This document is the approved product information for Alunbrig, with the changes since the previous procedure affecting the product information (EMEA/H/C/004248/R/0049) tracked.

For more information, see the European Medicines Agency’s website: <https://www.ema.europa.eu/en/medicines/human/EPAR/alunbrig>

**ANNEX I**

# SUMMARY OF PRODUCT CHARACTERISTICS

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 30 mg film‑coated tablets

Alunbrig 90 mg film‑coated tablets

Alunbrig 180 mg film‑coated tablets

**2. QUALITATIVE AND QUANTITATIVE COMPOSITION**

Alunbrig 30 mg film‑coated tablets

Each film‑coated tablet contains 30 mg of brigatinib.

*Excipient with known effect*

Each film‑coated tablet contains 56 mg of lactose monohydrate.

Alunbrig 90 mg film‑coated tablets

Each film‑coated tablet contains 90 mg of brigatinib.

*Excipient with known effect*

Each film‑coated tablet contains 168 mg of lactose monohydrate.

Alunbrig 180 mg film‑coated tablets

Each film‑coated tablet contains 180 mg of brigatinib.

*Excipient with known effec*t

Each film‑coated tablet contains 336 mg of lactose monohydrate.

For the full list of excipients, see section 6.1.

**3. PHARMACEUTICAL FORM**

Film‑coated tablet (tablet).

Alunbrig 30 mg film‑coated tablets

Round, white to off‑white film‑coated tablet of approximately 7 mm in diameter with debossed “U3” on one side and plain on the other side.

Alunbrig 90 mg film‑coated tablets

Oval, white to off‑white film‑coated tablet of approximately 15 mm in length with debossed “U7” on one side and plain on the other side.

Alunbrig 180 mg film‑coated tablets

Oval, white to off‑white film‑coated tablet of approximately 19 mm in length with debossed “U13” on one side and plain on the other side.

**4. CLINICAL PARTICULARS**

**4.1 Therapeutic indications**

Alunbrig is indicated as monotherapy for the treatment of adult patients with anaplastic lymphoma kinase (ALK)‑positive advanced non‑small cell lung cancer (NSCLC) previously not treated with an ALK inhibitor.

Alunbrig is indicated as monotherapy for the treatment of adult patients with ALK‑positive advanced NSCLC previously treated with crizotinib.

**4.2 Posology and method of administration**

Treatment with Alunbrig should be initiated and supervised by a physician experienced in the use of anticancer medicinal products.

ALK‑positive NSCLC status should be known prior to initiation of Alunbrig therapy. A validated ALK assay is necessary for the selection of ALK‑positive NSCLC patients (see section 5.1). Assessment for ALK‑positive NSCLC should be performed by laboratories with demonstrated proficiency in the specific technology being utilised.

Posology

The recommended starting dose of Alunbrig is 90 mg once daily for the first 7 days, then 180 mg once daily.

If Alunbrig is interrupted for 14 days or longer for reasons other than adverse reactions, treatment should be resumed at 90 mg once daily for 7 days before increasing to the previously tolerated dose.

If a dose is missed or vomiting occurs after taking a dose, an additional dose should not be administered and the next dose should be taken at the scheduled time.

Treatment should continue as long as clinical benefit is observed.

*Dose adjustments*

Dosing interruption and/or dose reduction may be required based on individual safety and tolerability.

Alunbrig dose reduction levels are summarised in Table 1.

**Table 1: Recommended Alunbrig dose reduction levels**

|  |  |  |  |
| --- | --- | --- | --- |
| **Dose** | **Dose reduction levels** | | |
| **First** | **Second** | **Third** |
| 90 mg once daily  (first 7 days) | reduce to 60 mg once daily | permanently discontinue | not applicable |
| 180 mg once daily | reduce to 120 mg once daily | reduce to 90 mg once daily | reduce to 60 mg once daily |

Alunbrig should be permanently discontinued if patient is unable to tolerate the 60 mg once daily dose.

Recommendations for dose modifications of Alunbrig for the management of adverse reactions are summarised in Table 2.

**Table 2: Recommended Alunbrig dose modifications for adverse reactions**

| **Adverse reaction** | **Severity**\* | **Dose modification** |
| --- | --- | --- |
| Interstitial lung disease (ILD)/pneumonitis | Grade 1 | * If event occurs during the first 7 days of treatment, Alunbrig should be withheld until recovery to baseline, then resumed at same dose level and not escalated to 180 mg once daily. * If ILD/pneumonitis occurs after the first 7 days of treatment, Alunbrig should be withheld until recovery to baseline, then resumed at same dose level. * If ILD/pneumonitis recurs, Alunbrig should be permanently discontinued. |
| Grade 2 | * If ILD/pneumonitis occurs during the first 7 days of treatment, Alunbrig should be withheld until recovery to baseline, then resumed at next lower dose level as described in Table 1 and not escalated to 180 mg once daily. * If ILD/pneumonitis occurs after the first 7 days of treatment, Alunbrig should be withheld until recovery to baseline. Alunbrig should be resumed at next lower dose level as described in Table 1. * If ILD/pneumonitis recurs, Alunbrig should be permanently discontinued. |
| Grade 3 or 4 | * Alunbrig should be permanently discontinued. |
| Hypertension | Grade 3 hypertension (SBP ≥ 160 mmHg or DBP ≥ 100 mmHg, medical intervention indicated, more than one anti‑hypertensive medicinal product, or more intensive therapy than previously used indicated) | * Alunbrig should be withheld until hypertension has recovered to Grade ≤ 1 (SBP < 140 mmHg and DBP < 90 mmHg), then resumed at same dose. * If Grade 3 hypertension recurs, Alunbrig should be withheld until hypertension has recovered to Grade ≤ 1 then resumed at the next lower dose level per Table 1 or permanently discontinued. |
| Grade 4 hypertension (life threatening consequences, urgent intervention indicated) | * Alunbrig should be withheld until hypertension has recovered to Grade ≤ 1 (SBP < 140 mmHg and DBP < 90 mmHg), then resumed at the next lower dose level per Table 1 or permanently discontinued. * If Grade 4 hypertension recurs, Alunbrig should be permanently discontinued. |
| Bradycardia (heart rate less than 60 bpm) | Symptomatic bradycardia | * Alunbrig should be withheld until recovery to asymptomatic bradycardia or to a resting heart rate of 60 bpm or above. * If a concomitant medicinal product known to cause bradycardia is identified and discontinued, or its dose is adjusted, Alunbrig should be resumed at same dose upon recovery to asymptomatic bradycardia or to a resting heart rate of 60 bpm or above. * If no concomitant medicinal product known to cause bradycardia is identified, or if contributing concomitant medicinal products are not discontinued or dose modified, Alunbrig should be resumed at the next lower dose level per Table 1 upon recovery to asymptomatic bradycardia or to a resting heart rate of 60 bpm or above. |
| Bradycardia with life‑threatening consequences, urgent intervention indicated | * If contributing concomitant medicinal product is identified and discontinued, or its dose is adjusted, Alunbrig should be resumed at the next lower dose level per Table 1 upon recovery to asymptomatic bradycardia or to a resting heart rate of 60 bpm or above, with frequent monitoring as clinically indicated. * Alunbrig should be permanently discontinued if no contributing concomitant medicinal product is identified. * Alunbrig should be permanently discontinued in case of recurrence. |
| Elevation of CPK | Grade 3 or 4 elevation of CPK (> 5.0 × ULN) with Grade ≥ 2 muscle pain or weakness | * Alunbrig should be withheld until recovery to Grade ≤ 1 (≤ 2.5 × ULN) elevation of CPK or to baseline, then resumed at the same dose. * If Grade 3 or 4 elevation of CPK recurs with Grade ≥ 2 muscle pain or weakness, Alunbrig should be withheld until recovery to Grade ≤ 1 (≤ 2.5 × ULN) elevation of CPK or to baseline, then resumed at the next lower dose level per Table 1. |
| Elevation of lipase or amylase | Grade 3 elevation of lipase or amylase (> 2.0 × ULN) | * Alunbrig should be withheld until recovery to Grade ≤ 1 (≤ 1.5 × ULN) or to baseline, then resumed at same dose. * If Grade 3 elevation of lipase or amylase recurs, Alunbrig should be withheld until recovery to Grade ≤ 1 (≤ 1.5 × ULN) or to baseline, then resumed at the next lower dose level per Table 1. |
| Grade 4 elevation of lipase or amylase (> 5.0 x ULN) | * Alunbrig should be withheld until recovery to Grade ≤ 1 (≤ 1.5 × ULN), then resumed at the next lower dose level per Table 1. |
| Hepatotoxicity | Grade ≥ 3 elevation (> 5.0 × ULN) of either alanine aminotransferase (ALT) or aspartate aminotransferase (AST) with bilirubin ≤ 2 × ULN | * Alunbrig should be withheld until recovery to baseline or less than or equal to 3 × ULN, then resumed at next lower dose per Table 1. |
| Grade ≥ 2 elevation (> 3 × ULN) of ALT or AST with concurrent total bilirubin elevation > 2 × ULN in the absence of cholestasis or haemolysis | * Alunbrig should be permanently discontinued. |
| Hyperglycaemia | For Grade 3 (greater than 250 mg/dL or 13.9 mmol/L) or greater | * If adequate hyperglycaemic control cannot be achieved with optimal medical management, Alunbrig should be withheld until adequate hyperglycaemic control is achieved. Upon recovery, Alunbrig may either be resumed at the next lower dose per Table 1 or permanently discontinued. |
| Visual disturbance | Grade 2 or 3 | * Alunbrig should be withheld until recovery to Grade 1 or baseline, then resumed at the next lower dose level per Table 1. |
| Grade 4 | * Alunbrig should be permanently discontinued. |
| Other adverse reactions | Grade 3 | * Alunbrig should be withheld until recovery to baseline, then resumed at the same dose level. * If the Grade 3 event recurs, Alunbrig should be withheld until recovery to baseline, then resumed at the next lower dose level as per Table 1 or permanently discontinued. |
| Grade 4 | * Alunbrig should be withheld until recovery to baseline, then resumed at the next lower dose level as per Table 1. * If the Grade 4 event recurs, Alunbrig should be withheld until recovery to baseline, then resumed at the next lower dose level as per Table 1 or permanently discontinued. |
| bpm = beats per minute; CPK = Creatine Phosphokinase; DBP = diastolic blood pressure; SBP = systolic blood pressure; ULN = upper limit of normal | | |

\*Graded per National Cancer Institute Common Terminology Criteria for Adverse Events. Version 4.0 (NCI CTCAE v4).

*Special populations*

*Elderly*

The limited data on the safety and efficacy of Alunbrig in patients aged 65 years and older suggest that a dose adjustment is not required in elderly patients (see section 4.8). There are no available data on patients over 85 years of age.

*Hepatic impairment*

No dose adjustment of Alunbrig is required for patients with mild hepatic impairment (Child‑Pugh class A) or moderate hepatic impairment (Child‑Pugh class B). A reduced starting dose of 60 mg once daily for the first 7 days, then 120 mg once daily is recommended for patients with severe hepatic impairment (Child‑Pugh class C) (see section 5.2).

*Renal impairment*

No dose adjustment of Alunbrig is required for patients with mild or moderate renal impairment (estimated glomerular filtration rate (eGFR) ≥ 30 mL/min). A reduced starting dose of 60 mg once daily for the first 7 days, then 90 mg once daily is recommended for patients with severe renal impairment (eGFR < 30 mL/min) (see section 5.2). Patients with severe renal impairment should be closely monitored for new or worsening respiratory symptoms that may indicate ILD/pneumonitis (e.g., dyspnoea, cough, etc.) particularly in the first week (see section 4.4).

