i>post-hoc Bonferroni</i> ** Breslow-Gehan-Wilcoxon				

Example 5

[0124] Two suspensions comprising Compound I were made in a manner similar to that in example 1 with the exception that 20 ml of a 9.375 mg/ml suspension was made using 19.4 ml of the vehicle and 644 mg of the blend and 20 ml of a 3.125 mg/ml suspension was made using 19.8 ml of the vehicle and 214.8 mg of the blend.

[0125] Cetuximab was purchased from ImClone Systems, Inc. (available as Erbitux®) as a 2 mg/ml solution.

[0126] HT-29 xenograft-containing mice as produced in the manner described in Example 2 were randomized into groups of 10 mice each according to tumor volume so that all groups had similar starting mean tumor volumes. The approximate starting mean tumor volume for this study was 135 mm³.

[0127] Treatment began on day 12 post-cell implant and ended at day 34 post cell implant. Each group was subjected to a different therapy as follows:

- (1) mice receiving Compound I vehicle bid po and cetuximab vehicle 2x/wk ip;
- (2) mice receiving cetuximab at 40 mg/kg 2x/wk ip;
- (3) mice receiving Compound I at 25 mg/kg bid po;
- (4) mice receiving Compound I at 75 mg/kg bid po;
- (5) mice receiving Compound I at 25 mg/kg bid po and cetuximab at 40 mg/kg 2x/wk ip;
- (6) mice receiving Compound I at 75 mg/kg bid po and cetuximab at 40 mg/kg 2x/wk ip.

[0128] The Compound I suspension and its corresponding vehicle were dosed using a sterile 1cc syringe and 18-gauge gavage needle (0.2 ml/animal) twice daily. The 9.375 mg/ml suspension was used for the Compound I at 75 mg/kg bid groups and the 3.125 mg/ml suspension was used for the Compound I at 25 mg/kg bid groups. Cetuximab and its corresponding vehicle were dosed intraperitoneally using a sterile 1cc syringe and 26-gauge needle (0.5 ml/animal) twice a week on a Monday/Thursday or Tuesday/Friday schedule. All dosing was based on an average mouse weight of 25 grams.

[0129] Tumor measurements were taken once or twice per week. All animals were individually followed throughout the experiment.

Toxicity

[0130] In general, no major signs of toxicity were noted in any dose group in this study described as assessed by measuring changes in body weight and gross observation of individual animals. See Table 6 and Figure 4. EGFR inhibitor related skin rash was common in cetuximab treated mice with a self-limiting nature even under continuous treatment. One mouse appeared to have a bacterial infection as a sequela to rash leading to progressive weight loss > 20% thereby requiring humane sacrifice. This mouse was censored from the overall tumor growth inhibition and survival analysis.

Table 6

Group	Frequency	Route	% Change in Body Weight at end of Study Day 34	Max % Weight Loss	Max % Weight Gain	# animals $\geq$ 20% BWL	Mortality
Combo Vehicle	bid, 2x/wk	po, ip	6.6	0.5	6.6	0	0
Cetuximab 40 mg/kg	2x/wk	ip	7.2	3.7	8.9	0	0
Compound I 25 mg/kg	bid	po	0.1	-2.2	0.3	0	0
Compound I 75 mg/kg	bid	po	-0.1	-1.4	0.7	0	0
Compound I 25 mg/kg + Cetuximab 40 mg/kg	bid, 2x/wk	po, ip	0.5	-2.1	2.9	1	0
Compound I 25 mg/kg + Cetuximab 40 mg/kg	bid, 2x/wk	po, ip	-0.8	-2.4	1.6	0	0

Tumor Growth Inhibition (TGI)

[0131] The group receiving Compound I monotherapy at 25 mg/kg bid exhibited 74 %TGI and the group receiving Compound I monotherapy at 75 mg/kg bid exhibited 93 %TGI. The group receiving cetuximab at 40 mg/kg 2x/wk achieved 51 %TGI. No tumor regression was observed with any of the aforementioned groups. Both combination therapy groups, however, exhibited >100 %TGI. The group receiving Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 7 out of 10 partial regressions (PRs) but no complete regressions (CRs). The group receiving Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 7 out of 10 PRs and 3 out of 10 CRs.

[0132] See Tables 7 and 8 and Figure 5.

Table 7

Group	Frequency	Route	Mean Tumor Volume (mm ³) Start Study	SEM	SD	Mean Tumor Volume (mm ³) End Study	SD	SEM
Combo Vehicle	bid, 2x/wk	po, ip	133.95	±2.31	±7.31	2011.75	±227.22	±71.85
Cetuximab 40 mg/kg	2x/wk	ip	133.13	±2.34	±7.39	1052.49	±249.18	±78.80
Compound I 25 mg/kg	bid	po	135.97	±2.75	±8.71	615.99	±148.01	±46.81
Compound I 75 mg/kg	bid	po	137.05	±2.35	±7.44	275.66	±56.34	±17.82
Compound I 25 mg/kg + Cetuximab 40 mg/kg	bid, 2x/wk	po, ip	134.56	±2.24	±7.07	90.98	±41.56	±13.85
Compound I 75 mg/kg + Cetuximab 40 mg/kg	bid, 2x/wk	po, ip	137.04	±3.20	±10.10	26.45	±20.60	±6.52

Table 8

Group	% T/C end of study Day: 34	% Inhibition end of study Day: 34	p value End of study Day: 34	Average % Regression per Group	Partial Regression	Complete Regression	Animals per Group	% Tumor Growth Inhibition
Combo Vehicle	---	---	---	---	0	0	10	---
Cetuximab 40 mg/kg 2x/wk	49	51	<0.001	---	0	0	10	51
Compound I 25 mg/kg bid	26	74	<0.001	---	0	0	10	74
Compound I 75 mg/kg bid	7	93	<0.001	---	0	0	10	93
Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	-2	regression	<0.001	39	7	0	9	>100
Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	-6	regression	<0.001	81	7	3	10	>100

Assessment of Survival

[0133] The group receiving Compound I monotherapy at 25 mg/kg bid exhibited 44 %ILS and the group receiving Compound I monotherapy at 75 mg/kg bid exhibited 75 %ILS. The group receiving cetuximab at 40 mg/kg 2x/wk achieved 16 %ILS. The group receiving Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 97 %ILS. The group receiving Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 122 %ILS. See Table 9 and Figure 6.

Table 9

Group	ILS Calculations		% ILS	p value
	50% Treatment Days	50% Vehicle Days		
Combo Vehicle	---	---	---	---
Cetuximab 40 mg/kg 2x/wk	37	32	16	< 0.0001
Compound I 25 mg/kg bid	46	32	44	< 0.0001
Compound I 75 mg/kg bid	56	32	75	< 0.0001
Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	63	32	97	< 0.0001
Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	71	32	122	< 0.0001

Statistical Analysis

[0134] The %TGI of both the Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy and the Compound I at 75 mg/kg bid and cetuximab at 40 mg/kg 2x/wk combination therapy were statistically superior to that of all monotherapy arms ($p < 0.05$). The %ILSs achieved in both monotherapies were also statistically superior to

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that of all monotherapy arms tested ($p < 0.05$ for all comparisons). See Table 10.

Table 10

	Treatment	versus	Treatment	TGI p value*	ILS p value **
5	Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid	<0.05	0.0010
	Cetuximab 40 mg/kg 2x/wk		Compound I 75 mg/kg bid	<0.05	<0.0001
	Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
10	Cetuximab 40 mg/kg 2x/wk		Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
	Compound I 25 mg/kg bid		Compound I 75 mg/kg bid	<0.05	0.0025
	Compound I 25 mg/kg bid		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
15	Compound I 25 mg/kg bid		Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
	Compound 175 mg/kg bid		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	0.0089
20	Compound I 75 mg/kg bid		Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	0.0002
	Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk		Compound I 75 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	>0.05	0.3210
25	*One-Way ANOVA, post-hoc Bonferroni ** Breslow-Gehan-Wilcoxon				

Example 6

[0135] A suspension comprising Compound I was made in a manner similar to that in Example 1 with the exception that 40 ml of a 3.125 mg/ml suspension was made using 39.6 ml of the vehicle and 429.6 mg of the blend.

[0136] Cetuximab was purchased from ImClone Systems, Inc. (available as Erbitux®) as a 2 mg/ml solution. Irinotecan HCl hydrochloride was purchased from Pfizer Inc. (available as Camptosar®) as a stock sterile solution of 20 mg/ml, which was diluted as required with sterile saline to 2 mg/ml.

[0137] HT-29 xenograft-containing mice as produced in the manner described in Example 2 were randomized into groups of 10 mice each according to tumor volume so that groups has similar starting mean tumor volumes. The approximate starting mean tumor volume for this study was 135 mm³.

