

ANNEX I
SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Imlytic 10^6 plaque forming units (PFU)/mL solution for injection
Imlytic 10^8 plaque forming units (PFU)/mL solution for injection

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

2.1 General description

Talimogene laherparepvec is an attenuated herpes simplex virus type-1 (HSV-1) derived by functional deletion of 2 genes (ICP34.5 and ICP47) and insertion of coding sequence for human granulocyte macrophage colony-stimulating factor (GM-CSF) (see section 5.1).

Talimogene laherparepvec is produced in Vero cells by recombinant DNA technology.

2.2 Qualitative and quantitative composition

Imlytic 10^6 plaque forming units (PFU)/mL solution for injection

Each vial contains 1 mL deliverable volume of Imlytic at a nominal concentration of 1×10^6 (1 million) plaque forming units (PFU)/mL.

Imlytic 10^8 plaque forming units (PFU)/mL solution for injection

Each vial contains 1 mL deliverable volume of Imlytic at a nominal concentration of 1×10^8 (100 million) plaque forming units (PFU)/mL.

Excipient with known effect

Each 1 mL vial contains 7.7 mg sodium and 20 mg sorbitol.

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Solution for injection (injection).

Imlytic 10^6 plaque forming units (PFU)/mL solution for injection

Clear to semi-translucent liquid following thaw from its frozen state.

It may contain white, visible, variously shaped, virus-containing particles.

Imlytic 10^8 plaque forming units (PFU)/mL solution for injection

Semi-translucent to opaque liquid following thaw from its frozen state.

It may contain white, visible, variously shaped, virus-containing particles.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Imlytic is indicated for the treatment of adults with unresectable melanoma that is regionally or distantly metastatic (Stage IIIB, IIIC and IVM1a) with no bone, brain, lung or other visceral disease (see sections 4.4 and 5.1).

4.2 Posology and method of administration

Treatment with talimogene laherparepvec should be initiated and supervised by a qualified physician experienced in the treatment of cancer.

Patients treated with Imlytic must be given the Patient Alert Card and be informed about the risks of the treatment (see also Package leaflet).

Posology

Imlytic is provided in single-use vials of 1 mL each in two different concentrations:

- 10^6 (1 million) PFU/mL - For initial dose only.
- 10^8 (100 million) PFU/mL - For all subsequent doses.

The total injection volume for each treatment visit should be up to a maximum of 4 mL. The initial recommended dose is up to a maximum of 4 mL of Imlytic at a concentration of 10^6 (1 million) PFU/mL. Subsequent doses should be administered up to 4 mL of Imlytic at a concentration of 10^8 (100 million) PFU/mL.

The recommended dosing schedule is shown in table 1.

Table 1. Recommended dosing schedule

Treatment visit	Treatment interval	Maximum total injection volume	Dose concentrations	Prioritisation of lesions to be injected
Initial	-	Up to 4 mL	10^6 (1 million) PFU/mL	<ul style="list-style-type: none">• Inject largest lesion(s) first.• Prioritise injection of remaining lesions based on lesion size until maximum injection volume is reached.
Second	3 weeks after initial treatment	Up to 4 mL	10^8 (100 million) PFU/mL	<ul style="list-style-type: none">• First inject any new lesions (lesions that may have developed since initial treatment).• Prioritise injection of remaining lesions based on lesion size until maximum injection volume is reached.

Treatment visit	Treatment interval	Maximum total injection volume	Dose concentrations	Prioritisation of lesions to be injected
All subsequent treatment visits (including re-initiation)	2 weeks after previous treatment	Up to 4 mL	10^8 (100 million) PFU/mL	<ul style="list-style-type: none"> First inject any new lesions (lesions that may have developed since previous treatment). Prioritise injection of remaining lesions based on lesion size until maximum injection volume is reached.

Determining Imlytic dose volume (per lesion)

The volume to be injected into each lesion is dependent on the size of the lesion and should be determined according to table 2. The total injection volume for each treatment session should be up to a maximum of 4 mL.

Table 2. Selection of Imlytic injection volume based on lesion size

Lesion size (longest dimension)	Imlytic injection volume
> 5 cm	up to 4 mL
> 2.5 cm to 5 cm	up to 2 mL
> 1.5 cm to 2.5 cm	up to 1 mL
> 0.5 cm to 1.5 cm	up to 0.5 mL
≤ 0.5 cm	up to 0.1 mL

Patients may experience increase in size of existing lesion(s) or the appearance of a new lesion prior to achieving a response. As long as there are injectable lesion(s) remaining, Imlytic should be continued for at least 6 months unless the physician considers that the patient is not benefitting from Imlytic treatment or that other treatment is required.

Imlytic treatment may be reinitiated if new lesions appear following a complete response and the physician considers that the patient will benefit from treatment.

Special populations

Elderly population

No adjustment of the dose is required in patients ≥ 65 years old (see section 5.1).

Hepatic and renal impairment

No clinical studies have been conducted to evaluate the effect of hepatic or renal impairment on the pharmacokinetics of talimogene laherparepvec. However, no adjustment in dosage is necessary for patients with hepatic or renal impairment.

Paediatric population

The safety and efficacy of Imlytic in paediatric patients has not been established. Currently available data for paediatric and young adult patients aged 7 to \leq 21 years with advanced non-central nervous system tumours amenable to direct injection are described in section 5.1.

Method of administration

Imlytic is to be administered by intralesional injection into cutaneous, subcutaneous, and/or nodal lesions that are visible, palpable or detectable by ultrasound guidance.

Precautions to be taken before manipulating or administering the medicinal product

This medicinal product contains genetically modified organisms. Personal protective equipment should be worn while preparing or administering talimogene laherparepvec (see section 6.6).

Healthcare professionals who are immunocompromised or pregnant should not administer Imlytic and should not come into direct contact with the injection site(s) or body fluids of treated patients (see sections 4.3 and 4.4).

Follow the instructions below to prepare and administer Imlytic to patients:

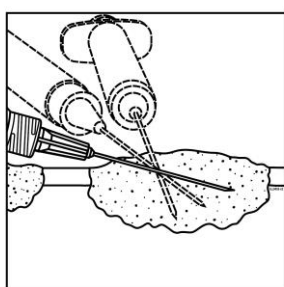
Pre-injection

- Thaw Imlytic vial(s) at room temperature. Thawed vials may be stored prior to administration (see section 6.3). For handling of thawed vials, see section 6.6.
- Draw the desired amount of Imlytic from the vial into a syringe using aseptic technique. A 22- to 26-gauge needle is recommended.
- The injection site may be treated with a topical anaesthetic agent. Injectable anaesthetic may be injected around the periphery of the lesion but should not be injected directly into the lesion.
- Clean the lesion and surrounding areas with an alcohol swab and let dry.

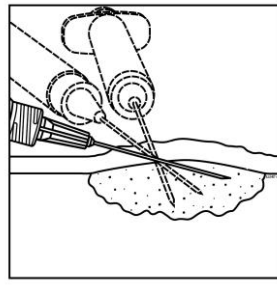
Injection

- Inject Imlytic intralesionally into cutaneous, subcutaneous, and/or nodal lesions that are visible, palpable or detectable by ultrasound guidance.
- Determine injection volume for each lesion using table 2 above.
- Using a single insertion point, inject Imlytic along multiple tracks as far as the radial reach of the needle allows within the lesion to achieve even and complete dispersion. Multiple insertion points may be used if a lesion is larger than the radial reach of the needle.