*Paediatric population*

The safety and efficacy of Alunbrig in patients less than 18 years of age have not been established. No data are available.

Method of administration

Alunbrig is for oral use. The tablets should be swallowed whole and with water. Alunbrig may be taken with or without food.

Grapefruit or grapefruit juice may increase plasma concentrations of brigatinib and should be avoided (see section 4.5).

**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**

Pulmonary adverse reactions

Severe, life‑threatening, and fatal pulmonary adverse reactions, including those with features consistent with ILD/pneumonitis, can occur in patients treated with Alunbrig (see section 4.8).

Most pulmonary adverse reactions were observed within the first 7 days of treatment. Grade 1‑2 pulmonary adverse reactions resolved with interruption of treatment or dose modification. Increased age and shorter interval (less than 7 days) between the last dose of crizotinib and the first dose of Alunbrig were independently associated with an increased rate of these pulmonary adverse reactions. These factors should be considered when initiating treatment with Alunbrig. Patients with a history of ILD or drug‑induced pneumonitis were excluded from the pivotal trials.

Some patients experienced pneumonitis later in treatment with Alunbrig.

Patients should be monitored for new or worsening respiratory symptoms (e.g., dyspnoea, cough,etc.), particularly in the first week of treatment. Evidence of pneumonitis in any patient with worsening respiratory symptoms should be promptly investigated. If pneumonitis is suspected, the dose of Alunbrig should be withheld, and the patient evaluated for other causes of symptoms (e.g., pulmonary embolism, tumour progression, and infectious pneumonia). The dose should be modified accordingly (see section 4.2).

Hypertension

Hypertension has occurred in patients treated with Alunbrig (see section 4.8).

Blood pressure should be monitored regularly during treatment with Alunbrig. Hypertension should be treated according to standard guidelines to control blood pressure. Heart rate should be monitored more frequently in patients if concomitant use of a medicinal product known to cause bradycardia cannot be avoided. For severe hypertension (≥ Grade 3), Alunbrig should be withheld until hypertension has recovered to Grade 1 or to baseline. The dose should be modified accordingly (see section 4.2).

Bradycardia

Bradycardia has occurred in patients treated with Alunbrig (see section 4.8). Caution should be exercised when administering Alunbrig in combination with other agents known to cause bradycardia. Heart rate and blood pressure should be monitored regularly.

If symptomatic bradycardia occurs, treatment with Alunbrig should be withheld and concomitant medicinal products known to cause bradycardia should be evaluated. Upon recovery, the dose should be modified accordingly (see section 4.2). In case of life‑threatening bradycardia, if no contributing concomitant medication is identified or in case of recurrence, treatment with Alunbrig should be discontinued (see section 4.2)*.*

Visual disturbance

Visual disturbance adverse reactions have occurred in patients treated with Alunbrig (see section 4.8). Patients should be advised to report any visual symptoms. For new or worsening severe visual symptoms, an ophthalmologic evaluation and dose reduction should be considered(see section 4.2).

Creatine phosphokinase (CPK) elevation

Elevations of CPK have occurred in patients treated with Alunbrig (see section 4.8). Patients should be advised to report any unexplained muscle pain, tenderness, or weakness. CPK levels should be monitored regularly during Alunbrig treatment. Based on the severity of the CPK elevation, and if associated with muscle pain or weakness, treatment with Alunbrig should be withheld, and the dose modified accordingly (see section 4.2).

Elevations of pancreatic enzymes

Elevations of amylase and lipase have occurred in patients treated with Alunbrig (see section 4.8). Lipase and amylase should be monitored regularly during treatment with Alunbrig. Based on the severity of the laboratory abnormalities, treatment with Alunbrig should be withheld, and the dose modified accordingly (see section 4.2).

Hepatotoxicity

Elevations of hepatic enzymes (aspartate aminotransferase, alanine aminotransferase) and bilirubin have occurred in patients treated with Alunbrig (see section 4.8). Liver function, including AST, ALT and total bilirubin should be assessed prior to the initiation of Alunbrig and then every 2 weeks during the first 3 months of treatment. Thereafter, monitoring should be performed periodically. Based on the severity of the laboratory abnormalities, treatment should be withheld, and the dose modified accordingly (see section 4.2).

Hyperglycaemia

Elevations of serum glucose have occurred in patients treated with Alunbrig. Fasting serum glucose should be assessed prior to initiation of Alunbrig and monitored periodically thereafter. Antihyperglycaemic treatment should be initiated or optimised as needed. If adequate hyperglycaemic control cannot be achieved with optimal medical management, Alunbrig should be withheld until adequate hyperglycaemic control is achieved; upon recovery reducing the dose as described in Table 1 may be considered or Alunbrig may be permanently discontinued.

Drug‑drug interactions

The concomitant use of Alunbrig with strong CYP3A inhibitors should be avoided. If concomitant use of strong CYP3A inhibitors cannot be avoided, the dose of Alunbrig should be reduced from 180 mg to 90 mg, or from 90 mg to 60 mg. After discontinuation of a strong CYP3A inhibitor, Alunbrig should be resumed at the dose that was tolerated prior to the initiation of the strong CYP3A inhibitor.

The concomitant use of Alunbrig with strong and moderate CYP3A inducers should be avoided (see section 4.5). If concomitant use of moderate CYP3A inducers cannot be avoided, the dose of Alunbrig may be increased in 30 mg increments after 7 days of treatment with the current Alunbrig dose as tolerated, up to a maximum of twice the Alunbrig dose that was tolerated prior to the initiation of the moderate CYP3A inducer. After discontinuation of a moderate CYP3A inducer, Alunbrig should be resumed at the dose that was tolerated prior to the initiation of the moderate CYP3A inducer.

Photosensitivity and photodermatosis

Photosensitivity to sunlight has occurred in patients treated with Alunbrig (see section 4.8). Patients should be advised to avoid prolonged sun exposure while taking Alunbrig, and for at least 5 days after discontinuation of treatment. When outdoors, patients should be advised to wear a hat and protective clothing, and to use a broad‑spectrum Ultraviolet A (UVA)/ Ultraviolet B (UVB) sunscreen and lip balm (SPF ≥ 30) to help protect against potential sunburn. For severe photosensitivity reactions (≥ Grade 3), Alunbrig should be withheld until recovery to baseline. The dose should be modified accordingly (see section 4.2).

Fertility

Women of childbearing potential should be advised to use effective non‑hormonal contraception during treatment with Alunbrig and for at least 4 months following the final dose. Men with female partners of childbearing potential should be advised to use effective contraception during treatment and for at least 3 months after the last dose of Alunbrig (see section 4.6).

Lactose

Alunbrig contains lactose monohydrate. Patients with rare hereditary problems of galactose intolerance, total lactase deficiency or glucose‑galactose malabsorption should not take this medicinal product.

Sodium

This medicinal product contains less than 1 mmol sodium (23 mg) per tablet, that is to say essentially ‘sodium‑free’.

**4.5 Interaction with other medicinal products and other forms of interaction**

Agents that may increase brigatinib plasma concentrations

*CYP3A inhibitors*

*In vitro* studies demonstrated that brigatinib is a substrate of CYP3A4/5. In healthy subjects, coadministration of multiple 200 mg twice daily doses of itraconazole, a strong CYP3A inhibitor, with a single 90 mg brigatinib dose increased brigatinib Cmax by 21%, AUC0‑INF by 101% (2‑fold), and AUC0‑120 by 82% (< 2‑fold), relative to a 90 mg brigatinib dose administered alone. The concomitant use of strong CYP3A inhibitors with Alunbrig, including but not limited to certain antivirals (e.g., indinavir, nelfinavir, ritonavir, saquinavir), macrolide antibiotics (e.g., clarithromycin, telithromycin, troleandomycin), antifungals (e.g., ketoconazole, voriconazole), and nefazodone should be avoided. If concomitant use of strong CYP3A inhibitors cannot be avoided, the dose of Alunbrig should be reduced by approximately 50% (i.e. from 180 mg to 90 mg, or from 90 mg to 60 mg). After discontinuation of a strong CYP3A inhibitor, Alunbrig should be resumed at the dose that was tolerated prior to the initiation of the strong CYP3A inhibitor.

Moderate CYP3A inhibitors (e.g., diltiazem and verapamil) may increase the AUC of brigatinib by approximately 40% based on simulations from a physiologically‑based pharmacokinetic model. No dose adjustment is required for Alunbrig in combination with moderate CYP3A inhibitors. Patients should be closely monitored when Alunbrig is coadministered with moderate CYP3A inhibitors.

Grapefruit or grapefruit juice may also increase plasma concentrations of brigatinib and should be avoided (see section 4.2).

*CYP2C8 inhibitors*

*In vitro* studies demonstrated that brigatinib is a substrate of CYP2C8. In healthy subjects, coadministration of multiple 600 mg twice daily doses of gemfibrozil, a strong CYP2C8 inhibitor, with a single 90 mg brigatinib dose reduced brigatinib Cmax by 41%, AUC0‑INF by 12%, and AUC0‑120 by 15%, relative to a 90 mg brigatinib dose administered alone. The effect of gemfibrozil on the pharmacokinetics of brigatinib is not clinically meaningful and the underlying mechanism for the decreased exposure of brigatinib is unknown. No dose adjustment is required during coadministration with strong CYP2C8 inhibitors.

*P‑gp and BCRP inhibitors*

Brigatinib is a substrate of P‑glycoprotein (P‑gp) and breast cancer resistance protein (BCRP) *in vitro*. Given that brigatinib exhibits high solubility and high permeability, inhibition of P‑gp and BCRP is not expected to result in a clinically meaningful change in the systemic exposure of brigatinib. No dose adjustment is required for Alunbrig during coadministration with P‑gp and BCRP inhibitors.

Agents that may decrease brigatinib plasma concentrations

*CYP3A inducers*

In healthy subjects, coadministration of multiple 600 mg daily doses of rifampicin, a strong CYP3A inducer, with a single 180 mg brigatinib dose decreased brigatinib Cmax by 60%, AUC0‑INF by 80% (5‑fold), and AUC0‑120 by 80% (5‑fold), relative to a 180 mg brigatinib dose administered alone. The concomitant use of strong CYP3A inducers with Alunbrig, including but not limited to rifampicin, carbamazepine, phenytoin, rifabutin, phenobarbital, and St. John’s wort should be avoided.

Moderate CYP3A inducers may decrease the AUC of brigatinib by approximately 50% based on simulations from a physiologically‑based pharmacokinetic model. The concomitant use of moderate CYP3A inducers with Alunbrig, including but not limited to efavirenz, modafinil, bosentan, etravirine, and nafcillin should be avoided. If concomitant use of moderate CYP3A inducers cannot be avoided, the dose of Alunbrig may be increased in 30 mg increments after 7 days of treatment with the current Alunbrig dose as tolerated, up to a maximum of twice the Alunbrig dose that was tolerated prior to the initiation of the moderate CYP3A inducer. After discontinuation of a moderate CYP3A inducer, Alunbrig should be resumed at the dose that was tolerated prior to the initiation of the moderate CYP3A inducer.

Agents that may have their plasma concentrations altered by brigatinib

*CYP3A substrates*

*In vitro* studies in hepatocytes have shown that brigatinib is an inducer of CYP3A4. In patients with cancer, coadministration of multiple 180 mg daily doses of Alunbrig with a single 3 mg oral dose of midazolam, a sensitive CYP3A substrate, decreased midazolam Cmax by 16%, AUC0‑INF by 26%, and AUC0‑last by 30%, relative to a 3 mg oral dose of midazolam administered alone. Brigatinib reduces plasma concentrations of coadministered medicinal products that are predominantly metabolised by CYP3A. Therefore, coadministration of Alunbrig with CYP3A substrates with a narrow therapeutic index (e.g., alfentanil, fentanyl, quinidine, cyclosporine, sirolimus, tacrolimus) should be avoided as their effectiveness may be reduced.

