ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Tasigna 50 mg hard capsules Tasigna 150 mg hard capsules Tasigna 200 mg hard capsules

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Tasigna 50 mg hard capsules

One hard capsule contains 50 mg nilotinib (as hydrochloride monohydrate).

Excipient with known effect

One hard capsule contains 39.03 mg lactose monohydrate.

Tasigna 150 mg hard capsules

One hard capsule contains 150 mg nilotinib (as hydrochloride monohydrate).

Excipient with known effect

One hard capsule contains 117.08 mg lactose monohydrate.

Tasigna 200 mg hard capsules

One hard capsule contains 200 mg nilotinib (as hydrochloride monohydrate).

Excipient with known effect

One hard capsule contains 156.11 mg lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Hard capsule.

Tasigna 50 mg hard capsules

White to yellowish powder in hard gelatin capsule with red opaque cap and light yellow opaque body, size 4 with black radial imprint "NVR/ABL" on cap.

Tasigna 150 mg hard capsules

White to yellowish powder in red opaque hard gelatin capsules, size 1 with black axial imprint "NVR/BCR".

Tasigna 200 mg hard capsules

White to yellowish powder in light yellow opaque hard gelatin capsules, size 0 with red axial imprint "NVR/TKI".

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Tasigna is indicated for the treatment of:

- adult and paediatric patients with newly diagnosed Philadelphia chromosome positive chronic myelogenous leukaemia (CML) in the chronic phase,
- adult patients with chronic phase and accelerated phase Philadelphia chromosome positive CML with resistance or intolerance to prior therapy including imatinib. Efficacy data in patients with CML in blast crisis are not available,
- paediatric patients with chronic phase Philadelphia chromosome positive CML with resistance or intolerance to prior therapy including imatinib.

4.2 Posology and method of administration

Therapy should be initiated by a physician experienced in the diagnosis and the treatment of patients with CML.

Posology

Treatment should be continued as long as clinical benefit is observed or until unacceptable toxicity occurs.

If a dose is missed the patient should not take an additional dose, but take the usual prescribed next dose.

Posology for Philadelphia chromosome positive CML adult patients

The recommended dose is:

- 300 mg twice daily in newly diagnosed patients with CML in the chronic phase,
- 400 mg twice daily in patients with chronic or accelerated phase CML with resistance or intolerance to prior therapy.

Posology for Philadelphia chromosome positive CML paediatric patients

Dosing in paediatric patients is individualised and is based on body surface area (mg/m²). The recommended dose of nilotinib is 230 mg/m² twice daily, rounded to the nearest 50 mg dose (to a maximum single dose of 400 mg) (see Table 1). Different strengths of Tasigna hard capsules can be combined to attain the desired dose.

There is no experience with treatment of paediatric patients below 2 years of age. There are no data in newly diagnosed paediatric patients below 10 years of age and limited data in imatinib-resistant or intolerant paediatric patients below 6 years of age.

Table 1 Paediatric dosing scheme of nilotinib 230 mg/m² twice daily

Body Surface Area	Dose in mg
(BSA)	(twice daily)
Up to 0.32 m^2	50 mg
$0.33 - 0.54 \text{ m}^2$	100 mg
$0.55 - 0.76 \text{ m}^2$	150 mg
$0.77 - 0.97 \text{ m}^2$	200 mg
$0.98 - 1.19 \text{ m}^2$	250 mg
$1.20 - 1.41 \text{ m}^2$	300 mg
$1.42 - 1.63 \text{ m}^2$	350 mg
$\geq 1.64 \text{ m}^2$	400 mg

Adult Philadelphia chromosome positive CML patients in chronic phase who have been treated with nilotinib as first-line therapy and who achieved a sustained deep molecular response (MR4.5)

Discontinuation of treatment may be considered in eligible adult Philadelphia chromosome positive (Ph+) CML patients in chronic phase who have been treated with nilotinib at 300 mg twice daily for a minimum of 3 years if a deep molecular response is sustained for a minimum of one year immediately prior to discontinuation of therapy. Discontinuation of nilotinib therapy should be initiated by a physician experienced in the treatment of patients with CML (see sections 4.4 and 5.1).

Eligible patients who discontinue nilotinib therapy must have their BCR-ABL transcript levels and complete blood count with differential monitored monthly for one year, then every 6 weeks for the second year, and every 12 weeks thereafter. Monitoring of BCR-ABL transcript levels must be performed with a quantitative diagnostic test validated to measure molecular response levels on the International Scale (IS) with a sensitivity of at least MR4.5 (BCR-ABL/ABL $\leq 0.0032\%$ IS).

For patients who lose MR4 (MR4=BCR-ABL/ABL \leq 0.01%IS) but not MMR (MMR=BCR-ABL/ABL \leq 0.1%IS) during the treatment-free phase, BCR-ABL transcript levels should be monitored every 2 weeks until BCR-ABL levels return to a range between MR4 and MR4.5. Patients who maintain BCR-ABL levels between MMR and MR4 for a minimum of 4 consecutive measurements can return to the original monitoring schedule.

Patients who lose MMR must re-initiate treatment within 4 weeks of when loss of remission is known to have occurred. Nilotinib therapy should be re-initiated at 300 mg twice daily or at a reduced dose level of 400 mg once daily if the patient had a dose reduction prior to discontinuation of therapy. Patients who re-initiate nilotinib therapy should have their BCR-ABL transcript levels monitored monthly until MMR is re-established and every 12 weeks thereafter (see section 4.4).

Adult Philadelphia chromosome positive CML patients in chronic phase who have achieved a sustained deep molecular response (MR 4.5) on nilotinib following prior imatinib therapy

Discontinuation of treatment may be considered in eligible adult Philadelphia chromosome positive (Ph+) CML patients in chronic phase who have been treated with nilotinib for a minimum of 3 years if a deep molecular response is sustained for a minimum of one year immediately prior to discontinuation of therapy. Discontinuation of nilotinib therapy should be initiated by a physician experienced in the treatment of patients with CML (see sections 4.4 and 5.1).

Eligible patients who discontinue nilotinib therapy must have their BCR-ABL transcript levels and complete blood count with differential monitored monthly for one year, then every 6 weeks for the second year, and every 12 weeks thereafter. Monitoring of BCR-ABL transcript levels must be performed with a quantitative diagnostic test validated to measure molecular response levels on the International Scale (IS) with a sensitivity of at least MR4.5 (BCR-ABL/ABL ≤0.0032% IS).

Patients with confirmed loss of MR4 (MR4= BCR-ABL/ABL $\leq 0.01\%$ IS) during the treatment-free phase (two consecutive measures separated by at least 4 weeks showing loss of MR4) or loss of major molecular response (MMR=BCR-ABL/ABL $\leq 0.1\%$ IS) must re-initiate treatment within 4 weeks of when loss of remission is known to have occurred. Nilotinib therapy should be re-initiated at either 300 mg or 400 mg twice daily. Patients who re-initiate nilotinib therapy should have their BCR-ABL transcript levels monitored monthly until previous major molecular response or MR4 level is re-established and every 12 weeks thereafter (see section 4.4).

Dose adjustments or modifications

Tasigna may need to be temporarily withheld and/or dose reduced for haematological toxicities (neutropenia, thrombocytopenia) that are not related to the underlying leukaemia (see Table 2).

Table 2 Dose adjustments for neutropenia and thrombocytopenia

Adult patients with newly diagnosed chronic phase CML at 300 mg twice daily and imatinib-resistant or intolerant CML in chronic phase at 400 mg twice daily	ANC* <1.0 x 10 ⁹ /l and/or platelet counts <50 x 10 ⁹ /l	 Treatment with nilotinib must be interrupted and blood count monitored. Treatment must be resumed within 2 weeks at prior dose if ANC >1.0 x 10⁹/l and/or platelets >50 x 10⁹/l. If blood counts remain low, a dose reduction to 400 mg once daily may be required.
Adult patients with imatinib-resistant or intolerant CML in accelerated phase at 400 mg twice daily	ANC* <0.5 x 10 ⁹ /l and/or platelet counts <10 x 10 ⁹ /l	 Treatment with nilotinib must be interrupted and blood count monitored. Treatment must be resumed within 2 weeks at prior dose if ANC >1.0 x 10⁹/l and/or platelets >20 x 10⁹/l. If blood counts remain low, a dose reduction to 400 mg once daily may be required.
Paediatric patients with newly diagnosed CML in chronic phase at 230 mg/m ² twice daily and imatinib-resistant or intolerant CML in chronic phase at 230 mg/m ² twice daily	ANC* <1.0 x 10 ⁹ /l and/or platelet counts <50 x 10 ⁹ /l	 Treatment with nilotinib must be interrupted and blood count monitored. Treatment must be resumed within 2 weeks at prior dose if ANC >1.5 x 10⁹/l and/or platelets >75 x 10⁹/l. If blood counts remain low, a dose reduction to 230 mg/m² once daily may be required. If event occurs after dose reduction, consider discontinuing treatment.

^{*}ANC = absolute neutrophil count

If clinically significant moderate or severe non-haematological toxicity develops, dosing should be interrupted, and patients should be monitored and treated accordingly. If the prior dose was 300 mg twice daily in adult newly diagnosed patients with CML in the chronic phase, or 400 mg twice daily in adult patients with imatinib-resistant or intolerant CML in chronic or accelerated phase, or 230 mg/m² twice daily in paediatric patients, dosing may be resumed at 400 mg once daily in adult patients and at 230 mg/m² once daily in paediatric patients once the toxicity has resolved. If the prior dose was 400 mg once daily in adult patients or 230 mg/m² once daily in paediatric patients, treatment should be discontinued. If clinically appropriate, re-escalation of the dose to the starting dose of 300 mg twice daily in adult newly diagnosed patients with CML in the chronic phase or to 400 mg twice daily in adult patients with imatinib-resistant or intolerant CML in chronic or accelerated phase or to 230 mg/m² twice daily in paediatric patients should be considered.

Elevated serum lipase: For Grade 3-4 serum lipase elevations, doses in adult patients should be reduced to 400 mg once daily or interrupted. In paediatric patients, treatment must be interrupted until the event returns to Grade ≤ 1 . Thereafter, if the prior dose was 230 mg/m² twice daily, treatment can be resumed at 230 mg/m² once daily. If the prior dose was 230 mg/m² once daily, treatment should be discontinued. Serum lipase levels should be tested monthly or as clinically indicated (see section 4.4).

Elevated bilirubin and hepatic transaminases: For Grade 3-4 bilirubin and hepatic transaminase elevations in adult patients, doses should be reduced to 400 mg once daily or interrupted. For Grade ≥ 2 bilirubin elevations or Grade ≥ 3 hepatic transaminase elevations in paediatric patients, treatment must be interrupted until the levels return to Grade ≤ 1 . Thereafter, if the prior dose was 230 mg/m² twice daily, treatment can be resumed at 230 mg/m² once daily. If the prior dose was 230 mg/m² once daily, and recovery to Grade ≤ 1 takes longer than 28 days, treatment should be discontinued. Bilirubin and hepatic transaminases levels should be tested monthly or as clinically indicated.

Special populations

Elderly

Approximately 12% of subjects in the Phase III study in patients with newly diagnosed CML in chronic phase and approximately 30% of subjects in the Phase II study in patients with imatinib-resistant or intolerant CML in chronic phase and accelerated phase were 65 years of age or over. No major differences were observed for safety and efficacy in patients ≥65 years of age as compared to adults aged 18 to 65 years.

Renal impairment

Clinical studies have not been performed in patients with impaired renal function. Since nilotinib and its metabolites are not renally excreted, a decrease in total body clearance is not anticipated in patients with renal impairment.

Hepatic impairment

Hepatic impairment has a modest effect on the pharmacokinetics of nilotinib. Dose adjustment is not considered necessary in patients with hepatic impairment. However, patients with hepatic impairment should be treated with caution (see section 4.4).

Cardiac disorders

In clinical studies, patients with uncontrolled or significant cardiac disease (e.g., recent myocardial infarction, congestive heart failure, unstable angina or clinically significant bradycardia) were excluded. Caution should be exercised in patients with relevant cardiac disorders (see section 4.4).

Increases in total serum cholesterol levels have been reported with nilotinib therapy (see section 4.4). Lipid profiles should be determined prior to initiating nilotinib therapy, assessed at month 3 and 6 after initiating therapy and at least yearly during chronic therapy.

Increases in blood glucose levels have been reported with nilotinib therapy (see section 4.4). Blood glucose levels should be assessed prior to initiating nilotinib therapy and monitored during treatment.

Paediatric population

The safety and efficacy of Tasigna in paediatric patients with Philadelphia chromosome positive CML in chronic phase from 2 to less than 18 years of age have been established (see sections 4.8, 5.1 and 5.2). There is no experience in paediatric patients below 2 years of age or in paediatric patients with Philadelphia chromosome positive CML in accelerated phase or blast crisis. There are no data in newly diagnosed paediatric patients below 10 years of age and limited data in imatinib-resistant or intolerant paediatric patients below 6 years of age.

Method of administration

Tasigna should be taken twice daily approximately 12 hours apart and must not be taken with food. The hard capsules should be swallowed whole with water. No food should be consumed for 2 hours before the dose is taken and no food should be consumed for at least one hour after the dose is taken.

For patients who are unable to swallow hard capsules, the content of each hard capsule may be dispersed in one teaspoon of apple sauce (puréed apple) and should be taken immediately. Not more than one teaspoon of apple sauce and no food other than apple sauce must be used (see sections 4.4 and 5.2).

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

4.4 Special warnings and precautions for use

Myelosuppression

Treatment with nilotinib is associated with (National Cancer Institute Common Toxicity Criteria grade 3 and 4) thrombocytopenia, neutropenia and anaemia. Occurrence is more frequent in patients with imatinib-resistant or intolerant CML, in particular in patients with accelerated-phase CML. Complete blood counts should be performed every two weeks for the first 2 months and then monthly thereafter, or as clinically indicated. Myelosuppression was generally reversible and usually managed by withholding Tasigna temporarily or dose reduction (see section 4.2).

QT prolongation

Nilotinib has been shown to prolong cardiac ventricular repolarisation as measured by the QT interval on the surface ECG in a concentration-dependent manner in adult and paediatric patients.

In the Phase III study in patients with newly diagnosed CML in chronic phase receiving 300 mg nilotinib twice daily, the change from baseline in mean time-averaged QTcF interval at steady state was 6 msec. No patient had a QTcF >480 msec. No episodes of torsade de pointes were observed.

In the Phase II study in imatinib-resistant and intolerant CML patients in chronic and accelerated phase receiving 400 mg nilotinib twice daily, the change from baseline in mean time-averaged QTcF interval at steady state was 5 and 8 msec, respectively. QTcF of >500 msec was observed in <1% of these patients. No episodes of torsade de pointes were observed in clinical studies.

In a healthy volunteer study with exposures that were comparable to the exposures observed in patients, the time-averaged mean placebo-subtracted QTcF change from baseline was 7 msec (CI \pm 4 msec). No subject had a QTcF >450 msec. Additionally, no clinically relevant arrhythmias were observed during the conduct of the trial. In particular, no episodes of torsade de pointes (transient or sustained) were observed.

Significant prolongation of the QT interval may occur when nilotinib is inappropriately taken with strong CYP3A4 inhibitors and/or medicinal products with a known potential to prolong the QT interval, and/or food (see section 4.5). The presence of hypokalaemia and hypomagnesaemia may further enhance this effect. Prolongation of the QT interval may expose patients to the risk of fatal outcome.

Tasigna should be used with caution in patients who have or who are at significant risk of developing prolongation of QTc, such as those:

- with congenital long QT prolongation
- with uncontrolled or significant cardiac disease including recent myocardial infarction, congestive heart failure, unstable angina or clinically significant bradycardia.
- taking anti-arrhythmic medicinal products or other substances that lead to QT prolongation.

Close monitoring for an effect on the QTc interval is advisable and a baseline ECG is recommended prior to initiating nilotinib therapy and as clinically indicated. Hypokalaemia or hypomagnesaemia must be corrected prior to Tasigna administration and should be monitored periodically during therapy.

