ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 75 mg film-coated tablets Kalydeco 150 mg film-coated tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Kalydeco 75 mg film-coated tablets

Each film-coated tablet contains 75 mg of ivacaftor.

Excipient with known effect

Each film-coated tablet contains 83.6 mg of lactose monohydrate.

Kalydeco 150 mg film-coated tablets

Each film-coated tablet contains 150 mg of ivacaftor.

Excipient with known effect

Each film-coated tablet contains 167.2 mg of lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet (tablet)

Kalydeco 75 mg film-coated tablets

Light blue, capsule-shaped film-coated tablets, printed with "V 75" in black ink on one side and plain on the other (12.7 mm \times 6.8 mm in modified tablet shape).

Kalydeco 150 mg film-coated tablets

Light blue, capsule-shaped film-coated tablets, printed with "V 150" in black ink on one side and plain on the other (16.5 mm \times 8.4 mm in modified tablet shape).

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Kalydeco tablets are indicated:

- As monotherapy for the treatment of adults, adolescents, and children aged 6 years and older and weighing 25 kg or more with cystic fibrosis (CF) who have an *R117H CFTR* mutation or one of the following gating (Class III) mutations in the cystic fibrosis transmembrane conductance regulator (*CFTR*) gene: *G551D*, *G1244E*, *G1349D*, *G178R*, *G551S*, *S1251N*, *S1255P*, *S549N* or *S549R* (see sections 4.4 and 5.1).
- In a combination regimen with tezacaftor/ivacaftor tablets for the treatment of adults, adolescents, and children aged 6 years and older with cystic fibrosis (CF) who are homozygous for the *F508del* mutation or who are heterozygous for the *F508del* mutation and have one of the following mutations in the *CFTR* gene: *P67L*, *R117C*, *L206W*, *R352Q*, *A455E*, *D579G*,

 $711+3A \rightarrow G$, S945L, S977F, R1070W, D1152H, $2789+5G \rightarrow A$, $3272-26A \rightarrow G$, and $3849+10kbC \rightarrow T$.

• In a combination regimen with ivacaftor/tezacaftor/elexacaftor tablets for the treatment of adults, adolescents, and children aged 6 years and older with cystic fibrosis (CF) who have at least one non-Class I mutation in the *CFTR* gene (see sections 4.2 and 5.1).

4.2 Posology and method of administration

Kalydeco should only be prescribed by physicians with experience in the treatment of cystic fibrosis. If the patient's genotype is unknown, an accurate and validated genotyping method should be performed before starting treatment to confirm the presence of an indicated mutation in the *CFTR* gene (see section 4.1). The phase of the poly-T variant identified with the *R117H* mutation should be determined in accordance with local clinical recommendations.

Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor

There are a limited number of patients who harbour mutations not listed in Table 6 that may be responsive to ivacaftor/tezacaftor/elexacaftor (IVA/TEZ/ELX). In these cases, ivacaftor (IVA) in combination with IVA/TEZ/ELX can be considered when the physician deems the potential benefits outweigh the potential risks and under close medical supervision. This excludes patients with two Class I (null) mutations (mutations that are known not to produce CFTR protein) as they are not expected to respond to modulator therapy (see sections 4.1, 4.4 and 5.1).

Posology

Adults, adolescents, and children aged 6 years and older should be dosed according to Table 1.

Table 1: Dosing recommendations

Age/weight	Morning dose	Evening dose			
Ivacaftor as monotherapy					
6 years and older,	One ivacaftor 150 mg tablet	One ivacaftor			
\geq 25 kg	One tvacation 130 mg tablet	150 mg tablet			
Ivacaftor in combinat	ion with tezacaftor/ivacaftor				
6 years to < 12 years,	One tezacaftor 50 mg/ivacaftor 75 mg tablet	One ivacaftor			
< 30 kg	One tezacation 50 mg/tvacation 75 mg tablet	75 mg tablet			
6 years to < 12 years,	One tezacaftor 100 mg/ivacaftor 150 mg tablet	One ivacaftor			
\geq 30 kg	One regarding 100 mg/tvacation 150 mg tablet	150 mg tablet			
12 years and older	One tezacaftor 100 mg/ivacaftor 150 mg tablet	One ivacaftor			
12 years and order	One tezacation 100 mg/tvacation 150 mg tablet	150 mg tablet			
Ivacaftor in combinat	ion with ivacaftor/tezacaftor/elexacaftor				
6 years to < 12 years,	Two ivacaftor 37.5 mg/tezacaftor 25 mg/elexacaftor	One ivacaftor			
< 30 kg	50 mg tablets	75 mg tablet			
6 years to < 12 years,	Two ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg	One ivacaftor			
\geq 30 kg	tablets	150 mg tablet			
12 years and older	Two ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg	One ivacaftor			
12 years and older	tablets	150 mg tablet			

The morning and evening dose should be taken approximately 12 hours apart with fat-containing food (see Method of administration).

Missed dose

If 6 hours or less have passed since the missed morning or evening dose, the patient should be advised to take it as soon as possible and then take the next dose at the regularly scheduled time. If more than

6 hours have passed since the time the dose is usually taken, the patient should be advised to wait until the next scheduled dose.

Patients receiving Kalydeco in a combination regimen should be advised not to take more than one dose of either medicinal product at the same time.

Concomitant use of CYP3A inhibitors

During concomitant administration with moderate or strong inhibitors of CYP3A, the ivacaftor dose should be adjusted as detailed in Table 2. Dosing intervals should be modified according to clinical response and tolerability (see sections 4.4 and 4.5).

Table 2: Dosing recommendations for concomitant use with moderate or strong CYP3A inhibitors

Age/	Moderate CYP3A inhibitors	Strong CYP3A inhibitors				
weight						
Ivacaftor as	Ivacaftor as monotherapy					
6 years and	One morning tablet of ivacaftor 150 mg	One morning tablet of ivacaftor 150 mg				
older,	once daily.	twice a week, approximately 3 to 4 days				
\geq 25 kg		apart.				
	No evening ivacaftor dose.					
		No evening ivacaftor dose.				
Ivacaftor in	a combination regimen with tezacaftor/iva	caftor				
6 years to	Alternate each day:	One morning tablet of tezacaftor				
< 12 years,	 one morning tablet of tezacaftor 	50 mg/ivacaftor 75 mg twice a week,				
< 30 kg	50 mg/ivacaftor 75 mg on the first	approximately 3 to 4 days apart.				
	day					
	 one morning tablet of ivacaftor 	No evening ivacaftor dose.				
	75 mg on the next day					
	No evening ivacaftor dose.					
6 years to	Alternate each day:	One morning tablet of tezacaftor				
< 12 years,	- one morning tablet of tezacaftor	100 mg/ivacaftor 150 mg twice a week,				
\geq 30 kg	100 mg/ivacaftor 150 mg on the	approximately 3 to 4 days apart.				
	first day					
	- one morning tablet of ivacaftor	No evening ivacaftor dose.				
	150 mg on the next day					
	N					
12	No evening ivacaftor dose.	0				
12 years	Alternate each day:	One morning tablet of tezacaftor				
and older	- one morning tablet of tezacaftor	100 mg/ivacaftor 150 mg twice a week,				
	100 mg/ivacaftor 150 mg on the first	approximately 3 to 4 days apart.				
	day - one morning tablet of ivacaftor	Ni i				
	150 mg on the next day	No evening ivacaftor dose.				
	150 mg on the next day					
	No evening ivacaftor dose.					

Age/ weight	Moderate CYP3A inhibitors	Strong CYP3A inhibitors					
	Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor						
6 years to < 12 years, < 30 kg	Alternate each day: - two morning tablets of ivacaftor 37.5 mg/tezacaftor 25 mg/ elexacaftor 50 mg on the first day - one morning tablet of ivacaftor 75 mg on the next day	Two morning tablets of ivacaftor 37.5 mg/tezacaftor 25 mg/elexacaftor 50 mg twice a week, approximately 3 to 4 days apart. No evening ivacaftor dose.					
	No evening ivacaftor dose.						
6 years to < 12 years, ≥ 30 kg	Alternate each day: - two morning tablets of ivacaftor 75 mg/tezacaftor 50 mg/ elexacaftor 100 mg on the first day - one morning tablet of ivacaftor 150 mg on the next day	Two morning tablets of ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg twice a week, approximately 3 to 4 days apart. No evening ivacaftor dose.					
12 years and older	No evening ivacaftor dose. Alternate each day: - two morning tablets of ivacaftor 75 mg/tezacaftor 50 mg/ elexacaftor 100 mg on the first day - one morning tablet of ivacaftor 150 mg on the next day No evening ivacaftor dose.	Two morning tablets of ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg twice a week, approximately 3 to 4 days apart. No evening ivacaftor dose.					

Special populations

Elderly

Very limited data are available for elderly patients treated with ivacaftor (administered as monotherapy or in a combination regimen). No dose adjustment specific to this patient population is required (see section 5.2).

Renal impairment

No dose adjustment is necessary for patients with mild to moderate renal impairment. Caution is recommended in patients with severe renal impairment (creatinine clearance less than or equal to 30 mL/min) or end-stage renal disease (see sections 4.4 and 5.2).

Hepatic impairment

No dose adjustment is necessary in patients with mild hepatic impairment (Child-Pugh Class A).

In patients with moderate hepatic impairment (Child-Pugh Class B) or severe hepatic impairment (Child-Pugh Class C), the ivacaftor dose should be adjusted as detailed in Table 3 (see sections 4.4, 4.8, and 5.2).

Table 3: Dosing recommendations for patients with moderate or severe hepatic impairment

Age/weigh	Moderate (Child-Pugh Class B)	Severe (Child-Pugh Class C)			
	Ivacaftor as monotherapy				
6 years and older, ≥ 25 kg	One morning tablet of ivacaftor 150 mg once daily.	Use is not recommended, unless the benefits are expected to outweigh the risks.			
	No evening ivacaftor dose.	If used, one morning tablet of ivacaftor 150 mg every other day or less frequently according to clinical response and tolerability.			
		No evening ivacaftor dose.			
Ivacaftor in	a combination regimen with teza	caftor/ivacaftor			
6 years to < 12 years, < 30 kg	One morning tablet of tezacaftor 50 mg/ivacaftor 75 mg once daily.	Use is not recommended, unless the benefits are expected to outweigh the risks.			
	No evening ivacaftor dose.	If used, one morning tablet of tezacaftor 50 mg/ivacaftor 75 mg once daily or less frequently according to clinical response and tolerability.			
		No evening ivacaftor dose.			
6 years to < 12 years, ≥ 30 kg	One morning tablet of tezacaftor 100 mg/ivacaftor 150 mg once daily.	Use is not recommended, unless the benefits are expected to outweigh the risks. If used, one morning tablet of tezacaftor			
	No evening ivacaftor dose.	100 mg/ivacaftor 150 mg once daily or less frequently according to clinical response and tolerability.			
12 years and older	One morning tablet of tezacaftor 100 mg/ivacaftor 150 mg once daily.	No evening ivacaftor dose. Use is not recommended, unless the benefits are expected to outweigh the risks. If used, one morning tablet of			
	No evening ivacaftor dose.	tezacaftor 100 mg/ivacaftor 150 mg once daily or less frequently according to clinical response and tolerability.			
		No evening ivacaftor dose.			

Ivacaftor in	a combination regimen with ivac	aftor/tezacaftor/elexacaftor
6 years to	Use is not recommended,	Should not be used.
< 12 years,	unless the benefits are expected	
< 30 kg	to outweigh the risks.	
	If used, the dose should be	
	adjusted as follows:	
	• Day 1: two	
	ivacaftor 37.5 mg/tezacaft	
	or	
	25 mg/elexacaftor 50 mg	
	tablets in the morning	
	Day 2: one ivacaftor 37.5 mg/tezacaft	
	or	
	25 mg/elexacaftor 50 mg	
	tablet in the morning	
	Continue alternating Day 1 and	
	Day 2 dosing thereafter.	
	No evening ivacaftor dose.	
6 years to	Use is not recommended,	Should not be used.
< 12 years,	unless the benefits are expected	
\geq 30 kg	to outweigh the risks.	
	If used, the dose should be	
	adjusted as follows:	
	• Day 1: two	
	ivacaftor 75 mg/tezacaftor	
	50 mg/elexacaftor 100 mg	
	tablets in the morning	
	• Day 2: one	
	ivacaftor 75 mg/tezacaftor	
	50 mg/elexacaftor 100 mg	
	tablet in the morning	
	Continue alternating Day 1 and	
	Day 2 dosing thereafter.	
	<i>y =y</i>	
	No evening ivacaftor dose.	
12 years	Use is not recommended,	Should not be used.
and older	unless the benefits are expected	
	to outweigh the risks.	
	T61 41- 1 111	
	If used, the dose should be	
	adjusted as follows:	
	Day 1: two	
	ivacaftor 75 mg/tezacaftor	
	50 mg/elexacaftor 100 mg	
	tablets in the morning	
	• Day 2: one	
	ivacaftor 75 mg/tezacaftor	
	50 mg/elexacaftor 100 mg	
	tablet in the morning	

Paediatric population

The safety and efficacy of ivacaftor as monotherapy have not been established in children less than 1 month of age or in children less than 6 months of age born prematurely (less than 37 weeks of gestational age), neither in combination with tezacaftor/ivacaftor in children less than 6 years of age or in combination with ivacaftor/tezacaftor/elexacaftor in children less than 2 years of age. No data are available.

Limited data are available in patients less than 6 years of age with an *R117H* mutation in the *CFTR* gene. Available data in patients aged 6 years and older are described in sections 4.8, 5.1, and 5.2.

Method of administration

For oral use.

Patients should be instructed to swallow the tablets whole. The tablets should not be chewed, crushed, or broken before swallowing because there are no clinical data currently available to support other methods of administration.

Ivacaftor tablets should be taken with fat-containing food.

Food or drink containing grapefruit should be avoided during treatment (see section 4.5).

4.3 Contraindications

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

4.4 Special warnings and precautions for use

Only patients with CF who had a *G551D*, *G1244E*, *G1349D*, *G178R*, *G551S*, *S1251N*, *S1255P*, *S549N*, *S549R* gating (Class III), *G970R* or *R117H* mutation in at least one allele of the *CFTR* gene were included in studies 770-102, 770-103, 770-111 and 770-110 (see section 5.1).

In study 770-111, four patients with the G970R mutation were included. In three of four patients the change in the sweat chloride test was < 5 mmol/L and this group did not demonstrate a clinically relevant improvement in FEV₁ after 8 weeks of treatment. Clinical efficacy in patients with the G970R mutation of the CFTR gene could not be established (see section 5.1).

Efficacy results from a phase 2 study in patients with CF who are homozygous for the F508del mutation in the CFTR gene showed no statistically significant difference in FEV_1 over 16 weeks of ivacaftor treatment compared to placebo (see section 5.1). Therefore, use of ivacaftor as monotherapy in these patients is not recommended.

Less evidence of a positive effect of ivacaftor has been shown for patients with an *R117H-7T* mutation associated with less severe disease in study 770-110 (see section 5.1).

Ivacaftor in a combination regimen with tezacaftor/ivacaftor should not be prescribed in patients with CF who are heterozygous for the *F508del* mutation and have a second *CFTR* mutation not listed in section 4.1.

Elevated transaminases and hepatic injury

In a patient with cirrhosis and portal hypertension, liver failure leading to transplantation has been reported while receiving ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor. This medicinal product should be used with caution in patients with pre-existing advanced liver disease (e.g., cirrhosis, portal hypertension) and only if the benefits are expected to outweigh the risks. If used in these patients, they should be closely monitored after the initiation of treatment (see sections 4.2, 4.8, and 5.2).

Moderate transaminase (alanine transaminase [ALT] or aspartate transaminase [AST]) elevations are common in subjects with CF. Transaminase elevations have been observed in some patients treated with ivacaftor as monotherapy and in combination regimens with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor. In patients taking ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor, these elevations have sometimes been associated with concomitant elevations in total bilirubin. Therefore, assessments of transaminases (ALT and AST) and total bilirubin are recommended for all patients prior to initiating ivacaftor, every 3 months during the first year of treatment and annually thereafter. For all patients with a history of liver disease or transaminase elevations, more frequent monitoring of liver function tests should be considered. In the event of significant elevations of transaminases (e.g., patients with ALT or AST > 5 × the upper limit of normal (ULN), or ALT or AST > 3 × ULN with bilirubin > 2 × ULN), dosing should be interrupted, and laboratory tests closely followed until the abnormalities resolve. Following resolution of transaminase elevations, the benefits and risks of resuming treatment should be considered (see sections 4.2, 4.8, and 5.2).

Hepatic impairment

Use of ivacaftor, either as monotherapy or in a combination regimen with tezacaftor/ivacaftor, is not recommended in patients aged 6 years and older with severe hepatic impairment unless the benefits are expected to outweigh the risks. These patients should not be treated with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor (see Table 3 in section 4.2, and sections 4.8 and 5.2).

For patients aged 6 years and older with moderate hepatic impairment, use of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor is not recommended. Treatment should only be considered when there is a clear medical need and the benefits are expected to outweigh the risks. If used, it should be used with caution at a reduced dose (see Table 3 in section 4.2, and sections 4.8 and 5.2).

Depression

Depression (including suicidal ideation and suicide attempt) has been reported in patients while receiving ivacaftor, mainly in a combination regimen with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor, usually occurring within three months of treatment initiation and in patients with a history of psychiatric disorders. In some cases, symptom improvement was reported after dose reduction or treatment discontinuation. Patients (and caregivers) should be alerted about the need to monitor for depressed mood, suicidal thoughts, or unusual changes in behaviour and to seek medical advice immediately if these symptoms present.

Renal impairment

Caution is recommended while using ivacaftor in patients with severe renal impairment or end-stage renal disease (see sections 4.2 and 5.2).

Mutations unlikely to respond to modulator therapy

Patients with a genotype consisting of two *CFTR* mutations that are known not to produce CFTR protein (i.e., two Class I mutations) are not expected to respond to CFTR modulator therapy.

Clinical studies comparing ivacaftor/tezacaftor/elexacaftor to tezacaftor/ivacaftor or ivacaftor

No clinical study has been conducted to directly compare ivacaftor/tezacaftor/elexacaftor to tezacaftor/ivacaftor or ivacaftor in patients not harbouring F508del variants.

Patients after organ transplantation

Ivacaftor has not been studied in patients with CF who have undergone organ transplantation. Therefore, use in transplanted patients is not recommended. See section 4.5 for interactions with ciclosporin or tacrolimus.

Rash events

The incidence of rash events with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor was higher in females than in males, particularly in females taking hormonal contraceptives. A role for hormonal contraceptives in the occurrence of rash cannot be excluded. For patients taking hormonal contraceptives who develop rash, interrupting treatment with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor and hormonal contraceptives should be considered. Following the resolution of rash, it should be considered if resuming ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor without hormonal contraceptives is appropriate. If rash does not recur, resumption of hormonal contraceptives can be considered (see section 4.8).

Interactions with medicinal products

CYP3A inducers

Exposure to ivacaftor is significantly decreased by the concomitant use of CYP3A inducers, potentially resulting in the loss of ivacaftor efficacy; therefore, co-administration of ivacaftor with strong CYP3A inducers is not recommended (see section 4.5).

CYP3A inhibitors

Exposure to ivacaftor, tezacaftor and elexacaftor are increased when co-administered with strong or moderate CYP3A inhibitors. The dose of ivacaftor must be adjusted when used concomitantly with strong or moderate CYP3A inhibitors (see Table 2 in section 4.2 and section 4.5).

Paediatric population

Cases of non-congenital lens opacities/cataracts without impact on vision have been reported in paediatric patients treated with ivacaftor and ivacaftor-containing regimens. Although other risk factors were present in some cases (such as corticosteroid use and exposure to radiation), a possible risk attributable to treatment with ivacaftor cannot be excluded. Baseline and follow-up ophthalmological examinations are recommended in paediatric patients initiating ivacaftor treatment (see section 5.3).

Excipients with known effect

Lactose

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

Sodium

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

4.5 Interaction with other medicinal products and other forms of interaction

Ivacaftor is a substrate of CYP3A4 and CYP3A5. It is a weak inhibitor of CYP3A and P-glycoprotein (P-gp) and a potential inhibitor of CYP2C9. *In vitro* studies showed that ivacaftor is not a substrate for P-gp.

Medicinal products affecting the pharmacokinetics of ivacaftor

CYP3A inducers

Co-administration of ivacaftor with rifampicin, a strong CYP3A inducer, decreased ivacaftor exposure (AUC) by 89% and decreased hydroxymethyl ivacaftor (M1) to a lesser extent than ivacaftor. Co-administration of ivacaftor with strong CYP3A inducers, such as rifampicin, rifabutin, phenobarbital, carbamazepine, phenytoin and St. John's wort (*Hypericum perforatum*), is not recommended (see section 4.4).

No dose adjustment is recommended when ivacaftor is used with moderate or weak CYP3A inducers.

CYP3A inhibitors

Ivacaftor is a sensitive CYP3A substrate. Co-administration with ketoconazole, a strong CYP3A inhibitor, increased ivacaftor exposure (measured as area under the curve [AUC]) by 8.5-fold and increased M1 to a lesser extent than ivacaftor. A reduction of the ivacaftor dose is recommended for co-administration with strong CYP3A inhibitors, such as ketoconazole, itraconazole, posaconazole, voriconazole, telithromycin and clarithromycin (see Table 2 in section 4.2 and section 4.4).

Co-administration with fluconazole, a moderate inhibitor of CYP3A, increased ivacaftor exposure by 3-fold and increased M1 to a lesser extent than ivacaftor. A reduction of the ivacaftor dose is recommended for patients taking concomitant moderate CYP3A inhibitors, such as fluconazole, erythromycin, and verapamil (see Table 2 in section 4.2 and section 4.4).

Co-administration of ivacaftor with grapefruit juice, which contains one or more components that moderately inhibit CYP3A, may increase exposure to ivacaftor. Food or drink containing grapefruit should be avoided during treatment with ivacaftor (see section 4.2).

Potential for ivacaftor to interact with transporters

In vitro studies showed that ivacaftor is not a substrate for OATP1B1 or OATP1B3. Ivacaftor and its metabolites are substrates of BCRP *in vitro*. Due to its high intrinsic permeability and low likelihood of being excreted intact, co-administration of BCRP inhibitors is not expected to alter exposure of ivacaftor and M1-IVA, while any potential changes in M6-IVA exposures are not expected to be clinically relevant.

Ciprofloxacin

Co-administration of ciprofloxacin with ivacaftor did not affect the exposure of ivacaftor. No dose adjustment is required when ivacaftor is co-administered with ciprofloxacin.

Medicinal products affected by ivacaftor

Administration of ivacaftor may increase systemic exposure of medicinal products that are sensitive substrates of CYP2C9, and/or P-gp, and/or CYP3A which may increase or prolong their therapeutic effect and adverse reactions.

CYP2C9 substrates

Ivacaftor may inhibit CYP2C9. Therefore, monitoring of the international normalised ratio (INR) is recommended during co-administration of warfarin with ivacaftor. Other medicinal products for which exposure may be increased include glimepiride and glipizide; these medicinal products should be used with caution.

Digoxin and other P-gp substrates

Co-administration with digoxin, a sensitive P-gp substrate, increased digoxin exposure by 1.3-fold, consistent with weak inhibition of P-gp by ivacaftor. Administration of ivacaftor may increase systemic exposure of medicinal products that are sensitive substrates of P-gp, which may increase or prolong their therapeutic effect and adverse reactions. When used concomitantly with digoxin or other substrates of P-gp with a narrow therapeutic index, such as ciclosporin, everolimus, sirolimus or tacrolimus, caution and appropriate monitoring should be used.

CYP3A substrates

Co-administration with (oral) midazolam, a sensitive CYP3A substrate, increased midazolam exposure 1.5-fold, consistent with weak inhibition of CYP3A by ivacaftor. No dose adjustment of CYP3A substrates, such as midazolam, alprazolam, diazepam or triazolam, is required when these are co-administered with ivacaftor.

Hormonal contraceptives

Ivacaftor has been studied with an oestrogen/progesterone oral contraceptive and was found to have no significant effect on the exposures of the oral contraceptive. Therefore, no dose adjustment of oral contraceptives is necessary.

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no or limited amount of data (less than 300 pregnancy outcomes) from the use of ivacaftor in pregnant women. Animals studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see section 5.3). As a precautionary measure, it is preferable to avoid the use of ivacaftor during pregnancy.

Breast-feeding

Limited data show that ivacaftor is excreted into human milk. A risk to the newborns/infants cannot be excluded. A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from ivacaftor therapy taking into account the benefit of breast-feeding for the child and the benefit of therapy for the woman.

Fertility

There are no data available on the effect of ivacaftor on fertility in humans. Ivacaftor had an effect on fertility in rats (see section 5.3).

4.7 Effects on ability to drive and use machines

Ivacaftor has minor influence on the ability to drive and use machines. Ivacaftor may cause dizziness (see section 4.8) and, therefore, patients experiencing dizziness should be advised not to drive or use machines until symptoms abate.

4.8 Undesirable effects

Summary of the safety profile

The most common adverse reactions experienced by patients aged 6 years and older who received ivacaftor are headache (23.9%), oropharyngeal pain (22.0%), upper respiratory tract infection (22.0%), nasal congestion (20.2%), abdominal pain (15.6%), nasopharyngitis (14.7%), diarrhoea (12.8%), dizziness (9.2%), rash (12.8%) and bacteria in sputum (12.8%). Transaminase elevations occurred in 12.8% of ivacaftor-treated patients versus 11.5% of placebo-treated patients.

In patients aged 2 to less than 6 years the most common adverse reactions were nasal congestion (26.5%), upper respiratory tract infection (23.5%), transaminase elevations (14.7%), rash (11.8%), and bacteria in sputum (11.8%).

Serious adverse reactions included abdominal pain (0.9%) and transaminase elevations (1.8%) in patients who received ivacaftor, while serious adverse reactions of rash were reported in 1.5% patients aged 12 years and older treated with a combination regimen with ivacaftor/tezacaftor/elexacaftor (see section 4.4).

