

Patient Safety & Pharmacovigilance

European Union Risk Management Plan (EU RMP)

Mekinist[®], Spexotras[®] (trametinib)
TMT212

Document status:	Final
Version number:	23.0
Data lock point for this RMP	04-Jun-2025
Date of final sign off	09-Jan-2026

Rationale for submitting an updated RMP

This update to the EU RMP is intended to provide a consolidated RMP which includes the updates proposed in RMP v21.0 and v22.0 submitted under procedure numbers EMA/VR/0000271728 (application for the age range extension for the melanoma indications) and EMA/VR/0000278305 (application for a new indication for the treatment of adults with differentiated thyroid cancer (DTC)), respectively). This particular RMP v23.0 is prepared utilizing the currently approved EU RMP v20.0 as its base.

Summary of significant changes in this RMP

This RMP update is carried out in the new RMP template in accordance with the GVP Module V (Rev.2) guidance.

Part	Major changes compared to RMP v 20.0
Part I	Table 1-1 updated to reflect the extension of the melanoma indication to include the adolescent age group and proposed indication (treatment of adults with DTC)
Part II	Section adapted to reflect the extension of the melanoma indication to include the adolescent age group Module SI updated to reflect the epidemiology data for the DTC indication Module SIII updated to reflect the clinical trial exposure data for the DTC indication Module SV updated to reflect the updated post-marketing exposure (until May-2025) Module SVII.2 updated to indicate there are no changes to the list of safety concerns since the last approved RMP v 20.0
Part III	No update
Part IV	No update
Part V	Table 5-2 updated to include the newly assigned EU Clinical Trial (CT) number for study CDRB436G2401
Part VI	Updated to reflect the extension of the melanoma indication to include the adolescent age group, proposed indication and a newly assigned EU CT Number for study CDRB436G2401
Part VII	

Other RMP versions under evaluation

RMP Version number	Submitted on	Submitted within procedure number
Version 21.0	04-Jun-2025	EMA/VR/0000271728
Version 22.0	25-Aug-2025	EMA/VR/0000278305

Details of the currently approved RMP

Version number	20.0
Approved with procedure	EMEA/H/C/PSUSA/00010262/202405
Date of approval (opinion date)	16-Jan-2025

QPPV name: Dr. Justin Daniels, PhD

QPPV oversight declaration: The content of this RMP has been reviewed and approved by the marketing authorization holder's QPPV. The electronic signature is available on file.

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List of abbreviations

ADR	Adverse drug reaction
AAIR	Age-adjusted incidence rate
AE	Adverse event
AHR	Adjusted hazard ratio
ALK	Anaplastic lymphoma kinase
ALT	Alanine aminotransferase
AST	Aspartate aminotransferase
AUC	Area under plasma concentration time curve
AOR	Adjusted odds ratio
ARR	Adjusted risk ratio
BCRP	Breast cancer resistant protein
bid	Twice daily
BRAF	B-Raf proto-oncogene, serine/threonine kinase
CNS	Central nervous system
CI	Confidence interval
CRC	Colorectal cancer
CSR	Clinical study report
CRVO	Central retinal vein occlusion
CTCAE	Common Terminology Criteria for Adverse Event
DMFS	Distant metastasis-free survival
DTC	Differentiated thyroid cancer
DVT	Deep vein thrombosis
ECOG	Eastern Cooperative Oncology Group
EF	Ejection fraction
EGFR	Epidermal growth factor receptor
ERK	Extracellular signal-regulated kinase
EU	European Union
FDA	Food and Drug Administration
FFR	Freedom from relapse
GI	Gastrointestinal
GSK	GlaxoSmithKline
HDI	High dose interferon alpha
HGG	High grade glioma
HPMC	Hydroxypropyl methylcellulose
IFN	Interferon
ILD	Interstitial lung disease
IL-2	Interleukin-2
LFT	Liver function test
LGG	Low grade glioma
LLN	Lower limit of normal
LVEF	Left ventricular ejection fraction

MedDRA	Medical Dictionary for Regulatory Activities
MAH	Marketing Authorisation Holder
MAPK	Mitogen-activated protein kinase
MEK	Mitogen-activated extracellular signal regulated kinase
MTD	Maximum tolerated dose
NCI	National Cancer Institute
NNH	Number needed to harm
NSCLC	Non-small cell lung cancer
NYHA	New York Heart Association
OAT	Organic anion transporter
OR	Overall response rate
OS	Overall survival
PD-1	Programmed cell death 1
PE	Pulmonary embolism
PFS	Progression free survival
P-gp	p-glycoprotein
PIP	Paediatric investigation plan
PK	Pharmacokinetics
PMR	Post-marketing requirement
PR	Partial response
PRAC	Pharmacovigilance Risk Assessment Committee
PSUR	Periodic safety update report
PSUSA	PSUR Single Assessment
PTC	Papillary thyroid cancer
PTY	Patient treatment years
qd	Once daily
QPPV	Qualified Person for Pharmacovigilance
QTc	Corrected QT
RFS	Relapse free survival
RMP	Risk management plan
RPED	Retinal pigment epithelial detachment
RP2D	Recommended Phase II dose
RVO	Retinal vein occlusion
SAE	Serious adverse event
SEER	Surveillance, Epidemiology and End Results
SmPC	Summary of Product Characteristics
SRS	Stereotactic radiosurgery
TE	Thromboembolic event
TKI	Tyrosine kinase inhibitor
ULN	Upper limit of normal
US	United States
VEGFR	Vascular endothelial growth factor receptor

VTE	Venous thromboembolic event
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1 Part I: Product(s) Overview

Table 1-1 Part I.1 – Product(s) Overview

Active substance(s) (INN or common name)	Trametinib
Pharmacotherapeutic group(s) (ATC Code)	L01EE01
Marketing Authorization Holder/Applicant	Novartis Europharm Limited
Medicinal products to which this RMP refers	2
Invented name(s) in the European Economic Area (EEA)	Mekinist®, Spexotras®
Marketing authorization procedure	Centralized Procedure
Brief description of the product	Chemical class: Antineoplastic agent – Mitogen-activated protein kinase (MEK) inhibitors
	Summary of mode of action: Trametinib is a reversible, allosteric inhibitor of mitogen-activated extracellular signal regulated kinase 1 (MEK1) and MEK2 activation and kinase activity. MEK proteins are critical components of the mitogen-activated protein kinase (MAPK) pathway. In melanoma and other cancers, this pathway is often activated by mutated forms of B-Raf proto-oncogene, serine/threonine kinase (BRAF) which activates MEK and stimulates tumour cell growth. Trametinib inhibits activation of MEK by BRAF and inhibits MEK kinase activity, thereby inhibiting tumor growth.
	Important information about its composition: Trametinib is a polycyclic, nitrogen containing heterocycle also possessing aromatic halide and amide functionality.
Hyperlink to the Product Information	[Proposed SmPC]
Indication(s) in the EEA	Current (Mekinist): - Trametinib as monotherapy or in combination with dabrafenib is indicated for the treatment of adult patients with unresectable or metastatic melanoma with a BRAF V600E mutation; - Trametinib monotherapy has not demonstrated clinical activity in patients who have progressed on a prior BRAF inhibitor therapy; - Trametinib in combination with dabrafenib is indicated for the treatment of adults with advanced non-small cell lung cancer (NSCLC) with a BRAF V600E mutation. - Trametinib in combination with dabrafenib is indicated for the adjuvant treatment of adult patients with Stage III melanoma with a BRAF V600 mutation, following complete resection. Current (Spexotras):

	• Trametinib in combination with dabrafenib is indicated for the treatment of paediatric patients aged 1 year and older with low-grade glioma (LGG) with a BRAF V600E mutation who require systemic therapy. • Trametinib in combination with dabrafenib is indicated for the treatment of paediatric patients aged 1 year and older with high-grade glioma (HGG) with a BRAF V600E mutation who have received at least one prior radiation and/or chemotherapy treatment. Proposed (Mekinist): • Trametinib as monotherapy or in combination with dabrafenib is indicated for the treatment of adults and adolescents aged 12 years and older with unresectable or metastatic melanoma with a BRAF V600 mutation. • Trametinib in combination with dabrafenib is indicated for the adjuvant treatment of adults and adolescents aged 12 years and older with Stage III melanoma with a BRAF V600 mutation, following complete resection. • Trametinib in combination with dabrafenib is indicated for the treatment of adult patients with locally or advanced differentiated thyroid cancer with a BRAF V600E mutation, refractory to or not eligible for radioactive iodine (RAI) who have progressed during or after prior systemic therapy.
Dosage in the EEA	Current (Mekinist): • 2 mg once daily in adults Current (Spexotras): • Weight-based dosing, once daily (powder for oral solution formulation; see Spexotras SmPC for details) in paediatric patients as described in the SmPC Proposed: • Weight-based dosing, once daily p.o administration (film-coated tablet formulation, adolescent patients with melanoma; see proposed Mekinist SmPC for details).
Pharmaceutical form(s) and strengths	Current (Mekinist): • 0.5 mg and 2 mg film-coated tablets Current (Spexotras): • 0.05 mg/mL powder for oral solution
Is/will the product be subject to additional monitoring in the EU?	No

2 Part II: Safety specification

2.1 Part II: Module SI: Epidemiology of the indication(s) and target population

2.1.1 Indications: Treatment of adults and adolescents aged 12 years and older with unresectable or metastatic melanoma with BRAF V600E mutation and adjuvant treatment of patients with Stage III melanoma with a BRAF V600 mutation, following complete resection

2.1.1.1 Indication: Treatment of adults and adolescents aged 12 years and older with unresectable or metastatic melanoma with BRAF V600E mutation

Incidence

Cutaneous melanoma is the most aggressive form of all skin cancers, with approximately 232000 new cases and approximately 55000 disease-related deaths worldwide each year ([Globocan 2012a](#)). In Europe, malignant melanoma is the 9th most common cancer, with more than 100000 new cases diagnosed in 2012 ([Ferlay et al 2013](#)).

According to EUCAN, the age-standardized incidence rate (per 100000) of skin melanoma, in 2012 was 11.1 (males – 11.4, females – 11.0) in Europe (40 countries), and 13.0 (males – 13.2, females – 13.1) in the European Union (EU) (27 countries in 2012) ([EUCAN 2012d](#), [EUCAN 2012e](#), [EUCAN 2012f](#)). In the US, it was estimated that 87110 individuals will be diagnosed with skin melanoma and an estimated 9730 people will die of this disease in 2017 ([SEER 2017a](#)). The age-standardized incidence rate was 22.3 per 100000 individuals per year based on cases diagnosed in 2010-2014 in the geographical area of the Surveillance, Epidemiology and End Results (SEER) registries ([SEER 2017a](#)).

Table 2-1 Incidence of cutaneous melanoma

Country/Region	Incidence			Source of data/ reference
	Number of patients	of Annual Rate (per 100000)		
Europe	100339	11.1		EUCAN*/Ferlay et al (2013)
France	9871	13.0		EUCAN*/Ferlay et al (2013)
Germany	16884	14.8		EUCAN*/Ferlay et al (2013)
Italy	10012	13.4		EUCAN*/Ferlay et al (2013)
Spain	5004	8.6		EUCAN*/Ferlay et al (2013)
United Kingdom	14445	19.0		EUCAN*/Ferlay et al (2013)
US	87110	22.3		SEER (2017a)**

*EUCAN incidence rates are age-adjusted to the European standard population;**SEER incidence rates are age-adjusted to the 2000 US population

Although the number of cases of incident malignant melanoma is large, a small percentage of subjects are considered to have unresectable or metastatic disease (unresectable Stage IIIC or Stage IV disease). According to the SEER Program in the US, Stage IIIC and Stage IV

melanoma, respectively, comprise 1.6% and 4.2% of all new melanoma cases with known stage information ([SEER 2017b](#)).

Prevalence

The 2012 estimated 1-year prevalence proportion of melanoma is 13.9 per 100000 for Europe, corresponding to 87280 prevalent cases; and 24.0 per 100000 for the US, corresponding to 60518 prevalent cases; and 53.6 per 100000 in Australia/New Zealand, corresponding to 11846 prevalent cases ([Globocan 2012c](#), [Globocan 2012d](#)).

Table 2-2 Prevalence of cutaneous melanoma

Country/Region	Number of prevalent cases			Source of data/ reference
	1-year	3-year	5-year	
Europe	87285	247837	391316	EUCAN 2012d
France	8601	24760	39533	EUCAN 2012d
Germany	14735	42207	66997	EUCAN 2012d
Italy	8719	25154	40248	EUCAN 2012d
Spain	4309	12425	19792	EUCAN 2012d
United Kingdom	12602	36005	57163	EUCAN 2012d
US	60518	175103	281577	GLOBOCAN 2012d

The 2012 unresectable and metastatic prevalent population of melanoma (i.e. unresectable Stage IIIC and IV) that is considered eligible for drug treatment is estimated to be 15120 patients in the US. Of these, 5380 patients are estimated to harbor the BRAF mutation and be eligible for 1st or 2nd line treatment ([Webster and Hughes 2012](#)). Similarly, the drug-treatable, unresectable and metastatic prevalent melanoma population in the EU-5 is expected to number 16414 cases, of which 5260 patients would have BRAF mutation and be eligible for 1st or 2nd line treatment ([Webster and Hughes 2012](#)).

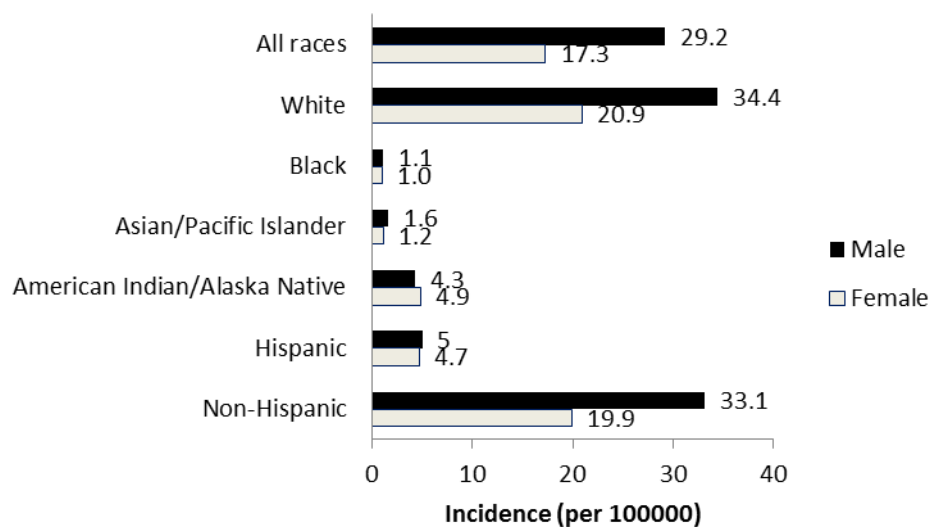
The frequency of BRAF mutations in melanoma has been reported to be approximately 50% (range: 27% to 70%) ([Garnett and Marais 2004](#), [Chapman 2011a](#)).

Demographics of the population in the authorized indication – age, gender, racial and/or ethnic origin and risk factors for the disease

Melanoma risk varies by age and gender. Based on 2006 to 2008 SEER cancer registry data among Whites, the probability of developing melanoma is higher among women than men from birth to age 39 (0.27 and 0.15, respectively), but reverses thereafter (0.56 and 0.63, respectively, from ages 40 to 59; 0.39 and 0.75, respectively, from ages 60 to 69; 0.82 and 1.94, respectively, from age 70 onwards) ([Siegel et al 2012](#)).

The overall incidence (per 100000) is 29.2 among men and 17.3 among women in the US. The corresponding incidence rates (male and females) across race/ethnicity is shown in [Figure 2-1 \(SEER 2017a\)](#).

Figure 2-1 Incidence of cutaneous melanoma by race/ethnicity and gender



SEER 18 registries (2010-2014), Age-adjusted rates

Based on SEER Cancer Stat Facts: Melanoma of the skin ([SEER 2017a](#))

From 2010-2014, the median age at diagnosis of skin melanoma was 64 years in the US. Approximately 22.2% were diagnosed between 55 and 64 years; 22.7% between 65 and 74 years; 17.1% between 75 and 84 years. About 8.2% were diagnosed over 84 years of age ([SEER 2017a](#)).

The US SEER cancer registry included 4210 melanoma patients diagnosed in 2005-2009 initially as Stage IIIC to Stage IV, which can be considered as a proxy for unresectable and metastatic melanoma patients. Of these, 48% of patients diagnosed with unresectable and metastatic melanoma were at age 65 years and above, 28% at 75 years and above, and 8% at 85 years and above ([Table 2-3](#)). Similar age distribution was observed among unresectable and metastatic melanoma subjects in Denmark.

Table 2-3 Age distribution of unresectable and metastatic melanoma subjects

Age at diagnosis	US SEER 2005-2009 ^a Unresectable/metastatic (Stage IIIC+IV) melanoma		Denmark 1997-2010 ^b Metastatic (Stage IV) melanoma	
	N	%	N	%
<55 years	1209	28.7	801	28.4
55-64 years	973	23.1	669	23.8
65-74 years	836	19.9	629	22.4
75-84 years	849	20.2	536	19.0
85+ years	343	8.1	179	6.4
Total	4210	100.0	2814	100.0

^a-Data Source: Software: Surveillance Research Program, SEER*Stat software (www.seer.cancer.gov/seerstat) version 8.0.1.Data: Surveillance, Epidemiology, and End Results (SEER) Program (www.seer.cancer.gov) SEER*Stat Database: Incidence – SEER 18 Regs Research Data + Hurricane Katrina Impacted Louisiana Cases, Nov 2011 Sub (1973-2009 varying) – Linked To County Attributes – Total US, 1969-2010 Counties,

Age at diagnosis	US SEER 2005-2009 ^a Unresectable/metastatic (Stage IIIC+IV) melanoma		Denmark 1997-2010 ^b Metastatic (Stage IV) melanoma	
	N	%	N	%
National Cancer Institute, DCCPS, Surveillance Research Program, Surveillance Systems Branch, released April 2012, based on the November 2011 submission.				
b-Data Source: Danish Cancer registry and Danish Pathology Registry. GSK sponsored study (ID: WEUSKOP6139), unpublished (Mekinist EU RMP V11-Annex 12).				

Cutaneous melanoma is a multi-factorial disease with both genetic and environmental risk factors – a personal or family history of melanoma, the presence of atypical or numerous moles (>50), sun sensitivity (sun burning easily, tanning minimally, natural blond or red hair color), a history of high intermittent sun exposure, including sunburns, use of tanning booths, diseases that suppress the immune system, and past history of basal or squamous cell carcinoma ([American Cancer Society 2012a](#)).

The main existing treatment options

The treatment choice for malignant melanoma depends on cancer stage, whether the tumour is resectable, BRAF mutation status, patient health status and drug toxicity profile ([Solanki et al 2012](#)). Patients who have unresectable Stage III melanoma are generally treated like those with metastatic disease ([Solanki et al 2012](#)).

Systemic treatment with chemotherapy has been the traditional way to treat unresectable and metastatic melanoma, although with little to no impact on survival for subjects. The alkylating agent, dacarbazine (DTIC), is the most widely used chemotherapy for advanced disease with a response rate of 5-12% and median duration of response of 6 months ([Solanki et al 2012](#), [Avril et al 2004](#), [Middleton et al 2000a](#), [Bedikian et al 2006](#), [Schadendorf et al 2006](#), [Chapman 2011a](#), [Robert et al 2011](#)).

Current treatment options for unresectable and metastatic melanoma include:

- immunotherapy,
- chemotherapy,
- targeted therapy.

The approvals of ipilimumab, vemurafenib, dabrafenib, trametinib, pembrolizumab and nivolumab in the US and EU in 2011 to 2014 for the treatment of unresectable and metastatic malignant melanoma as well as more recent approval of cobimetinib in combination with vemurafenib in US and EU in 2015, and talimogene laherparepvec (an oncolytic virus therapy, also known as T-vec) in US with positive opinion in EU marks the start of a new era for the treatment of this disease. Prior to these, immunotherapy and chemotherapy -both as single agents and as combination regimens-had failed to significantly improve survival for advanced malignant melanoma subjects. In patients with BRAF V600E or K mutation, the combination of a BRAF inhibitor and a MEK inhibitor, including dabrafenib plus trametinib, is the treatment with highest level of medical evidence (category 1) recommended by National Comprehensive Cancer Network (NCCN) guideline as first line treatment for unresectable or metastasis disease ([NCCN 2016](#)).

On 29-May-2013, Tafenlar® (dabrafenib) and Mekinist (trametinib) were approved by the US Food and Drug Administration (FDA) for the treatment of patients with unresectable or metastatic melanoma with BRAF V600E (dabrafenib monotherapy and trametinib monotherapy) or BRAF V600K (trametinib monotherapy only) mutations. Tafenlar and Mekinist were also approved in Canada as monotherapies (16-Jul-2013 and 18-Jul-2013, respectively), Australia (21-Aug-2013 and 11-Feb-2014, respectively) and in the EU (26-Aug-2013 and 30-Jun-2014, respectively). Tafenlar and Mekinist have subsequently been approved in multiple additional countries as single agents.

The combination of Tafenlar and Mekinist was first approved by the FDA on 08-Jan-2014 (accelerated approval), Australia TGA on 11-Feb-2014, Canada on 06-Mar-2015 (conditional approval), and New Zealand on 20-Mar-2015 for the treatment of patients with unresectable or metastatic melanoma with BRAF V600E or V600K mutations. The combination was approved on 25-Aug-2015 in the EU and the FDA granted conversion of the accelerated approval to regular approval on 20-Nov-2015 based on Phase III data. The combinations of Tafenlar and Mekinist have subsequently been approved in multiple additional countries.

The purpose of the medicinal product covered by this EU RMP, including trametinib monotherapy and trametinib in combination with dabrafenib, is to reduce progression of disease in patients with BRAF V600E mutation.

Natural history of the indicated condition in the population, including mortality and morbidity

The melanoma mortality rate in Europe is 1.5 per 100000 ([Forsea et al 2012](#)). Due to early detection, a majority of melanoma patients are cured with surgery alone. Historically, the median survival time for subjects with Stage IV melanoma was short, at approximately 6 months with 26% of subjects alive at 1 year, and a median progression-free survival (PFS) of 1.7 months with 14.5% of subjects progression-free at 6 months ([Korn et al 2008](#)). With the advances in immune check-point inhibitor, such as the combination treatment of nivolumab and ipilimumab, the median PFS for metastatic melanoma patients can reach 11.5 months and response rate reached 57.6% ([Larkin et al 2015](#)). The mean age at baseline was 59 years old. On average, an individual loses 20.4 years of potential life as a result of melanoma mortality, compare to 16.6 years for all malignancies ([NCCN 2016](#)).

In the US, the mortality rates for skin melanoma are 4.0 per 100000 men and 1.7 per 100000 women. The mortality rates (per 100000) in men and women are 4.6 and 1.9 among Caucasians, 0.5 and 0.4 among Blacks, 0.4 and 0.3 among Asian/Pacific Islanders, 1.4 and 0.5 among American Indian/Alaska Natives, and 1.0 and 0.6 among Hispanics ([SEER 2017a](#)).

