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VOLASERTIB IN COMBINATION WITH AZACITIDINE FOR THE TREATMENT OF ACUTE MYELOID LEUKEMIA AND MYELODYSPLASTIC SYNDROME II

(57) Abstract:  

The present invention relates to the use of Volasertib or a salt thereof or the hydrate thereof in combination with Azacitidine or a salt thereof or the hydrate thereof for treating patients suffering from acute myeloid leukemia (AML) or myelodysplastic syndrome (MDS).
The present invention relates to the use of Volasertib or a pharmaceutically acceptable salt thereof or the hydrate thereof in combination with Azacitidine or a pharmaceutically acceptable salt thereof or the hydrate thereof for treating patients suffering from acute myeloid leukemia (AML) and myelodysplastic syndrome (MDS).

Background of the invention

Acute myeloid leukemia (AML), also known as acute myelogenous leukemia, is a cancer of the myeloid line of blood cells, characterized by the rapid growth of abnormal white blood cells that accumulate in the bone marrow and interfere with the production of normal blood cells. As an acute leukemia, AML progresses rapidly and is typically fatal within weeks or months if left untreated. AML is the most prevalent form of adult acute leukemia, particularly among older adults and is slightly more common in men than women. There is an estimated prevalence of 30,000 cases of AML in the US and 47,000 in the EU.

The incidence of AML increases with age with a median age at diagnosis of 67 years. The global incidence CAGR for AML out to 2023 is 1.4%. An aging population, along with an increased incidence of treatment-related AML in cancer survivors, currently accounting for 10-20% of all AML cases, is expected to drive the incidence of AML. In addition, there is some geographic variation in the incidence of AML. In adults, the highest rates are seen in North America, Europe, and Oceania, while adult AML is rarer in Asia and Latin America.

AML accounts for approximately 1.2% of all cancer deaths. The 5 year survival rates for AML are low, driven by therapy failure and patients relapsing. Among patients <65 the 5 year survival rate is 34.4%, among patients >65 it is only 5%.

The WHO classification of myeloid neoplasms and acute leukemia is the current standard for classification of AML and incorporates genetic abnormalities into diagnostic algorithms. This classification is done by examining the appearance of the
malignant cells under light microscopy and by using cytogenetics and molecular genetics to characterize any underlying chromosomal abnormalities or genetic changes. The subtypes impact on prognoses, responses to therapy and treatment decisions.

**Myelodysplastic syndromes (MDS)** are clonal hematopoietic stem-cell disorders characterized by ineffective hematopoiesis, peripheral-blood cytopenias, and increased tendency to progress to acute myeloid leukemia (AML). The age-adjusted incidence of MDS is 3.3 cases per 100,000 people, and this rate appears to be increasing. MDS is primarily a disease of older adults, the median age of patients with MDS is approximately 70 years. This patient population is frequently affected by other comorbid conditions, which often influences treatment decisions. Treatment of MDS is based on prognostic factors that predict survival and progression to AML. Currently, the treatment of patients with MDS is guided by the International Prognostic Scoring System (IPSS). This system stratifies patients into four groups: low, intermediate-1, intermediate-2, and high-risk, based on number of cytopenias, percentage of bone marrow blasts, and karyotype. Low risk and intermediate-1 risk are usually grouped together as lower-risk disease, whereas intermediate-2 risk and high risk are grouped together as higher-risk disease. The survival of patients with higher-risk MDS is significantly different than that of patients with lower-risk disease. Without intervention, median survival of higher-risk patients is close to 12 months. Survival of patients with lower-risk disease is more diverse and ranges from a few months (poor-prognosis, lower-risk disease) to more than a decade. Therefore, the objectives of therapy are different in lower- versus higher-risk disease. While in lower-risk MDS, the goal is to relieve symptoms, manage cytopenias, and minimize the need for transfusions [eg: erythropoiesis-stimulating agents (ESA) and growth factors (GF)], in higher-risk MDS, disease-modifying therapies directed to slowing progression to AML and improving survival are used. These disease modifying therapies include hypomethylating agents (HMA, as e.g. azacitidine), intensive chemotherapy, and allogeneic stem cell transplantation (SCT), with SCT currently being the only known curative modality. Despite these treatment alternatives, the prognosis of patients with higher-risk MDS remains very poor owing to the disappointing activity of standard chemotherapy-based therapies, particularly those
with therapy-related MDS, the eventual loss of response to HMA, and the restriction of allogeneic SCT to younger patients with an appropriate donor.