[0138] Treatment began on day 11 post-cell implant and ended at day 32 post cell implant. Each group was subjected to a different therapy as follows:

- (1) mice receiving Compound I vehicle bid po, cetuximab vehicle 2x/wk ip, and irinotecan HCl vehicle q4d x5 ip;
- (2) mice receiving irinotecan HCl at 40 mg/kg q4d x5 ip;
- (3) mice receiving Compound I at 25 mg/kg bid po;
- (4) mice receiving cetuximab at 40 mg/kg 2x/wk ip;
- (5) mice receiving Compound I at 25 mg/kg bid po and irinotecan HCl at 40 mg/kg q4d x5 ip;
- (6) mice receiving cetuximab at 40 mg/kg 2x/wk ip and irinotecan HCl at 40 mg/kg q4d x5 ip;
- (7) mice receiving Compound I at 25 mg/kg bid po and cetuximab at 40 mg/kg 2x/wk ip;
- (8) mice receiving Compound I at 25 mg/kg bid po, cetuximab at 40 mg/kg 2x/wk ip, and irinotecan HCl at 40 mg/kg q4d x5 ip.

[0139] The Compound I suspension and its corresponding vehicle were dosed using a sterile 1cc syringe and 18-gauge gavage needle (0.2 ml/animal) twice daily. Cetuximab and its corresponding vehicle were dosed intraperitoneally using a sterile 1cc syringe and 26-gauge needle (0.2 ml/animal) twice a week on a Monday/Thursday or Tuesday/Friday schedule. Irinotecan HCl and its corresponding vehicle were dosed intraperitoneally using a sterile 1cc syringe and 26-gauge needle (0.2 ml/animal) on a q4d x5 schedule. All dosing was based on an average mouse weight of 25 grams.

[0140] Tumor measurements were taken once or twice per week. All animals were individually followed throughout the experiment.

Toxicity

[0141] In general, no major signs of toxicity were noted in any dose group in this study described as assessed by measuring changes in body weight and gross observation of individual animals. See Table 11 and Figure 7. EGFR inhibitor related skin rash was common in cetuximab treated mice with a self-limiting nature even under continuous treatment.

Table 11

Group	Frequency	Route	% Change in Body Weight at end of Study Day 32	Max % Weight Loss	Max % Weight Gain	# animals $\geq$ 20% BWL	Mortality
Combo Vehicle	bid, 2x/wk, q4d x5	po, ip, ip	3.2	1.3	4.4	0	0
Irinotecan HCl 40 mg/kg	q4d x5	ip	2.7	0.1	2.8	0	0
Compound I 25 mg/kg	bid	po	2.9	-0.6	3.8	0	0
Cetuximab 40 mg/kg	2x/wk	ip	4.0	0.1	4.0	0	0
Compound I 25 mg/kg + irinotecan HCl 40 mg/kg	bid, q4d x5	po, ip	1.2	-0.1	2.3	0	0
Cetuximab 40 mg/kg + irinotecan HCl 40 mg/kg	2x/wk, q4d x5	ip, ip	2.8	0.2	3.2	0	0
Compound I 25 mg/kg + cetuximab 40 mg/kg	bid, 2x/wk	po, ip	0.8	0.3	2.6	0	0
Compound I 25 mg/kg + cetuximab 40 mg/kg + irinotecan HCl 40 mg/kg	bid, 2x/wk, q4d x5	po, ip, ip	0.8	-0.7	2.4	0	0

Tumor Growth Inhibition (TGI)

[0142] The group receiving Compound I monotherapy at 25 mg/kg bid exhibited 76 %TGI. The group receiving cetuximab monotherapy at 40 mg/kg 2x/wk exhibited 58 %TGI. The group receiving irinotecan HCl monotherapy at 40 mg/kg q4dx5 exhibited 59 %TGI. The group receiving Compound I at 25 mg/kg bid and irinotecan HCl at 40 mg/kg q4dx5 exhibited 98 %TGI. The group receiving cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 exhibited 92 %TGI. The group receiving Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited >100 %TGI. The group receiving Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 exhibited >100 %TGI. No tumor regression was observed with any of the monotherapy groups. The group receiving Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 5 out of 10 partial regressions (PRs) but no complete regressions (CRs). The group receiving Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk, and irinotecan HCl at 40 mg/kg q4dx5 exhibited 9 out of 10 PRs and 1 out of 10 CRs.

[0143] See Tables 12 and 13 and Figure 8.

Table 12

Group	Frequency	Route	Mean Tumor Volume (mm ³) Start Study	SEM	SD	Mean Tumor Volume (mm ³) End Study	SD	SEM
Combo Vehicle	bid, 2x/wk, q4d x5	po, ip, ip	133.61	±5.44	±17.20	1920.46	±395.43	±125.05
Irinotecan HCl 40 mg/kg	q4d x5	ip	127.56	±4.44	±14.03	862.41	±321.20	±101.57
Compound I 25 mg/kg	bid	po	136.24	±6.05	±19.13	563.72	±140.24	±44.35
Cetuximab 40 mg/kg	2x/wk	ip	132.09	±5.80	±18.33	885.00	±406.03	±128.40
Compound I 25 mg/kg + irinotecan HCl 40 mg/kg	bid, q4d x5	po, ip	144.93	±5.35	±16.93	182.76	±69.45	±21.96
Cetuximab 40 mg/kg + irinotecan HCl 40 mg/kg	2x/wk, q4d x5	ip, ip	148.52	±6.75	±21.34	295.26	±113.09	±35.76
Compound I 25 mg/kg + cetuximab 40 mg/kg	bid, 2x/wk	po, ip	132.52	±6.39	±20.22	122.05	±35.99	±11.38
Compound I 25 mg/kg + cetuximab 40 mg/kg + irinotecan HCl 40 mg/kg	bid, 2x/wk, q4d x5	po, ip, ip	134.61	±6.88	±21.74	40.67	±23.89	±7.55

Table 13

Group	% T/C end of study Day: 32	% Inhibition end of study Day: 32	p value End of study Day: 32	Average % Regression per Group	Partial Regression	Complete Regression	Animals per Group	% Tumor Growth Inhibition
Combo Vehicle	---	---	---	---	0	0	10	---
Irinotecan HCl 40 mg/kg q4d x5	41	59	<0.001	---	0	0	10	59
Compound I 25 mg/kg bid	24	76	<0.001	---	0	0	10	76
Cetuximab 40 mg/kg 2x/wk	42	58	<0.001	---	0	0	10	58
Compound I 25 mg/kg bid + irinotecan HCl 40 mg/kg q4d x5	2	98	<0.001	---	4	0	10	98
Cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x5	8	92	<0.001	---	0	0	10	92
Compound I 25 mg/kg bid + cetuximab 40 mg/kg 2x/wk	-1	regression	<0.001	8	5	0	10	>100
Compound I 25 mg/kg bid + cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x5	-5	regression	<0.001	70	9	1	10	>100

Assessment of Survival

[0144] The group receiving Compound I monotherapy at 25 mg/kg bid exhibited 80 % ILS. The group receiving cetuximab monotherapy at 40 mg/kg 2x/wk exhibited 27 % ILS. The group receiving irinotecan HCl monotherapy at 40 mg/kg q4dx5 exhibited 17 % ILS. The group receiving Compound I at 25 mg/kg bid and irinotecan HCl at 40 mg/kg q4dx5 exhibited 163 % ILS. The group receiving cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 exhibited 80 % ILS. The group receiving Compound I at 25 mg/kg bid and cetuximab at 40 mg/kg 2x/wk exhibited 127 % ILS. The group receiving Compound I at 25 mg/kg bid, cetuximab at 40 mg/kg 2x/wk and irinotecan HCl at 40 mg/kg q4dx5 exhibited 259 % ILS. See Table 14 and Figure 9.

Table 14

	Group	ILS Calculations			
		50% Treatment Days	50% Vehicle Days	% ILS	p value
5	Combo Vehicle	---	---	---	---
	Irinotecan HCl 40 mg/kg q4d x5	35	30	17	< 0.0001
	Compound I 25 mg/kg bid	54	30	80	< 0.0001
10	Cetuximab 40 mg/kg 2x/wk	38	30	27	< 0.0001
	Compound I 25 mg/kg bid + irinotecan HCl 40 mg/kg q4d x5	79	30	163	< 0.0001
	Cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x5	54	30	80	< 0.0001
	Compound I 25 mg/kg bid + cetuximab 40 mg/kg 2x/wk	68	30	127	< 0.0001
15	Compound I 25 mg/kg bid + cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x5	105	30	250	< 0.0001

Statistical Analysis

20 **[0145]** The %TGI of the Compound I/cetuximab, the Compound I/irinotecan HCl, and the Compound I/cetuximab/irinotecan HCl combination therapies were statistically superior to that of all monotherapy arms ($p < 0.05$). The %TGI of the Compound I/cetuximab/irinotecan HCl combination therapy was also statistically superior to that of the Compound I/irinotecan HCl and cetuximab/irinotecan HCl combination therapies ($p < 0.05$).

25 **[0146]** The %ILS of the Compound I/cetuximab, the Compound I/irinotecan HCl, and the Compound I/cetuximab/irinotecan HCl combination therapies were statistically superior to that of all monotherapy arms ($p < 0.05$ for all comparisons). The %ILS of the Compound I/cetuximab/irinotecan HCl combination therapy was also statistically superior to that of the Compound I/irinotecan HCl and Compound I/cetuximab combination therapies. See Table 15.