Tabulated list of adverse reactions

Table 4 reflects the adverse reactions observed with ivacaftor monotherapy in clinical trials (placebo-controlled and uncontrolled studies) in which the length of exposure to ivacaftor ranged from 16 weeks to 144 weeks. Additional adverse reactions observed with ivacaftor in a combination regimen with tezacaftor/ivacaftor and/or in a combination regimen with ivacaftor/tezacaftor/elexacaftor are also provided in Table 4. The frequency of adverse reactions is defined as follows: very common ($\geq 1/100$); common ($\geq 1/100$); uncommon ($\geq 1/1000$); rare ($\geq 1/10000$); rare ($\geq 1/100000$); very rare (< 1/100000); not known (cannot be estimated from the available data). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 4: Adverse reactions

System organ class	Adverse reactions	Frequency
Infections and infestations	Upper respiratory tract infection	very common
	Nasopharyngitis	very common
	Influenza [†]	common
	Rhinitis	common
Metabolism and nutrition disorders	Hypoglycaemia [†]	common
Psychiatric disorders	Depression	not known
Nervous system disorders	Headache	very common
	Dizziness	very common
Ear and labyrinth disorders	Ear pain	common
	Ear discomfort	common

System organ class	Adverse reactions	Frequency
	Tinnitus	common
	Tympanic membrane hyperaemia	common
	Vestibular disorder	common
	Ear congestion	uncommon
Respiratory, thoracic and	Oropharyngeal pain	very common
mediastinal disorders	Nasal congestion	very common
	Abnormal breathing [†]	common
	Rhinorrhoea [†]	common
	Sinus congestion	common
	Pharyngeal erythema	common
	Wheezing [†]	uncommon
Gastrointestinal disorders	Abdominal pain	very common
	Diarrhoea	very common
	Abdominal pain upper [†]	common
	Flatulence [†]	common
	Nausea*	common
Hepatobiliary disorders	Transaminase elevations	very common
	Alanine aminotransferase	very common
	increased [†] Aspartate aminotransferase increased [†]	common
	Liver injury	not known
	Total bilirubin increase	not known
Skin and subcutaneous tissue	Rash	very common
disorders	Acne [†]	common
	Pruritus [†]	common
Reproductive system and breast	Breast mass	common
disorders	Breast inflammation	uncommon
	Gynaecomastia	uncommon
	Nipple disorder	uncommon
	Nipple pain	uncommon
Investigations	Bacteria in sputum	very common
-	Blood creatine phosphokinase increased [†]	common
	Blood pressure increased [†]	uncommon

^{*} Adverse reaction and frequency reported from clinical studies with ivacaftor in combination with tezacaftor/ivacaftor.

Description of selected adverse reactions

Transaminase elevations

During the 48-week placebo-controlled studies 770-102 and 770-103 of ivacaftor as monotherapy in patients aged 6 years and older, the incidence of maximum transaminase (ALT or AST) > 8, > 5 or $> 3 \times$ ULN was 3.7%, 3.7% and 8.3% in ivacaftor-treated patients and 1.0%, 1.9% and 8.7% in

[†] Adverse reaction and frequency reported from clinical studies with ivacaftor in combination with ivacaftor/tezacaftor/elexacaftor.

[^] Liver injury (ALT and AST and total bilirubin increase) reported from post-marketing data with ivacaftor in combination with ivacaftor/tezacaftor/elexacaftor. This also included liver failure leading to transplantation in a patient with pre-existing cirrhosis and portal hypertension. Frequency cannot be estimated from the available data.

placebo-treated patients, respectively. Two patients, one on placebo and one on ivacaftor permanently discontinued treatment for elevated transaminases, each $> 8 \times \text{ULN}$. No ivacaftor-treated patients experienced a transaminase elevation $> 3 \times \text{ULN}$ associated with elevated total bilirubin $> 1.5 \times \text{ULN}$. In ivacaftor-treated patients, most transaminase elevations up to $5 \times \text{ULN}$ resolved without treatment interruption. Ivacaftor dosing was interrupted in most patients with transaminase elevations $> 5 \times \text{ULN}$. In all instances where dosing was interrupted for elevated transaminases and subsequently resumed, ivacaftor dosing was able to be resumed successfully (see section 4.4).

During the placebo-controlled phase 3 studies (up to 24 weeks) of tezacaftor/ivacaftor, the incidence of maximum transaminase (ALT or AST) > 8, > 5, or $> 3 \times$ ULN were 0.2%, 1.0%, and 3.4% in tezacaftor/ivacaftor-treated patients, and 0.4%, 1.0%, and 3.4% in placebo-treated patients. One patient (0.2%) on therapy and 2 patients (0.4%) on placebo permanently discontinued treatment for elevated transaminases. No patients treated with tezacaftor/ivacaftor experienced a transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN.

During the 24-week, placebo-controlled, phase 3 study of ivacaftor/tezacaftor/elexacaftor, these figures were 1.5%, 2.5%, and 7.9% in ivacaftor/tezacaftor/elexacaftor-treated patients and 1.0%, 1.5%, and 5.5% in placebo-treated patients. The incidence of adverse reactions of transaminase elevations was 10.9% in ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor-treated patients and 4.0% in placebo-treated patients.

Post-marketing cases of treatment discontinuation due to elevated transaminases have been reported (see section 4.4).

Rash events

In study 445-102, the incidence of rash events (e.g., rash, rash pruritic) was 10.9% in ivacaftor/tezacaftor/elexacaftor-treated patients and 6.5% in placebo-treated patients. The rash events were generally mild to moderate in severity. The incidence of rash events by patient sex was 5.8% in males and 16.3% in females in ivacaftor/tezacaftor/elexacaftor-treated patients and 4.8% in males and 8.3% in females in placebo-treated patients. In patients treated with ivacaftor/tezacaftor/elexacaftor, the incidence of rash events was 20.5% in females taking hormonal contraceptive and 13.6% in females not taking hormonal contraceptive (see section 4.4).

Increased creatine phosphokinase

In study 445-102, the incidence of maximum creatine phosphokinase > 5 x the ULN was 10.4% in ivacaftor/tezacaftor/elexacaftor-treated patients and 5.0% in placebo-treated patients. The observed creatine phosphokinase elevations were generally transient and asymptomatic and many were preceded by exercise. No ivacaftor/tezacaftor/elexacaftor-treated patients discontinued treatment for increased creatine phosphokinase.

Increased blood pressure

In study 445-102, the maximum increase from baseline in mean systolic and diastolic blood pressure was 3.5 mmHg and 1.9 mmHg, respectively for ivacaftor/tezacaftor/elexacaftor-treated patients (baseline: 113 mmHg systolic and 69 mmHg diastolic) and 0.9 mmHg and 0.5 mmHg, respectively for placebo-treated patients (baseline: 114 mmHg systolic and 70 mmHg diastolic).

The proportion of patients who had systolic blood pressure > 140 mmHg or diastolic blood pressure > 90 mmHg on at least two occasions was 5.0% and 3.0%, respectively in ivacaftor/tezacaftor/elexacaftor-treated patients compared with 3.5% and 3.5%, respectively in placebo-treated patients.

Paediatric population

Ivacaftor as monotherapy

Safety of ivacaftor as monotherapy for 24 weeks was evaluated in 43 patients between 1 month to less than 24 months of age (with 7 of them less than 4 months old), 34 patients between 2 to less than 6 years of age, 61 patients between 6 to less than 12 years of age and 94 patients between 12 to less than 18 years of age.

The safety profile of ivacaftor (as monotherapy or in a combination regimen) is generally consistent among paediatric patients and is also consistent with adult patients.

The incidence of transaminase elevations (ALT or AST) observed in studies 770-103, 770-111 and 770-110 (patients aged 6 to less than 12 years), study 770-108 (patients aged 2 to less than 6 years), and study 770-124 (patients aged 1 to less than 24 months) are described in Table 5. In the placebo-controlled studies, the incidence of transaminase elevations were similar between treatment with ivacaftor (15.0%) and placebo (14.6%). Peak LFT elevations were generally higher in paediatric patients than in older patients. Across all populations, peak LFT elevations returned to baseline levels following interruption, and in almost all instances where dosing was interrupted for elevated transaminases and subsequently resumed, ivacaftor dosing was able to be resumed successfully (see section 4.4). Cases suggestive of positive rechallenge were observed.

In study 770-108 ivacaftor was permanently discontinued in one patient. In study 770-124, in the cohort of patients aged 1 month to less than 4 months, a 1-month old (14.3%) patient had transaminase values of ALT $> 8 \times$ ULN and AST > 3 to $\le 5 \times$ ULN, which led to discontinuation of ivacaftor treatment (see section 4.4 for management of elevated transaminases).

Table 5: Transaminase elevations in patients aged 1 month to < 12 years treated with ivacaftor as monotherapy

Age group	n	% of Patients > 3 × ULN	% of Patients > 5 × ULN	% of Patients > 8 × ULN
6 to < 12 years	40	15.0% (6)	2.5% (1)	2.5% (1)
2 to < 6 years	34	14.7% (5)	14.7% (5)	14.7% (5)
12 to < 24 months	18	27.8% (5)	11.1% (2)	11.1% (2)
1 to < 12 months	24	8.3% (2)	4.2% (1)	4.2% (1)

Ivacaftor in a combination regimen with tezacaftor/ivacaftor

The safety of tezacaftor/ivacaftor in combination with ivacaftor was evaluated in 124 patients between 6 to less than 12 years of age. The tezacaftor 100 mg/ivacaftor 150 mg and ivacaftor 150 mg dose has not been investigated in clinical studies in children aged 6 to less than 12 years weighing 30 to < 40 kg.

The safety profile is generally consistent among children and adolescents, and is also consistent with adult patients.

During the 24-week, open-label phase 3 study in patients aged 6 to less than 12 years (study 661-113 part B, n = 70), the incidence of maximum transaminase (ALT or AST) > 8, > 5, and > 3 × ULN were 1.4%, 4.3%, and 10.0%, respectively. No tezacaftor/ivacaftor treated patients experienced a transaminase elevation > 3 × ULN associated with elevated total bilirubin > 2 × ULN or discontinued tezacaftor/ivacaftor treatment due to transaminase elevations. One patient interrupted treatment due to elevated transaminases, and subsequently resumed tezacaftor/ivacaftor treatment successfully (see section 4.4 for management of elevated transaminases).

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

The safety data of ivacaftor/tezacaftor/elexacaftor in combination with ivacaftor in studies 445-102, 445-103, 445-104, 445-106, 445-111, and 445-124 was evaluated in 272 patients between 2 to less than 18 years of age. The safety profile is generally consistent among paediatric and adult patients.

During study 445-106 in patients aged 6 to less than 12 years, the incidence of maximum transaminase (ALT or AST) > 8, > 5, and $> 3 \times$ ULN were 0.0%, 1.5%, and 10.6%, respectively. No ivacaftor/tezacaftor/elexacaftor-treated patients had transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN or discontinued treatment due to transaminase elevations (see section 4.4).

During study 445-111 in patients aged 2 to less than 6 years, the incidence of maximum transaminase (ALT or AST) > 8, > 5, and $> 3 \times$ ULN were 1.3%, 2.7%, and 8.0%, respectively. No ivacaftor/tezacaftor/elexacaftor-treated patients had transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN or discontinued treatment due to transaminase elevations (see section 4.4).

Rash

During study 445-111 in patients aged 2 to less than 6 years, 15 (20.0%) subjects had at least 1 rash event, 4 (9.8%) females and 11 (32.4%) males.

Lenticular opacity

One patient had an adverse event of lenticular opacity.

Reporting of suspected adverse reactions

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

4.9 Overdose

No specific antidote is available for overdose with ivacaftor. Treatment of overdose consists of general supportive measures including monitoring of vital signs, liver function tests and observation of the clinical status of the patient.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other respiratory system products, ATC code: R07AX02

Mechanism of action

Ivacaftor as monotherapy

Ivacaftor is a potentiator of the CFTR protein, i.e., *in vitro* ivacaftor increases CFTR channel gating to enhance chloride transport in specified gating mutations (as listed in section 4.1) with reduced channel-open probability compared to normal CFTR. Ivacaftor also potentiated the channel-open probability of *R117H-CFTR*, which has both low channel-open probability (gating) and reduced channel current amplitude (conductance). The *G970R* mutation causes a splicing defect resulting in

little-to-no CFTR protein at the cell surface which may explain the results observed in subjects with this mutation in study 770-111 (see Pharmacodynamic effects and Clinical efficacy and safety).

In vitro responses seen in single channel patch clamp experiments using membrane patches from rodent cells expressing mutant CFTR forms do not necessarily correspond to *in vivo* pharmacodynamic response (e.g., sweat chloride) or clinical benefit. The exact mechanism leading ivacaftor to potentiate the gating activity of normal and some mutant CFTR forms in this system has not been completely elucidated.

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

Elexacaftor and tezacaftor are CFTR correctors that bind to different sites on the CFTR protein and have an additive effect in facilitating the cellular processing and trafficking of CFTR to increase the amount of CFTR protein delivered to the cell surface compared to either molecule alone. When ivacaftor is administered in combination with ivacaftor/tezacaftor/elexacaftor, the combined effect is increased quantity and function of CFTR at the cell surface, resulting in increased CFTR activity as measured by CFTR mediated chloride transport.

CFTR Chloride Transport Assay in Fischer Rat Thyroid (FRT) cells expressing mutant CFTR

The chloride transport response of mutant CFTR protein to ivacaftor/tezacaftor/elexacaftor was determined in Ussing chamber electrophysiology studies using a panel of FRT cell lines transfected with individual *CFTR* mutations. Ivacaftor/tezacaftor/elexacaftor increased chloride transport in FRT cells expressing select *CFTR* mutations.

The *in vitro* CFTR chloride transport response threshold was designated as a net increase of at least 10% of normal over baseline because it is predictive or reasonably expected to predict clinical response. For individual mutations, the magnitude of the net change over baseline in CFTR-mediated chloride transport *in vitro* is not correlated with the magnitude of clinical response.

In CF, the presence of one *CFTR* mutation responsive to ivacaftor/tezacaftor/elexacaftor based on *in vitro* data in FRT cells, will likely result in a clinical response.

Table 6 lists ivacaftor/tezacaftor/elexacaftor-responsive *CFTR* mutations. The occurrence of *CFTR* mutations listed in this table should not be used in lieu of a diagnosis of cystic fibrosis, nor as a sole determinant for prescribing purposes.

Table 6: CFTR mutations identified to be responsive to ivacaftor/tezacaftor/elexacaftor based on clinical and/or in vitro data

293A→G	E217G	H620Q	N900K	S50P
314del9	E264V	H939R	N1088D	S108F
546insCTA	E282D	H939R;H949L [‡]	N1195T	S158N
548insTAC	E292K	H954P	N1303I	S182R
$711+3A\rightarrow G^*$	E384K	H1054D	N1303K*	S308P
1140-1151dup	E403D	H1079P	P5L [†]	S341P
1336K	E474K	H1085P	P67L*	S364P
1461insGAT	E527G	H1085R	P111L	S434P
1507_1515del9	E588V	H1375N	P140S	S492F
2055del9	E822K	H1375P	P205S	S519G
2183A→G	E831X	I86M	P439S	S531P
2789+5G→A*	E1104K	I105N	P499A	S549I
2851A/G	E1104V	I125T	P574H	S549N
3007del6	E1126K	I148L	P750L	S549R*
3132T→G	E1221V	I148N	P798S	S557F

3141del9	E1228K	I175V	P988R	S589I
3143del9	E1228K E1409K	1331N	P1013H	S589N
$3272-26A \rightarrow G^{*\dagger}$	E1409K E1433K	1336L	P1013L	S624R
3331del6	F87L	1330L 1444S	P1021L	S686Y
3410T→C	F191V	14443 1497S	P1021T	S737F
3523A→G	F200I	1502T	P10211 P1372T	S821G
	F311del	1506L	Q30P	S898R
3601A→C		1506V	~	
3761T→G 3791C/T	F311L		Q98P	S912L
	F312del	I506V;D1168G [‡]	Q98R	S912L;G1244V [‡]
3849+10kbC→T*†	F433L	I521S	Q151K	S912T
3850G→A	F508C;S1251 N [‡]	I530N	Q179K	S945L*†
3978G→C	F508del*	I556V	Q237E	S955P
A46D	F508del;R1438W [‡]	I586V	Q237H	S977F
A62P	F575Y	I601F	Q237P	S977F;R1438W ‡
A107G	F587I	I618N	Q359K;T360K [‡]	-
A120T	F587L	I618T	Q359R	S1045Y
A141D	F693L(TTG)	I980K	Q372H	S1118F
A155P	F932S	I1023R	Q493L	S1159F
A234D	F1016S	I1139V	Q493R	S1159P
A234V	F1052V	I1203V	Q552P	S1188L
A238V	F1074L	I1234L	Q1012P	S1251N
A309D	F1078S	I1234V	Q1209P	S1255P
A349V	F1099L	I1269N	Q1291H	T338I
A357T	F1107L	I1366N	Q1291R	T351I
A455E*†	G27E	I1366T	Q1313K	T351S
A455V	G27R	K162E	Q1352H	T351S;R851L [‡]
A457T	G126D	K464E	R31L	T388M
A462P	G178E	K464N	R74Q	T465I
A534E	G178R	K522E	R74Q;R297Q [‡]	T501A
A554E	G194R	K522Q	R74Q;V201M;D1270N [‡]	T582S
A566D	G194V	K951E	R74W	T908N
A872E	G213E	K1060T	R74W;D1270N [‡]	T990I
A1006E	G213E;R668C [‡]	L15P	R74W;R1070W;D1270	T1036N*
A1025D	G213V	L15P;L1253F‡	N [‡]	T1057R
A1067P	G226R	L32P	R74W;S945L [‡]	T1086A
A1067T	G239R	L88S	R74W;V201M [‡]	T1086I
A1067V	G253R	L102R;F1016S [‡]	R74W;V201M;D1270N [‡]	T1246I
A1081V	G314E	L137P	R74W;V201M;L997F‡	T1299I
A1087P	G314R	L159S	R75L	T1299K
A1319E	G424S	L165S	R75Q;L1065P [‡]	V11I
A1374D	G437D	L167R	R75Q;N1088D [‡]	V93D
A1466S	G461R	L206W*†	R75Q;S549N [‡]	V201M
C225R	G461V	L210P	R117C [†]	V232A
C491R	G463V	L293P	R117C;G576A;R668C [‡]	V232D
C590Y	G480C	L327P	R117G	V317A
C866Y	G480D	L333F	R117H*	V322M
c.1367 1369dupTTG	G480S	L333H	R117L	V392G
D58H	G500D	L346P	R117L;L997F‡	V456A
D58V	G545R	L441P	R117P	V456F
D110E	G551A	L453S	R248K	V520I
D110H	G551D*	L467F	R258G	V562I;A1006E‡
D110N	G551R	L558F	R297Q	V562L
D192G	G551S	L619S	R334L	V591A
D192N	G576A;R668C [‡]	L633P	R334Q	V603F
D373N	G576A;S1359Y [‡]	L636P	R334W	V920L
			1	. , = = =

D426N	G622D	L927P	R347H*	V920M
D443Y	G622V	L967F;L1096R [‡]	R347L	V1008D
D443Y;G576A;R668C [‡]	G628A	L973F	R347P	V1000D V1010D
D529G	G628R	L1011S	R352Q	V1153E
D565G	G85E*†	L1065R	R352W	V1240G
D567N	G930E	L1077P*†	R516S	V1293G
D579G	G970D	L1227S	R553Q	V1293I
D614G	G970S	L1324P	R555G	V1415F
D651H	G970V	L1335P	R600S	W202C
D651N	G1047D	L1388P	R709Q	W361R
D806G	G1047R	L1480P	R751L	W496R
D924N	G1061R	M150K	R792G	W1098C
D979A	G1069R	M150R	R792Q	W1282G
D979V	G1123R	M152L	R810G	W1282R
D985H	G1173S	M152V	R851L	Y89C
D985Y	G1237V	M265R	R933G	Y109H
D993A	G1244E	M348K	R1048G	Y109N
D993G	G1244R	M394L	R1066C	Y122C
D993Y	G1247R	M469V	R1066G	Y161C
D1152A	G1249E	M498I	R1066H*†	Y161D
D1152H*†	G1249R	M952I	R1070P	Y161S
D1270N*	G1265V	M952T	R1070Q	Y301C
D1270Y	G1298V	M961L	R1070W	Y563N
D1312G	G1349D	M1101K*†	R1162Q	Y913S
D1377H	G149R;G576A;R668C [‡]	M1137R	R1239S	Y919C
D1445N	H139L	M1137V	R1283G	Y1014C
E56K	H139R	M1210K	R1283M	Y1032C
E60K	H146R	N186K	R1283S	Y1032N
E92K	H199Q	N187K	R1438W	Y1073C
E116K	H199Y	N396Y	S13F	Y1092H
E116Q	H609L	N418S	S13P	Y1381H
E193K	H620P		S18I	
			S18N	
TEL 1 14 CE		7500 1 1 CETTO		

There are people with CF harbouring two, rare non-F508del CFTR mutations not listed in Table 6. Provided that they do not harbour two class I (null) mutations (mutations that are known not to produce CFTR protein) (see section 4.1), they may respond to treatment. In these cases, Kaftrio can be considered when the physician deems the potential benefits outweigh the potential risks and under close medical supervision.

The individual diagnosis of CF should be based on diagnostic guidelines and clinical judgement as considerable variability exists in phenotype for patients harbouring the same genotype.

- * Mutations supported by clinical data.
- [†] Mutations supported by Real-World data in \geq 5 patients.
- [‡] Complex/compound mutations where a single allele of the *CFTR* gene has multiple mutations; these exist independent of the presence of mutations on the other allele.

Non-annotated mutations are included based on the FRT assay in which a positive response is indicative of a clinical response.

Pharmacodynamic effects

Ivacaftor as monotherapy

In studies 770-102 and 770-103 in patients with the *G551D* mutation in one allele of the *CFTR* gene, ivacaftor led to rapid (15 days), substantial (the mean change in sweat chloride from baseline through week 24 was -48 mmol/L [95% CI: -51, -45] and -54 mmol/L [95% CI: -62, -47], respectively) and sustained (through 48 weeks) reductions in sweat chloride concentration.

In study 770-111, part 1 in patients who had a non-*G551D* gating mutation in the *CFTR* gene, treatment with ivacaftor led to a rapid (15 days) and substantial mean change from baseline in sweat chloride of -49 mmol/L (95% CI: -57, -41) through 8 weeks of treatment. However, in patients with the *G970R-CFTR* mutation, the mean (SD) absolute change in sweat chloride at week 8 was -6.25 (6.55) mmol/L. Similar results to part 1 were seen in part 2 of the study. At the 4-week follow-up visit (4 weeks after dosing with ivacaftor ended), mean sweat chloride values for each group were trending to pre-treatment levels.

In study 770-110 in patients aged 6 years or older with CF who had an *R117H* mutation in the *CFTR* gene, the treatment difference in mean change in sweat chloride from baseline through 24 weeks of treatment was -24 mmol/L (95% CI: -28, -20). In subgroup analyses by age, the treatment difference was -21.87 mmol/L (95% CI: -26.46, -17.28) in patients aged 18 years or older, and -27.63 mmol/L (95% CI: -37.16, -18.10) in patients aged 6 to 11 years. Two patients 12 to 17 years of age were enrolled in this study.

Ivacaftor in a combination regimen with tezacaftor/ivacaftor

In study 661-106 (patients homozygous for the *F508del* mutation), the treatment difference between ivacaftor in combination with tezacaftor/ivacaftor and placebo in mean absolute change from baseline in sweat chloride through week 24, was -10.1 mmol/L (95% CI: -11.4, -8.8).

In study 661-108 (patients heterozygous for the *F508del* mutation and a second mutation associated with residual CFTR activity), the treatment difference in mean absolute change from baseline in sweat chloride through week 8 was -9.5 mmol/L (95% CI: -11.7, -7.3) between tezacaftor/ivacaftor in combination with ivacaftor and placebo, and -4.5 mmol/L (95% CI: -6.7, -2.3) between ivacaftor and placebo.

In study 661-115 (patients aged 6 to less than 12 years who were homozygous or heterozygous for the *F508del* mutation and a second mutation associated with residual CFTR activity), the within treatment mean absolute change in sweat chloride from baseline at week 8 was -12.3 mmol/L (95% CI: -15.3, -9.3) in the tezacaftor/ivacaftor group.

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

In study 445-102 (patients with an *F508del* mutation on one allele and a mutation on the second allele that predicts either no production of a CFTR protein or a CFTR protein that does not transport chloride and is not responsive to ivacaftor and tezacaftor/ivacaftor (minimal function mutation) *in vitro*), the treatment difference of ivacaftor/tezacaftor/elexacaftor compared to placebo for mean absolute change in sweat chloride from baseline through week 24 was -41.8 mmol/L (95% CI: -44.4, -39.3).

In study 445-103 (patients homozygous for the *F508del* mutation), the treatment difference of ivacaftor/tezacaftor/elexacaftor compared to tezacaftor/ivacaftor for mean absolute change in sweat chloride from baseline at week 4 was -45.1 mmol/L (95% CI: -50.1, -40.1).

In study 445-104 (patients heterozygous for the *F508del* mutation and a mutation on the second allele with a gating defect or residual CFTR activity), the treatment difference of ivacaftor/tezacaftor/elexacaftor compared to the control group (ivacaftor monotherapy group or tezacaftor/ivacaftor in combination with ivacaftor group) for mean absolute change in sweat chloride from baseline through week 8 was -23.1 mmol/L (95% CI: -26.1, -20.1).

In study 445-106 (patients aged 6 to less than 12 years, homozygous for the *F508del* mutation or heterozygous for the *F508del* mutation and a minimal function mutation), the mean absolute change in sweat chloride from baseline (n=62) through week 24 (n=60) was -60.9 mmol/L (95% CI: -63.7, -58.2)*. The mean absolute change in sweat chloride from baseline through week 12 (n=59) was -58.6 mmol/L (95% CI: -61.1, -56.1).

In study 445-116 (patients aged 6 to less than 12 years who are heterozygous for the *F508del* mutation and a minimal function mutation), the mean treatment difference for the ivacaftor/tezacaftor/elexacaftor in combination with ivacaftor group versus placebo for the absolute change in sweat chloride from baseline through week 24 was -51.2 mmol/L (95% CI: -55.3, -47.1).

In study 445-124 (patients aged 6 years and older with a qualifying non-F508del, ivacaftor/tezacaftor/elexacaftor-responsive CFTR mutation), the mean absolute change in sweat chloride from baseline through week 24 compared to placebo was -28.3 mmol/L (95% CI: -32.1, -24.5 mmol/L; P < 0.0001).

Clinical efficacy and safety

Ivacaftor as monotherapy

Studies 770-102 and 770-103: studies in patients with CF with G551D gating mutations

The efficacy of ivacaftor has been evaluated in two phase 3 randomised, double-blind, placebo-controlled, multi-centre studies of clinically stable patients with CF who had the G551D mutation in the CFTR gene on at least one allele and had $FEV_1 \ge 40\%$ predicted.

Patients in both studies were randomised 1:1 to receive either 150 mg of ivacaftor or placebo every 12 hours with food containing fat for 48 weeks in addition to their prescribed CF therapies (e.g., tobramycin, dornase alfa). The use of inhaled hypertonic sodium chloride was not permitted.