Alunbrig may also induce other enzymes and transporters (e.g., CYP2C, P‑gp) via the same mechanisms responsible for induction of CYP3A (e.g., pregnane X receptor activation).

*Transporter substrates*

Coadministration of brigatinib with substrates of P‑gp (e.g., digoxin, dabigatran, colchicine, pravastatin), BCRP (e.g., methotrexate, rosuvastatin, sulfasalazine), organic cation transporter 1 (OCT1), multidrug and toxin extrusion protein 1 (MATE1), and 2K (MATE2K) may increase their plasma concentrations. Patients should be closely monitored when Alunbrig is coadministered with substrates of these transporters with a narrow therapeutic index (e.g., digoxin, dabigatran, methotrexate).

**4.6 Fertility, pregnancy and lactation**

Women of childbearing potential/Contraception in males and females

Women of childbearing age being treated with Alunbrig should be advised not to become pregnant and men being treated with Alunbrig should be advised not to father a child during treatment. Women of reproductive potential should be advised to use effective non‑hormonal contraception during treatment with Alunbrig and for at least 4 months following the final dose. Men with female partners of reproductive potential should be advised to use effective contraception during treatment and for at least 3 months after the last dose of Alunbrig.

Pregnancy

Alunbrig may cause foetal harm when administered to a pregnant woman. Studies in animals have shown reproductive toxicity (see section 5.3). There are no clinical data on the use of Alunbrig in pregnant women. Alunbrig should not be used during pregnancy unless the clinical condition of the mother requires treatment. If Alunbrig is used during pregnancy, or if the patient becomes pregnant while taking this medicinal product, the patient should be apprised of the potential hazard to a foetus.

Breast‑feeding

It is unknown whether Alunbrig is excreted in human milk. Available data cannot exclude potential excretion in human milk. Breast‑feeding should be stopped during treatment with Alunbrig.

Fertility

No human data on the effect of Alunbrig on fertility are available. Based on repeat‑dose toxicity studies in male animals, Alunbrig may cause reduced fertility in males (see section 5.3). The clinical relevance of these findings to human fertility is unknown.

**4.7 Effects on ability to drive and use machines**

Alunbrig has minor influence on the ability to drive and use machines. However, caution should be exercised when driving or operating machines as patients may experience visual disturbance, dizziness, or fatigue while taking Alunbrig.

**4.8 Undesirable effects**

Summary of the safety profile

The most common adverse reactions (≥ 25%) reported in patients treated with Alunbrig at the recommended dosing regimen were increased AST, increased CPK, hyperglycaemia, increased lipase, hyperinsulinaemia, diarrhoea, increased ALT, increased amylase, anaemia, nausea, fatigue, hypophosphataemia, decreased lymphocyte count, cough, increased alkaline phosphatase, rash, increased APTT, myalgia, headache, hypertension, decreased white blood cell count, dyspnoea, and vomiting.

The most common serious adverse reactions (≥ 2%) reported in patients treated with Alunbrig at the recommended dosing regimen other than events related to neoplasm progression were pneumonia, pneumonitis, dyspnoea and pyrexia.

Tabulated list of adverse reactions

The data described below reflect exposure to Alunbrig at the recommended dosing regimen in three clinical trials: a Phase 3 trial (ALTA 1L) in patients with advanced ALK‑positive NSCLC previously not treated with an ALK‑inhibitor (N = 136), a Phase 2 trial (ALTA) in patients treated with Alunbrig with ALK‑positive NSCLC who previously progressed on crizotinib (N = 110), and a phase 1/2 dose escalation/expansion trial in patients with advanced malignancies (N = 28). Across these studies, the median duration of exposure in patients receiving Alunbrig at the recommended dosing regimen was 21.8 months.

Adverse reactions reported are presented in Table 3 and are listed by system organ class, preferred term and frequency. Frequency categories are very common (≥ 1/10), common (≥ 1/100 to < 1/10) and uncommon (≥ 1/1 000 to < 1/100). Within each frequency grouping, undesirable effects are presented in order of frequency.

**Table 3: Adverse reactions reported in patients treated with Alunbrig (per Common Terminology Criteria for Adverse Events (CTCAE) version 4.03) at the 180 mg regimen (N = 274)**

| **System organ class** | **Frequency**  **category** | **Adverse reactions†**  **all grades** | **Adverse reactions**  **Grade 3‑4** |
| --- | --- | --- | --- |
| Infections and infestations | Very common | Pneumoniaa,b  Upper respiratory tract infection |  |
| Common |  | Pneumoniaa |
| Blood and lymphatic system disorders | Very common | Anaemia  Lymphocyte count decreased  APTT increased  White blood cell count decreased  Neutrophil count decreased | Lymphocyte count decreased |
| Common | Decreased platelet count | APTT increased  Anaemia |
| Uncommon |  | Neutrophil count decreased |
| Metabolism and nutrition disorders | Very common | Hyperglycaemia  Hyperinsulinaemiac  Hypophosphataemia  Hypomagnesaemia  Hypercalcaemia  Hyponatraemia  Hypokalaemia  Decreased appetite |  |
| Common |  | Hypophosphataemia  Hyperglycaemia  Hyponatraemia  Hypokalaemia  Decreased appetite |
| Psychiatric disorders | Common | Insomnia |  |
| Nervous system disorders | Very common | Headached  Peripheral neuropathye  Dizziness |  |
| Common | Memory impairment  Dysgeusia | Headached  Peripheral neuropathye |
| Uncommon |  | Dizziness |
| Eye disorders | Very common | Visual disturbancef |  |
| Common |  | Visual disturbancef |
| Cardiac disorders | Common | Bradycardiag  Electrocardiogram QT prolonged  Tachycardiah  Palpitations | Electrocardiogram QT prolonged |
| Uncommon |  | Bradycardiag |
| Vascular disorders | Very common | Hypertensioni | Hypertensioni |
| Respiratory, thoracic and mediastinal disorders | Very common | Cough  Dyspnoeaj |  |
| Common | Pneumonitisk | Pneumonitisk  Dyspnoeaj |
| Gastrointestinal disorders | Very common | Lipase increased  Diarrhoea  Amylase increased  Nausea  Vomiting  Abdominal painl  Constipation  Stomatitism | Lipase increased |
| Common | Dry mouth  Dyspepsia  Flatulence | Amylase increased  Nausea  Abdominal painl  Diarrhoea |
| Uncommon | Pancreatitis | Vomiting  Stomatitism  Dyspepsia  Pancreatitis |
| Hepatobiliary disorders | Very common | AST increased  ALT increased  Alkaline phosphatase increased |  |
| Common | Blood lactate dehydrogenase increased  Hyperbilirubinaemia | ALT increased  AST increased  Alkaline phosphatase increased |
| Uncommon |  | Hyperbilirubinaemia |
| Skin and subcutaneous tissue disorders | Very common | Rashn  Prurituso |  |
| Common | Dry skin  Photosensitivity reactionp | Rashn  Photosensitivity reactionp |
| Uncommon |  | Dry skin  Prurituso |
| Musculoskeletal and connective tissue disorders | Very common | Blood CPK increased  Myalgiaq  Arthralgia | Blood CPK increased |
| Common | Musculoskeletal chest pain  Pain in extremity  Musculoskeletal stiffness |  |
| Uncommon |  | Pain in extremity  Musculoskeletal chest pain  Myalgiaq |
| Renal and urinary disorders | Very common | Blood creatinine increased |  |
| General disorders and administration site conditions | Very common | Fatiguer  Oedemas  Pyrexia |  |
| Common | Non‑cardiac chest pain  Chest discomfort  Pain | Fatiguer |
| Uncommon |  | Pyrexia  Oedemas  Non‑cardiac chest pain |
| Investigations | Common | Blood cholesterol increasedt  Weight decreased |  |
| Uncommon |  | Weight decreased |
| † The frequencies for ADR terms associated with chemistry and haematology laboratory changes were determined based on the frequency of abnormal laboratory shifts from baseline.  a Includes atypical pneumonia, pneumonia, pneumonia aspiration, pneumonia cryptococcal, lower respiratory tract infection, lower respiratory tract infection viral, lung infection  b Includes Grade 5 events  c Grade not applicable  d Includes headache, sinus headache, head discomfort, migraine, tension headache  e Includes paraesthesia, peripheral sensory neuropathy, dysaesthesia, hyperaesthesia, hypoaesthesia, neuralgia, neuropathy peripheral, neurotoxicity, peripheral motor neuropathy, polyneuropathy, burning sensation, post herpetic neuralgia  f Includes altered visual depth perception, cataract, colour blindness acquired, diplopia, glaucoma, intraocular pressure increased, macular oedema, photophobia, photopsia, retinal oedema, vision blurred, visual acuity reduced, visual field defect, visual impairment, vitreous detachment, vitreous floaters, amaurosis fugax  g Includes bradycardia, sinus bradycardia  h Includes sinus tachycardia, tachycardia, atrial tachycardia, heart rate increased  i Includes blood pressure increased, diastolic hypertension, hypertension, systolic hypertension  j Includes dyspnoea, dyspnoea exertional  k Includes interstitial lung disease, pneumonitis  l Includes abdominal discomfort, abdominal distension, abdominal pain, abdominal pain lower, abdominal pain upper, epigastric discomfort  m Includes aphthous stomatitis, stomatitis, aphthous ulcer, mouth ulceration, oral mucosal blistering  n Includes dermatitis acneiform, erythema, exfoliative rash, rash, rash erythematous, rash macular, rash maculo‑papular, rash papular, rash pruritic, rash pustular, dermatitis, dermatitis allergic, dermatitis contact, generalised erythema, rash follicular, urticaria, drug eruption, toxic skin eruption  o Includes pruritus, pruritus allergic, pruritus generalised, pruritus genital, vulvovaginal pruritus  p Includes photosensitivity reaction, polymorphic light eruption, solar dermatitis  q Includes musculoskeletal pain, myalgia, muscle spasms, muscle tightness, muscle twitching, musculoskeletal discomfort  r Includes asthenia, fatigue  s Includes eyelid oedema, face oedema, oedema peripheral, periorbital oedema, swelling face, generalised oedema, peripheral swelling, angioedema, lip swelling, periorbital swelling, skin swelling, swelling of eyelid  t Includes blood cholesterol increased, hypercholesterolemia | | | |

Description of selected adverse reactions

*Pulmonary adverse reactions*

In ALTA 1L, 2.9% of patients experienced any Grade ILD/pneumonitis early in treatment (within 8 days), with Grade 3‑4 ILD/pneumonitis in 2.2% of patients. There were no fatal ILD/pneumonitis. Additionally, 3.7% of patients experienced pneumonitis later in treatment.

In ALTA, 6.4% of patients experienced pulmonary adverse reactions of any grade, including ILD/pneumonitis, pneumonia and dyspnoea, early in treatment (within 9 days, median onset: 2 days); 2.7% of patients had Grade 3‑4 pulmonary adverse reactions and 1 patient (0.5%) had fatal pneumonia. Following Grade 1‑2 pulmonary adverse reactions, treatment with Alunbrig was either interrupted and then restarted or the dose was reduced. Early pulmonary adverse reactions also occurred in a dose escalation study in patients (N = 137) (Study 101) including three fatal cases (hypoxia, acute respiratory distress syndrome and pneumonia). Additionally, 2.3% of patients in ALTA experienced pneumonitis later in treatment, with 2 patients having Grade 3 pneumonitis (see sections 4.2 and 4.4).