Cutaneous lesions



Subcutaneous lesions



Nodal lesions

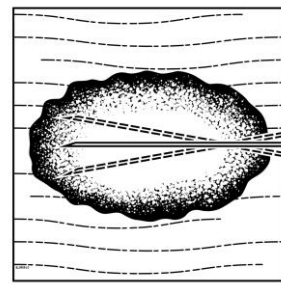


Figure 1.
Injection administration for cutaneous lesions

Figure 2.
Injection administration for subcutaneous lesions

Figure 3.
Injection administration for nodal lesions

- Disperse Imlytic evenly and completely within the lesion by pulling the needle back without exiting the lesion. Redirect the needle as many times as necessary while injecting the remainder of the dose. Continue until the full dose is evenly and completely dispersed.

- When removing the needle, withdraw it from the lesion slowly to avoid leakage or splash back of Imlytic at the insertion point.
- Repeat these steps for other lesions that need to be injected. Use a new needle anytime the needle is completely removed from a lesion and each time a different lesion is injected.

Post-injection

- Apply pressure to the injection site with a sterile gauze for at least 30 seconds.
- Swab the injection site and surrounding area with alcohol, and cover the injected lesion with an absorbent pad and dry occlusive dressing.

4.3 Contraindications

- Patients with a history of hypersensitivity to talimogene laherparepvec or any of its excipients.
- Patients who are severely immunocompromised (e.g. patients with severe congenital or acquired cellular and/or humoral immune deficiency) (see section 4.4).

4.4 Special warnings and precautions for use

Traceability

In order to improve the traceability of biological medicinal products, the name and the batch number of the administered product should be clearly recorded.

Previously treated patients

Efficacy data for Imlytic in the current second or later line treatment settings are limited.

Disseminated herpetic infection

Disseminated herpetic infection, including serious cases of disseminated herpetic infection, have been reported in patients treated with Imlytic (see section 4.8).

Imlytic has not been studied in immunocompromised patients. Based on epidemiological data, immunocompromised patients (such as those with HIV/AIDS, leukaemia, lymphoma, common variable immunodeficiency, or who require chronic high-dose steroids or other immunosuppressive agents) may be at increased risk of disseminated herpetic infection. Consider the risks and benefits of treatment before administering Imlytic to immunocompromised patients.

Based on animal data, patients who are severely immunocompromised may be at an increased risk of disseminated herpetic infection and should not be treated with Imlytic (see sections 4.3 and 5.3).

Accidental exposure to Imlytic

Accidental exposure may lead to transmission of Imlytic and herpetic infection. Healthcare professionals and close contacts (e.g. household members, caregivers, sex partners or persons sharing the same bed) should avoid direct contact with injected lesions or body fluids of treated patients during the entirety of the treatment period and up to 30 days after the last treatment administration (see section 6.6). Accidental needle stick and splash back have been reported in healthcare professionals during preparation and administration.

Close contacts who are pregnant or immunocompromised should not change the patient's dressing or clean their injection site. Pregnant women, neonates, and immunocompromised individuals should not be exposed to potentially contaminated materials.

Healthcare professionals should ensure that patients are able to comply with the requirement to cover injection sites with occlusive dressings (see section 6.6). Patients should also be advised to avoid

touching or scratching injection sites as this could lead to inadvertent transfer of Imlytic to other areas of their body or to their close contacts.

Although it is not known if Imlytic could be transmitted through sexual contact, it is known that wild-type HSV-1 can be transmitted through sexual contact. Patients should be advised to use a latex condom during sexual contact to prevent possible transmission of Imlytic. Women of childbearing potential should be advised to use an effective method of contraception to prevent pregnancy during treatment (see section 4.6).

Caregivers should be advised to wear protective gloves when assisting patients in applying or changing occlusive dressings and to observe safety precautions for disposal of used dressings and cleaning materials (see section 6.6).

In the event of an accidental exposure to Imlytic, follow instructions in section 6.6. If signs or symptoms of herpetic infection develop, exposed individuals should contact their healthcare professional. In case suspected herpetic lesions occur, patients, close contacts or healthcare providers have the option of follow-up testing by the Marketing Authorisation Holder for further characterisation of the infection.

Herpetic infection in Imlytic-treated patients

Herpetic infections (including but not limited to cold sores and herpes keratitis) and serious cases of disseminated herpetic infections have been reported in patients treated with Imlytic (see section 4.8). Symptoms of a local or systemic infection possibly related to Imlytic are anticipated to be similar to symptoms caused by wild-type HSV-1 infections.

Individuals with wild-type HSV-1 infection are known to be at a lifelong risk for symptomatic herpetic infection due to reactivation of latent wild-type HSV-1. Symptomatic herpetic infection due to possible reactivation of Imlytic should be considered.

Patients who develop herpetic infections should be advised to follow standard hygienic practices to prevent viral transmission.

Talimogene laherparepvec is sensitive to acyclovir. The risks and benefits of Imlytic treatment should be considered before administering acyclovir or other anti-viral agents indicated for management of herpetic infection. These agents may interfere with the effectiveness of the treatment if administered systemically or topically directly to the injection site.

Information on herpetic lesions is provided in the Patient Alert Card.

Cellulitis at the injection site

Necrosis or ulceration of tumour tissue may occur following Imlytic treatment. Cellulitis and systemic bacterial infection have been reported. Careful wound care and infection precautions are recommended, particularly if tissue necrosis results in open wounds.

Impaired healing at the injection site

In clinical studies, impaired healing at the injection site has been reported. Imlytic may increase the risk of impaired healing in patients with underlying risk factors (e.g. previous radiation at the injection site, or lesions in poorly vascularised areas).

The risks and benefits of Imlytic should be considered before continuing treatment if persistent infection or delayed healing develops.

Immune-mediated events

In clinical studies, immune-mediated events including as glomerulonephritis, vasculitis, pneumonitis, worsening psoriasis, and vitiligo have been reported in patients treated with Imlytic.

The risks and benefits of Imlytic should be considered before initiating treatment in patients who have underlying autoimmune disease or before continuing treatment in patients who develop immune-mediated events.

Plasmacytoma at injection site

Plasmacytoma has been reported in proximity to the injection site after administration of Imlytic. The risks and benefits of Imlytic should be considered in patients with multiple myeloma or in whom plasmacytoma develops during treatment.

Obstructive airway disorder

Obstructive airway disorder has been reported following Imlytic treatment. Caution should be used when injecting lesions close to major airways.

HSV-1 seronegative patients

Patients who were HSV-1 seronegative at baseline were reported to have a greater incidence of pyrexia, chills, and influenza-like illness compared with those who were HSV-1 seropositive at baseline, especially within the period of the first 6 treatments (see section 4.8).

Hepatic haemorrhage from transcutaneous intrahepatic route of administration

Imlytic is not indicated for transcutaneous intrahepatic route of administration. In clinical studies, cases of hepatic haemorrhage resulting in hospitalisation and death have been reported in patients receiving transcutaneous intrahepatic Imlytic injections.