Study 770-102 evaluated 161 patients who were 12 years of age or older; 122 (75.8%) patients had the *F508del* mutation in the second allele. At the start of the study, patients in the placebo group used some medicinal products at a higher frequency than the ivacaftor group. These medicinal products included dornase alfa (73.1% versus 65.1%), salbutamol (53.8% versus 42.2%), tobramycin (44.9% versus 33.7%) and salmeterol/fluticasone (41.0% versus 27.7%). At baseline, mean predicted FEV₁ was 63.6% (range: 31.6% to 98.2%) and mean age was 26 years (range: 12 to 53 years).

Study 770-103 evaluated 52 patients who were 6 to 11 years of age at screening; mean (SD) body weight was 30.9 (8.63) kg; 42 (80.8%) patients had the *F508del* mutation in the second allele. At baseline, mean predicted FEV₁ was 84.2% (range: 44.0% to 133.8%) and mean age was 9 years (range: 6 to 12 years); 8 (30.8%) patients in the placebo group and 4 (15.4%) patients in the ivacaftor group had an FEV₁ less than 70% predicted at baseline.

The primary efficacy endpoint in both studies was the mean absolute change from baseline in percent predicted FEV₁ through 24 weeks of treatment.

The treatment difference between ivacaftor and placebo for the mean absolute change (95% CI) in percent predicted FEV_1 from baseline through week 24 was 10.6 percentage points (8.6, 12.6) in study 770-102 and 12.5 percentage points (6.6, 18.3) in study 770-103. The treatment difference between ivacaftor and placebo for the mean relative change (95% CI) in percent predicted FEV_1 from baseline through week 24 was 17.1% (13.9, 20.2) in study 770-102 and 15.8% (8.4, 23.2) in study 770-103. The mean change from baseline through week 24 in FEV_1 (L) was 0.37 L in the ivacaftor group and 0.01 L in the placebo group in study 770-102 and 0.30 L in the ivacaftor group and 0.07 L in the placebo group in study 770-103. In both studies, improvements in FEV_1 were rapid in onset (day 15) and durable through 48 weeks.

The treatment difference between ivacaftor and placebo for the mean absolute change (95% CI) in percent predicted FEV₁ from baseline through week 24 in patients 12 to 17 years of age in study 770-102 was 11.9 percentage points (5.9, 17.9). The treatment difference between ivacaftor and placebo for

^{*} Not all participants included in the analyses had data available for all follow-up visits, especially from week 16 onwards. The ability to collect data at week 24 was hampered by the COVID-19 pandemic. Week 12 data were less impacted by the pandemic.

the mean absolute change (95% CI) in percent predicted FEV_1 from baseline through week 24 in patients with baseline predicted FEV_1 greater than 90% in study 770-103 was 6.9 percentage points (-3.8, 17.6).

The results for clinically relevant secondary endpoints are shown in Table 7.

Table 7: Effect of ivacaftor on other efficacy endpoints in studies 770-102 and 770-103

	Study 770-102		Study 770-103	
Endpoint	Treatment difference ^a (95% CI)	<i>P</i> -value	Treatment difference ^a (95% CI)	<i>P-</i> value
Mean absolute change from				
Through week 24	8.1	< 0.0001	6.1	0.1092
	(4.7, 11.4)		(-1.4, 13.5) 5.1	
Through week 48	8.6	< 0.0001	5.1	0.1354
_	(5.3, 11.9)		(-1.6, 11.8)	
Relative risk of pulmonar				
Through week 24	0.40^{d}	0.0016	NA	NA
Through week 48	0.46^{d}	0.0012	NA	NA
Mean absolute change from	m baseline in body	weight (kg)		
At week 24	2.8	< 0.0001	1.9	0.0004
	(1.8, 3.7)		(0.9, 2.9)	
At week 48	2.7	0.0001	2.8	0.0002
	(1.3, 4.1)		(1.3, 4.2)	
Mean absolute change from baseline in BMI (kg/m²)				
At week 24	0.94	< 0.0001	0.81	0.0008
	(0.62, 1.26)		(0.34, 1.28)	
At week 48	0.93	< 0.0001	1.09	0.0003
	(0.48, 1.38)		(0.51, 1.67)	
Mean change from baseline in z-scores				
Weight-for-age z-score at	0.33	0.0260	0.39	< 0.0001
week 48 ^e	(0.04, 0.62)		(0.24, 0.53)	
BMI-for-age z-score at	0.33	0.0490	0.45	< 0.0001
week 48 ^e	(0.002, 0.65)		(0.26, 0.65)	

CI: Confidence Interval; NA: not analysed due to low incidence of events

Study 770-111: study in patients with CF with non-G551D gating mutations

Study 770-111 was a phase 3, two-part, randomised, double-blind, placebo-controlled, crossover study (part 1) followed by a 16-week open-label extension period (part 2) to evaluate the efficacy and safety of ivacaftor in patients with CF aged 6 years and older who have a *G970R* or non-*G551D* gating mutation in the *CFTR* gene (*G178R*, *S549N*, *S549R*, *G551S*, *G1244E*, *S1251N*, *S1255P* or *G1349D*).

In part 1, patients were randomised 1:1 to receive either 150 mg of ivacaftor or placebo every 12 hours with fat-containing food for 8 weeks in addition to their prescribed CF therapies and crossed over to the other treatment for the second 8 weeks after a 4- to 8-week washout period. The use of inhaled hypertonic saline was not permitted. In part 2, all patients received ivacaftor as indicated in part 1 for 16 additional weeks. The duration of continuous ivacaftor treatment was 24 weeks for patients

^a Treatment difference = effect of ivacaftor – effect of placebo.

b CFQ-R: Cystic Fibrosis Questionnaire-Revised is a disease-specific, health-related quality-of-life measure for CF

^c Study 770-102 data were pooled from CFQ-R for adults/adolescents and CFQ-R for children 12 to 13 years of age; study 770-103 data were obtained from CFQ-R for children 6 to 11 years of age.

d Hazard ratio for time to first pulmonary exacerbation.

e In subjects under 20 years of age (CDC growth charts).

randomised to part 1 placebo/ivacaftor treatment sequence and 16 weeks for patients randomised to part 1 ivacaftor/placebo treatment sequence.

Thirty-nine patients (mean age 23 years) with baseline $FEV_1 \ge 40\%$ predicted (mean $FEV_1 78\%$ predicted [range: 43% to 119%]) were enrolled. Sixty-two percent (24/39) of them carried the F508del-CFTR mutation in the second allele. A total of 36 patients continued into part 2 (18 per treatment sequence).

In part 1 of study 770-111, the mean FEV₁ percent predicted at baseline in placebo-treated patients was 79.3% while in ivacaftor-treated patients this value was 76.4%. The mean overall post-baseline value was 76.0% and 83.7%, respectively. The mean absolute change from baseline through week 8 in percent predicted FEV₁ (primary efficacy endpoint) was 7.5% in the ivacaftor period and -3.2% in the placebo period. The observed treatment difference (95% CI) between ivacaftor and placebo was 10.7% (7.3, 14.1) (*P* < 0.0001).

The effect of ivacaftor in the overall population of study 770-111 (including the secondary endpoints absolute change in BMI at 8 weeks of treatment and absolute change in the respiratory domain score of the CFQ-R through 8 weeks of treatment) and by individual mutation (absolute change in sweat chloride and in percent predicted FEV₁ at week 8) is shown in Table 8. Based on clinical (percent predicted FEV₁) and pharmacodynamic (sweat chloride) responses to ivacaftor, efficacy in patients with the G970R mutation could not be established.

Table 8: Effect of ivacaftor for efficacy variables in the overall population and for specific CFTR mutations

Absolute change in percent	BMI	CFQ-R respiratory domain		
predicted FEV ₁	(kg/m^2)	score (points)		
Through week 8	At week 8	Through week 8		
All patients $(N = 39)$				
Results shown as mean (95% CI) change from baseline ivacaftor vs. placebo-treated patients:				
10.7 (7.3, 14.1)				
Patients grouped under mutation types (n)				

Results shown as mean (minimum, maximum) change from baseline for ivacaftor-treated patients at week 8*:

Mutation (n)	Absolute change in sweat chloride (mmol/L)	Absolute change in percent predicted FEV ₁ (percentage points)
	At week 8	At week 8
G1244E (5)	-55 (-75, -34)	8 (-1, 18)
G1349D (2)	-80 (-82, -79)	20 (3, 36)
G178R(5)	-53 (-65, -35)	8 (-1, 18)
G551S(2)	-68 [†]	3^{\dagger}
$G970R^{\#}(4)$	-6 (-16, -2)	3 (-1, 5)
S1251N (8)	-54 (-84, -7)	9 (-20, 21)
S1255P (2)	-78 (-82, -74)	3 (-1, 8)
S549N (6)	-74 (-93, -53)	11 (-2, 20)
S549R (4)	-61 ^{††} (-71, -54)	5 (-3, 13)

Statistical testing was not performed due to small numbers for individual mutations.

In part 2 of study 770-111, the mean (SD) absolute change in percent predicted FEV₁ following 16 weeks (patients randomised to the ivacaftor/placebo treatment sequence in part 1) of continuous ivacaftor treatment was 10.4% (13.2%). At the follow-up visit, 4 weeks after ivacaftor dosing had ended, the mean (SD) absolute change in percent predicted FEV₁ from part 2 week 16 was -5.9% (9.4%). For patients randomised to the placebo/ivacaftor treatment sequence in part 1 there was a

Reflects results from the one patient with the G551S mutation with data at the 8-week time point.

n = 3 for the analysis of absolute change in sweat chloride.

Causes a splicing defect resulting in little-to-no CFTR protein at the cell surface.

further mean (SD) change of 3.3% (9.3%) in percent predicted FEV₁ after the additional 16 weeks of treatment with ivacaftor. At the follow-up visit, 4 weeks after ivacaftor dosing had ended, the mean (SD) absolute change in percent predicted FEV₁ from part 2 week 16 was -7.4% (5.5%).

Study 770-104: study in patients with CF with the F508del mutation in the CFTR gene

Study 770-104 (part A) was a 16-week, 4:1 randomised, double-blind, placebo-controlled, parallel-group phase 2 study of ivacaftor (150 mg every 12 hours) in 140 patients with CF aged 12 years and older who were homozygous for the F508del mutation in the CFTR gene and who had $FEV_1 \ge 40\%$ predicted.

The mean absolute change from baseline through week 16 in percent predicted FEV₁ (primary efficacy endpoint) was 1.5 percentage points in the ivacaftor group and -0.2 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 1.7 percentage points (95% CI: -0.6, 4.1); this difference was not statistically significant (P = 0.15).

Study 770-105: open-label extension study

In study 770-105 patients who completed treatment in studies 770-102 and 770-103 with placebo were switched to ivacaftor while patients on ivacaftor continued to receive it for a minimum of 96 weeks, i.e., the length of treatment with ivacaftor was at least 96 weeks for patients in the placebo/ivacaftor group and at least 144 weeks for patients in the ivacaftor/ivacaftor group.

One hundred and forty-four (144) patients from study 770-102 were rolled over in study 770-105, 67 in the placebo/ivacaftor group and 77 in the ivacaftor/ivacaftor group. Forty-eight (48) patients from study 770-103 were rolled over in study 770-105, 22 in the placebo/ivacaftor group and 26 in the ivacaftor/ivacaftor group.

Table 9 shows the results of the mean (SD) absolute change in percent predicted FEV_1 for both groups of patients. For patients in the placebo/ivacaftor group baseline percent predicted FEV_1 is that of study 770-105 while for patients in the ivacaftor/ivacaftor group the baseline value is that of studies 770-102 and 770-103.

Table 9: Effect of ivacaftor on percent predicted FEV₁ in study 770-105

Original study and treatment group	Duration of ivacaftor treatment (weeks)	Absolute change from baseline in percent predicted FEV ₁ (percentage points)		
		N	Mean (SD)	
Study 770-102				
Ivacaftor	48*	77	9.4 (8.3)	
	144	72	9.4 (10.8)	
Placebo	0*	67	-1.2 (7.8) [†]	
	96	55	9.5 (11.2)	
Study 770-103				
Ivacaftor	48*	26	10.2 (15.7)	
	144	25	10.3 (12.4)	
Placebo	0*	22	-0.6 (10.1) [†]	
	96	21	10.5 (11.5)	

Treatment occurred during blinded, controlled, 48-week phase 3 study.

When the mean (SD) absolute change in percent predicted FEV_1 is compared from study 770-105 baseline for patients in the ivacaftor/ivacaftor group (n = 72) who rolled over from study 770-102, the mean (SD) absolute change in percent predicted FEV_1 was 0.0% (9.05), while for patients in the ivacaftor/ivacaftor group (n = 25) who rolled over from study 770-103 this figure was 0.6% (9.1). This shows that patients in the ivacaftor/ivacaftor group maintained the improvement seen at week 48 of the

Change from prior study baseline after 48 weeks of placebo treatment.

initial study (day 0 through week 48) in percent predicted FEV₁ through week 144. There were no additional improvements in study 770-105 (week 48 through week 144).

For patients in the placebo/ivacaftor group from study 770-102, the annualised rate of pulmonary exacerbations was higher in the initial study when patients were on placebo (1.34 events/year) than during the subsequent study 770-105 when patients rolled over to ivacaftor (0.48 events/year across day 1 to week 48, and 0.67 events/year across weeks 48 to 96). For patients in the ivacaftor/ivacaftor group from study 770-102, the annualised rate of pulmonary exacerbations was 0.57 events/year across day 1 to week 48 when patients were on ivacaftor. When they rolled over into study 770-105, the rate of annualised pulmonary exacerbations was 0.91 events/year across day 1 to week 48 and 0.77 events/year across weeks 48 to 96.

For patients who rolled over from study 770-103 the number of events was overall low.

Study 770-110: study in patients with CF with an R117H mutation in the CFTR gene

Study 770-110 evaluated 69 patients who were 6 years of age or older; 53 (76.8%) patients had the *F508del* mutation in the second allele. The confirmed *R117H* poly-T variant was *5T* in 38 patients and 7T in 16 patients. At baseline, mean predicted FEV₁ was 73% (range: 32.5% to 105.5%) and mean age was 31 years (range: 6 to 68 years). The mean absolute change from baseline through week 24 in percent predicted FEV₁ (primary efficacy endpoint) was 2.57 percentage points in the ivacaftor group and 0.46 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 2.1 percentage points (95% CI: -1.1, 5.4).

A pre-planned subgroup analysis was conducted in patients aged 18 years and older (26 patients on placebo and 24 patients on ivacaftor). Treatment with ivacaftor resulted in a mean absolute change in percent predicted FEV_1 through week 24 of 4.5 percentage points in the ivacaftor group versus -0.46 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 5.0 percentage points (95% CI: 1.1, 8.8).

In a subgroup analysis in patients with a confirmed R117H-5T genetic variant, the difference in the mean absolute change from baseline through week 24 in percent predicted FEV₁ between ivacaftor and placebo was 5.3% (95% CI: 1.3, 9.3). In patients with a confirmed R117H-7T genetic variant, the treatment difference between ivacaftor and placebo was 0.2% (95% CI: -8.1, 8.5).

For secondary efficacy variables, no treatment differences were observed for ivacaftor versus placebo in absolute change from baseline in BMI at week 24 or time to first pulmonary exacerbation. Treatment differences were observed in absolute change in CFQ-R respiratory domain score through week 24 (treatment difference of ivacaftor versus placebo was 8.4 [95% CI: 2.2, 14.6] points) and for the mean change from baseline in sweat chloride (see Pharmacodynamic effects).

Ivacaftor in a combination regimen with tezacaftor/ivacaftor or with ivacaftor/tezacaftor/elexacaftor

The efficacy and safety of ivacaftor in a combination regimen with tezacaftor/ivacaftor in patients with CF aged 12 years and older was assessed in two clinical studies; a 24-week, randomised, double-blind, placebo-controlled study with 504 patients who were homozygous for the *F508del* mutation (study 661-106); and a randomised, double-blind, placebo-controlled and ivacaftor-controlled, 2 period, 3 treatment, 8-week crossover study with 244 patients who were heterozygous for the *F508del* mutation and a second mutation associated with residual CFTR activity (study 661-108). The long-term safety and efficacy of the combination regimen was also assessed in both patient populations in a 96-week open-label, rollover, long-term extension study (study 661-110). Refer to the Summary of Product Characteristics of tezacaftor/ivacaftor for additional data.

The efficacy and safety of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor tablets in patients aged 12 years and older was demonstrated in three, phase 3, randomised, double-blind, placebo-controlled studies (patients were heterozygous for the F508del mutation and a mutation with minimal function on the second allele, n = 403) and active-controlled (patients were homozygous

for the F508del mutation, n = 107, or heterozygous for the F508del mutation and a gating or residual CFTR activity mutation on the second allele, n = 258) studies of 24 (study 445-102), 4 (study 445-103), and 8 weeks (study 445-104) of duration respectively. Patients from all studies were eligible to enter open-label, rollover, long-term extension studies (study 445-105 or study 445-110).

The efficacy and safety of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor tablets in patients aged 6 years and older with non-F508del, ivacaftor/tezacaftor/elexacaftor-responsive *CFTR* mutations was demonstrated in a phase 3, randomised, double-blind, placebo-controlled study (n = 307) and an observational retrospective study (CFD-016; n = 422) of Real-World clinical outcomes.

Refer to the Summary of Product Characteristics of ivacaftor/tezacaftor/elexacaftor for additional data.

Paediatric population

Ivacaftor in a combination regimen with tezacaftor/ivacaftor

The efficacy and safety in patients aged 6 to less than 12 years (mean age 8.6 years) were assessed in an 8-week, double-blind, phase 3 trial (study 661-115) with 67 patients who were randomised 4:1 to either ivacaftor in a combination regimen with tezacaftor/ivacaftor or a blinding group. Forty-two patients were homozygous for the *F508del* mutation (F/F) and 12 were heterozygous for the *F508del* mutation and a second mutation associated with residual CFTR activity (F/RF). Patients were eligible to enter an open-label, rollover, 96-week study (study 661-116 part A). Refer to the Summary of Product Characteristics of tezacaftor/ivacaftor for additional data.

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

The pharmacokinetics and safety in patients aged 6 to less than 12 years (n = 66) and in those from 2 to less than 6 years (n = 75) who have at least one F508del mutation were assessed in two 24-week open-label studies (study 445-106 and study 445-116). Refer to the Summary of Product Characteristics of ivacaftor/tezacaftor/elexacaftor for additional data.

The European Medicines Agency has deferred the obligation to submit the results of studies with Kalydeco in one or more subsets of the paediatric population in cystic fibrosis (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

The pharmacokinetics of ivacaftor are similar between healthy adult volunteers and patients with CF.

After oral administration of a single 150 mg dose to healthy volunteers in a fed state, the mean (\pm SD) for AUC and C_{max} were 10.60 (5.26) µg·h/mL and 0.768 (0.233) µg/mL, respectively. After every 12-hour dosing, steady-state plasma concentrations of ivacaftor were reached by days 3 to 5, with an accumulation ratio ranging from 2.2 to 2.9.

Absorption

Following multiple oral dose administrations of ivacaftor, the exposure of ivacaftor generally increased with dose from 25 mg every 12 hours to 450 mg every 12 hours. When given with fatcontaining food, the exposure of ivacaftor increased approximately 2.5- to 4-fold. When coadministered with tezacaftor and elexacaftor, the increase in AUC was similar (approximately 3-fold and 2.5- to 4-fold respectively). Therefore, ivacaftor, administered as monotherapy or in a combination regimen with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor, should be administered with fat-containing food. The median (range) t_{max} is approximately 4.0 (3.0; 6.0) hours in the fed state.

Ivacaftor granules (2×75 mg sachets) had similar bioavailability as the 150 mg tablet when given with fat-containing food to healthy adult subjects. The geometric least squares mean ratio (90% CI) for

the granules relative to tablets was 0.951 (0.839, 1.08) for $AUC_{0-\infty}$ and 0.918 (0.750, 1.12) for C_{max} . The effect of food on ivacaftor absorption is similar for both formulations, i.e., tablets and granules.

Distribution

Ivacaftor is approximately 99% bound to plasma proteins, primarily to alpha 1-acid glycoprotein and albumin. Ivacaftor does not bind to human red blood cells. After oral administration of ivacaftor 150 mg every 12 hours for 7 days in healthy volunteers in a fed state, the mean (\pm SD) apparent volume of distribution was 353 L (122).

Biotransformation

Ivacaftor is extensively metabolised in humans. *In vitro* and *in vivo* data indicate that ivacaftor is primarily metabolised by CYP3A. M1 and M6 are the two major metabolites of ivacaftor in humans. M1 has approximately one-sixth the potency of ivacaftor and is considered pharmacologically active. M6 has less than one-fiftieth the potency of ivacaftor and is not considered pharmacologically active.

The effect of the CYP3A4*22 heterozygous genotype on ivacaftor, tezacaftor, and elexacaftor exposure is consistent with the effect of co-administration of a weak CYP3A4 inhibitor, which is not clinically relevant. No dose-adjustment of ivacaftor, tezacaftor, or elexacaftor is considered necessary. The effect in CYP3A4*22 homozygous genotype patients is expected to be stronger. However, no data are available for such patients.

Elimination

Following oral administration in healthy volunteers, the majority of ivacaftor (87.8%) was eliminated in the faeces after metabolic conversion. The major metabolites M1 and M6 accounted for approximately 65% of the total dose eliminated with 22% as M1 and 43% as M6. There was negligible urinary excretion of ivacaftor as unchanged parent. The apparent terminal half-life was approximately 12 hours following a single dose in the fed state. The apparent clearance (CL/F) of ivacaftor was similar for healthy subjects and patients with CF. The mean (\pm SD) CL/F for a single 150 mg dose was 17.3 (8.4) L/hr in healthy subjects.

Linearity/non-linearity

The pharmacokinetics of ivacaftor are generally linear with respect to time or dose ranging from 25 mg to 250 mg.

Special populations

Hepatic impairment

Following a single dose of 150 mg of ivacaftor, adult subjects with moderately impaired hepatic function (Child-Pugh Class B, score 7 to 9) had similar ivacaftor C_{max} (mean [\pm SD] of 0.735 [0.331] μ g/mL) but an approximately 2-fold increase in ivacaftor $AUC_{0-\infty}$ (mean [\pm SD] of 16.80 [6.14] μ g·h/mL) compared with healthy subjects matched for demographics. Simulations for predicting the steady-state exposure of ivacaftor showed that by reducing the dose from 150 mg q12h to 150 mg once daily, adults with moderate hepatic impairment would have comparable steady-state C_{min} values as those obtained with a dose of 150 mg q12h in adults without hepatic impairment.

In subjects with moderately impaired hepatic function (Child-Pugh Class B, score 7 to 9), ivacaftor AUC increased approximately by 50% following multiple doses for 10 days of either tezacaftor and ivacaftor or of ivacaftor, tezacaftor and elexacaftor.

The impact of severe hepatic impairment (Child-Pugh Class C, score 10 to15) on the pharmacokinetics of ivacaftor has not been studied. The magnitude of increase in exposure in these patients is unknown but is expected to be higher than that observed in patients with moderate hepatic impairment.

For guidance on appropriate use and dose modification see Table 3 in section 4.2.

Renal impairment

Pharmacokinetic studies have not been performed with ivacaftor in patients with renal impairment. In a human pharmacokinetic study with ivacaftor monotherapy, there was minimal elimination of ivacaftor and its metabolites in urine (only 6.6% of total radioactivity was recovered in the urine). There was negligible urinary excretion of ivacaftor as unchanged parent (less than 0.01% following a single oral dose of 500 mg).

No dose adjustments are recommended for mild and moderate renal impairment. Caution is recommended when administering ivacaftor to patients with severe renal impairment (creatinine clearance less than or equal to 30 mL/min) or end-stage renal disease (see sections 4.2 and 4.4).

Race

Race had no clinically meaningful effect on the PK of ivacaftor in white (n = 379) and non-white (n = 29) patients based on a population PK analysis.

Gender

The pharmacokinetic parameters of ivacaftor are similar in males and females.

Elderly

Clinical studies of ivacaftor did not include sufficient numbers of patients aged 65 years and older to determine whether pharmacokinetic parameters are similar or not to those in younger adults.

The pharmacokinetic parameters of ivacaftor in combination with tezacaftor in the elderly patients (65-72 years) are comparable to those in younger adults.

Paediatric population

Predicted ivacaftor exposure based on observed ivacaftor concentrations in phase 2 and 3 studies as determined using compartmental analysis is presented by age group in Table 10.

Table 10: Mean (SD) ivacaftor exposure by age group

Age group	Dose	C _{min, ss} (μg/mL)	AUC _{0-12h} , ss (μg·h/mL)
1 month to less than 2 months (≥ 3 kg)*	13.4 mg q24h	0.300 (0.221) [†]	5.84 (2.98) [†]
2 months to less than 4 months (≥ 3 kg)*	13.4 mg q12h	0.406 (0.266) [†]	6.45 (3.43) [†]
4 months to less than 6 months (≥ 5 kg)*	25 mg q12h	0.371 (0.183)	6.48 (2.52)
6 months to less than 12 months (\geq 5 kg to $<$ 7 kg) [‡]	25 mg q12h	0.336	5.41
6 months to less than 12 months (7 kg to < 14 kg)	50 mg q12h	0.508 (0.252)	9.14 (4.20)
12 months to less than 24 months (7 kg to < 14 kg)	50 mg q12h	0.440 (0.212)	9.05 (3.05)

Age group	Dose	C _{min, ss} (μg/mL)	AUC _{0-12h} , ss (μg·h/mL)
12 months to less than 24 months (≥ 14 kg to < 25 kg)	75 mg q12h	0.451 (0.125)	9.60 (1.80)
2- to 5-year-olds (< 14 kg)	50 mg q12h	0.577 (0.317)	10.50 (4.26)
2- to 5-year-olds (≥ 14 kg to < 25 kg)	75 mg q12h	0.629 (0.296)	11.30 (3.82)
6- to 11-year-olds [§] (≥ 14 kg to < 25 kg)	75 mg q12h	0.641 (0.329)	10.76 (4.47)
6- to 11-year-olds [§] (≥ 25 kg)	150 mg q12h	0.958 (0.546)	15.30 (7.34)
12- to 17-year-olds	150 mg q12h	0.564 (0.242)	9.24 (3.42)
Adults (≥ 18 years old)	150 mg q12h	0.701 (0.317)	10.70 (4.10)

- * Patients 1 month to less than 6 months of age were ≥37 weeks gestational age.
- † Exposures for 1 month to less than 4 months of age are predictions based on simulations from the physiologically based PK model incorporating data from the given age group.
- [‡] Values based on data from a single patient; standard deviation not reported.
- Exposures in 6- to 11-year-olds are predictions based on simulations from the population PK model using data obtained for this age group.