From 2010-2014, the median age at death for cancer of melanoma was 70 years of age. Approximately 2.0% died between ages 20 and 34; 4.6% between 35 and 44; 11.2% between 45 and 54; 20.1% between 55 and 64; 22.8% between 65 and 74; 24.1% between 75 and 84; and 15.1% for those over 84 years of age ([SEER 2017a](#)).

The age-adjusted death rate was 2.7 per 100000 men and women per year based on data from patients who died between 2010 and 2014 in the US. Based on 2007-2013 data, 5-year survival rate of melanoma was 91.7% ([SEER 2017a](#)).

Cutaneous melanoma accounts for less than 5% of all skin cancers, which also includes basal cell carcinoma and squamous cell carcinoma ([American Cancer Society 2012a](#)), but it causes 75% of skin cancer deaths ([Jerant et al 2000](#)).

Mortality is worse among Whites compared with African Americans – the death rates (per 100000 persons) in males were 4.6 versus 0.5, respectively, and in females, 1.9 versus 0.4, respectively ([SEER 2017a](#)).

Overall, death rate has been decreasing among Whites, younger than age 50, by 2.9% annually in men and 2.3% annually in women from 2004 to 2008, whereas the rates among Whites, 50 years and older, has increased 1% per year for males and remained stable for females during the same time period ([American Cancer Society 2012a](#)).

Melanoma is highly curable if detected in its earliest stages and treated properly (usually surgery). However, the prognosis for metastatic melanoma patients has been historically poor because of limited treatment options and efficacy, which up until recently, included mainly alkylating agents, DTIC and temozolomide, and immunotherapy with IL-2 and/or interferon-alpha (IFN- α) ([American Cancer Society 2012b](#)).

Unresectable, locally advanced melanoma (Stage IIIC) is often treated in the same manner as metastatic (Stage IV) melanoma ([Webster and Hughes 2012](#)). Based on SEER cancer registry data, for melanoma subjects diagnosed from 2004 to 2009, the 1-year and 5-year survival rates are approximately 81% and 35%, respectively, for Stage IIIC melanoma, and 39% and 13%, respectively, for Stage IV melanoma ([NCI 2011](#)). Using data from the Surveillance, Epidemiology, and End Results (SEER) database, patients diagnosed with unresectable Stage IIIB/C and Stage IV (M1a, M1b, M1c) melanoma between 2004 and 2009 were selected. Patients at Stage IIIB/IIIC had a median overall survival (OS) of 24.3 months, with a survival rate of 67.2% at 1 year, 42.9% at 2 years, and 32.1% at 3 years. For patients at Stage M1a, the median OS was 22.3 months, 1 year, 2 year, and 3 year survival rates were 64.5%, 40.4%, and 26.4%, respectively; for patients at Stage M1b, median OS was 11.2 months, 1 year, 2 year, and 3 year survival rates were 43.8%, 23.4%, and 13.8%, respectively; for patients at Stage M1c, median OS was 5.1 months, and 1 year, 2 year, and 3 year survival rates were 22.3%, 8.9%, and 4.7%, respectively ([Song et al 2015](#)).

Adverse events can occur in patients with unresectable or metastatic melanoma. Population-based studies or clinical trials in patients with placebo only arms evaluating untreated patients with unresectable or metastatic melanoma were not available. The following information is based on Phase III clinical trials in patients with unresectable or metastatic melanoma that included a comparator arm (DTIC or carboplatin/paclitaxel combination) with a placebo. In a Phase III trial on patients with previously untreated unresectable or metastatic melanoma, among patients in the control arm (treated with DTIC and placebo), 94% had an adverse event with 27.5% reported to have a grade 3 or 4 adverse event. Grade 3 or 4 nausea was reported in 1.2%, vomiting in 1.6%, abdominal pain in 2.8%, fatigue in 4.8%, asthenia in 2.4%, back pain in 1.2%, decreased appetite in 1.6%, increased aspartate aminotransferase (AST) in 1.2% and immune-related adverse events in 6% of patients in the control arm ([Robert et al 2011](#), [Robert et al 2011a](#)). Similar information on patients unresectable or metastatic melanoma treated with carboplatin and paclitaxel as first ([Flaherty et al 2013](#)) or second line therapy ([Hauschild et al 2009](#)) is included in [Table 2-4](#).

Table 2-4 Adverse events reported in patients with unresected or metastatic melanoma (in a clinical trial control arm treated with placebo and carboplatin/paclitaxel)

Adverse Event (Grade 3 or higher)	Frequency
Total	69%² – 78.2%¹
Blood/bone marrow	60%²
Neutrophils	46% ² -49.1% ¹
Platelets	8.8% ¹ -12% ²
Hemoglobin	7.1% ¹ - 13% ²
Leukocytes	19% ² – 22.9% ¹
Constitutional symptoms	13%²
Fatigue	10% ² – 14.1% ¹
Anorexia	2.3%¹
Gastrointestinal	14%²
Diarrhea	3.0% ² – 3.8% ^{1*}
Infection	15%²
Febrile neutropenia	4.0% ¹ – 7.0% ²
Metabolic/laboratory	11%²
Lipase	2% ²
Neurology	20%²
Neuropathy, sensory	13.0% ² – 14.9% ¹
Pain	18%²
Pain, extremity	5.0% ²
Muscle pain	5.5%¹
Dermatology	4%²
Rash/desquamation	2.0% ¹
Hand-foot skin reaction	0.3% ¹
Hypertension	1.3%¹
Allergic reaction	2.8%¹
Lymphopenia	4.3%¹
Dehydration	4.8%¹
Hyponatremia	2.3%¹
Hyperglycemia	4.8%¹

Source: ¹Flaherty et al 2013; ²Hauschild et al 2009; *without prior colostomy

Important co-morbidities

To obtain background rates for the most commonly occurring co-morbidities in a real-world population of unresectable and metastatic melanoma subjects, a retrospective study was conducted using the US SEER-Medicare Linked Databases. This study included 1746 subjects (aged 65+ years; male, 61%; primarily Whites, 95%) with initial diagnoses of Stage IIIC unresectable or Stage IV metastatic melanoma during 1992 to 2005 (cases) and 1746 age-, gender-, race-, and region-matched non-cancer controls, drawn from SEER-Medicare Linkage Databases. Of these unresectable/metastatic melanoma subjects, 89.8% subjects died during the

follow-up and the median survival was 10 months ([SEER-Medicare Study](#), Mekinist EU RMP V12-Annex 12.1).

During the 12 months prior to unresectable and metastatic melanoma diagnosis, the most prevalent co-morbidities, with prevalence >20%, included essential hypertension (53.1%), other skin disorders (38.7%), disorders of lipid metabolism (37.5%), cataract (32.9%), connective tissue disease (30.9%), lower respiratory disease (30.0%), non-traumatic joint disorders (26.8%), coronary atherosclerosis (23.3%), and diabetes mellitus without complication (21.0%). Compared with age-, gender, race-, and region-matched non-cancer controls, unresectable/metastatic melanoma subjects had more than 2-fold higher prevalence of neoplasms of unspecified nature or uncertain behavior (19% versus 9%), pleurisy and pleural effusion (4.8% versus 2.2%), other non-epithelial cancer of skin (17% versus 8%), and open wounds of extremities (5.5% versus 2.6%). Comparable databases with oncology information encompassing countries comprising the EU are not available. Therefore, US estimates of important co-morbidities are presented as a surrogate for the EU.

2.1.1.2 Indication: Adjuvant treatment of adults and adolescents aged 12 years and older with Stage III melanoma with a BRAF V600 mutation, following complete resection

Incidence

According to the data from the 18 SEER (registries) in the US, among 102706 patients with melanoma diagnosed between 2010 and 2014, stage information ([AJCC stage groups, 7th ed](#)) was available on 93782 patients of whom 7.3% had Stage III disease (including 1.5% with Stage IIIA, 2.1% with Stage IIIB, 1.6% with Stage IIIC, and 2.1% listed as Stage III or Stage III NOS) ([SEER 2017b](#)). Applying this percentage to the number of new patients expected to be diagnosed with melanoma in the US in 2017, it is estimated that 6359 new patients with Stage III melanoma will be diagnosed in the US in 2017. Similar information on stage was not available in Europe. Applying the stage distribution from the SEER data to the incidence of melanoma in Europe in 2012, it is estimated that 7325 new patients are diagnosed with Stage III melanoma in a year in Europe ([EUCAN 2012d](#), [SEER 2017b](#)).

Prevalence

Data on the prevalence of Stage III melanoma are limited. Based on prevalence data from SEER and age-adjustment to the US population in 2016, it is estimated that there were 13322 patients who had been newly diagnosed to have melanoma with Stage III disease within the previous 3 years and were alive in 2016 in the US ([SEER 2017c](#), [CDC Wonder 2017](#)). The corresponding estimate, age-adjusted to the European population in 2016, is 34,160 ([SEER 2017c](#), [United Nations 2017](#)).

Demographics of the population in the authorized indication – age, gender, racial and/or ethnic origin and risk factors for the disease

According to data from the 18 SEER registries in the US, among 6868 patients diagnosed with Stage III melanoma between 2010 and 2014, 4496 (65%) were male and 2372 (35%) were female. About 34% were younger than 55 years of age, 24% were between 55 and 64 years,

21% were between 65 and 74 years, 15% were between 75 and 84 years and 6% were over the age of 84 years ([SEER 2017d](#)). Most patients were white (97%), 1.2% were black and the remaining patients were of unknown or other races ([SEER 2017e](#)). Risk factors for melanoma are listed in [Section 2.1.1.1](#).

Table 2-5 Age and gender distribution of patients with Stage III melanoma*

Age	Male	Female	Total
<55 years	1344 (30%)	994 (42%)	2338 (34%)
55-64 years	1141 (25%)	480 (20%)	1621 (24%)
65-74 years	1040 (23%)	415 (18%)	1455 (21%)
75-84 years	706 (16%)	332 (14%)	1038 (15%)
85+ years	265 (6%)	151 (6%)	416 (6%)
Total	4496 (100%)	2372 (100%)	6868 (100%)

*Data from SEER 18 registries, patients with Stage III melanoma ([AJCC Stage group, 7th ed](#)) diagnosed between 2010 and 2014, obtained using SEER*Stat version 8.3.4 ([SEER 2017d](#))

The main existing treatment options

Adjuvant therapy is indicated in patients with Stage III melanoma at high risk of recurrence following complete surgical resection with the intent of treating micrometastatic disease and reduce the risk of local and distant relapse ([Kirkwood et al 2001](#), [Van Akkooi et al 2009](#)).

Different therapies have been explored in the adjuvant setting, including interferon, interleukin-2, and vaccines, bevacizumab (a vascular endothelial growth factor inhibitor), as well as ipilimumab (an immune checkpoint inhibitor) in the last decade. Specifically, high dose interferon alpha (HDI) and PEGylated interferon have been approved for adjuvant melanoma treatment based on relapse free survival (RFS) improvement, without significant survival benefit in the majority of the studies conducted; only one study ([Kirkwood et al 2004](#)) demonstrated initial survival benefit, although it was not confirmed in the analysis performed with additional follow up. The unfavorable safety profile, as shown by the significant treatment related toxicities, has limited use in clinical practice and patient adherence ([Kirkwood 1996](#), [Kirkwood 2000](#), [Eggermont 2005](#)). Ipilimumab, an immune checkpoint inhibitor, used at a dosage of 10 mg/kg, has shown significant RFS improvement in high risk Stage III melanoma after complete resection (HR=0.76; 95% CI: 0.64, 0.89), and this has translated into a survival benefit (HR=0.72; 95% CI: 0.58, 0.88) after a median follow-up of 5.3 years; however, the treatment related toxicities are severe. Nearly half of the patients had toxicity equal to or greater than common terminology criteria for adverse events (CTCAE) grade 3. Five (1%) patients died due to drug related adverse event (AE) and all these events occurred within the first 12 weeks of treatment. A total of 52% patients discontinued treatment because of an AE. Only 7% of the patients completed the planned 3 year treatment ([Eggermont et al 2015](#), [Eggermont et al 2016](#)). Ipilimumab at a dosage of 10 mg/kg in the adjuvant setting was approved in the US in October 2015. In the AVAST-M Phase III study, bevacizumab treatment was assessed as adjuvant treatment in patients with Stage IIB, IIC and III melanoma. The primary endpoint, OS was not met; survival rate at 5 years was 64% on bevacizumab versus 63% on observation arm (HR=0.99; 95% CI: 0.84, 1.18; p=0.96) ([Corrie et al 2017](#)).

The poor clinical outcome observed in patients with Stage III melanoma reflects the need for effective adjuvant treatments to prevent relapse.

Natural history of the indicated condition in the population, including mortality and morbidity

Stage III melanoma accounts for approximately 10% of newly diagnosed melanomas, is treated with complete resection, however it is associated with a high risk of relapse. The risk of relapse and mortality is defined by independent predictive factors including, primary tumor thickness; ulceration; mitotic rate and lymph node burden ([Balch et al 2009](#)). The overall 5-year RFS observed for Stage IIIA, IIIB, and IIIC patients was 63%, 32%, and 11%, respectively ([Romano et al 2010](#)). The estimated 5-year survivals for stages IIIA, IIIB, and IIIC from time of first relapse were 20%, 20%, and 11%, respectively ([Romano et al 2010](#)).

Patients with Stage III melanoma who are treated with surgical resection may experience recurrence and other adverse events. In a Phase III trial evaluating ipilimumab versus placebo in patients who had undergone complete resection of Stage III melanoma, at a median follow-up of 5.3 years, the 5-year rate of recurrence-free survival was 30.3% in the placebo group ([Eggermont et al 2016](#)). The rate of OS was 54.4% and the rate of distant metastasis-free survival (DMFS) was 38.9% in the placebo group. Among the 474 patients who received placebo, 91.1% had an adverse event of any grade. Grade 3 or 4 events occurred in 26.2% of patients in the placebo group and included diarrhea (2.1%), abdominal pain (0.2%), vomiting (0.2%), colitis (0.2%), fatigue (1.5%), headache (0.2%), weight loss (0.4%), increased weight (0.4%), pyrexia (0.2%), and decreased appetite (0.2%).

Important co-morbidities

Data on comorbidities specifically in patients with Stage III melanoma were not available from population-based studies. However, according to a nationwide cohort study in Denmark on patients diagnosed with melanoma (n=23476) between 1987 and 2009, 19% of patients with melanoma suffered from one or more comorbidities with 9.9% of patients having one comorbidity, 5.8% having two comorbidities, 1.8% having 3 comorbidities and 1.4% having 4 or more comorbidities. Any cancer (excluding melanoma and non-melanoma skin cancer) was the most common comorbidity (3.9%), followed by cerebrovascular disease (3.4%) and chronic pulmonary disease (2.4%), and diabetes (2.0%). Other comorbidities that occurred in over 1% of the melanoma patients included myocardial infarction (1.7%), congestive heart failure (1.7%), peripheral vascular disease (1.4%), ulcer disease (1.5%), and connective tissue disease (1.4%) ([Grann et al 2013](#)).

2.1.2 Indication: Treatment of patients with advanced non-small cell lung cancer with a BRAF V600 mutation

Incidence

According to the [Globocan \(2012a\)](#) project of the World Health Organization and the International agency for Research on Cancer, lung cancer has been the most common cancer in the world for several decades, and in 2012, there were an estimated 1.8 million new cases worldwide, representing 12.9% of all new cancers. It was also the most common cause of death

from cancer, with 1.59 million deaths worldwide in 2012 (19.4% of the total) ([Globocan 2012b](#)). The estimated number of patients diagnosed with lung cancer in 2012 was 409911 in Europe and 309589 in the 27 member states of EU ([EUCAN 2012a](#), [EUCAN 2012b](#), [EUCAN 2012c](#)).

Table 2-6 Incidence of lung cancer and NSCLC

Country/Region	Incidence			Source of data
	Annual rate of lung cancer (per 100000)	Number of lung cancer patients	Number of NSCLC patients with BRAF V600E mutation***	
Europe	41.9	409911	6968	EUCAN*/Ferlay et al (2013)
France	49.2	40043	681	EUCAN*/Ferlay et al (2013)
Germany	39.8	50813	864	EUCAN*/Ferlay et al (2013)
Italy	36.6	37238	633	EUCAN*/Ferlay et al (2013)
Spain	43.5	26715	454	EUCAN*/Ferlay et al (2013)
United Kingdom	45.1	40382	686	EUCAN*/Ferlay et al (2013)
US	55.8	222500	3783	SEER (2017a)**

*EUCAN incidence rates are age-adjusted to the European standard population; **SEER incidence rates are age-adjusted to the 2000 US population; ***number of NSCLC patients with BRAF V600E mutation was estimated assuming that 85% of newly diagnosed lung cancer patients have NSCLC and that 2% of NSCLC patients harbor the V600E mutation.

NSCLC accounts for the majority of cases (~85%) of lung cancer. [Pagano et al 2010](#) analyzed data on incident lung cancer cases in a regional cancer registry in Italy from 2000 through 2003. There were 2572 cases of NSCLC which represented 90% of all incident lung cancers. A Spanish study of 481 lung cancers diagnosed in a defined health area from February 1997 through December 1999 reported that approximately 80% were NSCLC ([Prim et al 2010](#)).

Prevalence

The 5-year prevalence count of lung cancer, including trachea and bronchus, was 442810 in Europe (40 countries) and 336143 (230842 men and 105301 women) in the EU (27 countries) in 2012 ([EUCAN 2012a](#), [EUCAN 2012b](#), [EUCAN 2012c](#), [Bray et al 2013](#)). In the US, there were an estimated 527228 people living with lung and bronchus cancer in 2014 ([SEER 2017](#)).

Table 2-7 Prevalence of lung cancer

Country/Region	Number of prevalent cases			Source of data/reference
	1-year	3-year	5-year	
Europe	184032	356582	442810	EUCAN 2012a
France	21863	43732	54811	EUCAN 2012a
Germany	21666	43554	55783	EUCAN 2012a
Italy	17866	35159	43960	EUCAN 2012a
Spain	11551	22532	28148	EUCAN 2012a
United Kingdom	13430	24826	30298	EUCAN 2012a
US	103571	210138	268629	GLOBOCAN 2012d

BRAF mutations are observed in approximately 2% of NSCLC and occur most frequently in adenocarcinomas. Of the BRAF mutations, around half were BRAF V600 mutations. Of the BRAF V600 mutations, almost all are V600E mutations ([Chen et al 2014](#)). [Pratilis et al \(2008\)](#) evaluated 916 patients from Japan, Taiwan, US and Australia with NSCLC and reported that 17 patients had BRAF mutations, including 11 patients with V600E mutation (1.2%). Similarly, in a study by [Paik et al \(2011\)](#), among 697 patients with lung adenocarcinoma, BRAF mutations were present in 18 patients. Out of these 18 patients, 9 patients were diagnosed with the BRAF V600E mutation, with a frequency of 1.3% ([Paik et al 2011](#)). [Marchetti et al \(2011\)](#) selected a cohort of 1046 patients in Italy with NSCLC of whom 739 patients had adenocarcinoma and 307 had squamous cell carcinoma. BRAF mutations were present in 37 patients among whom twenty-one patients were identified with BRAF V600E mutation leading to a frequency of 2%. A similar frequency of 2% of BRAF V600E mutation was reported by [Cardarella et al \(2013\)](#) when they evaluated 883 patients with NSCLC in the US at a cancer institute among whom 36 tumors harbored the BRAF mutations (V600E in 18 and non-V600E in 18). According to a 1-year nationwide program of the French Cooperative Thoracic Intergroup (IFCT) on routine molecular profiling of patients with advanced NSCLC, BRAF mutations were reported in 262 (2%) of 13906 molecular analyses with available data among 17664 patients with NSCLC ([Barlesi et al 2016](#)).

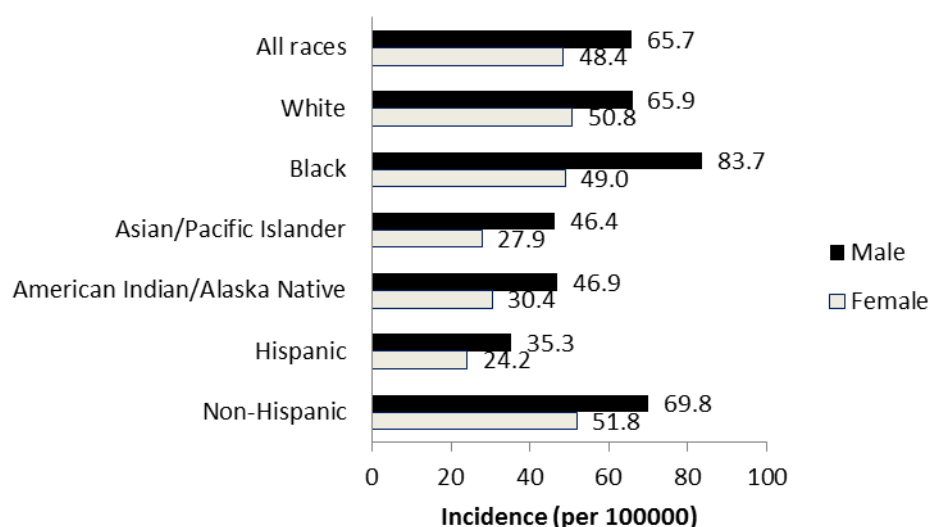
Based on the above, the proportion of patients with BRAF V600E NSCLC among all lung cancer patients is expected to be small. Of the estimated 222500 new cases of lung cancer in the US in 2017 ([SEER 2017](#)), assuming that around 85% of lung cancer patients have NSCLC ([Pagano et al 2010](#)) and up to 2% harbor the BRAF V600E mutation ([Cardarella et al 2013](#)), it is expected that up to 3783 patients have BRAF V600E mutant NSCLC. Similarly, of the estimated 409911 new cases of lung cancer in Europe in 2012 ([EUCAN 2012a](#)), up to 6968 patients are expected to have BRAF V600E mutant NSCLC.

Demographics of the population in the authorized indication – age, gender, racial and/or ethnic origin and risk factors for the disease

Lung cancer occurs in men more frequently than women. It is the most common cancer in men worldwide (1.2 million new cases worldwide in 2012, 16.7% of the total), with the highest estimated age-standardized incidence rates in Central and Eastern Europe, and Eastern Asia. In females, incidence rates are generally lower, (583000 cases and 491000 deaths worldwide in 2012) ([Globocan 2012b](#)). According to SEER data in the US, the overall incidence (per 100000) was 65.7 among men and 48.4 among women. The corresponding incidence rates across different races/ethnicities (male and females) are shown in [Figure 2-2 \(SEER 2017\)](#).

Lung cancer is a disease of elderly population. From 2010-2014, the median age at diagnosis of cancer of the lung and bronchus was 70 years in the US. Approximately 21.5% were diagnosed between 55 and 64 years; 32.9% between 65 and 74; 27.1% between 75 and 84 years. About 9.4% were diagnosed over 84 years of age ([SEER 2017](#)).