Sudden death

Uncommon cases (0.1 to 1%) of sudden deaths have been reported in patients with imatinib-resistant or intolerant CML in chronic phase or accelerated phase with a past medical history of cardiac disease or significant cardiac risk factors. Co-morbidities in addition to the underlying malignancy were also frequently present as were concomitant medicinal products. Ventricular repolarisation abnormalities may have been contributory factors. No cases of sudden death were reported in the Phase III study in newly diagnosed patients with CML in chronic phase.

Fluid retention and oedema

Severe forms of drug-related fluid retention such as pleural effusion, pulmonary oedema, and pericardial effusion were uncommonly (0.1 to 1%) observed in a Phase III study of newly diagnosed CML patients. Similar events were observed in post-marketing reports. Unexpected, rapid weight gain should be carefully investigated. If signs of severe fluid retention appear during treatment with nilotinib, the aetiology should be evaluated and patients treated accordingly (see section 4.2 for instructions on managing non-haematological toxicities).

Cardiovascular events

Cardiovascular events were reported in a randomised Phase III study in newly diagnosed CML patients and observed in post-marketing reports. In this clinical study with a median on-therapy time of 60.5 months, Grade 3-4 cardiovascular events included peripheral arterial occlusive disease (1.4% and 1.1% at 300 mg and 400 mg nilotinib twice daily, respectively), ischaemic heart disease (2.2% and 6.1% at 300 mg and 400 mg nilotinib twice daily, respectively) and ischaemic cerebrovascular events (1.1% and 2.2% at 300 mg and 400 mg nilotinib twice daily, respectively). Patients should be advised to seek immediate medical attention if they experience acute signs or symptoms of cardiovascular events. The cardiovascular status of patients should be evaluated and cardiovascular risk factors monitored and actively managed during nilotinib therapy according to standard guidelines. Appropriate therapy should be prescribed to manage cardiovascular risk factors (see section 4.2 for instructions on managing non-haematological toxicities).

Hepatitis B reactivation

Reactivation of hepatitis B in patients who are chronic carriers of this virus has occurred after these patients received BCR-ABL tyrosine kinase inhibitors. Some cases resulted in acute hepatic failure or fulminant hepatitis leading to liver transplantation or a fatal outcome.

Patients should be tested for HBV infection before initiating treatment with nilotinib. Experts in liver disease and in the treatment of hepatitis B should be consulted before treatment is initiated in patients with positive hepatitis B serology (including those with active disease) and for patients who test positive for HBV infection during treatment. Carriers of HBV who require treatment with nilotinib should be closely monitored for signs and symptoms of active HBV infection throughout therapy and for several months following termination of therapy (see section 4.8).

Special monitoring of adult Ph+ CML patients in chronic phase who have achieved a sustained deep molecular response

Eligibility for discontinuation of treatment

Eligible patients who are confirmed to express the typical BCR-ABL transcripts, e13a2/b2a2 or e14a2/b3a2, can be considered for treatment discontinuation. Patients must have typical BCR-ABL transcripts to allow quantitation of BCR-ABL, evaluation of the depth of molecular response, and determination of a possible loss of molecular remission after discontinuation of treatment with nilotinib.

Monitoring of patients who have discontinued therapy

Frequent monitoring of BCR-ABL transcript levels in patients eligible for treatment discontinuation must be performed with a quantitative diagnostic test validated to measure molecular response levels with a sensitivity of at least MR4.5 (BCR-ABL/ABL \leq 0.0032% IS). BCR-ABL transcript levels must be assessed prior to and during treatment discontinuation (see sections 4.2 and 5.1).

Loss of major molecular response (MMR=BCR-ABL/ABL \leq 0.1%IS) in CML patients who received nilotinib as first- or second-line therapy, or confirmed loss of MR4 (two consecutive measures separated by at least 4 weeks showing loss of MR4 (MR4=BCR-ABL/ABL \leq 0.01%IS)) in CML patients who received nilotinib as second-line therapy will trigger treatment re-initiation within

4 weeks of when loss of remission is known to have occurred. Molecular relapse can occur during the treatment-free phase, and long-term outcome data are not yet available. It is therefore crucial to perform frequent monitoring of BCR-ABL transcript levels and complete blood count with differential in order to detect possible loss of remission (see section 4.2). For patients who fail to achieve MMR after three months of treatment re-initiation, BCR-ABL kinase domain mutation testing should be performed.

Laboratory tests and monitoring

Blood lipids

In a Phase III study in newly diagnosed CML patients, 1.1% of the patients treated with 400 mg nilotinib twice daily showed a Grade 3-4 elevation in total cholesterol; no Grade 3-4 elevations were however observed in the 300 mg twice daily dose group (see section 4.8). It is recommended that the lipid profiles be determined before initiating treatment with nilotinib, assessed at month 3 and 6 after initiating therapy and at least yearly during chronic therapy (see section 4.2). If a HMG-CoA reductase inhibitor (a lipid-lowering agent) is required, please refer to section 4.5 before initiating treatment since certain HMG-CoA reductase inhibitors are also metabolised by the CYP3A4 pathway.

Blood glucose

In a Phase III study in newly diagnosed CML patients, 6.9% and 7.2% of the patients treated with 400 mg nilotinib and 300 mg nilotinib twice daily, respectively, showed a Grade 3-4 elevation in blood glucose. It is recommended that the glucose levels be assessed before initiating treatment with Tasigna and monitored during treatment, as clinically indicated (see section 4.2). If test results warrant therapy, physicians should follow their local standards of practice and treatment guidelines.

Interactions with other medicinal products

The administration of Tasigna with agents that are strong CYP3A4 inhibitors (including, but not limited to, ketoconazole, itraconazole, voriconazole, clarithromycin, telithromycin, ritonavir) should be avoided. Should treatment with any of these agents be required, it is recommended that nilotinib therapy be interrupted if possible (see section 4.5). If transient interruption of treatment is not possible, close monitoring of the individual for prolongation of the QT interval is indicated (see sections 4.2, 4.5 and 5.2).

Concomitant use of nilotinib with medicinal products that are potent inducers of CYP3A4 (e.g., phenytoin, rifampicin, carbamazepine, phenobarbital and St. John's Wort) is likely to reduce exposure to nilotinib to a clinically relevant extent. Therefore, in patients receiving nilotinib, co-administration of alternative therapeutic agents with less potential for CYP3A4 induction should be selected (see section 4.5).

Food effect

The bioavailability of nilotinib is increased by food. Tasigna must not be taken in conjunction with food (see sections 4.2 and 4.5) and should be taken 2 hours after a meal. No food should be consumed for at least one hour after the dose is taken. Grapefruit juice and other foods that are known to inhibit CYP3A4 should be avoided. For patients who are unable to swallow hard capsules, the content of each hard capsule may be dispersed in one teaspoon of apple sauce and should be taken immediately. Not more than one teaspoon of apple sauce and no food other than apple sauce must be used (see section 5.2).

Hepatic impairment

Hepatic impairment has a modest effect on the pharmacokinetics of nilotinib. Single dose administration of 200 mg of nilotinib resulted in increases in AUC of 35%, 35% and 19% in subjects with mild, moderate and severe hepatic impairment, respectively, compared to a control group of subjects with normal hepatic function. The predicted steady-state C_{max} of nilotinib showed an increase of 29%, 18% and 22%, respectively. Clinical studies have excluded patients with alanine transaminase

(ALT) and/or aspartate transaminase (AST) >2.5 (or >5, if related to disease) times the upper limit of the normal range and/or total bilirubin >1.5 times the upper limit of the normal range. Metabolism of nilotinib is mainly hepatic. Patients with hepatic impairment might therefore have increased exposure to nilotinib and should be treated with caution (see section 4.2).

Serum lipase

Elevation in serum lipase has been observed. Caution is recommended in patients with previous history of pancreatitis. In case lipase elevations are accompanied by abdominal symptoms, nilotinib therapy should be interrupted and appropriate diagnostic measures considered to exclude pancreatitis.

Total gastrectomy

The bioavailability of nilotinib might be reduced in patients with total gastrectomy (see section 5.2). More frequent follow-up of these patients should be considered.

Tumour lysis syndrome

Due to possible occurrence of tumour lysis syndrome (TLS) correction of clinically significant dehydration and treatment of high uric acid levels are recommended prior to initiating nilotinib therapy (see section 4.8).

Lactose

Tasigna hard capsules contain lactose. Patients with rare hereditary problems of galactose intolerance, the Lapp lactase deficiency or glucose-galactose malabsorption should not take this medicinal product.

Paediatric population

Laboratory abnormalities of mild to moderate transient elevations of aminotransferases and total bilirubin have been observed in children at a higher frequency than in adults, indicating a higher risk of hepatotoxicity in the paediatric population (see section 4.8). Liver function (bilirubin and hepatic transaminases levels) should be monitored monthly or as clinically indicated. Elevations of bilirubin and hepatic transaminases should be managed by withholding nilotinib temporarily, dose reduction and/or discontinuation of nilotinib (see section 4.2). In a study in the CML paediatric population, growth retardation has been documented in patients treated with nilotinib (see section 4.8). Close monitoring of growth in paediatric patients under nilotinib treatment is recommended.

4.5 Interaction with other medicinal products and other forms of interaction

Tasigna may be given in combination with haematopoietic growth factors such as erythropoietin or granulocyte colony-stimulating factor (G-CSF) if clinically indicated. It may be given with hydroxyurea or anagrelide if clinically indicated.

Nilotinib is mainly metabolised in the liver with CYP3A4 expected to be the main contributor to the oxidative metabolism. Nilotinib is also a substrate for the multi-drug efflux pump, P-glycoprotein (P-gp). Therefore, absorption and subsequent elimination of systemically absorbed nilotinib may be influenced by substances that affect CYP3A4 and/or P-gp.

Substances that may increase nilotinib serum concentrations

Concomitant administration of nilotinib with imatinib (a substrate and moderator of P-gp and CYP3A4), had a slight inhibitory effect on CYP3A4 and/or P-gp. The AUC of imatinib was increased by 18% to 39%, and the AUC of nilotinib was increased by 18% to 40%. These changes are unlikely to be clinically important.

The exposure to nilotinib in healthy subjects was increased 3-fold when co-administered with the strong CYP3A4 inhibitor ketoconazole. Concomitant treatment with strong CYP3A4 inhibitors, including ketoconazole, itraconazole, voriconazole, ritonavir, clarithromycin, and telithromycin, should therefore be avoided (see section 4.4). Increased exposure to nilotinib might also be expected with moderate CYP3A4 inhibitors. Alternative concomitant medicinal products with no or minimal CYP3A4 inhibition should be considered.

Substances that may decrease nilotinib serum concentrations

Rifampicin, a potent CYP3A4 inducer, decreases nilotinib C_{max} by 64% and reduces nilotinib AUC by 80%. Rifampicin and nilotinib should not be used concomitantly.

The concomitant administration of other medicinal products that induce CYP3A4 (e.g. phenytoin, carbamazepine, phenobarbital and St. John's Wort) is likewise likely to reduce exposure to nilotinib to a clinically relevant extent. In patients for whom CYP3A4 inducers are indicated, alternative agents with less enzyme induction potential should be selected.

Nilotinib has pH dependent solubility, with lower solubility at higher pH. In healthy subjects receiving esomeprazole at 40 mg once daily for 5 days, gastric pH was markedly increased, but nilotinib absorption was only decreased modestly (27% decrease in C_{max} and 34% decrease in AUC_0 - ∞). Nilotinib may be used concurrently with esomeprazole or other proton pump inhibitors as needed.

In a study in healthy subjects, no significant change in nilotinib pharmacokinetics was observed when a single 400 mg dose of nilotinib was administered 10 hours after and 2 hours before famotidine. Therefore, when the concurrent use of a H2 blocker is necessary, it may be administered approximately 10 hours before and approximately 2 hours after the dose of Tasigna.

In the same study as above, administration of an antacid (aluminium hydroxide/magnesium hydroxide/simethicone) 2 hours before or after a single 400 mg dose of nilotinib also did not alter nilotinib pharmacokinetics. Therefore, if necessary, an antacid may be administered approximately 2 hours before or approximately 2 hours after the dose of Tasigna.

Substances that may have their systemic concentration altered by nilotinib

In vitro, nilotinib is a relatively strong inhibitor of CYP3A4, CYP2C8, CYP2C9, CYP2D6 and UGT1A1, with Ki value being lowest for CYP2C9 (Ki=0.13 microM).

A single-dose drug-drug interaction study in healthy volunteers with 25 mg warfarin, a sensitive CYP2C9 substrate, and 800 mg nilotinib did not result in any changes in warfarin pharmacokinetic parameters or warfarin pharmacodynamics measured as prothrombin time (PT) and international normalised ratio (INR). There are no steady-state data. This study suggests that a clinically meaningful drug-drug interaction between nilotinib and warfarin is less likely up to a dose of 25 mg of warfarin. Due to lack of steady-state data, control of warfarin pharmacodynamic markers (INR or PT) following initiation of nilotinib therapy (at least during the first 2 weeks) is recommended.

In CML patients, nilotinib administered at 400 mg twice daily for 12 days increased the systemic exposure (AUC and C_{max}) of oral midazolam (a substrate of CYP3A4) 2.6-fold and 2.0-fold, respectively. Nilotinib is a moderate CYP3A4 inhibitor. As a result, the systemic exposure of other medicinal products primarily metabolised by CYP3A4 (e.g. certain HMG-CoA reductase inhibitors) may be increased when co-administered with nilotinib. Appropriate monitoring and dose adjustment may be necessary for medicinal products that are CYP3A4 substrates and have a narrow therapeutic index (including but not limited to alfentanil, cyclosporine, dihydroergotamine, ergotamine, fentanyl, sirolimus and tacrolimus) when co-administered with nilotinib.

The combination of nilotinib with those statins that are mainly eliminated by CYP3A4, may increase the potential for statin-induced myopathy, including rhabdomyolysis.

Anti-arrhythmic medicinal products and other substances that may prolong the QT interval

Nilotinib should be used with caution in patients who have or may develop prolongation of the QT interval, including those patients taking anti-arrhythmic medicinal products such as amiodarone, disopyramide, procainamide, quinidine and sotalol or other medicinal products that may lead to QT prolongation such as chloroquine, halofantrine, clarithromycin, haloperidol, methadone and moxifloxacin (see section 4.4).

Food interactions

The absorption and bioavailability of nilotinib are increased if it is taken with food, resulting in a higher serum concentration (see sections 4.2, 4.4 and 5.2). Grapefruit juice and other foods that are known to inhibit CYP3A4 should be avoided.

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential/Contraception

Women of childbearing potential have to use highly effective contraception during treatment with nilotinib and for up to two weeks after ending treatment.

Pregnancy

There are no or limited amount of data from the use of nilotinib in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Tasigna should not be used during pregnancy unless the clinical condition of the woman requires treatment with nilotinib. If it is used during pregnancy, the patient must be informed of the potential risk to the foetus.

If a woman who is being treated with nilotinib is considering pregnancy, treatment discontinuation may be considered based on the eligibility criteria for discontinuing treatment as described in sections 4.2 and 4.4. There is a limited amount of data on pregnancies in patients while attempting treatment-free remission (TFR). If pregnancy is planned during the TFR phase, the patient must be informed of a potential need to re-initiate nilotinib treatment during pregnancy (see sections 4.2 and 4.4).

Breast-feeding

It is unknown whether nilotinib is excreted in human milk. Available toxicological data in animals have shown excretion of nilotinib in milk (see section 5.3). Since a risk to the newborns/infants cannot be excluded, women should not breast-feed during Tasigna treatment and for 2 weeks after the last dose.

<u>Fertility</u>

Animal studies did not show an effect on fertility in male and female rats (see section 5.3).

4.7 Effects on ability to drive and use machines

Tasigna has no or negligible influence on the ability to drive and use machines. However, it is recommended that patients experiencing dizziness, fatigue, visual impairment or other undesirable effects with a potential impact on the ability to drive or use machines safely should refrain from these activities as long as the undesirable effects persist (see section 4.8).