5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, and carcinogenic potential.

Pregnancy and fertility

Ivacaftor was associated with slight decreases of the seminal vesicle weights, a decrease of overall fertility index and number of pregnancies in females mated with treated males and significant reductions in number of corpora lutea and implantation sites with subsequent reductions in the average litter size and average number of viable embryos per litter in treated females. The No-Observed-Adverse-Effect-Level (NOAEL) for fertility findings provides an exposure level of approximately 4 times the systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy in adult humans at the maximum recommended human dose (MRHD). Placental transfer of ivacaftor was observed in pregnant rats and rabbits.

Peri- and post-natal development

Ivacaftor decreased survival and lactation indices and caused a reduction in pup body weights. The NOAEL for viability and growth in the offspring provides an exposure level of approximately 3 times the systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy in adult humans at the MRHD.

Juvenile animal studies

Findings of cataracts were observed in juvenile rats dosed from postnatal day 7 through 35 at ivacaftor exposure levels of 0.22 times the MRHD based on systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy. This finding has not been observed in foetuses derived from rat dams treated with ivacaftor on gestation days 7 to 17, in rat pups exposed to ivacaftor through milk ingestion up to postnatal day 20, in 7-week old rats, nor in 3.5 to 5-month old dogs treated with ivacaftor. The potential relevance of these findings in humans is unknown.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Tablet core

Cellulose, microcrystalline Lactose monohydrate Hypromellose acetate succinate Croscarmellose sodium Sodium laurilsulfate (E487) Silica, colloidal anhydrous Magnesium stearate

Tablet film coat

Polyvinyl alcohol Titanium dioxide (E171) Macrogol (PEG 3350) Talc Indigo carmine aluminium lake (E132) Carnauba wax

Printing ink

Shellac Iron oxide black (E172) Propylene glycol (E1520) Ammonia solution, concentrated

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

4 years.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

Thermoform (PolyChloroTriFluoroEthylene [PCTFE]/foil) blister or a High-Density PolyEthylene (HDPE) bottle with a polypropylene child-resistant closure, foil-lined induction seal and molecular sieve desiccant.

Kalydeco 75 mg film-coated tablets

The following pack sizes are available:

• Blister card pack containing 28 film-coated tablets

Kalydeco 150 mg film-coated tablets

The following pack sizes are available:

- Blister card pack containing 28 film-coated tablets
- Blister pack containing 56 film-coated tablets
- Bottle containing 56 film-coated tablets

Not all pack sizes may be marketed.

6.6 Special precautions for disposal and other handling

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

7. MARKETING AUTHORISATION HOLDER

Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/12/782/001 EU/1/12/782/002 EU/1/12/782/005 EU/1/12/782/007

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 23 July 2012 Date of latest renewal: 29 April 2022

10. DATE OF REVISION OF THE TEXT

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

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 13.4 mg granules in sachet

Kalydeco 25 mg granules in sachet

Kalydeco 50 mg granules in sachet

Kalydeco 59.5 mg granules in sachet

Kalydeco 75 mg granules in sachet

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Kalydeco 13.4 mg granules in sachet

Each sachet contains 13.4 mg of ivacaftor.

Excipient with known effect

Each sachet contains 19.7 mg of lactose monohydrate.

Kalydeco 25 mg granules in sachet

Each sachet contains 25 mg of ivacaftor.

Excipient with known effect

Each sachet contains 36.6 mg of lactose monohydrate.

Kalydeco 50 mg granules in sachet

Each sachet contains 50 mg of ivacaftor.

Excipient with known effect

Each sachet contains 73.2 mg of lactose monohydrate.

Kalydeco 59.5 mg granules in sachet

Each sachet contains 59.5 mg of ivacaftor.

Excipient with known effect

Each sachet contains 87.3 mg of lactose monohydrate.

Kalydeco 75 mg granules in sachet

Each sachet contains 75 mg of ivacaftor.

Excipient with known effect

Each sachet contains 109.8 mg of lactose monohydrate.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Granule in sachet (granule)

White to off-white granules approximately 2 mm in diameter.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Kalydeco granules are indicated:

- As monotherapy for the treatment of infants aged at least 1 month, toddlers and children weighing 3 kg to less than 25 kg with cystic fibrosis (CF) who have an R117H CFTR mutation or one of the following gating (Class III) mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene: G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N or S549R (see sections 4.4 and 5.1).
- In a combination regimen with ivacaftor/tezacaftor/elexacaftor for the treatment of cystic fibrosis (CF) in paediatric patients aged 2 to less than 6 years who have at least one non-Class I mutation in the *CFTR* gene (see sections 4.2 and 5.1).

4.2 Posology and method of administration

Kalydeco should only be prescribed by physicians with experience in the treatment of cystic fibrosis. If the patient's genotype is unknown, an accurate and validated genotyping method should be performed before starting treatment to confirm the presence of an indicated mutation in at least one allele of the *CFTR* gene (see section 4.1). The phase of the poly-T variant identified with the *R117H* mutation should be determined in accordance with local clinical recommendation.

Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor

There are a limited number of patients who harbour mutations not listed in Table 6 that may be responsive to ivacaftor/tezacaftor/elexacaftor (IVA/TEZ/ELX). In these cases, ivacaftor (IVA) in combination with IVA/TEZ/ELX can be considered when the physician deems the potential benefits outweigh the potential risks and under close medical supervision. This excludes patients with two Class I (null) mutations (mutations that are known not to produce CFTR protein) as they are not expected to respond to modulator therapy (see sections 4.1, 4.4 and 5.1).

Posology

Dosing recommendations are shown in Table 1.

Table 1: Dosing recommendations

Age	Weight	Morning dose	Evening dose	
Ivacaftor as mon	Ivacaftor as monotherapy			
1 month to less	> 2 1ra	One sachet of ivacaftor	No avanina daga	
than 2 months	\geq 3 kg	13.4 mg granules	No evening dose	
2 months to less	> 2 1ra	One sachet of ivacaftor	One sachet of ivacaftor	
than 4 months	\geq 3 kg	13.4 mg granules	13.4 mg granules	
4 months to less	> 5 1ra	One sachet of ivacaftor	One sachet of ivacaftor	
than 6 months	\geq 5 kg	25 mg granules	25 mg granules	

\geq 5 kg to \leq 7 kg		One sachet of ivacaftor	One sachet of ivacaftor	
	≥ 3 kg to < 7 kg	25 mg granules	25 mg granules	
<i>c</i> 1 1	\geq 7 kg to < 14 kg	One sachet of ivacaftor	One sachet of ivacaftor	
6 months and	2 / kg to < 14 kg	50 mg granules	50 mg granules	
older	\geq 14 kg to \leq 25 kg	One sachet of ivacaftor	One sachet of ivacaftor	
	\geq 14 kg to \sim 23 kg	75 mg granules	75 mg granules	
	≥ 25 kg	See Kalydeco tablets SmPC for further details.		
Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor				
		One sachet of ivacaftor		
	10 lea to < 14 lea	60 mg/tezacaftor	One sachet of ivacaftor	
	10 kg to < 14 kg	40 mg/elexacaftor 80 mg	59.5 mg granules	
2 years to less		granules		
than 6 years		One sachet of ivacaftor		
	> 1.4 lrg	75 mg/tezacaftor	One sachet of ivacaftor	
	≥ 14 kg	50 mg/elexacaftor 100 mg	75 mg granules	
		granules		

The morning and evening dose should be taken approximately 12 hours apart with fat-containing food (see Method of administration).

Missed dose

If 6 hours or less have passed since the missed morning or evening dose, the patient should be advised to take it as soon as possible and then take the next dose at the regularly scheduled time. If more than 6 hours have passed since the time the dose is usually taken, the patient should be advised to wait until the next scheduled dose.

Patients receiving Kalydeco in a combination regimen should be advised not to take more than one dose of either medicinal product at the same time.

Concomitant use of CYP3A inhibitors

During concomitant administration with moderate or strong inhibitors of CYP3A, the ivacaftor dose should be adjusted as detailed in Table 2 (see sections 4.4 and 4.5).

Table 2: Dosing recommendations for concomitant use with moderate or strong CYP3A inhibitors

Age/weight	Moderate CYP3A inhibitors	Strong CYP3A inhibitors		
Ivacaftor as monotherapy				
1 month to	Use is not recommended.	Use is not recommended.		
less than				
6 months				
6 months	One sachet of ivacaftor 25 mg granules	One sachet of ivacaftor 25 mg granules		
and older,	once daily.	twice a week.		
\geq 5 kg to				
< 7 kg	No evening ivacaftor dose.	No evening ivacaftor dose.		
6 months	One sachet of ivacaftor 50 mg granules	One sachet of ivacaftor 50 mg granules		
and older,	once daily.	twice a week.		
\geq 7 kg to				
< 14 kg	No evening ivacaftor dose.	No evening ivacaftor dose.		
6 months	One sachet of ivacaftor 75 mg granules	One sachet of ivacaftor 75 mg granules		
and older,	once daily.	twice a week.		
\geq 14 kg to				
< 25 kg	No evening ivacaftor dose.	No evening ivacaftor dose.		

a combination regimen with ivacaftor/teza	
a compination regimen with tvacattor/teza	caftor/elexacaftor
 Alternate each day: One morning sachet of ivacaftor 60 mg/tezacaftor 40 mg/elexacaftor 80 mg granules on the first day One morning sachet of ivacaftor 59.5 mg granules on the next day 	One sachet of ivacaftor 60 mg/tezacaftor 40 mg/elexacaftor 80 mg granules twice a week, approximately 3 to 4 days apart. No evening ivacaftor dose.
No evening ivacaftor dose.	
 Alternate each day: One morning sachet of ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg granules on the first day One morning sachet of ivacaftor 75 mg granules on the next day 	One sachet of ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg granules twice a week, approximately 3 to 4 days apart. No evening ivacaftor dose.
	 One morning sachet of ivacaftor 60 mg/tezacaftor 40 mg/elexacaftor 80 mg granules on the first day One morning sachet of ivacaftor 59.5 mg granules on the next day No evening ivacaftor dose. Alternate each day: One morning sachet of ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg granules on the first day One morning sachet of ivacaftor

Special populations

Renal impairment

No dose adjustment is necessary for patients with mild to moderate renal impairment. Caution is recommended in patients with severe renal impairment (creatinine clearance less than or equal to 30 mL/min) or end-stage renal disease (see sections 4.4 and 5.2).

Hepatic impairment

No dose adjustment is necessary for patients aged 6 months and older with mild hepatic impairment (Child-Pugh Class A). Treatment with ivacaftor is not recommended in patients aged 1 month to less than 6 months with any level of hepatic impairment.

In patients with moderate hepatic impairment (Child-Pugh Class B), or severe hepatic impairment (Child-Pugh Class C), the ivacaftor dose should be adjusted as detailed in Table 3 (see sections 4.4, 4.8, and 5.2).

Table 3: Dosing recommendations for patients with moderate or severe hepatic impairment

Age/weight	Moderate (Child-Pugh Class B)	Severe (Child-Pugh Class C)		
Ivacaftor as	Ivacaftor as monotherapy			
1 month to	Use is not recommended.	Use is not recommended.		
less than				
6 months				
6 months	One sachet of ivacaftor 25 mg granules	Use is not recommended, unless the		
and older,	once daily.	benefits are expected to outweigh the risks.		
\geq 5 kg to				
< 7 kg	No evening ivacaftor dose.	If used, one sachet of ivacaftor 25 mg		
		granules every other day according to		
		clinical response and tolerability.		
		No evening ivacaftor dose.		

6 months	One sachet of ivacaftor 50 mg granules	Use is not recommended, unless the
and older,	once daily.	benefits are expected to outweigh the risks.
\geq 7 kg to		
< 14 kg	No evening ivacaftor dose.	If used, one sachet of ivacaftor 50 mg
11.118	The Granding True Miles Constitution and	granules every other day according to
		clinical response and tolerability.
		No evening ivacaftor dose.
6 months	One sachet of ivacaftor 75 mg granules	Use is not recommended, unless the
and older,	once daily.	benefits are expected to outweigh the risks.
\geq 14 kg to		
< 25 kg	No evening ivacaftor dose.	If used, one sachet of ivacaftor 75 mg
		granules every other day according to
		clinical response and tolerability.
T 0: 1		No evening ivacaftor dose.
	a combination regimen with ivacaftor/tezac	
2 years to	Use is not recommended, unless the	Should not be used.
less than	benefits are expected to outweigh the risks.	
6 years,	TO 1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.	
10 kg to	If used, the dose should be adjusted as	
< 14 kg	follows:	
	D 12 1 6 6	
	• Days 1-3: one sachet of ivacaftor	
	60 mg/tezacaftor 40 mg/elexacaftor	
	80 mg granules each day	
	• Day 4: no dose	
	• Days 5-6: one sachet of ivacaftor	
	60 mg/tezacaftor 40 mg/elexacaftor	
	80 mg granules each day	
	• Day 7: no dose	
	Repeat above dosing schedule each week.	
	Repeat above dosing schedule each week.	
	No evening ivacaftor dose.	
2 years to	Use is not recommended, unless the	Should not be used.
less than	benefits are expected to outweigh the risks.	
6 years,	70 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
≥ 14 kg	If used, the dose should be adjusted as	
	follows:	
	David 2	
	• Days 1-3: one sachet of ivacaftor	
	75 mg/tezacaftor 50 mg/elexacaftor	
	100 mg granules each day	
	• Day 4: no dose	
	• Days 5-6: one sachet of ivacaftor	
	75 mg/tezacaftor 50 mg/elexacaftor	
	100 mg granules each day	
	• Day 7: no dose	
	Repeat above dosing schedule each week.	
	No evening ivacaftor dose.	

Paediatric population

The safety and efficacy of ivacaftor as monotherapy have not been established in children less than 1 month of age or in children less than 6 months of age born prematurely (less than 37 weeks of gestational age), neither in combination with ivacaftor/tezacaftor/elexacaftor in children less than 2 years of age. No data are available.

Limited data are available in patients less than 6 years of age with an *R117H* mutation in the *CFTR* gene. Available data in patients aged 6 years and older are described in sections 4.8, 5.1, and 5.2.

Method of administration

For oral use.

Each sachet is for single use only.

Each sachet of granules should be mixed with 5 mL of age-appropriate soft food or liquid and completely and immediately consumed. Food or liquid should be at room temperature or below. If not immediately consumed, the mixture has been shown to be stable for one hour and therefore should be ingested during this period. A fat-containing meal or snack should be consumed just before or just after dosing.

Food or drink containing grapefruit should be avoided during treatment (see section 4.5).

4.3 Contraindications

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

4.4 Special warnings and precautions for use

Only patients with CF who had a *G551D*, *G1244E*, *G1349D*, *G178R*, *G551S*, *S1251N*, *S1255P*, *S549N* or *S549R* gating (Class III) or *G970R* mutation in at least one allele of the *CFTR* gene were included in studies 770-102, 770-103, 770-111 and 770-108 (see section 5.1).

Less evidence of a positive effect of ivacaftor has been shown for patients with an *R117H-7T* mutation associated with less severe disease in study 770-110 (see section 5.1).

In study 770-111, four patients with the G970R mutation were included. In three of four patients the change in the sweat chloride test was < 5 mmol/L and this group did not demonstrate a clinically relevant improvement in FEV₁ after 8 weeks of treatment. Clinical efficacy in patients with the G970R mutation of the CFTR gene could not be established (see section 5.1).

Efficacy results from a phase 2 study in patients with CF who are homozygous for the *F508del* mutation in the *CFTR* gene showed no statistically significant difference in FEV₁ over 16 weeks of ivacaftor treatment compared to placebo (see section 5.1). Therefore, use of ivacaftor as monotherapy in these patients is not recommended.

Elevated transaminases and hepatic injury

In a patient with cirrhosis and portal hypertension, liver failure leading to transplantation has been reported while receiving ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor. This medicinal product should be used with caution in patients with pre-existing advanced liver disease (e.g., cirrhosis, portal hypertension) and only if the benefits are expected to outweigh the risks. If used

in these patients, they should be closely monitored after the initiation of treatment (see sections 4.2, 4.8, and 5.2).

Moderate transaminase (alanine transaminase [ALT] or aspartate transaminase [AST]) elevations are common in subjects with CF. Transaminase elevations have been observed in some patients treated with ivacaftor as monotherapy and in combination regimens with ivacaftor/tezacaftor/elexacaftor. In patients taking ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor, these elevations have sometimes been associated with concomitant elevations in total bilirubin. Therefore, assessments of transaminases (ALT and AST) and total bilirubin are recommended for all patients prior to initiating ivacaftor, every 3 months during the first year of treatment and annually thereafter. For all patients with a history of liver disease or transaminase elevations, more frequent monitoring of liver function tests should be considered. In the event of significant elevations of transaminases (e.g., patients with ALT or AST > 5 × the upper limit of normal (ULN), or ALT or AST > 3 × ULN with bilirubin > 2 × ULN), dosing should be interrupted, and laboratory tests closely followed until the abnormalities resolve. Following resolution of transaminase elevations, the benefits and risks of resuming treatment should be considered (see sections 4.2, 4.8, and 5.2).

Hepatic impairment

Use of ivacaftor as monotherapy is not recommended in patients aged 1 month to less than 6 months of age with any level of hepatic impairment. Use of ivacaftor as monotherapy is not recommended in patients aged 6 months and older weighing less than 25 kg with severe hepatic impairment unless the benefits are expected to outweigh the risks. Patients with severe hepatic impairment should not be treated with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor (see Table 3 in section 4.2, and sections 4.8 and 5.2).

For patients with moderate hepatic impairment, use of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor is not recommended. Treatment should only be considered when there is a clear medical need and the benefits are expected to outweigh the risks. If used, it should be used with caution at a reduced dose (see Table 3 in section 4.2, and sections 4.8 and 5.2).

Depression

Depression (including suicidal ideation and suicide attempt) has been reported in patients while receiving ivacaftor, mainly in a combination regimen with ivacaftor/tezacaftor/elexacaftor, usually occurring within three months of treatment initiation and in patients with a history of psychiatric disorders. In some cases, symptom improvement was reported after dose reduction or treatment discontinuation. Patients (and caregivers) should be alerted about the need to monitor for depressed mood, suicidal thoughts, or unusual changes in behaviour and to seek medical advice immediately if these symptoms present.

Renal impairment

Caution is recommended while using ivacaftor in patients with severe renal impairment or end-stage renal disease (see sections 4.2 and 5.2).

Mutations unlikely to respond to modulator therapy

Patients with a genotype consisting of two *CFTR* mutations that are known not to produce CFTR protein (i.e., two Class I mutations) are not expected to respond to CFTR modulator therapy.

Clinical studies comparing ivacaftor/tezacaftor/elexacaftor to tezacaftor/ivacaftor or ivacaftor

No clinical study has been conducted to directly compare ivacaftor/tezacaftor/elexacaftor to tezacaftor/ivacaftor or ivacaftor in patients not harbouring *F508del* variant.

Patients after organ transplantation

Ivacaftor has not been studied in patients with CF who have undergone organ transplantation. Therefore, use in transplanted patients is not recommended. See section 4.5 for interactions with ciclosporin or tacrolimus.

Rash events

The incidence of rash events with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor was higher in females than in males, particularly in females taking hormonal contraceptives. A role for hormonal contraceptives in the occurrence of rash cannot be excluded. For patients taking hormonal contraceptives who develop rash, interrupting treatment with ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor and hormonal contraceptives should be considered. Following the resolution of rash, it should be considered if resuming ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor without hormonal contraceptives is appropriate. If rash does not recur, resumption of hormonal contraceptives can be considered (see section 4.8).

Interactions with medicinal products

CYP3A inducers

Exposure to ivacaftor is significantly decreased by the concomitant use of CYP3A inducers, potentially resulting in the loss of ivacaftor efficacy; therefore, co-administration of ivacaftor with strong CYP3A inducers is not recommended (see section 4.5).

CYP3A inhibitors

Exposure to ivacaftor is increased when co-administered with strong or moderate CYP3A inhibitors. The dose of ivacaftor must be adjusted when used concomitantly with strong or moderate CYP3A inhibitors (see Table 2 in section 4.2 and section 4.5). Treatment with ivacaftor monotherapy is not recommended in patients aged 1 month to less than 6 months taking strong or moderate CYP3A inhibitors.

Paediatric population

Cases of non-congenital lens opacities/cataracts without impact on vision have been reported in paediatric patients treated with ivacaftor and ivacaftor-containing regimens. Although other risk factors were present in some cases (such as corticosteroid use and exposure to radiation), a possible risk attributable to treatment with ivacaftor cannot be excluded. Baseline and follow-up ophthalmological examinations are recommended in paediatric patients initiating ivacaftor treatment (see section 5.3).

Excipients with known effect

Lactose

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

Sodium

This medicinal product contains less than 1 mmol sodium (23 mg) per sachet, that is to say essentially 'sodium-free'.

4.5 Interaction with other medicinal products and other forms of interaction

Ivacaftor is a substrate of CYP3A4 and CYP3A5. It is a weak inhibitor of CYP3A and P-glycoprotein (P-gp) and a potential inhibitor of CYP2C9. *In vitro* studies showed that ivacaftor is not a substrate for P-gp.

Medicinal products affecting the pharmacokinetics of ivacaftor

CYP3A inducers

Co-administration of ivacaftor with rifampicin, a strong CYP3A inducer, decreased ivacaftor exposure (AUC) by 89% and decreased hydroxymethyl ivacaftor (M1) to a lesser extent than ivacaftor. Co-administration of ivacaftor with strong CYP3A inducers, such as rifampicin, rifabutin, phenobarbital, carbamazepine, phenytoin and St. John's wort (*Hypericum perforatum*), is not recommended (see section 4.4).

No dose adjustment is recommended when ivacaftor is used with moderate or weak CYP3A inducers.

CYP3A inhibitors

Ivacaftor is a sensitive CYP3A substrate. Co-administration with ketoconazole, a strong CYP3A inhibitor, increased ivacaftor exposure (measured as area under the curve [AUC]) by 8.5-fold and increased M1 to a lesser extent than ivacaftor. Co-administration with fluconazole, a moderate inhibitor of CYP3A, increased ivacaftor exposure by 3-fold and increased M1 to a lesser extent than ivacaftor. A reduction of the ivacaftor dose is recommended for co-administration with strong CYP3A inhibitors, such as ketoconazole, itraconazole, posaconazole, voriconazole, telithromycin and clarithromycin as well as for co-administration with moderate CYP3A inhibitors, such as fluconazole, erythromycin, and verapamil. Treatment with ivacaftor monotherapy is not recommended when concomitantly used with strong or moderate CYP3A inhibitors in patients aged 1 month to less than 6 months (see Table 2 in section 4.2 and section 4.4).

Co-administration of ivacaftor with grapefruit juice, which contains one or more components that moderately inhibit CYP3A, may increase exposure to ivacaftor. Food or drink containing grapefruit should be avoided during treatment with ivacaftor (see section 4.2).

Potential for ivacaftor to interact with transporters

In vitro studies showed that ivacaftor is not a substrate for OATP1B1 or OATP1B3. Ivacaftor and its metabolites are substrates of BCRP *in vitro*. Due to its high intrinsic permeability and low likelihood of being excreted intact, co-administration of BCRP inhibitors is not expected to alter exposure of ivacaftor and M1-IVA, while any potential changes in M6-IVA exposures are not expected to be clinically relevant.

Ciprofloxacin

Co-administration of ciprofloxacin with ivacaftor did not affect the exposure of ivacaftor. No dose adjustment is required when ivacaftor is co-administered with ciprofloxacin.

Medicinal products affected by ivacaftor

Administration of ivacaftor may increase systemic exposure of medicinal products that are sensitive substrates of CYP2C9, and/or P-gp, and/or CYP3A which may increase or prolong their therapeutic effect and adverse reactions.

CYP2C9 substrates

Ivacaftor may inhibit CYP2C9. Therefore, monitoring of the international normalised ratio (INR) is recommended during co-administration of warfarin with ivacaftor. Other medicinal products for which exposure may be increased include glimepiride and glipizide; these medicinal products should be used with caution.

Digoxin and other P-gp substrates

Co-administration with digoxin, a sensitive P-gp substrate, increased digoxin exposure by 1.3-fold, consistent with weak inhibition of P-gp by ivacaftor. Administration of ivacaftor may increase systemic exposure of medicinal products that are sensitive substrates of P-gp, which may increase or prolong their therapeutic effect and adverse reactions. When used concomitantly with digoxin or other substrates of P-gp with a narrow therapeutic index, such as ciclosporin, everolimus, sirolimus or tacrolimus, caution and appropriate monitoring should be used.

CYP3A substrates

Co-administration with (oral) midazolam, a sensitive CYP3A substrate, increased midazolam exposure 1.5-fold, consistent with weak inhibition of CYP3A by ivacaftor. No dose adjustment of CYP3A substrates, such as midazolam, alprazolam, diazepam or triazolam, is required when these are co-administered with ivacaftor.

Hormonal contraceptives

Ivacaftor has been studied with an oestrogen/progesterone oral contraceptive and was found to have no significant effect on the exposures of the oral contraceptive. Therefore, no dose adjustment of oral contraceptives is necessary.

Paediatric population

Interaction studies have only been performed in adults.

4.6 Fertility, pregnancy and lactation

Pregnancy

There are no or limited amount of data (less than 300 pregnancy outcomes) from the use of ivacaftor in pregnant women. Animals studies do not indicate direct or indirect harmful effects with respect to reproductive toxicity (see section 5.3). As a precautionary measure, it is preferable avoid the use of ivacaftor during pregnancy.

Breast-feeding

Limited data show that ivacaftor is excreted into human milk. A risk to the newborns/infants cannot be excluded. A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from ivacaftor therapy taking into account the benefit of breast-feeding for the child and the benefit of therapy for the woman.

Fertility

There are no data available on the effect of ivacaftor on fertility in humans. Ivacaftor had an effect on fertility in rats (see section 5.3).

4.7 Effects on ability to drive and use machines

Ivacaftor has minor influence on the ability to drive and use machines. Ivacaftor may cause dizziness (see section 4.8) and, therefore, patients experiencing dizziness should be advised not to drive or use machines until symptoms abate.

4.8 Undesirable effects

Summary of the safety profile

The most common adverse reactions experienced by patients aged 6 years and older are headache (23.9%), oropharyngeal pain (22.0%), upper respiratory tract infection (22.0%), nasal congestion (20.2%), abdominal pain (15.6%), nasopharyngitis (14.7%), diarrhoea (12.8%), dizziness (9.2%), rash (12.8%) and bacteria in sputum (12.8%). Transaminase elevations occurred in 12.8% of ivacaftor-treated patients versus 11.5% of placebo-treated patients.