Table 15

	Treatment	versus	Treatment	TGI p value*	ILS p value **
30	Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid	>0.05	<0.0001
	Irinotecan HCl 40 mg/kg q4d x5		Cetuximab 40 mg/kg 2x/wk	>0.05	0.5370
35	Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
	Irinotecan HCl 40 mg/kg q4d x5		Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
	Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
40	Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
	Compound I 25 mg/kg bid		Cetuximab 40 mg/kg 2x/wk	>0.05	<0.0001
45	Compound I 25 mg/kg bid		Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5	<0.05	0.0004
	Compound I 25 mg/kg bid		Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk	>0.05	0.3457
	Compound I 25 mg/kg bid		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	0.0004
50	Compound I 25 mg/kg bid		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
	Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
55	Cetuximab 40 mg/kg 2x/wk		Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001

(continued)

	Treatment	versus	Treatment	TGI p value*	ILS p value **
5	Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	<0.0001
	Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
10	Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5		Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk	<0.05	0.0006
	Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	>0.05	0.0030
15	Compound I 25 mg/kg bid + Irinotecan HCl 40 mg/kg q4d x5		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	<0.05	0.0420
	Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	<0.05	0.0862
20	Irinotecan HCl 40 mg/kg q4d x5 + Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	<0.05	<0.0001
	Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk		Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + Irinotecan HCl 40 mg/kg q4d x5	>0.05	<0.0001
25	*One-Way ANOVA, post-hoc Bonferroni ** Breslow-Gehan-Wilcoxon				

Claims

1. A pharmaceutical product comprising (A) a first component which comprises, as an active agent, propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide (Compound I), or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor; as a combined preparation for the simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation, wherein the EGFR inhibitor is selected from erlotinib or a pharmaceutically acceptable salt thereof, or cetuximab.
2. The pharmaceutical product for use according to claim 1 for the treatment of colorectal cancer, melanoma and thyroid cancer, comprising b-Raf having the V600E mutation.
3. The pharmaceutical product for use according to any one of claims 1 and 2, further comprising a third component (C), which comprises as an active agent a topoisomerase inhibitor, wherein said topoisomerase inhibitor is irinotecan, or a pharmaceutically acceptable salt thereof.
4. The pharmaceutical product for use according to any one of claims 1 to 3 wherein propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluorophenyl}-amide, or a pharmaceutically-acceptable salt thereof, is in amorphous or substantially amorphous form.
5. A kit comprising: (A) a first component which comprises, as an active agent, propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluorophenyl}-amide, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, an EGFR inhibitor wherein the EGFR inhibitor is selected from erlotinib or a pharmaceutically acceptable salt thereof, or cetuximab.
6. A kit according to claim 5 further comprising a third component which comprises, as an active agent, irinotecan.
7. A kit according to claims 5 or 6, for use in the treatment of cancer comprising b-Raf having a V600 mutation.

8. The pharmaceutical product for use according to claim 1, wherein said propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide, or a pharmaceutically acceptable salt thereof, is administered twice daily in an amount of from about 850 mg to about 1050 mg; and said erlotinib, or a pharmaceutically acceptable salt thereof, is administered daily in an amount of from about 100 mg to about 200 mg.
9. The pharmaceutical product for use according to claim 1, wherein said propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide, or a pharmaceutically acceptable salt thereof, is administered twice daily in an amount of from about 850 mg to about 1050 mg; and said cetuximab is administered weekly with the first administration being in an amount of from about 400 mg/m² to about 500 mg/m² and each subsequent administration being in an amount of from about 200 mg/m² to about 300 mg/m².
10. The pharmaceutical product for use according to claims 8 and 9, further comprising irinotecan, or a pharmaceutically acceptable salt thereof.
11. The pharmaceutical product for use according to claim 10, wherein said irinotecan, or a pharmaceutically acceptable salt thereof, is administered weekly in an amount of from about 50 mg/m² to about 200 mg/m².
12. The pharmaceutical product for use according to claim 10, wherein irinotecan, or a pharmaceutically acceptable salt thereof, is administered in a six week cycle at about 75 to about 175 mg/m² weekly for the first four weeks.
13. The pharmaceutical product for use according to claim 10, wherein irinotecan, or a pharmaceutically acceptable salt thereof, is administered once every three weeks at about 300 to about 400 mg/m²; or once every two weeks at about 130 to about 230 mg/m².
14. A pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, erlotinib or a pharmaceutically acceptable salt thereof, as a combined preparation for simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation, wherein
 - (A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day, and
 - (B) is administered in an amount of from about 20 mg/day to about 500 mg/day, from about 100 mg/day to about 400 mg/day, or from about 100 mg/day to about 200 mg/day.
15. A pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I, or a pharmaceutically-acceptable salt thereof; and (B) a second component which comprises, as an active agent, cetuximab; as a combined preparation for simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation, wherein
 - (A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day; and
 - (B) is administered in an amount of from about 50 mg/m²/week to about 700 mg/m²/week, from about 100 mg/m²/week to about 600 mg/m²/week, or from about 200 mg/m²/week to about 500 mg/m²/week.
16. A pharmaceutical product comprising (A) a first component which comprises, as an active agent, Compound I or a pharmaceutically-acceptable salt thereof; (B) a second component which comprises, as an active agent, cetuximab; and (C) a third component which comprises, as an active agent, irinotecan or a pharmaceutically-acceptable salt thereof; as a combined preparation for simultaneous or sequential use in the treatment of cancer comprising b-Raf having a V600 mutation, wherein
 - (A) is administered in an amount of from about 200 mg/day to about 3000 mg/day, from about 1000 mg/day to about 2500 mg/day, from about 1700 mg/day to about 2100 mg/day or about 1920 mg/day;
 - (B) is administered in an amount of from about 50 mg/m²/week to about 700 mg/m²/week, from about 100 mg/m²/week to about 600 mg/m²/week, or from about 200 mg/m²/week to about 500 mg/m²/week; and
 - (C) is administered in an amount of from about 1 to about 250 mg/m²/week, about 50 to about 200 mg/m²/week, or about 125 mg/m²/week.

17. The use of propane-1-sulfonic acid {3-[5-(4-chloro-phenyl)-1H-pyrrolo [2,3-b] pyridine-3-carbonyl]-2,4-difluoro-phenyl}-amide, or a pharmaceutically-acceptable salt thereof, and an EGFR inhibitor for the manufacture of medicaments for the treatment of cancer comprising b-Raf having a V600 mutation, wherein the EGFR inhibitor is selected from erlotinib or a pharmaceutically acceptable salt thereof, or cetuximab.

5

Patentansprüche

1. Ein pharmazeutisches Produkt, umfassend (A) eine erste Komponente, die als einen Wirkstoff Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluor-phenyl}-amid (Verbindung I) oder ein pharmazeutisch verträgliches Salz davon umfasst; und (B) eine zweite Komponente, die als einen Wirkstoff einen EGFR-Inhibitor umfasst; als Kombinationszubereitung für die gleichzeitige oder sequenzielle Verwendung in der Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation, wobei der EGFR-Inhibitor ausgewählt ist aus Erlotinib oder einem pharmazeutisch verträgliches Salz davon, oder Cetuximab.
2. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 1 zur Behandlung von kolorektalem Krebs, Melanom und Schilddrüsenkrebs umfassend b-Raf mit der V600E Mutation.
3. Das pharmazeutische Produkt zur Verwendung gemäß einem der Ansprüche 1 und 2, ferner umfassend eine dritte Komponente (C), die als einen Wirkstoff einen Topoisomeraseinhibitor umfasst, wobei der Topoisomeraseinhibitor Irinotecan oder ein pharmazeutisch verträgliches Salz davon ist.
4. Das pharmazeutische Produkt zur Verwendung gemäß einem der Ansprüche 1 bis 3, wobei Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluor-phenyl}-amid oder ein pharmazeutisch verträgliches Salz davon in amorpher oder im wesentlichen amorpher Form vorliegt.
5. Ein Kit, umfassend: (A) eine erste Komponente, die als einen Wirkstoff Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluorphenyl}-amid oder ein pharmazeutisch verträgliches Salz davon umfasst; und (B) eine zweite Komponente, die als einen Wirkstoff einen EGFR-Inhibitor umfasst, wobei der EGFR-Inhibitor ausgewählt ist aus Erlotinib oder einem pharmazeutisch verträgliches Salz davon oder Cetuximab.
6. Ein Kit gemäß Anspruch 5, ferner umfassend eine dritte Komponente, die als einen Wirkstoff Irinotecan umfasst.
7. Ein Kit gemäß Anspruch 5 oder 6, zur Verwendung in der Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation.
8. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 1, wobei das Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluorphenyl}-amid oder ein pharmazeutisch verträgliches Salz davon zweimal täglich in einer Menge von etwa 850 mg bis 1050 mg verabreicht wird; und das Erlotinib oder ein pharmazeutisch verträgliches Salz davon täglich in einer Menge von etwa 100 mg bis etwa 200 mg verabreicht wird.
9. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 1, wobei das Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluor-phenyl}-amid oder ein pharmazeutisch verträgliches Salz davon zweimal täglich in einer Menge von etwa 850 mg bis 1050 mg verabreicht wird; und das Cetuximab wöchentlich verabreicht wird, wobei die erste Verabreichung in einer Menge von etwa 400 mg/m² bis etwa 500 mg/m² erfolgt und jede nachfolgende Verabreichung in einer Menge von etwa 200 mg/m² bis etwa 300 mg/m² erfolgt.
10. Das pharmazeutische Produkt zur Verwendung gemäß Ansprüchen 8 und 9, ferner umfassend Irinotecan oder ein pharmazeutisch verträgliches Salz davon.
11. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 10, wobei das Irinotecan oder ein pharmazeutisch verträgliches Salz davon wöchentlich in einer Menge von etwa 50 mg/m² bis etwa 200 mg/m² verabreicht wird.
12. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 10, wobei Irinotecan oder ein pharmazeutisch verträgliches Salz davon in einem 6-Wochen-Zyklus mit etwa 75 bis etwa 175 mg/m² wöchentlich für die ersten vier Wochen verabreicht wird.
13. Das pharmazeutische Produkt zur Verwendung gemäß Anspruch 10, wobei Irinotecan oder ein pharmazeutisch

verträgliches Salz davon einmal alle 3 Wochen mit etwa 300 bis etwa 400 mg/m² verabreicht wird, oder einmal alle 2 Wochen mit etwa 130 bis etwa 230 mg/m².