Treatment of higher-risk patients is dependent on whether they are considered to be candidates for intensive therapy (e.g., allogeneic SCT or intensive chemotherapy). Clinical features relevant for this determination include the patient's age, performance status, comorbidities, patient's preference and availability of suitable donor and caregiver. The access to allogeneic SCT is restricted to approximately 8% of patients with MDS, owing to advanced age, concomitant comorbidities and/or donor availability. For higher-risk patients who are not candidates for high-intensity therapy, the use of HMA is considered the standard of care.

The efficacy of chemotherapeutic agents can be improved by using combination therapies with other compounds and/or improving the dosage schedule. Even if the concept of combining several therapeutic agents or improved dosage schedules already has been suggested, there is still a need for new and efficient therapeutic concepts for the treatment of cancer diseases, which show advantages over standard therapies.

Volasertib is a highly potent and selective inhibitor of the serine-threonine Polo like kinase (Plk), a key regulator of cell-cycle progression. Volasertib is a second-generation dihydropteridinone derivative with distinct pharmacokinetic (PK) properties. The problem underlying this invention was to develop a combination treatment and improved dosage schedules for combination therapy of Volasertib and Azacitidine in AML or MDS with maximal activity and limited toxicity.

Volasertib (I) is known as the compound N-[trans-4-[4-(cyclopropylmethyl)-1-piperazinyl][cyclohexyl]-4-[[7R]-7-ethyl-5,6,7,8-tetrahydro-5-methyl-8-(1-methylethyl)-6-oxo-2-pteridinyl]amino]-3-methoxy-benzamide,
This compound is disclosed in WO 04/076454. Furthermore, trihydrochloride salt forms and hydrates thereof are known from WO 07/090844. They possess properties which make those forms especially suitable for pharmaceutical use. The above mentioned patent applications further disclose the use of this compound or its monoethanesulfonate salt for the preparation of pharmaceutical compositions intended especially for the treatment of diseases characterized by excessive or abnormal cell proliferation.

Document WO 2006/018182 discloses other combinations for the treatment of diseases involving cell proliferation.

Azacitidine is a hypomethylating agent inhibiting DNA methyltransferases and known e.g. by the brand name Vidaza. Azacitidine has been studied in the treatment of previously treated and untreated, young adult and older AML and MDS patients.

**Summary of the Invention**

In animal experiments it has been found that a cancer treatment with Volasertib and Azacitidine comprise a synergistic efficacy profile (e.g. reduced tumor growth and beneficial side effect profile) compared to the monotherapy of both compounds.

Accordingly, a first object of the present invention refers to a pharmaceutical combination comprising Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, and Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for simultaneous, separate or sequential use of the active ingredients.
Another object of the present invention relates to a kit comprising one pharmaceutical composition comprising Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, and another pharmaceutical composition comprising Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof.

Another object of the present invention relates to a pharmaceutical kit, comprising a first compartment which comprises an effective amount of Volasertib and a second compartment which comprises Azacitidine, optionally together with an instruction for administration of both active ingredients to a patient suffering from AML or MDS, wherein according to said instruction Volasertib (in one embodiment 250, 300, 350, 400, 450 or 500 mg, in another embodiment 300 or 350 mg) and Azacitidine (in one embodiment 50 to 100 mg/m² BSA, in another embodiment 75 mg/m² BSA) is to be administered according to below mentioned dosage schedules.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS, characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, wherein both active ingredients can be administered simultaneously, separately or sequentially.