In patients aged 2 to less than 6 years the most common adverse reactions were nasal congestion (26.5%), upper respiratory tract infection (23.5%), transaminase elevations (14.7%), rash (11.8%) and bacteria in sputum (11.8%).

Serious adverse reactions included abdominal pain (0.9%) and transaminase elevations (1.8%) in patients who received ivacaftor, while serious adverse reactions of rash were reported in 1.5% patients aged 12 years and older treated with a combination regimen with ivacaftor/tezacaftor/elexacaftor (see section 4.4).

Tabulated list of adverse reactions

Table 4 reflects the adverse reactions observed with ivacaftor in clinical trials (placebo-controlled and uncontrolled studies) in which the length of exposure to ivacaftor ranged from 16 weeks to 144 weeks. The frequency of adverse reactions is defined as follows: very common ($\geq 1/10$); common ($\geq 1/100$) to < 1/10); uncommon ($\geq 1/1,000$ to < 1/100); rare ($\geq 1/10,000$ to < 1/1,000); very rare (< 1/10,000); not known (cannot be estimated from the available data). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 4: Adverse reactions

System organ class	Adverse reactions	Frequency
Infections and infestations	Upper respiratory tract	very common
	infection	
	Nasopharyngitis	very common
	Influenza*	common
	Rhinitis	common
Metabolism and nutrition	Hypoglycaemia*	common
disorders		
Psychiatric disorders	Depression	not known
Nervous system disorders	Headache	very common
	Dizziness	very common
Ear and labyrinth	Ear pain	common
disorders	Ear discomfort	common
	Tinnitus	common
	Tympanic membrane	common
	hyperaemia	
	Vestibular disorder	common
	Ear congestion	uncommon
Respiratory, thoracic and	Oropharyngeal pain	very common
mediastinal disorders	Nasal congestion	very common

System organ class	Adverse reactions	Frequency
	Abnormal breathing*	common
	Rhinorrhoea*	common
	Sinus congestion	common
	Pharyngeal erythema	common
	Wheezing*	uncommon
Gastrointestinal disorders	Abdominal pain	very common
	Diarrhoea	very common
	Abdominal pain upper*	common
	Flatulence*	common
Hepatobiliary disorders	Transaminase elevations	very common
	Alanine aminotransferase increased*	very common
	Aspartate aminotransferase increased*	common
	Liver injury [†]	not known
	Total bilirubin increase [†]	not known
Skin and subcutaneous tissue disorders	Rash	very common
tissue disorders	Acne*	common
	Pruritus*	common
Reproductive system and	Breast mass	common
breast disorders	Breast inflammation	uncommon
	Gynaecomastia	uncommon
	Nipple disorder	uncommon
	Nipple pain	uncommon
Investigations	Bacteria in sputum	very common
	Blood creatine phosphokinase increased*	common
	Blood pressure increased*	uncommon

^{*} Adverse reaction and frequency reported from clinical studies with ivacaftor in combination with ivacaftor/tezacaftor/elexacaftor.

Description of selected adverse reactions

Transaminase elevations

During the 48-week placebo-controlled studies 770-102 and 770-103 in patients aged 6 years and older, the incidence of maximum transaminase (ALT or AST) > 8, > 5 or $> 3 \times$ ULN was 3.7%, 3.7% and 8.3% in ivacaftor-treated patients and 1.0%, 1.9% and 8.7% in placebo-treated patients, respectively. Two patients, one on placebo and one on ivacaftor, permanently discontinued treatment for elevated transaminases, each $> 8 \times$ ULN. No ivacaftor-treated patients experienced a transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 1.5 \times$ ULN. In ivacaftor-treated patients, most transaminase elevations up to $5 \times$ ULN resolved without treatment interruption. Ivacaftor dosing

[†] Liver injury (ALT and AST and total bilirubin increase) reported from post-marketing data with ivacaftor in combination with ivacaftor/tezacaftor/elexacaftor. This also included liver failure leading to transplantation in a patient with pre-existing cirrhosis and portal hypertension. Frequency cannot be estimated from the available data.

was interrupted in most patients with transaminase elevations $> 5 \times \text{ULN}$. In all instances where dosing was interrupted for elevated transaminases and subsequently resumed, ivacaftor dosing was able to be resumed successfully (see section 4.4).

During the placebo-controlled phase 3 studies (up to 24 weeks) of tezacaftor/ivacaftor, the incidence of maximum transaminase (ALT or AST) > 8, > 5, or $> 3 \times$ ULN were 0.2%, 1.0%, and 3.4% in tezacaftor/ivacaftor-treated patients, and 0.4%, 1.0%, and 3.4% in placebo-treated patients. One patient (0.2%) on therapy and 2 patients (0.4%) on placebo permanently discontinued treatment for elevated transaminases. No patients treated with tezacaftor/ivacaftor experienced a transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN.

During the 24-week, placebo-controlled, phase 3 study of ivacaftor/tezacaftor/elexacaftor, these figures were 1.5%, 2.5%, and 7.9% in ivacaftor/tezacaftor/elexacaftor-treated patients and 1.0%, 1.5%, and 5.5% in placebo-treated patients. The incidence of adverse reactions of transaminase elevations was 10.9% in ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor-treated patients and 4.0% in placebo-treated patients.

Post-marketing cases of treatment discontinuation due to elevated transaminases have been reported with ivacaftor/tezacaftor/elexacaftor (see section 4.4).

Rash events

In study 445-102, the incidence of rash events (e.g., rash, rash pruritic) was 10.9% in ivacaftor/tezacaftor/elexacaftor-treated patients and 6.5% in placebo-treated patients. The rash events were generally mild to moderate in severity. The incidence of rash events by patient sex was 5.8% in males and 16.3% in females in ivacaftor/tezacaftor/elexacaftor-treated patients and 4.8% in males and 8.3% in females in placebo-treated patients. In patients treated with ivacaftor/tezacaftor/elexacaftor, the incidence of rash events was 20.5% in females taking hormonal contraceptive and 13.6% in females not taking hormonal contraceptive (see section 4.4).

Increased creatine phosphokinase

In study 445-102, the incidence of maximum creatine phosphokinase > 5 x the ULN was 10.4% in ivacaftor/tezacaftor/elexacaftor-treated patients and 5.0% in placebo-treated patients. The observed creatine phosphokinase elevations were generally transient and asymptomatic and many were preceded by exercise. No ivacaftor/tezacaftor/elexacaftor-treated patients discontinued treatment for increased creatine phosphokinase.

Increased blood pressure

In study 445-102, the maximum increase from baseline in mean systolic and diastolic blood pressure was 3.5 mmHg and 1.9 mmHg, respectively for ivacaftor/tezacaftor/elexacaftor-treated patients (baseline: 113 mmHg systolic and 69 mmHg diastolic) and 0.9 mmHg and 0.5 mmHg, respectively for placebo-treated patients (baseline: 114 mmHg systolic and 70 mmHg diastolic).

The proportion of patients who had systolic blood pressure > 140 mmHg or diastolic blood pressure > 90 mmHg on at least two occasions was 5.0% and 3.0%, respectively in ivacaftor/tezacaftor-treated patients compared with 3.5% and 3.5%, respectively in placebo-treated patients.

Paediatric population

Ivacaftor as monotherapy

Safety of ivacaftor as monotherapy for 24 weeks was evaluated in 43 patients between 1 month to less than 24 months of age (with 7 of them less than 4 months old), 34 patients between 2 to less than

6 years of age, 61 patients between 6 to less than 12 years of age and 94 patients between 12 to less than 18 years of age.

The safety profile is generally consistent among paediatric patients aged 1 month and older and is also consistent with adult patients.

The incidence of transaminase elevations (ALT or AST) observed in studies 770-103, 770-111 and 770-110 (patients aged 6 to less than 12 years), study 770-108 (patients aged 2 to less than 6 years), and study 770-124 (patients aged 1 to less than 24 months) are described in Table 5. In the placebocontrolled studies, the incidence of transaminase elevations were similar between treatment with ivacaftor (15.0%) and placebo (14.6%). Peak LFT elevations were generally higher in paediatric patients than in older patients. Across all populations, peak LFT elevations returned to baseline levels following interruption, and in almost all instances where dosing was interrupted for elevated transaminases and subsequently resumed, ivacaftor dosing was able to be resumed successfully (see section 4.4). Cases suggestive of positive rechallenge were observed.

In study 770-108 ivacaftor was permanently discontinued in one patient. In study 770-124, in the cohort of patients aged 1 month to less than 4 months, a 1-month old (14.3%) patient had transaminase values of ALT $> 8 \times ULN$ and AST of > 3 to $\le 5 \times ULN$, which led to discontinuation of ivacaftor treatment (see section 4.4 for management of elevated transaminases).

Table 5: Transaminase elevations in patients aged 1 month to < 12 years treated with ivacaftor as monotherapy

	n	% of Patients > 3 × ULN	% of Patients > 5 × ULN	% of Patients > 8 × ULN
6 to < 12 years	40	15.0% (6)	2.5% (1)	2.5% (1)
2 to < 6 years	34	14.7% (5)	14.7% (5)	14.7% (5)
12 to < 24 months	18	27.8% (5)	11.1% (2)	11.1% (2)
1 to < 12 months	24	8.3% (2)	4.2% (1)	4.2% (1)

Ivacaftor in a combination regimen with tezacaftor/ivacaftor

The safety of tezacaftor/ivacaftor in combination with ivacaftor was evaluated in 124 patients between 6 to less than 12 years of age. The tezacaftor 100 mg/ivacaftor 150 mg and ivacaftor 150 mg dose has not been investigated in clinical studies in children aged 6 to less than 12 years weighing 30 to $<40\ kg$.

The safety profile is generally consistent among children and adolescents, and is also consistent with adult patients.

During the 24-week, open-label phase 3 study in patients aged 6 to less than 12 years (study 661-113 part B, n = 70), the incidence of maximum transaminase (ALT or AST) > 8, > 5, and > 3 \times ULN were 1.4%, 4.3%, and 10.0%, respectively. No tezacaftor/ivacaftor-treated patients experienced a transaminase elevation > 3 \times ULN associated with elevated total bilirubin > 2 \times ULN or discontinued tezacaftor/ivacaftor treatment due to transaminase elevations. One patient interrupted treatment due to elevated transaminases, and subsequently resumed tezacaftor/ivacaftor treatment successfully (see section 4.4 for management of elevated transaminases).

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

The safety data of ivacaftor/tezacaftor/elexacaftor in combination with ivacaftor in studies 445-102, 445-103, 445-104, 445-106, 445-111, and 445-124 was evaluated in 272 patients between 2 to less than 18 years of age. The safety profile is generally consistent among paediatric and adult patients.

During study 445-106 in patients aged 6 to less than 12 years, the incidence of maximum transaminase (ALT or AST) > 8, > 5, and $> 3 \times$ ULN were 0.0%, 1.5%, and 10.6%, respectively. No ivacaftor/tezacaftor/elexacaftor-treated patients had transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN or discontinued treatment due to transaminase elevations (see section 4.4).

During study 445-111 in patients aged 2 to less than 6 years, the incidence of maximum transaminase (ALT or AST) > 8, > 5, and $> 3 \times$ ULN were 1.3%, 2.7%, and 8.0%, respectively. No ivacaftor/tezacaftor/elexacaftor-treated patients had transaminase elevation $> 3 \times$ ULN associated with elevated total bilirubin $> 2 \times$ ULN or discontinued treatment due to transaminase elevations (see section 4.4).

Rash

During study 445-111 in patients aged 2 to less than 6 years, 15 (20.0%) subjects had at least 1 rash event, 4 (9.8%) females and 11 (32.4%) males.

Lenticular opacity

One patient had an adverse event of lenticular opacity.

Reporting of suspected adverse reactions

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

4.9 Overdose

No specific antidote is available for overdose with ivacaftor. Treatment of overdose consists of general supportive measures including monitoring of vital signs, liver function tests and observation of the clinical status of the patient.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other respiratory system products, ATC code: R07AX02

Mechanism of action

Ivacaftor as monotherapy

Ivacaftor is a potentiator of the CFTR protein, i.e., *in vitro* ivacaftor increases CFTR channel gating to enhance chloride transport in specified gating mutations (as listed in section 4.1) with reduced channel-open probability compared to normal CFTR. Ivacaftor also potentiated the channel-open probability of *R117H-CFTR*, which has both low channel-open probability (gating) and reduced channel current amplitude (conductance). The *G970R* mutation causes a splicing defect resulting in little-to-no CFTR protein at the cell surface which may explain the results observed in subjects with this mutation in study 770-111 (see Pharmacodynamic effects and Clinical efficacy and safety).

In vitro responses seen in single channel patch clamp experiments using membrane patches from rodent cells expressing mutant CFTR forms do not necessarily correspond to *in vivo* pharmacodynamic response (e.g., sweat chloride) or clinical benefit. The exact mechanism leading ivacaftor to potentiate

the gating activity of normal and some mutant CFTR forms in this system has not been completely elucidated.

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

Elexacaftor and tezacaftor are CFTR correctors that bind to different sites on the CFTR protein and have an additive effect in facilitating the cellular processing and trafficking of CFTR to increase the amount of CFTR protein delivered to the cell surface compared to either molecule alone. When ivacaftor is administered in combination with ivacaftor/tezacaftor/elexacaftor, the combined effect is increased quantity and function of CFTR at the cell surface, resulting in increased CFTR activity as measured by CFTR mediated chloride transport.

CFTR Chloride Transport Assay in Fischer Rat Thyroid (FRT) cells expressing mutant CFTR

The chloride transport response of mutant CFTR protein to ivacaftor/tezacaftor/elexacaftor was determined in Ussing chamber electrophysiology studies using a panel of FRT cell lines transfected with individual *CFTR* mutations. Ivacaftor/tezacaftor/elexacaftor increased chloride transport in FRT cells expressing select *CFTR* mutations.

The *in vitro* CFTR chloride transport response threshold was designated as a net increase of at least 10% of normal over baseline because it is predictive or reasonably expected to predict clinical response. For individual mutations, the magnitude of the net change over baseline in CFTR mediated chloride transport *in vitro* is not correlated with the magnitude of clinical response.

In CF, the presence of one *CFTR* mutation responsive to ivacaftor/tezacaftor/elexacaftor based on *in vitro* data in FRT cells, will likely result in a clinical response.

Table 6 lists *CFTR* mutations included in the indication for treatment in combination with ivacaftor/tezacaftor/elexacaftor. The occurrence of *CFTR* mutations listed in this table should not be used in lieu of a diagnosis of cystic fibrosis, nor as a sole determinant for prescribing purposes.

Table 6: CFTR mutations identified to be responsive to ivacaftor/tezacaftor/elexacaftor based on clinical and/or in vitro data

293A→G	E217G	H620Q	N900K	S50P
314del9	E264V	H939R	N1088D	S108F
546insCTA	E282D	H939R;H949L [‡]	N1195T	S158N
548insTAC	E292K	H954P	N1303I	S182R
$711+3A\rightarrow G^*$	E384K	H1054D	N1303K*	S308P
1140-1151dup	E403D	H1079P	P5L [†]	S341P
1336K	E474K	H1085P	P67L*	S364P
1461insGAT	E527G	H1085R	P111L	S434P
1507_1515del9	E588V	H1375N	P140S	S492F
2055del9	E822K	H1375P	P205S	S519G
2183A→G	E831X	I86M	P439S	S531P
2789+5G→A*	E1104K	I105N	P499A	S549I
2851A/G	E1104V	I125T	P574H	S549N
3007del6	E1126K	I148L	P750L	S549R*
3132T→G	E1221V	I148N	P798S	S557F
3141del9	E1228K	I175V	P988R	S589I
3143del9	E1409K	I331N	P1013H	S589N
3272-26A→G* [†]	E1433K	I336L	P1013L	S624R
3331del6	F87L	I444S	P1021L	S686Y
3410T→C	F191V	I497S	P1021T	S737F
3523A→G	F200I	I502T	P1372T	S821G
3601A→C	F311del	I506L	Q30P	S898R
3761T→G	F311L	I506V	Q98P	S912L

3791C/T	F312del	I506V;D1168G‡	Q98R	S912L;G1244V [‡]
3/91C/1 3849+10kbC→T*†	F433L	1500 V,D1108G*	Q151K	S912L,G1244V*
3850G→A	F508C;S1251 N [‡]	1521S 1530N	Q179K	S945L*†
3978G→C	F508del*	1556V	Q237E	S955P
3978G→C A46D		1586V		S977F
	F508del;R1438W [‡]		Q237H	
A62P	F575Y	I601F	Q237P	\$977F;R1438W
A107G	F587I	I618N	Q359K;T360K [‡]	* C104537
A120T	F587L	I618T	Q359R	S1045Y
A141D	F693L(TTG)	I980K	Q372H	S1118F
A155P	F932S	I1023R	Q493L	S1159F
A234D	F1016S	I1139V	Q493R	S1159P
A234V	F1052V	I1203V	Q552P	S1188L
A238V	F1074L	I1234L	Q1012P	S1251N
A309D	F1078S	I1234V	Q1209P	S1255P
A349V	F1099L	I1269N	Q1291H	T338I
A357T	F1107L	I1366N	Q1291R	T351I
A455E*†	G27E	I1366T	Q1313K	T351S
A455V	G27R	K162E	Q1352H	T351S;R851L [‡]
A457T	G126D	K464E	R31L	T388M
A462P	G178E	K464N	R74Q	T465I
A534E	G178R	K522E	R74Q;R297Q [‡]	T501A
A554E	G194R	K522Q	R74Q;V201M;D1270N [‡]	T582S
A566D	G194V	K951E	R74W	T908N
A872E	G213E	K1060T	R74W;D1270N [‡]	T990I
A1006E	G213E;R668C [‡]	L15P	R74W;R1070W;D1270	T1036N*
A1025D	G213V	L15P;L1253F [‡]	N [‡]	T1057R
A1067P	G226R	L32P	R74W;S945L [‡]	T1086A
A1067T	G239R	L88S	R74W;V201M [‡]	T1086I
A1067V	G253R	L102R;F1016S [‡]	R74W;V201M;D1270N [‡]	T1246I
A1081V	G314E	L137P	R74W;V201M;L997F [‡]	T1299I
A1087P	G314R	L159S	R75L	T1299K
A1319E	G424S	L165S	R75Q;L1065P [‡]	V11I
A1374D	G437D	L167R	R75Q;N1088D [‡]	V93D
A1466S	G461R	L206W*†	R75Q;S549N [‡]	V201M
C225R	G461V	L210P	R117C [†]	V232A
C491R	G463V	L293P	R117C;G576A;R668C [‡]	V232D
C590Y	G480C	L327P	R117G	V317A
C866Y	G480D	L333F	R117H*	V322M
c.1367_1369dupTTG	G480S	L333H	R117L	V392G
D58H	G500D	L346P	R117L;L997F [‡]	V456A
D58V	G545R	L441P	R117P	V456F
D110E	G551A	L453S	R248K	V520I
D110H	G551D*	L467F	R258G	V562I;A1006E [‡]
D110N	G551R	L558F	R297Q	V562L
D192G	G551S	L619S	R334L	V591A
D192N	G576A;R668C [‡]	L633P	R334Q	V603F
D373N	G576A;S1359Y‡	L636P	R334W	V920L
D426N	G622D	L927P	R347H*	V920M
D443Y	G622V	L967F;L1096R‡	R347L	V1008D
D443Y;G576A;R668C [‡]	G628A	L973F	R347P	V1010D
D529G	G628R	L1011S	R352Q	V1153E
D565G	G85E*†	L1065R	R352W	V1240G
D567N	G930E	L1077P*†	R516S	V1293G
D579G	G970D	L1227S	R553Q	V1293I
D614G	G970S	L1324P	R555G	V1415F
		1 11	1	

D651H	G970V	L1335P	R600S	W202C
D651N	G1047D	L1388P	R709Q	W361R
D806G	G1047R	L1480P	R751L	W496R
D924N	G1061R	M150K	R792G	W1098C
D979A	G1069R	M150R	R792Q	W1282G
D979V	G1123R	M152L	R810G	W1282R
D985H	G1173S	M152V	R851L	Y89C
D985Y	G1237V	M265R	R933G	Y109H
D993A	G1244E	M348K	R1048G	Y109N
D993G	G1244R	M394L	R1066C	Y122C
D993Y	G1247R	M469V	R1066G	Y161C
D1152A	G1249E	M498I	R1066H*†	Y161D
D1152H*†	G1249R	M952I	R1070P	Y161S
D1270N*	G1265V	M952T	R1070Q	Y301C
D1270Y	G1298V	M961L	R1070W	Y563N
D1312G	G1349D	M1101K*†	R1162Q	Y913S
D1377H	G149R;G576A;R668C [‡]	M1137R	R1239S	Y919C
D1445N	H139L	M1137V	R1283G	Y1014C
E56K	H139R	M1210K	R1283M	Y1032C
E60K	H146R	N186K	R1283S	Y1032N
E92K	H199Q	N187K	R1438W	Y1073C
E116K	H199Y	N396Y	S13F	Y1092H
E116Q	H609L	N418S	S13P	Y1381H
E193K	H620P		S18I	
			S18N	

There are people with CF harbouring two, rare non-*F508del* mutations not listed in Table 6. Provided that they do not harbour two class I (null) mutations (mutations that are known not to produce CFTR protein) (see section 4.1), they may respond to treatment. In these cases, IVA in combination with IVA/TEZ/ELX can be considered when the physician deems the potential benefits outweigh the potential risks and under close medical supervision. The individual diagnosis of CF should be based on diagnostic guidelines and clinical judgement as considerable variability exists in phenotype for patients harbouring the same genotype.

Non-annotated mutations are included based on the FRT assay in which a positive response is indicative of a clinical response.

Pharmacodynamic effects

Ivacaftor as monotherapy

In studies 770-102 and 770-103 in patients with the *G551D* mutation in one allele of the *CFTR* gene, ivacaftor led to rapid (15 days), substantial (the mean change in sweat chloride from baseline through week 24 was -48 mmol/L [95% CI: -51, -45] and -54 mmol/L [95% CI: -62, -47], respectively) and sustained (through 48 weeks) reductions in sweat chloride concentration.

In study 770-111, part 1 in patients who had a non-*G551D* gating mutation in the *CFTR* gene, treatment with ivacaftor led to a rapid (15 days) and substantial mean change from baseline in sweat chloride of -49 mmol/L (95% CI: -57, -41) through 8 weeks of treatment. However, in patients with the *G970R-CFTR* mutation, the mean (SD) absolute change in sweat chloride at week 8 was -6.25 (6.55) mmol/L. Similar results to part 1 were seen in part 2 of the study. At the 4-week follow-up visit (4 weeks after dosing with ivacaftor ended), mean sweat chloride values for each group were trending to pre-treatment levels.

^{*} Mutations supported by clinical data.

[†] Mutations supported by Real-World data in \geq 5 patients.

[‡] Complex/compound mutations where a single allele of the *CFTR* gene has multiple mutations; these exist independent of the presence of mutations on the other allele.

In study 770-110 in patients aged 6 years or older with CF who had an *R117H* mutation in the *CFTR* gene, the treatment difference in mean change in sweat chloride from baseline through 24 weeks of treatment was -24 mmol/L (95% CI: -28, -20). In subgroup analyses by age, the treatment difference was -21.87 mmol/L (95% CI: -26.46, -17.28) in patients aged 18 years or older, and -27.63 mmol/L (95% CI: -37.16, -18.10) in patients aged 6 to 11 years. Two patients 12 to 17 years of age were enrolled in this study.

In study 770-108 in patients aged 2 to less than 6 years with a gating mutation on at least one allele of the *CFTR* gene administered either 50 mg or 75 mg of ivacaftor twice daily, the mean absolute change from baseline in sweat chloride was -47 mmol/L (95% CI: -58, -36) at week 24.

In study 770-124 in patients with CF aged 1 month to less than 24 months, the mean absolute change from baseline in sweat chloride was -62.0 mmol/L (95% CI: -71.6, -52.4) at week 24. Results were consistent in the 12 months to less than 24 months, 6 months to less than 12 months, and 4 months to less than 6 months age cohorts.

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

In study 445-111 in patients aged 2 to less than 6 years who are homozygous for the *F508del* mutation or heterozygous for the *F508del* mutation and a minimal function mutation, the mean absolute change in sweat chloride from baseline through week 24 was -57.9 mmol/L (95% CI: -61.3, -54.6).

In study 445-124 (patients aged 6 years and older with a qualifying non-F508del, ivacaftor/tezacaftor/elexacaftor-responsive CFTR mutation), the mean absolute change in sweat chloride from baseline through week 24 compared to placebo was -28.3 mmol/L (95% CI: -32.1, -24.5 mmol/L; P < 0.0001).

Clinical efficacy and safety

Ivacaftor as monotherapy

Studies 770-102 and 770-103: studies in patients with CF with G551D gating mutations

The efficacy of ivacaftor has been evaluated in two phase 3 randomised, double-blind, placebo-controlled, multi-centre studies of clinically stable patients with CF who had the G551D mutation in the CFTR gene on at least one allele and had $FEV_1 \ge 40\%$ predicted.

Patients in both studies were randomised 1:1 to receive either 150 mg of ivacaftor or placebo every 12 hours with food containing fat for 48 weeks in addition to their prescribed CF therapies (e.g., tobramycin, dornase alfa). The use of inhaled hypertonic sodium chloride was not permitted.

Study 770-102 evaluated 161 patients who were 12 years of age or older; 122 (75.8%) patients had the F508del mutation in the second allele. At the start of the study, patients in the placebo group used some medicinal products at a higher frequency than the ivacaftor group. These medicinal products included dornase alfa (73.1% versus 65.1%), salbutamol (53.8% versus 42.2%), tobramycin (44.9% versus 33.7%) and salmeterol/fluticasone (41.0% versus 27.7%). At baseline, mean predicted FEV₁ was 63.6% (range: 31.6% to 98.2%) and mean age was 26 years (range: 12 to 53 years).

Study 770-103 evaluated 52 patients who were 6 to 11 years of age at screening; mean (SD) body weight was 30.9 (8.63) kg; 42 (80.8%) patients had the F508del mutation in the second allele. At baseline, mean predicted FEV₁ was 84.2% (range: 44.0% to 133.8%) and mean age was 9 years (range: 6 to 12 years); 8 (30.8%) patients in the placebo group and 4 (15.4%) patients in the ivacaftor group had an FEV₁ less than 70% predicted at baseline.

The primary efficacy endpoint in both studies was the mean absolute change from baseline in percent predicted FEV₁ through 24 weeks of treatment.