- 5 14. Ein pharmazeutisches Produkt, umfassend (A) eine erste Komponente, die als einen Wirkstoff Verbindung I oder ein pharmazeutisch verträgliches Salz davon umfasst; und (B) eine zweite Komponente, die als einen Wirkstoff Erlotinib oder ein pharmazeutisch verträgliches Salz davon umfasst; als Kombinationszubereitung für die gleichzeitige oder sequenzielle Verwendung in der Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation, wobei

10 (A) in einer Menge von etwa 200 mg/Tag bis etwa 3000 mg/Tag, von etwa 1000 mg/Tag bis etwa 2500 mg/Tag, von etwa 1700 mg/Tag bis etwa 2100 mg/Tag oder etwa 1920 mg/Tag verabreicht wird, und

(B) in einer Menge von etwa 20 mg/Tag bis etwa 500 mg/Tag, von etwa 100 mg/Tag bis etwa 400 mg/Tag, oder von etwa 100 mg/Tag bis etwa 200 mg/Tag verabreicht wird.

- 15 15. Ein pharmazeutisches Produkt, umfassend (A) eine erste Komponente, die als einen Wirkstoff Verbindung I oder ein pharmazeutisch verträgliches Salz davon umfasst; und (B) eine zweite Komponente, die als einen Wirkstoff Cetuximab umfasst; als Kombinationszubereitung für die gleichzeitige oder sequenzielle Verwendung in der Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation, wobei

20 (A) in einer Menge von etwa 200 mg/Tag bis etwa 3000 mg/Tag, von etwa 1000 mg/Tag bis etwa 2500 mg/Tag, von etwa 1700 mg/Tag bis etwa 2100 mg/Tag oder etwa 1920 mg/Tag verabreicht wird, und

(B) in einer Menge von etwa 50 mg/m²/Woche bis etwa 700 mg/m²/Woche, von etwa 100 mg/m²/Woche bis etwa 600 mg/m²/Woche, oder von etwa 200 mg/m²/Woche bis etwa 500 mg/m²/Woche verabreicht wird.

- 25 16. Ein pharmazeutisches Produkt, umfassend (A) eine erste Komponente, die als einen Wirkstoff Verbindung I oder ein pharmazeutisch verträgliches Salz davon umfasst; (B) eine zweite Komponente, die als einen Wirkstoff Cetuximab umfasst; und (C) eine dritte Komponente, die als einen Wirkstoff Irinotecan oder ein pharmazeutisch verträgliches Salz davon umfasst; als Kombinationszubereitung für die gleichzeitige oder sequenzielle Verwendung in der Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation, wobei

30 (A) in einer Menge von etwa 200 mg/Tag bis etwa 3000 mg/Tag, von etwa 1000 mg/Tag bis etwa 2500 mg/Tag, von etwa 1700 mg/Tag bis etwa 2100 mg/Tag oder etwa 1920 mg/Tag verabreicht wird,

(B) in einer Menge von etwa 50 mg/m²/Woche bis etwa 700 mg/m²/Woche, von etwa 100 mg/m²/Woche bis etwa 600 mg/m²/Woche, oder von etwa 200 mg/m²/Woche bis etwa 500 mg/m²/Woche verabreicht wird, und

35 (C) in einer Menge von etwa 1 bis etwa 250 mg/m²/Woche, etwa 50 bis etwa 200 mg/m²/Woche oder etwa 125 mg/m²/Woche verabreicht wird.

- 40 17. Die Verwendung von Propan-1-sulfonsäure-{3-[5-(4-chlor-phenyl)-1H-pyrrolo[2,3-b]pyridin-3-carbonyl]-2,4-difluor-phenyl}-amid oder eines pharmazeutisch verträglichen Salzes davon und eines EGFR-Inhibitors zur Herstellung eines Medikaments zur Behandlung von Krebs umfassend b-Raf mit einer V600 Mutation, wobei der EGFR-Inhibitor ausgewählt ist aus Erlotinib oder einem pharmazeutisch verträglichen Salz davon, oder Cetuximab.

Revendications

- 45 1. Produit pharmaceutique comprenant (A) un premier composant qui comprend, à titre d'agent actif, du {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propane-1-sulfonique (composé I), ou un sel pharmaceutiquement acceptable de celui-ci ; et (B) un deuxième composant qui comprend, à titre d'agent actif, un inhibiteur d'EGFR ; sous la forme d'une préparation combinée pour utilisation simultanée ou séquentielle dans le traitement d'un cancer comprenant b-Raf ayant une mutation V600, dans lequel l'inhibiteur d'EGFR est choisi parmi l'erlotinib, un sel pharmaceutiquement acceptable de celui-ci, et le cétuximab.

- 50 2. Produit pharmaceutique pour utilisation selon la revendication 1 pour le traitement d'un cancer colorectal, d'un mélanome et d'un cancer de la thyroïde, comprenant b-Raf ayant la mutation V600E.

- 55 3. Produit pharmaceutique pour utilisation selon l'une quelconque des revendications 1 et 2, comprenant en outre un troisième composant (C) qui comprend, à titre d'agent actif, un inhibiteur de topoisomérase, dans lequel ledit inhibiteur de topoisomérase est l'irinotécan ou un sel pharmaceutiquement acceptable de celui-ci.

4. Produit pharmaceutique pour utilisation selon l'une quelconque des revendications 1 à 3, dans lequel le {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propane-1-sulfonique ou un sel pharmaceutiquement acceptable de celui-ci est sous une forme amorphe ou pratiquement amorphe.
- 5 5. Trousse comprenant : (A) un premier composant qui comprend, à titre d'agent actif, du {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propane-1-sulfonique ou un sel pharmaceutiquement acceptable de celui-ci ; et (B) un deuxième composant qui comprend, à titre d'agent actif, un inhibiteur d'EGFR, dans laquelle l'inhibiteur d'EGFR est choisi parmi l'erlotinib, un sel pharmaceutiquement acceptable de celui-ci, et le cétuximab.
- 10 6. Trousse selon la revendication 5, comprenant en outre un troisième composant qui comprend, à titre d'agent actif, de l'irinotécan.
- 15 7. Trousse selon la revendication 5 ou 6, pour utilisation dans le traitement d'un cancer comprenant b-Raf ayant une mutation V600.
- 20 8. Produit pharmaceutique pour utilisation selon la revendication 1, dans lequel ledit {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propane-1-sulfonique ou un sel pharmaceutiquement acceptable de celui-ci est administré deux fois par jour en une quantité d'environ 850 mg à environ 1050 mg ; et ledit erlotinib ou un sel pharmaceutiquement acceptable de celui-ci est administré chaque jour en une quantité d'environ 100 mg à environ 200 mg.
- 25 9. Produit pharmaceutique pour utilisation selon la revendication 1, dans lequel ledit {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propane-1-sulfonique ou un sel pharmaceutiquement acceptable de celui-ci est administré deux fois par jour en une quantité d'environ 850 mg à environ 1050 mg ; et ledit cétuximab est administré chaque semaine, la première administration étant une quantité d'environ 400 mg/m² à environ 500 mg/m² et chaque administration subséquente étant une quantité d'environ 200 mg/m² à environ 300 mg/m².
- 30 10. Produit pharmaceutique pour utilisation selon les revendications 8 et 9, comprenant en outre de l'irinotécan, ou un sel pharmaceutiquement acceptable de celui-ci.
- 35 11. Produit pharmaceutique pour utilisation selon la revendication 10, dans lequel ledit irinotécan, ou un sel pharmaceutiquement acceptable de celui-ci, est administré chaque semaine en une quantité d'environ 50 mg/m² à environ 200 mg/m².
- 40 12. Produit pharmaceutique pour utilisation selon la revendication 10, dans lequel ledit irinotécan, ou un sel pharmaceutiquement acceptable de celui-ci, est administré selon un cycle de six semaines à raison d'environ 75 à environ 175 mg/m² chaque semaine pendant les quatre premières semaines.
- 45 13. Produit pharmaceutique pour utilisation selon la revendication 10, dans lequel ledit irinotécan, ou un sel pharmaceutiquement acceptable de celui-ci, est administré une fois toutes les trois semaines à raison d'environ 300 à environ 400 mg/m² ; ou une fois toutes les deux semaines à raison d'environ 130 à environ 230 mg/m².
- 50 14. Produit pharmaceutique comprenant (A) un premier composant qui comprend, à titre d'agent actif, le composé I ou un sel pharmaceutiquement acceptable de celui-ci ; et (B) un deuxième composant qui comprend, à titre d'agent actif, de l'erlotinib ou un sel pharmaceutiquement acceptable de celui-ci, sous la forme d'une préparation combinée pour utilisation simultanée ou séquentielle dans le traitement d'un cancer comprenant b-Raf ayant une mutation V600, dans lequel