*Elderly*

Early pulmonary adverse reaction was reported in 10.1% of patients ≥ 65 years of age compared with 3.1% of patients < 65 years of age.

*Hypertension*

Hypertension was reported in 30% of patients treated with Alunbrig at the 180 mg regimen with 11% having Grade 3 hypertension. Dose reduction for hypertension occurred in 1.5% at the 180 mg regimen. Mean systolic and diastolic blood pressure, in all patients, increased over time (see sections 4.2 and 4.4).

*Bradycardia*

Bradycardia was reported in 8.4% of patients treated with Alunbrig at the 180 mg regimen.

Heart rates of less than 50 beats per minute (bpm) were reported in 8.4% of patients at the 180 mg regimen. (see sections 4.2 and 4.4).

*Visual disturbance*

Visual disturbance adverse reactions were reported in 14% of patients treated with Alunbrig at the 180 mg regimen. Of these, three Grade 3 adverse reactions (1.1%) including macular oedema and cataract were reported.

Dose reduction for visual disturbance occurred in two patients (0.7%) at the 180 mg regimen (see sections 4.2 and 4.4).

*Peripheral neuropathy*

Peripheral neuropathy adverse reactions were reported in 20% of patients treated at the 180 mg regimen. Thirty‑three percent of patients had resolution of all peripheral neuropathy adverse reactions. The median duration of peripheral neuropathy adverse reactions was 6.6 months, with a maximum duration of 28.9 months.

*Creatine phosphokinase (CPK) elevation*

In ALTA 1L and ALTA, elevations of CPK were reported in 64% of patients treated with Alunbrig at the 180 mg regimen. The incidence of Grade 3‑4 elevations of CPK was 18%. The median time to onset for CPK elevations was 28 days.

Dose reduction for CPK elevation occurred in 10% of patients at the 180 mg regimen (see sections 4.2 and 4.4).

*Elevations of pancreatic enzymes*

Elevations of amylase and lipase were reported in 47% and 54% of patients treated with Alunbrig, respectively at the 180 mg regimen. For elevations to Grade 3 and 4, the incidences for amylase and lipase were 7.7% and 15%, respectively. The median time to onset for amylase elevations and lipase elevations was 16 days and 29 days, respectively.

Dose reduction for elevation of lipase and amylase occurred in 4.7% and 2.9% of patients, respectively at the 180 mg regimen (see sections 4.2 and 4.4).

*Elevation of hepatic enzymes*

Elevations of ALT and AST were reported in 49% and 68% of patients treated with Alunbrig, respectively at the 180 mg regimen. For elevations to Grade 3 and 4, the incidences for ALT and AST were 4.7% and 3.6%, respectively.

Dose reduction for elevation of ALT and AST occurred in 0.7% and 1.1% of patients, respectively at the 180 mg regimen (see sections 4.2 and 4.4).

*Hyperglycaemia*

Sixty one percent of patients experienced hyperglycaemia. Grade 3 hyperglycemia occurred in 6.6% of patients.

No patients had dose reductions due to hyperglycaemia.

*Photosensitivity and photodermatosis*

A pooled analysis from seven clinical trials with data from 804 patients, treated with Alunbrig at different dosing regimens, showed that photosensitivity and photodermatosis was reported in 5.8% of patients and Grade 3‑4 occurred in 0.7% of patients. Dose reduction occurred in 0.4% of patients (see sections 4.2 and 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](http://www.ema.europa.eu/docs/en_GB/document_library/Template_or_form/2013/03/WC500139752.doc).

**4.9 Overdose**

There is no specific antidote for overdose with Alunbrig. In the event of an overdose, the patient should be monitored for adverse reactions (see section 4.8) and appropriate supportive care should be provided.

**5. PHARMACOLOGICAL PROPERTIES**

**5.1 Pharmacodynamic properties**

Pharmacotherapeutic group: antineoplastic agent, protein kinase inhibitors, ATC code: L01ED04

Mechanism of action

Brigatinib is a tyrosine kinase inhibitor that targets ALK, c‑ros oncogene 1 (ROS1), and insulin‑like growth factor 1 receptor (IGF‑1R). Brigatinib inhibited autophosphorylation of ALK and ALK‑mediated phosphorylation of the downstream signalling protein STAT3 in *in vitro* and *in vivo* assays.

Brigatinib inhibited the *in vitro* proliferation of cell lines expressing EML4‑ALK and NPM‑ALK fusion proteins and demonstrated dose‑dependent inhibition of EML4‑ALK‑positive NSCLC xenograft growth in mice. Brigatinib inhibited the *in vitro* and *in vivo* viability of cells expressing mutant forms of EML4‑ALK associated with resistance to ALK inhibitors, including G1202R and L1196M.

Cardiac electrophysiology

In Study 101, the QT interval prolongation potential of Alunbrig was assessed in 123 patients with advanced malignancies following once daily brigatinib doses of 30 mg to 240 mg. The maximum mean QTcF (corrected QT by the Fridericia method) change from baseline was less than 10 msec. An exposure‑QT analysis suggested no concentration‑dependent QTc interval prolongation.

Clinical efficacy and safety

*ALTA 1L*

The safety and efficacy of Alunbrig was evaluated in a randomised (1:1), open‑label, multicentre trial (ALTA 1L) in 275 adult patients with advanced ALK‑positive NSCLC who had not previously received an ALK‑targeted therapy. Eligibility criteria permitted enrolment of patients with a documented ALK rearrangement based on a local standard of care testing and an ECOG Performance status of 0‑2. Patients were allowed to have up to 1 prior regimen of chemotherapy in the locally advanced or metastatic setting. Neurologically stable patients with treated or untreated central nervous system (CNS) metastases, including leptomeningeal metastases, were eligible. Patients with a history of pulmonary interstitial disease, drug‑related pneumonitis, or radiation pneumonitis were excluded.

Patients were randomised in a 1:1 ratio to receive Alunbrig 180 mg once daily with a 7‑day lead‑in at 90 mg once daily (N = 137) or crizotinib 250 mg orally twice daily (N = 138). Randomisation was stratified by brain metastases (present, absent) and prior chemotherapy use for locally advanced or metastatic disease (yes, no).

Patients in the crizotinib arm who experienced disease progression were offered crossover to receive treatment with Alunbrig. Among all 121 patients who were randomised to the crizotinib arm and discontinued study treatment by the time of the final analysis, 99 (82%) patients received subsequent ALK tyrosine kinase inhibitors (TKIs). Eighty (66%) patients who were randomised to the crizotinib arm received subsequent Alunbrig treatment, including 65 (54%) patients who crossed over in the study.

The major outcome measure was progression‑free survival (PFS) according to Response Evaluation Criteria in Solid Tumours (RECIST v1.1) as evaluated by a Blinded Independent Review Committee (BIRC). Additional outcome measures as evaluated by the BIRC include confirmed objective response rate (ORR), duration of response (DOR), time to response, disease control rate (DCR), intracranial ORR, intracranial PFS, and intracranial DOR. Investigator‑assessed outcomes include PFS and overall survival.

Baseline demographics and disease characteristics in ALTA 1L were median age 59 years old (range 27 to 89; 32% 65 and over), 59% White and 39% Asian, 55% female, 39% ECOG PS 0, and 56% ECOG PS 1, 58% never smokers, 93% Stage IV disease, 96% adenocarcinoma histology, 30% CNS metastases at baseline, 14% prior radiotherapy to the brain, and 27% prior chemotherapy. Sites of extra‑thoracic metastases include brain (30% of patients), bone (31% of patients), and liver (20% of patients). The median relative dose intensity was 97% for Alunbrig and 99% for crizotinib.

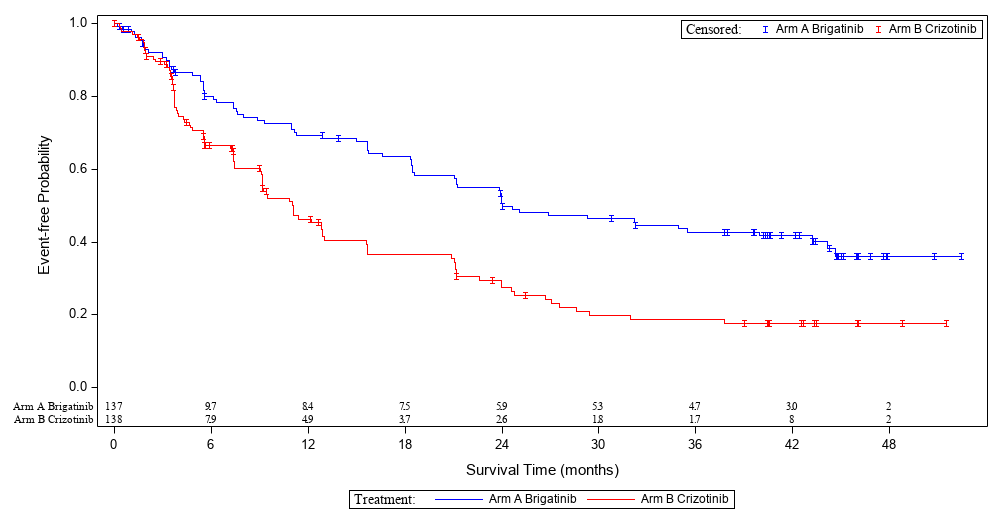
At the primary analysis performed at a median follow‑up duration of 11 months in the Alunbrig arm, the ALTA 1L study met its primary endpoint demonstrating a statistically significant improvement in PFS by BIRC.

A protocol‑specified interim analysis with cut‑off date of 28 June 2019 was performed at a median follow‑up duration of 24.9 months in the Alunbrig arm. The median PFS by BIRC in the ITT population was 24 months in the Alunbrig arm and 11 months in the crizotinib arm (HR = 0.49 [95% CI (0.35, 0.68)], p < 0.0001).

The results from the protocol‑specified final analysis with last patient last contact date of 29 January 2021 performed at a median follow‑up duration of 40.4 months in the Alunbrig arm are presented below.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Table 4: Efficacy results in ALTA IL (ITT population)** | | | | |
| **Efficacy parameters** | **Alunbrig**  **N = 137** | | **Crizotinib**  **N = 138** | |
| **Median duration of follow-up (months)a** | 40.4  (range: 0.0–52.4) | | 15.2  (range: 0.1–51.7) | |
| ***Primary efficacy parameters*** | | | | |
| **PFS (BIRC)** | | | | |
| Number of patients with events, n (%) | 73 (53.3%) | | 93 (67.4%) | |
| Progressive disease, n (%) | 66 (48.2%)**b** | | 88 (63.8%)**c** | |
| Death, n (%) | 7 (5.1%) | | 5 (3.6%) | |
| Median (in months) (95% CI) | 24.0 (18.5, 43.2) | | 11.1 (9.1, 13.0) | |
| Hazard ratio (95% CI) | 0.48 (0.35, 0.66) | | | |
| Log-rank p-valued | < 0.0001 | | | |
| ***Secondary efficacy parameters*** | | | | |
| **Confirmed objective response rate (BIRC)** | | | | |
| Responders, n (%)  (95% CI) | 102 (74.5%)  (66.3, 81.5) | | 86 (62.3%)  (53.7, 70.4) | |
| p-value**d,e** | 0.0330 | | | |
| Complete response, % | 24.1% | | 13.0% | |
| Partial response, % | 50.4% | | 49.3% | |
| **Duration of confirmed response (BIRC)** | | | | |
| Median (months) (95% CI) | 33.2 (22.1, NE) | 13.8 (10.4, 22.1) | | |
| **Overall survivalf** | | | | |
| Number of events, n (%) | 41 (29.9%) | 51 (37.0%) | | |
| Median (in months) (95% CI) | NE (NE, NE) | NE (NE, NE) | | |
| Hazard ratio (95% CI) | 0.81 (0.53, 1.22) | | | |
| Log-rank p-valued | 0.3311 | | | |
| Overall survival at 36 months | 70.7% | | | 67.5% |
| BIRC = Blinded Independent Review Committee; NE = Not Estimable; CI = Confidence Interval  Results in this table are based on final efficacy analysis with last patient last contact date of 29 January 2021.  a duration of follow up for the whole study  b includes 3 patients with palliative radiotherapy to the brain  c includes 9 patients with palliative radiotherapy to the brain  d Stratified by presence of iCNS metastases at baseline and prior chemotherapy for locally advanced or metastatic disease for log-rank test and Cochran Mantel-Haenszel test, respectively  e From a Cochran Mantel-Haenszel test  f Patients in the crizotinib arm who experienced disease progression were offered crossover to receive treatment with Alunbrig. | | | | |

**Figure 1: Kaplan‑Meier plot of progression‑free survival by BIRC in ALTA 1L**



Results in this figure are based on final efficacy analysis with last patient last contact date of 29 January 2021.