The treatment difference between ivacaftor and placebo for the mean absolute change (95% CI) in percent predicted FEV_1 from baseline through week 24 was 10.6 percentage points (8.6, 12.6) in study 770-102 and 12.5 percentage points (6.6, 18.3) in study 770-103. The treatment difference between ivacaftor and placebo for the mean relative change (95% CI) in percent predicted FEV_1 from baseline through week 24 was 17.1% (13.9, 20.2) in study 770-102 and 15.8% (8.4, 23.2) in study 770-103. The mean change from baseline through week 24 in FEV_1 (L) was 0.37 L in the ivacaftor group and 0.01 L in the placebo group in study 770-102 and 0.30 L in the ivacaftor group and 0.07 L in the placebo group in study 770-103. In both studies, improvements in FEV_1 were rapid in onset (day 15) and durable through 48 weeks.

The treatment difference between ivacaftor and placebo for the mean absolute change (95% CI) in percent predicted FEV₁ from baseline through week 24 in patients 12 to 17 years of age in study 770-102 was 11.9 percentage points (5.9, 17.9). The treatment difference between ivacaftor and placebo for the mean absolute change (95% CI) in percent predicted FEV₁ from baseline through week 24 in patients with baseline predicted FEV₁ greater than 90% in study 770-103 was 6.9 percentage points (-3.8, 17.6).

The results for clinically relevant secondary endpoints are shown in Table 7.

Table 7: Effect of ivacaftor on other efficacy endpoints in studies 770-102 and 770-103

	Study 770-102		Study 7	70-103
Endpoint	Treatment difference ^a (95% CI)	<i>P</i> -value	Treatment difference ^a (95% CI)	<i>P</i> -value
Mean absolute change fr		Q-R ^b respirator		oints) ^c
Through week 24	8.1	< 0.0001	6.1	0.1092
	(4.7, 11.4)		(-1.4, 13.5)	
Through week 48	8.6	< 0.0001	5.1	0.1354
	(5.3, 11.9)		(-1.6, 11.8)	
Relative risk of pulmona	ry exacerbation			
Through week 24	0.40^{d}	0.0016	NA	NA
Through week 48	0.46^{d}	0.0012	NA	NA
Mean absolute change fr	om baseline in bod	y weight (kg)		
At week 24	2.8	< 0.0001	1.9	0.0004
	(1.8, 3.7)		(0.9, 2.9)	
At week 48	2.7	0.0001	2.8	0.0002
	(1.3, 4.1)		(1.3, 4.2)	
Mean absolute change fr	om baseline in BM	$I (kg/m^2)$		
At week 24	0.94	< 0.0001	0.81	0.0008
	(0.62, 1.26)		(0.34, 1.28)	
At week 48	0.93	< 0.0001	1.09	0.0003
	(0.48, 1.38)		(0.51, 1.67)	
Mean change from basel	ine in z-scores			
Weight-for-age z-score	0.33	0.0260	0.39	< 0.0001
at week 48e	(0.04, 0.62)		(0.24, 0.53)	

	Study 770-102		Study 7	70-103
	Treatment difference ^a		Treatment difference ^a	
Endpoint	(95% CI)	<i>P</i> -value	(95% CI)	<i>P</i> -value
BMI-for-age z-score at	0.33	0.0490	0.45	< 0.0001
week 48 ^e	(0.002, 0.65)		(0.26, 0.65)	

CI: Confidence Interval; NA: not analysed due to low incidence of events

- ^a Treatment difference = effect of ivacaftor effect of placebo.
- b CFQ-R: Cystic Fibrosis Questionnaire-Revised is a disease-specific, health-related quality-of-life measure for CF
- Study 770-102 data were pooled from CFQ-R for adults/adolescents and CFQ-R for children 12 to 13 years of age; study 770-103 data were obtained from CFQ-R for children 6 to 11 years of age.
- d Hazard ratio for time to first pulmonary exacerbation.
- e In subjects under 20 years of age (CDC growth charts).

Study 770-111: study in patients with CF with non-G551D gating mutations

Study 770-111 was a phase 3, two-part, randomised, double-blind, placebo-controlled, crossover study (part 1) followed by a 16-week open-label extension period (part 2) to evaluate the efficacy and safety of ivacaftor in patients with CF aged 6 years and older who have a *G970R* or non-*G551D* gating mutation in the *CFTR* gene (*G178R*, *S549N*, *S549R*, *G551S*, *G1244E*, *S1251N*, *S1255P* or *G1349D*).

In part 1, patients were randomised 1:1 to receive either 150 mg of ivacaftor or placebo every 12 hours with fat-containing food for 8 weeks in addition to their prescribed CF therapies and crossed over to the other treatment for the second 8 weeks after a 4- to 8-week washout period. The use of inhaled hypertonic saline was not permitted. In part 2, all patients received ivacaftor as indicated in part 1 for 16 additional weeks. The duration of continuous ivacaftor treatment was 24 weeks for patients randomised to the part 1 placebo/ivacaftor treatment sequence and 16 weeks for patients randomised to part 1 ivacaftor/placebo treatment sequence.

Thirty-nine patients (mean age 23 years) with baseline $FEV_1 \ge 40\%$ predicted (mean FEV_1 78% predicted [range: 43% to 119%]) were enrolled. Sixty-two percent (24/39) of them carried the *F508del-CFTR* mutation in the second allele. A total of 36 patients continued into part 2 (18 per treatment sequence).

In part 1 of study 770-111, the mean FEV₁ percent predicted at baseline in placebo-treated patients was 79.3% while in ivacaftor-treated patients this value was 76.4%. The mean overall post-baseline value was 76.0% and 83.7%, respectively. The mean absolute change from baseline through week 8 in percent predicted FEV₁ (primary efficacy endpoint) was 7.5% in the ivacaftor period and -3.2% in the placebo period. The observed treatment difference (95% CI) between ivacaftor and placebo was 10.7% (7.3, 14.1) (P < 0.0001).

The effect of ivacaftor in the overall population of study 770-111 (including the secondary endpoints absolute change in BMI at 8 weeks of treatment and absolute change in the respiratory domain score of the CFQ-R through 8 weeks of treatment) and by individual mutation (absolute change in sweat chloride and in percent predicted FEV_1 at week 8) is shown in Table 8. Based on clinical (percent predicted FEV_1) and pharmacodynamic (sweat chloride) responses to ivacaftor, efficacy in patients with the G970R mutation could not be established.

Table 8: Effect of ivacaftor for efficacy variables in the overall population and for specific *CFTR* mutations

Absolute change in percent predicted FEV ₁	BMI (kg/m²)	CFQ-R respiratory domain score (points)
Through week 8	At week 8	Through week 8

All patients (N = 39)			
Results shown as mean (95% CI) change from baseline ivacaftor vs. placebo-treated patients:			
10.7 (7.3, 14.1)	0.66 (0.34, 0.99)	9.6 (4.5, 14.7)	

Patients grouped under mutation types (n)

Results shown as mean (minimum, maximum) change from baseline for ivacaftor-treated patients at week 8^* :

	Absolute change in sweat chloride (mmol/L)	Absolute change in percent predicted FEV ₁ (percentage points)
Mutation (n)	At week 8	At week 8
G1244E (5)	-55 (-75, -34)	8 (-1, 18)
G1349D(2)	-80 (-82, -79)	20 (3, 36)
G178R(5)	-53 (-65, -35)	8 (-1, 18)
G551S(2)	-68 [†]	3^{\dagger}
$G970R^{\#}(4)$	-6 (-16, -2)	3 (-1, 5)
S1251N(8)	-54 (-84, -7)	9 (-20, 21)
S1255P(2)	-78 (-82, -74)	3 (-1, 8)
S549N(6)	-74 (-93, -53)	11 (-2, 20)
S549R (4)	-61 ^{††} (-71, -54)	5 (-3, 13)

Statistical testing was not performed due to small numbers for individual mutations.

In part 2 of study 770-111, the mean (SD) absolute change in percent predicted FEV₁ following 16 weeks (patients randomised to the ivacaftor/placebo treatment sequence in part 1) of continuous ivacaftor treatment was 10.4% (13.2%). At the follow-up visit 4 weeks after ivacaftor dosing had ended, the mean (SD) absolute change in percent predicted FEV₁ from part 2 week 16 was -5.9% (9.4%). For patients randomised to the placebo/ivacaftor treatment sequence in part 1 there was a further mean (SD) change of 3.3% (9.3%) in percent predicted FEV₁ after the additional 16 weeks of treatment with ivacaftor. At the follow-up visit 4 weeks after ivacaftor dosing had ended, the mean (SD) absolute change in percent predicted FEV₁ from part 2, week 16 was -7.4% (5.5%).

Study 770-104: study in patients with CF with the F508del mutation in the CFTR gene

Study 770-104 (part A) was a 16-week, 4:1 randomised, double-blind, placebo-controlled, parallel-group phase 2 study of ivacaftor (150 mg every 12 hours) in 140 patients with CF aged 12 years and older who were homozygous for the F508del mutation in the CFTR gene and who had $FEV_1 \ge 40\%$ predicted.

The mean absolute change from baseline through week 16 in percent predicted FEV_1 (primary efficacy endpoint) was 1.5 percentage points in the ivacaftor group and -0.2 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 1.7 percentage points (95% CI: -0.6, 4.1); this difference was not statistically significant (P = 0.15).

Study 770-105: open-label extension study

In study 770-105, patients who completed treatment in studies 770-102 and 770-103 with placebo were switched to ivacaftor while patients on ivacaftor continued to receive it for a minimum of 96 weeks, i.e., the length of treatment with ivacaftor was at least 96 weeks for patients in the placebo/ivacaftor group and at least 144 weeks for patients in the ivacaftor/ivacaftor group.

One hundred and forty-four (144) patients from study 770-102 were rolled over in study 770-105, 67 in the placebo/ivacaftor group and 77 in the ivacaftor/ivacaftor group. Forty-eight (48) patients from study 770-103 were rolled over in study 770-105, 22 in the placebo/ivacaftor group and 26 in the ivacaftor/ivacaftor group.

[†] Reflects results from the one patient with the G551S mutation with data at the 8-week time point.

 $^{^{\}dagger\dagger}$ n = 3 for the analysis of absolute change in sweat chloride.

[#] Causes a splicing defect resulting in little-to-no CFTR protein at the cell surface.

Table 9 shows the results of the mean (SD) absolute change in percent predicted FEV_1 for both groups of patients. For patients in the placebo/ivacaftor group baseline percent predicted FEV_1 is that of study 770-105 while for patients in the ivacaftor/ivacaftor group the baseline value is that of studies 770-102 and 770-103.

Table 9: Effect of ivacaftor on percent predicted FEV₁ in study 770-105

Original study and	Duration of ivacaftor	Absolute change from baseline in percent predicted FEV ₁ (percentage points)				
treatment group	treatment (weeks)	N Mean (SD)				
Study 770-102						
Ivacaftor	48*	77	9.4 (8.3)			
	144	72	9.4 (10.8)			
Placebo	0*	67	-1.2 (7.8) [†]			
	96	55	9.5 (11.2)			
Study 770-103						
Ivacaftor	48*	26	10.2 (15.7)			
	144	25	10.3 (12.4)			
Placebo	0*	22	-0.6 (10.1) [†]			
	96	21	10.5 (11.5)			

^{*} Treatment occurred during blinded, controlled, 48-week phase 3 study.

When the mean (SD) absolute change in percent predicted FEV_1 is compared from study 770-105 baseline for patients in the ivacaftor/ivacaftor group (n = 72) who rolled over from study 770-102, the mean (SD) absolute change in percent predicted FEV_1 was 0.0% (9.05), while for patients in the ivacaftor/ivacaftor group (n = 25) who rolled over from study 770-103 this figure was 0.6% (9.1). This shows that patients in the ivacaftor/ivacaftor group maintained the improvement seen at week 48 of the initial study (day 0 through week 48) in percent predicted FEV_1 through week 144. There were no additional improvements in study 770-105 (week 48 through week 144).

For patients in the placebo/ivacaftor group from study 770-102, the annualised rate of pulmonary exacerbations was higher in the initial study when patients were on placebo (1.34 events/year) than during the subsequent study 770-105 when patients rolled over to ivacaftor (0.48 events/year across day 1 to week 48, and 0.67 events/year across weeks 48 to 96). For patients in the ivacaftor/ivacaftor group from study 770-102, the annualised rate of pulmonary exacerbations was 0.57 events/year across day 1 to week 48 when patients were on ivacaftor. When they rolled over into study 770-105, the rate of annualised pulmonary exacerbations was 0.91 events/year across day 1 to week 48 and 0.77 events/year across weeks 48 to 96.

For patients who rolled over from study 770-103 the number of events was overall low.

Study 770-110: study in patients with CF with an R117H mutation in the CFTR gene

Study 770-110 evaluated 69 patients who were 6 years of age or older; 53 (76.8%) patients had the *F508del* mutation in the second allele. The confirmed *R117H* poly-T variant was *5T* in 38 patients and 7T in 16 patients. At baseline, mean predicted FEV₁ was 73% (range: 32.5% to 105.5%) and mean age was 31 years (range: 6 to 68 years). The mean absolute change from baseline through week 24 in percent predicted FEV₁ (primary efficacy endpoint) was 2.57 percentage points in the ivacaftor group and 0.46 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 2.1 percentage points (95% CI: -1.1, 5.4).

A pre-planned subgroup analysis was conducted in patients aged 18 years and older (26 patients on placebo and 24 patients on ivacaftor). Treatment with ivacaftor resulted in a mean absolute change in percent predicted FEV_1 through week 24 of 4.5 percentage points in the ivacaftor group

[†] Change from prior study baseline after 48 weeks of placebo treatment.

versus -0.46 percentage points in the placebo group. The estimated treatment difference for ivacaftor versus placebo was 5.0 percentage points (95% CI: 1.1, 8.8).

In a subgroup analysis in patients with a confirmed R117H-5T genetic variant, the difference in the mean absolute change from baseline through week 24 in percent predicted FEV₁ between ivacaftor and placebo was 5.3% (95% CI: 1.3, 9.3). In patients with a confirmed R117H-7T genetic variant, the treatment difference between ivacaftor and placebo was 0.2% (95% CI: -8.1, 8.5).

For secondary efficacy variables, no treatment differences were observed for ivacaftor versus placebo in absolute change from baseline in BMI at week 24 or time to first pulmonary exacerbation. Treatment differences were observed in absolute change in CFQ-R respiratory domain score through week 24 (treatment difference of ivacaftor versus placebo was 8.4 [95% CI: 2.2, 14.6] points) and for the mean change from baseline in sweat chloride (see Pharmacodynamic effects).

Study 770-108: study in paediatric patients with CF aged 2 to less than 6 years with G551D or another gating mutation

The pharmacokinetic profile, safety and efficacy of ivacaftor in 34 patients aged 2 to less than 6 years with CF who had a *G551D*, *G1244E*, *G1349D*, *G178R*, *G551S*, *S1251N*, *S1255P*, *S549N* or *S549R* mutation in the *CFTR* gene were assessed in a 24-week uncontrolled study with ivacaftor (patients weighing less than 14 kg received ivacaftor 50 mg and patients weighing 14 kg or more received ivacaftor 75 mg). Ivacaftor was administered orally every 12 hours with fat-containing food in addition to their prescribed CF therapies.

Patients in study 770-108 were aged 2 to less than 6 years (mean age 3 years). Twenty-six patients out of the 34 enrolled (76.5%) had a *CFTR* genotype G551D/F508del with only 2 patients with a non-G551D mutation (S549N). The mean (SD) sweat chloride at baseline (n = 25) was 97.88 mmol/L (14.00). The mean (SD) faecal elastase-1 value at baseline (n = 27) was 28 μ g/g (95).

The primary endpoint of safety was evaluated through week 24 (see section 4.8). Secondary and exploratory efficacy endpoints evaluated were absolute change from baseline in sweat chloride through 24 weeks of treatment, absolute change from baseline in weight, body mass index (BMI) and stature (supported by weight, BMI and stature z-scores) at 24 weeks of treatment, and measures of pancreatic function such as faecal elastase-1. Data on percent predicted FEV₁ (exploratory endpoint) were available for 3 patients in the ivacaftor 50 mg group and 17 patients in the 75 mg dosing group.

The mean (SD) overall (both ivacaftor dosing groups combined) absolute change from baseline in BMI at week 24 was 0.32 kg/m² (0.54) and the mean (SD) overall change in BMI-for-age z-score was 0.37 (0.42). The mean (SD) overall change in stature-for-age z-score was -0.01 (0.33). The mean (SD) overall change from baseline in faecal elastase-1 (n = 27) was 99.8 μ g/g (138.4). Six patients with initial levels below 200 μ g/g achieved, at week 24, a level of \geq 200 μ g/g. The mean (SD) overall change in percent predicted FEV₁ from baseline at week 24 (exploratory endpoint) was 1.8 (17.81).

Study 770-124: study in paediatric patients with CF aged less than 24 months

The pharmacokinetics and safety of ivacaftor in patients with CF aged 1 month to less than 24 months were assessed in a 24-week (part B only), open-label single arm study where 19 patients aged 12 months to less than 24 months (mean age 15.2 months at baseline), 11 patients aged 6 months to less than 12 months, 6 patients aged 4 months to less than 6 months, and 7 patients (part A/B) aged 1 month to less than 4 months were enrolled and treated with ivacaftor according to their age and body weight. Mean age at baseline of each cohort was 15.2 months, 9.0 months, 4.5 months, and 1.9 months, respectively.

Primary endpoint in part B and part A/B was safety through 24 weeks. Pharmacokinetics and the absolute change from baseline in sweat chloride through 24 weeks (see Pharmacodynamic effects) were secondary endpoints. Tertiary endpoints included efficacy measures such as faecal elastase-1 and growth parameters.

For patients aged 1 month to less than 24 months, with both baseline and week 24 values available, mean (SD) weight-for-age, length-for-age, and weight-for-length z-scores are provided in Table 10.

Table 10: Effect of ivacaftor on growth parameters in patients aged 1 month to less than 24 months with baseline and week 24 values

Danamatan	Number of	Baseline		Absolute change at week 24	
Parameter	patients	Mean (SD)	Median (min, max)	Mean (SD)	Median (min, max)
Weight-for-age z-score	41	0.00	0.07	0.45	0.30
		(0.94)	[-1.93, 1.79]	(0.64)	[-0.54, 2.66]
Length-for-age z-score	40	-0.03	-0.03	0.44	0.52
		(1.11)	[-1.99, 2.79]	(0.92)	[-1.81, 3.38]
Weight-for-length z-score	40	0.07	0.14	0.32	0.32
		(1.02)	[-1.72, 2.16]	(0.99)	[-2.04, 2.22]

In patients aged 1 month to less than 24 months, out of the 24 subjects who were pancreatic insufficient at baseline (defined as faecal elastase-1 < 200 μ g/g), 14 had faecal elastase-1 values above 200 μ g/g at week 24. In the overall population of part B and part A/B, the median (min, max) value of faecal elastase-1 (μ g/g) was 55.5 (7.5, 500.0) at baseline. The median (min, max) absolute change in faecal elastase-1 from baseline (n = 40) to week 24 (n = 33) was 126.0 (-23.0, 423.5).

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

The efficacy and safety of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor tablets in patients aged 12 years and older was demonstrated in three phase 3, randomised, double-blind, placebo-controlled studies (patients were heterozygous for the F508del mutation and a mutation with minimal function on the second allele, n = 403) and active-controlled (patients were homozygous for the F508del mutation, n = 107, or heterozygous for the F508del mutation and a gating or residual CFTR activity mutation on the second allele, n = 258) studies of 24 (study 445-102), 4 (study 445-103), and 8 weeks (study 445-104) of duration respectively. Patients from all studies were eligible to enter open-label, rollover, long-term extension studies (study 445-105 or study 445-110).

The efficacy and safety of ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor tablets in patients aged 6 years and older with non-F508del, ivacaftor/tezacaftor/elexacaftor-responsive *CFTR* mutations was demonstrated in a phase 3, randomised, double-blind, placebo-controlled study (n = 307) and an observational retrospective study (CFD-016; n = 422) of Real-World clinical outcomes.

Refer to the Summary of Product Characteristics of ivacaftor/tezacaftor/elexacaftor for additional data.

Paediatric population

Ivacaftor in a combination regimen with ivacaftor/tezacaftor/elexacaftor

The pharmacokinetics and safety in patients aged 6 to less than 12 years (n = 66) and in those from 2 to less than 6 years (n = 75) who have at least one F508del mutation were assessed in two 24-week open-label studies (study 445-106 and study 445-111), respectively. Refer to the Summary of Product Characteristics of ivacaftor/tezacaftor/elexacaftor for additional data.

The European Medicines Agency has deferred the obligation to submit the results of studies with Kalydeco in one or more subsets of the paediatric population in cystic fibrosis (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

The pharmacokinetics of ivacaftor are similar between healthy adult volunteers and patients with CF.

After oral administration of a single 150 mg dose to healthy volunteers in a fed state, the mean (\pm SD) for AUC and C_{max} were 10.60 (5.26) µg·h/mL and 0.768 (0.233) µg/mL, respectively. After every 12-hour dosing, steady-state plasma concentrations of ivacaftor were reached by days 3 to 5, with an accumulation ratio ranging from 2.2 to 2.9.

Absorption

Following multiple oral dose administrations of ivacaftor, the exposure of ivacaftor generally increased with dose from 25 mg every 12 hours to 450 mg every 12 hours. When given with fatcontaining food the exposure of ivacaftor increased approximately 2.5- to 4-fold. When coadministered with tezacaftor and elexacaftor, the increase in AUC was similar (approximately 3-fold and 2.5- to 4-fold, respectively). Therefore, ivacaftor, administered as monotherapy or in a combination regimen with ivacaftor/tezacaftor/elexacaftor, should be administered with fat-containing food. The median (range) t_{max} is approximately 4.0 (3.0; 6.0) hours in the fed state.

Ivacaftor granules (2 \times 75 mg sachets) had similar bioavailability as the 150 mg tablet when given with fat-containing food to healthy adult subjects. The geometric least squares mean ratio (90% CI) for the granules relative to tablets was 0.951 (0.839, 1.08) for AUC_{0- ∞} and 0.918 (0.750, 1.12) for C_{max}. The effect of food on ivacaftor absorption is similar for both formulations, i.e., tablets and granules.

Distribution

Ivacaftor is approximately 99% bound to plasma proteins, primarily to alpha 1-acid glycoprotein and albumin. Ivacaftor does not bind to human red blood cells. After oral administration of ivacaftor 150 mg every 12 hours for 7 days in healthy volunteers in a fed state, the mean (\pm SD) apparent volume of distribution was 353 L (122).

Biotransformation

Ivacaftor is extensively metabolised in humans. *In vitro* and *in vivo* data indicate that ivacaftor is primarily metabolised by CYP3A. M1 and M6 are the two major metabolites of ivacaftor in humans. M1 has approximately one-sixth the potency of ivacaftor and is considered pharmacologically active. M6 has less than one-fiftieth the potency of ivacaftor and is not considered pharmacologically active.

The effect of the CYP3A4*22 heterozygous genotype on ivacaftor, tezacaftor, and elexacaftor exposure is consistent with the effect of co-administration of a weak CYP3A4 inhibitor, which is not clinically relevant. No dose adjustment of ivacaftor, tezacaftor, or elexacaftor is considered necessary. The effect in CYP3A4*22 homozygous genotype patients is expected to be stronger. However, no data are available for such patients.

Elimination

Following oral administration in healthy volunteers, the majority of ivacaftor (87.8%) was eliminated in the faeces after metabolic conversion. The major metabolites M1 and M6 accounted for approximately 65% of the total dose eliminated with 22% as M1 and 43% as M6. There was negligible urinary excretion of ivacaftor as unchanged parent. The apparent terminal half-life was approximately 12 hours following a single dose in the fed state. The apparent clearance (CL/F) of ivacaftor was similar for healthy subjects and patients with CF. The mean (±SD) CL/F for a single 150 mg dose was 17.3 (8.4) L/hr in healthy subjects.

Linearity/non-linearity

The pharmacokinetics of ivacaftor are generally linear with respect to time or dose ranging from 25 mg to 250 mg.

Special populations

Hepatic impairment

Following a single dose of 150 mg of ivacaftor, adult subjects with moderately impaired hepatic function (Child-Pugh Class B, score 7 to 9) had similar ivacaftor C_{max} (mean [\pm SD] of 0.735 [0.331] μ g/mL) but an approximately 2-fold increase in ivacaftor AUC_{0-∞} (mean [\pm SD] of 16.80 [6.14] μ g·h/mL) compared with healthy subjects matched for demographics. Simulations for predicting the steady-state exposure of ivacaftor showed that by reducing the dose from 150 mg q12h to 150 mg once daily, adults with moderate hepatic impairment would have comparable steady-state C_{min} values as those obtained with a dose of 150 mg q12h in adults without hepatic impairment.

In subjects with moderately impaired hepatic function (Child-Pugh Class B, score 7 to 9), ivacaftor AUC increased approximately by 50% following multiple doses for 10 days of ivacaftor, tezacaftor and elexacaftor.

The impact of severe hepatic impairment (Child-Pugh Class C, score 10 to 15) on the pharmacokinetics of ivacaftor or in a combination regimen with ivacaftor/tezacaftor/elexacaftor have not been studied. The magnitude of increase in exposure in these patients is unknown but is expected to be higher than that observed in patients with moderate hepatic impairment.

For guidance on appropriate use and dose modification see Table 3 in section 4.2.

Renal impairment

Pharmacokinetic studies have not been performed with ivacaftor in patients with renal impairment. In a human pharmacokinetic study, there was minimal elimination of ivacaftor and its metabolites in urine (only 6.6% of total radioactivity was recovered in the urine). There was negligible urinary excretion of ivacaftor as unchanged parent (less than 0.01% following a single oral dose of 500 mg).

No dose adjustments are recommended for mild and moderate renal impairment. However, caution is recommended when administering ivacaftor to patients with severe renal impairment (creatinine clearance less than or equal to 30 mL/min) or end-stage renal disease (see sections 4.2 and 4.4).

Race

Race had no clinically meaningful effect on the PK of ivacaftor in white (n = 379) and non-white (n = 29) patients based on a population PK analysis.

Gender

The pharmacokinetic parameters of ivacaftor are similar in males and females.

Elderly

Clinical studies of ivacaftor as monotherapy did not include sufficient numbers of patients aged 65 years and older to determine whether pharmacokinetic parameters are similar or not to those in younger adults.

Paediatric population

Predicted ivacaftor exposure based on observed ivacaftor concentrations in phase 2 and 3 studies as determined using compartmental analysis is presented by age group in Table 11.