Figure 2-2 Incidence of lung cancer by race/ethnicity and gender



SEER 18 registries (2010-2014), Age-adjusted rates

Based on SEER Cancer Stat Facts: Lung and Bronchus Cancer ([SEER 2017](#))

There were no significant differences between the age and stage of the tumor at initial diagnosis between patients with BRAF mutations and wild type tumors ([Cardarella et al 2013](#); [Marchetti et al 2011](#)). The gender difference of NSCLC with the BRAF V600 mutation is unclear. Some studies reported that it is more frequent in women than men ([Li et al 2015](#), [Marchetti et al 2011](#)). However, other studies did not report a significant difference between males and females with respect to BRAF mutations ([Luk et al 2015](#), [Paik et al 2011](#), [Schmid et al 2009](#)). According to IFCT data, among 262 NSCLC patients with a BRAF mutation, 61% were male and 39% were female (compared to 65% males and 34% females in the wild type NSCLC group). The median age was 65.9 years among patients with the BRAF mutation and 64.7 years among those in the wild-type NSCLC group ([Barlesi et al 2016](#)).

Several risk factors contribute to the development of lung cancer, including cigarette, pipe, or cigar smoking; exposure to second-hand smoke, radon, arsenic, asbestos, chromates, chloromethyl ethers, nickel, polycyclic aromatic hydrocarbons, radon progeny, other agents, air pollution, and radiation therapy to the breast or chest ([NCI 2016a](#)). Smoking is considered the single most important risk factor for the development of lung cancer.

The BRAF V600E mutation can be diagnosed both in smokers and non-smokers. In a study by [Litvak et al \(2014\)](#), majority of patients with BRAF mutations were smokers (92%), although patients with V600 mutations were more likely to be light/never smokers compared to patients with non-V600 mutations (42% versus 11%). [Marchetti et al \(2011\)](#) reported that the V600E mutations were more frequent in patients who never smoked than in smokers or former smokers (10 of 197 patients [5.1%] versus 11 of 542 patients [2%]). [Pratils et al \(2008\)](#) reported that majority of NSCLC patients with BRAF mutations were current or former smokers, although information specifically on patients with BRAF V600 mutations was not available. In a study from France, the proportion of never, former, and current smokers was 25%, 38%, and 37%

among NSCLC patients with the BRAF mutation and 18%, 42%, and 40% among those with wild-type NSCLC ([Barlesi et al 2016](#)).

The main existing treatment options

There are different treatment options available for patients with NSCLC ([NCI 2016b](#)). These vary depending on the stage of the disease. According to the National Cancer Institute (NCI), results of standard treatment in NSCLC are poor except for the most localized cancers. Surgery is the most potentially curative therapeutic option for early stage disease. For advanced or metastatic NSCLC, systemic treatment is needed.

In advanced-stage and metastatic NSCLC, systemic chemotherapy with four to six cycles of a platinum-based doublet is widely used as the standard first-line therapy. In those NSCLC patients with good performance status who have experienced tumor regression or achieved at least disease stabilization, maintenance treatment with anticancer agents including pemetrexed or erlotinib has been validated as an effective treatment. Before programmed cell death (PD-1) antibodies were introduced in the clinic, for patients without an actionable mutation, further treatment options upon disease progression included single-agent chemotherapy such as pemetrexed or docetaxel or molecularly targeted therapy, such as erlotinib ([Schiller et al 2002](#), [Borghaei et al 2015](#), [Mok et al 2009](#), [Solomon et al 2014](#), [Hanna et al 2004](#), [Rosell et al 2012](#)). The median PFS usually is about 4-6 months for patients receiving platinum doublet chemotherapy as first line, and 2-3 months for single agent chemotherapy as second line. The median OS is less than one year.

Major advances in the definition of the molecular pathology of NSCLC have led to the development of targeted agents attacking cancer-cell specific attributes essential for growth or survival, such as epidermal growth factor receptor (EGFR), anaplastic lymphoma kinase (ALK) or vascular endothelial growth factor receptor (VEGFR) tyrosine kinases, and antibodies that inhibit the immune checkpoint and restore antitumor immunity, while avoiding some of the severe side effects of conventional cytotoxic chemotherapy. The subgroup of non-squamous NSCLC patients who benefit most from systemic treatment are those who receive targeted therapies based on the presence of a specific actionable oncogenic driver mutation. For patients where this option is not available and who have progressed during or after platinum based chemotherapy, treatment using an immune checkpoint blocker (PD-1 antibody) is an option. However, the clinical benefit remains modest. Non-small cell lung carcinoma patients with non-squamous cancer histology treated with nivolumab as second line treatment had a PFS of 2.3 months, and response rate was 19% by RECIST1.1 criteria. The median survival time was 12.2 months ([Borghaei et al 2015](#)).

Recently pembrolizumab was approved as monotherapy for first line treatment in patients with $\geq 50\%$ PD-L1 overexpression, based on data showing significantly improved survival vs chemotherapy (HR=0.60; 95% CI: 0.41,0.89; p=0.005) ([Reck et al 2016](#)).

A Phase II study also reported pembrolizumab plus chemotherapy has better efficacy than chemotherapy alone in non-selected patients as first line treatment in NSCLC ([Langer et al 2016](#)). But this combination use is not yet approved in European region. Despite the advances in immune checkpoint inhibitors in NSCLC, data suggested patients with EGFR mutation may not benefit as much as those with wild type EGFR from these type of treatment ([Rittmeyer et](#)

al 2017). Small molecule targeted therapy remains as the backbone treatment for NSCLC patients with an actionable mutation.

Natural history of the indicated condition in the population, including mortality and morbidity

In the US, the mortality rates for cancer of the lung and bronchus were 55.9 per 100000 men and 36.3 per 100000 women. The mortality rates (per 100000) in men and women were 55.9 and 37.5 among Caucasians, 68.0 and 34.6 among Blacks, 31.7 and 18.0 among Asian/Pacific Islanders, 46.3 and 30.8 among American Indian/Alaska Natives, and 27.3 and 13.4 among Hispanics (SEER 2017).

From 2010-2014, the median age at death for cancer of the lung and bronchus was 72 years. Approximately 0.1% died between 20 and 34; 0.8% between 35 and 44; 7.0% between 45 and 54; 20.0% between 55 and 64; 31.3% between 65 and 74; 28.7% between 75 and 84; and 12.0% for those over 84 years of age (SEER 2017).

The age-adjusted death rate was 44.7 per 100000 men and women per year based on data from patients who died between 2010 and 2014 in the US. The overall 5-year relative survival for 2007-2013 from 18 SEER geographic areas was 18.1% (SEER 2017).

Litvak et al (2014) investigated the OS of patients who were diagnosed with BRAF mutant lung adenocarcinomas between 2009 and 2013. The study included 36 patients who had a V600 mutation and 27 patients with a non-V600 mutation. In patients with Stage IIIB or IV BRAF mutant lung adenocarcinomas, those with V600 mutations had a longer 3-year OS as compared to patients with non-V600 mutations (24% versus 0%, $p<0.001$). The 3-year OS after resection of early stage lung cancer was similar for patients with V600 mutant tumors compared to non-V600 mutant tumors (67% vs 75%, $p=0.42$). Marchetti et al (2011) reported that in a series of 331 patients with lung adenocarcinoma including 21 patients with the BRAF V600E mutation and 310 patients with wild type tumors, patients with V600E BRAF mutations had shorter median disease-free survival (DFS) and OS than patients with wild type tumors (15.2 versus 52.1 months; $p<0.001$ and 29.3 versus 72.4 months; $p<0.001$, respectively). Cardarella et al (2013) reported no significant difference in OS between patients with advanced NSCLC who had BRAF mutations and those who had wild-type tumors. Barlesi et al (2016) reported the median OS for a group of 230 patients with mixed stage (mainly Stage III and IV) NSCLC who had BRAF mutation was 13.8 months.

Various adverse events can occur in patients with advanced or metastatic NSCLC. Data on adverse events in untreated patients were not available from population-based studies. The following information on patients receiving placebo is based on completed Phase III clinical trials reported in Clinicaltrials.gov.

In a randomized trial in patients with advanced or metastatic NSCLC that has not responded to standard therapy for advanced or metastatic cancer, serious adverse events (SAEs) occurred in 36.4% of patients in the placebo arm (NCT01000025). Such events with a frequency of over 1% included abdominal pain and vomiting (1.26% each), lung infection (5.02%), sepsis (1.26%), other neoplasms (17.99%), stroke (1.26%), bronchopulmonary hemorrhage (1.67%), dyspnea (5.86%), pleural effusion (1.26%), and thromboembolic event (1.26%).

In Study NCT00556712, among patients with advanced, recurrent, or metastatic NSCLC with previous platinum-based chemotherapy who have not had disease progression or unacceptable toxicity during chemotherapy, SAEs were reported in 7.64% of patients in the placebo arm. No specific SAEs were reported to occur in more than 1% of patients in the placebo arm.

In Study NCT00404924 on patients with NSCLC, whose disease has recurred after previous chemotherapy and an EGFR Tyrosine Kinase Inhibitor (EGFR TKI), 20.79% of patients in the placebo arm experienced SAEs. Reported SAEs with a frequency of over 1% included dyspnea, pleural effusion and pulmonary embolism (1.65% of patients each) and pneumonia (1.98% of patients).

Important co-morbidities

[Islam et al \(2015\)](#) evaluated 5683 newly diagnosed lung cancer patients in the US and reported that the most common comorbidities were chronic pulmonary disease (52.5%), diabetes (15.7%), congestive heart failure (12.9%), peripheral vascular disease (8.8%), cerebrovascular disease (7%), myocardial infarction (6.8%) and renal disease (5.7%). [Janssen-Heijnen et al \(1998\)](#) evaluated 3864 lung cancer patients in the Netherlands between 1993 and 1995. The most frequent concomitant diseases reported were cardiovascular diseases (23%), chronic obstructive pulmonary diseases (COPD) (22%), other malignancies (15%), hypertension (12%), and diabetes (7%).

2.1.3 Indication: Treatment of adult patients with differentiated thyroid cancer with BRAF V600E mutation

Incidence

The incidence of DTC has risen substantially across various global regions in recent decades, with consistently higher rates in women than in men. Reported age-standardized or age-adjusted incidence rates (AAIR) range is from approximately 15.2 ([Abe et al 2024](#)) to 79.6 ([Cho et al 2013](#)) per 100,000 in women and 3.7 ([Pellegriti et al 2009](#)) to 15.4 ([Cho et al 2013](#)) per 100,000 in men. In Italy, a population-based study from Sicily that included 534 histologically confirmed DTC cases reported incidence rates of 17.8 per 100,000 per year in women and 3.7 in men during 2002-2004. Notably, in the province of Catania, these figures increased to 31.7 in women and 6.4 in men. The BRAF V600E mutation was detected in 52% of tumors in Catania compared to 33% in other Sicilian regions, though incidence data were not stratified by mutation status ([Pellegriti et al 2009](#)).

In the United States (US), data from the Multiethnic Cohort Study involving 118,344 women, reported 373 incident DTC cases over a mean follow-up of 19.8 years. This corresponds to a crude incidence rate of 15.9 per 100,000 person-years. The AAIR was 17.4 per 100,000 in women and 6.8 in men, while the AAIR for the DTC subtype specifically was 15.2 in women and 5.5 in men. The study did not provide stratification by BRAF mutation or subtype beyond DTC ([Abe et al 2024](#)).

Prevalence

While many studies provide insight into the molecular epidemiology of DTC, population-based prevalence estimates of DTC, either overall or stratified by mutation status, were not identified

in the literature. Most available data are derived from clinical or hospital-based cohorts and cannot be extrapolated to the general population due to the absence of denominator data. Nevertheless, BRAF V600E mutation is consistently reported as highly prevalent in adult DTC patients, with rates ranging from 62% to 79% across large-scale cohorts and pooled analyses (Hong et al 2014, Brumfield et al 2025, Wei et al 2022, Henke et al 2015, Kowalska et al 2016). Higher prevalence has been associated with increasing age and, in some cases, male sex (Hong et al 2014, Wei et al 2022).

Demographics of the population in the authorized indication – age, gender, racial and/or ethnic origin and risk factors for the disease

Age

Differentiated thyroid cancer predominantly affects adults in middle age, with most studies reporting a mean or median age at diagnosis between 42 and 48 years (Zhou et al 2024, De Sousa et al 2024, Shi et al 2015, Kowalska et al 2016). Population-based data consistently show that peak incidence occurs between the ages of 40 and 59, particularly among women (Lam et al 2022, Jung et al 2014, Zhou et al 2024). In South Korea, the highest age-specific incidence was observed in the 40-49 age group (Jung et al 2014), while recent SEER data indicate a median age of 51 years, with most diagnoses occurring between 45 and 64 years (SEER 2025). In a large US cohort studied by Abe et al (2024), the mean age was 59.7 years, reflecting an older baseline population aged 45-75 at enrollment. Brumfield et al (2025) similarly reported a mean age of 48.1 years in their clinical sample. Over time, there appears to be a shift toward older age at diagnosis, for instance, Romei et al (2012) documented an increase in median age from 39.4 years during 1996-2000 to 47.8 years in 2006-2010. Likewise, Hong et al (2014) observed a rise in mean age from 45.3 years in the 1995-2003 period to 47.8 years during 2009-2012.

Multiple studies have found a strong association between increasing age especially age ≥ 45 years and the presence of the BRAF V600E mutation. Kim et al (2012) reported higher mutation prevalence among patients aged ≥ 45 , while Daliri et al (2014) found that BRAF-positive patients had a mean age of 48.5 years, significantly older than the BRAF-negative group (mean 34.8 years). This trend was further supported by Wei et al (2022), who analyzed data from 9,908 DTC patients and identified age ≥ 45 as a significant risk factor for BRAF mutation (OR=1.39).

Gender

Differentiated thyroid cancer shows a consistent and marked female predominance across all age groups and study populations. Women represent between 70% and over 90% of diagnosed cases, with individual studies reporting female proportions ranging from 72% to 91% (Abe et al 2024, Henke et al 2015, De Sousa et al 2024, Shi et al 2015). Current SEER data show incidence rates of 19.7 per 100,000 for females and 7.3 per 100,000 for males in the US (SEER 2025).

While women are more commonly affected by DTC, several studies have suggested that male sex may be modestly associated with worse prognostic features, including a higher prevalence of the BRAF V600E mutation. Kim et al (2012) identified male gender as a predictor of BRAF mutation (OR=1.834), and Wei et al (2022) confirmed this association with a smaller effect size (OR=1.13).

Racial and Ethnic Origin

The racial and ethnic distribution of DTC shows considerable variability across populations. In the US based Multiethnic Cohort Study, [Abe et al \(2024\)](#) found that Filipino and Latino women were overrepresented among DTC cases, comprising 8.3% and 29.5% of the cohort, respectively, compared to their proportions in the general population (3.8% and 20.8%). Similarly, [Henke et al \(2015\)](#) reported that 83.3% of DTC patients in their US cohort were White, followed by 9.8% Black, 4.9% Asian, and 2.0% Hispanic. [Kitahara et al \(2022\)](#) also observed that thyroid cancer incidence was highest among non-Hispanic White populations and lowest among Black and Native American/Alaska Native groups. [SEER 2025](#) data confirm these disparities: among women, the highest DTC incidence rates were observed in non-Hispanic Asian/Pacific Islanders (21.8 per 100,000), followed by Hispanic (20.7), non-Hispanic White (20.6), non-Hispanic American Indian/Alaska Native (18.0), and non-Hispanic Black women (11.8 per 100,000) ([SEER 2025](#)).

Asian countries, particularly South Korea and China, consistently report the highest thyroid cancer incidence rates globally, largely driven by the papillary subtype. In Korea, [Jung et al \(2014\)](#) documented a 15-fold increase in thyroid cancer incidence between 1993 and 2011, with nearly all cases being papillary carcinoma. [Ahn et al \(2016\)](#) supported this finding and attributed the dramatic rise primarily to widespread use of diagnostic ultrasonography, which led to the detection of many subclinical or indolent tumors. In China, [Zhou et al \(2024\)](#) reported an age-standardized incidence rate of 10.7 per 100,000, again with higher rates in women than in men.

Risk factors

Age is one of the most consistent risk factors associated with BRAF V600E mutation in DTC. Several studies demonstrate a clear trend of increasing mutation prevalence with advancing age. [Kim et al \(2012\)](#) reported a significant association between age ≥ 45 years and BRAF mutation. [Daliri et al \(2014\)](#) observed that the mean age was 48.5 years in BRAF-positive patients versus 34.8 years in mutation-negative ones ($p=0.023$). [Wei et al \(2022\)](#), in a large cohort of 9,908 patients, confirmed this with an odds ratio (OR) of 1.39 (95% CI: 1.21-1.60) for those aged ≥ 45 years compared to other ages. Pessoa-Pereira et al (2019) also found that older age increased the likelihood of BRAF mutation (OR=1.15, 95% CI: 1.00-1.33).

Sex-based associations with BRAF mutation are less consistent but noteworthy. While DTC occurs more frequently in women, some studies suggest that male gender may be a modest independent risk factor for BRAF positivity or worse outcomes. [Kim et al \(2012\)](#) reported an OR of 1.834 (95% CI: 1.021–3.463) for males.

Environmental and lifestyle exposures are additional contributors. [Kitahara et al \(2022\)](#) found that obesity (BMI ≥ 30) was more strongly associated with BRAF-mutant DTC, while smoking was inversely associated, though no ORs were provided. [Romei et al \(2012\)](#) and [Jung et al \(2014\)](#) emphasized that ionizing radiation exposure, while a known risk factor for DTC, is more commonly linked to Rearranged during transfection receptor (RET)/ Papillary thyroid cancer (PTC) rearrangements than BRAF. However, high iodine intake was associated with increased BRAF mutation prevalence. [Pellegriti et al \(2009\)](#) further reported a higher incidence and BRAF mutation rate in urban areas of Sicily, with no link to industrial pollution or iodine deficiency.

The main existing treatment options

Differentiated thyroid cancer is typically treated with surgery and subsequent therapy is dependent on the risk of disease recurrence. Most patients will require suppressive hormone therapy to prevent hypothyroidism and to minimize tumor growth stimulation through thyroid stimulating hormone (TSH). Radioiodine therapy may be administered to (1) ablate residual normal thyroid tissue, (2) to eliminate subclinical adjuvant therapy, or (3) to treat clinically apparent residual metastatic disease. The decision to treat with RAI therapy is dependent on the risk of recurrence. For patients who present as recurrent metastatic disease RAI therapy is a critical mainstay of therapy.

Approximately 10-15% of patients do not trap iodine and are considered RAI-refractory. For these cases, the 10-year survival rate is 10% from the time of detection of metastasis ([Gianoukakis et al 2018](#), [Ancker et al 2020](#)).

If disease progression cannot be controlled with local therapies such as surgery or radiotherapy, systemic therapy with chemotherapy can be considered. However, treatment with cytotoxic chemotherapy has shown disappointing results as complete remissions are rare and long-term responses are limited.

Advances in the understanding of the molecular pathogenesis of thyroid cancer have led to the development of targeted agents aimed at combating advanced disease. The novel treatment with multi-targeted tyrosine kinase inhibitors (MKIs) has shown favourable results in otherwise treatment-resistant thyroid cancer. The primary target for all of these effective tyrosine kinase inhibitors is angiogenic signaling in the tumor microenvironment, particularly the VEGFR family. The MKIs block signaling from the tyrosine kinase receptors, thus preventing phosphorylation and, ultimately angiogenesis and tumor growth. The first targeted agents which have shown to improve PFS was the multikinase inhibitor, sorafenib, approved for use in patients with RAI-refractory DTC. Sorafenib is a protein kinase inhibitor with activity against multiple protein kinases, including VEGFR, PDGFR and RAF kinases. Of the RAF kinases, Sorafenib is more selective for C-RAF than B-RAF. In the DECISION trial, patients who had RAI-refractory locally advanced or metastatic DTC that had progressed within the past 14 months were randomly assigned to sorafenib or placebo. Median PFS was significantly longer in the sorafenib group (10.8 months) than in the placebo group (5.8 months; HR 0.59, 95%CI 0.45-0.76, $p < 0.0001$). Improvement of PFS in the sorafenib group compared to the placebo group has been observed in all prespecified clinical and genetic biomarker subgroups. The partial response rate was 12.2 % and stable disease for 6 months or longer was achieved in 41.8% of patients ([Brose et al 2014](#)). More recently, lenvatinib, another multi-kinase inhibitor, was approved for the treatment of locally recurrent or metastatic, progressive RAI-refractory DTC. Lenvatinib acts as a multi-tyrosine kinase inhibitor and inhibits vascular endothelial growth factor receptor family (VEGFR1-3), fibroblast growth factor receptor family (FGFR1-4), platelet-derived growth factor receptor- α (PDGFR α), tyrosine-kinase receptor (KIT) and RET. In the SELECT study, Lenvatinib was shown to prolong PFS compared with placebo (median 18.3 vs 3.6 months; HR 0.21; 99% CI 0.14-0.31) ([Schlumberger et al 2015](#)). Cabozantinib is an oral multikinase inhibitor targeting MET in addition to VEGFR and is approved for medullary thyroid cancer. A single-arm open-label Phase II study of cabozantinib in patients with metastatic, RAI-refractory DTC and evidence of progression on prior VEGFR-targeted therapy showed of the 25 patients, 10 (40%) had a partial response, 13 (52%) had stable

disease, and 2 (8%) had non-evaluable disease. The median PFS and overall survival were 12.7 months and 34.7 months, respectively ([Cabanillas et al 2017](#)). Newly available targeted therapies show improvements in PFS for patients with RAI-refractory DTC. Sorafenib and lenvatinib are approved as first line indication in refractory RAI DTC in the US, China, Japan, and other countries. Adverse effects of VEGFR-targeted therapy may include hypertension, renal toxicity, proteinuria, arthralgia/myalgia, headache, bleeding, myelosuppression, arterial thromboembolism, cardiotoxicity including prolonged QT intervals and risk for arrhythmia, thyroid dysfunction, cutaneous toxicity including hand-foot skin reaction, delayed wound healing, hepatotoxicity, nausea, vomiting, diarrhea, muscle wasting, fistula formation, osteonecrosis of the jaw.

Natural history of the indicated condition in the population, including mortality and morbidity

Differentiated thyroid cancer generally has a good prognosis, with 10-year survival rates exceeding 90% ([Henke et al 2015](#), [Kitahara et al 2022](#), [Lam et al 2022](#)). While BRAF V600E is linked to aggressive features such as lymph node metastasis and extrathyroidal extension ([Henke et al 2015](#), [Wei et al 2022](#)), its impact on recurrence and survival is inconsistent: some studies show no association ([Brumfield et al 2025](#), [Daliri et al 2014](#)), while others report increased risk, especially when co-mutated with Telomerase reverse transcriptase (TERT) ([Kitahara et al 2022](#), [Wei et al 2022](#)). Specifically, [Brumfield et al \(2025\)](#) studied 301 DTC patients over a median follow-up of 23.4 months. Among them, 79% were BRAF mutation-positive. Recurrence occurred in 15%, distant metastases in 3%, and death in 1%. No statistically significant differences were seen in outcomes between mutation-positive and negative groups. Follow-up duration was longer in the BRAF-positive group (24.5 vs. 19.5 months), but this did not impact event rates ([Brumfield et al 2025](#)). [Daliri et al \(2014\)](#) followed 69 DTC patients over a mean of 63.9 months. Incomplete response at 1 year was 32.1% in BRAF-positive patients versus 39% in BRAF-negative ones ($p=0.8$), and final remission rates were comparable (78% vs. 78.6%, $p=0.6$). Among those with early incomplete response, later remission occurred in 57.1% of BRAF-positive and 70% of BRAF-negative patients ($p=0.44$), suggesting that BRAF did not significantly impact long-term outcome ([Daliri et al 2014](#)).