Another object of the present invention relates to Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS, characterized in that Azacitidine is administered in combination with Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, wherein both active ingredients can be administered simultaneously, separately or sequentially.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS characterized in that Volasertib is administered in combination with
Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to a dosage schedule (I) comprising or consisting of

a) administration of an effective amount of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof on minimally one day, preferably on two days, during a 4 week treatment cycle and

b) administration of an effective amount of Azacitidine on at least one day of the said 4 week treatment cycle

to a patient suffering from AML or MDS.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (II)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to dosage schedule (I), wherein Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof is administered on day 1 and on one of the days 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20 or 21 during a 4 week treatment cycle. Preferably, equal doses of Volasertib are administered on both days of administration.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (III)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I) or (II)) wherein in one embodiment 250 to 500 mg, in another embodiment 250, 300, 350, 400, 450 or 500 mg, yet in another embodiment 300 or 350 mg of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof are administered per day of administration.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (IV)), characterized in that Volasertib is administered
in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 5 days of the said 4 week treatment cycle, preferably from day 1 to 5.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (V)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 6 days of the said 4 week treatment cycle, preferably from day 1 to 6.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (VI)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 7 days of the said 4 week treatment cycle, preferably from day 1 to 7, alternatively from day 1 to 5 and day 8 to 9.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (VII)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 8 days of the said 4 week treatment cycle, preferably from day 1 to 8.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (VIII)), characterized in that Volasertib is
administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 9 days of the said 4 week treatment cycle, preferably from day 1 to 9.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (IX)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II) or (III)) wherein Azacitidine is administered on 10 days of the said 4 week treatment cycle, preferably from day 1 to 10.

Preferably Azacitidine is administered on 7 days of the said 4 week treatment cycle.

Another object of the present invention relates to Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML or MDS (dosage schedule (X)), characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to one of the above dosage schedules (dosage schedule (I), (II), (III), (IV), (V), (VI), (VII), (VIII) or (IX)) wherein in one embodiment 50 to 100 mg/m\(^2\) BSA, in another embodiment 75 mg/m\(^2\) BSA of Azacitidine are administered per day of administration.

Another object of the invention refers to a method of treating AML or MDS characterized in that Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, and Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, are administered according to one of the dosage schedules (I) to (X).

Another object of the invention refers to the use of Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for the manufacture
of a medicament for treating AML or MDS in patients suffering from AML or MDS wherein the medicament is prepared for administration according to one of the dosage schedules (I) to (X).

Another object of the invention refers to the use of Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for the manufacture of a medicament for treating AML or MDS in patients suffering from AML or MDS wherein the medicament is prepared for administration according to one of the dosage schedules (I) to (X).

Another object of the invention is a pharmaceutical composition comprising an effective amount of Volasertib and an effective amount of Azacitidine, optionally together with an instruction for administration of both active ingredients to a patient suffering from AML or MDS, wherein according to said instruction Volasertib is to be administered according to the above mentioned dosage schedules (I) to (X).

**Brief Description of the Figures**

**Figure 1** shows the tumor growth kinetics in a nude mouse xenograft model derived from human AML cell line MV4;1 1. Tumor-bearing mice were treated for 3 weeks either with vehicle or with 10 mg/kg Volasertib (BI 6727) once a week i.v., 40 mg/kg azacitidine once a week i.v. or a combination of Volasertib and azacitidine. Median tumor volumes are plotted over time. Day 1 was the first day, day 19 the last day of the experiment. Efficacy results from this xenograft models are considered valid for AML as well as MDS.

**Figure 2** shows the change of body weight over time in a nude mouse xenograft model derived from human AML cell line MV4;1 1. Tumor-bearing mice were treated for 3 weeks either with vehicle or with 10 mg/kg Volasertib once a week i.v., 40 mg/kg azacitidine once a week i.v. or a combination of Volasertib and azacitidine. Median changes in body weight compared to day 1 are plotted over time. Day 1 was the first day, day 19 the last day of the experiment.
Figure 3 shows the tumor growth kinetics in a nude mouse xenograft model derived from human AML cell line MV4;1 1. Tumor-bearing mice were treated for 3 weeks either with vehicle or with 20 mg/kg Volasertib once a week i.v., 40 mg/kg azacitidine once a week i.v. or a combination of Volasertib and azacitidine. Median tumor volumes are plotted over time. Day 1 was the first day, day 19 the last day of the experiment. Efficacy results from this xenograft models are considered valid for AML as well as MDS.