Table 11: Mean (SD) ivacaftor exposure by age group

Age group	Dose	C _{min, ss} (μg/mL)	AUC _{0-12h} , ss (μg·h/mL)
1 month to less than 2 months	13.4 mg q24h	0.300	5.84
$(\geq 3 \text{ kg})^*$		$(0.221)^{\dagger}$	$(2.98)^{\dagger}$
2 months to less than 4 months (\geq 3 kg)*	13.4 mg q12h	0.406	6.45
		$(0.266)^{\dagger}$	$(3.43)^{\dagger}$
4 months to less than 6 months ($\geq 5 \text{ kg}$)*	25 mg q12h	0.371	6.48
		(0.183)	(2.52)
6 months to less than 12 months (\geq 5 kg to < 7 kg) [‡]	25 mg q12h	0.336	5.41
6 months to less than 12 months (7 kg to < 14 kg)	50 mg q12h	0.508	9.14
, ,		(0.252)	(4.20)
12 months to less than 24 months (7 kg to < 14 kg)	50 mg q12h	0.440	9.05
		(0.212)	(3.05)
12 months to less than 24 months (≥ 14 kg to < 25 kg)	75 mg q12h	0.451	9.60
		(0.125)	(1.80)
2- to 5-year-olds (< 14 kg)	50 mg q12h	0.577	10.50
		(0.317)	(4.26)
2- to 5-year-olds (≥ 14 kg to < 25 kg)	75 mg q12h	0.629	11.30
		(0.296)	(3.82)
(to 1111-8 (> 14 l to < 25 l)	75 mg q12h	0.641	10.76
6- to 11-year-olds $(\ge 14 \text{ kg to} < 25 \text{ kg})$		(0.329)	(4.47)
6- to 11-year-olds [§] (≥ 25 kg)	150 mg q12h	0.958	15.30
, , ,		(0.546)	(7.34)
12- to 17-year-olds	150 mg q12h	0.564	9.24
		(0.242)	(3.42)
Adults (≥ 18 years old)	150 mg q12h	0.701	10.70
		(0.317)	(4.10)

^{*} Patients 1 month to less than 6 months of age were ≥37 weeks gestational age.

5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, and carcinogenic potential.

[†] Exposures for 1 month to less than 4 months of age are predictions based on simulations from the physiologically based PK model incorporating data from the given age group.

[‡] Values based on data from a single patient; standard deviation not reported.

Exposures in 6- to 11-year-olds are predictions based on simulations from the population PK model using data obtained for this age group.

Pregnancy and fertility

Ivacaftor was associated with slight decreases of the seminal vesicle weights, a decrease of overall fertility index and number of pregnancies in females mated with treated males and significant reductions in number of corpora lutea and implantation sites with subsequent reductions in the average litter size and average number of viable embryos per litter in treated females. The No-Observed-Adverse-Effect-Level (NOAEL) for fertility findings provides an exposure level of approximately 4 times the systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy in adult humans at the maximum recommended human dose (MRHD). Placental transfer of ivacaftor was observed in pregnant rats and rabbits.

Peri- and post-natal development

Ivacaftor decreased survival and lactation indices and caused a reduction in pup body weights. The NOAEL for viability and growth in the offspring provides an exposure level approximately 3 times the systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy in adult humans at the MRHD.

Juvenile animal studies

Findings of cataracts were observed in juvenile rats dosed from postnatal day 7 through 35 at ivacaftor exposure levels of 0.22 times the MRHD based on systemic exposure of ivacaftor and its metabolites when administered as ivacaftor monotherapy. This finding has not been observed in foetuses derived from rat dams treated with ivacaftor on gestation days 7 to 17, in rat pups exposed to ivacaftor through milk ingestion up to postnatal day 20, in 7-week old rats, nor in 3.5 to 5-month old dogs treated with ivacaftor. The potential relevance of these findings in humans is unknown.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Silica, colloidal anhydrous Croscarmellose sodium Hypromellose acetate succinate Lactose monohydrate Magnesium stearate Mannitol Sucralose Sodium laurilsulfate (E487)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

4 years.

Once mixed, the mixture has been shown to be stable for one hour.

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

Biaxially Oriented Polyethylene Terephthalate/Polyethylene/Foil/Polyethylene (BOPET/PE/Foil/PE) sachet.

<u>Kalydeco 13.4 mg granules in sachet, Kalydeco 25 mg granules in sachet, Kalydeco 50 mg granules in sachet, and Kalydeco 75 mg granules in sachet</u>

Pack size of 56 sachets (contains 4 individual wallets with 14 sachets per wallet).

Kalydeco 13.4 mg granules in sachet, Kalydeco 59.5 mg granules in sachet, and Kalydeco 75 mg granules in sachet

Pack size of 28 sachets (contains 4 individual wallets with 7 sachets per wallet).

Not all pack sizes may be marketed.

6.6 Special precautions for disposal and other handling

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

7. MARKETING AUTHORISATION HOLDER

Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/12/782/003

EU/1/12/782/004

EU/1/12/782/006

EU/1/12/782/008

EU/1/12/782/009

EU/1/12/782/010

EU/1/12/782/011

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 23 July 2012 Date of latest renewal: 29 April 2022

10. DATE OF REVISION OF THE TEXT

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

ANNEX II

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

A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer(s) responsible for batch release

Almac Pharma Services (Ireland) Limited Finnabair Industrial Estate Dundalk Co. Louth A91 P9KD Ireland

Almac Pharma Services Limited Seagoe Industrial Estate Craigavon Northern Ireland BT63 5UA United Kingdom

Millmount Healthcare Limited Block-7, City North Business Campus Stamullen Co. Meath K32 YD60 Ireland

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

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

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

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

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

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

• Risk management plan (RMP)

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

An updated RMP should be submitted:

• At the request of the European Medicines Agency;

		being reached	

Whenever the risk management system is modified, especially as the result of new information

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

OUTER CARTON FOR BLISTER – 56-TABLET PACK 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 150 mg film-coated tablets ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 150 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 56 tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use Instructions for use Take with fat-containing food. Do not break, chew or dissolve the tablets. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT 6. OF THE SIGHT AND REACH OF CHILDREN Keep out of the sight and reach of children. 7. OTHER SPECIAL WARNING(S), IF NECESSARY

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

8.

EXP

EXPIRY DATE

9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	ex Pharmaceuticals (Ireland) Limited 49, Block 5, Northwood Court, Northwood Crescent, in 9, D09 T665, and
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/12/782/002
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Kalyo	deco 150 mg tablets
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	arcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS
BLISTERS – 56-TABLET PACK
1. NAME OF THE MEDICINAL PRODUCT
Kalydeco 150 mg tablets ivacaftor
2. NAME OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals
3. EXPIRY DATE
EXP
4. BATCH NUMBER
Lot
5. OTHER

PARTICULARS TO APPEAR ON THE OUTER PACKAGING **OUTER CARTON FOR BLISTER CARD – 28-TABLET PACK** 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 150 mg film-coated tablets ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 150 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 28 tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use Instructions for use Always take this medicine exactly as your doctor has told you to. Take with fat-containing food. Do not break, chew or dissolve the tablets. Insert tab below to close

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

Keep out of the sight and reach of children.

Open

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8.	EXPIRY DATE
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9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	2 Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, 19, D09 T665, 1
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/	12/782/005
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Kalyd	eco 150 mg tablets
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D bar	rcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGING BLISTER CARD – 28-TABLET PACK 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 150 mg film-coated tablets ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 150 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 7 tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use Instructions for use Always take this medicine exactly as your doctor has told you to. Take with fat-containing food. Do not break, chew or dissolve the tablets.

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

Keep out of the sight and reach of children.

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9.	SPECIAL STORAGE CONDITIONS
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11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 49	Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, 9, D09 T665,
12.	MARKETING AUTHORISATION NUMBER(S)
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13. Lot 14.	BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE
13. Lot 14.	BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE
13. Lot 14. 15.	BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE INFORMATION IN BRAILLE

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
BLISTERS – 28-TABLET PACK	
1. NAME OF THE MEDICINAL PRODUCT	
Kalydeco 150 mg tablets ivacaftor	
2. NAME OF THE MARKETING AUTHORISATION HOLDER	
Vertex	
3. EXPIRY DATE	
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4. BATCH NUMBER	
Lot	
5. OTHER	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING **OUTER CARTON FOR BLISTER CARD – 28-TABLET PACK** 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 75 mg film-coated tablets ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 75 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 28 tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use Instructions for use Always take this medicine exactly as your doctor has told you to. Take with fat-containing food. Do not break, chew or dissolve the tablets. Insert tab below to close

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

Keep out of the sight and reach of children.

Open

8.	EXPIRY DATE
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9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	2 Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, 1 9, D09 T665, 1
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/	12/782/007
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Kalyd	eco 75 mg tablets
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D bar	rcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

BLISTER CARD – 28-TABLET PACK 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 75 mg film-coated tablets ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE(S) Each tablet contains 75 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. 4. PHARMACEUTICAL FORM AND CONTENTS 7 tablets 5. METHOD AND ROUTE(S) OF ADMINISTRATION Read the package leaflet before use. Oral use Instructions for use Always take this medicine exactly as your doctor has told you to. Take with fat-containing food. Do not break, chew or dissolve the tablets.

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGING

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

Keep out of the sight and reach of children.

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9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS
	OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
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12.	MARKETING AUTHORISATION NUMBER(S)
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13.	BATCH NUMBER
Lot	
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
17.	UNIQUE IDENTIFIER – 2D BARCODE
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA

MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS	
BLISTERS – 28-TABLET PACK	
1. NAME OF THE MEDICINAL PRODUCT	
TWINE OF THE MEETING OF	
Kalydeco 75 mg tablets	
ivacaftor	
2. NAME OF THE MARKETING AUTHORISATION HOLDER	
Vertex	
3. EXPIRY DATE	
EXP	
4. BATCH NUMBER	
Lot	
5. OTHER	

PARTICULARS TO APPEAR ON THE OUTER PACKAGING
OUTER CARTON FOR BOTTLE
1. NAME OF THE MEDICINAL PRODUCT
Kalydeco 150 mg film-coated tablets ivacaftor
2. STATEMENT OF ACTIVE SUBSTANCE(S)
Each tablet contains 150 mg of ivacaftor.
3. LIST OF EXCIPIENTS
Contains lactose.
See leaflet for further information.
4. PHARMACEUTICAL FORM AND CONTENTS
56 tablets
5. METHOD AND ROUTE(S) OF ADMINISTRATION
Read the package leaflet before use.
Oral use
Instructions for use
Take with fat-containing food.
Do not break, chew or dissolve the tablets.
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN
Keep out of the sight and reach of children.
7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
EXP

9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	x Pharmaceuticals (Ireland) Limited 49, Block 5, Northwood Court, Northwood Crescent, n 9, D09 T665, d
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/	12/782/001
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Kalyd	leco 150 mg tablets
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D ba	rcode carrying the unique identifier included.
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

PARTICULARS TO APPEAR ON THE IMMEDIATE PACKAGING	
BOTTLE LABEL	
1. NAME OF THE MEDICINAL PRODUCT	
Kalydeco 150 mg film-coated tablets ivacaftor	
2. STATEMENT OF ACTIVE SUBSTANCE(S)	
Each tablet contains 150 mg of ivacaftor.	
3. LIST OF EXCIPIENTS	
Contains lactose.	
See leaflet for further information.	
4. PHARMACEUTICAL FORM AND CONTENTS	
56 tablets	
5. METHOD AND ROUTE(S) OF ADMINISTRATION	
Read the package leaflet before use.	
Oral use	
6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN	
Keep out of the sight and reach of children.	
7. OTHER SPECIAL WARNING(S), IF NECESSARY	
8. EXPIRY DATE	
EXP	
9. SPECIAL STORAGE CONDITIONS	

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/12/782/001
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
17. UNIQUE IDENTIFIER – 2D BARCODE
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

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

10.

APPROPRIATE

PARTICULARS TO APPEAR ON THE OUTER PACKAGING OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 13.4 mg granules in sachet ivacaftor

For patients aged 1 to less than 2 months

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 13.4 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

28 sachets

4 individual wallets with 7 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

7.	OTHER SPECIAL WARNING(S), IF NECESSARY
8.	EXPIRY DATE
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9.	SPECIAL STORAGE CONDITIONS
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10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF
	APPROPRIATE
	THO THE TE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
T. 7	
	Pharmaceuticals (Ireland) Limited
	9, Block 5, Northwood Court, Northwood Crescent, 19, D09 T665,
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Irciano	
12.	MARKETING AUTHORISATION NUMBER(S)
	MARKETING AUTHORISATION NUMBER(S) 2/782/010
EU/1/1	.2/782/010
EU/1/1	.2/782/010
EU/1/7	.2/782/010
EU/1/2 13. Lot	2/782/010 BATCH NUMBER
EU/1/7	.2/782/010
EU/1/2 13. Lot	2/782/010 BATCH NUMBER
EU/1/2 13. Lot	2/782/010 BATCH NUMBER
EU/1/2 13. Lot 14.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY
EU/1/1 13. Lot 14.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE
EU/1/2 13. Lot 14.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY
EU/1/1 13. Lot 14. 15.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE INFORMATION IN BRAILLE
EU/1/1 13. Lot 14. 15.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE
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EU/1/1 13. Lot 14. 15.	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE INFORMATION IN BRAILLE
13. Lot 14. 15. Kalydo	2/782/010 BATCH NUMBER GENERAL CLASSIFICATION FOR SUPPLY INSTRUCTIONS ON USE INFORMATION IN BRAILLE 200 13.4 mg granules

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC SN

NN

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING

WALLET FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 13.4 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 13.4 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

7 sachets

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Use all 7 days' doses before starting a new wallet.

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

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

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY
8. EXPIRY DATE
9. SPECIAL STORAGE CONDITIONS
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/12/782/010
13. BATCH NUMBER
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
17. UNIQUE IDENTIFIER – 2D BARCODE
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS
SACHETS
1. NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION
Kalydeco 13.4 mg granules ivacaftor Oral use
2. METHOD OF ADMINISTRATION
3. EXPIRY DATE
EXP
4. BATCH NUMBER
Lot
5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT
6. OTHER
Vertex Pharmaceuticals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 13.4 mg granules in sachet ivacaftor For patients aged 2 to less than 4 months

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 13.4 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

56 sachets

4 individual wallets with 14 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

7.	OTHER SPECIAL WARNING(S), IF NECESSARY	
8.	EXPIRY DATE	
EXP		
9.	SPECIAL STORAGE CONDITIONS	
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE	
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER	
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland		
12.	MARKETING AUTHORISATION NUMBER(S)	
EU/1/12/782/011		
13.	BATCH NUMBER	
Lot		
14.	GENERAL CLASSIFICATION FOR SUPPLY	
15.	INSTRUCTIONS ON USE	
16.	INFORMATION IN BRAILLE	
Kalydeco 13.4 mg granules		
17.	UNIQUE IDENTIFIER – 2D BARCODE	
2D ba	2D barcode carrying the unique identifier included.	

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC SN

NN

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING WALLET FOR SACHET 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 13.4 mg granules in sachet ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE Each sachet of granules contains 13.4 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. PHARMACEUTICAL FORM AND CONTENTS 4. Granules in sachet 14 sachets METHOD AND ROUTE(S) OF ADMINISTRATION 5. Read the package leaflet before use. Oral use Instructions for use Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely. Use within one hour after mixing, just before or after a fat-containing meal or snack. Use all 7 days' doses before starting a new wallet. Morning

Evening

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

Keep out of the sight and reach of children.	
7.	OTHER SPECIAL WARNING(S), IF NECESSARY
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8.	EXPIRY DATE
L	
9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland	
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/12/782/011
13.	BATCH NUMBER
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
17.	UNIQUE IDENTIFIER – 2D BARCODE
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA

SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT

OF THE SIGHT AND REACH OF CHILDREN

6.

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS		
SACHETS		
1.	NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION	
Kalydeco 13.4 mg granules ivacaftor Oral use		
2.	METHOD OF ADMINISTRATION	
3.	EXPIRY DATE	
EXP		
4.	BATCH NUMBER	
Lot		
5.	CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT	
6.	OTHER	

Vertex Pharmaceuticals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 25 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 25 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

56 sachets

4 individual wallets with 14 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

7.	OTHER SPECIAL WARNING(S), IF NECESSARY
8.	EXPIRY DATE
0.	EATIKI DATE
EXP	
9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS
	OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF
	APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex	Pharmaceuticals (Ireland) Limited
	9, Block 5, Northwood Court, Northwood Crescent,
Dublin Ireland	19, D09 T665,
ireiand	
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/1	2/782/006
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
15.	INSTRUCTIONS ON USE
15.	INSTRUCTIONS ON USE INFORMATION IN BRAILLE
16.	INFORMATION IN BRAILLE
16.	
16.	INFORMATION IN BRAILLE
16.	INFORMATION IN BRAILLE

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC

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PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING WALLET FOR SACHET 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 25 mg granules in sachet ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE Each sachet of granules contains 25 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. PHARMACEUTICAL FORM AND CONTENTS 4. Granules in sachet 14 sachets METHOD AND ROUTE(S) OF ADMINISTRATION 5. Read the package leaflet before use. Oral use Instructions for use Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely. Use within one hour after mixing, just before or after a fat-containing meal or snack. Use all 7 days' doses before starting a new wallet. Morning Evening

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

Keep out of the sight and reach of children.	
7. OTHER SPECIAL WARNING(S), IF NECESSARY	
8. EXPIRY DATE	
9. SPECIAL STORAGE CONDITIONS	
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE	
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER	
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland	
12. MARKETING AUTHORISATION NUMBER(S)	
EU/1/12/782/006	
13. BATCH NUMBER	
14. GENERAL CLASSIFICATION FOR SUPPLY	
15. INSTRUCTIONS ON USE	
16. INFORMATION IN BRAILLE	
17. UNIQUE IDENTIFIER – 2D BARCODE	
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA	

SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT

OF THE SIGHT AND REACH OF CHILDREN

6.

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS		
SACHETS		
1. NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION		
Kalydeco 25 mg granules ivacaftor Oral use		
2. METHOD OF ADMINISTRATION		
3. EXPIRY DATE		
EXP		
4. BATCH NUMBER		
Lot		
5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT		
6. OTHER		

Vertex Pharmaceuticals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 50 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 50 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

56 sachets

4 individual wallets with 14 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

8. EXPIRY DATE	
EXP	
9. SPECIAL STORAGE CONDITIONS	
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE	
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER	
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland	
12. MARKETING AUTHORISATION NUMBER(S)	
EU/1/12/782/003	
13. BATCH NUMBER	
Lot	
14. GENERAL CLASSIFICATION FOR SUPPLY	
15. INSTRUCTIONS ON USE	
16. INFORMATION IN BRAILLE	
Kalydeco 50 mg granules	
17. UNIQUE IDENTIFIER – 2D BARCODE	
2D barcode carrying the unique identifier included.	
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA	
PC SN NN	

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING WALLET FOR SACHET 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 50 mg granules in sachet ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE Each sachet of granules contains 50 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. PHARMACEUTICAL FORM AND CONTENTS 4. Granules in sachet 14 sachets METHOD AND ROUTE(S) OF ADMINISTRATION 5. Read the package leaflet before use. Oral use Instructions for use Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely. Use within one hour after mixing, just before or after a fat-containing meal or snack. Use all 7 days' doses before starting a new wallet. Morning Evening

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

Keep out of the sight and reach of children.	
7.	OTHER SPECIAL WARNING(S), IF NECESSARY
8.	EXPIRY DATE
9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	x Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, n 9, D09 T665, d
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/	12/782/003
13.	BATCH NUMBER
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
17.	UNIQUE IDENTIFIER – 2D BARCODE
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA

SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT

OF THE SIGHT AND REACH OF CHILDREN

6.

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS	
C A CI	
SACI	HETS
1.	NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION
Valve	dono 50 ma granulas
ivaca	deco 50 mg granules
Oral ı	
J 1 4 1 1	
2.	METHOD OF ADMINISTRATION
3.	EXPIRY DATE
EXD	
EXP	
4.	BATCH NUMBER
Lot	
5.	CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT
6.	OTHER
Vonto	x Pharmaceuticals
v CI lC	A I Hallilacculicals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 59.5 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 59.5 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

28 sachets

4 individual wallets with 7 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/12/782/008
44 D. J. T. C. V. N. V. V. D. D. D.
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
Kalydeco 59.5 mg granules
17. UNIQUE IDENTIFIER – 2D BARCODE
2D barcode carrying the unique identifier included.
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING

WALLET FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 59.5 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 59.5 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

7 sachets

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Use all 7 days' doses before starting a new wallet.

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

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

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8.	EXPIRY DATE		
9.	SPECIAL STORAGE CONDITIONS		
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE		
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER		
Unit 4	x Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, n 9, D09 T665, d		
12.	MARKETING AUTHORISATION NUMBER(S)		
EU/1/	12/782/008		
13.	BATCH NUMBER		
14.	GENERAL CLASSIFICATION FOR SUPPLY		
15.	INSTRUCTIONS ON USE		
16.	INFORMATION IN BRAILLE		
17.	UNIQUE IDENTIFIER – 2D BARCODE		
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA		

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS		
SACI	HETS	
1.	NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION	
Kalyd ivacaf Oral u		
2.	METHOD OF ADMINISTRATION	
3.	EXPIRY DATE	
EXP		
4.	BATCH NUMBER	
Lot		
5.	CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT	
-		
6.	OTHER	

Vertex Pharmaceuticals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON FOR SACHET

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 75 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 75 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

56 sachets

4 individual wallets with 14 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE
EXP
9. SPECIAL STORAGE CONDITIONS
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland
12. MARKETING AUTHORISATION NUMBER(S)
EU/1/12/782/004
13. BATCH NUMBER
Lot
14. GENERAL CLASSIFICATION FOR SUPPLY
15. INSTRUCTIONS ON USE
16. INFORMATION IN BRAILLE
Kalydeco 75 mg granules
17. UNIQUE IDENTIFIER – 2D BARCODE
2D barcode carrying the unique identifier included.
18. UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING WALLET FOR SACHET 1. NAME OF THE MEDICINAL PRODUCT Kalydeco 75 mg granules in sachet ivacaftor 2. STATEMENT OF ACTIVE SUBSTANCE Each sachet of granules contains 75 mg of ivacaftor. 3. LIST OF EXCIPIENTS Contains lactose. See leaflet for further information. PHARMACEUTICAL FORM AND CONTENTS 4. Granules in sachet 14 sachets METHOD AND ROUTE(S) OF ADMINISTRATION 5. Read the package leaflet before use. Oral use Instructions for use Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely. Use within one hour after mixing, just before or after a fat-containing meal or snack. Use all 7 days' doses before starting a new wallet. Morning Evening

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

Keep	Keep out of the sight and reach of children.		
7.	OTHER SPECIAL WARNING(S), IF NECESSARY		
8.	EXPIRY DATE		
9.	SPECIAL STORAGE CONDITIONS		
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE		
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER		
Unit 4	R Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, 19, D09 T665, d		
12.	MARKETING AUTHORISATION NUMBER(S)		
EU/1/	12/782/004		
13.	BATCH NUMBER		
14.	GENERAL CLASSIFICATION FOR SUPPLY		
15.	INSTRUCTIONS ON USE		
16.	INFORMATION IN BRAILLE		
17.	UNIQUE IDENTIFIER – 2D BARCODE		
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA		

SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT

OF THE SIGHT AND REACH OF CHILDREN

6.

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS		
SACH	ETS	
1.	NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION	
	co 75 mg granules	
2.	METHOD OF ADMINISTRATION	
3.	EXPIRY DATE	
EXP		
4.	BATCH NUMBER	
Lot		
5.	CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT	
J.	CONTENTS DI MEIGHI, DI VOLUME ON DI UMI	
6.	OTHER	

Vertex Pharmaceuticals

PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON FOR SACHET – 28-COUNT

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 75 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 75 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

28 sachets

4 individual wallets with 7 sachets per wallet

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Lift here to open

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

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8.	EXPIRY DATE
EXP	
9.	SPECIAL STORAGE CONDITIONS
OR W	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS (ASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF OPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	Pharmaceuticals (Ireland) Limited 9, Block 5, Northwood Court, Northwood Crescent, 19, D09 T665,
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1/1	12/782/009
13.	BATCH NUMBER
Lot	
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
Kalydo	eco 75 mg granules
17.	UNIQUE IDENTIFIER – 2D BARCODE
2D baı	code carrying the unique identifier included.
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA
PC SN NN	

PARTICULARS TO APPEAR ON THE INTERMEDIATE PACKAGING

WALLET FOR SACHET – 7-COUNT

1. NAME OF THE MEDICINAL PRODUCT

Kalydeco 75 mg granules in sachet ivacaftor

2. STATEMENT OF ACTIVE SUBSTANCE

Each sachet of granules contains 75 mg of ivacaftor.

3. LIST OF EXCIPIENTS

Contains lactose.

See leaflet for further information.

4. PHARMACEUTICAL FORM AND CONTENTS

Granules in sachet

7 sachets

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.

Oral use

Instructions for use

Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid that is at or below room temperature and consume it completely.

Use within one hour after mixing, just before or after a fat-containing meal or snack.

Use all 7 days' doses before starting a new wallet.

Mon. Tue. Wed. Thu. Fri. Sat. Sun.

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

Keep out of the sight and reach of children.

7.	OTHER SPECIAL WARNING(S), IF NECESSARY
8.	EXPIRY DATE
9.	SPECIAL STORAGE CONDITIONS
10.	SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE
11.	NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER
Unit 4	ex Pharmaceuticals (Ireland) Limited 49, Block 5, Northwood Court, Northwood Crescent, in 9, D09 T665, ad
12.	MARKETING AUTHORISATION NUMBER(S)
EU/1	/12/782/009
13.	BATCH NUMBER
14.	GENERAL CLASSIFICATION FOR SUPPLY
15.	INSTRUCTIONS ON USE
16.	INFORMATION IN BRAILLE
17.	UNIQUE IDENTIFIER – 2D BARCODE
18.	UNIQUE IDENTIFIER – HUMAN READABLE DATA

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS
SACHETS – 28-COUNT
1. NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION
Kalydeco 75 mg granules ivacaftor Oral use
2. METHOD OF ADMINISTRATION
3. EXPIRY DATE
EXP
4. BATCH NUMBER
Lot
5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT
J. CONTENTS D1 WEIGHT, D1 VOLUME OR D1 UNIT
6. OTHER

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Kalydeco 75 mg film-coated tablets Kalydeco 150 mg film-coated tablets ivacaftor

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

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

What is in this leaflet

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

1. What Kalydeco is and what it is used for

Kalydeco contains the active substance ivacaftor. Ivacaftor acts at the level of the cystic fibrosis transmembrane conductance regulator (CFTR), a protein that forms a channel at the cell surface that allows the movement of particles such as chloride in and out of the cell. Due to mutations in the *CFTR* gene (see below), chloride movement is reduced in those with cystic fibrosis (CF). Ivacaftor helps certain abnormal CFTR proteins open more often to improve chloride movement in and out of the cell.