BIRC assessment of intracranial efficacy according to RECIST v1.1 in patients with any brain metastases and patients with measurable brain metastases (≥ 10 mm in longest diameter) at baseline are summarised in Table 5.

| **Table** **5: BIRC‑assessed intracranial efficacy in patients in ALTA 1L** | | |
| --- | --- | --- |
| **Efficacy parameters** | **Patients with measurable brain metastases at baseline** | |
| **Alunbrig**  **N = 18** | **Crizotinib**  **N = 23** |
| **Confirmed intracranial objective response rate** | | |
| Responders, n (%)  (95% CI) | 14 (77.8%)  (52.4, 93.6) | 6 (26.1%)  (10.2, 48.4) |
| p-valuea,b | 0.0014 | |
| Complete response % | 27.8% | 0.0% |
| Partial response % | 50.0% | 26.1% |
| **Duration of confirmed intracranial response**c | | |
| Median (months) (95% CI) | 27.9 (5.7, NE) | 9.2 (3.9, NE) |
|  | **Patients with any brain metastases at baseline** | |
| **Alunbrig**  **N = 47** | **Crizotinib**  **N = 49** |
| **Confirmed intracranial objective response rate** | | |
| Responders, n (%)  (95% CI) | 31 (66.0%)  (50.7, 79.1) | 7 (14.3%)  (5.9, 27.2) |
| p-valuea,b | < 0.0001 | |
| Complete response (%) | 44.7% | 2.0% |
| Partial response (%) | 21.3% | 12.2% |
| **Duration of confirmed intracranial response**c | | |
| Median (months) (95% CI) | 27.1 (16.9, 42.8) | 9.2 (3.9, NE) |
| **Intracranial PFS**d |  |  |
| Number of patients with events, n (%) | 27 (57.4%) | 35 (71.4%) |
| Progressive disease, n (%) | 27 (57.4%)e | 32 (65.3%)f |
| Death, n (%) | 0 (0.0%) | 3 (6.1%) |
| Median (in months) (95% CI) | 24.0 (12.9, 30.8) | 5.5 (3.7, 7.5) |
| Hazard ratio (95% CI) | 0.29 (0.17, 0.51) | |
| Log-rank p-valuea | < 0.0001 | |
| CI = Confidence Interval; NE = Not Estimable  Results in this table are based on final efficacy analysis with last patient last contact date of 29 January 2021.  a Stratified by presence prior chemotherapy for locally advanced or metastatic disease for log-rank test and Cochran Mantel‑Haenszel test, respectively  bFrom a Cochran Mantel‑Haenszel test  c measured from date of first confirmed intracranial response until date of intracranial disease progression (new intracranial lesions, intracranial target lesion diameter growth ≥ 20% from nadir, or unequivocal progression of intracranial nontarget lesions) or death or censoring  d measured from date of randomisation until date of intracranial disease progression (new intracranial lesions, intracranial target lesion diameter growth ≥ 20% from nadir, or unequivocal progression of intracranial nontarget lesions) or death or censoring.  e includes 1 patient with palliative radiotherapy to the brain  f includes 3 patients with palliative radiotherapy to the brain | | |

*ALTA*

The safety and efficacy of Alunbrig was evaluated in a randomised (1:1), open‑label, multicenter trial (ALTA) in 222 adult patients with locally advanced or metastatic ALK‑positive NSCLC who had progressed on crizotinib. Eligibility criteria permitted enrolment of patients with a documented ALK rearrangement based on a validated test, ECOG Performance Status of 0‑2, and prior chemotherapy. Additionally, patients with central nervous system (CNS) metastases were included, provided they were neurologically stable and did not require an increasing dose of corticosteroids. Patients with a history of pulmonary interstitial disease or drug‑related pneumonitis were excluded.

Patients were randomised in a 1:1 ratio to receive Alunbrig either 90 mg once daily (90 mg regimen, N = 112) or 180 mg once daily with 7‑day lead‑in at 90 mg once daily (180 mg regimen, N = 110). The median duration of follow‑up was 22.9 months. Randomisation was stratified by brain metastases (present, absent) and best prior response to crizotinib therapy (complete or partial response, any other response/unknown).

The major outcome measure was confirmed objective response rate (ORR) according to Response Evaluation Criteria in Solid Tumours (RECIST v1.1) as evaluated by investigator. Additional outcome measures included confirmed ORR as evaluated by an Independent Review Committee (IRC); time to response; progression free survival (PFS); duration of response (DOR); overall survival; and intracranial ORR and intracranial DOR as evaluated by an IRC.

Baseline demographics and disease characteristics in ALTA were median age 54 years old (range 18 to 82; 23% 65 and over), 67% White and 31% Asian, 57% female, 36% ECOG PS 0 and 57% ECOG PS 1, 7% ECOG PS2, 60% never smoker, 35% former smoker, 5% current smoker, 98% Stage IV, 97% adenocarcinoma, and 74% prior chemotherapy. The most common sites of extra‑thoracic metastasis included 69% brain (of whom 62% had received prior radiation to the brain), 39% bone, and 26% liver.

Efficacy results from ALTA analysis are summarised in Table 6. and the Kaplan‑Meier (KM) curve for investigator‑assessed PFS is shown in Figure 2

**Table 6: Efficacy results in ALTA (ITT population)**

| **Efficacy parameter** | **Investigator assessment** | | **IRC assessment** | |
| --- | --- | --- | --- | --- |
| **90 mg regimen\* N = 112** | **180 mg**  **regimen**† **N = 110** | **90 mg**  **regimen\* N = 112** | **180 mg**  **regimen**† **N = 110** |
| **Objective response rate** | | | | |
| (%) | 46% | 56% | 51% | 56% |
| CI‡ | (35, 57) | (45, 67) | (41, 61) | (47, 66) |
| **Time to response** | | | | |
| Median (months) | 1.8 | 1.9 | 1.8 | 1.9 |
| **Duration of response** | | | | |
| Median (months) | 12.0 | 13.8 | 16.4 | 15.7 |
| 95% CI | (9.2,17.7) | (10.2,19.3) | (7.4, 24.9) | (12.8, 21.8) |
| **Progression‑free survival** | | | | |
| Median (months) | 9.2 | 15.6 | 9.2 | 16.7 |
| 95% CI | (7.4, 11.1) | (11.1, 21) | (7.4, 12.8) | (11.6, 21.4) |
| **Overall survival** | | | | |
| Median (months) | 29.5 | 34.1 | NA | NA |
| 95% CI | (18.2, NE) | (27.7, NE) | NA | NA |
| 12‑month survival probability (%) | 70.3% | 80.1% | NA | NA |

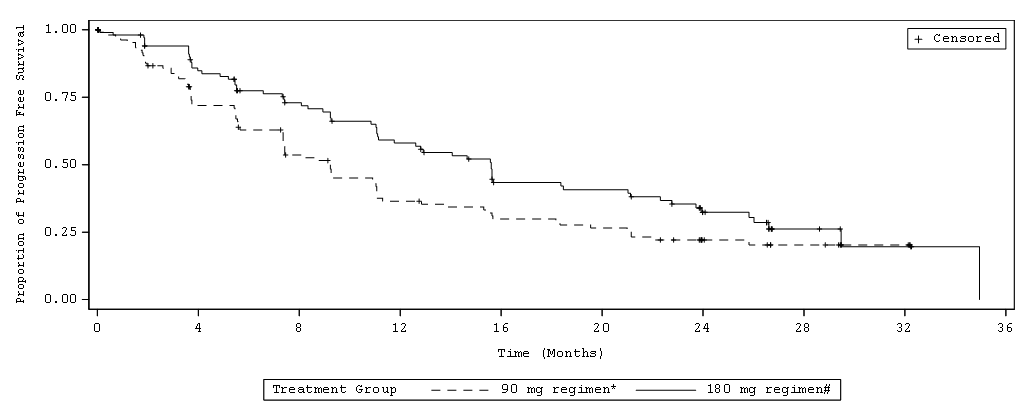
CI = Confidence Interval; NE = Not Estimable; NA = Not Applicable

\*90 mg once daily regimen

†180 mg once daily with 7‑day lead‑in at 90 mg once daily

‡Confidence Interval for investigator assessed ORR is 97.5% and for IRC assessed ORR is 95%

**Figure 2:** **Investigator‑assessed systemic progression‑free survival: ITT population by treatment arm (ALTA)**

Abbreviations: ITT = Intent‑to‑treat

Note: Progression‑Free survival was defined as time from initiation of treatment until the date at which disease progression was first evident or death, whichever comes first.

\*90 mg once daily regimen

†180 mg once daily with 7‑day lead‑in at 90 mg once daily

IRC assessments of intracranial ORR and duration of intracranial response in patients from ALTA with measurable brain metastases (≥ 10 mm in longest diameter) at baseline are summarised in Table 7.

**Table 7 Intracranial efficacy in patients with measurable brain metastases at baseline in ALTA**

| **IRC‑assessed efficacy parameter** | **Patients with measurable**  **brain metastases at baseline** | |
| --- | --- | --- |
| **90 mg regimen**\* **(N = 26)** | **180 mg regimen**† **(N = 18)** |
| **Intracranial objective response rate** | | |
| (%) | 50% | 67% |
| 95% CI | (30, 70) | (41, 87) |
| **Intracranial disease control rate** | | |
| (%) | 85% | 83% |
| 95% CI | (65, 96) | (59, 96) |
| **Duration of intracranial response**‡**,** | | |
| Median (months) | 9.4 | 16.6 |
| 95% CI | (3.7, 24.9) | (3.7, NE) |

% CI = Confidence Interval; NE = Not Estimable

\*90 mg once daily regimen

†180 mg once daily with 7‑day lead‑in at 90 mg once daily

‡Events include intracranial disease progression (new lesions, intracranial target lesion diameter growth ≥ 20% from nadir, or unequivocal progression of intracranial non‑target lesions) or death.

In patients with any brain metastases at baseline, intracranial disease control rate was 77.8% (95% CI 67.2‑86.3) in the 90 mg arm (N = 81) and 85.1% (95% CI 75‑92.3) in the 180 mg arm (N = 74).

*Study 101*

In a separate dose finding study, 25 patients with ALK‑positive NSCLC that progressed on crizotinib were administered Alunbrig at 180 mg once daily with 7‑day lead‑in at 90 mg once daily regimen. Of these, 19 patients had an investigator‑assessed confirmed objective response (76%; 95% CI: 55, 91) and the KM estimate median duration of response among the 19 responders was 26.1 months (95% CI: 7.9, 26.1). The KM median PFS was 16.3 months (95% CI: 9.2, NE) and the 12‑month probability of overall survival was 84.0% (95% CI: 62.8, 93.7).