All patients

This medicinal product contains 20 mg sorbitol per 1 mL vial. The additive effect of concomitantly administered products containing sorbitol (or fructose) and dietary intake of sorbitol (or fructose) should be taken into account.

This medicinal product contains 7.7 mg sodium per 1 mL vial, equivalent to 0.4% of the WHO recommended maximum daily intake of 2 g sodium for an adult.

4.5 Interaction with other medicinal products and other forms of interaction

No interaction studies have been conducted with Imlytic. Acyclovir and other anti-viral agents may interfere with the effectiveness of the treatment if administered systemically or topically directly to the injection site. Consider the risks and benefits of Imlytic treatment before administering acyclovir or other anti-viral agents indicated for management of herpetic infection.

4.6 Fertility, pregnancy and lactation

Women of childbearing potential/contraception

Women of childbearing potential should be advised to use an effective method of contraception to prevent pregnancy during treatment.

All patients should be advised to use a latex condom during sexual contact to prevent possible transmission of Imlytic (see section 4.4).

Pregnancy

Adequate and well controlled studies with talimogene laherparepvec have not been conducted in pregnant women.

If a pregnant woman has an infection with wild-type HSV-1 (primary or reactivation), there is potential for the virus to cross the placental barrier, and also a risk of transmission during birth due to viral shedding. Infections with wild-type HSV-1 have been associated with serious adverse effects, including multi-organ failure and death, if a foetus or neonate contracts the wild-type herpes infection. While there are no clinical data to date on talimogene laherparepvec infections in pregnant women, there could be a risk to the foetus or neonate if talimogene laherparepvec were to act in the same manner. No effects on embryo-foetal development have been observed in animal studies (see section 5.3). As a precautionary measure, it is preferable to avoid the use of talimogene laherparepvec during pregnancy.

Transplacental metastases of malignant melanoma can occur. Because talimogene laherparepvec is designed to enter and replicate in the tumour tissue, there could be a risk of foetal exposure to talimogene laherparepvec from tumour tissue that has crossed the placenta.

If Imlytic is used during pregnancy, or if the patient becomes pregnant while taking the medicinal product, the patient should be apprised of the potential hazards to the foetus and/or neonate.

Breast-feeding

It is unknown whether talimogene laherparepvec is transferred into human milk. A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from Imlytic therapy taking into account the benefit of breast-feeding for the child and the benefit of therapy for the woman.

Fertility

No clinical studies have been performed to evaluate the effects of talimogene laherparepvec on fertility (see section 5.3).

4.7 Effects on ability to drive and use machines

Talimogene laherparepvec may have a minor influence on the ability to drive and use machines. Because of potential adverse reactions such as dizziness and confusional state (see section 4.8), patients should be advised to use caution when driving or operating machinery until they are certain that talimogene laherparepvec does not adversely affect them.

4.8 Undesirable effects

Summary of the safety profile

The safety of Imlytic was evaluated in the pivotal study where 292 patients received at least 1 dose of Imlytic (see section 5.1). The median duration of exposure to Imlytic was 23 weeks (5.3 months). Twenty-six (26) patients were exposed to Imlytic for at least one year.

The most commonly reported adverse reactions ($\geq 25\%$) in Imlytic-treated patients were fatigue (50.3%), chills (48.6%), pyrexia (42.8%), nausea (35.6%), influenza-like illness (30.5%), and injection site pain (27.7%). Overall, 98% of these adverse reactions reported were mild or moderate in severity. The most common grade 3 or higher adverse reaction was cellulitis (2.1%) (see section 4.4).

Tabulated list of adverse reactions

Adverse reactions were determined based on clinical trials in patients with melanoma treated with Imlytic compared to GM-CSF and post-marketing experience. Incidence of adverse reactions are presented by system organ class and by frequency. Frequencies are defined as: very common ($\geq 1/10$), common ($\geq 1/100$ to $< 1/10$) and uncommon ($\geq 1/1\,000$ to $< 1/100$). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness.

Table 3. Adverse reactions from clinical trials in patients with melanoma and post-marketing experience

Infections and infestations	
Common	Cellulitis*, Herpes infections**
Uncommon	Incision site infection
Neoplasms benign, malignant and unspecified (including cysts and polyps)	
Common	Tumour pain, Infected neoplasm
Uncommon	Plasmacytoma at injection site*
Blood and lymphatic system disorders	
Very common	Oedema peripheral
Common	Anaemia
Immune system disorders	
Common	Immune-mediated events ^{†*}
Uncommon	Hypersensitivity
Metabolism and nutrition disorders	
Common	Dehydration
Nervous system disorders	
Very common	Headache
Common	Confusional state, Anxiety, Depression, Dizziness, Insomnia
Eye disorders	
Uncommon	Keratitis herpetic
Ear and labyrinth disorders	
Common	Ear pain
Cardiac disorders	
Common	Tachycardia
Vascular disorders	
Common	Deep vein thrombosis, Hypertension, Flushing
Respiratory, thoracic and mediastinal disorders	
Very common	Cough
Common	Dyspnoea, Oropharyngeal pain, Upper respiratory tract infection
Uncommon	Obstructive airways disorder
Gastrointestinal disorders	
Very common	Vomiting, Diarrhoea, Constipation, Nausea
Common	Abdominal pain, Abdominal discomfort
Skin and subcutaneous tissue disorders	
Common	Vitiligo, Rash, Dermatitis
Uncommon	Granulomatous dermatitis
Musculoskeletal and connective tissue disorders	
Very common	Myalgia, Arthralgia, Pain in extremity
Common	Back pain, Groin pain
General disorders and administration site conditions	
Very common	Influenza-like illness*, Pyrexia, Chills, Fatigue, Pain, Injection site reactions [§]
Common	Malaise, Axillary pain
Investigations	
Common	Weight decreased

Injury, poisoning and procedural complications	
Common	Wound complication, Wound secretion, Contusion, Procedural pain

[§] Injection site reactions include: very common term of injection site pain, common terms of injection site erythema, injection site haemorrhage, injection site swelling, injection site reaction, injection site inflammation, secretion discharge, injection site discharge, uncommon term of injection site warmth.

[†] Immune-mediated events include: uncommon terms of vasculitis, pneumonitis, worsening psoriasis and glomerulonephritis.

^{*} See Description of selected adverse reactions.

^{**} Herpetic infections (including, but not limited to Oral herpes).

Description of selected adverse reactions

Immune-mediated events

Immune-mediated events reported in the pivotal clinical study included a case of worsening psoriasis in a patient with a prior history of psoriasis, one case of pneumonitis in a patient with a prior history of autoimmune disease, one case of vasculitis, and two cases of glomerulonephritis of which one presented with acute renal failure.

Plasmacytoma

In clinical trials, one case of plasmacytoma at injection site was observed in a patient who was found to have multiple myeloma.

Cellulitis

In the pivotal clinical trial (study 005/05), events of cellulitis were recorded, some of them being considered as serious adverse events. However, none lead to permanent discontinuation of Imlytic treatment. Careful wound care and infection precautions are recommended, particularly if tissue necrosis results in open wounds.