4.8 Undesirable effects

Summary of the safety profile

The safety profile is based on pooled data from 3,422 patients treated with Tasigna in 13 clinical studies in the approved indications: adults and paediatric patients with newly diagnosed Philadelphia chromosome positive chronic myelogenous leukaemia (CML) in the chronic phase (5 clinical studies with 2,414 patients), adult patients with chronic phase and accelerated phase Philadelphia chromosome positive CML with resistance or intolerance to prior therapy including imatinib (6 clinical studies with 939 patients) and paediatric patients with chronic phase Philadelphia chromosome positive CML with resistance or intolerance to prior therapy including imatinib (2 clinical studies with 69 patients). These pooled data represents 9,039.34 patient-years of exposure.

The safety profile of nilotinib is consistent across indications.

The most common adverse reactions (incidence $\geq 15\%$) from the pooled safety data were: rash (26.4%), upper respiratory tract infection (including pharyngitis, nasopharyngitis, rhinitis) (24.8%) headache (21.9%), hyperbilirubinaemia (including blood bilirubin increased) (18.6%), arthralgia (15.8%), fatigue (15.4%), nausea (16.8%), pruritus (16.7%) and thrombocytopenia (16.4%).

Tabulated list of adverse reactions

Adverse reactions from clinical studies and post-marketing reports (Table 3) are listed by MedDRA system organ class and frequency category. Frequency categories are defined using the following convention: very common ($\geq 1/10$); common ($\geq 1/100$ to < 1/10); uncommon ($\geq 1/1,000$); rare ($\leq 1/10,000$); very rare (< 1/10,000); not known (cannot be estimated from the available data).

Table 3 Adverse drug reactions

Infections and infes	tations	
Very common:	Upper respiratory tract infection (including pharyngitis, nasopharyngitis, rhinitis)	
Common:	Folliculitis, bronchitis, candidiasis (including oral candidiasis), pneumonia, gastroenteritis, urinary tract infection	
Uncommon:	Herpes virus infection, anal abscess, candidiasis (candida infection), furuncle, sepsis, subcutaneous abscess, tinea pedis	
Rare:	Hepatitis B reactivation	
Neoplasms benign,	malignant and unspecified (including cysts and polyps)	
Uncommon:	Skin papilloma	
Rare:	Oral papilloma, paraproteinaemia	
Blood and lymphati	c system disorders	
Very common:	Anaemia, thrombocytopenia	
Common:	Leukopenia, leukocytosis, neutropenia, thrombocythaemia	
Uncommon:	Eosinophilia, febrile neutropenia, lymphopenia, pancytopenia	
Immune system disc	orders	
Uncommon:	Hypersensitivity	
Endocrine disorders	8	
Very common:	Growth retardation	
Common:	Hypothyroidism	
Uncommon:	Hyperthyroidism	
Rare:	Hyperparathyroidism secondary, thyroiditis	

Metabolism and nutrition	on disorders		
Common:	Electrolyte imbalance (including hypomagnesaemia, hyperkalaemia,		
	hypokalaemia, hyponatraemia, hypocalcaemia, hypercalcaemia,		
	hyperphosphataemia), diabetes mellitus, hyperglycaemia,		
	hypercholesterolaemia, hyperlipidaemia, hypertriglyceridaemia,		
	decreased appetite, gout, hyperuricaemia, hypophosphataemia (including		
	blood phosphorus decreased)		
Uncommon:	Dehydration, increased appetite, dyslipidaemia, hypoglycaemia		
Rare:	Appetite disorder, tumour lysis syndrome		
Psychiatric disorders			
Common:	Depression, insomnia, anxiety		
Uncommon:	Amnesia, confusional state, disorientation		
Rare:	Dysphoria		
Nervous system disorde			
Very common:	Headache		
Common:	Dizziness, hypoaesthesia, paraesthesia, migraine		
Uncommon:	Cerebrovascular accident, intracranial/cerebral haemorrhage, ischaemic		
	stroke, transient ischaemic attack, cerebral infarction, loss of		
	consciousness (including syncope), tremor, disturbance in attention,		
	hyperaesthesia, dysaesthesia, lethargy, peripheral neuropathy, restless		
	legs syndrome, facial paralysis		
Rare:	Basilar artery stenosis, brain oedema, optic neuritis		
Eye disorders	/ 1		
Common:	Conjunctivitis, dry eye (including xerophthalmia), eye irritation,		
	hyperaemia (scleral, conjunctival, ocular), vision blurred		
Uncommon:	Visual impairment, conjunctival haemorrhage, visual acuity reduced,		
eyelid oedema, blepharitis, photopsia, conjunctivitis allergic, d			
	eye haemorrhage, eye pain, eye pruritus, eye swelling, ocular surface		
	disease, periorbital oedema, photophobia		
Rare:	Chorioretinopathy, papilloedema		
Ear and labyrinth disor	ders		
Common:	Vertigo, ear pain, tinnitus		
Uncommon:	Hearing impaired (hypoacusis)		
Cardiac disorders			
Common:	Angina pectoris, arrhythmia (including atroventricular block, cardiac		
	flutter, ventricular extrasystoles, tachycardia, atrial fibrillation,		
	bradycardia), palpitations, electrocardiogram QT prolonged, coronary		
	artery disease		
Uncommon:	Myocardial infarction, cardiac murmur, pericardial effusion, cardiac		
	failure, diastolic dysfunction, left bundle branch block, pericarditis		
Rare:	Cyanosis, ejection fraction decreased		
Not known:	Ventricular dysfunction		
Vascular disorders			
Common:	Hypertension, flushing, peripheral arterial occlusive disease		
Uncommon:	Hypertensive crisis, intermittent claudication, peripheral artery stenosis,		
	Hypertensive crisis, intermittent claudication, peripheral artery stenosis,		
	haematoma, arteriosclerosis, hypotension, thrombosis		
Rare:			
	haematoma, arteriosclerosis, hypotension, thrombosis		
	haematoma, arteriosclerosis, hypotension, thrombosis Shock haemorrhagic		
Respiratory, thoracic ar	haematoma, arteriosclerosis, hypotension, thrombosis Shock haemorrhagic d mediastinal disorders		
Respiratory, thoracic ar	haematoma, arteriosclerosis, hypotension, thrombosis Shock haemorrhagic nd mediastinal disorders Cough		
Respiratory, thoracic and Very common: Common:	haematoma, arteriosclerosis, hypotension, thrombosis Shock haemorrhagic d mediastinal disorders Cough Dyspnoea, dyspnoea exertional, epistaxis, oropharyngeal pain		
Respiratory, thoracic and Very common: Common:	haematoma, arteriosclerosis, hypotension, thrombosis Shock haemorrhagic d mediastinal disorders Cough Dyspnoea, dyspnoea exertional, epistaxis, oropharyngeal pain Pulmonary oedema, pleural effusion, interstitial lung disease, pleuritic		

Gastrointestinal diso	rders
Very common:	Nausea, upper abdominal pain, constipation, diarrhoea, vomiting
Common:	Pancreatitis, abdominal discomfort, abdominal distension, flatulence,
Common.	abdominal pain, dyspepsia, gastritis, gastroesophageal reflux,
	haemorrhoids, stomatitis
Uncommon:	Gastrointestinal haemorrhage, melaena, mouth ulceration, oesophageal
encommon.	pain, dry mouth, sensitivity of teeth (hyperaesthesia teeth), dysgeusia,
	enterocolitis, gastric ulcer, gingivitis, hiatus hernia, rectal haemorrhage
Rare:	Gastrointestinal ulcer perforation, haematemesis, oesophageal ulcer,
Ture.	oesophagitis ulcerative, retroperitoneal haemorrhage, subileus
Hepatobiliary disord	
Very common:	Hyperbilirubinaemia (including blood bilirubin increased)
Common:	Hepatic function abnormal
Uncommon:	Hepatotoxicity, toxic hepatitis, jaundice, cholestasis, hepatomegaly
Skin and subcutaneo	
Very common:	Rash, pruritus, alopecia
Common:	Night sweats, eczema, urticaria, hyperhidrosis, contusion, acne,
	dermatitis (including allergic, exfoliative and acneiform), dry skin,
	erythema
Uncommon:	Exfoliative rash, drug eruption, skin pain, ecchymosis, swelling face,
	blister, dermal cysts, erythema nodosum, hyperkeratosis, petechiae,
	photosensitivity, psoriasis, skin discolouration, skin exfoliation, skin
	hyperpigmentation, skin hypertrophy, skin ulcer
Rare:	Erythema multiforme, palmar-plantar erythrodysaesthesia syndrome,
	sebaceous hyperplasia, skin atrophy
Musculoskeletal and	connective tissue disorders
Very common	Myalgia, arthralgia, back pain, pain in extremity
Common:	Musculoskeletal chest pain, musculoskeletal pain, neck pain, muscular
	weakness, muscle spasms, bone pain
Uncommon:	Musculoskeletal stiffness, joint swelling, arthritis, flank pain
Renal and urinary di	sorders
Common:	Pollakiuria, dysuria
Uncommon:	Micturition urgency, nocturia, chromaturia, haematuria, renal failure,
	urinary incontinence
Reproductive system	and breast disorders
Common:	Erectile dysfunction, menorrhagia
Uncommon:	Breast pain, gynaecomastia, nipple swelling
Rare:	Breast induration
General disorders an	d administration site conditions
Very common	Fatigue, pyrexia
Common:	Chest pain (including non-cardiac chest pain), pain, chest discomfort,
	malaise, asthenia and oedema peripheral, chills, influenza-like illness
Uncommon:	Face oedema, gravitational oedema, feeling body temperature change
	(including feeling hot, feeling cold), localised oedema
Rare:	Sudden death

Investigations		
Very common:	Alanine aminotransferase increased, lipase increased	
Common:	Haemoglobin decreased, blood amylase increased, aspartate	
	aminotransferase increased, blood alkaline phosphatase increased,	
	gamma-glutamyltransferase increased, blood creatinine phosphokinase	
	increased, weight decreased, weight increased, elevated creatinine, total	
	cholesterol increased	
Uncommon:	Blood lactate dehydrogenase increased, blood urea increased, blood	
	bilirubin unconjugated increased, blood parathyroid hormone increased,	
	blood triglycerides increased, globulins decreased, lipoprotein	
	cholesterol (including low density and high density) increased, troponin	
	increased	
Rare:	Blood glucose decreased, blood insulin decreased, blood insulin	
	increased, insulin C-peptide decreased	

Note: Not all adverse drug reactions were observed in paediatric studies.

Description of selected adverse reactions

Sudden death

Uncommon cases (0.1 to 1%) of sudden deaths have been reported in Tasigna clinical studies and/or compassionate use programs in patients with imatinib -resistant or intolerant CML in chronic phase or accelerated phase with a past medical history of cardiac disease or significant cardiac risk factors (see section 4.4).

Hepatitis B reactivation

Hepatitis B reactivation has been reported in association with BCR-ABL TKIs. Some cases resulted in acute hepatic failure or fulminant hepatitis leading to liver transplantation or a fatal outcome (see section 4.4).

Paediatric population

The safety of nilotinib in paediatric patients (from 2 to <18 years of age) with Philadelphia chromosome positive CML in chronic phase (n=58) has been investigated in one main study over a period of 60 months (see section 5.1). In paediatric patients, the frequency, type and severity of adverse reactions observed have been generally consistent with those observed in adults, with the exception of hyperbilirubinaemia/blood bilirubin increase (Grade 3/4: 10.3%) and transaminase elevation (AST Grade 3/4: 1.7%, ALT Grade 3/4: 12.1%) which were reported at a higher frequency than in adult patients. Bilirubin and hepatic transaminase levels should be monitored during treatment (see sections 4.2 and 4.4).

Growth retardation in paediatric population

In a study conducted in the CML paediatric population, with a median exposure of 51.9 months in newly diagnosed patients and 59.9 months in imatinib/dasatinib-resistant or imatinib-intolerant Ph+CML-CP patients, growth deceleration (crossing at least two main percentile lines from baseline) was observed in eight patients: five (8.6%) crossed two main percentile lines from baseline and three (5.2%) crossed three main percentile lines from baseline. Growth retardation related events were reported in 3 patients (5.2%). Close monitoring of growth in paediatric patients under nilotinib treatment is recommended (see section 4.4).

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

Isolated reports of intentional overdose with nilotinib were reported, where an unspecified number of Tasigna hard capsules were ingested in combination with alcohol and other medicinal products. Events included neutropenia, vomiting and drowsiness. No ECG changes or hepatotoxicity were reported. Outcomes were reported as recovered.

In the event of overdose, the patient should be observed and appropriate supportive treatment given.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antineoplastic agents, BCR-ABL tyrosine kinase inhibitors, ATC code: L01EA03.

Mechanism of action

Nilotinib is a potent inhibitor of the ABL tyrosine kinase activity of the BCR-ABL oncoprotein both in cell lines and in primary Philadelphia-chromosome positive leukaemia cells. The substance binds with high affinity to the ATP-binding site in such a manner that it is a potent inhibitor of wild-type BCR-ABL and maintains activity against 32/33 imatinib-resistant mutant forms of BCR-ABL. As a consequence of this biochemical activity, nilotinib selectively inhibits the proliferation and induces apoptosis in cell lines and in primary Philadelphia-chromosome positive leukaemia cells from CML patients. In murine models of CML, as a single agent nilotinib reduces tumour burden and prolongs survival following oral administration.

Pharmacodynamic effects

Nilotinib has little or no effect against the majority of other protein kinases examined, including Src, except for the PDGF, KIT and Ephrin receptor kinases, which it inhibits at concentrations within the range achieved following oral administration at therapeutic doses recommended for the treatment of CML (see Table 4).

Table 4 Kinase profile of nilotinib (phosphorylation IC₅₀ nM)

BCR-ABL	PDGFR	KIT
20	69	210

Clinical efficacy

Clinical studies in newly diagnosed CML in chronic phase

An open-label, multicentre, randomised Phase III study was conducted to determine the efficacy of nilotinib versus imatinib in 846 adult patients with cytogenetically confirmed newly diagnosed Philadelphia chromosome positive CML in the chronic phase. Patients were within six months of diagnosis and were previously untreated, with the exception of hydroxyurea and/or anagrelide. Patients were randomised 1:1:1 to receive either nilotinib 300 mg twice daily (n=282), nilotinib 400 mg twice daily (n=281) or imatinib 400 mg once daily (n=283). Randomisation was stratified by Sokal risk score at the time of diagnosis.

Baseline characteristics were well balanced between the three treatment arms. Median age was 47 years in both nilotinib arms and 46 years in the imatinib arm, with 12.8%, 10.0% and 12.4% of patients were \geq 65 years of age in the nilotinib 300 mg twice daily, nilotinib 400 mg twice daily and imatinib 400 mg once daily treatment arms, respectively. There were slightly more male than female patients (56.0%, 62.3% and 55.8%, in the nilotinib 300 mg twice daily, 400 mg twice daily and

imatinib 400 mg once daily arm, respectively). More than 60% of all patients were Caucasian and 25% of all patients were Asian.

The primary data analysis time point was when all 846 patients completed 12 months of treatment (or discontinued earlier). Subsequent analyses reflect when patients completed 24, 36, 48, 60 and 72 months of treatment (or discontinued earlier). The median time on treatment was approximately 70 months in the nilotinib treatment groups and 64 months in the imatinib group. The median actual dose intensity was 593 mg/day for nilotinib 300 mg twice daily, 772 mg/day for nilotinib 400 mg twice daily and 400 mg/day for imatinib 400 mg once daily. This study is ongoing.

The primary efficacy endpoint was major molecular response (MMR) at 12 months. MMR was defined as \leq 0.1% BCR-ABL/ABL% by international scale (IS) measured by RQ-PCR, which corresponds to a \geq 3 log reduction of BCR-ABL transcript from standardised baseline. The MMR rate at 12 months was statistically significantly higher for nilotinib 300 mg twice daily compared to imatinib 400 mg once daily (44.3% versus 22.3%, p<0.0001). The rate of MMR at 12 months, was also statistically significantly higher for nilotinib 400 mg twice daily compared to imatinib 400 mg once daily (42.7% versus 22.3%, p<0.0001).