Important co-morbidities

Hashimoto thyroiditis appears to be inversely associated with BRAF V600E mutation in DTC, with several studies showing significantly lower mutation rates in patients with thyroiditis ([Kyeong et al 2009](#), [Wang et al 2012](#), [Kim et al 2018](#)). Other comorbidities such as hypertension and autoimmune conditions (e.g., lupus, Basedow disease) were also noted, with mixed or unclear associations to BRAF status ([Yan et al 2019](#), [Kitahara et al 2022](#), [Zbitou et al 2022](#)).

[Kitahara et al \(2022\)](#) used US population-based data to examine associations between DTC and several benign comorbidities, including breast disorders, uterine fibroids, asthma, and autoimmune diseases like lupus and Sjögren's syndrome. Smoking showed an inverse association with DTC, particularly in BRAF-mutant tumors, suggesting possible behavioral or hormonal modulation of mutation risk ([Kitahara et al 2022](#)).

2.1.4 Indication: Treatment of paediatric glioma with BRAF V600E mutation (low and high grade glioma with BRAF V600E mutation)

Epidemiology data on paediatric gliomas, including grading (high grade vs. low grade glioma) and presence of BRAF V600E mutation, are limited. However, there is a greater degree of epidemiology data for paediatric central nervous system (CNS) tumors or for gliomas of specific histology. Therefore, many estimates reported in this review have been estimations based on either, (1) the rates of paediatric CNS tumors and the proportion of these that are paediatric gliomas, or (2) the sums of data broken down by specific histologies that comprise all paediatric gliomas. All reported values that are based on estimations, are noted as such in the tables.

Methods

For population-based registry databases reporting the incidence of all paediatric CNS tumors, the incidence of paediatric gliomas was estimated from incidence of all brain and CNS tumors in children and the proportion of all paediatric CNS tumors that are gliomas in Europe and North America (range 45-65%, unweighted average of 56.2% used for estimations) ([Ostrom et al 2021](#), [Erdmann et al 2020](#), [Desandes et al 2014](#), [Rosychuk et al 2012](#)).

The incidence of paediatric low grade glioma (LGG) and high grade glioma (HGG) are estimated from the proportion of all paediatric gliomas that are LGG and HGG in the US and Australia (range 61-72% LGG, 28-39% HGG, unweighted average 66.5% LGG and 33.5% HGG, used for estimations) ([Ostrom et al 2021](#), [Youlden et al 2021](#)).

The estimation of the proportion of paediatric LGG and HGG that are BRAF V600E mutation positive was calculated as follows. The unweighted average of two estimates of the proportion of paediatric LGG that are BRAF V600E mutation positive is 10.5% (range 9-12%), and from this estimate and the proportion of BRAF V600E mutation positive paediatric gliomas that are HGG (16%), it is possible to estimate the proportion of paediatric HGG tumors that are BRAF V600E mutation positive (estimated to be 4%) ([Nobre et al 2020](#), [Gierke et al 2016](#), [Horbinski et al 2012](#)).

Incidence

[Table 2-8](#) provides the incidence of paediatric gliomas by world region or country. Worldwide the age-adjusted incidence is 0.7 to 1.04 per 100,000 for all paediatric gliomas (0.5 to 0.7 for LGG, 0.2 to 0.34 for HGG, 0.05 to 0.07 for BRAF V600E mutation positive LGG, and 0.01 for BRAF V600E mutation positive HGG) ([GBD 2022](#), [Globocan 2022](#)).

Estimates of the age-adjusted incidence of paediatric glioma for Europe range from 0.71 to 1.49 per 100,000 persons, and specifically, 0.47 to 0.99 per 100,000 persons for LGG and 0.24 to 0.50 per 100,000 persons for HGG, and 0.05 to 0.10 per 100,000 persons for BRAF V600E mutation positive LGG, and 0.01 to 0.02 per 100,000 persons for BRAF V600E mutation positive HGG ([GBD 2022](#), [Globocan 2022](#), [Rarecarenet 2022](#)). Within Europe, age-adjusted incidence per 100,000 persons is highest in Germany and the Nordic countries (2.4 to 2.7 for all paediatric gliomas, 1.6 to 1.8 for LGG, 0.8 to 0.9 for HGG, 0.17 to 0.19 for BRAF V600E mutation positive LGG, and 0.03 to 0.04 for BRAF V600E mutation positive HGG) and lowest in France (1.96 to 2.24 for all paediatric gliomas, 1.30 to 1.49 for LGG, 0.69 to 0.75 for HGG,

0.14 to 0.16 for BRAF V600E mutation positive LGG, and 0.03 for BRAF V600E mutation positive HGG) ([Nordcan 2022](#), [Erdmann et al 2020](#), [Coll et al 2015](#), [Desandes et al 2014](#)).

The age-adjusted incidence of paediatric glioma in the US is 2.6 to 3.12 per 100,000 persons (1.7 to 2.07 for LGG and 0.9 to 1.05 for HGG, 0.18-0.22 for BRAF V600E mutation positive LGG, and 0.04 for BRAF V600E mutation positive HGG) ([SEER 2022](#), [Ostrom et al 2021](#)).

Table 2-8 Incidence of paediatric CNS tumors, paediatric gliomas, LGG, HGG and BRAF V600E

Setting	Incidence per 100,000 persons, age-standardized						References
	All paediatric CNS tumors	All paediatric gliomas	LGG	BRAF V600E+ LGG	HGG	BRAF V600E+ HGG	
Europe (WHO region)	2.2-2.65	1.2-1.49*	0.80-0.99†	0.08-0.10‡	0.40-0.50**	0.02††	GBD (2022) , Globocan (2022)
EU27	1.26	0.71*	0.47†	0.05‡	0.23**	0.01††	
Southern and Southeastern Europe	-	-	-	-	0.98-1.34**	0.04-0.05††	Papathoma et al (2015)
Nordic countries	4.3-4.8	2.4-2.7*	1.6-1.8†	0.17-0.19‡	0.8-0.9**	0.03-0.04††	Nordcan (2022)
France	-	1.96-2.24§	1.30-1.49†	0.14-0.16‡	0.66-0.75**	0.03††	Coll et al (2015) , Desandes et al (2014)
Germany	-	2.49§	1.58-1.66†	0.17‡	0.83**	0.03††	
Netherlands	-	2.30§	1.53†	0.16‡	0.81**	0.03††	Reedijk et al (2020)
UK	-	2.29§	1.56†	0.16‡	0.77**	0.03††	Stiller et al (2019)
US and Canada	2.77-3.2	1.56-1.8*	1.04-1.2†	0.11-0.13‡	0.52-0.6**	0.02††	GBD (2022) , Globocan (2022)
US	-	2.60-3.12§	1.73-2.07†	0.18-0.22‡	0.87-1.05**	0.03-0.04††	
Canada	4.3	2.88§	1.92†	0.20‡	0.96**	0.04††	Rosychuk et al (2012)
Worldwide	1.2-1.85	0.7-1.04*	0.5-0.69†	0.05-0.07‡	0.2-0.35**	0.01-0.02††	GBD (2022) , Globocan (2022)

* All paediatric gliomas estimated as Incidence of all CNS tumors $\times$ 0.562, § All paediatric gliomas estimated as sum of astrocytomas, ependymomas oligodendrogliomas and other gliomas, † LGG estimated as Incidence of all paediatric gliomas $\times$ 0.665, ** HGG estimated as Incidence of all paediatric gliomas $\times$ 0.335, ‡ BRAF V600E mutation positive LGG estimated as Incidence of LGG $\times$ 0.105, †† BRAF V600E mutation positive HGG estimated as Incidence of HGG $\times$ 0.04, CNS: central nervous system, EU27: the 27 EU states in 2012 (including UK), GBD: global burden of disease, HGG: high grade glioma, LGG: low * grade glioma, SEER: Surveillance, Epidemiology and End Results, UK: United Kingdom, US: United States

Prevalence

For population-based registry databases reporting the prevalence of all paediatric CNS tumors, the prevalence of paediatric gliomas was estimated from prevalence of all brain and CNS tumors in children and the proportion of all incident paediatric CNS tumors that are gliomas in Europe and North America (range 45-65%, unweighted average of 56.2% used for estimations) ([Ostrom et al 2021](#), [Erdmann et al 2020](#), [Desandes et al 2014](#), [Rosychuk et al 2012](#)). Where each type of glioma is reported separately, the prevalence for all gliomas is summed across histological subtypes. The prevalence of paediatric LGG and HGG are estimated from the proportion of all incident paediatric gliomas that are LGG and HGG in the US and Australia (range 61-72% LGG, 28-39% HGG, unweighted average 66.5% LGG and 33.5% HGG, used for estimations) ([Ostrom et al 2021](#), [Youlden et al 2021](#)). This method of estimation may overestimate the prevalence of HGG and underestimate the prevalence of LGG, due to differences in survival between LGG and HGG.

The estimation of the proportion of paediatric LGG and HGG that are BRAF V600E mutation positive was calculated as follows. The unweighted average of two estimates of the proportion of paediatric LGG that are BRAF V600E mutation positive is 10.5% (range 9-12%), and from this estimate and the proportion of BRAF V600E mutation paediatric gliomas that are HGG (16%), it is possible to estimate the proportion of paediatric HGG tumors that are BRAF V600E mutation positive (estimated to be 4%) ([Nobre et al 2020](#), [Gierke et al 2016](#), [Horbinski et al 2012](#)).

For Europe, the estimated prevalence of LGG in children in 2019 was 5.81 per 100,000 persons, and the estimated prevalence of HGG in children in 2019 was 2.93 per 100,000 persons (GBD 2022). The estimated 10-year period prevalence in children in Nordic countries is 6.5 to 7.5 per 100,000 persons for LGG and 3.2 to 3.9 per 100,000 persons for HGG ([Nordcan 2022](#)). In Europe, the estimated prevalence of BRAF V600E mutation positive LGG in children in 2019 was 0.61 per 100,000 persons and the estimated prevalence of BRAF V600E mutation positive HGG in children in 2019 was 0.12 per 100,000 persons ([GBD 2022](#)). For the Nordic countries, the estimated 10-year period prevalence in children is 0.7 to 0.8 per 100,000 for BRAF V600E mutation positive LGG and 0.1 to 0.2 per 100,000 for BRAF V600E mutation positive HGG ([Nordcan 2022](#)).

For the US and Canada, the estimated prevalence of LGG in children in 2019 was 7.09 per 100,000 persons, and for HGG it was 3.57 per 100,000 persons ([GBD 2022](#)). In the US, the 26-year period prevalence in children is estimated to be 10 per 100,000 persons for LGG and 5 per 100,000 persons for HGG ([SEER 2022](#)). The 26-year period prevalence of BRAF V600E mutation positive LGG in US children is estimated to be 0.74 per 100,000 persons, and the 26-year period prevalence of BRAF V600E mutation positive HGG in US children is estimated to be 0.14 per 100,000 persons ([SEER 2022](#)). [Table 2-9](#) provides the prevalence of paediatric gliomas by region or country.

Table 2-9 Prevalence of paediatric CNS tumors, LGG, HGG and BRAF V600E

Setting	Outcome definition	5-y period prevalence (per 100,000)	10-y period prevalence (per 100,000)	26-y period prevalence (per 100,000)	Prevalence in 2019 (per 100,000)	References
Europe (WHO region)	All CNS tumors	7.0	-	-	15.56	GBD (2022), Globocan (2022)
Europe (WHO region)	LGG	2.6†	-	-	5.81†	GBD (2022), Globocan (2022)
Europe (WHO region)	HGG	1.3**	-	-	2.93**	GBD (2022), Globocan (2022)
Europe (WHO region)	BRAFV600E mutation+ LGG	0.3‡	-	-	0.61‡	GBD (2022), Globocan (2022)
Europe (WHO region)	BRAFV600E mutation+ HGG	0.05††	-	-	0.12††	GBD (2022), Globocan (2022)
Nordic countries	All CNS tumors	17.2-20.1	33.7-39.2	-	-	Nordcan (2022)
Nordic countries	LGG	6.5-7.5†	12.6-14.6†	-	-	Nordcan (2022)
Nordic countries	HGG	3.2-3.9**	6.3-7.4**	-	-	Nordcan (2022)
Nordic countries	BRAFV600E mutation+ LGG	0.7-0.8‡	1.3-1.5‡	-	-	Nordcan (2022)
Nordic countries	BRAFV600E mutation+ HGG	0.1-0.2††	0.3††	-	-	Nordcan (2022)
US and Canada	All CNS tumors	11.3	-	-	18.96	GBD (2022), Globocan (2022)
US and Canada	LGG	4.3†	-	-	7.09†	GBD (2022), Globocan (2022)
US and Canada	HGG	2.1**	-	-	3.57**	GBD (2022), Globocan (2022)

Setting	Outcome definition	5-y period prevalence (per 100,000)	10-y period prevalence (per 100,000)	26-y period prevalence (per 100,000)	Prevalence in 2019 (per 100,000)	References
US and Canada	BRAFV600E mutation+ LGG	0.5‡	-	-	0.75‡	GBD (2022), Globocan (2022)
US and Canada	BRAFV600E mutation+ HGG	0.1††	-	-	0.14††	GBD (2022), Globocan (2022)
US	LGG	-	-	10*	-	SEER (2022)
US	HGG	-	-	5***	-	SEER (2022)
US	BRAFV600E mutation+ LGG	-	-	1.1‡	-	SEER (2022)
US	BRAFV600E mutation+ HGG	-	-	0.2††	-	SEER (2022)
Worldwide	All CNS tumors	3.3	-	-	8.24	GBD (2022), Globocan (2022)
Worldwide	LGG	1.2†	-	-	3.08†	GBD (2022), Globocan (2022)
Worldwide	HGG	0.6**	-	-	1.56**	GBD (2022), Globocan (2022)
Worldwide	BRAFV600E mutation+ LGG	0.1‡	-	-	0.32‡	GBD (2022), Globocan (2022)
Worldwide	BRAFV600E mutation+ HGG	0.02††	-	-	0.06††	GBD (2022), Globocan (2022)

†LGG estimated as Prevalence of all CNS tumors × 0.562 × 0.665, * LGG estimated as sum of all paediatric gliomas × 0.665, ** HGG estimated as Prevalence of all CNS tumors × 0.562 × 0.335, *** HGG estimated as sum of all paediatric gliomas × 0.335, ‡ BRAF V600E mutation positive LGG estimated as Prevalence of LGG × 0.105, †† BRAF V600E mutation positive HGG estimated as Prevalence of HGG × 0.04, CNS: central nervous system, GBD: global burden of disease, HGG: high grade glioma, LGG: low grade glioma, SEER: Surveillance, Epidemiology and End Results, US: United States, y: year

Demographics of the population in the authorized indication – age, gender, racial and/or ethnic origin and risk factors for the disease

Globally, between 30% and 40% of paediatric gliomas are diagnosed before the age of five years, and there is a slight male preponderance (51-57% males) ([Youlden et al 2021](#), [Erdmann et al 2020](#), [Rosychuk et al 2012](#)). In the US, 57-81% of paediatric gliomas are in Non-Hispanic whites or whites including white Hispanics, 18-24% are in Hispanics (white and non-white), 13-14% are in blacks, 5-6% are in Asians or Pacific Islanders, and 1-2% are in American Indians and Alaska Natives ([Ostrom et al 2021](#), [Jiang et al 2020](#)). Table 2-10 Demographic characteristics of paediatric gliomas provides the demographic characteristics of paediatric glioma patients in Europe and the US and Canada.

Table 2-10 Demographic characteristics of paediatric gliomas

Setting	Outcome definition	Age (%)	Sex (%)	Race (%)	References
Germany	All gliomas	Age < 1 y: 6%* Age 1-4 y: 27%* Age 5-9 y: 28%* Age 10-14 y: 27%* Age 15-17 y: 12%*	Males: 54%* Females: 48%*	-	Erdmann et al (2020)
US	All gliomas	Age 0-4 y: 29%* Age 5-9 y: 27%* Age 10-14 y: 24%* Age 15-19 y: 20%*	-	White: 81%* Black: 13%* Asian or Pacific Islander: 5%* American Indian or Alaska Native: 1%* Non-Hispanic: 82%* Hispanic: 18%*	Ostrom et al (2021)
US	All gliomas and medulloblastomas	-	-	57% Non-Hispanic Whites 24% Hispanic whites 12% Blacks 6% Asians or Pacific Islanders 2% American Indians or Alaska Natives	Jiang et al (2020)
Canada	All brain and CNS tumors	Age < 1 y: 6% Age 1-4 y: 24% Age 5-9 y: 28% Age 10-14 y: 21%	Male: 57% Female: 43%	-	Rosychuk et al (2012)

Setting	Outcome definition	Age (%)	Sex (%)	Race (%)	References
Australia	All gliomas	Age 15-19 y: 21% Age 0-4 y: 40%* Age 5-9 y: 32%* Age 10-14 y: 27%*	Male: 51%* Female: 49%*	-	Youlden et al (2021)

* Proportions are taken for all paediatric gliomas, which are summed across the histologies that comprise gliomas, CNS: central nervous system, US: United States, y: year

In the US, the incidence of malignant brainstem gliomas among children is significantly higher in Hispanics than in non-Hispanic whites (Patil et al 2021). Non-white race is associated with reduced risk of incident ependymal tumors in US children (Zhang et al 2020). Prenatal pesticide and diesel exhaust exposures are associated with an increased risk of incident astrocytomas and ependymomas (Lombardi et al 2021, Volk et al 2019). For instance, Danish children whose mothers are employed in industries with diesel exhaust exposure have an increased risk of incident astrocytomas with an odds ratio (OR) of 1.5, with a 95% Confidence Interval (CI) of 1.0-2.1 (Volk et al 2019).

Natural history of the indicated condition in the population, including mortality and morbidity

Survival is closely linked to histology, tumor site, age at diagnosis and tumor grade. OS in paediatric gliomas ranges from 1-year OS of 91-99% in LGG to a 1-year OS of 50 to 60% in HGG (Youlden et al 2021, Tabash et al 2019, Ostrom et al 2021), and a 5-year OS of 78 to 98% in LGG and 16 to 70% in HGG (SEER 2022, Gnekow et al 2021, Ostrom et al 2021, Youlden et al 2021, Napieralska et al 2021a, Tabash et al 2019). Long-term survival is good for paediatric LGG with a 10-year OS of 75 to 98% (Gnekow et al 2021, Ostrom et al 2021, Youlden et al 2021), but poor for paediatric HGG with a 10-year OS of 14 to 69% (SEER 2022, Napieralska et al 2021a, Ostrom et al 2021, Youlden et al 2021). Survival outcomes in BRAF V600E mutation-positive paediatric HGG are poor compared to other HGG, but in paediatric LGG the mutation is not associated with differences in prognosis (Nobre et al 2020, Horbinski et al 2012). Table 2-11 provides the 1-year, 2-year, 5-year and 10-year OS or relative survival (RS) of paediatric glioma patients by country and histological subtype. Only the SEER data report RS, all other sources in Table 2-11 report OS.

Table 2-11 Survival in paediatric gliomas

Setting	Grade	Glioma histology	1-y OS or RS	2-y OS	5-y OS or RS	10-y OS	References
Germany	LGG	All LGG	--	--	98%	98%	Gnekow et al (2021)
Poland	HGG	Primary HGG	78%	48%	30%	17%	Napieralska et al (2021a)
Poland	Both	Ependymomas	98%	95%	83%	73%	Napieralska et al (2021b)

Setting	Grade	Glioma histology	1-y OS or RS	2-y OS	5-y OS or RS	10-y OS	References
US	HGG	Glioblastomas	57- 58%*	-	15- 19%*	16%	SEER (2022), Ostrom et al (2021)
US	Both	Brainstem gliomas	91%-	87%-	86%	--	Khalid et al (2019)
US	Both	Ependymal tumors	96%-	--	80%	72%-	Ostrom et al (2021)
US	LGG	Pilocytic astrocytomas	99%	--	95- 97%	96%	Tabash et al (2019), Ostrom et al (2021)
US	HGG	Anaplastic astrocytomas	66%	-	25%	19%	Ostrom et al (2021)
US	Both	Diffuse and anaplastic astrocytomas	83%*	-	45%*	-	SEER (2022)
US	Both	Diffuse astrocytomas	92%-	--	82%	80%-	Ostrom et al (2021)
US	Both	Oligodendrogliomas	97%	-	94%	92%	Ostrom et al (2021)
US	Both	Malignant gliomas, NOS	82%	-	70%	69%	Ostrom et al (2021)
US	Both	Other gliomas	92%*	-	86%*		SEER (2022)
Australia	LGG	Grade I astrocytomas	98%	-	96%	94%	Youlden et al (2021)
Australia	LGG	Grade II gliomas	91%	-	78%	75%	Youlden et al (2021)
Australia	HGG	Grade III ependymomas	90%	-	56%	51%	Youlden et al (2021)
Australia	HGG	Grade III astrocytomas	59%	-	30%	25%	Youlden et al (2021)
Australia	HGG	Grade IV astrocytomas	50%	-	16%	14%	Youlden et al (2021)
Australia	HGG	Grade III/IV gliomas, NOS	60%	-	44%	42%	Youlden et al (2021)

*SEER data are relative survival, all other data are overall survival, CNS: central nervous system, PNET: primitive neuroendocrine tumors, OS: overall survival, RS: relative survival, SEER: Surveillance, Epidemiology and End Results, US: United States, y: year

After adjusting for confounding factors, histology is strongly associated with OS or mortality risk in paediatric gliomas. The adjusted odds ratio (AOR) and adjusted hazard ratios (AHR) are as follows (if the histological type in the following list is always HGG or always LGG, this is noted in parentheses): for ependymoma (both LGG and HGG) vs. astrocytoma (both LGG and HGG) the AHR is 0.6 (95% CI 0.5-0.8), for anaplastic glioma (HGG) vs. pilocytic astrocytoma (LGG) the AOR is 7.8 (95% CI 3.2-19.2), for glioblastoma (HGG) vs. pilocytic astrocytoma

(LGG) the AOR is 36.5 (95% CI 18.3-72.7), for oligodendroglioma (both LGG and HGG) vs. pilocytic astrocytoma (LGG) the AOR is 3.8 (95% CI 1.4-9.8), for glioma not otherwise specified (NOS) (both LGG and HGG) vs. ependymomas (both LGG and HGG) the AHR is 13.0 (95% CI 2.5-67.5), and for astrocytoma (both LGG and HGG) NOS vs. ependymomas (both LGG and HGG) the AHR is 12.5 (95% CI 1.9-80.9) ([Jiang et al 2020](#), [Zhou et al 2020](#), [Khalid et al 2019](#)).