Influenza-like symptoms

Ninety percent (90%) of patients treated with Imlytic experienced influenza-like symptoms. Pyrexia, chills, and influenza-like illness, which can occur any time during treatment, generally resolved within 72 hours. These events were reported more frequently within the period of the first 6 treatments, particularly in patients who were HSV-1 negative at baseline.

Paediatric population

A paediatric phase 1 clinical trial (study 20110261) was conducted in 15 paediatric and young adult patients aged 7 to \leq 21 years with advanced non-central nervous system tumours amenable to direct injection (see section 5.1). The safety data was consistent with the patients' underlying disease and the known safety profile of talimogene laherparepvec in adults.

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

There is no clinical experience with overdose with Imlytic. Doses up to 4 mL at a concentration of 10^8 PFU/mL every 2 weeks have been administered in clinical trials with no evidence of dose limiting toxicity. The maximum dose that can be safely administered has not been determined. In the event of a suspected overdose or inadvertent intravenous administration, the patient should be treated

symptomatically, e.g. with acyclovir or other anti-viral agents (see section 4.4) and supportive measures instituted as required.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Antineoplastic cell and gene therapy, ATC code: L01XL02.

Mechanism of action

Talimogene laherparepvec is an oncolytic immunotherapy that is derived from HSV-1. Talimogene laherparepvec has been modified to replicate within tumours and to produce the immune stimulatory protein human GM-CSF. Talimogene laherparepvec causes the death of tumour cells and the release of tumour-derived antigens. It is thought that together with GM-CSF, it will promote a systemic anti-tumour immune response and an effector T-cell response. Mice that had complete regression of their primary tumours following treatment were resistant to subsequent tumour re-challenge.

The modifications to talimogene laherparepvec from HSV-1 include deletion of ICP34.5 and ICP47. Whereas anti-viral immune responses defend normal cells following infection by talimogene laherparepvec, tumours have been shown to be susceptible to injury and cell death from ICP34.5-deficient HSV-1 viruses, including talimogene laherparepvec. Deletion of ICP47 prevents down-regulation of antigen presentation molecules and increases the expression of HSV US11 gene, thereby enhancing viral replication in tumour cells.

Clinical efficacy and safety

Study 005/05

The safety and efficacy of Imlytic monotherapy compared with subcutaneously administered GM-CSF were evaluated in a phase 3, multinational, open-label, and randomised clinical study of patients with stage IIIB, IIIC, and IV melanoma that was not considered to be surgically resectable. Previous systemic treatment for melanoma was allowed but not required. Patients with active brain metastases, bone metastases, extensive visceral disease, primary ocular or mucosal melanoma, evidence of immunosuppression, or receiving treatment with a systemic anti-herpetic agent were excluded from the study.

Patients were randomised in a 2:1 ratio to receive either Imlytic or GM-CSF (N = 436; 295 Imlytic, 141 GM-CSF). Imlytic was administered by intralesional injection at an initial concentration of 10^6 (1 million) PFU/mL on day 1, followed by a concentration of 10^8 (100 million) PFU/mL on day 21 and every 2 weeks thereafter at a dose of up to 4 mL. GM-CSF was administered subcutaneously at 125 mcg/m² delivered daily for 14 days followed by a 14-day rest period in repeating intervals.

To allow for delayed immune-mediated anti-tumour effects to occur, patients were treated for a minimum of 6 months or until there were no longer any injectable lesions. During this period, treatment was to continue despite an increase in size in existing lesion(s) and/or development of new lesion(s) unless the patient developed intolerable toxicity or the investigator believed that it was in the best interest of the patient to stop treatment or to be given other therapy for melanoma. After 6 months of treatment, patients were to continue treatment until clinically relevant disease progression (i.e. disease progression associated with a decline in performance status and/or alternative therapy was required in the opinion of the investigator). Patients experiencing a response at 12 months of treatment could continue treatment for up to an additional 6 months. The mean (SD) treatment duration for the intent-to-treat (ITT) population was 15.76 weeks (15.79) in the GM-CSF arm and 26.83 weeks (18.39) in the Imlytic arm. The primary endpoint was durable response rate (DRR) [defined as the percent of patients with complete response (CR) or partial response (PR) maintained continuously for a minimum of 6 months] per blinded central review. The secondary endpoints included overall survival (OS),

overall response rate (ORR) [PR+CR], time to response, duration of response, and time to treatment failure (time from randomisation until the first episode of clinically relevant disease progression where there is no response achieved after the progression event, or until death).

The mean age was 63 (range: 22 to 94) years; 26.5% were over 65 years old and 23.3% were over 74 years old. The majority of patients, 98%, were caucasian. Male patients comprised 57% of study population and 70% of patients were baseline ECOG 0 performance status. Of the enrolled patients, 22% had stage IV M1c disease and 53% of patients had received prior therapy for melanoma such as chemotherapy and cytokine-based immunotherapy in addition to surgery, adjuvant therapy or radiation. Overall, 58% of all patients enrolled into the study were seropositive for wild-type HSV-1 at baseline and 32.6% were seronegative; the HSV-1 serostatus of the remaining 9.4% was unknown.

The difference in DRR between Imlytic and GM-CSF in the ITT population was statistically significant (see table 4) in favour of Imlytic.

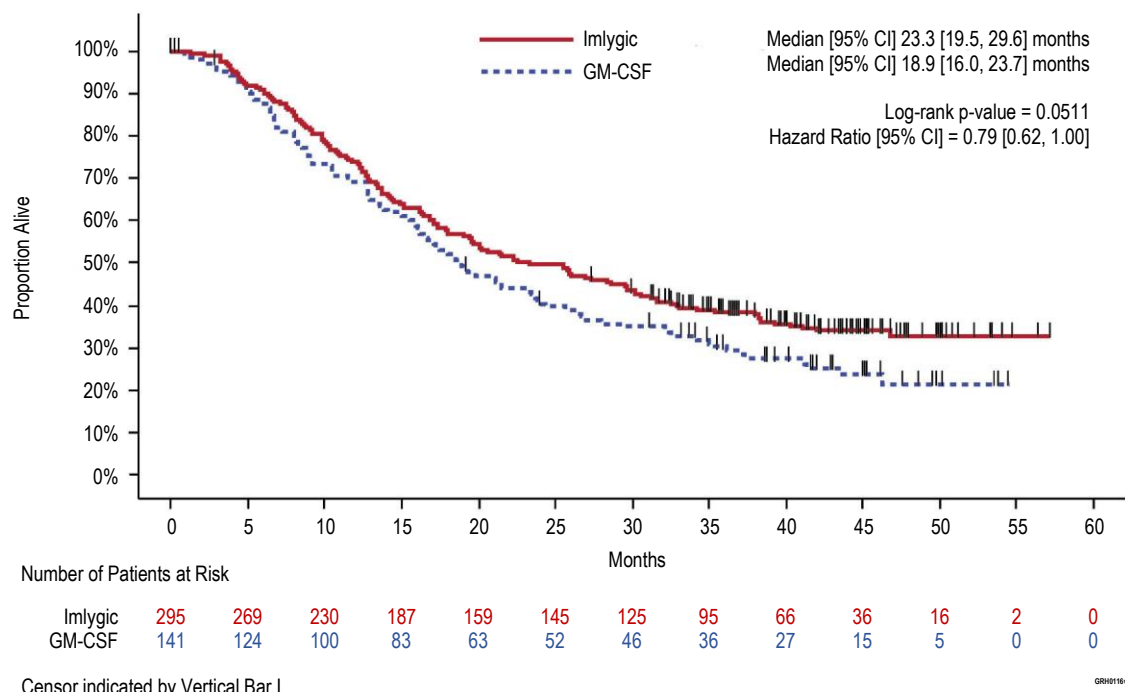
Table 4. Summary of results for the ITT population from Imlytic study 005/05