Figure 4 shows the change of body weight over time in a nude mouse xenograft model derived from human AML cell line MV4;1 1. Tumor-bearing mice were treated for 3 weeks either with vehicle or with 20 mg/kg Volasertib once a week i.v., 40 mg/kg azacitidine once a week i.v. or a combination of Volasertib and azacitidine. Median changes in body weight compared to day 1 are plotted over time. Day 1 was the first day, day 19 the last day of the experiment.

Detailed Description of the Invention

In case Volasertib is administered on minimally two days during a 4 week treatment cycle, then Volasertib is administered on two non-consecutive days during a 4 week treatment cycle.

The administration of an effective amount of Azacitidine on at least one day of the said 4 week treatment cycle means that during the 4 week treatment cycle in which Volasertib is administered minimally one time, also Azacitidine is administered at least one day.

The administration of Volasertib on day 1 and 15 during a 4 week treatment cycle means that one dosage of Volasertib or a pharmaceutically acceptable salt or a hydrate thereof is administered on day one and the second dosage is administered on day 15 to the patient suffering from AML or MDS in the four week treatment cycle.
The administration of Azacitidine from days 1 to 5, days 1 to 6, days 1 to 7, days 1 to 8, days 1 to 9 or from days 1 to 10, respectively, during a 4 week treatment cycle means that a daily dosage of Azacitidine or a pharmaceutically acceptable salt thereof is administered to the patient suffering from AML or MDS beginning on day one and ending with the last dosage on day 5, on day 6, on day 7, on day 8, on day 9 or on day 10, respectively, in the four week treatment cycle.

Accordingly, a complete four week treatment cycle according to one of the above mentioned dosage schedules may comprise the following administrations:

- Day 1: one dosage of Volasertib (e.g. 300 or 350 mg) and one dosage of Azacitidine (e.g. 75 mg/m² BSA);
- Day 2 to day 7 (including): one dosage of Azacitidine (e.g. 75 mg/m² BSA) per day;
- Day 8 to day 14 (including): no administration of Volasertib and Azacitidine;
- Day 15: one dosage of Volasertib (e.g. 300 or 350 mg);
- Day 16 to day 28 (including): no administration of Volasertib and Azacitidine.

This treatment cycle can be repeated as long as patients are eligible for repeated cycles, i.e. until progression of disease and as long as neither patient nor investigator requests treatment discontinuation.

The instruction for coadministration may be in any form suitable for pharmaceuticals, e.g. in form of a leaflet added to the dosage form within secondary packaging or an imprint on the primary or secondary packaging.

**Dosages / Volasertib:**

For intravenous treatment Volasertib may be administered to the human patient in a daily dose of 250 to 500 mg/application, in another embodiment 250, 300, 350, 400, 450 or 500 mg/application, yet in another embodiment 300 or 350 mg/application. For instance, Volasertib can be administered as a slow intravenous infusion over several hours, e.g. over about 1, 2, 4, 6, 10, 12 or 24 hours, preferably about 1 or 2 hours.

**Dosages / Azacitidine:**
Azacitidine may be administered in a total daily dose of 50 to 100 mg/m² BSA, e.g. in a total daily dose of 50, 55, 60, 65, 70, 75, 80, 85, 90, 95 or 100 mg/m² BSA one time daily. The total daily dose may also be divided into two or three subdoses to be taken within one day. Preferably, the daily dose is administered in a single dose of 75 mg/m² BSA.

However, it may optionally be necessary to deviate from the dosage amounts specified for Volasertib and Azacitidine, depending on the body weight on the method of administration, the individual response to the medication, the nature of the formulation used and the time or interval over which it is administered. Thus, in some cases, it may be sufficient to use less than the minimum quantity specified above, while in other cases the upper limit specified will have to be exceeded. When large amounts are administered it may be advisable to spread them over the day in a number of single doses.