(A) est administré en une quantité d'environ 200 mg/jour à environ 3000 mg/jour, d'environ 1000 mg/jour à environ 2500 mg/jour, d'environ 1700 mg/jour à environ 2100 mg/jour ou d'environ 1920 mg/jour, et

(B) est administré en une quantité d'environ 20 mg/jour à environ 500 mg/jour, d'environ 100 mg/jour à environ 400 mg/jour, ou d'environ 100 mg/jour à environ 200 mg/jour.
- 55 15. Produit pharmaceutique comprenant (A) un premier composant qui comprend, à titre d'agent actif, le composé I ou un sel pharmaceutiquement acceptable de celui-ci ; et (B) un deuxième composant qui comprend, à titre d'agent actif, du cétuximab ; sous la forme d'une préparation combinée pour utilisation simultanée ou séquentielle dans le

traitement d'un cancer comprenant b-Raf ayant une mutation V600, dans lequel

(A) est administré en une quantité d'environ 200 mg/jour à environ 3000 mg/jour, d'environ 1000 mg/jour à environ 2500 mg/jour, d'environ 1700 mg/jour à environ 2100 mg/jour ou d'environ 1920 mg/jour ; et

(B) est administré en une quantité d'environ 50 mg/m²/semaine à environ 700 mg/m²/semaine, d'environ 100 mg/m²/semaine à environ 600 mg/m²/semaine, ou d'environ 200 mg/m²/semaine à environ 500 mg/m²/semaine.

16. Produit pharmaceutique comprenant (A) un premier composant qui comprend, à titre d'agent actif, le composé I ou un sel pharmaceutiquement acceptable de celui-ci ; (B) un deuxième composant qui comprend, à titre d'agent actif, du cétuximab ; et (C) un troisième composant qui comprend, à titre d'agent actif, de l'irinotécan ou un sel pharmaceutiquement acceptable de celui-ci ; sous la forme d'une préparation combinée pour utilisation simultanée ou séquentielle dans le traitement d'un cancer comprenant b-Raf ayant une mutation V600, dans lequel

(A) est administré en une quantité d'environ 200 mg/jour à environ 3000 mg/jour, d'environ 1000 mg/jour à environ 2500 mg/jour, d'environ 1700 mg/jour à environ 2100 mg/jour ou d'environ 1920 mg/jour ;

(B) est administré en une quantité d'environ 50 mg/m²/semaine à environ 700 mg/m²/semaine, d'environ 100 mg/m²/semaine à environ 600 mg/m²/semaine, ou d'environ 200 mg/m²/semaine à environ 500 mg/m²/semaine ; et

(C) est administré en une quantité d'environ 1 à environ 250 mg/m²/semaine, d'environ 50 à environ 200 mg/m²/semaine, ou d'environ 125 mg/m²/semaine.

17. Utilisation de {3-[5-(4-chlorophényl)-1H-pyrrolo[2,3-b]pyridine-3-carbonyl]-2,4-difluorophényl}amide d'acide propa- ne-1-sulfonique ou d'un sel pharmaceutiquement acceptable de celui-ci et d'un inhibiteur d'EGFR pour la fabrication de médicaments destinés au traitement d'un cancer comprenant b-Raf ayant une mutation V600, dans laquelle l'inhibiteur d'EGFR est choisi parmi l'erlotinib, un sel pharmaceutiquement acceptable de celui-ci, et le cétuximab.

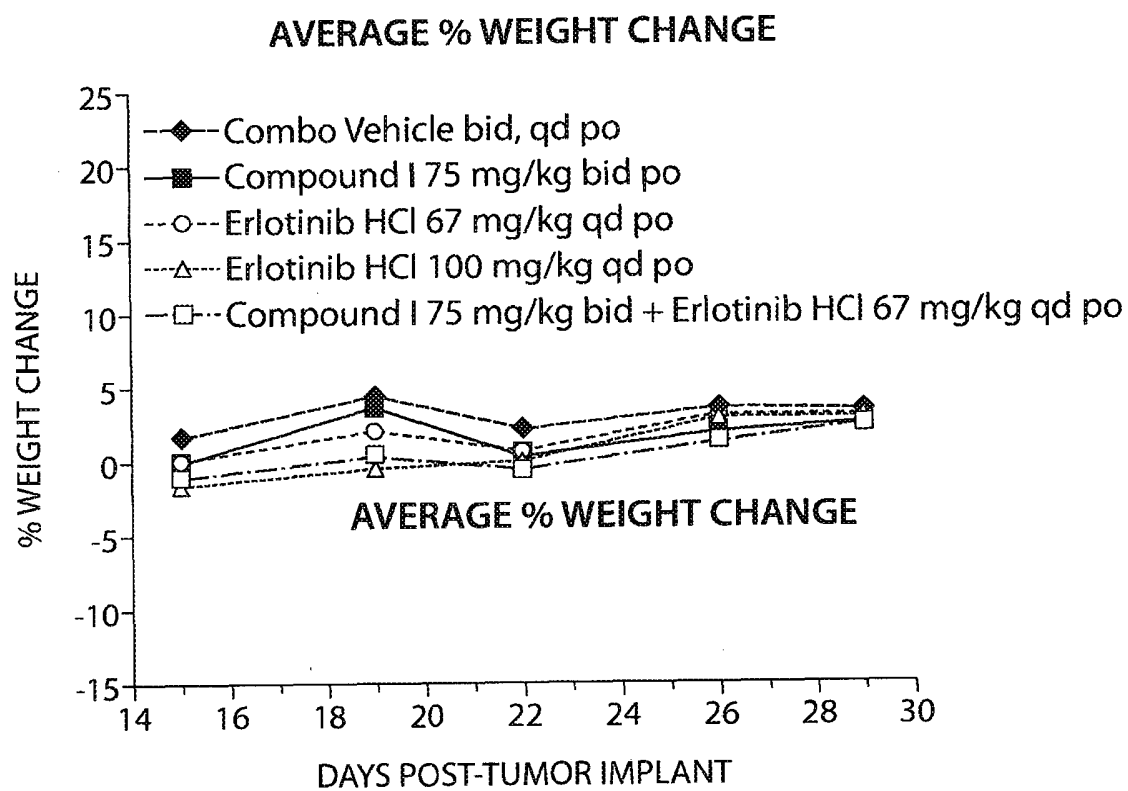


Fig. 1

- ◆ Combo Vehicle bid, qd
- Compound I 75 mg/kg bid
- ▲ Erlotinib HCl 67 mg/kg qd
- △ Erlotinib HCl 100 mg/kg qd
- Compound I 75 mg/kg bid + Erlotinib HCl 67 mg/kg qd