	Study endpoint	Imlytic N = 295	GM-CSF N = 141
Durable response rate	Primary	16.3% (n = 48) (95% CI: 12.1, 20.5)	2.1% (n = 3) (95% CI: 0.0, 4.5)
		Odds ratio 8.9; (95% CI: 2.7, 29.2) p < 0.0001	
Overall response rate (% CR, % PR)	Secondary	26.4% (n = 78) (95% CI: 21.4%, 31.5%) (10.8% CR, 15.6% PR)	5.7% (n = 8) (95% CI: 1.9%, 9.5%) (0.7% CR, 5% PR)
Overall survival	Secondary	Median 23.3 (95% CI: 19.5, 29.6) months	Median 18.9 (95% CI: 16.0, 23.7) months
		HR: 0.79; (95% CI: 0.62, 1.00) p = 0.051	
Duration of response (ongoing response at last tumour evaluation)	Secondary	Not reached (Range: > 0.0 to > 16.8 months)	Median 2.8 months (Range: 1.2 to > 14.9 months)
		HR: 0.46; (95% CI: 0.35, 0.60)	
Time to response (median)	Secondary	4.1 months	3.7 months
Time to treatment failure (median)	Secondary	8.2 months (95% CI: 6.5, 9.9)	2.9 months (95% CI: 2.8, 4.0)
		HR: 0.42; (95% CI: 0.32, 0.54)	

Among the Imlytic-treated responders, 56 (72%) responses were still ongoing at the time of primary analysis. Of the responders, 42 (54%) experienced a $\geq 25\%$ increase in overall size of existing lesion(s) and/or developed a new lesion(s) prior to ultimately achieving a response.

In an analysis to evaluate systemic activity of Imlytic, 27 of 79 patients (34.2%) had a $\geq 50\%$ overall decrease in non-visceral lesions that were not injected with Imlytic, and 8 of 71 patients (11.3%) had a $\geq 50\%$ overall decrease in visceral lesions that were not injected with Imlytic.

Figure 4. Kaplan-Meier plot –overall survival (ITT population)



No overall differences in safety or efficacy were observed between elderly (≥ 65 years old) and younger adult patients.

Exploratory subgroups

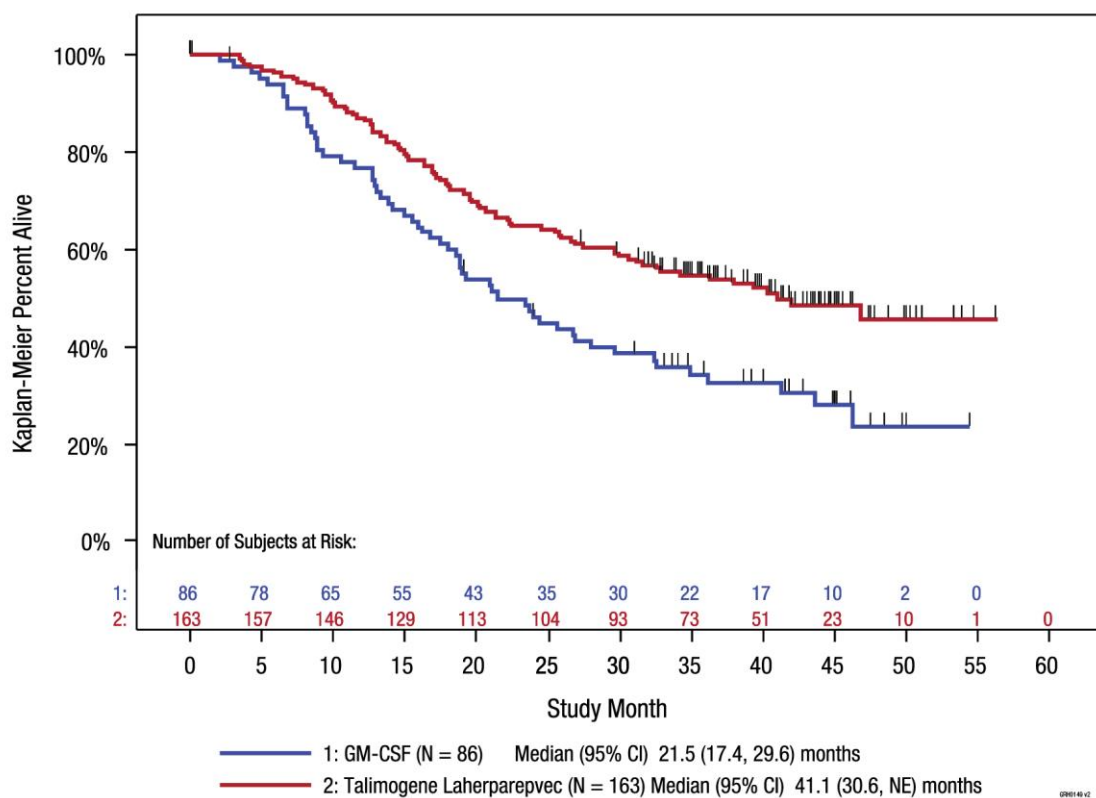
Exploratory subgroup analyses for DRR and overall survival by stage of disease were also carried out (see figure 5 and table 5). While the pivotal study was not powered to evaluate efficacy in these individual subgroups, patients with no visceral disease derived greater benefit from Imlytic treatment than those with more advanced disease.

Table 5. Summary of results from exploratory subgroup analysis from Imlytic study 005/05

	DRR, (%)		ORR, (%)		OS (hazard ratio)
	Imlytic	GM-CSF	Imlytic	GM-CSF	Imlytic vs GM-CSF
Stage [§] IIIB/IIIC/ stage IVM1a (Imlytic, n = 163; GM-CSF, n = 86)	25.2	1.2	40.5	2.3	0.57, (95% CI: 0.40, 0.80);
Stage [§] IVM1B/ IVM1C (Imlytic, n = 131; GM-CSF, n = 55)	5.3	3.6	9.2	10.9	1.07, (95% CI: 0.75, 1.52);

[§] American Joint Committee on Cancer (AJCC) staging 6th edition.

Figure 5. Kaplan-Meier estimate of overall survival by randomised treatment arm for disease stage IIIB/IIIC/ stage IVM1a (exploratory subgroup analysis)



Censor indicated by vertical bar |

NE = not estimable

Due to the exploratory nature of the analysis and based on the current evidence, it has not been established that Imlytic is associated with an effect on overall survival.