Dosage Forms and Formulation Aspects

Regarding any aspects of the invention for Volasertib pharmaceutically acceptable salts or hydrates thereof may be used, preferably trihydrochloride salt forms and hydrates thereof as disclosed in WO 07/090844. Dosages or amounts of the active ingredient provided in the context of this invention refer in any case to the free base equivalent, that is Volasertib in the free base form.

The term "therapeutically effective amount" shall mean that amount of a drug or pharmaceutical agent that will elicit the biological or medical response of a tissue system, animal or human that is being sought by a researcher or clinician, resulting in a beneficial effect for at least a statistically significant fraction of patients, such as an improvement of symptoms, improvement of peripheral blood cell counts, a cure, a reduction in disease load, reduction in tumor mass or leukaemia cell numbers, extension of life, or improvement in quality of life.

Day 1 of a 4 week treatment cycle is defined as that day on which the first dose of Volasertib administered.
Within the present invention the term "AML" is to be understood to encompass all forms of acute myeloid leukemia and related neoplasms according to the 2008 revision of the World Health Organization (WHO) classification of myeloid neoplasms and acute leukemia. These are:

- Acute myeloid leukemia with recurrent genetic abnormalities
  - AML with t(8;21)(q22;q22); RUNX1-RUNX1T1
  - AML with inv(16)(p13q22) or t(16;16)(p13.1;q22); CBFB-MYH11
  - AML with t(9;11)(p22;q23); MLLT3-MLL
  - AML with t(6;9)(p23;q34); DEK-NUP214
  - AML with inv(3)(q21q22) or t(3;3)(q21;q26.2); RPN1-EVI1
  - AML (megakaryoblastic) with t(1;22)(p13;q13); RBM15-MKL1
  - Provisional entity: AML with mutated NPM1
  - Provisional entity: AML with mutated CEBPA

- Acute myeloid leukemia with myelodysplasia-related changes
- Therapy-related myeloid neoplasms
- Acute myeloid leukemia, not otherwise specified
  - AML with minimal differentiation
  - AML without maturation
  - AML with maturation
  - Acute myelomonocytic leukemia
  - Acute monoblastic/monocytic leukemia
  - Acute erythroid leukemia
    - Pure erythroid leukemia
    - Erythroleukemia, erythroid/myeloid
  - Acute megakaryoblastic leukemia
  - Acute basophilic leukemia
  - Acute panmyelosis with myelofibrosis

- Myeloid sarcoma
- Myeloid proliferations related to Down syndrome
  - Transient abnormal myelopoiesis
  - Myeloid leukemia associated with Down syndrome
- Blastic plasmacytoid dendritic cell neoplasm
Within the present invention the term "MDS" is to be understood to encompass all forms of myelodysplastic/myeloproliferative neoplasms (MDS/MPN) and myelodysplastic syndromes according to the 2008 revision of the World Health Organization (WHO) classification of myeloid neoplasms and acute leukemia. These are:

- Myelodysplastic/myeloproliferative neoplasms (MDS/MPN)
  - Chronic myelomonocytic leukemia
  - Atypical chronic myeloid leukemia, BCR-ABL1-negative
  - Juvenile myelomonocytic leukemia
  - Myelodysplastic/myeloproliferative neoplasm, unclassifiable
  - Provisional entity: refractory anemia with ring sideroblasts and thrombocytosis

- Myelodysplastic syndrome (MDS)
  - Refractory cytopenia with unilineage dysplasia
  - Refractory anemia
  - Refractory neutropenia
  - Refractory thrombocytopenia
  - Refractory anemia with ring sideroblasts
  - Refractory cytopenia with multilineage dysplasia
  - Refractory anemia with excess blasts
  - Myelodysplastic syndrome with isolated del(5q)
  - Myelodysplastic syndrome, unclassifiable
  - Childhood myelodysplastic syndrome
  - Provisional entity: refractory cytopenia of childhood

In accordance with the present invention Volasertib may be administered by parenteral (e.g. intramuscular, intraperitoneal, intravenous, transdermal or subcutaneous injection), and may be formulated, alone or together, in suitable dosage unit formulations containing conventional non-toxic pharmaceutically
acceptable carriers, adjuvants and vehicles appropriate for each route of administration. Dosage forms and formulations of both active ingredients suitable within the present invention are known in the art. For instance, such dosage forms and formulations include those disclosed for Volasertib in WO 2006/018221.