Kalydeco tablets are indicated:

- As monotherapy for patients aged 6 years and older and weighing 25 kg or more with cystic fibrosis (CF) who have an R117H CFTR mutation or one of the following gating mutations in the CFTR gene: G551D, G1244E, G1349D, G178R, G551S, S1251N, S1255P, S549N or S549R.
- In combination with tezacaftor/ivacaftor tablets for patients aged 6 years and older with CF who have two F508del mutations in the CFTR gene (homozygous for the F508del mutation) or who have an F508del mutation and certain other second mutations that result in reduced amount and/or function of the CFTR protein (heterozygous for the F508del mutation with a residual function (RF) mutation). If you have been prescribed Kalydeco to be taken with tezacaftor/ivacaftor, read the package leaflet of the latter. It contains important information about how to take these two medicines.
- In combination with ivacaftor/tezacaftor/elexacaftor tablets for patients aged 6 years and over who have CF, with at least one mutation in the *CFTR* gene that is responsive to Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor. If you have been prescribed Kalydeco to be taken with ivacaftor/tezacaftor/elexacaftor, read the package leaflet of the latter. It contains important information about how to take these two medicines.

2. What you need to know before you take Kalydeco

Do not take Kalydeco

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

Warnings and precautions

Talk to your doctor or pharmacist before taking Kalydeco.

- Talk to your doctor if you have liver problems or have previously had them. Your doctor may need to adjust your dose.
- Increased liver enzymes in the blood have been seen in some people receiving Kalydeco (alone or in combination with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor). Tell your doctor right away if you have any of these symptoms, which may be a sign of liver problems:
 - Pain or discomfort in the upper right stomach (abdominal) area
 - Yellowing of the skin or the white part of the eyes
 - Loss of appetite
 - Nausea or vomiting
 - Dark urine
- Your doctor will do some blood tests to check your liver before and during treatment, particularly during the first year and especially if your blood tests showed high liver enzymes in the past.
- Depression (including suicidal thoughts and behaviours) has been reported in patients while taking Kalydeco, mainly in a combination regimen with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor, usually starting within the first three months of treatment. Talk to a doctor straightaway if you (or someone taking this medicine) experience any of the following symptoms: sad or altered mood, anxiety, feelings of emotional discomfort or thoughts of harming or killing yourself, which may be signs of depression.
- Talk to your doctor if you have kidney problems or have previously had them.
- If you have two Class I mutations (mutations known not to make CFTR protein), you should not take Kalydeco, as you are not expected to respond to this medicine.
- Kalydeco is not recommended if you have undergone an organ transplant.
- Talk to your doctor if you are using hormonal contraception for example, women using the contraceptive pill. This may mean you are more likely to get a rash while taking Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor.
- Abnormality of the eye lens (cataract) without any effect on vision has been noted in some children and adolescents treated with Kalydeco (alone or in combination with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor). Your doctor may perform some eye examinations prior to and during treatment.
- Kalydeco should only be used if you have one of the mutations in the *CFTR* gene indicated in section 1 (What Kalydeco is and what it is used for).

Children and adolescents

Do not give this medicine to children under 1 month of age as it is not known if ivacaftor is safe and effective in these children.

Do not give this medicine in combination with tezacaftor/ivacaftor to children under 6 years of age or in combination with ivacaftor/tezacaftor/elexacaftor to children under 2 years of age as it is not known if they are safe and effective for them.

Other medicines and Kalydeco

Tell your doctor or pharmacist if you are using, have recently used or might use any other medicines. Some medicines can affect how Kalydeco works or make side effects more likely. In particular, tell your doctor if you are taking any of the medicines listed below. Your doctor may decide to adjust your dose or that you need extra check-ups.

- **Antifungal medicines** (used for the treatment of fungal infections). These include fluconazole, itraconazole, ketoconazole, posaconazole, and voriconazole.
- **Antibiotic medicines** (used for the treatment of bacterial infections). These include clarithromycin, erythromycin, rifabutin, rifampicin, and telithromycin.
- **Epilepsy medicines** (used for the treatment of epileptic seizures or fits). These include carbamazepine, phenobarbital, and phenytoin.
- **Herbal medicines.** These include St. John's wort (*Hypericum perforatum*).
- **Immunosuppressants** (used after an organ transplantation). These include ciclosporin, everolimus, sirolimus, and tacrolimus.
- Cardiac glycosides (used for the treatment of some heart conditions). These include digoxin.
- Anticoagulant medicines (used to prevent blood clots). These include warfarin.
- Medicines for diabetes. These include glimepiride and glipizide.
- Medicines for lowering blood pressure. These include verapamil.

Kalydeco with food and drink

Avoid food or drink containing grapefruit during treatment as these may increase the side effects of Kalvdeco by increasing the amount of ivacaftor in your body.

Pregnancy and breast-feeding

If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor for advice before taking this medicine. It may be better to avoid using Kalydeco during pregnancy, if possible, and your doctor will help you decide what is best for you and your child.

Ivacaftor passes into breast milk. If you plan to breast-feed, ask your doctor for advice before taking Kalydeco. Your doctor will decide whether to recommend that you stop breast-feeding or for you to stop ivacaftor therapy. Your doctor will take into account the benefit of breast-feeding for the child and the benefit of therapy for you.

Driving and using machines

Kalydeco can make you dizzy. If you feel dizzy, do not drive, cycle or use machines.

Kalydeco contains lactose and sodium

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

Kalydeco contains less than 1 mmol sodium (23 mg) per dose, that is to say essentially 'sodium-free'.

3. How to take Kalydeco

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

Your doctor will determine which medicine and dose is right for you.

Kalydeco dosing recommendations are provided in Table 1.

Table 1: Dosing recommendations

Age/weight	Morning dose	Evening dose		
Kalydeco as monotherapy				
6 years and older, ≥ 25 kg	One Kalydeco 150 mg tablet	One Kalydeco 150 mg tablet		
Kalydeco in combination with	Kalydeco in combination with tezacaftor/ivacaftor			
6 years to less than 12 years, < 30 kg	One tezacaftor 50 mg/ivacaftor 75 mg tablet	One Kalydeco 75 mg tablet		
6 years to less than 12 years, ≥ 30 kg	One tezacaftor 100 mg/ivacaftor 150 mg tablet	One Kalydeco 150 mg tablet		
12 years and older	One tezacaftor 100 mg/ivacaftor 150 mg tablet	One Kalydeco 150 mg tablet		
Kalydeco in combination with	Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor			
6 years to less than 12 years, < 30 kg	Two ivacaftor 37.5 mg/tezacaftor 25 mg/elexacaftor 50 mg tablets	One Kalydeco 75 mg tablet		
6 years to less than 12 years,	Two ivacaftor 75 mg/tezacaftor	One Kalydeco 150 mg		
≥ 30 kg	50 mg/elexacaftor 100 mg tablets	tablet		
12 years and older	Two ivacaftor 75 mg/tezacaftor 50 mg/elexacaftor 100 mg tablets	One Kalydeco 150 mg tablet		

Take the morning and evening doses approximately 12 hours apart with food that contains fat.

You must keep using all other medicines you use, unless your doctor tells you to stop using any.

If you have liver problems, either moderate or severe, your doctor may need to reduce the dose of your tablets, because your liver will not clear the medicine as fast as in people who have normal liver function.

This medicine is for oral use.

Swallow the tablet whole. Do not break, chew or dissolve the tablets. Take Kalydeco tablets with food that contains fat.

Meals or snacks that contain fat include those prepared with butter or oils or those containing eggs. Other fat-containing foods are:

- Cheese, whole milk, whole-milk dairy products, yogurt, chocolate
- Meats, oily fish
- Avocados, hummus, soy-based products (tofu)
- Nuts, fat-containing nutritional bars or drinks

If you take more Kalydeco than you should

You may experience side effects, including those mentioned in section 4 below. If so, contact your doctor or pharmacist to ask for advice. If possible, have your medicine and this leaflet with you.

If you forget to take Kalydeco

Take the missed dose if less than 6 hours have passed since the time you missed the dose. Otherwise, wait until your next scheduled dose as you normally would. Do not take a double dose to make up for a forgotten dose.

If you stop taking Kalydeco

Take Kalydeco for as long as your doctor recommends. Do not stop unless your doctor advises you to.

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

4 Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Serious side effects

Stomach (abdominal) ache and increased liver enzymes in the blood.

Possible signs of liver problems

Increased liver enzymes in the blood are common in patients with CF and have also been reported in patients taking Kalydeco alone or in combination with tezacaftor/ivacaftor or ivacaftor/tezacaftor/elexacaftor.

In patients taking Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor, liver damage and worsening of liver function in people with severe liver disease has been reported. The worsening of liver function can be serious and may require transplantation.

These may be signs of liver problems:

- Pain or discomfort in the upper right area of the stomach (abdominal) area
- Yellowing of the skin or white part of the eyes
- Loss of appetite
- Nausea or vomiting
- Dark urine

Depression

Signs of this include sad or altered mood, anxiety, feelings of emotional discomfort.

Tell your doctor straight away if you have any of these symptoms.

Very common side effects (may affect more than 1 in 10 people)

- Upper respiratory tract infection (the common cold), including sore throat and nasal congestion
- Headache
- Dizziness
- Diarrhoea
- Stomach or abdominal pain
- Changes in the type of bacteria in mucus
- Increased liver enzymes (signs of stress on the liver)
- Rash

Common side effects (may affect up to 1 in 10 people)

- Runny nose
- Ear pain, ear discomfort
- Ringing in the ears

- Redness inside the ear
- Inner ear disorder (feeling dizzy or spinning)
- Sinus problems (sinus congestion)
- Redness in the throat
- Breast mass
- Feeling sick (nausea)
- Flu
- Low blood sugar (hypoglycaemia)
- Abnormal breathing (shortness of breath or difficulty breathing)
- Wind (flatulence)
- Spots (acne)
- Itchy skin
- Increased creatine phosphokinase (sign of muscle breakdown) seen in blood tests

Uncommon side effects (may affect up to 1 in 100 people)

- Ear congestion
- Breast inflammation
- Enlargement of the breast in males
- Nipple changes or pain
- Wheezing
- Increased blood pressure

Not known (frequency cannot be estimated from the available data)

- Damage to the liver (liver injury)
- Raised bilirubin measurement (liver blood test)

Additional side effects in children and adolescents

Side effects seen in children and adolescents are similar to those observed in adults. However, increased liver enzymes in the blood are more frequently seen in young children.

Reporting of side effects

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

5. How to store Kalydeco

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

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

This medicine does not require any special storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Kalydeco contains

The active substance is ivacaftor.

Kalydeco 75 mg film-coated tablets

Each film-coated tablet contains 75 mg of ivacaftor.

Kalydeco 150 mg film-coated tablets

Each film-coated tablet contains 150 mg of ivacaftor.

The other ingredients are:

- Tablet core: cellulose, microcrystalline, lactose monohydrate, hypromellose acetate succinate, croscarmellose sodium, sodium laurilsulfate (E487), silica, colloidal anhydrous, and magnesium stearate.
- Tablet film coating: polyvinyl alcohol, titanium dioxide (E171), macrogol (PEG 3350), talc, indigo carmine aluminium lake (E132) and carnauba wax.
- Printing ink: shellac, iron oxide black (E172), propylene glycol (E1520) and ammonia solution, concentrated.

See the end of section 2 – Kalydeco contains lactose and sodium.

What Kalydeco looks like and contents of the pack

Kalydeco 75 mg film-coated tablets are light blue, capsule-shaped, $12.7 \text{ mm} \times 6.8 \text{ mm}$, and printed with "V 75" in black ink on one side and plain on the other.

The following pack sizes are available:

• Blister card pack containing 28 film-coated tablets

Kalydeco 150 mg film-coated tablets are light blue, capsule-shaped, $16.5 \text{ mm} \times 8.4 \text{ mm}$, and printed with "V 150" in black ink on one side and plain on the other.

The following pack sizes are available:

- Blister card pack containing 28 film-coated tablets
- Blister pack containing 56 film-coated tablets
- Bottle containing 56 film-coated tablets

Marketing Authorisation Holder

Vertex Pharmaceuticals (Ireland) Limited Unit 49, Block 5, Northwood Court, Northwood Crescent, Dublin 9, D09 T665, Ireland

Tel: +353 (0)1 761 7299

Manufacturer

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Almac Pharma Services Limited Seagoe Industrial Estate Craigavon Northern Ireland BT63 5UA United Kingdom

Millmount Healthcare Limited Block-7, City North Business Campus Stamullen Co. Meath K32 YD60 Ireland

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

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This leaflet was last revised in

Other sources of information

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

Package leaflet: Information for the patient

Kalydeco 13.4 mg granules in sachet Kalydeco 25 mg granules in sachet Kalydeco 50 mg granules in sachet Kalydeco 59.5 mg granules in sachet Kalydeco 75 mg granules in sachet ivacaftor

Read all of this leaflet carefully before your child starts taking this medicine because it contains important information for your child.

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

What is in this leaflet

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

1. What Kalydeco is and what it is used for

Kalydeco contains the active substance ivacaftor. Ivacaftor acts at the level of the cystic fibrosis transmembrane conductance regulator (CFTR), a protein that forms a channel at the cell surface that allows the movement of particles such as chloride in and out of the cell. Due to mutations in the *CFTR* gene (see below), chloride movement is reduced in those with cystic fibrosis (CF). Ivacaftor helps certain abnormal CFTR proteins open more often to improve chloride movement in and out of the cell.

Kalydeco granules are indicated:

- As monotherapy for the treatment of babies and children aged 1 month and older and weighing 3 kg to less than 25 kg with cystic fibrosis (CF) who have an *R117H CFTR* mutation or one of the following gating mutations in the *CFTR* gene: *G551D*, *G1244E*, *G1349D*, *G178R*, *G551S*, *S1251N*, *S1255P*, *S549N* or *S549R*.
- In combination with ivacaftor/tezacaftor/elexacaftor granules for patients aged 2 to 6 years who have CF, with at least one mutation in the *CFTR* gene that is responsive to Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor. If you have been prescribed Kalydeco to be taken with ivacaftor/tezacaftor/elexacaftor, read the package leaflet of the latter. It contains important information about how to take these two medicines.

2. What you need to know before your child takes Kalydeco

Do not give your child Kalydeco

• if your child is allergic to ivacaftor or any of the other ingredients of this medicine (listed in section 6).

Warnings and precautions

Talk to your child's doctor before your child takes Kalydeco.

- Talk to your child's doctor if your child has liver problems or has had them previously. Your child's doctor may need to adjust your child's dose.
- Increased liver enzymes in the blood have been seen in some people receiving Kalydeco (alone or in combination with ivacaftor/tezacaftor/elexacaftor). Tell your child's doctor right away if your child has any of these symptoms, which may be a sign of liver problems:
 - Pain or discomfort in the upper right stomach (abdominal) area
 - Yellowing of the skin or the white part of the eyes
 - Loss of appetite
 - Nausea or vomiting
 - Dark urine
- Your child's doctor will do some blood tests to check your child's liver before and during treatment, particularly during the first year and especially if blood tests showed high liver enzymes in the past.
- Depression (including suicidal thoughts and behaviours) has been reported in patients while taking Kalydeco, mainly in a combination regimen with ivacaftor/tezacaftor/elexacaftor, usually starting within the first three months of treatment. Talk to a doctor straightaway if you (or someone taking this medicine) experience any of the following symptoms: sad or altered mood, anxiety, feelings of emotional discomfort or thoughts of harming or killing yourself, which may be signs of depression.
- Talk to your child's doctor if you have been told your child has kidney problems or has previously had them.
- If you have two Class I mutations (mutations known not to make CFTR protein), you should not take Kalydeco, as you are not expected to respond to this medicine.
- Kalydeco is not recommended for patients who have undergone an organ transplant.
- Abnormality of the eye lens (cataract) without any effect on vision has been noted in some children and adolescents during treatment (alone or in combination with ivacaftor/tezacaftor/elexacaftor). Your child's doctor may perform some eye examinations prior to and during treatment.
- Kalydeco should only be used if your child has one of the mutations in the *CFTR* gene indicated in section 1 (What Kalydeco is and what it is used for).

Children

Do not give this medicine to children under 1 month of age as it is not known if ivacaftor is safe and effective in these children.

Do not give this medicine in combination with ivacaftor/tezacaftor/elexacaftor to children under 2 years of age as it is not known if they are safe and effective for them.

Other medicines and Kalydeco

Tell your child's doctor or pharmacist if your child is using, has recently used or might use any other medicines. Some medicines can affect how Kalydeco works or make side effects more likely. In

particular, tell your child's doctor if your child is taking any of the medicines listed below. Your child's doctor may decide to adjust your child's dose or if extra check-ups are needed.

- **Antifungal medicines** (used for the treatment of fungal infections). These include fluconazole, itraconazole, ketoconazole, posaconazole, and voriconazole.
- **Antibiotic medicines** (used for the treatment of bacterial infections). These include clarithromycin, erythromycin, rifabutin, rifampicin and telithromycin.
- **Epilepsy medicines** (used for the treatment of epileptic seizures or fits). These include carbamazepine, phenobarbital, and phenytoin.
- **Herbal medicines**. These include St. John's wort (*Hypericum perforatum*).
- **Immunosuppressants** (used after an organ transplantation). These include ciclosporin, everolimus, sirolimus, and tacrolimus.
- Cardiac glycosides (used for the treatment of some heart conditions). These include digoxin.
- Anticoagulant medicines (used to prevent blood clots). These include warfarin.
- Medicines for diabetes. These include glimepiride and glipizide.
- Medicines for lowering blood pressure. These include verapamil.

Kalydeco with food and drink

Avoid giving your child food or drink containing grapefruit during treatment as these may increase the side effects of Kalydeco by increasing the amount of ivacaftor in your child's body.

Driving and using machines

Kalydeco can make your child dizzy. If your child feels dizzy, it is advised that your child does not ride his/her bike or do anything else that needs his/her full attention.

Kalvdeco contains lactose and sodium

If you have been told by your child's doctor that your child has an intolerance to some sugars, contact your child's doctor before your child takes this medicine.

Kalydeco contains less than 1 mmol sodium (23 mg) per dose, that is to say essentially 'sodium-free'.

3. How to take Kalydeco

Always give your child this medicine exactly as your child's doctor has told you to. Check with your child's doctor if you are not sure.

Your child's doctor will determine the correct dose for your child. Your child must keep using all other medicines, unless your child's doctor advises to stop using any.

Kalydeco dosing recommendations are provided in Table 1.

Table 1: Dosing recommendations

Age/weight	Morning dose	Evening dose	
Kalydeco as monotherapy			
1 month to less than 2 months, \geq 3 kg	One Kalydeco sachet of	No evening dose	
	13.4 mg granules		
2 months to less than 4 months, \geq 3 kg	One Kalydeco sachet of	One Kalydeco sachet of	
	13.4 mg granules	13.4 mg granules	
4 months to less than 6 months, \geq 5 kg	One Kalydeco sachet of	One Kalydeco sachet of	
	25 mg granules	25 mg granules	
6 months and older, \geq 5 kg to $<$ 7 kg	One Kalydeco sachet of	One Kalydeco sachet of	
	25 mg granules	25 mg granules	

6 months and older, \geq 7 kg to \leq 14 kg	One Kalydeco sachet of	One Kalydeco sachet of		
	50 mg granules	50 mg granules		
6 months and older, \geq 14 kg to \leq 25 kg	One Kalydeco sachet of	One Kalydeco sachet of		
	75 mg granules	75 mg granules		
6 months and older, \geq 25 kg	Please refer to Kalydeco tablets Package Leaflet			
Kalydeco in combination with ivacaft	Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor			
2 years to less than 6 years, 10 kg to	One sachet of ivacaftor	One Kalydeco sachet of		
< 14 kg	60 mg/tezacaftor	59.5 mg granules		
	40 mg/elexacaftor 80 mg			
	granules			
2 years to less than 6 years, ≥ 14 kg	One sachet of ivacaftor	One Kalydeco sachet of		
	75 mg/tezacaftor	75 mg granules		
	50 mg/elexacaftor 100 mg			
	granules			

Give your child the morning and evening granules about 12 hours apart.

If your child has liver problems, your child's doctor may need to reduce the dose of Kalydeco as your child's liver will not clear the medicine as fast as in children who have normal liver function.

- Moderate liver problems in children 6 months of age or older: the dose may be reduced to one half of the indicated dose in the table above, that is one sachet once daily.
- Severe liver problems in children 6 months of age or older: the use is not recommended but your child's doctor will decide if it is appropriate for your child to use this medicine in which case the dose (as indicated in the table above) must be reduced to one sachet every other day.
- Liver problems in children between 1 month and 6 months of age: the use is not recommended.

Kalydeco is for oral use.

Each sachet is for single use only.

Giving Kalydeco to your child:

- Hold sachet of granules with cut line on top.
- Shake sachet gently to settle contents.
- Tear or cut sachet open along cut line.
- Mix the entire content of a sachet with 5 mL of age-appropriate soft food or liquid. Food or liquid should be at room temperature or below. Some examples of age-appropriate soft foods or liquids include puréed fruits or vegetables, yogurt, applesauce, water, milk, breast milk, infant formula, or juice.
- Once mixed, give the product to your child immediately. If this is not possible, give it within the following hour after mixing. Ensure that the mixture is completely and immediately consumed.
- A fat-containing meal or snack should be given to your child just before or just after dosing (some examples are provided below).

Meals or snacks that contain fat include those prepared with butter or oils or those containing eggs. Other fat-containing foods are:

- Cheese, whole milk, whole-milk dairy products, yogurt, breast milk, infant formula, chocolate
- Meats, oily fish
- Avocados, hummus, soy-based products (tofu)
- Nuts, fat-containing nutritional bars or drinks

If your child takes more Kalydeco than he/she should

Your child may experience side effects, including those mentioned in section 4 below. If so, contact your child's doctor or pharmacist to ask for advice. If possible, have your child's medicine and this leaflet with you.

If you forget to give your child Kalydeco

Give the missed dose if less than 6 hours have passed since the time your child missed the dose. Otherwise, wait until your child's next scheduled dose as you normally would. Do not give your child a double dose to make up for a forgotten dose.

If you stop giving your child Kalydeco

Give Kalydeco to your child for as long as your child's doctor recommends. Do not stop unless your child's doctor advises you to. If you have any further questions on the use of this medicine, ask your child's doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Serious side effects

Stomach (abdominal) ache and increased liver enzymes in the blood.

Possible signs of liver problems

Increased liver enzymes in the blood are common in patients with CF and have also been reported in patients taking Kalydeco alone or in combination with ivacaftor/tezacaftor/elexacaftor.

In patients taking Kalydeco in combination with ivacaftor/tezacaftor/elexacaftor, liver damage and worsening of liver function in people with severe liver disease has been reported. The worsening of liver function can be serious and may require transplantation.

These may be signs of liver problems:

- Pain or discomfort in the upper right area of the stomach (abdominal) area
- Yellowing of the skin or white part of the eyes
- Loss of appetite
- Nausea or vomiting
- Dark urine

Depression

Signs of this include sad or altered mood, anxiety, feelings of emotional discomfort.

Tell your child's doctor straight away if he/she gets any of these.

Very common side effects (may affect more than 1 in 10 people)

- Upper respiratory tract infection (the common cold), including sore throat and nasal congestion
- Headache
- Dizziness
- Diarrhoea
- Stomach or abdominal pain
- Changes in the type of bacteria in mucus
- Increased liver enzymes (signs of stress on the liver)
- Rash

Common side effects (may affect up to 1 in 10 people)

- Runny nose
- Ear pain, ear discomfort
- Ringing in the ears
- Redness inside the ear
- Inner ear disorder (feeling dizzy or spinning)
- Sinus problems (sinus congestion)
- Redness in the throat
- Breast mass
- Feeling sick (nausea)
- Flu
- Low blood sugar (hypoglycaemia)
- Abnormal breathing (shortness of breath or difficulty breathing)
- Wind (flatulence)
- Spots (acne)
- Itchy skin
- Increased creatine phosphokinase (sign of muscle breakdown) seen in blood tests

Uncommon side effects (may affect up to 1 in 100 people)

- Ear congestion
- Breast inflammation
- Enlargement of the breast in males
- Nipple changes or pain
- Wheezing
- Increased blood pressure

Not known (frequency cannot be estimated from the available data)

- Damage to the liver (liver injury)
- Raised bilirubin measurement (liver blood test)

Additional side effects in children and adolescents

Side effects seen in children and adolescents are similar to those observed in adults. However, increased liver enzymes in the blood are more frequently seen in young children.

Reporting of side effects

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

5. How to store Kalydeco

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

Do not use this medicine after the expiry date which is stated on the carton, wallet and sachet after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

Once mixed, the mixture has been shown to be stable for one hour.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. Contents of the pack and other information

What Kalydeco contains

The active substance is ivacaftor.

Kalydeco 13.4 mg granules in sachet:

Each sachet contains 13.4 mg of ivacaftor.

Kalydeco 25 mg granules in sachet:

Each sachet contains 25 mg of ivacaftor.

Kalydeco 50 mg granules in sachet:

Each sachet contains 50 mg of ivacaftor.

Kalydeco 59.5 mg granules in sachet:

Each sachet contains 59.5 mg of ivacaftor.

Kalydeco 75 mg granules in sachet:

Each sachet contains 75 mg of ivacaftor.

The other ingredients are: silica, colloidal anhydrous, croscarmellose sodium, hypromellose acetate succinate, lactose monohydrate, magnesium stearate, mannitol, sucralose and sodium laurilsulfate (E487).

See the end of section 2 - Kalydeco contains lactose and sodium.

What Kalydeco looks like and contents of the pack

Kalydeco 13.4 mg granules in sachet are white to off-white granules.

Kalydeco 25 mg granules in sachet are white to off-white granules.

Kalydeco 50 mg granules in sachet are white to off-white granules.

Kalydeco 59.5 mg granules in sachet are white to off-white granules.

Kalydeco 75 mg granules in sachet are white to off-white granules.

The granules are supplied in sachets.

Kalydeco 13.4 mg granules in sachet, Kalydeco 25 mg granules in sachet, Kalydeco 50 mg granules in sachet, and Kalydeco 75 mg granules in sachet:

Pack size of 56 sachets (contains 4 individual wallets with 14 sachets per wallet)

Kalydeco 13.4 mg granules in sachet, Kalydeco 59.5 mg granules in sachet, and Kalydeco 75 mg granules in sachet:

Pack size of 28 sachets (contains 4 individual wallets with 7 sachets per wallet)

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Other sources of information

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