TGI(%)	N	PR	CR	ILS(%)
91	10	0	0	100
38	10	0	0	35
51	10	0	0	38
>100	10	9	0	142

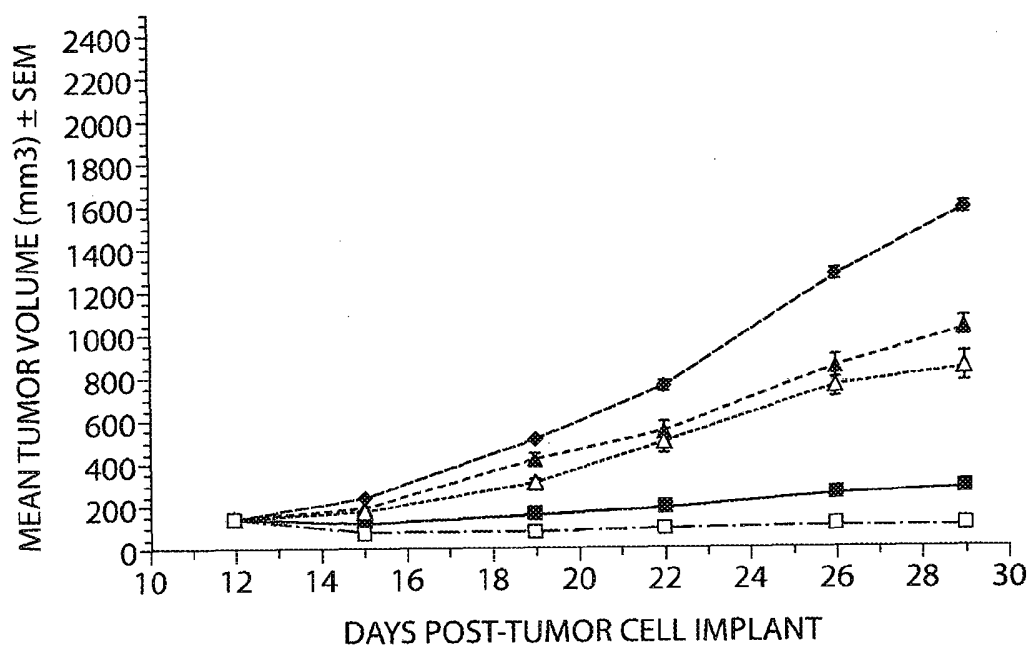


Fig. 2

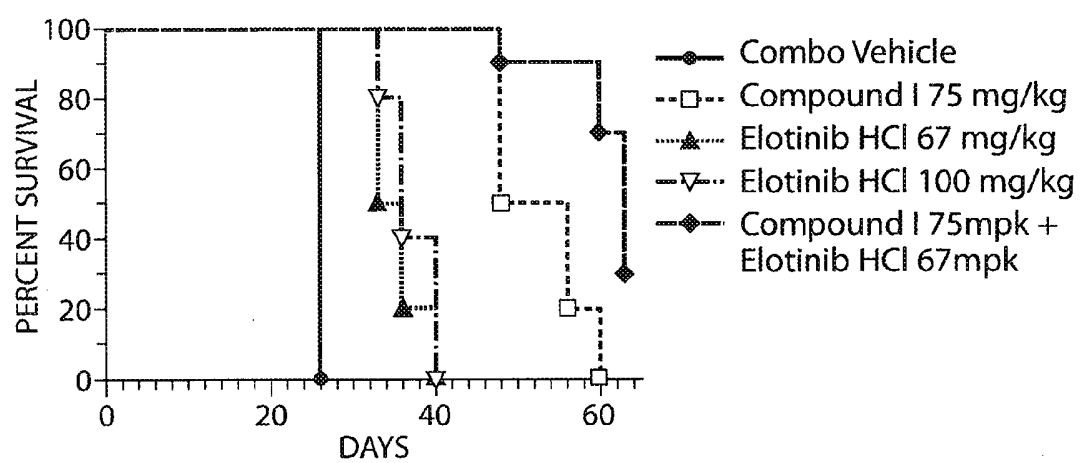


Fig. 3

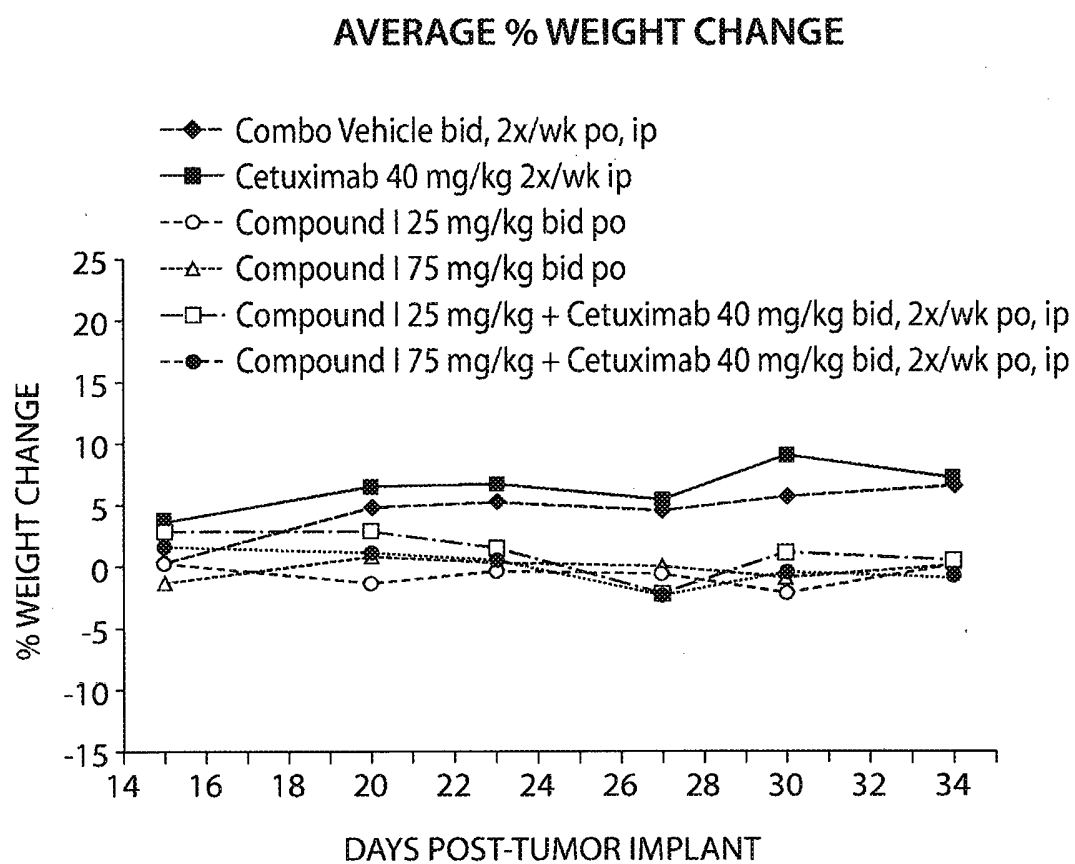


Fig. 4

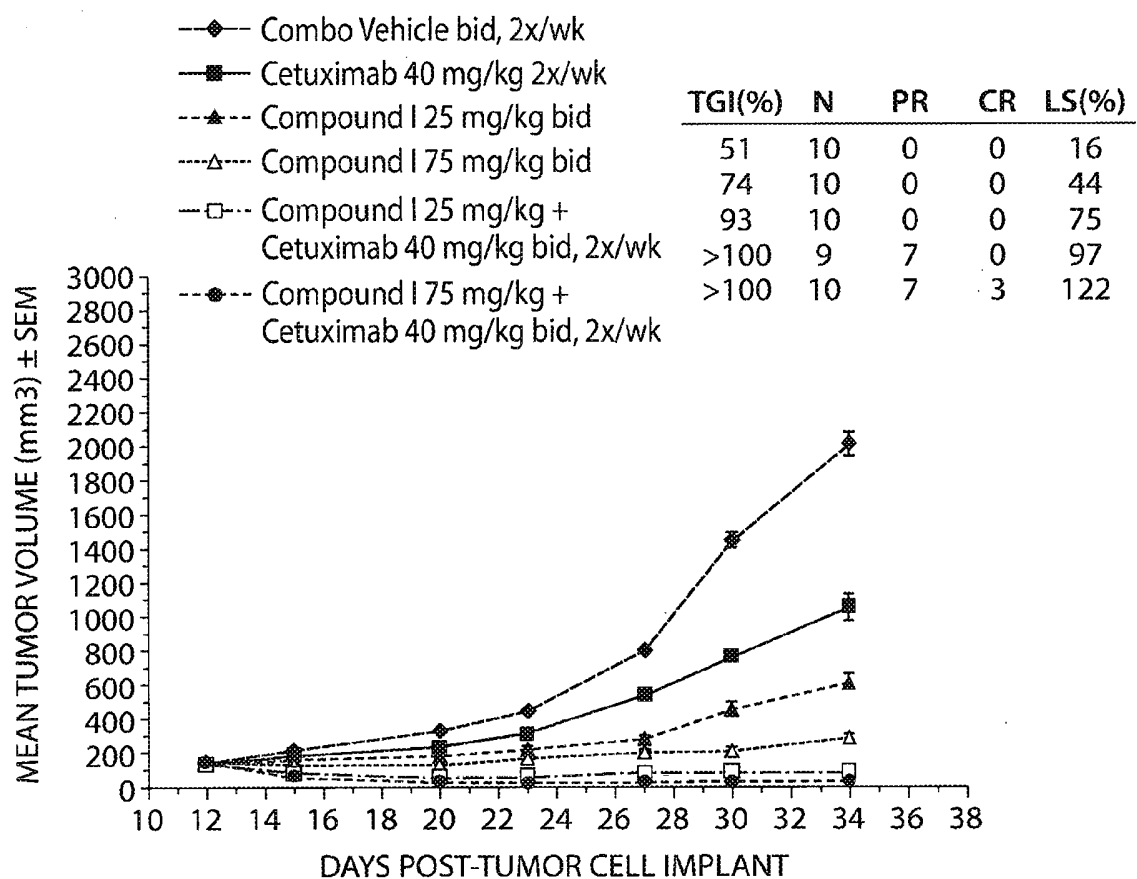


Fig. 5

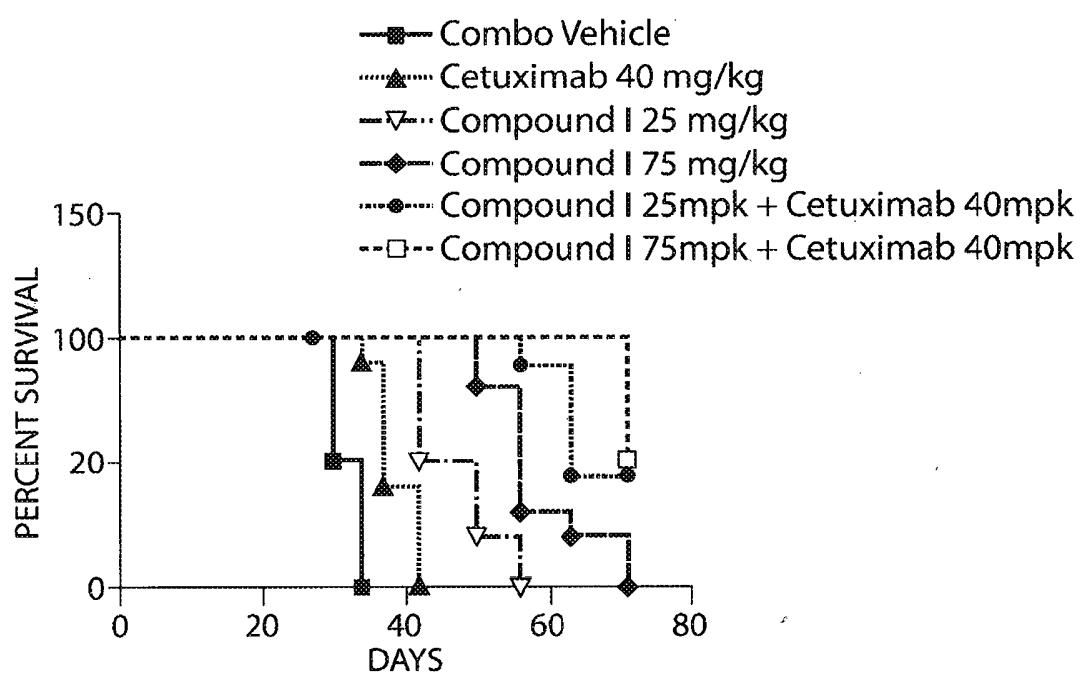


Fig. 6

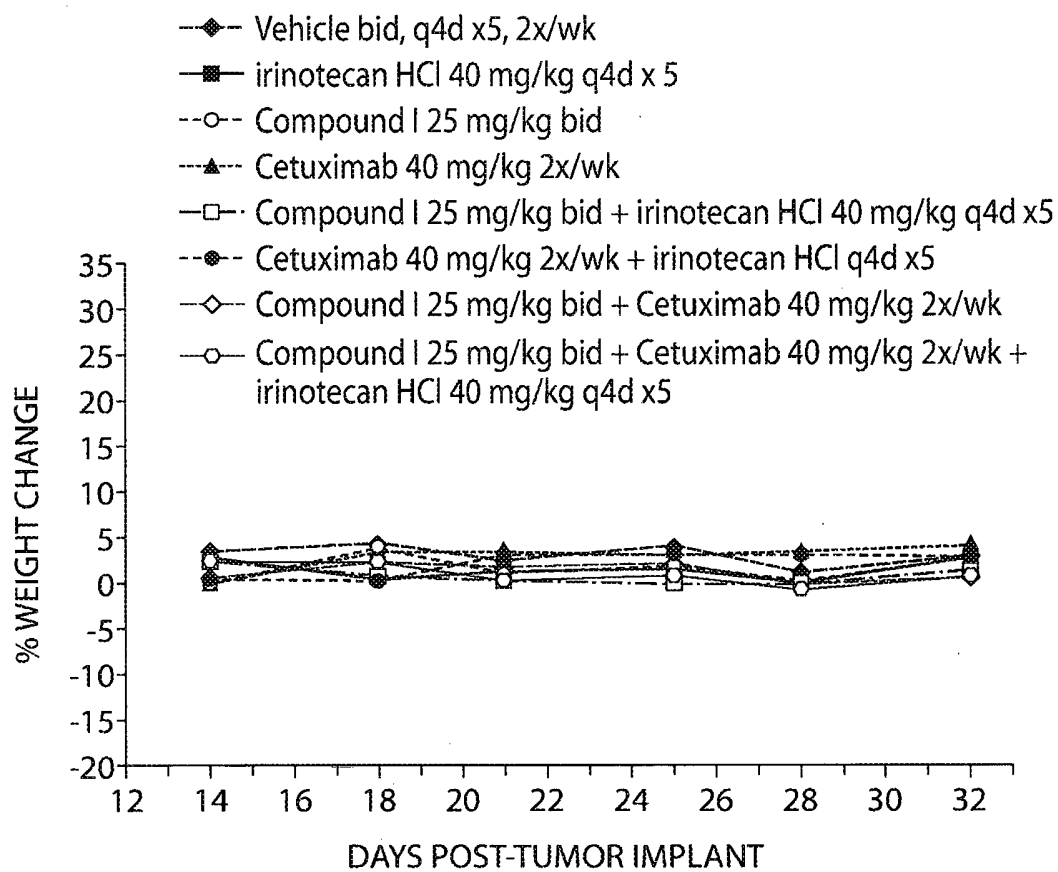


Fig. 7

	TGI(%)	N	PR	CR	ILS(%)
Vehicle bid, q4d x5, 2x/wk	59	10	0	0	17
irinotecan HCl 40 mg/kg q4d x 5	76	10	0	0	80
Compound I 25 mg/kg bid	58	10	0	0	27
Cetuximab 40 mg/kg bid	98	10	4	0	163