The rates of MMR at 3, 6, 9 and 12 months were 8.9%, 33.0%, 43.3% and 44.3% for nilotinib 300 mg twice daily, 5.0%, 29.5%, 38.1% and 42.7% for nilotinib 400 mg twice daily and 0.7%, 12.0%, 18.0% and 22.3% for imatinib 400 mg once daily.

The MMR rate at 12, 24, 36, 48, 60 and 72 months is presented in Table 5.

Table 5 MMR rate

	•	•	
	Nilotinib	Nilotinib	Imatinib
	300 mg twice daily	400 mg twice daily	400 mg once daily
	n=282	n=281	n=283
	(%)	(%)	(%)
MMRat 12 months			
Response (95% CI)	44.31 (38.4; 50.3)	42.71 (36.8; 48.7)	22.3 (17.6; 27.6)
MMR at 24 months			
Response (95% CI)	61.71 (55.8; 67.4)	59.11(53.1; 64.9)	37.5 (31.8; 43.4)
MMR at 36 months ²			
Response (95% CI)	58.51 (52.5; 64.3)	57.3 ¹ (51.3; 63.2)	38.5 (32.8; 44.5)
MMR at 48 months ³			
Response (95% CI)	59.91 (54.0; 65.7)	55.2 (49.1; 61.1)	43.8 (38.0; 49.8)
MMR at 60 months ⁴			
Response (95% CI)	62.8 (56.8; 68.4)	61.2 (55.2; 66.9)	49.1 (43.2; 55.1)
MMR at 72 months ⁵			
Response (95% CI)	52.5 (46.5; 58.4)	57.7 (51.6; 63.5)	41.7 (35.9; 47.7)
1	(C) (TT)		100 \ 0.0001

¹Cochran-Mantel-Haenszel (CMH) test p-value for response rate (vs. imatinib 400 mg) <0.0001 ²Only patients who were in MMR at a specific time point are included as responders for that time point. A total of 199 (35.2%) of all patients were not evaluable for MMR at 36 months (87 in the nilotinib 300 mg twice daily group and 112 in the imatinib group) due to missing/unevaluable PCR assessments (n=17), atypical transcripts at baseline (n=7), or discontinuation prior to the 36-month time point (n=175).

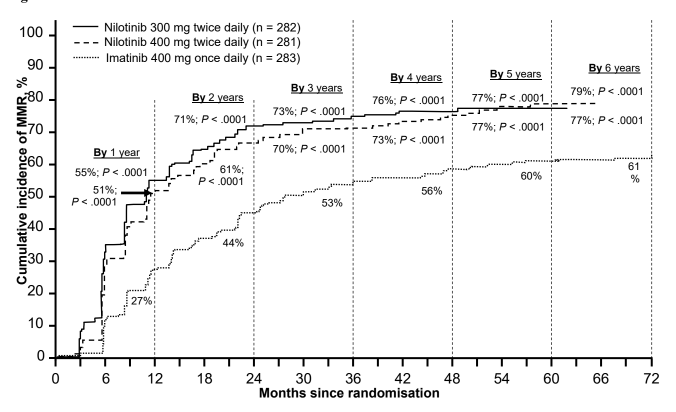
MMR rates by different time points (including patients who achieved MMR at or before those time points as responders) are presented in the cumulative incidence of MMR (see Figure 1).

³ Only patients who were in MMR at a specific time point are included as responders for that time point. A total of 305 (36.1%) of all patients were not evaluable for MMR at 48 months (98 in the nilotinib 300 mg BID group, 88 in the nilotinib 400 mg BID group and 119 in the imatinib group) due to missing/unevaluable PCR assessments (n=18), atypical transcripts at baseline (n=8), or discontinuation prior to the 48-month time point (n=279).

⁴ Only patients who were in MMR at a specific time point are included as responders for that time point. A total of 322 (38.1%) of all patients were not evaluable for MMR at 60 months (99 in the nilotinib 300 mg twice daily group, 93 in the nilotinib 400 mg twice daily group and 130 in the imatinib group) due to missing/unevaluable PCR assessments (n=9), atypical transcripts at baseline (n=8) or discontinuation prior to the 60-month time point (n=305).

Only patients who were in MMR at a specific time point are included as responders for that time point. A total of 395 (46.7%) of all patients were not evaluable for MMR at 72 months (130 in the nilotinib 300 mg twice daily group, 110 in the nilotinib 400 mg twice daily group and 155 in the imatinib group) due to missing/unevaluable PCR assessments (n=25), atypical transcripts at baseline (n=8) or discontinuation prior to the 72-month time point (n=362).

Figure 1 Cumulative incidence of MMR



For all Sokal risk groups, the MMR rates at all time points remained consistently higher in the two nilotinib groups than in the imatinib group.

In a retrospective analysis, 91% (234/258) of patients on nilotinib 300 mg twice daily achieved BCR-ABL levels \leq 10% at 3 months of treatment compared to 67% (176/264) of patients on imatinib 400 mg once daily. Patients with BCR-ABL levels \leq 10% at 3 months of treatment show a greater overall survival at 72 months compared to those who did not achieve this molecular response level (94.5% vs. 77.1% respectively [p=0.0005]).

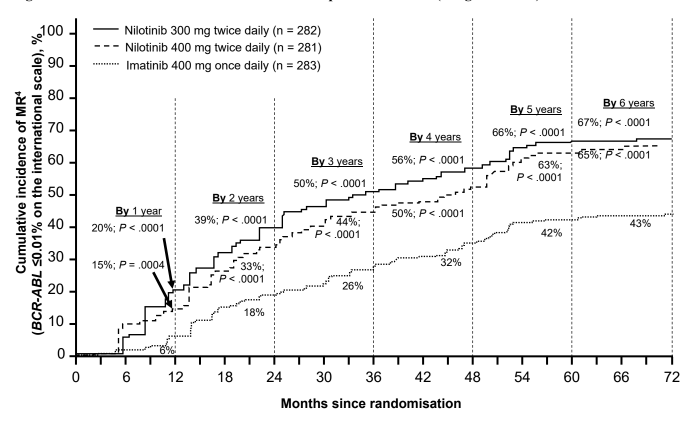
Based on the Kaplan-Meier analysis of time to first MMR the probability of achieving MMR at different time points was higher for both nilotinib at 300 mg and 400 mg twice daily compared to imatinib 400 mg once daily (HR=2.17 and stratified log-rank p<0.0001 between nilotinib 300 mg twice daily and imatinib 400 mg once daily, HR=1.88 and stratified log-rank p<0.0001 between nilotinib 400 mg twice daily and imatinib 400 mg once daily).

The proportion of patients who had a molecular response of $\leq 0.01\%$ and $\leq 0.0032\%$ by IS at different time points are presented in Table 6 and the proportion of patients who had a molecular response of $\leq 0.01\%$ and $\leq 0.0032\%$ by IS by different time points are presented in Figures 2 and 3. Molecular responses of $\leq 0.01\%$ and $\leq 0.0032\%$ by IS correspond to a ≥ 4 log reduction and ≥ 4.5 log reduction, respectively, of BCR-ABL transcripts from a standardised baseline.

Table 6 Proportions of patients who had molecular response of $\leq 0.01\%$ (4 log reduction) and $\leq 0.0032\%$ (4.5 log reduction)

	Nil	otinib	Ni	lotinib	Im	atinib
	300 mg	twice daily	400 mg	twice daily	400 mg	once daily
	n=	=282	n:	=281	n=	=283
		(%)		(%)	((%)
	≤0.01%	≤0.0032%	≤0.01%	≤ 0.0032%	≤0.01%	≤0.0032%
At 12 months	11.7	4.3	8.5	4.6	3.9	0.4
At 24 months	24.5	12.4	22.1	7.8	10.2	2.8
At 36 months	29.4	13.8	23.8	12.1	14.1	8.1
At 48 months	33.0	16.3	29.9	17.1	19.8	10.2
At 60 months	47.9	32.3	43.4	29.5	31.1	19.8
At 72 months	44.3	31.2	45.2	28.8	27.2	18.0

Figure 2 Cumulative incidence of molecular response of ≤0.01% (4-log reduction)



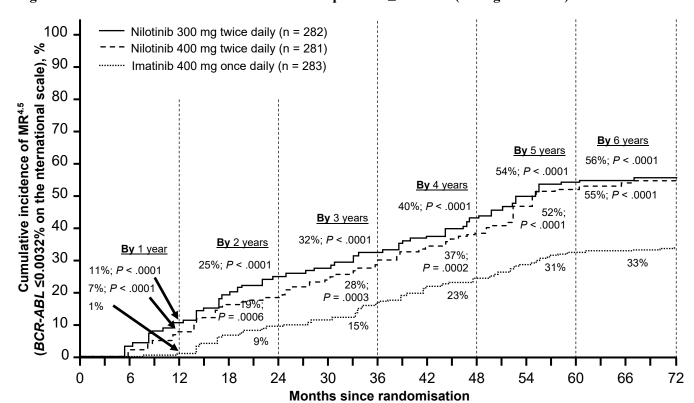


Figure 3 Cumulative incidence of molecular response of ≤0.0032% (4.5 log reduction)

Based on Kaplan-Meier estimates of the duration of first MMR, the proportions of patients who were maintaining response for 72 months among patients who achieved MMR were 92.5% (95% CI: 88.6-96.4%) in the nilotinib 300 mg twice daily group, 92.2% (95% CI: 88.5-95.9%) in the nilotinib 400 mg twice daily group and 88.0% (95% CI: 83.0-93.1%) in the imatinib 400 mg once daily group.

Complete cytogenetic response (CCyR) was defined as 0% Ph+ metaphases in the bone marrow based on a minimum of 20 metaphases evaluated. Best CCyR rate by 12 months (including patients who achieved CCyR at or before the 12 month time point as responders) was statistically higher for both nilotinib 300 mg and 400 mg twice daily compared to imatinib 400 mg once daily, see Table 7.

CCyR rate by 24 months (includes patients who achieved CCyR at or before the 24 month time point as responders) was statistically higher for both the nilotinib 300 mg twice daily and 400 mg twice daily groups compared to the imatinib 400 mg once daily group.

Table 7 Best CCyR rate

	Nilotinib	Nilotinib	Imatinib
	300 mg twice	400 mg twice	400 mg once daily
	daily	daily	n=283
	n=282	n=281	(%)
	(%)	(%)	
By 12 months			
Response (95% CI)	80.1 (75.0; 84.6)	77.9 (72.6; 82.6)	65.0 (59.2; 70.6)
No response	19.9	22.1	35.0
CMH test p-value for response rate	< 0.0001	0.0005	
(versus imatinib 400 mg once			
daily)			
By 24 months			
Response (95% CI)	86.9 (82.4; 90.6)	84.7 (79.9; 88.7)	77.0 (71.7; 81.8)
No response	13.1	15.3	23.0
CMH test p-value for response rate	0.0018	0.0160	
(versus imatinib 400 mg once			
daily)			

Based on Kaplan-Meier estimates, the proportions of patients who were maintaining response for 72 months among patients who achieved CCyR were 99.1% (95% CI: 97.9-100%) in the nilotinib 300 mg twice daily group, 98.7% (95% CI: 97.1-100%) in the nilotinib 400 mg twice daily group and 97.0% (95% CI: 94.7-99.4%) in the imatinib 400 mg once daily group.

Progression to accelerated phase (AP) or blast crisis (BC) on treatment is defined as the time from the date of randomisation to the first documented disease progression to accelerated phase or blast crisis or CML-related death. Progression to accelerated phase or blast crisis on treatment was observed in a total of 17 patients: 2 patients on nilotinib 300 mg twice daily, 3 patients on nilotinib 400 mg twice daily and 12 patients on imatinib 400 mg once daily. The estimated rates of patients free from progression to accelerated phase or blast crisis at 72 months were 99.3%, 98.7% and 95.2%, respectively (HR=0.1599 and stratified log-rank p=0.0059 between nilotinib 300 mg twice daily and imatinib once daily, HR=0.2457 and stratified log-rank p=0.0185 between nilotinib 400 mg twice daily and imatinib once daily). No new events of progression to AP/BC were reported on-treatment since the 2-year analysis.

Including clonal evolution as a criterion for progression, a total of 25 patients progressed to accelerated phase or blast crisis on treatment by the cut-off date (3 in the nilotinib 300 mg twice daily group, 5 in the nilotinib 400 mg twice daily group and 17 in the imatinib 400 mg once daily group). The estimated rates of patients free from progression to accelerated phase or blast crisis including clonal evolution at 72 months were 98.7%, 97.9% and 93.2%, respectively (HR=0.1626 and stratified log-rank p=0.0009 between nilotinib 300 mg twice daily and imatinib once daily, HR=0.2848 and stratified log-rank p=0.0085 between nilotinib 400 mg twice daily and imatinib once daily).

A total of 55 patients died during treatment or during the follow-up after discontinuation of treatment (21 in the nilotinib 300 mg twice daily group, 11 in the nilotinib 400 mg twice daily group and 23 in the imatinib 400 mg once daily group). Twenty-six (26) of these 55 deaths were related to CML (6 in the nilotinib 300 mg twice daily group, 4 in the nilotinib 400 mg twice daily group and 16 in the imatinib 400 mg once daily group). The estimated rates of patients alive at 72 months were 91.6%, 95.8% and 91.4%, respectively (HR=0.8934 and stratified log-rank p=0.7085 between nilotinib 300 mg twice daily and imatinib, HR=0.4632 and stratified log-rank p=0.0314 between nilotinib 400 mg twice daily and imatinib). Considering only CML-related deaths as events, the estimated rates of overall survival at 72 months were 97.7%, 98.5% and 93.9%, respectively (HR=0.3694 and stratified log-rank p=0.0302 between nilotinib 300 mg twice daily and imatinib, HR=0.2433 and stratified log-rank p=0.0061 between nilotinib 400 mg twice daily and imatinib).

Clinical studies in imatinib-resistant or intolerant CML in chronic phase and accelerated phase. An open-label, uncontrolled, multicentre Phase II study was conducted to determine the efficacy of nilotinib in adult patients with imatinib resistant or intolerant CML with separate treatment arms for chronic and accelerated phase disease. Efficacy was based on 321 CP patients and 137 AP patients enrolled. Median duration of treatment was 561 days for CP patients and 264 days for AP patients (see Table 8). Tasigna was administered on a continuous basis (twice daily 2 hours after a meal and with no food for at least one hour after administration) unless there was evidence of inadequate response or disease progression. The dose was 400 mg twice daily and dose escalation to 600 mg twice daily was allowed.

Table 8 Duration of exposure with nilotinib

	Chronic phase n=321	Accelerated phase n=137
Median duration of therapy in days	561	264
(25th-75th percentiles)	(196-852)	(115-595)

Resistance to imatinib included failure to achieve a complete haematological response (by 3 months), cytogenetic response (by 6 months) or major cytogenetic response (by 12 months) or progression of disease after a previous cytogenetic or haematological response. Imatinib intolerance included patients who discontinued imatinib because of toxicity and were not in major cytogenetic response at time of study entry.

Overall, 73% of patients were imatinib-resistant, while 27% were imatinib-intolerant. The majority of patients had a long history of CML that included extensive prior treatment with other antineoplastic agents, including imatinib, hydroxyurea, interferon, and some had even failed organ transplant (Table 9). The median highest prior imatinib dose had been 600 mg/day. The highest prior imatinib dose was \geq 600 mg/day in 74% of all patients, with 40% of patients receiving imatinib doses \geq 800 mg/day.