In LGGs, no survival difference was found between paediatric patients with and without BRAF V600E mutations ([Horbinski et al 2012](#)). In HGGs, no observational studies reported differences in survival by BRAF V600E mutation status, but one study reported 1-y PFS for BRAF V600E mutation positive paediatric HGGs (n=11) to be 27% (95% CI 10-72%) ([Nobre et al 2020](#), [Youlden et al 2021](#)). In this same small study, the 1-y PFS for BRAF V600E mutation positive paediatric LGGs (n=56) was 86% (95% CI 78-96%) ([Nobre et al 2020](#)). [Table 2-12](#) shows frequency of BRAF V600E mutations in paediatric gliomas and survival outcomes.

Neurocognitive impairments are common in paediatric gliomas, with a 25-y cumulative incidence (CuI) of 26% for at least one grade 3-5 neurological condition in survivors of paediatric astrocytomas in the US and Canada, and a 25-y CuI of 7% for paralysis ([Effinger et al 2019](#)). Other common complications in paediatric gliomas include visual acuity deficits (68% of neurofibromatosis-associated optic pathway gliomas, 25-y CuI of 19% in astrocytomas), auditory impairments (25-y CuI of 17% in astrocytomas), post-operative speech impairment (30% of posterior fossa tumors), subsequent neoplasms (25-y CuI of 7% in astrocytomas), cardiac conditions (25-y CuI of 8% in astrocytomas), endocrine conditions (25-y CuI of 6% in astrocytomas) and stroke (25-y CuI of 13% in astrocytomas) ([Kotch et al 2022](#), [Gronbaek et al 2021](#), [Effinger et al 2019](#)).

Neurofibromatosis type 1 (NF1) is associated with increased risk of subsequent neoplasms in survivors of paediatric glioma with a relative risk (RR) of 4.0 (95% CI 2.1-7.6) ([De Blank et al 2020](#)). Familial NF1 inheritance is associated with increased risk of relapsed/refractory optic-pathway paediatric gliomas with an adjusted risk ratio (ARR) of 2.2 (95% CI 1.2-3.9) ([Kotch et al 2022](#)). Other risk factors for relapsed/refractory optic pathway paediatric gliomas include age < 2 y at initial therapy (ARR 3.2, 95% CI 1.2-5.2) and posterior tumor location (ARR 2.2, 95% CI 1.1-4.1) ([Kotch et al 2022](#)).

Some tumor locations are associated with a reduced risk of post-operative speech impairments in paediatric posterior fossa gliomas: cerebellar vermis vs. fourth ventricle AOR 0.3 (95% CI 0.1-0.8), and hemispheric vs fourth ventricle AOR 0.2 (95% CI 0.1-0.7) ([Gronbaek et al 2021](#)). Survivors of paediatric ependymomas who become mothers are at increased risk of preterm birth, with an AOR of 2.8 (95% CI 1.2-6.5) ([Huang et al 2020](#)).

Table 2-12 Frequency of BRAF V600E mutations in paediatric gliomas and survival outcomes

Reference	Setting and study period	Design and data source	Study population (N)	Glioma grade and histology (n)	Frequency (%) BRAF V600E mutation positive	PFS (95% CI) or HR for PFS (95% CI)	FU / comments
Nobre et al (2020)	International, study period NR	Case series, chart review from 29 institutions	Age < 25 y, BRAF V600E mutation positive gliomas (N=67) treated with BRAF inhibitors, excluding those w/missing data or < 6 months FU, median age 4.8 y (range 0.1-22.3 y), 55%* male (37/67)	HGG (n=11) LGG (n=56)	100% 100%	1-y PFS: 27% (10-72%) 1-y PFS: 86% (78-96%)	FU ≥ 0.5 y (6/12) * % male: (37/67)
Gierke et al (2016)	Germany, study period NR	Retrospective observational, data source NR	Age 0-18 paediatric brain tumors (N=170), mean age NR, 56%* male (431/765)	Grade IV glioblastoma (n=10) Grade III anaplastic astrocytoma (n=3) Grade III anaplastic ependymoma (n=4)	0% 0% 0%	NR NR NR	* % male: (431/765)

Reference	Setting and study period	Design and data source	Study population (N)	Glioma grade and histology (n)	Frequency (%) BRAF V600E mutation positive	PFS (95% CI) or HR for PFS (95% CI)	FU / comments
				Grade II diffuse astrocytoma (n=6)	34%	NR	
				Grade II ependymoma (n=3)	0%	NR	
				Grade I pilocytic astrocytoma- (n=45)	2%	NR	
				Grade II pilomyxoid astrocytoma (n=3)	0%	NR	
				Grade II pleomorphic xanthoastrocytoma (n=5)	60%	NR	
				Grade I ganglioglioma / gangliocytoma (n=22)	55%	NR	
				All HGG	0%	NR	
Koelsche et al (2014)	Germany and Italy, study period NR	Case series, archives of three institutions	Desmoplastic infantile ganglioglioma. (N=16), Age < 24 months at diagnosis,	All LGG	9% (16/170)	NR	FU NR
				Grade I Desmoplastic infantile ganglioglioma.	13% (2/16)	NR	

Reference	Setting and study period	Design and data source	Study population (N)	Glioma grade and histology (n)	Frequency (%) BRAF V600E mutation positive	PFS (95% CI) or HR for PFS (95% CI)	FU / comments
			Median age at surgery 10.5 months (range 1-60 months), Female to male ratio 0.8				
Horbinski et al (2012)	US, study period NR	Retrospective cohort study	Paediatric non-NF1-related LGGs (N=198) (157 successfully analyzed for BRAF V600E mutation), median age 8.2 y, 56% male (111/198)	Grade I/II Pilocytic astrocytomas (n=110) Grade I/II gangliogliomas (n=22) Grade I/II pilomyxoid astrocytomas (n=5) Grade I/II pleomorphic xanthoastrocytomas (n=5) Other LGG (n=12) All LGG (n=154)	9% 23% 20% 40% 8% 12%* (19/154)	NR NR NR NR NR HR 2.4 (0.9-6.2) for BRAF V600E vs no BRAF V600E mutation	Median FU 6.3 y * % male: (111/198)

CI: confidence interval, FU: follow up, HGG: high grade glioma, HR: hazard ratio, LGG: low grade glioma, N: sample size, n: number of patients per histology/grade group, NF1: neurofibromatosis type 1, NR: not reported, PFS: progression free survival, y: years

Important comorbidities

The prevalence of any comorbidity in paediatric glioma patients ranges from 7% in posterior fossa tumors to 37% in astrocytomas ([Gronbaek et al 2021](#), [Effinger 2019](#)). Common comorbidities include neurological, psychiatric or speech problems (12% of posterior fossa tumors) ([Gronbaek et al 2021](#)). Compared to matched siblings, paediatric astrocytoma patients are more likely to experience poor general health (ARR 2.3, 95% CI 1.9-2.7), poor mental health (ARR 1.6, 95% CI 1.4-1.8), functional impairments (ARR 5.3, 95% CI 4.4-6.4) and activity limitations (ARR 1.9, 95% CI 1.6-2.3) ([Effinger et al 2019](#)).

In paediatric patients with LGG, the prevalence of NF1 is 17% ([Gnekow et al 2021](#)). Among paediatric patients with NF1-associated optic pathway gliomas, common comorbidities include central precocious puberty (72%), growth hormone deficiency (9%), diencephalic syndrome (12%), and growth hormone hyper-secretion (6%) ([Santoro et al 2020](#)). For more details on comorbidities in paediatric gliomas, see [Table 2-13](#).

Table 2-13 Morbidity and complications in paediatric glioma

Reference	Setting and study period	Design and data source	Study population (N)	Glioma definition	Complications and frequency Cul or OR	AEs: (%), FU comments
Kotch et al (2022)	US, 2005-2014	Retrospective cohort, chart review at 7 hospitals	Age ≤ 18 y, NF1-associated optic pathway gliomas (N=103), excluding those who received CT before 2005 or received radiation therapy as initial treatment, median age at diagnosis 2.1* y (25/12)	NF1-associated optic pathway gliomas	Visual acuity deficit at initiation of therapy: 68% Worsened visual acuity at last FU: 35% Relapsed/refractory OPG: 44%	Median FU 7.9* y (range 1.1* -15.4* y) * FU: (95/12), (13/12 and 185/12)
Gronbaek et al (2021)	Europe, 2014-2020	Prospective observational, 26 centers in 9 countries	Age < 18 y, posterior fossa tumors undergoing primary surgery (N=426), 55% male	Posterior fossa tumors	Post-operative speech impairment: 30% Post-operative mutism: 14% Post-operative reduced speech: 16%	Maximum FU 1 year
Huang et al (2020)	Sweden, 1973-2014	Population-based cohort study, linked Swedish Medical Birth Register and Swedish Cancer Registry	Singleton live births to parents w/CNS tumor (N=1,369), excluding those born within 1 y of parental diagnosis, and 5:1 matched controls (N=6,845)	Any CNS tumor	Adjusted OR for preterm birth: Reference: matched controls Ependymoma: 2.8 (1.2-6.5)	FU NR Controls matched on birth year, gender, maternal and paternal age at birth, and region of birth Adjusted OR:

Reference	Setting and study period	Design and data source	Study population (N)	Glioma definition	Complications and frequency Cul or OR	AEs: (%),	FU / comments
							adjusted for year of childbirth, gender, maternal and paternal age at birth, region at birth, parity, maternal birth country, maternal highest education, maternal pregnancy BMI, maternal smoking, gestational hypertensive disorder and gestational diabetes
Effinger et al (2019)	US and Canada, 1970-1986	Retrospective cohort, Childhood Cancer Survivor Study	Astrocytoma patients diagnosed before age 21 who survived > 5 y (N=1,182), 54% male	Astrocytoma	25-y Cul for at least one grade 3-5 chronic condition: 57% 25-y Cul for at least one grade 3-5 neurological condition: 26% Neurologic conditions include paralysis Cul 7%* 25-y Cul for at least one grade 3-5 visual condition: 19%, including legally blind in one or both eyes or loss of an eye Cul 3%* 25-y Cul for at least one grade 3-5 auditory		Median FU 23.4 y (range 7.3-38.9 y) *Cul paralysis: (83/1,182) * Cul for legally blind in one or both eyes or loss of an eye: (33/1,182) * Cul for hearing loss

Reference	Setting and study period	Design and data source	Study population (N)	Glioma definition	Complications and frequency Cul or OR	AEs: (%),	FU / comments
					condition: 17%, including hearing loss requiring a hearing aid Cul 8%*		requiring a hearing aid: (100/1,182)
					25-y Cul for stroke: 13%		
					25-y Cul for at least one grade 3-5 cardiac condition: 8%		
					25-y Cul for subsequent neoplasm: 7%		
					25-y Cul for at least one grade 3-5 endocrine condition: 6%, including ovarian failure Cul 3%*		* Cul for ovarian failure: (30/1,182)

2.1.4.1 Low-grade Glioma with BRAF V600E mutation

In LGGs, the frequency of BRAF V600E mutations ranges from 0% in ependymomas, pilomyxoid astrocytomas, and choroid plexus tumors to 2 to 9% in pilocytic astrocytomas, 13 to 55% in gangliogliomas and gangliocytomas, and 40 to 60% in pleomorphic xanthroastrocytomas ([Gierke et al 2016](#), [Koelsche et al 2014](#), [Horbinski et al 2012](#)).

Main existing treatment options

Regardless of the molecular profile, surgical removal is often the treatment of choice, if practical. The extent of resection is predictive of PFS. Only those patients with LGGs that can be completely resected can anticipate a median PFS of 10 years or more. Most patients will eventually experience progression of their disease and require post-surgical therapy with focal irradiation to the tumor bed plus additional chemotherapy.

For paediatric patients with molecularly unselected LGG, who could not be cured by surgical resection and were enrolled into studies of cytotoxic chemotherapy with carboplatin plus vincristine regimens, the ORR at 6 months was 29% (CR+PR), the 5-year PFS rate was 46% and 5-year OS was 89% ([Gnekow et al 2017](#)). In another large study, the ORR (CR+PR) by central review was 35% in paediatric patients with molecularly unselected LGG requiring postoperative systemic therapy with carboplatin and vincristine; 5-year OS was 86% ([Ater et al 2012](#)). In this setting of disease requiring systemic therapy after optimal surgical resection, the treatment goals generally are to prolong OS and PFS while minimizing morbidity of disease and treatment. Because of the typical young age of paediatric LGG patients and the potential

for long term neurocognitive effects of radiotherapy, this modality is often avoided where possible.

An analysis revealed that patients with paediatric LGG harboring the BRAF V600E mutation had worse PFS and OS ([Lassaletta et al 2017](#), [Ryall et al 2020](#)) than BRAF V600 wild type patients. In paediatric LGG patients with a BRAF V600E mutation, the 10-year OS was 89% and the 10-year PFS rate was 30% ([Ryall et al 2020](#)). In Lassaletta et al (2017), the 10-year PFS rate for BRAF V600E-mutant LGG was 27% (95% CI: 12.1, 41.9) compared to 60.2% (95% CI, 53.3% to 67.1%) for wild type. The Lassaletta work suggests a lower ORR of 11% (PR+CR) for these patients when treated with chemotherapy ([Lassaletta et al 2017](#)) versus 35% for the molecularly unselected population treated with chemotherapy ([Ater et al 2012](#)). There is evidence of poorer outcomes when deletion of CDKN2A is coupled with the BRAF V600E mutation ([Mistry et al 2015](#), [Lassaletta et al 2017](#), [Ryall et al 2020](#)). Patients with LGG who have progressed to secondary HGG (sHGG) are more likely to have had BRAF V600E mutation in their LGG at initial diagnosis ([Mistry et al 2015](#)), contributing to the poor prognosis upon initial diagnosis of BRAF V600 mutant paediatric LGG.

The survival outcomes have changed little over the past several decades, and thus improved treatment options are needed in the majority of patients with paediatric gliomas.

2.1.4.2 High-grade glioma with BRAF V600E mutation

Main existing treatment options

Current therapies for paediatric patients with HGGs are limited. Agents that have demonstrated activity in adult patients with HGG have not demonstrated similar benefit to paediatric patients with HGG ([Sturm et al 2017](#)). Current standard of care for newly diagnosed paediatric patients with HGGs include:

- Gross total surgical resection
- followed by focal irradiation to the tumor bed
- plus additional chemotherapy ([MacDonald et al 2011](#))

The majority of patients develop recurrent disease and in these cases there are no effective systemic treatment options. Chemotherapy regimens have been used, but they often have burdensome toxicity and provide limited benefit. Temozolomide is currently the only anticancer substance authorized specifically for HGG; it is most often used in the recurrent disease setting in adults but has proven to be of limited benefit for paediatric patients. In 5 trials evaluating temozolomide monotherapy or temozolomide-based combinations, the ORR in recurrent or refractory, paediatric molecularly unselected HGGs ranged from 0-12% ([Lashford et al 2002](#), [Ruggiero et al 2006](#), [Nicholson et al 2007](#), [Warren et al 2012](#), [Hummel et al 2013](#)). An OS of 4.7 months was estimated in [Lashford et al \(2002\)](#) and a 6-month PFS rate of 16% was estimated in [Warren et al \(2012\)](#). Treatment of relapsed, refractory paediatric molecularly unselected HGG with several other chemotherapies and/or targeted agents has shown a similar lack of benefit.

For paediatric HGG, the BRAF V600E mutation is more frequently found in favorable prognosis subgroups of this disease, such as those lacking H3K27 mutations, and is not found

in some of the worst prognostic subgroups, such as those arising from the brainstem ([Mackay et al 2017](#)). Thus, a paediatric patient diagnosed with a BRAF V600E mutation positive HGG may expect an improved OS versus those paediatric HGG that are wild type at BRAF V600. It is not known if this improvement in outcome would also be seen in those same patients at the time their disease has relapsed or become refractory to their initial treatment.

Overall, the treatment of children with HGG reflects a significant unmet need, with almost no improvement in survival outcomes in recent years.

2.2 Part II: Module SII: Non-clinical part of the safety specification

Key safety findings from non-clinical studies that are associated with combination study with dabrafenib are also described in the dabrafenib EU RMP.

Table 2-14 Key safety findings from non-clinical studies and relevance to human usage

Key Safety findings (from non-clinical studies)	Relevance to human usage
Skin effects In animals given trametinib, adverse effects on skin (consisted of scabs, acanthosis, erosion and ulcerations) were dose-limiting, but reversible toxicities. In dogs given trametinib+dabrafenib combination for up to 4 weeks, scabs were observed, consistent with animals given trametinib alone. Hyperproliferative skin effects were not observed.	Trametinib Monotherapy Skin-related toxicities were observed in over three-quarters of subjects in human clinical studies with trametinib; the most frequently reported were rash, dermatitis acneiform and erythema. Trametinib+Dabrafenib Combination Therapy Skin-related toxicities were observed in approximately one-half of subjects in the Phase III human clinical studies with trametinib+dabrafenib combination therapy; the most frequently reported were rash, dermatitis acneiform and erythema. While hyperproliferative skin effects were not observed preclinically for the combination of trametinib and dabrafenib, proliferative and hyperkeratotic skin lesions including squamous cell carcinoma (including keratoacanthoma), papilloma and hyperkeratosis, which are known adverse effects of BRAF-kinase inhibitors (including dabrafenib), were seen in human clinical studies with trametinib+dabrafenib combination therapy albeit at a frequencies lower than those observed with comparator arms with BRAF inhibitors.
Gastrointestinal effects In mice given trametinib monotherapy, perforation of the colon with secondary peritonitis (abdominal inflammation) and degeneration/necrosis of the glandular mucosa of the stomach were observed and were dose-limiting. In rats and dogs given trametinib,	Trametinib Monotherapy Gastrointestinal effects were observed in human clinical studies with trametinib; the most frequently reported were diarrhea, nausea and vomiting. Trametinib+Dabrafenib Combination Therapy

Key Safety findings (from non-clinical studies)	Relevance to human usage
erosions and ulcerations in the GI tract with local and systemic inflammatory responses were dose-limiting and reversible. In dogs given trametinib and dabrafenib in combination for up to 4 weeks, GI effects (minimal granulomatous inflammation of the stomach) were observed at a lower dose.	Gastrointestinal effects were observed in human clinical studies with trametinib+dabrafenib combination therapy; the most frequently reported were diarrhea, nausea and vomiting.
Cardiovascular effects Mice given trametinib monotherapy in excess of clinical exposures demonstrated lower heart rate, reduced heart weight, and lower left ventricular functional parameters, without microscopic effects in the myocardium. At non-tolerated doses of trametinib in rats, myocardial mineralization and necrosis were observed that were secondary to increased serum phosphorus causing metastatic mineralization in multiple tissues. In a 4-week toxicity study in dogs given trametinib and dabrafenib in combination, there were no cardiovascular findings.	Trametinib Monotherapy Subjects receiving trametinib monotherapy were monitored for cardiovascular effects with electrocardiograms and serial echocardiogram assessments for left ventricular ejection fraction (LVEF). Cardiovascular effects including decreases in ejection fraction (EF) and hypertension were observed in human clinical studies with trametinib. Trametinib+Dabrafenib Combination Therapy Subjects receiving trametinib+ dabrafenib combination therapy were monitored for cardiovascular effects with echocardiogram/electrocardiogram monitoring. Cardiovascular effects including decreases in EF and hypertension were seen in human clinical studies with dabrafenib + trametinib combination therapy.
Hepatic effects In mice and rats given trametinib monotherapy, hepatocellular vacuolation (rats only), increased transaminases (rats only) and necrosis (at non-tolerated doses) were observed. In dogs given non-tolerated doses of trametinib, mild increases in transaminases, bilirubin and cholesterol were observed, which were considered secondary to a systemic inflammatory response to GI toxicity. In a 4-Week Oral Capsule/Oral Gavage Investigational Pyrexia Study (Study No. 1970520) in Male Dogs, administration of dabrafenib and trametinib for up to 18 days resulted in earlier euthanization of one animal due to hepatic and GI toxicity	Trametinib Monotherapy Hepatic effects including increases in alanine aminotransferase (ALT) and AST were seen in human clinical studies with trametinib. Trametinib+Dabrafenib Combination Therapy Hepatic effects including increases in ALT and AST were seen in human clinical studies with trametinib+dabrafenib combination therapy.
Hematologic and bone marrow effects In animals given trametinib monotherapy, dose-dependent effects in the hematopoietic system were seen in mice (bone marrow degeneration/necrosis), rats (hematopoietic cell and lymphoid necrosis, bone marrow	Trametinib Monotherapy Haematologic effects including anaemia (9%), thrombocytopenia (2%), lymphopenia (2%), and neutropenia (2%) have been observed in human clinical studies with trametinib and are considered manageable and not clinically significant.

Key Safety findings (from non-clinical studies)	Relevance to human usage
hypocellularity, splenic necrosis) and dogs (reversible decreased reticulocytes, bone marrow hypocellularity, and lymphoid necrosis). In vitro, trametinib monotherapy also inhibited proliferation of cultured human granulocyte-macrophage colony forming cells and megakaryocyte colony forming cells in a concentration-dependent manner. In vitro, combination treatment of trametinib with dabrafenib inhibited proliferation of cultured human granulocyte-macrophage colony forming cells and megakaryocyte colony forming cells in a concentration-dependent manner at higher potency than with each as a single agent.	Trametinib+Dabrafenib Combination Therapy Hematologic effects of the combination therapy were observed in human clinical trials. In Study MEK115306 (Novartis trial ID: Study CDRB436B2301) haematologic effects including neutropenia (10%) anaemia (6%), thrombocytopenia (4%), and lymphopenia (2%) were observed with trametinib+dabrafenib combination therapy. Similar results were observed on combination therapy in Study MEK116513 (Novartis trial ID: Study CDRB436B2302). With the exception of neutropenia, the frequencies of these events are similar to those reported with trametinib monotherapy and did not require dose modifications. Therefore, these haematologic effects for the trametinib+dabrafenib combination therapy are considered manageable. Neutropenia (listed as an Adverse drug reaction (ADR)) is also considered manageable.
Bone effects Bone effects consisted primarily of physeal thickening in the femur in rats given trametinib doses ≥ 0.1 mg/kg/day). At higher non-tolerated trametinib doses of ≥ 1 mg/kg/day, subepiphyseal infarcts were observed. Physeal thickening was reversible. Reduced long bone length was observed in a juvenile rat study. No bone findings were observed in studies with dogs.	Trametinib Monotherapy and Trametinib+Dabrafenib Combination Therapy There have been no bone effects observed in adult patients taking trametinib. Bone effects in rats occur in actively growing bones and therefore would not pose a risk to adult human patients with closed physes. The effects on bone growth may be relevant for the paediatric population, and growth is monitored in clinical trials with paediatric patients during treatment.
Developmental toxicity and reproductive effects In rats and rabbits given trametinib monotherapy, maternal and developmental toxicity (decreased fetal body weights and increased ossification variations) were observed at exposures below exposures achieved at the recommended clinical dose of 2 mg per day. Additionally, decreased corpora lutea were observed in rats given trametinib, which may impact female fertility.	Trametinib Monotherapy There are no data in pregnant women. Trametinib+Dabrafenib Combination Therapy No pregnancies were reported in Study MEK115306. Two pregnancies have been reported in subjects receiving combination therapy with trametinib and dabrafenib in Study MEK116513; both subjects chose to have an elective abortion.
Ocular effects Trametinib Monotherapy: In mice, rats and dogs given trametinib monotherapy at or exceeding maximum tolerated doses (MTD) for up to 26 weeks, no ophthalmological or retinal microscopic findings were observed. In a juvenile	Trametinib Monotherapy Ocular effects, most commonly blurry vision, dry eyes and impaired vision were observed in human clinical studies with trametinib. Additionally, cases of retinal vein occlusion and retinal pigment epithelial detachment were reported in subjects receiving trametinib.