Paediatric population

One phase 1, multi-centre, open-label, dose de-escalation clinical trial (study 20110261) evaluated the safety and efficacy of talimogene laherparepvec in paediatric patients with advanced non-central nervous system tumours that were amenable to direct injection. A total of 15 paediatric and young adult patients aged 7 to \leq 21 years, divided in two cohorts; cohort A1 (13 patients aged 12 to \leq 21 years) and cohort B1 (2 patients aged 7 to $<$ 12 years) received talimogene laherparepvec during the study. The dosing schedule was consistent with the recommended adult talimogene laherparepvec dose.

Thirteen patients were included in the dose limiting toxicity (DLT) analysis set. No patient had a DLT during the DLT evaluation period. Overall, all patients (15 patients, 100.0%) had at least 1 treatment-emergent adverse event, and 8 patients (53.3 %) had grade \geq 3 adverse events.

No responses were observed, ORR per modified irRC-RECIST was 0% (95% CI: 0.0, 21.8).

5.2 Pharmacokinetic properties

Talimogene laherparepvec is a genetically modified and replication-competent HSV-1 virus. Therefore, its pharmacokinetics and biodistribution are driven by the site of intralesional injection, tumour-selective replication, and release from tumour tissue.

Absorption

Cellular uptake of talimogene laherparepvec occurs through HSV-1 receptors on tumours and non-tumour cells following local injection into tumours. As talimogene laherparepvec is injected and replicates intratumourally, bioavailability and systemic concentration of talimogene laherparepvec are not predictive of drug substance activity and therefore have not been evaluated.

Metabolism/elimination

Talimogene laherparepvec is cleared through general host-defence mechanisms (e.g. autophagy, adaptive immune responses). Talimogene laherparepvec is degraded by typical endogenous protein and DNA catabolic pathways. As with other wild-type HSV-1 infections, a latent pool of talimogene laherparepvec DNA may persist in neuronal cell bodies innervating the injection sites; therefore, the occurrence of latent infection with talimogene laherparepvec cannot be excluded.

Biodistribution (within the body) and viral shedding (excretion/secretion)

Talimogene laherparepvec DNA was quantified with a highly sensitive and specific quantitative polymerase chain reaction (qPCR) assay which may not correlate with viral infectivity risk.

Talimogene laherparepvec was also quantified in selected patient samples in clinical studies using viral infectivity assays at the injection sites and in some cases of potential herpetic lesions.

Clinical biodistribution, elimination, and shedding

The biodistribution and shedding of intralesionally administered talimogene laherparepvec were investigated in a clinical study that measured talimogene laherparepvec DNA in blood, urine, injection site, exterior of the occlusive dressings, oral mucosa, anogenital area, and suspected herpetic lesions. Sixty patients with melanoma received Imlytic intralesional injection at a dose and schedule same as clinical study 005/05 (see section 5.1). Occlusive dressings samples were collected during treatment. Blood and urine samples were collected during treatment and for up to 30 days after the end of treatment. Injection site, oral mucosa, and anogenital area samples were collected during treatment and for up to 60 days after the end of treatment. Suspected herpetic lesion samples were collected any time a patient experienced lesions of suspected herpetic origin. If the qPCR testing for talimogene laherparepvec DNA was positive, then a TCID₅₀ assay was performed to measure viral infectivity. In the 60 patients treated, data indicate that talimogene laherparepvec DNA was present in all sites during the study (see table 6).

Table 6. Patients with detectable DNA during treatment

Body fluid/site	Patients with detectable DNA during treatment (n = 60)
Blood	59 (98%)
Urine	19 (32%)
Injection site	60 (100%)
Exterior of the occlusive dressing	48 (80%)
Oral mucosa	8 (13%)
Anogenital area	5 (19%) ^a

^a For the anogenital area, 26 patients were tested for Imlytic DNA.

The proportion of samples and subjects with talimogene laherparepvec DNA was highest during cycle 2 of treatment for the blood, urine, injection site, and occlusive dressings; highest in cycle 1 of treatment for the oral mucosa; and highest in cycles 1 and 2 for the anogenital area. Among patients with detectable talimogene laherparepvec DNA in the blood, urine, oral mucosa, and anogenital area, no samples had detectable talimogene laherparepvec DNA 30 days after the end of treatment. For patients with detectable DNA in injected lesions, no samples had detectable talimogene laherparepvec DNA 60 days after the end of treatment.

Overall 3 of 19 patients with lesions of suspected herpetic origin had talimogene laherparepvec DNA present at any time during the study. Viral activity was measured in samples that were positive for talimogene laherparepvec DNA from the injection site, occlusive dressings, oral mucosa, anogenital area, and suspected herpetic lesions. No viral activity was detected in samples of the occlusive dressings, oral mucosa, anogenital area, and suspected herpetic lesions. Infectious talimogene laherparepvec virus was detected at the site of injection in 7 (11%) patients at multiple time points in the study; no samples were positive for viral infectivity after cycle 2 or after the end of treatment.

Pharmacokinetics in special populations

No pharmacokinetic studies using talimogene laherparepvec have been conducted in special populations.

5.3 Preclinical safety data

At doses up to 4×10^8 PFU/kg or 10^7 PFU/dose (60-fold over the highest proposed clinical dose), single or repeated doses of talimogene laherparepvec administered by SC, IV, or intratumoural injection were well tolerated in immunocompetent mice, rats, and dogs. No neuropathology or adverse neurological effects were observed. In an *in vivo* study of intracerebral injection, talimogene laherparepvec was 10 000-fold less neurovirulent as compared to the wild-type HSV-1 dose that results in death 50% of the time in mice.

Talimogene laherparepvec was injected into various xenograft tumours at doses up to 2×10^8 PFU/kg (30-fold over the highest proposed clinical dose) in immunodeficient mice (nude and SCID). Lethal systemic viral infection was observed in up to 20% of nude mice (primarily deficient in T lymphocyte function) and 100% of SCID mice (devoid of both T and B lymphocytes).

Across studies, fatal disseminated viral infection was observed in 14% of nude mice following treatment with talimogene laherparepvec at doses that are 10- to 100-fold higher than those that result in 100% lethality with wild-type HSV-1.

Mutagenicity

The genotoxic potential of talimogene laherparepvec has not been evaluated in long-term animal or human studies. Because wild-type HSV-1 does not integrate into the host genome, the risk of insertional mutagenesis with talimogene laherparepvec is negligible.

Carcinogenicity

The carcinogenic potential of talimogene laherparepvec has not been evaluated in long-term animal or human studies. However, available data for talimogene laherparepvec and wild-type HSV-1 do not indicate a carcinogenic risk in humans.

Reproductive and development toxicity

There were no impacts to male or female reproductive tissues following treatment of adult mice at doses up to 4×10^8 PFU/kg (60-fold higher, on a PFU/kg basis, compared to the maximum clinical dose). No effects on embryo-foetal development were observed when talimogene laherparepvec was administered during organogenesis to pregnant mice at doses up to 4×10^8 (400 million) PFU/kg (60-fold higher, on a PFU/kg basis, compared to the maximum clinical dose). Negligible amounts (< 0.001% of maternal blood levels) of talimogene laherparepvec DNA were found in foetal blood.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Di-sodium phosphate dihydrate
Sodium dihydrogen phosphate dihydrate
Sodium chloride
Myo-inositol
Sorbitol (E420)
Water for injections

6.2 Incompatibilities

In the absence of compatibility studies, this medicinal product must not be mixed with other medicinal products.