Paediatric population

The European Medicines Agency has waived the obligation to submit the results of studies with Alunbrig in all subsets of the paediatric population in lung carcinoma (small cell and non‑small cell carcinoma) (see section 4.2 for information on paediatric use).

**5.2 Pharmacokinetic properties**

Absorption

In Study 101, following administration of a single oral dose of brigatinib (30‑240 mg) in patients, the median time to peak concentration (Tmax) was 1‑4 hours postdose. After a single dose and at steady state, systemic exposure was dose proportional over the dose range of 60‑240 mg once daily. Modest accumulation was observed upon repeated dosing (geometric mean accumulation ratio: 1.9 to 2.4). The geometric mean steady state Cmax of brigatinib at doses of 90 mg and 180 mg once daily was 552 and 1,452 ng/mL, respectively, and the corresponding AUC0‑τ was 8,165 and 20,276 h∙ng/mL, respectively. Brigatinib is a substrate of the transporter proteins P‑gp and BCRP.

In healthy subjects, compared to overnight fasting, a high fat meal reduced brigatinib Cmax by 13% with no effect on AUC. Brigatinib can be administered with or without food.

Distribution

Brigatinib was moderately bound (91%) to human plasma proteins and binding was not concentration‑dependent. The blood‑to‑plasma concentration ratio is 0.69. In patients given brigatinib 180 mg once daily, the geometric mean apparent volume of distribution (Vz/F) of brigatinib at steady state was 307 L, indicating moderate distribution into tissues.

Biotransformation

*In vitro* studies demonstrated that brigatinib is primarily metabolised by CYP2C8 and CYP3A4, and to a much lesser extent by CYP3A5.

Following oral administration of a single 180 mg dose of [14C]brigatinib to healthy subjects, N‑demethylation and cysteine conjugation were the two major metabolic clearance pathways. In urine and faeces combined, 48%, 27%, and 9.1% of the radioactive dose was excreted as unchanged brigatinib, N‑desmethyl brigatinib (AP26123), and brigatinib cysteine conjugate, respectively. Unchanged brigatinib was the major circulating radioactive component (92%) along with AP26123 (3.5%), the primary metabolite also observed *in vitro*. In patients, at steady state, the plasma AUC of AP26123 was < 10% of brigatinib exposure. In *in vitro* kinase and cellular assays, the metabolite, AP26123, inhibited ALK with approximately 3‑fold lower potency than brigatinib.

Elimination

In patients given brigatinib 180 mg once daily, the geometric mean apparent oral clearance (CL/F) of brigatinib at steady state was 8.9 L/h and the median plasma elimination half‑life was 24 h.

The primary route of excretion of brigatinib is in faeces. In six healthy male subjects given a single 180 mg oral dose of [14C]brigatinib, 65% of the administered dose was recovered in faeces and 25% of the administered dose was recovered in urine. Unchanged brigatinib represented 41% and 86% of the total radioactivity in faeces and urine, respectively, the remainder being metabolites.

Specific populations

*Hepatic impairment*

The pharmacokinetics of brigatinib was characterised in healthy subjects with normal hepatic function (N = 9), and patients with mild hepatic impairment (Child‑Pugh class A, N = 6), moderate hepatic impairment (Child‑Pugh class B, N = 6), or severe hepatic impairment (Child‑Pugh class C, N = 6). The pharmacokinetics of brigatinib was similar between healthy subjects with normal hepatic function and patients with mild (Child‑Pugh class A) or moderate (Child‑Pugh class B) hepatic impairment. Unbound AUC0‑INF was 37% higher in patients with severe hepatic impairment (Child‑Pugh class C) as compared to healthy subjects with normal hepatic function (see section 4.2).

*Renal impairment*

The pharmacokinetics of brigatinib is similar in patients with normal renal function and in patients with mild or moderate renal impairment (eGFR ≥ 30 mL/min) based on the results of population pharmacokinetic analyses. In a pharmacokinetic study, unbound AUC0‑INF was 94% higher in patients with severe renal impairment (eGFR < 30 mL/min, N = 6) as compared to patients with normal renal function (eGFR ≥ 90 mL/min, N = 8) (see section 4.2).

*Race and gender*

Population pharmacokinetic analyses showed that race and gender had no impact on the pharmacokinetics of brigatinib.

*Age, body weight, and albumin concentrations*

The population pharmacokinetic analyses showed that body weight, age, and albumin concentration had no clinically relevant impact on the pharmacokinetics of brigatinib.

**5.3 Preclinical safety data**

Safety pharmacology studies with brigatinib identified potential for pulmonary effects (altered respiration rate; 1‑2 times the human Cmax), cardiovascular effects (altered heart rate and blood pressure; at 0.5 times the human Cmax), and renal effects (reduced renal function; at 1‑2.5 times the human Cmax), but did not indicate any potential for QT prolongation or neurofunctional effects.

Adverse reactions seen in animals at exposure levels similar to clinical exposure levels with possible relevance to clinical use were as follows: gastrointestinal system, bone marrow, eyes, testes, liver, kidney, bone, and heart. These effects were generally reversible during the non‑dosing recovery period; however, effects in the eyes and testes were notable exceptions due to lack of recovery.

In repeated dose toxicity studies, lung changes (foamy alveolar macrophages) were noted in monkeys at ≥ 0.2 times the human AUC; however, these were minimal and similar to those reported as background findings in naive monkeys, and there was no clinical evidence of respiratory distress in these monkeys.

Carcinogenicity studies have not been performed with brigatinib.

Brigatinib was not mutagenic *in vitro* in the bacterial reverse mutation (Ames) or the mammalian cell chromosomal aberration assays, but slightly increased the number of micronuclei in a rat bone marrow micronucleus test. The mechanism of micronucleus induction was abnormal chromosome segregation (aneugenicity) and not a clastogenic effect on chromosomes. This effect was observed at approximately five fold the human exposure at the 180 mg once daily dose.

Brigatinib may impair male fertility. Testicular toxicity was observed in repeat‑dose animal studies. In rats, findings included lower weight of testes, seminal vesicles and prostate gland, and testicular tubular degeneration; these effects were not reversible during the recovery period. In monkeys, findings included reduced size of testes along with microscopic evidence of hypospermatogenesis; these effects were reversible during the recovery period. Overall, these effects on the male reproductive organs in rats and monkeys occurred at exposures ≥ 0.2‑times the AUC observed in patients at the 180 mg once daily dose. No apparent adverse effects on female reproductive organs were observed in general toxicology studies in rats and monkeys.

In an embryo‑foetal development study in which pregnant rats were administered daily doses of brigatinib during organogenesis; dose‑related skeletal anomalies were observed at doses as low as approximately 0.7‑times the human exposure by AUC at the 180 mg once daily dose. Findings included embryo‑lethality, reduced foetal growth, and skeletal variations.

**6. PHARMACEUTICAL PARTICULARS**

**6.1 List of excipients**

Tablet core

Lactose monohydrate

Microcrystalline cellulose

Sodium starch glycolate (type A)

Silica colloidal hydrophobic

Magnesium stearate

Tablet coating

Talc

Macrogol

Polyvinyl alcohol

Titanium dioxide

**6.2 Incompatibilities**

Not applicable.

**6.3 Shelf life**

3 years

**6.4 Special precautions for storage**

This medicinal product does not require any special storage conditions.

**6.5 Nature and contents of container**

Alunbrig 30 mg film‑coated tablets

Round wide mouth high density polyethylene (HDPE) bottles with two‑piece polypropylene child resistant screw cap closures with foil induction seal liner, containing either 60 or 120 film‑coated tablets, together with one HDPE canister containing a molecular sieve desiccant.

Clear thermoformable poly‑chloro‑tri‑fluoro‑ethylene (PCTFE) blister with heat sealable paper‑laminated foil lidding in a carton, containing either 28, 56 or 112 film‑coated tablets.

Alunbrig 90 mg film‑coated tablets

Round wide mouth high density polyethylene (HDPE) bottles with two‑piece polypropylene child resistant screw cap with foil induction seal liner closures, containing either 7 or 30 film‑coated tablets, together with one HDPE canister containing a molecular sieve desiccant.

Clear thermoformable poly‑chloro‑tri‑fluoro‑ethylene (PCTFE) blister with heat sealable paper‑laminated foil lidding in a carton, containing either 7 or 28 film‑coated tablets.

Alunbrig 180 mg film‑coated tablets

Round wide mouth high density polyethylene (HDPE) bottles with two‑piece polypropylene child resistant screw cap with foil induction seal liner closures, containing 30 film‑coated tablets, together with one HDPE canister containing a molecular sieve desiccant.

Clear thermoformable poly‑chloro‑tri‑fluoro‑ethylene (PCTFE) blister with heat sealable paper‑laminated foil lidding in a carton, containing 28 film‑coated tablets.

Treatment initiation pack Alunbrig 90 mg and 180 mg film‑coated tablets

Each pack consists of an outer carton with two inner cartons containing:

* Alunbrig 90 mg film‑coated tablets

1 clear thermoformable poly‑chloro‑tri‑fluoro‑ethylene (PCTFE) blister with heat sealable paper‑laminated foil lidding in a carton, containing 7 film‑coated tablets.

* Alunbrig 180 mg film‑coated tablets

3 clear thermoformable poly‑chloro‑tri‑fluoro‑ethylene (PCTFE) blisters with heat sealable paper‑laminated foil lidding in a carton, containing 21 film‑coated tablets.

Not all pack sizes may be marketed.

**6.6 Special precautions for disposal and other handling**

Patients should be advised to keep the desiccant canister in the bottle and not to swallow it.

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

**7. MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**8. MARKETING AUTHORISATION NUMBER(S)**

Alunbrig 30 mg film‑coated tablets

EU/1/18/1264/001 60 tablets in bottle

EU/1/18/1264/002 120 tablets in bottle

EU/1/18/1264/011 28 tablets in carton

EU/1/18/1264/003 56 tablets in carton

EU/1/18/1264/004 112 tablets in carton

Alunbrig 90 mg film‑coated tablets

EU/1/18/1264/005 7 tablets in bottle

EU/1/18/1264/006 30 tablets in bottle

EU/1/18/1264/007 7 tablets in carton

EU/1/18/1264/008 28 tablets in carton

Alunbrig 180 mg film‑coated tablets

EU/1/18/1264/009 30 tablets in bottle

EU/1/18/1264/010 28 tablets in carton

Alunbrig treatment initiation pack

EU/1/18/1264/012 7 x 90 mg + 21 x 180 mg tablets in carton

**9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION**

Date of first authorisation: 22 November 2018

Date of latest renewal: 24 July 2023

**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. MANUFACTURERS 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. MANUFACTURERS RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturers responsible for batch release

Takeda Austria GmbH

St. Peter‑Strasse 25

4020 Linz

Austria

Takeda Ireland Limited  
Bray Business Park  
Kilruddery   
Co. Wicklow   
A98 CD36  
Ireland

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.
* **Additional risk minimisation measures**

Not applicable.

**ANNEX III**

**LABELLING AND PACKAGE LEAFLET**

# A. LABELLING

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING AND THE IMMEDIATE PACKAGING**

**OUTER CARTON AND BOTTLE LABEL**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 30 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 30 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

60 film‑coated tablets

120 film‑coated tablets

**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**

Outer carton:

Do not swallow the desiccant canister found in the bottle.