Table 9 CML disease history characteristics

	Chronic phase	Accelerated phase
	(n=321)	(n=137)*
Median time since diagnosis in months	58	71
(range)	(5–275)	(2–298)
Imatinib		
Resistant	226 (70%)	109 (80%)
Intolerant without MCyR	95 (30%)	27 (20%)
Median time of imatinib treatment in	975	857
days	(519-1,488)	(424-1,497)
(25 th -75 th percentiles)		
Prior hydroxyurea	83%	91%
Prior interferon	58%	50%
Prior bone marrow transplant	7%	8%
* Missing information on imatinib-resista	nt/intolerant status for or	ne patient.

The primary endpoint in the CP patients was major cytogenetic response (MCyR), defined as elimination (CCyR, complete cytogenetic response) or significant reduction to <35% Ph+ metaphases (partial cytogenetic response) of Ph+ haematopoietic cells. Complete haematological response (CHR) in CP patients was evaluated as a secondary endpoint. The primary endpoint in the AP patients was overall confirmed haematological response (HR), defined as either a complete haematological response, no evidence of leukaemia or return to chronic phase.

Chronic phase

The MCyR rate in 321 CP patients was 51%. Most responders achieved their MCyR rapidly within 3 months (median 2.8 months) of starting nilotinib treatment and these were sustained. The median time to achieve CCyR was just past 3 months (median 3.4 months). Of the patients who achieved MCyR, 77% (95% CI: 70% - 84%) were maintaining response at 24 months. Median duration of MCyR has not been reached. Of the patients who achieved CCyR, 85% (95% CI: 78% - 93%) were maintaining response at 24 months. Median duration of CCyR has not been reached. Patients with a CHR at baseline achieved a MCyR faster (1.9 versus 2.8 months). Of CP patients without a baseline CHR, 70% achieved a CHR, median time to CHR was 1 month and median duration of CHR was 32.8 months. The estimated 24-month overall survival rate in CML-CP patients was 87%.

Accelerated phase

The overall confirmed HR rate in 137 AP patients was 50%. Most responders achieved a HR early with nilotinib treatment (median 1.0 months) and these have been durable (median duration of confirmed HR was 24.2 months). Of the patients who achieved HR, 53% (95% CI: 39% - 67%) were maintaining response at 24 months. MCyR rate was 30% with a median time to response of 2.8 months. Of the patients who achieved MCyR, 63% (95% CI: 45% - 80%) were maintaining response at 24 months. Median duration of MCyR was 32.7 months. The estimated 24-month overall survival rate in CML-AP patients was 70%.

The rates of response for the two treatment arms are reported in Table 10.

Table 10 Response in CML

(Best response rate)	Chronic phase			Accelerated phase		
	Intoleran t (n=95)	Resistant (n=226)	Total (n=321)	Intoleran t (n=27)	Resistant (n=109)	Total* (n=137)
Haematological			•			
Response (%)						
Overall (95%CI)	-	-	-	48	51	50
Complete	87	65	70^{1}	(29-68)	(42-61)	(42-59)
NEL	(74-94)	(56-72)	(63-76)	37	28	30
Return to CP	_	-	_	7	10	9
	-	-		4	13	11
Cytogenetic			•			
Response (%)						
Major (95%CI)	57	49	51 (46-57)	33	29	30
Complete	(46-67)	(42-56)	37	(17-54)	(21-39)	(22-38)
Partial	41	35	15	22	19	20
	16	14		11	10	10

NEL = no evidence of leukaemia/marrow response

Efficacy data in patients with CML-BC are not yet available. Separate treatment arms were also included in the Phase II study to investigate Tasigna in a group of CP and AP patients who had been extensively pre-treated with multiple therapies including a tyrosine kinase inhibitor agent in addition to imatinib. Of these patients 30/36 (83%) were treatment resistant not intolerant. In 22 CP patients evaluated for efficacy nilotinib induced a 32% MCyR rate and a 50% CHR rate. In 11 AP patients, evaluated for efficacy, treatment induced a 36% overall HR rate.

After imatinib failure, 24 different BCR-ABL mutations were noted in 42% of chronic phase and 54% of accelerated phase CML patients who were evaluated for mutations. Tasigna demonstrated efficacy

¹ 114 CP patients had a CHR at baseline and were therefore not assessable for complete haematological response

^{*} Missing information on imatinib-resistant/intolerant status for one patient.

in patients harboring a variety of BCR-ABL mutations associated with imatinib resistance, except T315I.

Treatment discontinuation in adult Ph+ CML patients in chronic phase who have been treated with nilotinib as first-line therapy and who have achieved a sustained deep molecular response

In an open-label, single-arm study, 215 adult patients with Ph+ CML in chronic phase treated with nilotinib in first-line for ≥2 years who achieved MR4.5 as measured with the MolecularMD MRDx BCR-ABL test were enrolled to continue nilotinib treatment for additional 52 weeks (nilotinib consolidation phase). 190 of 215 patients (88.4%) entered the TFR phase after achieving a sustained deep molecular response during the consolidation phase, defined by the following criteria:

- the 4 last quarterly assessments (taken every 12 weeks) were at least MR4.0 (BCR-ABL/ABL ≤0.01% IS), and maintained for one year
- the last assessment being MR4.5 (BCR-ABL/ABL ≤0.0032% IS)
- no more than two assessments falling between MR4.0 and MR4.5 (0.0032% IS < BCR-ABL/ABL \leq 0.01% IS).

The primary endpoint was the percentage of patients in MMR at 48 weeks after starting the TFR phase (considering any patient who required re-initiation of treatment as non-responder).

Table 11 Treatment-free remission after nilotinib first-line treatment

Patients entered TFR phase	190		
weeks after starting TFR phase	48 weeks	264 weeks	
patients remaining in MMR or	98 (51.6%, [95% CI: 44.2,	79 ^[2] (41.6%, 95% CI: 34.5,	
better	58.9])	48.9)	
Patients discontinued TFR phase	93 [1]	109	
due to loss of MMR	88 (46.3%)	94 (49.5%)	
due to other reasons	5	15	
Patients restarted treatment after loss of	86	91	
MMR			
regaining MMR	85 (98.8%)	90 (98.9%)	
regaining MR4.5	76 (88.4%)	84 (92,3%)	

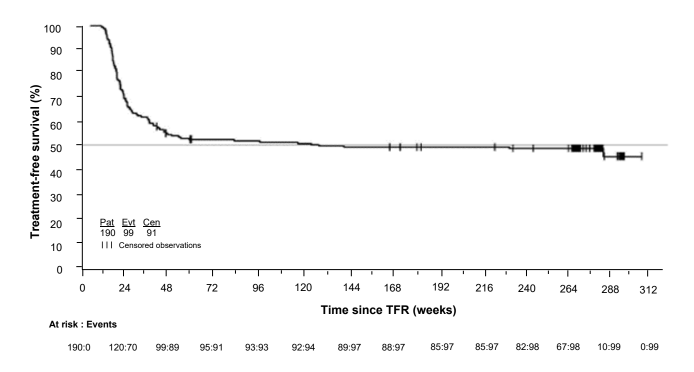
^[1] One patient did not lose MMR by week 48 but discontinued TFR phase.

The time by which 50% of all retreated patients regained MMR and MR4.5 was 7 and 12.9 weeks, respectively. The cumulative rate of MMR regained at 24 weeks after treatment re-initiation was 97.8% (89/91 patients) and MR4.5 regained at 48 weeks was 91.2% (83/91 patients).

The Kaplan-Meier estimate of median treatment-free survival (TFS) was 120.1 weeks (95% CI: 36.9, not estimable [NE]) (Figure 4); 91 of 190 patients (47.9%) did not have a TFS event.

^[2] For 2 patients, PCR assessment was not available at week 264 therefore their response was not considered for the week 264 data cut-off analysis.

Figure 4 Kaplan-Meier estimate of treatment-free survival after start of TFR (full analysis set)



<u>Treatment discontinuation in adult CML patients in chronic phase who have achieved a sustained deep molecular response on nilotinib treatment following prior imatinib therapy</u>

In an open-label, single-arm study, 163 adult patients with Ph+ CML in chronic phase taking tyrosine kinase inhibitors (TKIs) for ≥3 years (imatinib as initial TKI therapy for more than 4 weeks without documented MR4.5 on imatinib at the time of switch to nilotinib, then switched to nilotinib for at least two years), and who achieved MR4.5 on nilotinib treatment as measured with the MolecularMD MRDx BCR-ABL test were enrolled to continue nilotinib treatment for additional 52 weeks (nilotinib consolidation phase). 126 of 163 patients (77.3%) entered the TFR phase after achieving a sustained deep molecular response during the consolidation phase, defined by the following criterion:

- The 4 last quarterly assessments (taken every 12 weeks) showed no confirmed loss of MR4.5 (BCR-ABL/ABL ≤0.0032% IS) during one year.

The primary endpoint was the proportion of patients without confirmed loss of MR4.0 or loss of MMR within 48 weeks following treatment discontinuation.

Table 12 Treatment-free remission after nilotinib treatment following prior imatinib therapy

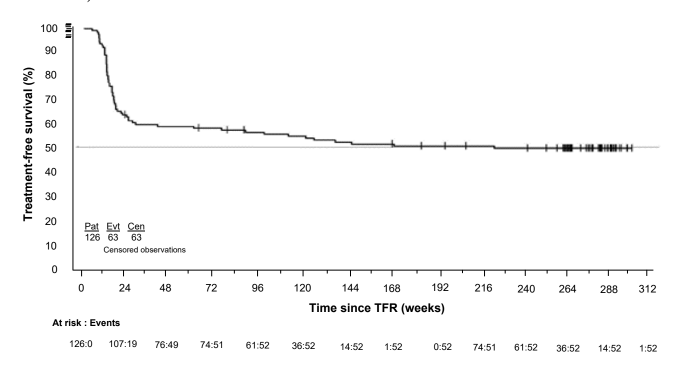
Patients entered TFR phase	126		
weeks after starting TFR phase	48 weeks	264 weeks	
patients remaining in MMR, no	73 (57.9%, [95% CI: 48.8,	54 (42.9% [54/126, 95%	
confirmed loss of MR4.0, and no	66.7])	CI: 34.1, 52.0])	
re-initiation of nilotinib	·		
Patients discontinued TFR Phase	53	74 [1]	
due to confirmed loss of MR4.0 or	53 (42.1%)	61 (82.4%)	
loss of MMR			
due to other reasons	0	13	
Patients restarted treatment after loss of	51	59	
MMR or confirmed loss of MR4.0			
regaining MR4.0	48 (94.1%)	56 (94.9%)	
regaining MR4.5	47 (92.2%)	54 (91.5%)	
517	A + 264 1 1 + 11	. 11 . 11 1	

[1] two patients had MMR (PCR assessment) at 264 weeks but were discontinued later and had no further PCR assessment.

The Kaplan-Meier estimated median time on nilotinib to regain MR4.0 and MR4.5 was 11.1 weeks (95% CI:8.1, 12.1) and 13.1 weeks (95% CI:12.0, 15.9), respectively. The cumulative rate of MR4 and MR4.5 regained by 48 weeks after treatment re-initiation was 94.9% (56/59 patients) and 91.5% (54/59 patients), respectively.

The median TFS Kaplan-Meier estimate is 224 weeks (95% CI: 39.9, NE) (Figure 5); 63 of 126 patients (50.0%) did not have a TFS event.

Figure 5 Kaplan-Meier estimate of treatment-free survival after start of TFR (full analysis set)



Paediatric population

In the main paediatric study conducted with nilotinib, a total of 58 patients from 2 to <18 years of age (25 patients newly diagnosed Ph+ CML in chronic phase and 33 patients imatinib/dasatinib-resistant or imatinib-intolerant Ph+ CML in chronic phase) received nilotinib treatment at a dose of 230 mg/m² twice daily, rounded to the nearest 50 mg dose (to a maximum single dose of 400 mg). Key study data are summarised in Table 13.

Table 13 Summary data for the main paediatric study conducted with nilotinib

	Newly diagnosed Ph+	resistant or intolerant Ph+
	CML-CP	CML-CP
	(n=25)	(n=33)
Median time on treatment in	51.9 (1.4 - 61.2)	60.5 (0.7 - 63.5)
month, (range)		
Median (range) actual dose	377.0 (149 - 468)	436.9 (196 - 493)
intensity (mg/m²/day)		
Relative dose intensity (%)		
compared to the planned dose		
of 230 mg/m ² twice daily		
Median (range)	82.0 (32-102)	95.0 (43-107)
Number of patients with	12 (48.0%)	19 (57.6%)
>90%		
MMR (BCR-ABL/ABL	60%, (38.7, 78.9)	48.5%, (30.8, 66.5)
\leq 0.1%) IS at 12 cycles, (95%		
CI)		
MMR by cycle 12, (95% CI)	64.0%, (42.5, 82.0)	57.6%, (39.2, 74.5)
MMR by cycle 66, (95% CI)	76.0%, (54.9, 90.6)	60.6%, (42.1, 77.1)
Median time to MMR in month	5.56 (5.52, 10.84)	2.79 (0.03, 5.75)
(95% CI)		
No. of patients (%) achieved	14 (56.0%)	9 (27.3%)
MR4.0 (BCR-ABL/ABL		
≤0.01% IS) by cycle 66		
No. of patients (%) achieved	11 (44.0%)	4 (12.1%)
MR4.5 (BCR-ABL/ABL		
\leq 0.0032% IS) by cycle 66		
Confirmed loss of MMR	3 out of 19	None out of 20
among patients who achieved		
MMR		
Emergent mutation while on	None	None
treatment		
Disease progression while on	1 patient temporarily matched	1 patient progressed to AP/BC
treatment	the technical definition for	after 10.1 months on treatment
	progression to AP/BC *	
Overall survival		
No. of events	0	0
Death on treatment	3 (12%)	1 (3%)
Death during survival	Not estimable	Not estimable
follow up	1.4 1 1 . 1 . 6	

^{*} one patient temporarily matched the technical definition for progression to AP/BC (due to increased basophil cell count), one month after the start of nilotinib (with temporary treatment interruption of 13 days during first cycle). The patient remained in the study, went back to CP and was in CHR and CCyR by 6 cycles of nilotinib treatment.

5.2 Pharmacokinetic properties

Absorption

Peak concentrations of nilotinib are reached 3 hours after oral administration. Nilotinib absorption following oral administration was approximately 30%. The absolute bioavailability of nilotinib has not been determined. As compared to an oral drink solution (pH of 1.2 to 1.3), relative bioavailability of nilotinib capsule is approximately 50%. In healthy volunteers, C_{max} and area under the serum concentration-time curve (AUC) of nilotinib are increased by 112% and 82%, respectively, compared to fasting conditions when Tasigna is given with food. Administration of Tasigna 30 minutes or

2 hours after food increased bioavailability of nilotinib by 29% or 15%, respectively (see sections 4.2, 4.4 and 4.5).

Nilotinib absorption (relative bioavailability) might be reduced by approximately 48% and 22% in patients with total gastrectomy and partial gastrectomy, respectively.

Distribution

The blood-to-plasma ratio of nilotinib is 0.71. Plasma protein binding is approximately 98% on the basis of *in vitro* experiments.

Biotransformation

Main metabolic pathways identified in healthy subjects are oxidation and hydroxylation. Nilotinib is the main circulating component in the serum. None of the metabolites contribute significantly to the pharmacological activity of nilotinib. Nilotinib is primarily metabolised by CYP3A4, with possible minor contribution from CYP2C8.

Elimination

After a single dose of radiolabelled nilotinib in healthy subjects, more than 90% of the dose was eliminated within 7 days, mainly in faeces (94% of the dose). Unchanged nilotinib accounted for 69% of the dose.

The apparent elimination half-life estimated from the multiple-dose pharmacokinetics with daily dosing was approximately 17 hours. Inter-patient variability in nilotinib pharmacokinetics was moderate to high.