6.3 Shelf life

Unopened vial

5 years.

Preparation and storage prior to administration

After thawing, administer Imlytic as soon as practically feasible.

Thawed Imlytic is stable when stored at temperatures of 2°C up to 25°C protected from light in its original vial, in a syringe, or in the original vial followed by a syringe. Do not exceed the storage times specified in table 7 and table 8.

If storing thawed Imlytic in the original vial followed by a syringe:

- The same temperature range should be maintained throughout the duration of storage until administration.
- The storage time in the syringe at ambient temperature up to 25°C cannot exceed 2 hours for 10^6 (1 million) PFU/mL and 4 hours for 10^8 (100 million) PFU/mL (see table 7).
- The maximum cumulative storage time (storage time in vial plus storage time in syringe) cannot exceed the durations in table 8.

Imlytic must not be refrozen once it has thawed. Discard any thawed Imlytic in the vial or syringe stored longer than the specified times below.

Table 7. Maximum storage time for thawed Imlytic in syringe

	10^6 (1 million) PFU/mL	10^8 (100 million) PFU/mL
2°C to 8°C	8 hours	8 hours
up to 25°C	2 hours	4 hours

Table 8. Maximum cumulative storage time (storage time in vial plus storage time in syringe) for thawed Imlytic

	10^6 (1 million) PFU/mL	10^8 (100 million) PFU/mL
2°C to 8°C	24 hours	1 week (7 days)
up to 25°C	12 hours	24 hours

6.4 Special precautions for storage

Store and transport frozen (-90°C to -70°C).

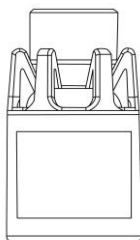
Store in the original carton in order to protect from light.

For storage conditions after thawing of the medicinal product, see section 6.3.

6.5 Nature and contents of container

Imlytic is provided as a one mL preservative-free solution in a single-use vial (cyclic olefin polymer plastic resin) with stopper (chlorobutyl elastomer) and seal (aluminium) with flip-off cap (polypropylene) in two different presentations:

Figure 6. Single-use vial permanently inserted into a clear copolyester plastic sleeve



OR

Figure 7. Single-use vial without a clear plastic sleeve



The vial cap is colour coded: 10^6 (1 million) PFU/mL is light green and 10^8 (100 million) PFU/mL is royal blue.

6.6 Special precautions for disposal and other handling

Thawing Imlytic vials

- Before use, thaw frozen Imlytic vials at room temperature (20°C to 25°C) until Imlytic is liquid. The time to achieve complete vial thaw is expected to be 30 to 70 minutes, depending on the ambient temperature. Gently swirl. Do NOT shake.
- Vials should be thawed and stored in the original carton until administration in order to protect from light.

Handling and Administration

Follow local guidelines for handling and administration, personal protective equipment, accidental spills, and waste disposal.

- Wear protective gown or laboratory coat, safety glasses, or face shield and gloves while preparing or administering Imlytic. Cover any exposed wounds before administering. Avoid contact with skin, eyes or mucous membranes.

- After administration, change gloves prior to applying occlusive dressings to injected lesions. Wipe the exterior of occlusive dressing with an alcohol wipe. It is recommended to keep injection sites covered with airtight and watertight dressings at all times, if possible. To minimise the risk of viral transmission, patients should keep their injection site covered for at least 8 days from the last treatment or longer if the injection site is weeping or oozing. Advise patients to apply dressing as instructed by the healthcare professional and to replace the dressing if it falls off.
- Dispose of all materials that have come in contact with Imlytic (e.g. vial, syringe, needle, any cotton or gauze) in accordance with local procedures.

Accidental exposure

- In the event of an accidental occupational exposure to Imlytic (e.g. through a splash to the eyes or mucous membranes) during preparation or administration, flush with clean water for at least 15 minutes. In the event of exposure to broken skin or needle stick, clean the affected area thoroughly with soap and water and/or disinfectant.
- Treat all Imlytic spills with a virucidal agent and absorbent materials.
- Advise patients to place used dressings and cleaning materials in a sealed plastic bag as they may be potentially contaminated, and to dispose of the bag in household waste.

This medicine contains genetically modified organisms.

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

7. MARKETING AUTHORISATION HOLDER

Amgen Europe B.V.
Minervum 7061
4817 ZK Breda
The Netherlands

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/15/1064/001
EU/1/15/1064/002

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 16 December 2015
Date of latest renewal: 18 November 2020

10. DATE OF REVISION OF THE TEXT

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

ANNEX II

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

A. MANUFACTURERS OF THE BIOLOGICAL ACTIVE SUBSTANCE AND MANUFACTURERS RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturers of the biological active substance

BioVex LLC - Subsidiary of Amgen, Inc.
34 Commerce Way
Woburn
Massachusetts
01801
United States

IDT Biologika GmbH
Am Pharmapark
Dessau-Roßlau
Sachsen-Anhalt 06861
Germany

Name and address of the manufacturers responsible for batch release

Amgen Europe B.V.
Minervum 7061
4817 ZK Breda
Netherlands

Amgen NV
Telecomlaan 5-7
1831 Diegem
Belgium

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

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

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

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

• Periodic safety update reports (PSURs)

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

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

- Risk management plan (RMP)**

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

An updated RMP should be submitted:

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

- Additional risk minimisation measures**

Prior to launch of Imlytic in each Member State the Marketing Authorisation Holder (MAH) must agree about the content and format of the educational and controlled distribution programme, including communication media, distribution modalities, and any other aspects of the programme, with the National Competent Authority.

The educational programme is aimed to inform about important risks associated with Imlytic:

- Herpetic infection occurring throughout the entire body (disseminated herpetic infection) in immunocompromised individuals (those with any congenital or acquired cellular and/or humoral immune deficiency, i.e. HIV/AIDS, leukaemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)
- Accidental exposure of Healthcare Professionals (HCPs) to Imlytic
- Spread of Imlytic to close contacts or healthcare providers after direct contact with injected lesions or body fluids
- Symptomatic herpetic infection due to latency and reactivation of Imlytic or herpes (wild-type HSV-1) in patients
- Patients with a weakened immune system (immunocompromised patients) treated with Imlytic and suffering from concomitant infection
- Combination with other therapies like chemotherapy or immunosuppressive agents
- Pregnant and lactating women

The MAH shall ensure that in each Member State where Imlytic is marketed, all healthcare professionals and patients/carers who are expected to prescribe, dispense and use Imlytic have access to/are provided with the following educational package:

- Physician educational material
- Patient information pack

The physician educational material should contain:

- The Summary of Product Characteristics
- Guide for healthcare professionals
- Patient Alert Card
- **Guide for healthcare professionals** shall contain the following key elements:
 - Information on the risk of herpetic infection in patients treated by Imlytic
 - Information on the risk of disseminated herpetic infection in immunocompromised individuals treated by Imlytic
 - Recommendation regarding accidental exposure of Imlytic to HCPs
 - To always wear protective gown/laboratory coat, safety glasses and gloves while preparing or administrating Imlytic