**8. EXPIRY DATE**

EXP

**9. SPECIAL STORAGE CONDITIONS**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/001 60 tablets

EU/1/18/1264/002 120 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Outer Carton:

Alunbrig 30 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

Outer Carton:

PC

SN

NN

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 30 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 30 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

28 film‑coated tablets

56 film‑coated tablets

112 film‑coated tablets

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/011 28 tablets

EU/1/18/1264/003 56 tablets

EU/1/18/1264/004 112 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 30 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

PC

SN

NN

**MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS**

**BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 30 mg film‑coated tablets

brigatinib

**2. NAME OF THE MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S (as Takeda logo)

**3. EXPIRY DATE**

EXP

**4. BATCH NUMBER**

Lot

**5. OTHER**

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING AND THE IMMEDIATE PACKAGING**

**OUTER CARTON AND BOTTLE LABEL**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 90 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

7 film‑coated tablets

30 film‑coated tablets

**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**

Outer carton:

Do not swallow the desiccant canister found in the bottle.

**8. EXPIRY DATE**

EXP

**9. SPECIAL STORAGE CONDITIONS**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/005 7 tablets

EU/1/18/1264/006 30 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Outer Carton:

Alunbrig 90 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

Outer Carton

PC

SN

NN

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 90 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

7 film‑coated tablets

28 film‑coated tablets

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/007 7 tablets

EU/1/18/1264/008 28 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 90 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

PC

SN

NN

**MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS**

**BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

brigatinib

**2. NAME OF THE MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S (as Takeda logo)

**3. EXPIRY DATE**

EXP

**4. BATCH NUMBER**

Lot

**5. OTHER**

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR TREATMENT INITIATION PACK (INCLUDING BLUE BOX)**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each 90 mg film‑coated tablet contains 90 mg brigatinib.

Each 180 mg film‑coated tablet contains 180 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

Treatment initiation packEach pack contains two cartons in an outer carton.

7 film‑coated tablets of Alunbrig 90 mg

21 film‑coated tablets of Alunbrig 180 mg

**5. METHOD AND ROUTE(S) OF ADMINISTRATION**

Read the package leaflet before use.

Oral use

Take only one tablet per day.

Alunbrig 90 mg once daily for the first 7 days, then Alunbrig 180 mg once daily.

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/012 7 x 90 mg + 21 x 180 mg tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 90 mg, 180 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

PC

SN

NN

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**INNER CARTON FOR TREATMENT INITIATION PACK – 7 TABLETS, 90 MG – 7 DAY TREATMENT (WITHOUT BLUE BOX)**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 90 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

Treatment initiation packEach pack contains 7 film‑coated tablets of Alunbrig 90 mg

**5. METHOD AND ROUTE(S) OF ADMINISTRATION**

Read the package leaflet before use.

Oral use

Take only one tablet per day.

Day 1 to Day 7

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/012 7 x 90 mg + 21 x 180 mg tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 90 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

**MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS**

**BLISTER – TREATMENT INITIATION PACK – 90 MG**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 90 mg film‑coated tablets

brigatinib

**2. NAME OF THE MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S (as Takeda logo)

**3. EXPIRY DATE**

EXP

**4. BATCH NUMBER**

Lot

**5. OTHER**

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**INNER CARTON FOR TREATMENT INITIATION PACK – 21 TABLETS, 180 MG – 21 DAY TREATMENT (WITHOUT BLUE BOX)**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 180 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

Treatment initiation packEach pack contains 21 film‑coated tablets of Alunbrig 180 mg

**5. METHOD AND ROUTE(S) OF ADMINISTRATION**

Read the package leaflet before use.

Oral use

Take only one tablet per day.

Day 8 to Day 28

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/012 7 x 90 mg + 21 x 180 mg tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 180 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

**MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS**

**BLISTER – TREATMENT INITIATION PACK – 180 MG**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. NAME OF THE MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S (as Takeda logo)

**3. EXPIRY DATE**

EXP

**4. BATCH NUMBER**

Lot

**5. OTHER**

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING AND THE IMMEDIATE PACKAGING**

**OUTER CARTON AND BOTTLE LABEL**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 180 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

30 film‑coated tablets

**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**

Outer carton:

Do not swallow the desiccant canister found in the bottle.

**8. EXPIRY DATE**

EXP

**9. SPECIAL STORAGE CONDITIONS**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/009 30 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Outer Carton:

Alunbrig 180 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

Outer Carton

PC

SN

NN

**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film‑coated tablet contains 180 mg brigatinib.

**3. LIST OF EXCIPIENTS**

Contains lactose. See the package leaflet for further information.

**4. PHARMACEUTICAL FORM AND CONTENTS**

Film‑coated tablets

28 film‑coated tablets

**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**

**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**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**12. MARKETING AUTHORISATION NUMBER(S)**

EU/1/18/1264/010 28 tablets

**13. BATCH NUMBER**

Lot

**14. GENERAL CLASSIFICATION FOR SUPPLY**

**15. INSTRUCTIONS ON USE**

**16. INFORMATION IN BRAILLE**

Alunbrig 180 mg

**17. UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.

**18. UNIQUE IDENTIFIER - HUMAN READABLE DATA**

PC

SN

NN

**MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS**

**BLISTER**

**1. NAME OF THE MEDICINAL PRODUCT**

Alunbrig 180 mg film‑coated tablets

brigatinib

**2. NAME OF THE MARKETING AUTHORISATION HOLDER**

Takeda Pharma A/S (as Takeda logo)

**3. EXPIRY DATE**

EXP

**4. BATCH NUMBER**

Lot

**5. OTHER**

# B. PACKAGE LEAFLET

**Package leaflet: Information for the patient**

**Alunbrig 30 mg film‑coated tablets**

**Alunbrig 90 mg film‑coated tablets**

**Alunbrig 180 mg film‑coated tablets**

brigatinib

**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 Alunbrig is and what it is used for

2. What you need to know before you take Alunbrig

3. How to take Alunbrig

4. Possible side effects

5. How to store Alunbrig

6. Contents of the pack and other information

**1. What Alunbrig is and what it is used for**

Alunbrig contains the active substance brigatinib, a type of cancer medicine called a kinase inhibitor. Alunbrig is used to treat adults with advanced stages of a **lung cancer** called non‑small cell lung cancer. It is given to patients whose lung cancer is related to an abnormal form of a gene called anaplastic lymphoma kinase (*ALK*).

**How Alunbrig works**

The abnormal gene produces a protein known as a kinase that stimulates the growth of the cancer cells. Alunbrig blocks the action of this protein and thus slows down the growth and spread of the cancer.

**2. What you need to know before you take Alunbrig**

**Do not take Alunbrig**

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

**Warnings and precautions**

Talk to your doctor before taking Alunbrig or during treatment if you have:

* **lung or breathing problems**

Lung problems, some severe, are more frequent within the first 7 days of treatment. Symptoms may be similar to symptoms from lung cancer. Tell your doctor of any new or worsening symptoms including breathing discomfort, shortness of breath, chest pain, cough and fever.

* **high blood pressure**
* **a slow heartbeat (bradycardia)**
* **vision disturbance**

Inform your doctor of any visual disturbance that occurs during treatment, such as seeing flashes of light, blurry vision or light hurting your eyes.

* **muscle problems**

Report any unexplained muscle pain, tenderness or weakness to your doctor.

* **pancreas problems**

Tell your doctor if you have upper abdominal pain, including abdominal pain that gets worse with eating and may spread to the back, weight loss or nausea.

* **liver problems**

Tell your doctor if you have pain on the right side of your stomach area, yellowing of your skin or the whites of your eyes, or dark urine.

* **high blood sugar**
* **sensitivity to sunlight**

Limit your time in the sun during treatment and for at least 5 days after your last dose. When you are in the sun, wear a hat, protective clothing, a broad‑spectrum Ultraviolet A (UVA)/ Ultraviolet B (UVB) sunscreen and lip balm with a Sun Protection Factor (SPF) of 30 or greater. These will help to protect against potential sunburn.

Tell your doctor if you have kidney problems or you are on dialysis. Symptoms of kidney problems may include, nausea, changes in volume or frequency of urination, abnormal blood tests (see section 4).

Your doctor may need to adjust your treatment or stop Alunbrig temporarily or permanently. See also the beginning of section 4.

**Children and adolescents**

Alunbrig has not been studied in children or adolescents. Treatment with Alunbrig is not recommended in persons under 18 years of age.

**Other medicines and Alunbrig**

Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines.

The following medicines can affect or be affected by Alunbrig:

* **ketoconazole, itraconazole, voriconazole:** medicines to treat fungal infections
* **indinavir, nelfinavir, ritonavir, saquinavir:** medicines to treat HIV infection
* **clarithromycin, telithromycin, troleandomycin:** medicines to treat bacterial infections
* **nefazodone:** a medicine to treat depression
* **St. John’s wort:** a herbal product used to treat depression
* **carbamazepine:** a medicine to treat epilepsy, euphoric/depressive episodes and certain pain conditions
* **phenobarbital, phenytoin:** medicines to treat epilepsy
* **rifabutin, rifampicin:** medicines to treat tuberculosis or certain other infections
* **digoxin:** a medicine to treat heart problems
* **dabigatran:** a medicine to inhibit blood clotting
* **colchicine:** a medicine to treat gout attacks
* **pravastatin, rosuvastatin:** medicines to lower elevated cholesterol levels
* **methotrexate:** a medicine to treat severe joint inflammation, cancer and the skin disease psoriasis
* **sulfasalazine:** a medicine to treat severe bowel and rheumatic joint inflammation
* **efavirenz**, **etravirine:** medicines to treat HIV infection
* **modafinil:** a medicine to treat narcolepsy
* **bosentan:** a medicine to treat pulmonary hypertension
* **nafcillin:** a medicine to treat bacterial infections
* **alfentanil, fentanyl:** medicines to treat pain
* **quinidine:** a medicine to treat irregular heart rhythm
* **cyclosporine, sirolimus, tacrolimus:** medicines to suppress the immune system

**Alunbrig with food and drink**

Avoid any grapefruit products during treatment as they may change the amount of brigatinib in your body.

**Pregnancy**

Alunbrig is **not recommended** during pregnancy unless the benefit outweighs the risk to the baby. If you are pregnant or think you may be pregnant or are planning to have a baby, talk to your doctor to discuss the risks of taking Alunbrig during pregnancy.

Women of childbearing age being treated with Alunbrig should avoid becoming pregnant. Effective non‑hormonal contraception must be used during treatment and for 4 months after stopping Alunbrig. Ask your doctor about the birth control methods that may be right for you.

**Breast‑feeding**

**Do not breast‑feed** during treatment with Alunbrig. It is unknown if brigatinib passes into breast milk and could potentially harm the baby.

**Fertility**

Men receiving treatment with Alunbrig are advised not to father a child during treatment and to use effective contraception during treatment and for 3 months after stopping treatment.

**Driving and using machines**

Alunbrig may cause visual disturbances, dizziness or tiredness. Do not drive or use machines during treatment if such signs occur.

**Alunbrig contains lactose**

If you have been told by your doctor that you have an intolerance to some sugars, contact your doctor before taking this medicine.

**Alunbrig contains** **sodium**

This medicine contains less than 1 mmol sodium (23 mg) per tablet, that is to say essentially ‘sodium‑free’.

**3. How to take Alunbrig**

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

**The recommended dose is**

One 90 mg tablet once daily for the first 7 treatment days; thereafter, one 180 mg tablet once daily.

Do not change the dose without talking to your doctor. Your doctor may adjust your dose according to your needs and this may require use of a 30 mg tablet to achieve the new recommended dose.

**Treatment initiation pack**

At the beginning of your treatment with Alunbrig your doctor may prescribe a treatment initiation pack. To help you start treatment each treatment initiation pack consists of an outer pack with two inner packs containing

* 7 Alunbrig 90 mg film‑coated tablets
* 21 Alunbrig 180 mg film‑coated tablets

The required dose is printed on the treatment initiation pack.