In accordance with the present invention Azacitidine may be administered by enteral or parenteral (e.g. intramuscular, intraperitoneal, intravenous, transdermal or subcutaneous injection, or implant), routes of administration and may be formulated, alone or together, in suitable dosage unit formulations containing conventional non-toxic pharmaceutically acceptable carriers, adjuvants and vehicles appropriate for each route of administration.

The following Examples serve to illustrate the invention without restricting it:

**Cells**

MV4;11 (CRL-9591) cells were obtained from ATCC. According to the Catalogue of Somatic Mutations in Cancer of the Wellcome Trust Sanger Institute, UK, this cell line carries a mutation in the FLT3 gene. Cells were cultured in T175 tissue culture flasks at 37°C and 5% CO2. The medium used was IMDM supplemented with 10% fetal calf serum, 1% NEAA, 1% sodium pyruvate and 1% glutamine.

**Mice**

Mice were 8-9 week-old athymic female BomTac: NMRI-Foxn1 nu purchased from Taconic, Denmark. After arrival in the animal facility, mice were allowed to adjust to ambient conditions for at least 3 days before they were used for experiments. They were housed in Macrolon® type II cages in groups of 5 under standardized conditions at 21.5 ± 1.5°C temperature and 55 ± 10% humidity. Standardized diet (PROVIMI KLIBA) and autoclaved tap water were provided ad libitum.

*Establishment of tumors, randomization*

To establish subcutaneous tumors, MV4;11 cells were harvested and resuspended in PBS + 5% FCS at 5 x 10^7 cells/ml. 50 µl of the cell suspension containing 2.5 x 10^6
cells was then injected subcutaneously into the right flank of the mice (1 site per mouse). Growth factor reduced BD Matrigel™ Matrix (BD Biosciences) was added to the cell suspension at a ratio of 1:1 before the injection. When tumors were well established and had reached a tumor volume of ~120 mm³, mice were randomly distributed between the treatment and the vehicle control groups 14 days after injecting the cells.

Administration of test compounds
Volasertib (BL 6727) was dissolved in hydrochloric acid (0.1 N), diluted with 0.9% NaCl and injected intravenously into the tail vein. An administration volume of 10 ml per kg body weight was used. The solution was freshly made up each injection day. Azacitidine was dissolved in 0.9% NaCl and administered i.v.. An administration volume of 10 ml per kg body weight was used. The application solution was prepared on each injection day.

Monitoring tumor growth and side effects
The tumor diameter was measured three times a week with a caliper. The volume of each tumor [in mm³] was calculated according to the formula "tumor volume = length * diameter² * π / 6". To monitor side effects of treatment, mice were inspected daily for abnormalities and body weight was determined three times a week. Animals were sacrificed at the end of the study when the control tumors reached a size of approximately 1100 mm³ on average. In addition animals with tumor sizes exceeding 2000 mm³ were sacrificed early during the studies for ethical reasons.

Example 1: nude mouse xenograft model derived from human AML cell line MV4;1
Results of an experiment comparing treatment of xenografts in mice with Volasertib alone (10 mg/kg), administered once weekly, Azacitidine alone (40 mg/kg), administered once weekly, and the combination of Volasertib / Azacitidine (10 mg/kg / 40 mg/kg) are shown in Figure 1. Animals were treated for 19 days. A combination of 10 mg/kg Volasertib plus 40 mg/kg azacitidine (T/C = 52%; T/C: ratio of median tumor volume of treated vs. control tumors) showed reduced tumor
growth compared to either single agent (Volasertib: T/C = 79%; azacitidine: T/C = 78%). Beneficial side effect profile was demonstrated as body weight gain in the combination group was comparable to single-agent azacitidine as shown in Figure 2.