Key Safety findings (from non-clinical studies)	Relevance to human usage
rat study, findings in the eye including corneal mineralization and corneal dystrophy were observed in treated juvenile animals. The eye findings were still present after a 6-week recovery period.	Trametinib+Dabrafenib Combination Therapy Ocular effects, most commonly blurry vision and impaired vision were observed in the Phase III human clinical studies with trametinib+dabrafenib combination therapy. Additionally, cases of retinal pigment epithelial detachment were reported in subjects receiving trametinib+dabrafenib combination therapy.
Potential for Phototoxicity In an in vitro 3T3 NRU-PT assay, the cytotoxicity IC50 values in the presence and absence of UV-A were 2.92 µg/mL and 18.98 µg/mL, respectively, translating to a PIF of 6.5 indicating that trametinib was weakly phototoxic under the conditions of this assay.	Trametinib Monotherapy Photosensitivity reactions have been reported in 3% of subjects treated with trametinib. Trametinib+Dabrafenib Combination Therapy Photosensitivity reactions have been reported in 2% and 4% of subjects treated with trametinib+dabrafenib combination therapy in Study MEK115306 and Study MEK116513, respectively. Based on the clinical data available to date with few reports of photosensitivity reactions, there is low risk for phototoxicity with monotherapy or combination therapy. Thus, photosensitivity is not included as an identified or potential risk for either trametinib monotherapy or trametinib+dabrafenib combination therapy.
Paediatric effects The principal toxicities associated with trametinib administration to juvenile rats were observed on growth (body weight and long bone length) and in bone, phosphate homeostasis, eye, skin, liver, heart and female reproductive system effects, consisting of a delay in a physical landmark of sexual maturity and mammary gland development, lower corpora lutea and lower ovarian weights. With the exception of corneal mineralization/dystrophy and increased heart weight, similar effects have been observed in adult animals given trametinib.	Trametinib Monotherapy The safety of trametinib in children below 1 year of age has not been established. Trametinib+Dabrafenib Combination Therapy The safety of dabrafenib + trametinib combination therapy in children below 1 year of age has not been established.

2.3 Part II: Module SIII: Clinical trial exposure

Trametinib is an orally administered, allosteric inhibitor of MEK1 and MEK2 activation and kinase activity. Trametinib is approved for the treatment of melanoma and NSCLC and is under development for the treatment of additional malignancies with activation of the MAPK pathway.

BRAF V600 mutations in melanoma lead to a constitutive activation of the RAS/RAF/MEK/ERK (MAP-kinase) signal transduction pathway.

In combination, trametinib and dabrafenib inhibit two critical kinases and provide a more pronounced inhibition of the MAP-kinase pathway. The combination of trametinib with dabrafenib is synergistic in BRAF V600 mutation positive melanoma cell lines in vitro and delays the emergence of resistance to dabrafenib monotherapy in BRAF V600 mutation positive melanoma xenografts in vivo. The combination also synergistically inhibits phosphorylation of ERK and induces apoptosis in the MV522 NSCLC cell line that harbours the BRAF V600E mutation.

Trametinib Monotherapy

Advanced/Metastatic Melanoma

Trametinib was evaluated in three clinical trials listed below.

Study MEK111054 was an open label, multiple dose, Phase I, first time in human, multicenter study conducted in three parts to identify the MTD, and to determine the recommended Phase II dose (RP2D) and regimen for trametinib monotherapy. Part 1 consisted of dose escalation and identification of MTD. Part 2 was an expansion cohort in selected tumour types evaluating RP2D regimen. Part 3 of the study was to evaluate pharmacodynamic markers. Subjects with melanoma, pancreatic, non-small cell lung, colorectal cancer (CRC), lymphoma or any BRAF mutation-positive cancer were enrolled in the study. Among these, there were 81 subjects with melanoma who received at least one dose of trametinib. These subjects had not received a prior therapy with a BRAF small molecule inhibitor. This study was completed.

Study MEK113583 (Novartis trial ID: Study CTMT212A2201) was an open label, single arm, Phase II, multicenter, international study designed to evaluate the objective response rate in subjects with histologically and cytologically confirmed metastatic cutaneous BRAF V600E/K/D mutation-positive melanoma treated with trametinib at 2 mg once daily dose. A total of 97 subjects were enrolled into 2 cohorts. Cohort A – 40 subjects who had prior treatment with BRAF inhibitor and Cohort B – 57 subjects who had received a prior standard chemotherapy or immunotherapy but were naive to BRAF inhibitor therapy. This study was completed.

Study MEK114267 (Novartis trial ID: Study CTMT212A2301) is a randomised, two-arm, open label, multicenter, international Phase III study to evaluate the efficacy and safety of trametinib monotherapy (2 mg once daily) compared with chemotherapy (either DTIC or paclitaxel) in subjects with histologically confirmed cutaneous unresectable or metastatic melanoma (Stage IIIC or Stage IV), with centrally confirmed BRAF V600E/K mutation-positive melanoma. Prior to study enrolment, subjects received no more than one prior regimen of chemotherapy for advanced or metastatic melanoma. In addition, eligible subjects were stratified according to: a) lactate dehydrogenase (above upper limit of normal (ULN) versus equal to or below ULN) and b) prior chemotherapy for advanced or metastatic disease. PFS was the primary endpoint of the study and subjects were also followed for OS.

This study enrolled 310 subjects who received at least one dose of treatment and of these 211 were randomised to receive trametinib monotherapy and 99 to receive chemotherapy. Primary analysis of PFS was completed. This study is still ongoing to report mature OS data.

Integrated safety data from the above-mentioned trametinib monotherapy trials as well as safety data from Phase III Study MEK114267 provides comprehensive evaluation of the incidence of adverse events as well as trametinib monotherapy safety profile.

Trametinib+Dabrafenib Combination Therapy

Adjuvant treatment of patients with Stage III melanoma with a BRAF V600 mutation, following complete resection

PASS Study BRF115532: a randomized, double-blind study of dabrafenib in combination with trametinib versus 2 placebos as adjuvant treatment of high risk BRAF V600 E/K mutant melanoma after surgical resection. Subjects with completely resected, histologically confirmed, BRAF V600E/K mutation-positive, high-risk [Stage IIIA (lymph node metastasis >1 mm), IIIB or IIIC] cutaneous melanoma were screened for eligibility. Approximately 852 subjects were planned to be randomized at 1:1 ratio, stratified by BRAF mutation status (V600E, V600K) and stage of disease (Stage IIIA, IIIB, IIIC). A total of 870 subjects were finally randomized. The primary endpoint of the study is RFS; OS is defined as key secondary endpoint. Other secondary endpoints include DMFS and freedom from relapse (FFR). The primary analysis has been conducted with a cut-off date of 30-Jun-2017. The study is currently ongoing with OS follow up ongoing. The final OS analysis is planned for when approximately 299 deaths are reached.

Adult patients with advanced NSCLC with BRAF V600E mutation

Study BRF113928 (Novartis trial ID: Study CDRB436E2201) is a Phase II, open-label, non-randomized, sequentially enrolled, multi-center study. This study consists of three patient cohorts with metastatic NSCLC that had BRAF V600E mutation tested from a certified local laboratory:

- Cohort A (dabrafenib monotherapy)-dabrafenib 150 mg twice daily (bid) for patients at second-or later lines treatments;
- Cohort B (combination therapy for second-, third-, or fourth-line) -dabrafenib 150 mg bid and trametinib 2 mg once daily for patients at second-, third-, or fourth-line treatment;
- Cohort C (combination therapy for first-line) -dabrafenib 150 mg bid and trametinib 2 mg once daily for patients at first-line treatment for the metastatic disease.

The primary endpoint was the objective response rate by Investigator assessment, which was completed for Cohorts A and B. This study is ongoing to assess the Investigator-assessed overall response rate for Cohort C, long-term time-to-event data for Cohorts A-C, and long-term safety.

Advanced/Metastatic Melanoma

Trametinib+dabrafenib combination therapy was evaluated in three clinical trials listed below. The OS analyses from Study MEK115306 and Study MEK116513 provides data for the combination indication, supported by data from Study BRF113220 (Novartis trial ID: Study CDRB436B2201) Part C.

Study BRF113220 is a Phase I/II, open-label, dose escalation study to investigate the safety, pharmacokinetics, pharmacodynamics and clinical efficacy of dabrafenib + trametinib combination treatment. This study comprised 4 parts: designated as Parts A, B, C, and D. Parts

A, B, and D were Phase I evaluations to investigate if repeat doses of trametinib had an effect on the pharmacokinetics of single-dose dabrafenib (Part A); identify appropriate doses for combination-therapy using a dose-escalation procedure, and assess whether concomitant repeat dosing of dabrafenib and trametinib affected the pharmacokinetics of either investigational product (Part B); and evaluate the pharmacokinetics and safety of dabrafenib administered in hydroxypropyl methylcellulose (HPMC) capsules alone and in combination with trametinib (Part D).

Part C is a randomised open-label Phase II portion of the study in subjects with BRAF mutation-positive metastatic melanoma. Subjects were randomised to receive either one of 2 combination dosing regimens (dabrafenib 150 mg bid and trametinib 1 mg once daily or dabrafenib 150 mg bid and trametinib 2 mg once daily) or dabrafenib 150 mg bid as monotherapy.

Study MEK115306 is a Phase III, two-arm, double-blinded, randomised study comparing dabrafenib and trametinib as first-line combination therapy (150 mg bid dabrafenib + 2 mg once daily trametinib) to dabrafenib monotherapy (150 mg bid + trametinib placebo) in subjects with unresectable or metastatic BRAF V600E/K mutation-positive cutaneous melanoma. Primary analysis of PFS and final comparative analysis of OS have both completed. This study is ongoing to assess long-term OS and safety.

Study MEK116513 is a Phase III, two-arm, open label, randomised study comparing dabrafenib + trametinib combination therapy (150 mg bid dabrafenib + 2 mg once daily trametinib) to vemurafenib (960 mg bid) in subjects with unresectable or metastatic BRAF V600E/K mutation-positive cutaneous melanoma. The final comparative analysis of the primary endpoint of OS is complete. This study is ongoing to assess long-term OS and safety.

Adult patients with differentiated thyroid cancer with BRAF V600E mutation

Study CDRB436J12301 (hereafter referred to as Study J12301) is an ongoing Phase III, two-arm, double-blind, placebo-controlled, randomized study evaluating the efficacy and safety of dabrafenib + trametinib (150 mg bid dabrafenib + 2 mg once daily trametinib) versus placebo in previously treated adult patients with locally advanced or metastatic, radio-active iodine refractory BRAF V600E mutation-positive DTC. The primary endpoint is PFS by RECIST 1.1 as assessed by a Blinded Independent Review Committee. Secondary endpoints include overall response rate, OS, duration of response, and safety. Patients in the placebo arm with confirmed disease progression may cross over to open-label dabrafenib plus trametinib. The study enrolled a total of 153 patients (101 patients were randomized to dabrafenib + trametinib and 52 patients were randomized to placebo).

Paediatric gliomas

CDRB436G2201 study: Phase II open-label global study to evaluate the effect of dabrafenib in combination with trametinib in children and adolescent patients with BRAF V600 mutation positive LGG or relapsed or refractory HGG. Paediatric patients (≥ 12 months and < 18 years of age) with BRAF V600 mutation-positive LGG or relapsed or refractory HGG were enrolled. LGG cohort is a randomized, open label part in children and adolescent patients with BRAF V600 mutation. HGG cohort is a single arm, open label part in which children and adolescent patients with BRAF V600 mutation- positive, refractory or relapsed HGG tumors received

dabrafenib+trametinib. A total of 151 patients were enrolled in the study; 110 in the LGG cohort and 41 in the HGG cohort. Of the 110 patients in LGG cohort 73 patients were randomized to dabrafenib+trametinib combination therapy arm and 37 patients were randomized to chemotherapy arm (carboplatin+vincristine). Four patients in chemotherapy arm discontinued prior to first dose and were not treated in this study. A total of 9 patients initially randomized to the chemotherapy arm and treated with chemotherapy later crossed-over to receive dabrafenib+trametinib combination therapy. The HGG part of the study was single-arm and enrolled 41 patients who received targeted therapy (D+T). A total of 123 patients (LGG: 73, LGG crossover: 9, and HGG: 41) received dabrafenib+trametinib combination therapy.

CTMT212X2101 study: This study was a dose escalation Phase I/IIa, multi center, open-label clinical study to investigate the safety, pharmacokinetics, pharmacodynamics and clinical activity of trametinib and dabrafenib + trametinib in paediatric patients with refractory or recurrent solid tumors with presumed MAPK pathway activation. This study had 4 parts:

Part A (≥ 1 month and <18 years) was a trametinib monotherapy dose escalation phase (0.0125, 0.025, 0.032, and 0.04 mg/kg/day).

Part B (≥ 1 month and <18 years) was a monotherapy disease cohort expansion phase (ages <6 years: 0.032 mg/kg/d; ages ≥ 6 years: 0.025 mg/kg/d).

Part C (≥ 12 months and <18 years) was a limited dose escalation phase of dabrafenib + trametinib in BRAF V600 mutant tumors (trametinib 0.025 mg/kg/day + 50% dabrafenib RP2D; trametinib 0.025 mg/kg/day + 100% dabrafenib RP2D; trametinib 0.032 mg/kg/day + 100% dabrafenib RP2D).

Part D (≥ 12 months and <18 years) was a cohort expansion phase of D+T in children and adolescents with BRAF V600 mutated tumors (LGG and LCH) (trametinib 0.032 mg/kg/day + 100% dabrafenib RP2D for patients < 6 years old and trametinib 0.025 mg/kg/day + 100% dabrafenib RP2D for patients ≥ 6 years old).

A total of 139 paediatric patients were enrolled into this study: 91 patients in trametinib monotherapy arms (A and B) and 48 patients in dabrafenib + trametinib combination therapy arms (C and D: LGG patients: 20 and LCH patients: 10). This study is completed.

Clinical trial exposure

Trametinib Monotherapy

There were no randomised, blinded trials conducted with trametinib.

Table 2-15 Duration of Exposure – Integrated Safety Population

Duration	Trametinib N=329	Subject time (months)
Less than 1 month	19	12.0
At least 1 month	310	1744.3
At least 3 months	197	1518.0
At least 6 months	107	1123.3
At least 9 months	63	804.5

Duration	Trametinib N=329	Subject time (months)
At least 12 months	30	459.7
At least 15 months	15	264.0
At least 18 months	7	140.6
At least 21 months	1	24.5
At least 24 months	1	24.5
Subject-time (months)	329	1756.3

Subject-time is the sum of each subject's treatment exposure in month.

Source: Table 8.76 MEK_120 Update; data as of 25-Jun-2012

Table 2-16 Exposure by age group and gender – Integrated Safety Population

Age	Sex	Trametinib N=329 Subjects n (%)	Subject-time (months)
Total	Total	329 (100)	1756.3
	Male	198 (60.2)	974.9
	Female	131 (39.8)	781.4
<65	Total	249 (75.7)	1383.5
65-74	Total	64 (19.4)	313.1
75-84	Total	15 (4.6)	57.0
At least 75	Total	16 (4.9)	59.7
At least 85	Total	1 (0.3)	2.7

Subject-time is the sum of each subject's treatment exposure in months. Subject-time is based on the number of subjects in each category.

Source: Table 8.78 MEK_120 Update; data as of 25-Jun-2012; Table 8.77 MEK_120 Update; data as of 25-Jun-2012

Trametinib + Dabrafenib Combination Therapy

Treatment of adult patients with unresectable or metastatic melanoma with a BRAF V600 mutation (Study MEK115306 and Study MEK116513)

Table 2-17 Duration of exposure – Study MEK115306

Duration	Dabrafenib N=420 n (%)	Trametinib N=209 n (%)	Trametinib dabrafenib N=209 n (%)	+
Less than 1 month	11 (2.6)	4 (1.9)	4 (1.9)	
At least 1 month	409 (97.4)	205 (98.1)	205 (98.1)	
At least 3 months	371 (88.3)	193 (92.3)	193 (92.3)	
At least 6 months	287 (68.3)	156 (74.6)	156 (74.6)	
At least 9 months	226 (53.8)	124 (59.3)	124 (59.3)	
At least 12 months	183 (43.6)	103 (49.3)	103 (49.3)	
At least 15 months	156 (37.1)	91 (43.5)	91 (43.5)	

Duration	Dabrafenib N=420 n (%)	Trametinib N=209 n (%)	Trametinib dabrafenib N=209 n (%)	+
At least 18 months	137 (32.6)	84 (40.2)	84 (40.2)	
At least 21 months	123 (29.3)	78 (37.3)	78 (37.3)	
At least 24 months	111 (26.4)	73 (34.9)	73 (34.9)	
At least 27 months	47 (11.2)	29 (13.9)	29 (13.9)	
At least 30 months	8 (1.9)	4 (1.9)	4 (1.9)	
Subject-time (months)	5332	3012	3012	

Source: RMP version 13.1 Attachment to Annex 12 MEK115306 Table 3.702

Table 2-18 Duration of exposure – Study MEK116513

Duration	Dabrafenib N=350 n (%)	Trametinib N=350 n (%)	Trametinib dabrafenib N=350 n (%)	+	Vemurafenib N=349 n (%)
Less than 1 month	7 (2.0)	6 (1.7)	7 (2.0)		27 (7.7)
At least 1 month	343 (98.0)	344 (98.3)	343 (98.0)		322 (92.3)
At least 3 months	318 (90.9)	317 (90.6)	317 (90.6)		268 (76.8)
At least 6 months	245 (70.0)	247 (70.6)	248 (70.9)		184 (52.7)
At least 9 months	199 (56.9)	199 (56.9)	200 (57.1)		132 (37.8)
At least 12 months	120 (34.3)	118 (33.7)	120 (34.3)		68 (19.5)
At least 15 months	48 (13.7)	46 (13.1)	48 (13.7)		21 (6.0)
At least 18 months	13 (3.7)	13 (3.7)	13 (3.7)		1 (0.3)
At least 21 months	1 (0.3)	1 (0.3)	1 (0.3)		0
At least 24 months	0	0	0		0
Subject-time (months)	3183	3171	3186		2368

Source: RMP version 13.1 Attachment to Annex 12 MEK116513-Table 3.702

Table 2-19 Exposure by age group and gender – Study MEK115306

Age	Sex	Dabrafenib N=420		Trametinib N=209		Trametinib dabrafenib N=209		+
		Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	
Total	Total	420 (100.0)	5332	209 (100.0)	3012	209 (100.0)	3012	
	Male	225 (53.6)	2653	111 (53.1)	1435	111 (53.9)	1435	
	Female	195 (46.4)	2679	98 (46.9)	1577	98 (46.9)	1577	
<65	Total	304 (72.4)	3916	153 (73.2)	2249	153 (73.2)	2249	
65-74	Total	88 (21.0)	1205	45 (21.5)	701	45 (21.5)	702	
75-84	Total	26 (6.2)	204	10 (4.8)	59	10 (4.8)	58	
At least 75	Total	28 (6.7)	211	11 (5.3)	62	11 (5.3)	61	

		Dabrafenib N=420		Trametinib N=209		Trametinib dabrafenib N=209		+	
Age	Sex	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)		
At least 85	Total	2 (0.5)	7	1 (0.5)	3	1 (0.5)	3		

Source: RMP version 13.1 Attachment to Annex 12 MEK115306-Table 3.703 and Annex 12 MEK115306-Table 3.704

Table 2-20 Exposure by age group and gender – Study MEK116513

Age	Sex	Dabrafenib N=350		Trametinib N=350		Trametinib + dabrafenib N=350		Vemurafenib N=349	
		Subjects n (%)	Subject-time (months)	Subjects n (%)	Subject-time (months)	Subjects n (%)	Subject-time (months)	Subjects n (%)	Subject-time (months)
Total	Total	350 (100.0)	3183	350 (100.0)	3171	350 (100.0)	3186	349 (100.0)	2368
	Male	207 (59.1)	1821	207 (59.1)	1795	207 (59.1)	1816	179 (51.3)	1215
	Female	143 (40.9)	1362	143 (40.9)	1376	143 (40.9)	1370	170 (48.7)	1153
<65	Total	273 (78.0)	2511	273 (78.0)	2520	273 (78.0)	2524	262 (75.1)	1755
65-74	Total	56 (16.0)	511	56 (16.0)	486	56 (16.0)	497	61 (17.5)	485
75-84	Total	18 (5.1)	159	18 (5.1)	163	18 (5.1)	163	25 (7.2)	128
At least 75	Total	21 (6.0)	161	21 (6.0)	165	21 (6.0)	165	26 (7.4)	128
At least 85	Total	3 (0.9)	2	3 (0.9)	2	3 (0.9)	2	1 (0.3)	0

Source: RMP version 13.1 Attachment to Annex 12 MEK116513 Table 3.703 and Annex 12 MEK116513 Table 3.704

Adjuvant treatment of patients with Stage III melanoma with a BRAF V600 mutation, following complete resection (Study BRF115532)

Table 2-21 Duration of exposure – Study BRF115532

Duration	Dabrafenib N=435 n(%)	Trametinib N=435 n(%)	Trametinib + dabrafenib N=435 n(%)
Less than 1 month	25 (6.0%)	25 (6%)	26 (6%)
At least 1 month	410 (94.0%)	410 (94%)	409 (94%)
At least 3 months	363 (83.0%)	363 (83%)	360 (83%)
At least 6 months	319 (73.0%)	324 (74%)	316 (73%)
At least 9 months	296 (68.0%)	299 (69%)	293 (67%)
Subject-time (months)	3567	3601	3540

Note: For 'Dabrafenib + Trametinib', when deriving person time, duration of exposure for each subject is calculated as: Min(Last Dose of Dabrafenib, Last Dose of Trametinib) – Max(First Dose of Dabrafenib, First Dose of Trametinib)+1

Source: RMP version 14 Attachment to Annex 7-Table 14.3-1.3

Table 2-22 Exposure by age group and gender – Study BRF115532

		Dabrafenib N=435		Trametinib N=435		Trametinib + dabrafenib N=435	
Age	Sex	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)
Total	Total	435 (100.0)		435 (100.0)		435 (100.0)	
	Male	242 (56.0)	2054	242 (56.0)	2059	242 (56.0)	2047
	Female	193 (42.0)	1513	193 (42.0)	1542	193 (42.0)	1493
<65	Total	350		350		350	
	Male	191 (44.0)	1663	191 (44.0)	1676	191 (44.0)	1666
	Female	159 (37.0)	1267	159 (37.0)	1299	159 (37.0)	1250
65-74	Total	73		73		73	
	Male	46 (11.0)	343	46 (11.0)	336	46 (11.0)	333
	Female	27 (6.0)	201	27 (6.0)	200	27 (6.0)	200
75-84	Total	11		11		11	
	Male	4 (<1.0)	37	4 (<1.0)	36	4 (<1.0)	37
	Female	7 (2.0)	45	7 (2.0)	43	7 (2.0)	43
At least 75	Total	12		12		12	
	Male	5 (1.0)	48	5 (1.0)	47	5 (1.0)	48
	Female	7 (2.0)	45	7 (2.0)	43	7 (2.0)	43
At least 85	Total	1		1		1	
	Male	1 (<1)	11	1 (<1)	11	1 (<1)	11

Age	Sex	Dabrafenib N=435		Trametinib N=435		Trametinib dabrafenib N=435		+
		Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	
	Female	0	0	0	0	0	0	

Subject-time is the sum of each subject's treatment exposure in <unit>. <Subject-time> is based on the number of subjects in each category.