**Method of use**

* Take Alunbrig once daily at the same time each day.
* Swallow the tablets whole, with a glass of water. Do not crush or dissolve the tablets.
* The tablets can be taken with or without food.
* If you vomit after taking Alunbrig, do not take any more tablets until your next scheduled dose.

Do not swallow the desiccant canister contained in the bottle.

**If you take more Alunbrig than you should**

Tell your doctor or pharmacist right away if you have taken more tablets than recommended.

**If you forget to take Alunbrig**

Do not take a double dose to make up for a forgotten dose. Take your next dose at your regular time.

**If you stop taking Alunbrig**

Do not stop taking Alunbrig before talking to your doctor.

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.

**Tell your doctor or pharmacist immediately** if you have any of the following serious side effects:

**Very common** (may affect more than 1 in 10 people):

* **high blood pressure**

Tell your doctor if you get headaches, dizziness, blurred vision, chest pain or shortness of breath.

* **vision problems**

Tell your doctor if you experience any visual disturbances, such as seeing flashes of light, blurry vision or light hurting eyes. Your doctor may stop Alunbrig treatment and refer you to an ophthalmologist.

* **increased blood level of creatine phosphokinase in tests** – may indicate muscle damage, such as of the heart. Tell your doctor if you have any unexplained muscle pain, tenderness or weakness.
* **increased blood levels of amylase or lipase in tests** – may indicate inflammation of the pancreas

Tell your doctor if you have upper abdominal pain, including abdominal pain that gets worse with eating and may spread to the back, weight loss or nausea.

* **increased blood levels of liver enzymes (aspartate aminotransferase, alanine aminotransferase) in tests** ‑may indicate liver cell damage. Tell your doctor if you have pain on the right side of your stomach area, yellowing of your skin or the whites of your eyes, or dark urine.
* **increased blood sugar**

Tell your doctor if you are feeling very thirsty, need to urinate more than usual, feeling very hungry, sick to your stomach, weak or tired, or confused.

**Common** (may affect up to 1 in 10 people):

* **lung inflammation**

Tell your doctor if you have any new or worsening lung or breathing problems, including chest pain, cough, and fever, especially within the first week of taking Alunbrig, as they may be a sign of serious lung problems.

* **slow heartbeat**

Tell your doctor if you have chest pain or discomfort, changes in heartbeat, dizziness, light‑headedness or fainting.

* **sensitivity to sunlight**

Tell your doctor if you develop any skin reaction.

See also section 2, “Warnings and precautions”.

**Uncommon** (may affect up to 1 in 100 people)

* inflammation of pancreas which may cause severe and persistent stomach pain, with or without nausea and vomiting (pancreatitis)

**Other possible side effects are:**

Tell your doctor or pharmacist if you notice any of the following side effects

**Very common** (mayaffect more than 1 in 10 people):

* lung infection (pneumonia)
* cold‑like symptoms (upper respiratory tract infection)
* reduced number of red blood cells (anaemia), in blood tests
* reduced number of white blood cells, called neutrophils and lymphocytes, in blood tests
* increased blood clotting time shown by test of activated partial thromboplastin time
* blood tests may show increased blood level of;

- insulin

- calcium

* blood tests may show reduced blood level of;
* phosphorus
* magnesium
* sodium
* potassium
* decreased appetite
* headache
* symptoms such as numbness, tingling, prickling sensation, weakness or pain in hands or feet (peripheral neuropathy)
* dizziness
* cough
* shortness of breath
* diarrhoea
* nausea
* vomiting
* abdominal (belly) pain
* constipation
* inflammation of the mouth and lips (stomatitis)
* increased level of the enzyme alkaline phosphatase in blood tests– may indicate organ malfunction or injury
* rash
* skin itching
* joint or muscle pain (including muscle spasms)
* increased level of creatinine in blood tests– may indicate reduced kidney function
* fatigue
* tissue swelling caused by excess fluid
* fever

**Common** (may affect up to 1 in 10 people):

* low platelet counts in blood tests, which may increase the risk of bleeding and bruising
* difficulty sleeping (insomnia)
* memory impairment
* change in sense of taste
* abnormal electrical activity of the heart (prolonged electrocardiogram QT interval)
* rapid heartbeat (tachycardia)
* palpitations
* dry mouth
* indigestion
* flatulence
* increased level of lactate dehydrogenase in blood tests – may indicate tissue breakdown
* increased level of bilirubin in blood tests
* dry skin
* musculoskeletal chest pain
* pain in arms and legs
* muscle and joint stiffness
* chest pain and discomfort
* pain
* increased level of cholesterol in blood tests
* weight loss

**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 [Appendix V](http://www.ema.europa.eu/docs/en_GB/document_library/Template_or_form/2013/03/WC500139752.doc). By reporting side effects you can help provide more information on the safety of this medicine.

**5. How to store Alunbrig**

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on either the bottle label or blister and carton after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

Do not throw away any medicines via waste water 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 Alunbrig contains**

* The active substance is brigatinib.

Each 30 mg film‑coated tablet contains 30 mg brigatinib.

Each 90 mg film‑coated tablet contains 90 mg brigatinib.

Each 180 mg film‑coated tablet contains 180 mg brigatinib.

* The other excipients are lactose monohydrate, microcrystalline cellulose, sodium starch glycolate (type A), silica colloidal hydrophobic, magnesium stearate, talc, macrogol, polyvinyl alcohol, and titanium dioxide (see also section 2 ‘Alunbrig contains lactose’ and ‘Alunbrig contains sodium’).

**What Alunbrig looks like and contents of the pack**

Alunbrig film‑coated tablets are white to off‑white, oval (90 mg and 180 mg) or round (30 mg). They are convex on the upper and lower side.

Alunbrig 30 mg:

* Each 30 mg tablet contains 30 mg brigatinib.
* The film‑coated tablets are approximately 7 mm in diameter with “U3” on one side and plain on the other side.

Alunbrig 90 mg:

* Each 90 mg tablet contains 90 mg brigatinib.
* The film‑coated tablets are approximately 15 mm long with “U7” on one side and plain on the other side.

Alunbrig 180 mg:

* Each 180 mg tablet contains 180 mg brigatinib.
* The film‑coated tablets are approximately 19 mm long with “U13” on one side and plain on the other side.

Alunbrig is available in plastic foil strips (blisters) packed in a carton with:

* Alunbrig 30 mg: 28, 56 or 112 film‑coated tablets
* Alunbrig 90 mg: 7 or 28 film‑coated tablets
* Alunbrig 180 mg: 28 film‑coated tablets

Alunbrig is available in plastic bottles with child resistant screw top closures. Each bottle contains one canister of a desiccant and is packed in a carton with:

* Alunbrig 30 mg: 60 or 120 film‑coated tablets
* Alunbrig 90 mg: 7 or 30 film‑coated tablets
* Alunbrig 180 mg: 30 film‑coated tablets

Keep the desiccant canister in the bottle.

Alunbrig is available as a treatment initiation pack. Each pack consists of an outer carton with two inner cartons containing:

* Alunbrig 90 mg film‑coated tablets

1 plastic foil strip (blister), containing 7 film‑coated tablets

* Alunbrig 180 mg film‑coated tablets

3 plastic foil strips (blisters), containing 21 film‑coated tablets

Not all pack sizes may be marketed.

**Marketing Authorisation Holder**

Takeda Pharma A/S

Delta Park 45

2665 Vallensbaek Strand

Denmark

**Manufacturer**

Takeda Austria GmbH

St. Peter‑Strasse 25

4020 Linz

Austria

Takeda Ireland Limited  
Bray Business Park  
Kilruddery   
Co. Wicklow   
A98 CD36  
Ireland

For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

|  |  |
| --- | --- |
| **België/Belgique/Belgien**  Takeda Belgium NV  Tél/Tel: +32 2 464 06 11  medinfoEMEA@takeda.com | **Lietuva**  Takeda, UAB  Tel: +370 521 09 070  medinfoEMEA@takeda.com |
| **България**  Такеда България ЕООД  Тел.: +359 2 958 27 36  medinfoEMEA@takeda.com | **Luxembourg/Luxemburg**  Takeda Belgium NV  Tél/Tel: +32 2 464 06 11  medinfoEMEA@takeda.com |
| **Česká republika**  Takeda Pharmaceuticals Czech Republic s.r.o.  Tel: +420 234 722 722  medinfoEMEA@takeda.com | **Magyarország**  Takeda Pharma Kft.  Tel.: +36 1 270 7030  medinfoEMEA@takeda.com |
| **Danmark**  Takeda Pharma A/S  Tlf: +45 46 77 10 10  medinfoEMEA@takeda.com | **Malta**  Drugsales Ltd  Tel: +356 21419070  safety@drugsalesltd.com |
| **Deutschland**  Takeda GmbH  Tel: +49 (0)800 825 3325  medinfoEMEA@takeda.com | **Nederland**  Takeda Nederland B.V.  Tel: +31 20 203 5492  medinfoEMEA@takeda.com |
| **Eesti**  Takeda Pharma AS  Tel: +372 6177 669  medinfoEMEA@takeda.com | **Norge**  Takeda AS  Tlf: +47 800 800 30  medinfoEMEA@takeda.com |
| **Ελλάδα**  Τakeda ΕΛΛΑΣ Α.Ε.  Tηλ: +30 210 6387800  medinfoEMEA@takeda.com | **Österreich**  Takeda Pharma Ges.m.b.H.  Tel: +43 (0) 800‑20 80 50  medinfoEMEA@takeda.com |
| **España**  Takeda Farmacéutica España, S.A.  Tel: +34 917 90 42 22  medinfoEMEA@takeda.com | **Polska**  Takeda Pharma Sp. z o.o.  Tel.: +48223062447  medinfoEMEA@takeda.com |
| **France**  Takeda France SAS  Tél: + 33 1 40 67 33 00  medinfoEMEA@takeda.com | **Portugal**  Takeda Farmacêuticos Portugal, Lda.  Tel: + 351 21 120 1457  medinfoEMEA@takeda.com |
| **Hrvatska**  Takeda Pharmaceuticals Croatia d.o.o.  Tel: +385 1 377 88 96  medinfoEMEA@takeda.com | **România**  Takeda Pharmaceuticals SRL  Tel: +40 21 335 03 91  medinfoEMEA@takeda.com |
| **Ireland**  Takeda Products Ireland Ltd  Tel: 1800 937 970  medinfoEMEA@takeda.com | **Slovenija**  Takeda Pharmaceuticals farmacevtska družba d.o.o.  Tel: + 386 (0) 59 082 480  medinfoEMEA@takeda.com |
| **Ísland**  Vistor hf.  Sími: +354 535 7000  medinfoEMEA@takeda.com | **Slovenská republika**  Takeda Pharmaceuticals Slovakia s.r.o.  Tel: +421 (2) 20 602 600  medinfoEMEA@takeda.com |
| **Italia**  Takeda Italia S.p.A.  Tel: +39 06 502601  medinfoEMEA@takeda.com | **Suomi/Finland**  Takeda Oy  Puh/Tel: 0800 774 051  medinfoEMEA@takeda.com |
| **Κύπρος**  A.POTAMITIS MEDICARE LTD  Τηλ: +357 22583333  a.potamitismedicare@cytanet.com.cy | **Sverige**  Takeda Pharma AB  Tel: 020 795 079  medinfoEMEA@takeda.com |
| **Latvija**  Takeda Latvia SIA  Tel: +371 67840082  medinfoEMEA@takeda.com | **United Kingdom (Northern Ireland)**  Takeda UK Ltd  Tel: +44 (0) 3333 000 181  medinfoEMEA@takeda.com |

**This leaflet was last revised in**

**Other sources of information**

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