**Example 2:** nude mouse xenograft model derived from human AML cell line MV4;11

Results of an experiment comparing treatment of xenografts in mice with Volasertib alone (20 mg/kg), administered once weekly, Azacitidine alone (40 mg/kg), administered once weekly, and the combination of Volasertib / Azacitidine (20 mg/kg / 40 mg/kg) are shown in Figure 3. Animals were treated for 19 days. A combination of 20 mg/kg Volasertib plus 40 mg/kg azacitidine (T/C = 18%) showed reduced tumor growth compared to either single agent (Volasertib: T/C = 39%; azacitidine: T/C = 78%). Beneficial side effect profile was demonstrated as body weight gain in the combination group was comparable to single-agent azacitidine as shown in Figure 4.
Claims

1. Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML and/or MDS, characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, wherein both active ingredients are administered simultaneously, separately or sequentially.

2. Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML and/or MDS, characterized in that Azacitidine is administered in combination with Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, wherein both active ingredients are administered simultaneously, separately or sequentially.

3. Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML and/or MDS according to claim 1 or 2, characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, according to a dosage schedule (I) comprising or consisting of
   a) administration of an effective amount of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof on minimally one day during a 4 week treatment cycle and
   b) administration of an effective amount of Azacitidine on at least one day of the said 4 week treatment cycle
to a patient suffering from AML and/or MDS.

4. Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for use of treating AML and/or MDS according to claim 1 or 2, characterized in that Volasertib is administered in combination with Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof
or a hydrate thereof, according to a dosage schedule (I) comprising or consisting of
  
a) administration of an effective amount of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof on minimally two days during a 4 week treatment cycle and
  
b) administration of an effective amount of Azacitidine on at least one day of the said 4 week treatment cycle
to a patient suffering from AML and/or MDS.

5. Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof for the use according to one or more of claims 1 to 4, wherein 250 to 500 mg of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof are administered per day of administration.

6. Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof for the use according to one or more of claims 1 to 5, wherein 50 to 100 mg/m² BSA of Azacitidine are administered per day of administration.

7. Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof for the use according to one or more of claims 1 to 6, wherein Azacitidine is administered on 5 days of the said 4 week treatment cycle.

8. Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof for the use according to one or more of claims 1 to 6, wherein Azacitidine is administered on 7 days of the said 4 week treatment cycle.

9. Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof for the use according to one or more of claims 1 to 6, wherein Azacitidine is administered on 10 days of the said 4 week treatment cycle.

10. Pharmaceutical composition comprising an effective amount of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof, and an effective
amount of Azacitidine or a pharmaceutically acceptable salt thereof or a hydrate thereof.

11. Pharmaceutical kit, comprising a first compartment which comprises an effective amount of Volasertib or a pharmaceutically acceptable salt thereof or a hydrate thereof and a second compartment which comprises an effective amount of Azacitidine or a pharmaceutically acceptable salt thereof or a hydrate thereof.

12. A pharmaceutical composition or pharmaceutical kit according to claim 10 or 11 for simultaneous, separate or sequential use as medicament for treating AML and/or MDS.

13. A pharmaceutical combination, characterized in that it comprises Volasertib, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, and Azacitidine, optionally in the form of a pharmaceutically acceptable salt thereof or a hydrate thereof, for simultaneous, separate or sequential use of the active ingredients as medicament for treating AML and/or MDS.
Figure 1: MV4;11 tumor growth kinetics
Figure 2: Change of body weight over time

- NaCl 0.9%
- azacitidine (40 mg/kg)
- BI 6727 (10 mg/kg)
- azacitidine (40 mg/kg) BI 6727 (10 mg/kg)
Figure 3: MV4;11 tumor growth kinetics
Figure 4: Change of body weight over time
### INTERNATIONAL SEARCH REPORT

**International application No.**

PCT/EP2014/065938

#### A. CLASSIFICATION OF SUBJECT MATTER

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#### ADD.

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

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

- A61K
- A61P

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

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

- EPO-Internal
- BIOSIS
- WPI Data
- EMBASE

#### C. DOCUMENTS CONSIDERED TO BE RELEVANT

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

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Date of the actual completion of the international search: 26 August 2014

Date of mailing of the international search report: 09/09/2014

Name and mailing address of the ISA:

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- NL - 2280 HV Rijswijk
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- Fax: (+31-31) 340-3016

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