Linearity/non-linearity

Steady-state nilotinib exposure was dose-dependent, with less than dose-proportional increases in systemic exposure at dose levels higher than 400 mg given as once-daily dosing. Daily systemic exposure to nilotinib with 400 mg twice-daily dosing at steady state was 35% higher than with 800 mg once-daily dosing. Systemic exposure (AUC) of nilotinib at steady state at a dose level of 400 mg twice daily was approximately 13.4% higher than at a dose level of 300 mg twice daily. The average nilotinib trough and peak concentrations over 12 months were approximately 15.7% and 14.8% higher following 400 mg twice-daily dosing compared to 300 mg twice daily. There was no relevant increase in exposure to nilotinib when the dose was increased from 400 mg twice daily to 600 mg twice daily.

Steady-state conditions were essentially achieved by day 8. An increase in serum exposure to nilotinib between the first dose and steady state was approximately 2-fold for daily dosing and 3.8-fold for twice-daily dosing.

Bioavailability/bioequivalence studies

Single-dose administration of 400 mg nilotinib, using 2 hard capsules of 200 mg whereby the content of each hard capsule was dispersed in one teaspoon of apple sauce, was shown to be bioequivalent with a single-dose administration of 2 intact hard capsules of 200 mg.

Paediatric population

Following administration of nilotinib in paediatric patients at 230 mg/m² twice daily, rounded to the nearest 50 mg dose (to a maximum single dose of 400 mg), steady-state exposure and clearance of nilotinib were found to be similar (within 2-fold) to adult patients treated with 400 mg twice daily. The pharmacokinetic exposure of nilotinib following a single or multiple doses appeared to be comparable between paediatric patients from 2 years to <10 years and from \geq 10 years to <18 years.

5.3 Preclinical safety data

Nilotinib has been evaluated in safety pharmacology, repeated-dose toxicity, genotoxicity, reproductive toxicity, phototoxicity and carcinogenicity (rats and mice) studies.

Safety pharmacology studies

Nilotinib did not have effects on CNS or respiratory functions. *In vitro* cardiac safety studies demonstrated a preclinical signal for QT prolongation, based upon block of hERG currents and prolongation of the action potential duration in isolated rabbit hearts by nilotinib. No effects were seen in ECG measurements in dogs or monkeys treated for up to 39 weeks or in a special telemetry study in dogs.

Repeated-dose toxicity studies

Repeated-dose toxicity studies in dogs of up to 4 weeks' duration and in cynomolgus monkeys of up to 9 months' duration revealed the liver as the primary target organ of toxicity of nilotinib. Alterations included increased alanine aminotransferase and alkaline phosphatase activity and histopathology findings (mainly sinusoidal cell or Kupffer cell hyperplasia/hypertrophy, bile duct hyperplasia and periportal fibrosis). In general the changes in clinical chemistry were fully reversible after a four-week recovery period and the histological alterations showed partial reversibility. Exposures at the lowest dose levels at which the liver effects were seen were lower than the exposure in humans at a dose of 800 mg/day. Only minor liver alterations were seen in mice or rats treated for up to 26 weeks. Mainly reversible increases in cholesterol levels were seen in rats, dogs and monkeys.

Genotoxicity studies

Genotoxicity studies in bacterial *in vitro* systems and in mammalian *in vitro* and *in vivo* systems with and without metabolic activation did not reveal any evidence for a mutagenic potential of nilotinib.

Carcinogenicity studies

In the 2-year rat carcinogenicity study, the major target organ for non-neoplastic lesions was the uterus (dilatation, vascular ectasia, endothelial cell hyperplasia, inflammation and/or epithelial hyperplasia). There was no evidence of carcinogenicity upon administration of nilotinib at 5, 15 and 40 mg/kg/day. Exposures (in terms of AUC) at the highest dose level represented approximately 2x to 3x human daily steady-state exposure (based on AUC) to nilotinib at the dose of 800 mg/day.

In the 26-week Tg.rasH2 mouse carcinogenicity study, in which nilotinib was administered at 30, 100 and 300 mg/kg/day, skin papillomas/carcinomas were detected at 300 mg/kg, representing approximately 30 to 40 times (based on AUC) the human exposure at the maximum approved dose of 800 mg/day (administered as 400 mg twice daily). The No-Observed-Effect-Level for the skin neoplastic lesions was 100 mg/kg/day, representing approximately 10 to 20 times the human exposure at the maximum approved dose of 800 mg/day (administered as 400 mg twice daily). The major target organs for non-neoplastic lesions were the skin (epidermal hyperplasia), the growing teeth (degeneration/atrophy of the enamel organ of upper incisors and inflammation of the gingiva/odontogenic epithelium of incisors) and the thymus (increased incidence and/or severity of decreased lymphocytes).

Reproductive toxicity and fertility studies

Nilotinib did not induce teratogenicity, but did show embryo- and foetotoxicity at doses that also showed maternal toxicity. Increased post-implantation loss was observed in both the fertility study, which involved treatment of both males and females, and the embryotoxicity study, which involved treatment of females. Embryo-lethality and foetal effects (mainly decreased foetal weights, premature fusion of the facial bones (fused maxilla/zygomatic) visceral and skeletal variations) in rats and increased resorption of foetuses and skeletal variations in rabbits were present in the embryotoxicity

studies. In a pre- and postnatal development study in rats, maternal exposure to nilotinib caused reduced pup body weight with associated changes in physical development parameters as well as reduced mating and fertility indices in the offspring. Exposure to nilotinib in females at No-Observed-Adverse-Effect-Levels was generally less or equal to that in humans at 800 mg/day.

No effects on sperm count/motility or on fertility were noted in male and female rats up to the highest tested dose, approximately 5 times the recommended dosage for humans.

Juvenile animal studies

In a juvenile development study, nilotinib was administered via oral gavage to juvenile rats from the first week post partum through young adult (day 70 post partum) at doses of 2, 6 and 20 mg/kg/day. Besides standard study parameters, evaluations of developmental landmarks, CNS effects, mating and fertility were performed. Based on a reduction in body weight in both genders and a delayed preputial separation in males (which may be associated with the reduction in weight), the No-Observed-Effect-Level in juvenile rats was considered to be 6 mg/kg/day. The juvenile animals did not exert increased sensitivity to nilotinib relative to adults. In addition, the toxicity profile in juvenile rats was comparable to that observed in adult rats.

Phototoxicity studies

Nilotinib was shown to absorb light in the UV-B and UV-A range, is distributed into the skin and showed a phototoxic potential *in vitro*, but no effects have been observed *in vivo*. Therefore the risk that nilotinib causes photosensitisation in patients is considered very low.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Tasigna 50 mg hard capsules

<u>Capsule content</u>
Lactose monohydrate
Crospovidone Type A
Poloxamer 188
Colloidal anhydrous silica
Magnesium stearate

<u>Capsule shell</u>
Gelatin
Titanium dioxide (E171)
Red iron oxide (E172)
Yellow iron oxide (E172)

Printing ink
Shellac
Black iron oxide (E172)
Propylene glycol
Ammonium hydroxide

Tasigna 150 mg hard capsules

Capsule content

Lactose monohydrate Crospovidone Type A Poloxamer 188 Colloidal anhydrous silica Magnesium stearate

Capsule shell

Gelatin
Titanium dioxide (E171)
Red iron oxide (E172)
Yellow iron oxide (E172)

Printing ink

Shellac
Black iron oxide (E172)
n-Butyl alcohol
Propylene glycol
Dehydrated ethanol
Isopropyl alcohol
Ammonium hydroxide

Tasigna 200 mg hard capsules

Capsule content

Lactose monohydrate Crospovidone Type A Poloxamer 188 Colloidal anhydrous silica Magnesium stearate

Capsule shell

Gelatin
Titanium dioxide (E171)
Yellow iron oxide (E172)

Printing ink

Shellac (E904)
Dehydrated alcohol
Isopropyl alcohol
Butyl alcohol
Propylene glycol
Strong ammonia solution
Potassium hydroxide
Red iron oxide (E172)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years.

6.4 Special precautions for storage

Do not store above 30°C.

Store in the original package in order to protect from moisture.

6.5 Nature and contents of container

Tasigna is available in the following pack sizes:

Tasigna 50 mg hard capsules

PVC/PVDC/Alu blisters

Pack containing 120 (3 packs of 40) hard capsules.

Tasigna 150 mg hard capsules

PVC/PVDC/Alu blisters

- Unit packs containing 28 hard capsules (7 daily blisters, each containing 4 hard capsules) or 40 hard capsules (5 blisters, each containing 8 hard capsules).
- Multipacks containing 112 (4 packs of 28) hard capsules, 120 (3 packs of 40) hard capsules or 392 (14 packs of 28) hard capsules.

Not all pack sizes may be marketed.

Tasigna 200 mg hard capsules

PVC/PVDC/Alu blisters

- Unit packs containing 28 hard capsules in a wallet.
- Unit packs containing 28 hard capsules (7 daily blisters, each containing 4 hard capsules) or 40 hard capsules (5 blisters, each containing 8 hard capsules).
- Multipacks containing 112 (4 wallets of 28) hard capsules.
- Multipacks containing 112 (4 packs of 28) hard capsules, 120 (3 packs of 40) hard capsules or 392 (14 packs of 28) hard capsules.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORISATION HOLDER

Novartis Europharm Limited Vista Building Elm Park, Merrion Road Dublin 4 Ireland

8. MARKETING AUTHORISATION NUMBER(S)

Tasigna 50 mg hard capsules

EU/1/07/422/015

Tasigna 150 mg hard capsules

EU/1/07/422/005-006 EU/1/07/422/009-010 EU/1/07/422/013

Tasigna 200 mg hard capsules

EU/1/07/422/001 EU/1/07/422/003 EU/1/07/422/007-008 EU/1/07/422/011-012 EU/1/07/422/014

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 19 November 2007 Date of latest renewal: 20 September 2012

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu

ANNEX II

- A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE
- B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE
- C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION
- D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

50 mg hard capsules

Novartis Pharmaceutical Manufacturing LLC Verovškova ulica 57 1000 Ljubljana Slovenia

Novartis Farmacéutica SA Gran Via de les Corts Catalanes, 764 08013 Barcelona Spain

Lek d.d., PE PROIZVODNJA LENDAVA Trimlini 2D Lendava, 9220 Slovenia

Novartis Pharma GmbH Sophie-Germain-Strasse 10 90443 Nuremberg Germany

150 mg hard capsules and 200 mg hard capsules

Novartis Pharmaceutical Manufacturing LLC Verovškova ulica 57 1000 Ljubljana Slovenia

Lek d.d., PE PROIZVODNJA LENDAVA Trimlini 2D Lendava, 9220 Slovenia

Novartis Pharma GmbH Sophie-Germain-Strasse 10 90443 Nuremberg Germany

The printed package leaflet of the medicinal product must state the name and address of the manufacturer responsible for the release of the concerned batch.

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to restricted medical prescription (See Annex I: Summary of Product Characteristics, section 4.2).

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

The requirements for submission of PSURs for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

• Risk management plan (RMP)

The marketing authorisation holder (MAH) shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the marketing authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:

- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached.

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
OUTER CARTON
1. NAME OF THE MEDICINAL PRODUCT
Tasigna 50 mg hard capsules nilotinib
2. STATEMENT OF ACTIVE SUBSTANCE(S)
One hard capsule contains 50 mg nilotinib (as hydrochloride monohydrate).
3. LIST OF EXCIPIENTS
Contains lactose – see the package leaflet for further information.
4. PHARMACEUTICAL FORM AND CONTENTS
Hard capsule
120 (3 packs of 40) hard capsules.
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use. Oral use.
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
8. EXPIRY DATE EXP

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10.	OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vista	
12.	MARKETING AUTHORISATION NUMBER(S)
EU/	1/07/422/015 120 hard capsules
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Tasig	gna 50 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA
PC SN NN	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING		
INTERMEDIATE CARTON		
1. NAME OF THE MEDICINAL PRODUCT		
Tasigna 50 mg hard capsules nilotinib		
2. STATEMENT OF ACTIVE SUBSTANCE(S)		
One hard capsule contains 50 mg nilotinib (as hydrochloride monohydrate).		
3. LIST OF EXCIPIENTS		
Contains lactose – see the package leaflet for further information.		
4. PHARMACEUTICAL FORM AND CONTENTS		
Hard capsule		
40 hard capsules. Not to be sold separately.		
5. METHOD AND ROUTE(S) OF ADMINISTRATION		
Read the package leaflet before use. Oral use.		
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN		
Keep out of the sight and reach of children.		
7. OTHER SPECIAL WARNING(S), IF NECESSARY		
8. EXPIRY DATE		
EXP		
9. SPECIAL STORAGE CONDITIONS		

Do not store above 30°C.

Store in the original package in order to protect from moisture.

	APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vista	
12.	MARKETING AUTHORISATION NUMBER(S)
EU/	1/07/422/015 120 hard capsules
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Tasig	na 50 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA

SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
BLISTERS	
1. NAME OF THE MEDICINAL PRODUCT	
Tasigna 50 mg hard capsules nilotinib	
2. NAME OF THE MARKETING AUTHORISATION HOLDER	
Novartis Europharm Limited	
3. EXPIRY DATE	
EXP	
4. BATCH NUMBER	
Lot	
5. OTHER	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING	
CARTON OF UNIT PACK	
1. NAME OF THE MEDICINAL PRODUCT	
Tasigna 150 mg hard capsules nilotinib	
2. STATEMENT OF ACTIVE SUBSTANCE(S)	
One hard capsule contains 150 mg nilotinib (as hydrochloride monohydrate).	
3. LIST OF EXCIPIENTS	
Contains lactose – see the package leaflet for further information.	
4. PHARMACEUTICAL FORM AND CONTENTS	
Hard capsule	
28 hard capsules 40 hard capsules	
5. METHOD AND ROUTE(S) OF ADMINISTRATION	
Read the package leaflet before use. Oral use.	
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN	
Keep out of the sight and reach of children.	
7. OTHER SPECIAL WARNING(S), IF NECESSARY	
8. EXPIRY DATE	
EXP	
9. SPECIAL STORAGE CONDITIONS	

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vista	
12.	MARKETING AUTHORISATION NUMBER(S)
	/1/07/422/005 28 hard capsules /1/07/422/009 40 hard capsules
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Tasig	gna 150 mg
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D b	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER - HUMAN READABLE DATA
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MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
BLISTERS	
1. NAME OF THE MEDICINAL PRODUCT	
Tasigna 150 mg hard capsules nilotinib	
2. NAME OF THE MARKETING AUTHORISATION HOLDER	
Novartis Europharm Limited	
3. EXPIRY DATE	
EXP	
4. BATCH NUMBER	
Lot	
5 OTHER	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

CARTON OF MULTIPACK (INCLUDING BLUE BOX)

1. NAME OF THE MEDICINAL PRODUCT

Tasigna 150 mg hard capsules nilotinib

2. STATEMENT OF ACTIVE SUBSTANCE(S)

One hard capsule contains 150 mg nilotinib (as hydrochloride monohydrate).

3. LIST OF EXCIPIENTS

Contains lactose – see the package leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Hard capsule

Multipack: 112 (4 packs of 28) hard capsules. Multipack: 120 (3 packs of 40) hard capsules. Multipack: 392 (14 packs of 28) hard capsules.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use. Oral use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP

9. SPECIAL STORAGE CONDITIONS

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Novartis Europharm Limited Vista Building Elm Park, Merrion Road Dublin 4 Ireland

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/07/422/006	112 hard capsules
EU/1/07/422/010	120 hard capsules
EU/1/07/422/013	392 hard capsules

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Tasigna 150 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC

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PARTICULARS TO APPEAR ON THE OUTER PACKAGING INTERMEDIATE CARTON OF MULTIPACK (WITHOUT BLUE BOX) 1. NAME OF THE MEDICINAL PRODUCT Tasigna 150 mg hard capsules nilotinib 2. STATEMENT OF ACTIVE SUBSTANCE(S) One hard capsule contains 150 mg nilotinib (as hydrochloride monohydrate). 3. LIST OF EXCIPIENTS Contains lactose – see the package leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS Hard capsule 28 hard capsules. Component of a multipack. Not to be sold separately. 40 hard capsules. Component of a multipack. Not to be sold separately. 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT 6. OF THE SIGHT AND REACH OF CHILDREN Keep out of the sight and reach of children. 7. OTHER SPECIAL WARNING(S), IF NECESSARY 8. **EXPIRY DATE EXP**

9. SPECIAL STORAGE CONDITIONS

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10.		FIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS RIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF
11.	NAME AND ADDRI	ESS OF THE MARKETING AUTHORISATION HOLDER
Vista		
12.	MARKETING AUTI	HORISATION NUMBER(S)
EU/	1/07/422/006 1/07/422/010 1/07/422/013	112 hard capsules 120 hard capsules 392 hard capsules
13.	BATCH NUMBER	
Lot		
14.	GENERAL CLASSI	FICATION FOR SUPPLY
15.	INSTRUCTIONS OF	NUSE
16.	INFORMATION IN	BRAILLE
Tasig	na 150 mg	

17.