Compound I 25 mg/kg bid + irinotecan HCl 40 mg/kg q4d x 5	92	10	0	0	80
Cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x 5	>100	10	5	0	127
Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk	>100	10	9	1	250
Compound I 25 mg/kg bid + Cetuximab 40 mg/kg 2x/wk + irinotecan HCl 40 mg/kg q4d x 5					

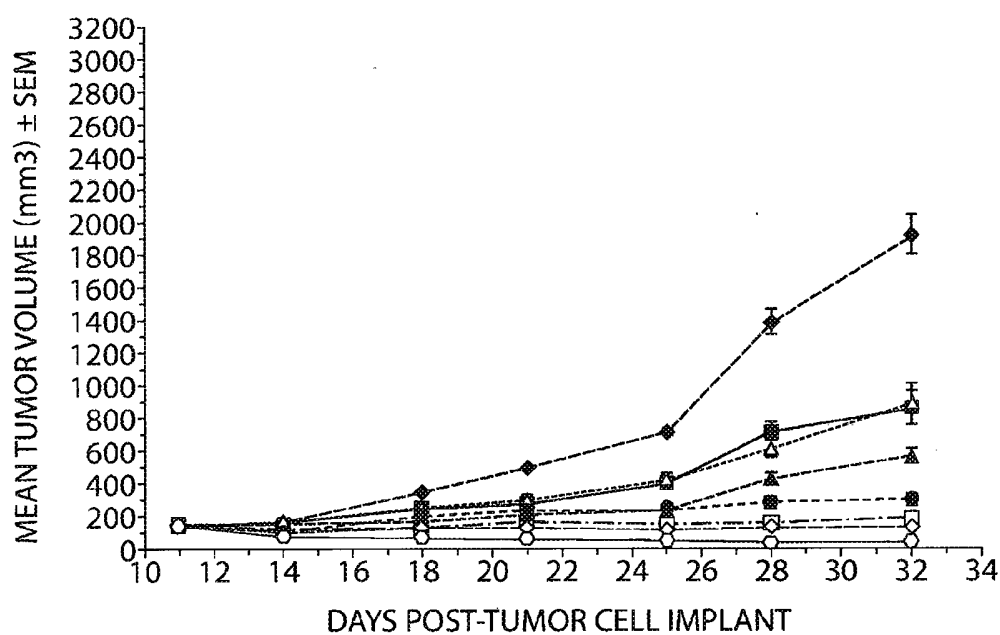


Fig. 8

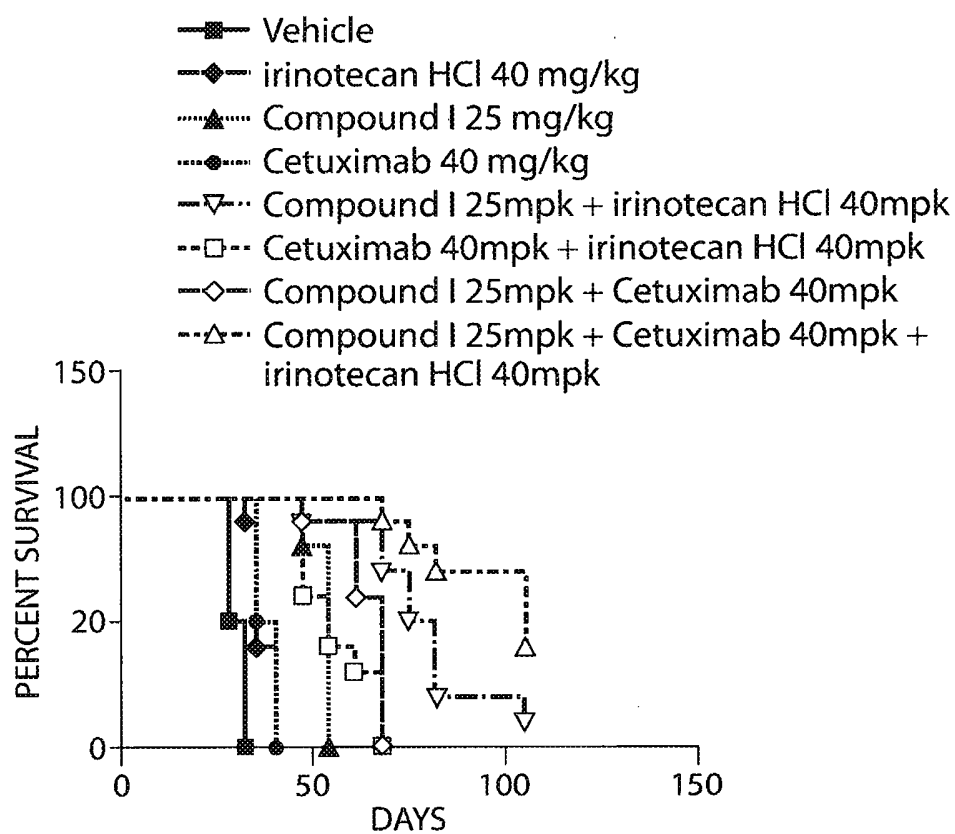


Fig. 9

REFERENCES CITED IN THE DESCRIPTION

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Az EP 2 605 764 lajstromszámú európai szabadalom igénypontjainak magyar fordítása:

1. Gyógyszerészeti termék, amely tartalmaz (A) egy első komponenst, amely hatóanyagként propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-fenil}-amidot (I vegyület) vagy gyógyászati lag elfogadható sóját tartalmazza; és (B) egy második komponenst, amely hatóanyagként EGFR inhibitor tartalmaz; mint kombinált készítmény egyidejű vagy egymást követő alkalmazásra olyan rák kezelésében, amely V600 mutációval rendelkező b-Raf-ot tartalmaz, ahol az EGFR inhibitor az erlotinib vagy annak gyógyászati lag elfogadható sója vagy a cetuximab közül választjuk.
2. Gyógyszerészeti termék az 1. igénypont szerinti alkalmazásra olyan vastag-végbél rák, melanoma és pajzsmirigy rák kezelésére, amely a V600E mutációval rendelkező b-Rafot tartalmazza.
3. Gyógyszerészeti termék az 1. és 2. igénypont szerinti alkalmazásra, amely tartalmaz továbbá (C) egy harmadik komponenst, amely hatóanyagként topoizomeráz inhibitor tartalmaz, ahol az említett topoizomeráz inhibitor irinotekán vagy gyógyászati lag elfogadható sója.
4. Gyógyszerészeti termék az 1-3. igénypontok bármelyike szerinti alkalmazásra, ahol a propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-fenil}-amid vagy gyógyászati lag elfogadható sója amorf vagy lényegében amorf formában van.
5. Kit, amely tartalmaz: (A) egy első komponenst, amely hatóanyagként propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-fenil}-amidot vagy gyógyászati lag elfogadható sóját tartalmazza; és (B) egy második komponenst, amely hatóanyagként EGFR inhibitor tartalmaz, ahol az EGFR inhibitor az erlotinib vagy gyógyászati lag elfogadható sója vagy a cetuximab közül választjuk.
6. Az 5. igénypont szerinti kit, amely tartalmaz továbbá egy harmadik komponenst, amely hatóanyagként irinotekánt tartalmaz.
7. Az 5. vagy 6. igénypont szerinti kit olyan rák kezelésében való alkalmazásra, amely tartalmaz V600 mutációval rendelkező b-Raf-ot.
8. Gyógyszerészeti termék az 1. igénypont szerinti alkalmazásra, ahol az említett propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-

fenil}-amidot vagy gyógyászatilag elfogadható sóját naponta kétszer adagoljuk kb. 850 mg – kb. 1050 mg mennyiségben; és az említett erlotinibet vagy gyógyászatilag elfogadható sóját naponta adagoljuk kb. 100 mg – kb. 200 mg mennyiségben.

9. Gyógyszerészeti termék az 1. igénypont szerinti alkalmazásra, ahol az említett propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-fenil}-amidot vagy gyógyászatilag elfogadható sóját naponta kétszer adagoljuk kb. 850 mg – kb. 1050 mg mennyiségben; és az említett cetuximabot hetenként adagoljuk, az első alkalommal a mennyisége kb. 400 mg/m^2 – kb. 500 mg/m^2 , és minden következő adagolás alkalmával a mennyisége kb. 200 mg/m^2 – kb. 300 mg/m^2 .

10. Gyógyszerészeti termék a 8. vagy 9. igénypont szerinti alkalmazásra, amely továbbá irinotekánt vagy gyógyászatilag elfogadható sóját tartalmazza.

11. Gyógyszerészeti termék a 10. igénypont szerinti alkalmazásra, ahol az említett irinotekánt vagy gyógyászatilag elfogadható sóját hetenként adagoljuk, kb. 50 mg/m^2 – kb. 200 mg/m^2 mennyiségben.

12. Gyógyszerészeti termék a 10. igénypont szerinti alkalmazásra, ahol az irinotekánt vagy gyógyászatilag elfogadható sóját hathetes ciklusban adagoljuk, hetente kb. 75 – kb. 175 mg/m^2 mennyiségben az első négy héten.

13. Gyógyszerészeti termék a 10. igénypont szerinti alkalmazásra, ahol az irinotekánt vagy gyógyászatilag elfogadható sóját minden harmadik héten egyszer adagoljuk kb. 300 – kb. 400 mg/m^2 mennyiségben; vagy minden második héten egyszer kb. 130 – kb. 230 mg/m^2 mennyiségben.

14. Gyógyszerészeti termék, amely tartalmaz (A) egy első komponenst, amely hatóanyagként I vegyületet vagy I vegyület gyógyászatilag elfogadható sóját tartalmazza; és (B) egy második komponenst, amely hatóanyagként erlotinibet vagy gyógyászatilag elfogadható sóját tartalmazza; mint kombinált készítmény egyidejű vagy egymást követő alkalmazásra olyan rák kezelésében, amely V600 mutációval rendelkező b-Raf-ot tartalmaz, ahol az (A) komponenst adagoljuk kb. 200 mg/nap-tól – kb. 3000 mg/nap-ig, kb. 1000 mg/nap-tól kb. 2500 mg/nap-ig, kb. 1700 mg/nap-tól kb. 2100 mg/napig terjedő vagy kb. 1920 mg/nap mennyiségben, és a (B) komponenst adagoljuk kb. 20 mg/nap-tól kb. 500 mg/nap-ig, kb. 100 mg/nap-tól kb. 400 mg/nap-ig, vagy kb. 100 mg/nap-tól kb. 200 mg/nap-ig terjedő mennyiségben.

15. Gyógyszerészeti termék, amely tartalmaz (A) egy első komponenst, amely hatóanyagként I vegyületet vagy I vegyület gyógyászatiilag elfogadható sóját tartalmazza; és (B) egy második komponenst, amely hatóanyagként cetuximabot tartalmaz; mint kombinált készítmény egyidejű vagy egymást követő alkalmazásra olyan rák kezelésében, amely V600 mutációval rendelkező b-Raf-ot tartalmaz, ahol az (A) komponenst adagoljuk kb. 200 mg/nap-tól – kb. 3000 mg/nap-ig, kb. 1000 mg/nap-tól kb. 2500 mg/nap-ig, kb. 1700 mg/nap-tól kb. 2100 mg/napig terjedő vagy kb. 1920 mg/nap mennyiségben; és

a (B) komponenst adagoljuk kb. 50 mg/m²/hét-től kb. 700 mg/m²/hét-ig, kb. 100 mg/m²/hét-től kb. 600 mg/m²/hét-ig, vagy kb. 200 mg/m²/hét-től kb. 500 mg/m²/hét-ig terjedő mennyiségben.

16. Gyógyszerészeti termék, amely tartalmaz (A) egy első komponenst, amely hatóanyagként I vegyületet vagy I vegyület gyógyászatiilag elfogadható sóját tartalmazza; (B) egy második komponenst, amely hatóanyagként cetuximabot tartalmaz;

és (C) egy harmadik komponenst, amely hatóanyagként irinotekánt vagy gyógyászatiilag elfogadható sóját tartalmazza; mint kombinált készítmény egyidejű vagy egymást követő alkalmazásra olyan rák kezelésében, amely V600 mutációval rendelkező b-Raf-ot tartalmaz, ahol

az (A) komponenst adagoljuk kb. 200 mg/nap-tól kb. 3000 mg/nap-ig, kb. 1000 mg/nap-tól kb. 2500 mg/nap-ig, kb. 1700 mg/nap-tól kb. 2100 mg/napig terjedő vagy kb. 1920 mg/nap mennyiségben;

a (B) komponenst adagoljuk 50 mg/m²/hét-től kb. 700 mg/m²/hét-ig, kb. 100 mg/m²/hét-től kb. 600 mg/m²/ig, vagy kb. 200 mg/m²/hét-től kb. 500 mg/m²/hét-ig terjedő mennyiségben; és

a (C) komponenst adagoljuk kb. 1 – kb. 250 mg/m²/hét, kb. 50 – kb. 200 mg/m²/hét, vagy kb. 125 mg/m²/hét mennyiségben.

17. Propán-1-szulfonsav-{3-[5-(4-klór-fenil)-1H-pirrolo[2,3-b]piridin-3-karbonil]-2,4-difluor-fenil}-amid vagy gyógyászatiilag elfogadható sója és EGFR inhibitor alkalmazása gyógyszerek előállítására olyan rák kezelésére, amely V600 mutációval rendelkező b-Raf-ot tartalmaz, ahol az EGFR inhibitort az erlotinib vagy gyógyászatiilag elfogadható sója vagy a cetuximab közül választjuk.

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