Note: For 'Dabrafenib + Trametinib', when deriving person time, duration of exposure for each subject is calculated as: Min(Last Dose of Dabrafenib, Last Dose of Trametinib) – Max(First Dose of Dabrafenib, First Dose of Trametinib)+1.

Source: RMP version 14 Attachment to Annex 7-Table 14.3-1.4

Treatment of adult patients with advanced non-small cell lung cancer (NSCLC) with a BRAF V600 mutation (Study BRF113928)

Table 2-23 Duration of exposure – Study BRF113928

Duration	Dabrafenib N=166 n (%)	Trametinib N=82 n (%)	Trametinib dabrafenib N=82 n (%)	+
Less than 1 month	16 (9.6)	8 (9.8)	9 (11.0)	
At least 1 month	150 (90.4)	74 (90.2)	73 (89.0)	
At least 3 months	108 (65.1)	57 (69.5)	56 (68.3)	
At least 6 months	74 (44.6)	38 (46.6)	39 (47.6)	
At least 9 months	58 (34.9)	32 (39.0)	32 (39.0)	
At least 12 months	38 (22.9)	19 (23.2)	16 (19.5)	
At least 15 months	28 (16.9)	12 (14.6)	12 (14.6)	
At least 18 months	21 (12.7)	6 (7.3)	4 (4.9)	
At least 21 months	14 (8.4)	1 (1.2)	0	
At least 24 months	8 (4.8)	0	0	
Subject-time (months)	1344	620	607	

Source: RMP version 13.1 Attachment to Annex 12 Table 3.0901 and Annex 12 Table 3.0902

Table 2-24 Exposure by age group and gender – Study BRF113928

Age	Sex	Dabrafenib N=166		Trametinib N=82		Trametinib dabrafenib N=82		+
		Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	
Total	Total	166 (100.0)	1344	82 (100.0)	620	82 (100.0)	607	
	Male	79 (47.6)	634	39 (47.6)	311	39 (47.6)	304	
	Female	87 (52.4)	710	43 (52.4)	309	43 (52.4)	303	
<65	Total	71 (42.8)	586	35 (42.7)	302	35 (42.7)	297	
65-74	Total	66 (39.8)	527	29 (35.4)	184	29 (35.4)	180	

		Dabrafenib N=166		Trametinib N=82		Trametinib dabrafenib N=82		+
Age	Sex	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	Subjects n (%)	Subject- time (months)	
75-84	Total	22 (13.3)	178	13 (15.9)	96	13 (15.9)	92	
At least 75	Total	29 (17.5)	231	18 (22.0)	134	18 (22.0)	130	
At least 85	Total	7 (4.2)	54	5 (6.1)	39	5 (6.1)	38	

Source: RMP version 13.1 Attachment to Annex 12 Table 3.0901, Annex 12 Table 3.0903, and Annex 12 Table 3.0904

Treatment of adult patients with DTC with BRAF V600 mutation (Study J12301)

Table 2-25 Duration of exposure - Study J12301 (primary analysis)

Duration of exposure	Dabrafenib + trametinib (N=101)		Placebo (N=52)	
	Dabrafenib n (%)	Trametinib n (%)	Placebo Dabrafenib n (%)	Placebo Trametinib n (%)
<1 month	1 (1.0)	2 (2.0)	7 (13.5)	7 (13.5)
At least 1 month	100 (99.0)	99 (98.0)	45 (86.5)	45 (86.5)
At least 3 months	93 (92.1)	90 (89.1)	37 (71.2)	37 (71.2)
At least 6 months	83 (82.2)	81 (80.2)	26 (50.0)	26 (50.0)
At least 9 months	76 (75.2)	72 (71.3)	22 (42.3)	22 (42.3)
At least 12 months	57 (56.4)	54 (53.5)	14 (26.9)	14 (26.9)
At least 15 months	47 (46.5)	43 (42.6)	8 (15.4)	8 (15.4)
At least 18 months	42 (41.6)	40 (39.6)	6 (11.5)	6 (11.5)
At least 21 months	29 (28.7)	27 (26.7)	2 (3.8)	2 (3.8)
At least 24 months	25 (24.8)	23 (22.8)	2 (3.8)	2 (3.8)
Subject time (months)	1619.0	1539.3	428.5	428.2

Source: [SCS Appendix 1-Table 1-5 & Table 1-6]

Table 2-26 Exposure by age group and gender - Study J12301 (primary analysis)

		Dabrafenib + trametinib (N=101)		Placebo (N=52)	
Age	Sex	Subjects n (%)	Subject time (months)	Subjects n (%)	Subject time (months)
<65	Male	32 (31.7)	504.6	13 (25.0)	110.8
65 to 74	Male	17 (16.8)	229.0	5 (9.6)	55.5
75 to 84	Male	5 (5.0)	75.5	1 (1.9)	9.5
75+	Male	5 (5.0)	75.5	1 (1.9)	9.5
85+	Male	0	0	0	0
<65	Female	18 (17.8)	348.1	18 (34.6)	123.4
65 to 74	Female	24 (23.8)	315.1	14 (26.9)	112.9
75 to 84	Female	5 (5.0)	67.0	1 (1.9)	16.2

		Dabrafenib + trametinib (N=101)		Placebo (N=52)	
Age	Sex	Subjects n (%)	Subject time (months)	Subjects n (%)	Subject time (months)
75+	Female	5 (5.0)	67.0	1 (1.9)	16.2
85+	Female	0	0	0	0

Source: [SCS Appendix 1-Table 1-4](#)

Exposure in paediatric combination therapy safety pool

Table 2-27 Duration of exposure to trametinib in combination therapy in paediatric patients

	All N=171 n (%)	Subjects
Total number of subjects receiving Combination-n (%)	171 (100.0)	
Duration of exposure categories-n (%)		
< 3 Weeks	3 (1.8)	
3 - < 6 Weeks	0	
6 - < 12 Weeks	7 (4.1)	
12 - < 24 Weeks	12 (7.0)	
24 - < 48 Weeks	25 (14.6)	
> 48 Weeks	124 (72.5)	
Subject Time (Week)	13697.7	
Subject Time (PY)*	262.51	

Subject-time is the sum of each subject's treatment exposure in Weeks.

*subjects Time in PY is calculated by time in weeks * 7/365.25

Source: [\[SCS Appendix 1-Table 2.1-1\]](#)

2.4 Part II: Module SIV: Populations not studied in clinical trials

2.4.1 Part II Module SIV.1. Exclusion criteria in pivotal clinical studies within the development program

Table 2-28 Important exclusion criteria in pivotal studies in the development program

Criteria	Reason exclusion	for	Is it considered to be included as missing information?	Rationale
Melanomas that have BRAF mutations that are non- V600 or do not have a BRAF mutation	The efficacy and safety of trametinib as monotherapy or in combination with dabrafenib have not been established in patients with wild-type BRAF melanoma.	No	No	Trametinib is an agent targeting the MAPK pathway; the efficacy of this agent, as monotherapy or in combination with dabrafenib was assessed in patients with BRAFV600 mutated melanoma (the population with wild-type BRAF melanoma was excluded).
NSCLC that have BRAF mutations that are non- V600 or do not have a BRAF mutation	The efficacy and safety of trametinib in combination with dabrafenib was not established in	No	No	Trametinib is an agent targeting the MAPK pathway; the efficacy of this agent, as monotherapy or in combination with dabrafenib

Criteria	Reason for exclusion	Is it considered to be included as missing information?	Rationale
	patients with wild-type BRAF NSCLC.		was assessed in patients with BRAFV600 mutated melanoma (the population with wild-type BRAF melanoma was excluded).
Age of <1 year old	Patients below 1 year of age were excluded from trametinib clinical trials.	No	There is no data to support a contraindication in this population. This is discussed under Posology and Administration section of labelling.
History of retinal vein occlusion (RVO) or central serous retinopathy (CSR)/Retinal pigment epithelial detachment (RPED), or predisposing factors to RVO or CSR/RPED (e.g. uncontrolled glaucoma or ocular hypertension, uncontrolled systemic disease such as hypertension, diabetes mellitus, or history of hyperviscosity or hypercoagulability syndromes). Visible retinal pathology as assessed by ophthalmic exam that is considered a risk factor for RVO or CSR/RPED such as: Evidence of new optic disc cupping Evidence of new visual field defects Intraocular pressure >21 mmHg as measured by tonography	Trametinib is associated with ocular toxicities which appear to be class effects, including CSR/RPED and RVO. Symptoms such as blurred vision, decreased acuity, and other visual phenomena have been reported in the clinical trials with trametinib as monotherapy and in combination with dabrafenib. History of RVO or CSR/RPED or risk factors may identify populations who may be at risk to develop these specific ocular adverse events.	No	RVO and RPED are class effects and labeled event, Patients with history of RVO and RPED are excluded from clinical studies. In the post-marketing setting, no significant data was retrieved.
Subjects treated with trametinib with a history of pneumonitis or interstitial lung disease (ILD).	Events of pneumonitis and ILD occurring prior to trametinib treatment have been considered as a potential risk factor	No	Pneumonitis or ILD are labeled events. In the post-marketing setting, no significant data was retrieved.

Criteria	Reason for exclusion	Is it considered to be included as missing information?	Rationale
	for recurrence of these specific adverse events during treatment with trametinib as monotherapy or in combination with dabrafenib.		
History or evidence of cardiovascular risk including any of the following: History of acute coronary syndromes, coronary angioplasty, or stenting Class II, III, or IV heart failure LVEF <lower limit of normal (LLN) Treatment refractory hypertension (blood pressure of systolic >140 mmHg and/or diastolic >90 mmHg which cannot be controlled by anti-hypertensive therapy History or evidence of current clinically significant uncontrolled arrhythmias. Exception: Subjects with controlled atrial fibrillation for >1 month prior to randomization are eligible.	Left ventricular dysfunction was reported with trametinib as well as with other MEK inhibitors in clinical development. In clinical trials, the average time to onset of left ventricular dysfunction and LVEF decreased was between 2 to 3 months.	No	Trametinib has been reported to decrease LVEF, when used as monotherapy or in combination with dabrafenib. Therefore, patients with left ventricular dysfunction, New York Heart Association Class II, III, or IV heart failure, acute coronary syndrome within the past 6 months, clinically significant uncontrolled arrhythmias, and uncontrolled hypertension were excluded from clinical trials; safety of use in this population is therefore unknown.

2.4.2 Part II Module SIV.2. Limitations to detect adverse reactions in clinical trial development programs

The clinical development program is unlikely to detect certain types of adverse reactions such as rare adverse reactions, adverse reactions with a long latency, specific to other than Caucasian population or those caused by prolonged or cumulative exposure.

2.4.3 Part II Module SIV.3. Limitations in respect to populations typically underrepresented in clinical trial development programs

Table 2-29 SIV.2: Exposure of special populations included or not in clinical trial development programs

Type of special population	Exposure
Pregnant women	Not included in the clinical development program
Breastfeeding women	Not included in the clinical development program
Patients with relevant comorbidities:	
• Patients with hepatic impairment	Not included in the clinical development program
• Patients with renal impairment	Not included in the clinical development program
• Patients with cardiovascular impairment were not included in the clinical development program	Not included in the clinical development program
• Immunocompromised patients were not included in the clinical development program	Not included in the clinical development program
• Patients with a disease severity different from inclusion criteria in clinical trials	Not included in the clinical development program
Population with relevant different ethnic origin	Data on patients of different racial and/or ethnic origins is limited.
Subpopulations carrying relevant genetic polymorphisms	Not included in the clinical development program
Other:	
Paediatric patients (<18 years old)	Paediatric patients 1 year to <18 years of age have been included in the clinical development program
Elderly (≥ 65 years old)	Included in the clinical development program (Section 2.3).

2.5 Part II: Module SV: Post-authorization experience

2.5.1 Part II Module SV.1. Post-authorization exposure

2.5.1.1 Part II Module SV.1.1 Method used to calculate exposure

Trametinib as monotherapy and in combination therapy

An estimate of patient exposure is calculated based on worldwide sales volume in kilogram (kg) of active substance sold and the Defined Daily Dose (DDD) i.e., 2 mg trametinib.

2.5.1.2 Part II Module SV.1.2. Exposure

The estimated interval exposure during reporting interval for Periodic safety update report (PSUR 30-May-2024 to 29-May-2025) was approximately 26,901 patient treatment years (PTY). The cumulative exposure estimate until 29-May-2025 is 148,668 PTY. [REDACTED]

2.6 Part II: Module SVI: Additional EU requirements for the safety specification

2.6.1 Potential for misuse for illegal purposes

Based on the mechanism of action of trametinib, the potential for misuse for illegal purposes, abuse or dependence has not been identified and is considered unlikely from the knowledge of the compound to date.

2.7 Part II: Module SVII: Identified and potential risks

2.7.1 Part II Module SVII.1. Identification of safety concerns in the initial RMP submission

2.7.1.1 Part II Module SVII.1.1. Risks not considered important for inclusion in the list of safety concerns in the RMP

This section is not applicable; the RMP was already approved.

2.7.1.2 Part II Module SVII.1.2. Risks considered important for inclusion in the list of safety concerns in the RMP

There are no additional risks considered important for inclusion in the list of safety concerns, since the RMP is already approved.

2.7.2 Part II Module SVII.2: New safety concerns and reclassification with a submission of an updated RMP

There were no changes to the list of safety concerns since the last approved RMP version 20.0.

2.7.3 Part II Module SVII.3: Details of important identified risks, important potential risks, and missing information

2.7.3.1 Part II Module SVII.3.1. Presentation of important identified risks and important potential risks

2.7.3.1.1 Important Identified Risk: ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)

Trametinib monotherapy

Table 2-30 Clinical trial data of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)

Frequency with 95% CI		Trametinib (N=211)	Chemotherapy (N=99)
Study MEK114267	Number of Subjects with Event	21 (10%)	4 (4%)
	Number of Events	31	4
	Event Characteristics		
	N	21	4
	Serious	2 (10%)	0
	Drug-related	17 (81%)	3 (75%)
	Leading to Withdrawal	0	0
	Severe	0	0
	Fatal	0	0
	Maximum Grade		
	N	21	4
	Grade 1	15 (71%)	3 (75%)
	Grade 2	3 (14%)	1 (25%)
	Grade 3	2 (10%)	0
	Grade 4	1 (5%)	0
	Grade 5	0	0

NA= In Study MEK114267, information on whether or not an event was reported as a 'Single episode' or as 'Intermittent' was not captured.

Any Ocular Event Relative Risk 95% CI: 2.46 (0.86, 6.98), Risk difference 95% CI: 6% (0%, 12%).

Most of the events noted above were grade 1 including non-specific terms such as blurry vision. As of 26-Sep-2012, there were 14 cases of chorioretinopathy or retinal pigment epithelial detachment (RPED) reported (including 4 from integrated studies). All 14 RPED events resolved. As of 26-Sep-2012, a total of 4 cases of RVO reported across the program (including 2 cases from Study MEK114267). All 4 cases of RVO occurred in one eye only. Study drug was stopped at time of diagnosis in all cases. There was a decrease of visual acuity in the 2 subjects with central retinal vein occlusion (CRVO) while the other two subjects experienced no meaningful decrease of visual acuity (one line—from 20/20 to 20/25 in both subjects). In the two subjects experiencing CRVO, local treatment with intravitreal injections of anti-VEGF antibodies was initiated within 2 weeks after RVO diagnosis and visual acuity was improved in one subject or restored to baseline conditions in other subject, at the time of the data cut-off date.

An post-marketing requirement report, PMR report 2014 provided the following information:

Including the event after crossover, a total of four subjects in the 150/2 combination and trametinib monotherapy treatment arms of the studies included in this report had RPED events.

No new risk factors were identified during the clinical review of subjects with RPED events.

Based on this review, there are no recommended changes to the current trametinib monitoring recommendation and dose modification guidelines for RPED.

In the Study MEK114653, visual disorders were reported in five subjects (6%) treated with trametinib monotherapy and all were grade 1-2. The preferred term included visual impairment, diplopia, dry eye and vision blurred (CSR Study MEK114653 Table 8.1401).

Table 2-31 Characterization of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)

Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Details				
Potential mechanisms	Potential mechanism is unknown.				
Evidence source(s) and strength of evidence	Blurred vision, decreased acuity, and other visual phenomena have been reported in the clinical trials with trametinib. In clinical trials, uveitis and iridocyclitis have also been reported in patients treated with trametinib in combination with dabrafenib.				
Characterization of the risk:	Ocular events were reported in three trametinib studies: MEK111054, MEK113583 and MEK114267. Trametinib is not recommended in patients with a history of RVO. The safety of trametinib in subjects with predisposing factors for RVO, including uncontrolled glaucoma or ocular hypertension, uncontrolled hypertension, uncontrolled diabetes mellitus, or a history of hyperviscosity or hypercoagulability syndromes, has not been established. In the SmPC prompt ophthalmological assessment is recommended for patients with new visual disturbances, such as diminished central vision, blurred vision or loss of vision at any time while on trametinib therapy. No studies (excluding case reports) of unresectable or metastatic melanoma patients were found reporting the incidence of RPED. SEER-Medicare Study for RPED: Among 1746 unresectable or metastatic melanoma patients who had no history of RPED during 1-year prior to unresectable or metastatic melanoma diagnosis, no patient had RPED post-cancer diagnosis. A similar analysis was conducted, expanding to all patients with melanoma (n=29948), as RPED is a rare event. Among melanoma patients who had no history of RPED during 1-year prior to melanoma diagnosis, very few had incident RPED during 6 months post diagnosis. No control developed incident RPED during the 6-month period post the index date.				
		Total N at risk	Event (n)	Cumulative Incidence = n/total N*100%	Incidence rate, per 1000 PTY (95% CI) = n/total PTY
	Melanoma Unresectable or metastatic melanoma	29948 1746	<11* 0	<0.6% 0	-- 0

Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Details
	*The number of events was less than 11. According to the privacy guidelines outlined by SEER-Medicare, no cell may be displayed and no use of percentages or other mathematical formulas may be used if they result in the display of a cell 10 or less. The incidence of branch RVO and central RVO among the general population have been reported in several population-based studies. The Beaver Dam Eye Study (n=4068) in Wisconsin reported 15-year cumulative incidences of 1.8% for branch RVO and 0.5% for central RVO (Klein et al 2008). The 10-year cumulative incidences for RVO was reported at 1.6%, branch RVO at 1.2% and central RVO at 0.4% in the Blue Mountain Eye Study in Australia (n=3654) (Cugati et al 2006). Rogers et al (2010) summarized the prevalence of RVO from population-based studies in the US, Europe, Asia, and Australia. The combined pooled data contained 68751 individuals across 15 studies, with participants' ages ranging from 30 to 101 years. The age and sex standardized prevalence was 4.40 per 1000 for any retinal vein occlusion, 3.77 per 1000 for branch RVO, and 0.65 per 1000 for central RVO. When confined to the 11 studies that assessed 2 or more fields of both eyes, the corresponding prevalence's were 4.42 per 1000 for branch RVO and 0.8 per 1000 for central RVO. Across individual studies, the standardized prevalence varied from 0.26 to 9.32 per 1000 for branch RVO and 0.04 to 1.59 per 1000 for central RVO.
Risk factors and risk groups	None identified yet for trametinib.
Preventability	Treatment with trametinib should not be recommended for patients with a history or current evidence/risk of RVO. Patients on or about to start trametinib therapy with any changes in visual acuity should have a full ophthalmologic evaluation. Information and management guidelines are included in the product labelling.
Impact on the benefit-risk balance of the product	The benefit-risk balance of trametinib is not impacted by this risk considering the nature of the treatment indication(s) and if any the impact it is low as the SmPC provides adequate information on timely identification and management of this risk. Furthermore, data in the recent PSUR for trametinib monotherapy also confirm no impact on the benefit–risk of trametinib+dabrafenib combination treatment with regards to this risk.
Public health impact	RPED has been reversible with drug interruption and subjects were able to restart trametinib at reduced dose. All 4 cases of RVO, reported across the program, occurred in one eye only. Study drug was stopped at time of diagnosis in all cases. Of these 4 cases, there was a decrease of visual acuity in two. The other two subjects experienced no meaningful decrease of visual acuity. These limited outcome data indicate that appropriate diagnosis and adequate treatment (e.g., local treatment with intravitreal injections of anti-VEGF antibodies) are essential for the clinical management of RVO in cancer patients taking trametinib. However, full recovery of visual acuity may not occur in all patients developing an RVO on trametinib and immediate cessation of trametinib therapy is recommended.

Trametinib + dabrafenib combination therapy

Table 2-32 Clinical trial data of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment): Study BRF115532 (COMBI-AD)

	Dabrafenib+Trametinib N=435 n (%)	Placebo N=432 n (%)
Number of subjects with at least one event	69 (16.0)	41 (9.0)
Maximum grade		
Grade 3	3 (4.0)	0
Grade 4	1 (1.0)	0
Grade 5	0	0
SAEs	6 (9.0)	0
AE leading to death	0	0
AE outcome ¹		
Recovered/resolved	59 (86.0)	36 (88.0)
Recovering/resolving	2 (3.0)	0
Not recovered/not resolved	7 (10.0)	5 (12.0)
Recovered/resolved with sequelae	1 (1.0)	0

Numbers (n) represent counts of subjects.

Subjects may be included in more than one category

One subject may report more than one AE

¹ Outcome worst case hierarchy: Fatal > Not Recovered/Not Resolved > Recovered/Resolved with sequelae >

Recovering/Resolving > Recovered/Resolved

Source: RMP version 14 Attachment to Annex 7-Table 14.3.1-3.2

Table 2-33 Clinical trial data of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment): Study BRF113928

	Dabrafenib 150 mg bid/trametinib 2 mg qd N=82 n (%)
All AEs	11 (13)
Total Number of Events	16
CTC grade 3/4 AEs	1 (1)
Related AEs	7 (9)
AE requiring dose reduction	0 (0)
AE requiring dose interruption	3 (4)
AE leading to discontinuation	0 (0)
SAEs	2 (2)
Deaths	0 (0)

A subject with multiple occurrences is counted only once with the maximum reported severity.