UNIQUE IDENTIFIER – 2D BARCODE

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PARTICULARS TO APPEAR ON THE OUTER PACKAGING	
CARTON OF UNIT PACK (WALLET) CARTON OF UNIT PACK (CARTON)	
1. NAME OF THE MEDICINAL PRODUCT	
Tasigna 200 mg hard capsules nilotinib	
2. STATEMENT OF ACTIVE SUBSTANCE(S)	
One hard capsule contains 200 mg nilotinib (as hydrochloride monohydrate).	
3. LIST OF EXCIPIENTS	
Contains lactose – see the package leaflet for further information.	
4. PHARMACEUTICAL FORM AND CONTENTS	
Hard capsule 28 hard capsules 40 hard capsules	
5. METHOD AND ROUTE(S) OF ADMINISTRATION	
Read the package leaflet before use. Oral use.	
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN	
Keep out of the sight and reach of children.	
7. OTHER SPECIAL WARNING(S), IF NECESSARY	
8. EXPIRY DATE	
EXP	

9. SPECIAL STORAGE CONDITIONS

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Novartis Europharm Limited Vista Building Elm Park, Merrion Road Dublin 4 Ireland

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/07/422/001	PVC/PVDC/Alu [in wallet] 28 hard capsules
EU/1/07/422/007	PVC/PVDC/Alu [in carton] 28 hard capsules
EU/1/07/422/011	PVC/PVDC/Alu [in carton] 40 hard capsules

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Tasigna 200 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC

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MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS			
BLISTERS			
1. NAME OF THE MEDICINAL PRODUCT			
Tasigna 200 mg hard capsules nilotinib			
2. NAME OF THE MARKETING AUTHORISATION HOLDER			
Novartis Europharm Limited			
3. EXPIRY DATE			
EXP			
4. BATCH NUMBER			
Lot			
5. OTHER			

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

CARTON OF MULTIPACK (WALLET) (INCLUDING BLUE BOX) CARTON OF MULTIPACK (CARTON) (INCLUDING BLUE BOX)

1. NAME OF THE MEDICINAL PRODUCT

Tasigna 200 mg hard capsules nilotinib

2. STATEMENT OF ACTIVE SUBSTANCE(S)

One hard capsule contains 200 mg nilotinib (as hydrochloride monohydrate).

3. LIST OF EXCIPIENTS

Contains lactose – see the package leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Hard capsule

Multipack: 112 (4 wallets of 28) hard capsules. Multipack: 112 (4 packs of 28) hard capsules. Multipack: 120 (3 packs of 40) hard capsules. Multipack: 392 (14 packs of 28) hard capsules.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use. Oral use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP

9. SPECIAL STORAGE CONDITIONS

Do not store above 30°C.

Store in the original package in order to protect from moisture.

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Novartis Europharm Limited Vista Building Elm Park, Merrion Road Dublin 4 Ireland

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/07/422/003	PVC/PVDC/Alu [in wallet] 112 hard capsules
EU/1/07/422/008	PVC/PVDC/Alu [in carton] 112 hard capsules
EU/1/07/422/012	PVC/PVDC/Alu [in carton] 120 hard capsules
EU/1/07/422/014	PVC/PVDC/Alu [in carton] 392 hard capsules

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Tasigna 200 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

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PARTICULARS TO APPEAR ON THE OUTER PACKAGING

INTERMEDIATE WALLET OF MULTIPACK (WITHOUT BLUE BOX) INTERMEDIATE CARTON OF MULTIPACK (WITHOUT BLUE BOX)

1. NAME OF THE MEDICINAL PRODUCT

Tasigna 200 mg hard capsules nilotinib

2. STATEMENT OF ACTIVE SUBSTANCE(S)

One hard capsule contains 200 mg nilotinib (as hydrochloride monohydrate).

3. LIST OF EXCIPIENTS

Contains lactose – see the package leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Hard capsule

- 28 hard capsules. Component of a multipack comprising 4 wallets. Not to be sold separately.
- 28 hard capsules. Component of a multipack comprising 4 cartons. Not to be sold separately.
- 40 hard capsules. Component of a multipack comprising 3 cartons. Not to be sold separately.
- 28 hard capsules. Component of a multipack comprising 14 cartons. Not to be sold separately.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use. Oral use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP

9.	SPECIAL STORAGE CONDITIONS			
	ot store above 30°C in the original pac	C. kage in order to protect from moisture.		
10.	0. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE			
11.	NAME AND AD	DRESS OF THE MARKETING AUTHORISATION HOLDER		
Nova	artis Europharm Lir	mited		
	Building	inica		
	Park, Merrion Road	4		
Dubl				
Irela				
11 0 100				
12.	MARKETING A	AUTHORISATION NUMBER(S)		
EII	/1/07/422/003	PVC/PVDC/Alu [in wallet] 112 hard capsules		
	/1/07/422/008	PVC/PVDC/Alu [in carton] 112 hard capsules		
	/1/07/422/012	PVC/PVDC/Alu [in carton] 112 hard capsules		
	/1/07/422/014	PVC/PVDC/Alu [in carton] 120 hard capsules		
ĽU	1/0//422/014	1 VC/1 VDC/Atu [iii carton] 392 hard capsules		
13.	BATCH NUMB	FD		
15.	DATCH NUMB.	EK		
Lot				
Lot				
14.	GENERAL CLA	ASSIFICATION FOR SUPPLY		
15.	INSTRUCTION	S ON USE		
16.	INFORMATION	N IN BRAILLE		
Tasio	gna 200 mg			
- 3018	- · · · · · · · · · · · · · · · · · · ·			

17.

UNIQUE IDENTIFIER – 2D BARCODE

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

B. PACKAGE LEAFLET

Package leaflet: Information for the user

Tasigna 50 mg, 150 mg and 200 mg hard capsules nilotinib

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.

- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor or pharmacist.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet

- 1. What Tasigna is and what it is used for
- 2. What you need to know before you take Tasigna
- 3. How to take Tasigna
- 4. Possible side effects
- 5. How to store Tasigna
- 6. Contents of the pack and other information

1. What Tasigna is and what it is used for

What Tasigna is

Tasigna is a medicine containing an active substance called nilotinib.

What Tasigna is used for

Tasigna is used to treat a type of leukaemia called Philadelphia chromosome positive chronic myeloid leukaemia (Ph-positive CML). CML is a cancer of the blood which makes the body produce too many abnormal white blood cells.

Tasigna is used in adult and paediatric patients with newly diagnosed CML or in patients with CML who are no longer benefiting from previous treatment including imatinib. It is also used in adult and paediatric patients who experienced serious side effects with previous treatment and are not able to continue taking it.

How Tasigna works

In patients with CML, a change in DNA (genetic material) triggers a signal that tells the body to produce abnormal white blood cells. Tasigna blocks this signal, and thus stops the production of these cells.

Monitoring during Tasigna treatment

Regular tests, including blood tests, will be performed during treatment. These tests will monitor:

- the amount of blood cells (white blood cells, red blood cells and platelets) in the body to see how Tasigna is tolerated.
- pancreas and liver function in the body to see how Tasigna is tolerated.
- the electrolytes in the body (potassium, magnesium). These are important in the functioning of the heart.
- the level of sugar and fats in the blood.

The heart rate will also be checked using a machine that measures electrical activity of the heart (a test called an "ECG").

Your doctor will regularly evaluate your treatment and decide whether you should continue to take Tasigna. If you are told to discontinue this medicine, your doctor will continue to monitor your CML and may tell you to re-start Tasigna if your condition indicates that this is necessary.

If you have any questions about how Tasigna works or why it has been prescribed for you or your child, ask your doctor.

2. What you need to know before you take Tasigna

Follow all the doctor's instructions carefully. They may differ from the general information contained in this leaflet.

Do not take Tasigna

- if you are allergic to nilotinib or any of the other ingredients of this medicine (listed in section 6).

If you think you may be allergic, tell your doctor before taking Tasigna.

Warnings and precautions

Talk to your doctor or pharmacist before taking Tasigna:

- if you have suffered prior cardiovascular events such as a heart attack, chest pain (angina), problems with the blood supply to your brain (stroke) or problems with the blood flow to your leg (claudication) or if you have risk factors for cardiovascular disease such as high blood pressure (hypertension), diabetes or problems with the level of fats in your blood (lipid disorders).
- if you have a **heart disorder**, such as an abnormal electrical signal called "prolongation of the OT interval".
- if you are being **treated with medicines** that lower your blood cholesterol (statins), or affect the heartbeat (anti-arrhythmics) or the liver (see **Other medicines and Tasigna**).
- if you suffer from lack of potassium or magnesium.
- if you have a liver or pancreas disorder.
- if you have symptoms such as easy bruising, feeling tired or short of breath or have experienced repeated infections.
- if you have had a surgical procedure involving the removal of the entire stomach (total gastrectomy).
- if you have ever had or might now have a hepatitis B infection. This is because Tasigna could cause hepatitis B to become active again, which can be fatal in some cases. Patients will be carefully checked by their doctor for signs of this infection before treatment is started.

If any of these apply to you or your child, tell your doctor.

During treatment with Tasigna

- if you faint (loss of consciousness) or have an irregular heart beat while taking this medicine, tell your doctor immediately as this may be a sign of a serious heart condition. Prolongation of the QT interval or an irregular heart beat may lead to sudden death. Uncommon cases of sudden death have been reported in patients taking Tasigna.
- if you have sudden heart palpitations, severe muscle weakness or paralysis, seizures or sudden changes in your thinking or level of alertness, **tell your doctor immediately** as this may be a sign of a fast breakdown of cancer cells called tumour lysis syndrome. Rare cases of tumour lysis syndrome have been reported in patients treated with Tasigna.
- if you develop chest pain or discomfort, numbness or weakness, problems with walking or with your speech, pain, discolouration or a cool feeling in a limb, **tell your doctor immediately** as this may be a sign of a cardiovascular event. Serious cardiovascular events including problems with the blood flow to the leg (peripheral arterial occlusive disease), ischaemic heart disease and problems with the blood supply to the brain (ischaemic cerebrovascular disease) have been reported in patients taking Tasigna. Your doctor should assess the level of fats (lipids) and sugar in your blood before initiating treatment with Tasigna and during treatment.

- if you develop swelling of the feet or hands, generalised swelling or rapid weight gain tell your doctor as these may be signs of severe fluid retention. Uncommon cases of severe fluid retention have been reported in patients treated with Tasigna.

If you are the parent of a child who is being treated with Tasigna, tell the doctor if any of the above conditions apply to your child.

Children and adolescents

Tasigna is a treatment for children and adolescents with CML. There is no experience with the use of this medicine in children below 2 years of age. There is no experience with the use of Tasigna in newly diagnosed children below 10 years of age and limited experience in patients below 6 years of age who are no longer benefiting from previous treatment for CML.

Some children and adolescents taking Tasigna may have slower than normal growth. The doctor will monitor growth at regular visits.

Other medicines and Tasigna

Tasigna may interfere with some other medicines.

Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines. This includes in particular:

- anti-arrhythmics used to treat irregular heart beat;
- chloroquine, halofantrine, clarithromycin, haloperidol, methadone, moxifloxacin medicines that may have an unwanted effect on the electrical activity of the heart;
- ketoconazole, itraconazole, voriconazole, clarithromycin, telithromycin used to treat infections;
- ritonavir a medicine from the class "antiproteases" used to treat HIV;
- carbamazepine, phenobarbital, phenytoin used to treat epilepsy;
- rifampicin used to treat tuberculosis;
- St. John's Wort a herbal product used to treat depression and other conditions (also known as *Hypericum perforatum*);
- midazolam used to relieve anxiety before surgery;
- alfentanil and fentanyl used to treat pain and as a sedative before or during surgery or medical procedures:
- cyclosporine, sirolimus and tacrolimus medicines that suppress the "self-defence" ability of the body and fight infections and are commonly used to prevent the rejection of transplanted organs such as the liver, heart and kidney;
- dihydroergotamine and ergotamine used to treat dementia;
- lovastatin, simvastatin used to treat high level of fats in blood;
- warfarin used to treat blood coagulation disorders (such as blood clots or thromboses);
- astemizole, terfenadine, cisapride, pimozide, quinidine, bepridil or ergot alkaloids (ergotamine, dihydroergotamine).

These medicines should be avoided during your treatment with Tasigna. If you are taking any of these, your doctor might prescribe other alternative medicines.

If you are taking a statin (a type of medicine to lower your blood cholesterol), talk to your doctor or pharmacist. If used with certain statins, Tasigna may increase the risk of statin-related muscle problems, which on rare occasions can lead to serious muscle breakdown (rhabdomyolysis) resulting in kidney damage.

In addition, tell your doctor or pharmacist before taking Tasigna if you are taking any antacids, which are medicines against heartburn. These medicines need to be taken separately from Tasigna:

- H2 blockers, which decrease the production of acid in the stomach. H2 blockers should be taken approximately 10 hours before and approximately 2 hours after you take Tasigna;
- antacids such as those containing aluminium hydroxide, magnesium hydroxide and simethicone, which neutralise high acidity in the stomach. These antacids should be taken approximately 2 hours before or approximately 2 hours after you take Tasigna.

You should also tell your doctor **if you are already taking Tasigna** and you are prescribed a new medicine that you have not taken previously during Tasigna treatment.

Tasigna with food and drink

Do not take Tasigna with food. Food may enhance the absorption of Tasigna and therefore increase the amount of Tasigna in the blood, possibly to a harmful level. Do not drink grapefruit juice or eat grapefruit. It may increase the amount of Tasigna in the blood, possibly to a harmful level.

Pregnancy and breast-feeding

- **Tasigna is not recommended during pregnancy** unless clearly necessary. If you are pregnant or think that you may be, tell your doctor who will discuss with you whether you can take this medicine during your pregnancy.
- **Women who might get pregnant** are advised to use highly effective contraception during treatment and for up to two weeks after ending treatment.
- **Breast-feeding is not recommended** during treatment with Tasigna and for two weeks after the last dose. Tell your doctor if you are breast-feeding.

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

Driving and using machines

If you experience side effects (such as dizziness or visual disorders) with a potential impact on the ability to safely drive or use any tools or machines after taking this medicine, you should refrain from these activities until the effect has disappeared.

Tasigna contains lactose

This medicine contains lactose (also known as milk sugar). If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicine.

3. How to take Tasigna

Always take this medicine exactly as your doctor or pharmacist has told you. Check with your doctor or pharmacist if you are not sure.

How much Tasigna to take

Use in adults

- **Patients newly diagnosed with CML**: The recommended dose is 600 mg per day. This dose is achieved by taking two hard capsules of 150 mg twice a day.
- Patients who are no longer benefiting from previous treatment for CML: The recommended dose is 800 mg per day. This dose is achieved by taking two hard capsules of 200 mg twice a day.

Use in children and adolescents

The dose given to your child will depend on your child's body weight and height. The doctor will calculate the correct dose to use and tell you which and how many capsules of Tasigna to give to your child. The total daily dose you give to your child must not exceed 800 mg.

Your doctor may prescribe a lower dose depending on how you respond to treatment.

Older people (age 65 years and over)

Tasigna can be used by people aged 65 years and over at the same dose as for other adults.

When to take Tasigna

Take the hard capsules:

- twice a day (approximately every 12 hours);

- at least 2 hours after any food;
- then wait 1 hour before eating again.

If you have questions about when to take this medicine, talk to your doctor or pharmacist. Taking Tasigna at the same time each day will help you remember when to take your hard capsules.

How to take Tasigna

- Swallow the hard capsules whole with water.
- Do not take any food together with the hard capsules.
- Do not open the hard capsules unless you are unable to swallow them. If so, you may sprinkle the content of each hard capsule in **one** teaspoon of apple sauce and take it immediately. Do not use more than one teaspoon of apple sauce for each hard capsule and do not use any food other than apple sauce.

How long to take Tasigna

Continue taking Tasigna every day for as long as your doctor tells you. This is a long-term treatment. Your doctor will regularly monitor your condition to check that the treatment is having the desired effect

Your doctor may consider discontinuing your treatment with Tasigna based on specific criteria. If you have questions about how long to take Tasigna, talk to your doctor.

If you take more Tasigna than you should

If you have taken more Tasigna than you should have, or if someone else accidentally takes your hard capsules, contact a doctor or hospital for advice straight away. Show them the pack of hard capsules and this package leaflet. Medical treatment may be necessary.

If you forget to take Tasigna

If you miss a dose, take your next dose as scheduled. Do not take a double dose to make up for a forgotten hard capsule.

If you stop taking Tasigna

Do not stop taking this medicine unless your doctor tells you to do so. Stopping Tasigna without your doctor's recommendation places you at risk for worsening of your disease which could have life-threatening consequences. Be sure to discuss with your doctor, nurse, and/or pharmacist if you are considering stopping Tasigna.

If your doctor recommends that you discontinue treatment with Tasigna

Your doctor will regularly evaluate your treatment with a specific diagnostic test and decide whether you should continue to take this medicine. If you are told to discontinue Tasigna, your doctor will continue to carefully monitor your CML before, during and after you have discontinued Tasigna and may tell you to re-start Tasigna if your condition indicates that this is necessary.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them. Most of the side effects are mild to moderate and will generally disappear after a few days to a few weeks of treatment.

Some side effects could be serious.

- signs of musculoskeletal pain: pain in joints and muscles
- signs of heart disorders: chest pain or discomfort, high or low blood pressure, irregular heart rhythm (fast or slow), palpitations (sensation of rapid heartbeat), fainting, blue discolouration of the lips, tongue or skin
- signs of artery blockage: pain, discomfort, weakness or cramping in the leg muscles, which may be due to decreased blood flow, ulcers on the legs or arms that heal slowly or not at all and

- noticeable changes in colour (blueness or paleness) or temperature (coolness) of the affected leg, arm, toes or fingers
- signs of underactive thyroid gland: weight gain, tiredness, hair loss, muscle weakness, feeling cold
- signs of overactive thyroid gland: fast heartbeat, bulging eyes, weight loss, swelling at the front of the neck
- signs of kidney or urinary tract disorders: thirst, dry skin, irritability, dark urine, decreased urine output, difficulty and pain when urinating, exaggerated sense of needing to urinate, blood in urine, abnormal urine colour
- signs of high blood level of sugar: excessive thirst, high urine output, increased appetite with weight loss, tiredness
- signs of vertigo: dizziness or spinning sensation
- signs of pancreatitis: severe upper (middle or left) abdominal pain
- signs of skin disorders: painful red lumps, skin pain, skin reddening, peeling or blisters
- signs of water retention: rapid weight gain, swelling of hands, ankles, feet or face
- signs of migraine: severe headache often accompanied by nausea, vomiting and sensitivity to light
- signs of blood disorders: fever, easy bruising or unexplained bleeding, severe or frequent infections, unexplained weakness
- signs of clotting within a vein: swelling and pain in one part of the body
- signs of nervous system disorders: weakness or paralysis of the limbs or face, difficulty speaking, severe headache, seeing, feeling or hearing things that are not there, changes in eyesight, loss of consciousness, confusion, disorientation, trembling, sensation of tingling, pain or numbness in fingers and toes
- signs of lung disorders: difficulty breathing or painful breathing, cough, wheezing with or without fever, swelling of the feet or legs
- signs of gastrointestinal disorders: abdominal pain, nausea, vomiting of blood, black or bloody stools, constipation, heartburn, stomach acid reflux, swollen abdomen
- signs of liver disorders: yellow skin and eyes, nausea, loss of appetite, dark-coloured urine
- signs of liver infection: recurrence (reactivation of hepatitis B infection)
- signs of eye disorders: visual disturbances including blurred vision, double vision, or perceived flashes of light, decreased sharpness or loss of vision, blood in eye, increased sensitivity of the eyes to light, eye pain, redness, itching or irritation, dry eye, swelling or itching of the eyelids
- signs of electrolyte imbalance: nausea, shortness of breath, irregular heartbeat, clouding of urine, tiredness and/or joint discomfort associated with abnormal results of blood tests (such as high levels of potassium, uric acid and phosphorous and low levels of calcium)

Contact your doctor immediately if you notice any of the above side effects.

Some side effects are very common (may affect more than 1 in 10 people)

- diarrhoea
- headache
- lack of energy
- muscle pain
- itching, rash
- nausea
- constipation
- vomiting
- hair loss
- pain in limbs, bone pain and spinal pain on discontinuing treatment with Tasigna
- slowing of growth in children and adolescents
- upper respiratory tract infection including sore throat and runny or stuffy nose, sneezing
- low level of blood cells (red cells, platelets) or haemoglobin
- high blood level of lipase (pancreas function)
- high blood level of bilirubin (liver function)
- high blood level of alanine aminotransferases (liver enzymes)

Some side effects are common (may affect up to 1 in 10 people)

- pneumonia
- abdominal pain, stomach discomfort after meals, flatulence, swelling or bloating of the abdomen
- bone pain, muscle spasms
- pain (including neck pain)
- dry skin, acne, decreased skin sensitivity
- weight decrease or increase
- insomnia, depression, anxiety
- night sweats, excessive sweating
- generally feeling unwell
- nose bleed
- signs of gout: painful and swollen joints
- inability to achieve or maintain an erection
- flu-like symptoms
- sore throat
- bronchitis
- ear pain, hearing noises (e.g. ringing, humming) in the ears that have no external source (also called tinnitus)
- haemorrhoids
- heavy periods
- itching at the hair follicles
- oral or vaginal thrush
- signs of conjunctivitis: discharge from the eye with itching, redness and swelling
- eye irritation, red eyes
- signs of hypertension: high blood pressure, headache, dizziness
- flushing
- signs of peripheral arterial occlusive disease: pain, discomfort, weakness or cramping in the leg muscles, which may be due to decreased blood flow, ulcers on the legs or arms that heal slowly or not at all and noticeable changes in colour (blueness or paleness) or temperature (coolness) of the legs or arms (possible signs of a blocked artery in the affected leg, arm, toes or fingers)
- shortness of breath (also called dyspnoea)
- mouth sores with gum inflammation (also called stomatitis)
- high blood level of amylase (pancreas function)
- high blood level of creatinine (kidney function)
- high blood level of alkaline phosphatase or creatine phosphokinase
- high blood level of aspartate aminotransferases (liver enzymes)
- high blood level of gamma glutamyltransferases (liver enzymes)
- signs of leukopenia or neutropenia: low level of white blood cells
- increase in the number of platelets or white cells in the blood
- low blood level of magnesium, potassium, sodium, calcium or phosphorus
- increased blood level of potassium, calcium or phosphorus
- high blood level of fats (including cholesterol)
- high blood level of uric acid

Some side effects are uncommon (may affect up to 1 in 100 people)

- allergy (hypersensitivity to Tasigna)
- dry mouth
- breast pain
- pain or discomfort on the side of the body
- increased appetite
- breast enlargement in men
- herpes virus infection
- muscle and joint stiffness, joint swelling
- feeling body temperature change (including feeling hot, feeling cold)
- disturbed sense of taste
- frequent urine output

- signs of inflammation of the stomach lining: abdominal pain, nausea, vomiting ,diarrhoea, bloating of the abdomen
- memory loss
- skin cyst, thinning or thickening of the skin, thickening of the outermost layer of the skin, skin discolouration
- signs of psoriasis: thickened patches of red/silver skin
- increased sensitivity of the skin to light
- difficulty hearing
- joint inflammation
- urinary incontinence
- inflammation of the intestine (also called enterocolitis)
- anal abscess
- nipple swelling
- symptoms of restless legs syndrome (an irresistable urge to move a part of the body, usually the leg, accompanied by uncomfortable sensations)
- signs of sepsis: fever, chest pain, elevated/increased heart rate, shortness of breath or rapid breathing
- skin infection (subcutaneous abscess)
- skin wart
- increase in specific types of white blood cells (called eosinophils)
- signs of lymphopenia: low level of white blood cells
- high blood level of parathyroid hormone (a hormone regulating calcium and phosphorus levels)
- high blood level of lactate dehydrogenase (an enzyme)
- signs of low blood level of sugar: nausea, sweating, weakness, dizziness, trembling, headache
- dehydration
- abnormal blood level of fat
- involuntary shaking (also called tremor)
- difficulty concentrating
- unpleasant and abnormal feeling when touched (also called dysaesthesia)
- tiredness (also called fatigue)
- sensation of numbness or tingling in the fingers and toes (also called peripheral neuropathy)
- paralysis of any muscle of the face
- red patch in the white of the eye caused by broken blood vessels (also called conjunctival haemorrhage
- blood in eyes (also called eye haemorrhage)
- eye irritation
- signs of heart attack (also called myocardial infarction): sudden and crushing chest pain, tiredness, irregular heartbeat
- signs of heart murmur: tiredness, chest discomfort, light-headedness, chest pain, palpitations
- fungal infection of the feet
- signs of heart failure: breathlessness, difficulty breathing when lying down, swelling of the feet or legs
- pain behind the breast bone (also called pericarditis)
- signs of hypertensive crisis: severe headache, dizziness, nausea
- leg pain and weakness brought on by walking (also called intermittent claudication)
- signs of narrowing of the arteries of the limbs: possible high blood pressure, painful cramping in one or both hips, thighs or calf muscles after certain activities such as walking or climbing stairs, leg numbness or weakness
- bruising (when you have not hurt yourself)
- fatty deposits in the arteries that can cause blockage (also called arteriosclerosis)
- signs of low blood pressure (also called hypotension): light-headedness, dizziness or fainting
- signs of pulmonary oedema: breathlessness
- signs of pleural effusion: fluid collection between the layers of tissue that line the lungs and chest cavity (which, if severe, can decrease the heart's ability to pump blood), chest pain, cough, hiccups, rapid breathing
- signs of interstitial lung disease: cough, difficulty breathing, painful breathing
- signs of pleuritic pain: chest pain

- signs of pleurisy: cough, painful breathing
- hoarse voice
- signs of pulmonary hypertension: high blood pressure in the arteries of the lungs
- wheezing
- sensitive teeth
- signs of inflammation (also called gingivitis): gum bleeding, tender or enlarged gums
- high blood level of urea (kidney function)
- change in blood proteins (low level of globulins or presence of paraprotein)
- high blood level of unconjugated bilirubin
- high blood level of troponins

Some side effects are rare (may affect up to people 1 in 1,000)

- reddening and/or swelling and possibly peeling on the palms of the hands and soles of the feet (so-called hand-foot syndrome)
- warts in the mouth
- feeling of hardening or stiffness in the breasts
- inflammation of the thyroid gland (also called thyroiditis)
- disturbed or depressed mood
- signs of secondary hyperparathyroidism: bone and joint pain, excessive urination, abdominal pain, weakness, tiredness
- signs of narrowing of the arteries in the brain: loss of vision in part or all of both eyes, double vision, vertigo (spinning sensation), numbness or tingling, loss of coordination, dizziness or confusion
- swelling of the brain (possible headache and/or mental status changes)
- signs of optic neuritis: blurred vision, loss of vision
- signs of heart dysfunction (ejection fraction decreased): tiredness, chest discomfort, light-headedness, pain, palpitations
- low or high blood level of insulin (a hormone regulating blood sugar level)
- low blood level of insulin C peptide (pancreas function)
- sudden death

The following other side effects have been reported with frequency not known (cannot be estimated from the available data):

- signs of heart dysfunction (ventricular dysfunction): shortness of breath, exertion at rest, irregular heartbeat, chest discomfort, light-headedness, pain, palpitations, excessive urination, swelling in the feet, ankles and abdomen.

Reporting of side effects

If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in <u>Appendix V</u>. By reporting side effects you can help provide more information on the safety of this medicine.

5. How to store Tasigna

- Keep this medicine out of the sight and reach of children.
- Do not use this medicine after the expiry date which is stated on the carton and blister after EXP. The expiry date refers to the last day of that month.
- Do not store above 30°C.
- Store in the original package in order to protect from moisture.
- Do not use this medicine if you notice that the pack is damaged or shows signs of tampering.
- Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Tasigna contains

- The active substance is nilotinib.
- Each 50 mg hard capsule contains 50 mg nilotinib (as hydrochloride monohydrate). The other ingredients are:

Capsule content: Lactose monohydrate, crospovidone type A, poloxamer 188, colloidal anhydrous silica, magnesium stearate

Capsule shell: Gelatin, titanium dioxide (E171), red iron oxide (E172), yellow iron oxide (E172)

Printing ink: Shellac (E904), black iron oxide (E172), propylene glycol, ammonium hydroxide

- Each 150 mg hard capsule contains 150 mg nilotinib (as hydrochloride monohydrate). The other ingredients are:

Capsule content: Lactose monohydrate, crospovidone type A, poloxamer 188, colloidal anhydrous silica, magnesium stearate

Capsule shell: Gelatin, titanium dioxide (E171), red iron oxide (E172), yellow iron oxide (E172)

Printing ink: Shellac (E904), black iron oxide (E172), n-butyl alcohol, propylene glycol, dehydrated ethanol, isopropyl alcohol, ammonium hydroxide

- Each 200 mg hard capsule contains 200 mg nilotinib (as hydrochloride monohydrate). The other ingredients are:

Capsule content: Lactose monohydrate, crospovidone type A, poloxamer 188, colloidal anhydrous silica, magnesium stearate

Capsule shell: Gelatin, titanium dioxide (E171), yellow iron oxide (E172)

Printing ink: Shellac (E904), dehydrated alcohol, isopropyl alcohol, butyl alcohol, propylene glycol, strong ammonia solution, potassium hydroxide, red iron oxide (E172)

What Tasigna looks like and contents of the pack

Tasigna 50 mg is supplied as hard capsules. The hard capsules are red/light yellow. A black imprint is stamped on each hard capsule ("NVR/ABL").

Tasigna 150 mg is supplied as hard capsules. The hard capsules are red. A black imprint is stamped on each hard capsule ("NVR/BCR").

Tasigna 200 mg is supplied as hard capsules. The hard capsules are light yellow. A red imprint is stamped on each hard capsule ("NVR/TKI").

Tasigna 50 mg hard capsules are available in a pack containing 120 hard capsules (3 packs of 40 hard capsules).

Tasigna 150 mg hard capsules are available in packs containing 28 or 40 hard capsules and in multipacks of 112 hard capsules (comprising 4 cartons, each containing 28 hard capsules), 120 hard capsules (comprising 3 cartons, each containing 40 hard capsules) or 392 hard capsules (comprising 14 cartons, each containing 28 hard capsules).

Tasigna 200 mg hard capsules are available in a wallet containing 28 hard capsules and in a carton containing 28 or 40 hard capsules. Tasigna is also available in multipacks of 112 hard capsules (comprising 4 wallets, each containing 28 hard capsules), 112 hard capsules (comprising 4 cartons, each containing 28 hard capsules), 120 hard capsules (comprising 3 cartons, each containing 40 hard capsules) or 392 hard capsules (comprising 14 cartons, each containing 28 hard capsules).

Not all packs may be marketed in your country.

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Other sources of information

Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu. There are also links to other websites about rare diseases and treatments.