MedDRA version 18.1 and (Final formatted Trametinib-Dabrafenib combo AESI terms_NSCLC_MedDRA 17 0_19Jun2014) have been used for the reporting of adverse events of special interest.

Source: Study BRF113928, Table 3.1824 (Subcomb reporting effort)

Table 2-34 Characterization of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)

Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Details				
Potential mechanisms	Potential mechanism is unknown.				
Evidence source(s) and strength of evidence	Disorders associated with visual disturbance, including RPED and RVO, may occur with trametinib as monotherapy and in combination with dabrafenib. Symptoms such as blurred vision, decreased acuity, and other visual phenomena have been reported in the clinical trials with trametinib. In clinical trials uveitis and iridocyclitis have also been reported in patients treated with trametinib in combination with dabrafenib.				
Characterization of the risk:	Adjuvant treatment of patients with Stage III melanoma with a BRAF V600 mutation, following complete resection Refer to Table 2-32 for additional information. Treatment of adult patients with advanced NSCLC with a BRAF V600 mutation Refer to Table 2-33 Clinical trial data of ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment) for additional information.				
Risk factors and risk groups	None identified yet for trametinib.				
Preventability	Treatment with trametinib should not be recommended for patients with a history or current evidence of RVO. Trametinib is not recommended in patients with a history of RVO or risk factors for RVO. Information and management guidelines are included in the proposed label.				
Impact on the benefit-risk balance of the product	The benefit-risk balance of trametinib+dabrafenib is not impacted by this risk considering the nature of the treatment indication(s) and if any it is low. The SmPC provides adequate information on timely identification and management of this risk. Data review for this risk in the latest PSUR for trametinib also confirm no impact on the benefit–risk of trametinib treatment with regards to this risk.				
Public health impact	Dabrafenib+Trametinib (Rate1, %)	Dabrafenib monotherapy (Rate2, %)	Relative risk^a (95% CI)	Risk Diff. (Rate 1- Rate 2) (95% CI)	NNH = 1/(Rate1 -Rate2)
	27 (13%)	23 (11%)	1.19 (0.70, 2.00)	2% (-4.27%, 8.37%)	50
	^a Study MEK115306 data: the combination arm (150 mg bid dabrafenib+ 2 mg once daily trametinib, N=209) relative to dabrafenib alone arm (150 mg bid, N=211). NNH: number needed to harm.				
	In MEK115306, 27 (13%) subjects who received the combination therapy (n=209), experienced ocular events, compared with 23 subjects (11%) in the dabrafenib monotherapy arm (n=211). Compared with those treated with dabrafenib alone, subjects who received the combination therapy had a 20% increased relative risk of ocular events. The absolute risk difference was 2%, which suggests that approximately 50 subjects need to be exposed to the				

Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Details
	combination therapy to cause ocular event in one subject that would not otherwise occur, if they were treated with dabrafenib alone (NNH). With appropriate management, the public health impact of ocular events is considered low.

2.7.3.1.2 Important potential risk: Developmental toxicity

Table 2-35 Clinical trial data of Developmental toxicity: Study BRF115532 (COMBI-AD)

	Dabrafenib+Trametinib N=435 n (%)	Placebo N=432 n (%)
Number of subjects with at least one event	3 (<1.0)	2 (<1.0)
Maximum grade		
Grade 3	0	0
Grade 4	0	0
Grade 5	0	0
SAEs	0	0
AE leading to death	0	0
AE outcome ¹		
Recovered/resolved	1 (33.0)	1 (50.0)
Recovering/resolving	0	0
Not recovered/not resolved	2 (67.0)	1 (50.0)
Recovered/resolved with sequelae	0	0

Numbers (n) represent counts of subjects.

Subjects may be included in more than one category

One subject may report more than one AE

¹ Outcome worst case hierarchy: Fatal > Not Recovered/Not Resolved > Recovered/Resolved with sequelae > Recovering/Resolving > Recovered/Resolved

Source: RMP version 14 Attachment to Annex 7-Table 8.01155

Table 2-36 Characterization of Developmental toxicity

Developmental toxicity	Details
Potential mechanisms	Potential mechanism is unknown.
Evidence source(s) and strength of evidence	In rats and rabbits given trametinib monotherapy, maternal and developmental toxicity (decreased fetal body weights and increased ossification variations) were observed at exposures below the exposures achieved at the recommended clinical dose of 2 mg per day. Additionally, decreased corpora lutea were observed in rats given trametinib, which may impact female fertility. It is not known whether these effects will also be seen in humans.
Characterization of the risk:	Pregnant women are excluded from participation in clinical studies with trametinib.

Developmental toxicity	Details
	Refer to Table 2-35 for additional information.
Risk factors and risk groups	Women of child-bearing potential.
Preventability	Trametinib should not be administered to pregnant women.
Impact on the benefit-risk balance of the product	In the latest PSUR, none of the cases reported developmental toxicity. In addition, no new information related to this risk was identified from published literature or clinical studies during the reporting interval of the PSUR. No impact to the benefit-risk is identified.
Public health impact	Potential public health impact is considered to be low.

2.7.3.1.3 Important potential risk: Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)

The safety of trametinib plus dabrafenib combination therapy has been evaluated in a pooled safety set of 171 paediatric patients across two studies in patients with BRAF V600 mutation-positive advanced solid tumors. The overall safety profile of combination therapy is presented in [Table 2-37](#). No adverse effects on skeletal maturation and sexual maturation were reported.

Table 2-37 Clinical trial data of Safety in patients <18 years of age (including potential adverse effects on skeletal maturation and sexual maturation) (Safety analysis set)

Category	All N=171	Patients	
	All n (%)	grades Grade n (%)	≥3
Adverse events	169 (98.8)	98 (57.3)	
Treatment-related	154 (90.1)	50 (29.2)	
SAEs	79 (46.2)	58 (33.9)	
Treatment-related	29 (17.0)	16 (9.4)	
Fatal SAEs	3 (1.8)	3 (1.8)	
Treatment-related	0	0	
AEs leading to discontinuation	13 (7.6)	6 (3.5)	
Treatment-related	11 (6.4)	5 (2.9)	
AEs leading to dose adjustment/interruption	125 (73.1)	64 (37.4)	
AEs requiring additional therapy	135 (78.9)	56 (32.7)	

Numbers (n) represent counts of patients.

A patient with multiple severity grades for an AE is only counted under the maximum grade.

MedDRA version 24.0, CTCAE version 4.03

Source: [\[SCS Appendix 1-Table 3.1-1\]](#)

Table 2-38 Characterization of Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)

Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)	Details
Potential mechanisms	Potential mechanism is unknown.
Evidence source(s) and strength of evidence	Studies in juvenile animals have shown adverse effects of trametinib which had not been observed in adult animals. The principal toxicities associated with trametinib administration to juvenile rats consisted of a delay in a physical landmark of sexual maturity and mammary gland development, lower corpora lutea and lower ovarian weights. With the exception of corneal mineralization/dystrophy and increased heart weight, similar effects have been observed in adult animals given trametinib. Bone effects consisted primarily of physeal thickening in the femur in rats given trametinib doses ≥ 0.1 mg/kg/day). Reduced long bone length was observed in a juvenile rat study. In juvenile rats, increased heart weight with no histopathology was observed at 0.35 mg/kg/day (approximately twice the adult human clinical exposure based on AUC).
Characterization of the risk:	The safety data are consistent with the currently labeled safety profiles of dabrafenib and trametinib in adults and no new signals were observed.
Risk factors and risk groups	Patients under 18 years of age.
Preventability	Preventability is addressed by communication in labeling. Currently the pediatric label clearly states that long-term data on growth and skeletal maturation in paediatric patients are currently limited.
Impact on the benefit-risk balance of the product	There was no new relevant safety concern identified from the cases retrieved during the reporting interval of the latest PSUR. In addition, no new information related to this safety concern was identified from the completed studies. The available data during the review period remains in accordance with our current understanding of the important potential risk. No impact on the benefit-risk.
Public health impact	Public health impact is considered to be low.

2.7.3.2 Part II Module SVII.3.2. Presentation of the missing information

There is no missing information for trametinib.

2.8 Part II: Module SVIII: Summary of the safety concerns

Table 2-39 Part II SVIII.1: Summary of safety concerns

Important identified risks for trametinib	Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)
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Important potential risks for trametinib	• Developmental toxicity • Long-term safety in patients < 18 years old (including potential adverse effects on skeletal maturation and sexual maturation)
Missing information for trametinib	• None

3 Part III: Pharmacovigilance plan (including post-authorization safety studies)

3.1 Part III.1. Routine pharmacovigilance activities

3.1.1 Routine pharmacovigilance activities beyond ADR reporting and signal detection

Specific adverse reaction follow-up checklists:

There is no specific adverse reaction follow-up checklist for Mekinist or Spexotras.

Other forms of routine pharmacovigilance activities:

There are no other forms of routine pharmacovigilance activities.

3.2 Part III.2. Additional pharmacovigilance activities

A long-term follow up roll-over study is ongoing; CDRB436G2401, details of which are provided below.

Study CDRB436G2401- An open label, multi-center roll-over study to assess long-term effect in pediatric patients treated with Tafinlar (dabrafenib) and/or Mekinist (trametinib).

Study short name and title:

A roll-over study to assess long-term effect in pediatric patients treated with dabrafenib and/or trametinib.

Rationale and study objectives:

This study will facilitate data collection of the long-term outcomes of pediatric subjects who have been treated in clinical trials with dabrafenib, trametinib or the combination, to assess the long-term effect on growth, development and general health of these subjects. Further, for those subjects currently on treatment in the parent protocol and would benefit from continued treatment (per investigator determination), this study will offer a mechanism to continue treatment outside the parent protocols. The primary objective is to assess the long-term safety of treatment with dabrafenib, trametinib or the combination. The secondary objectives are to assess the long-term effect of treatment with dabrafenib, trametinib or the combination on general health, growth and development; and to assess efficacy as determined by institutional standard of care procedures.

Study design:

This is a global single-arm, open-label, multi-center study to collect data on the long-term effects of dabrafenib, trametinib or the combination in pediatric subjects who have been treated on Novartis sponsored trials. No formal hypothesis will be tested. Additionally, this study will provide continued access to study medication(s) for subjects who have previously participated in dabrafenib and/or trametinib treatment studies (parent studies).

Parent studies include:

- CDRB436A2102:
Phase I/IIa, 2-part, multi-center, single-arm, open-label study to determine the safety, tolerability and pharmacokinetics of oral dabrafenib in children and adolescent patients with advanced BRAF V600-mutation positive solid tumors.
- CTMT212X2101:
Pharmacodynamics and clinical activity of the MEK inhibitor trametinib in children and adolescents patients with cancer or plexiform neurofibromas and trametinib in combination with dabrafenib in children and adolescents with cancers harboring V600 mutation.
- CDRB436G2201:
Phase II open-label global study to evaluate the effect of dabrafenib in combination with trametinib in children and adolescent patients with BRAF V600-mutation positive Low Grade Glioma (LGG) or relapsed or refractory High Grade Glioma (HGG).

Study population:

Pediatric patients (or young adults at the time of consent to this study) who have participated in an eligible parent protocol will be eligible to enroll into the observational period of this study. In addition, those patients who are currently eligible to receive treatment with dabrafenib and/or trametinib in the parent protocol, and who in the opinion of the investigator, would benefit from continued treatment will be eligible to take part in the treatment period of this study.

The investigator or designee must ensure that only patients who meet all the following inclusion and none of the exclusion criteria are offered treatment in the study.

Milestones:

Final CSR: May-2027 (Planned)

3.3 Part III.3. Summary Table of additional pharmacovigilance activities

Table 3-1 Part III.1: Ongoing and planned additional pharmacovigilance activities

Study Status	Summary objectives	of	Safety concerns addressed	Milestones	Due dates
Category 1 – Imposed mandatory additional pharmacovigilance activities which are conditions of the marketing authorization					
None					
Category 2 – Imposed mandatory additional pharmacovigilance activities which are Specific Obligations in the context of a conditional marketing authorization or a marketing authorization under exceptional circumstances					

Study Status	Summary objectives of	Safety concerns addressed	Milestones	Due dates
None				
Category 3 – Required additional pharmacovigilance activities				
CDRB436G2401	The primary objective: - To assess the long-term safety of treatment with dabrafenib, trametinib or the combination. The secondary objectives: - To assess the long-term effect of treatment with dabrafenib, trametinib or the combination on general health, growth and development. - To assess efficacy as determined by institutional standard of care procedures.	Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)	Final CSR	May-2027 (Planned)

4 Part IV: Plans for post-authorization efficacy studies

There are no plans for post-authorization efficacy studies.

5 Part V: Risk minimization measures (including evaluation of the effectiveness of risk minimization activities)

5.1 Part V.1. Routine risk minimization measures

Table 5-1 Part V.1: Description of routine risk minimization measures by safety concern

Safety concern	Routine risk minimization activities
Important identified risks	
Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Routine risk communication SmPC section 4.8. Routine risk minimization activities recommending specific clinical measures to address the risk: Trametinib is not recommended in patients with a history of RVO. Recommendations for use in patients diagnosed with RVO are included in SmPC section 4.4. Other routine risk minimization measures beyond the Product Information: None.
Important potential risks	
Developmental toxicity	Routine risk communication SmPC section 5.3. Routine risk minimization activities recommending specific clinical measures to address the risk: None. Other routine risk minimization measures beyond the Product Information: None.
Long term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)	Routine risk communication SmPC section 4.2. Routine risk minimization activities recommending specific clinical measures to address the risk: None. Other routine risk minimization measures beyond the Product Information: None.
Missing information	
None	

5.2 Part V.2. Additional Risk minimization measures

Routine risk minimization activities as described in Part V.1 are sufficient to manage the safety concerns of the medicinal product.

5.3 Part V.3. Summary of risk minimization measures

Table 5-2 Summary of pharmacovigilance activities and risk minimization activities by safety concerns

Safety concern	Risk minimization measures	Pharmacovigilance activities
Important identified risks for trametinib		
Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)	Routine risk minimization measures SmPC section 4.8. Additional risk minimization measures None.	Routine pharmacovigilance activities beyond adverse reactions reporting and signal detection: None. Additional pharmacovigilance activities: None.
Important potential risks for trametinib		
Developmental toxicity	Routine risk minimization measures SmPC section 5.3. Additional risk minimization measures None.	Routine pharmacovigilance activities beyond adverse reactions reporting and signal detection: None. Additional pharmacovigilance activities: None.
Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)	Routine risk minimization measures SmPC section 4.2. Additional risk minimization measures None.	Routine pharmacovigilance activities beyond adverse reactions reporting and signal detection: None. Additional pharmacovigilance activities: CDRB436G2401 (EU CT number 2023-509276-42-00).
Missing information for trametinib		
None		

6 Part VI: Summary of the risk management plan for Mekinist and Spexotras (trametinib)

This is a summary of the risk management plan (RMP) for Mekinist and Spexotras (trametinib). The RMP details important risks of Mekinist and Spexotras, how these risks can be minimized, and how more information will be obtained about Mekinist and Spexotras' risks and uncertainties (missing information).

Mekinist and Spexotras' summary of product characteristics (SmPC) and package leaflet give essential information to healthcare professionals and patients on how Mekinist and Spexotras should be used.

This summary of the RMP for Mekinist and Spexotras should be read in the context of all this information including the assessment reports of the evaluation and their plain-language summaries, all which are part of the European Public Assessment Report (EPAR).

Important new concerns or changes to the current ones will be included in updates of Mekinist and Spexotras' RMP.

6.1 Part VI: I. The medicine and what it is used for

Mekinist film-coated tablets contain trametinib as active substance and are authorized in the following indications:

- Trametinib as monotherapy or in combination with dabrafenib for the treatment of adults and adolescents aged 12 years and older with unresectable or metastatic melanoma with a BRAF V600 mutation;
 - Trametinib monotherapy has not demonstrated clinical activity in patients who have progressed on a prior BRAF inhibitor therapy
- Trametinib in combination with dabrafenib is indicated for the treatment of adults with advanced non-small cell lung cancer (NSCLC) with a BRAF V600 mutation;
- Trametinib in combination with dabrafenib is indicated for the adjuvant treatment of adults and adolescents aged 12 years and older with Stage III melanoma with a BRAF V600 mutation, following complete resection.
- Trametinib in combination with dabrafenib is indicated for the treatment of adult patients with locally advanced or metastatic differentiated thyroid cancer with BRAF V600E mutation, refractory to or not eligible for RAI who have progressed on prior systemic therapy.

The recommended dose of Mekinist tablets for adult patients is 2 mg once daily. The recommended dose of Mekinist tablets for adolescent patients (aged 12 years to less than 18 years) is body weight based and should be administered once daily.

Spexotras powder for oral solution contains trametinib as active substance and it is used in the following indications:

- Trametinib in combination with dabrafenib is indicated for the treatment of paediatric patients aged 1 year and older with low-grade glioma with a BRAF V600E mutation who require systemic therapy.
- Trametinib in combination with dabrafenib is indicated for the treatment of paediatric patients aged 1 year and older with high-grade glioma with a BRAF V600E mutation who have received at least one prior radiation and/or chemotherapy treatment.

The recommended dose of Spexotras powder for oral solution is body weight based and should be administered once daily.

Further information about the evaluation of Mekinist and Spexotras' benefits can be found in Mekinist and Spexotras' EPAR, including in its plain-language summary, available on the EMA website, under the medicine's webpages:

<https://www.ema.europa.eu/en/medicines/human/EPAR/mekinist>

<https://www.ema.europa.eu/en/medicines/human/EPAR/spexotras>

6.2 Part VI: II. Risks associated with the medicine and activities to minimize or further characterize the risks

Important risks of Mekinist and Spexotras, together with measures to minimize such risks and the proposed studies for learning more about Mekinist and Spexotras' risks, are outlined below.

Measures to minimize the risks identified for medicinal products can be:

- Specific information, such as warnings, precautions, and advice on correct use, in the package leaflet and SmPC addressed to patients and healthcare professionals;
- Important advice on the medicine's packaging;
- The authorised pack size — the amount of medicine in a pack is chosen so to ensure that the medicine is used correctly;
- The medicine's legal status — the way a medicine is supplied to the patient (e.g. with or without prescription) can help to minimize its risks.

Together, these measures constitute *routine risk minimization measures*.

In addition to these measures, information about adverse reactions is collected continuously and regularly analysed, including PSUR assessment so that immediate action can be taken as necessary. These measures constitute *routine pharmacovigilance activities*.

If important information that may affect the safe use of trametinib is not yet available, it is listed under 'missing information' below.

6.2.1 Part VI: II.A: List of important risks and missing information

Important risks of Mekinist and Spexotras are risks that need special risk management activities to further investigate or minimize the risk, so that the medicinal product can be safely taken. Important risks can be regarded as identified or potential. Identified risks are concerns for which there is sufficient proof of a link with the use of Mekinist and Spexotras. Potential risks are concerns for which an association with the use of this medicine is possible based on available

data, but this association has not been established yet and needs further evaluation. Missing information refers to information on the safety of the medicinal product that is currently missing and needs to be collected (e.g. on the long-term use of the medicine).

Table 6-1 List of important risks and missing information

List of important risks and missing information	
Important identified risks for trametinib	Ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)
Important potential risks for trametinib	- Developmental toxicity - Long-term safety in patients < 18 years old (including potential adverse effects on skeletal maturation and sexual maturation)
Missing information for trametinib	None

6.2.2 Part VI: II.B: Summary of important risks

Important identified risks for trametinib

Table 6-2 Important identified risk – ocular events (e.g., retinal vein occlusion, retinal pigment epithelial detachment)

Evidence for linking the risk to the medicine	Blurred vision, decreased acuity, and other visual phenomena have been reported in the clinical trials with trametinib. In clinical trials uveitis and iridocyclitis have also been reported in patients treated with trametinib in combination with dabrafenib.
Risk factors and risk groups	None identified yet for trametinib.
Risk minimization measures	Routine risk minimization measures SmPC section 4.8. Additional risk minimization measures No risk minimization measures.
Additional pharmacovigilance activities	Additional pharmacovigilance activities None.

Important Potential Risks for trametinib

Table 6-3 Important potential risk – Developmental toxicity

Evidence for linking the risk to the medicine	In rats and rabbits given trametinib monotherapy, maternal and developmental toxicity (decreased fetal body weights and increased ossification variations) were observed at exposures below the exposures achieved at the recommended clinical dose of 2 mg per day. Additionally, decreased corpora lutea were observed in rats given trametinib, which may impact female fertility. It is not known whether these effects will also be seen in humans.
Risk factors and risk groups	Women of child-bearing potential
Risk minimization measures	Routine risk minimization measures SmPC section 5.3.

Additional risk minimization measures
No risk minimization measures.

Table 6-4 Important potential risk – Long-term safety in patients <18 years old (including potential adverse effects on skeletal maturation and sexual maturation)

Evidence for linking the risk to the medicine	Studies in juvenile animals have shown adverse effects of trametinib which had not been observed in adult animals. In juvenile rats, increased heart weight with no histopathology was observed at 0.35 mg/kg/day (approximately twice the adult human clinical exposure based on AUC).
Risk factors and risk groups	Children under 18 years of age.
Risk minimization measures	Routine risk minimization measures SmPC section 4.2. Additional risk minimization measures No risk minimization measures.
Additional pharmacovigilance activities	Additional pharmacovigilance activities CDRB436G2401 (EU CT number 2023-509276-42-00)

Missing information for trametinib

There is no missing information for trametinib.

6.2.3 Part VI: II.C: Post-authorization development plan

6.2.3.1 Part VI: II.C.1 Studies which are conditions of the marketing authorization

There are no studies which are conditions of the marketing authorization or specific obligation of trametinib.

6.2.3.2 Part VI: II.C.2. Other studies in post-authorization development plan

CDRB436G2401 study is a post-authorization development plan for trametinib.

Table 6-5 Other studies in the post-authorization development plan

Study Short name	Rationale and study objectives
CDRB436G2401	This study will facilitate data collection of the long-term outcomes of pediatric subjects who have been treated in clinical trials with dabrafenib, trametinib or the combination, to assess the long-term effect on growth, development and general health of these subjects. Further, for those subjects currently on treatment in the parent protocol and would benefit from continued treatment (per investigator determination), this study will offer a mechanism to continue treatment outside the parent protocols. The primary objective is to assess the long-term safety of treatment with dabrafenib, trametinib or the combination. The secondary objectives are to assess the long-term effect of

Study Short name	Rationale and study objectives
	treatment with dabrafenib, trametinib or the combination on general health, growth and development; and to assess efficacy as determined by institutional standard of care procedures.

7 Part VII: Annexes

Annex 4 – Specific adverse drug reaction follow-up forms

There are no specific follow-up forms for Mekinist or Spexotras.

Annex 6 – Details of proposed additional risk minimization activities (if applicable)

There are no proposed additional risk minimization activities.