- To avoid contact with skin, eyes, mucous membranes and ungloved direct contact with injected lesions or body fluids of treated patients
- Instruction on first aid after accidental exposure
- Immunocompromised and pregnant healthcare professionals should not prepare and administer Imlytic
- Recommendation regarding the accidental transmission of Imlytic from patient to close contacts or HCPs
- Instruction on how to behave after administration/accidental transmission and how and how often the dressing has to be changed and who should not change the dressing
- Instructions to minimise the risk of exposure of blood and body fluids to close contacts for the duration of Imlytic treatment through 30 days after the last administration of Imlytic. The following activities should be avoided:
 - Sexual intercourse without a latex condom
 - Kissing if either party has an open mouth sore
 - Common usage of cutlery, crockery, and drinking vessels
 - Common usage of injection needles, razorblades and toothbrushes
- Adequate waste disposal and decontamination, following the recommendations for disposal of biohazardous waste
- Information on Imlytic use in pregnancy
- Instructions how to handle possible adverse events including providing of batch number when reporting adverse drug reactions

- **The Patient Alert Card** shall contain the following key messages:
 - A warning message for HCPs treating the patient at any time, including in conditions of emergency, that the patient is using Imlytic
 - Contact details of the Imlytic prescriber
 - Details about Imlytic treatment start date, batch number, date administered, product manufacturer and license holder
 - Information of herpetic lesions
- The patient information pack should contain:
 - Patient information leaflet
 - A patient/carer and close contacts guide
- **The Patient/carer and close contacts guide** shall contain the following key messages:
 - A description of the important risks associated with the use of Imlytic
 - Instruction on how to behave after administration and how and how often the dressing has to be changed and who should not change the dressing
 - Information of the sign and symptoms of the risk of herpetic infection
 - Information on Imlytic use in pregnancy
 - Recommendation regarding the accidental transmission of Imlytic from patient to close contacts or HCPs
 - Instructions to minimise the risk of exposure of blood and body fluids to close contacts for the duration of Imlytic treatment through 30 days after the last administration of Imlytic. The following activities should be avoided:
 - Sexual intercourse without a latex condom
 - Kissing if either party has an open mouth sore
 - Common usage of cutlery, crockery, and drinking vessels
 - Common usage of injection needles, razorblades and toothbrushes
 - Adequate waste disposal and decontamination, following the recommendations for disposal of biohazardous waste
 - Instruction on how to behave after accidental transmission

The controlled distribution programme is aimed to manage the product supply chain to ensure that cold storage requirements are observed and to control the distribution of Imlytic to qualified centres and up to the patients.

The MAH shall ensure that in each Member State where Imlytic is marketed, a system aimed to control distribution to Imlytic beyond the level of control ensured by routine risk minimisation measures. The following requirements need to be fulfilled before the product is dispensed:

- Adequately trained and experienced HCPs in order to minimise the risk of occurrence of specified adverse drug reactions in patients, HCPs, and close contacts of the patients
- Trained HCPs and support personnel regarding safe and appropriate storage, handling, and administration of Imlytic, and clinical follow-up for patients treated with Imlytic
- Provide specified safety information to patients and communicate to patients the importance for sharing this information with family and caregivers
- Trained HCPs to record batch number information in patient's charts and on the patient's alert card for all injections and to provide the batch number when reporting adverse drug reactions.

ANNEX III
LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGING**CARTON****1. NAME OF THE MEDICINAL PRODUCT**

Imlytic 10^6 plaque forming units (PFU)/mL solution for injection
talimogene laherparepvec

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each vial contains 1 mL of 1×10^6 (1 million) plaque forming units (PFU) of talimogene laherparepvec.

3. LIST OF EXCIPIENTS

Di-sodium phosphate dihydrate, sodium dihydrogen phosphate dihydrate, sodium chloride, myo-inositol, sorbitol (E420), water for injections.

4. PHARMACEUTICAL FORM AND CONTENTS

Solution for injection

1 vial.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Intralesional use.

Read the package leaflet before 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

Store and transport frozen at -90°C to -70°C .

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

This medicine contains genetically modified organisms.
Unused medicine must be disposed of in compliance with the local guidelines.

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Amgen Europe B.V.
Minervum 7061
4817 ZK Breda
The Netherlands

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/15/1064/001

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Justification for not including Braille accepted.

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC
SN
NN

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS**VIAL LABEL****1. NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION**

Imlytic 10⁶ PFU/mL injection
talimogene laherparepvec
Intraleisional use

2. METHOD OF ADMINISTRATION**3. EXPIRY DATE**

EXP

4. BATCH NUMBER

Lot

5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT

1 mL

6. OTHER

PARTICULARS TO APPEAR ON THE OUTER PACKAGING**CARTON****1. NAME OF THE MEDICINAL PRODUCT**

Imlytic 10⁸ plaque forming units (PFU)/mL solution for injection
talimogene laherparepvec

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each vial contains 1 mL of 1 × 10⁸ (100 million) plaque forming units (PFU) of talimogene laherparepvec.

3. LIST OF EXCIPIENTS

Di-sodium phosphate dihydrate, sodium dihydrogen phosphate dihydrate, sodium chloride, myo-inositol, sorbitol (E420), water for injections.

4. PHARMACEUTICAL FORM AND CONTENTS

Solution for injection

1 vial.

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Intralesional use.

Read the package leaflet before 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

Store and transport frozen at -90°C to -70°C.

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

This medicine contains genetically modified organisms.
Unused medicine must be disposed of in compliance with the local guidelines.

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Amgen Europe B.V.
Minervum 7061
4817 ZK Breda
The Netherlands

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/15/1064/002

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Justification for not including Braille accepted.

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC
SN
NN

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS**VIAL LABEL****1. NAME OF THE MEDICINAL PRODUCT AND ROUTE(S) OF ADMINISTRATION**

Imlytic 10⁸ PFU/mL injection

talimogene laherparepvec

Intralesional use

2. METHOD OF ADMINISTRATION**3. EXPIRY DATE**

EXP

4. BATCH NUMBER

Lot

5. CONTENTS BY WEIGHT, BY VOLUME OR BY UNIT

1 mL

6. OTHER

B. PACKAGE LEAFLET

Package leaflet: Information for the patient

Imlytic 10^6 plaque forming units (PFU)/mL solution for injection Imlytic 10^8 plaque forming units (PFU)/mL solution for injection talimogene laherparepvec

Read all of this leaflet carefully before you start using 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 healthcare professional (doctor or nurse).
- If you get any side effects, talk to your healthcare professional. This includes any possible side effects not listed in this leaflet. See section 4.
- Your doctor will give you a Patient Alert Card. Read it carefully and follow the instructions on it.
- Always show the Patient Alert Card to your doctor or nurse when you see them or if you go to hospital.

What is in this leaflet

1. What Imlytic is and what it is used for
2. What you need to know before and during Imlytic treatment
3. How Imlytic is given
4. Possible side effects
5. How Imlytic is stored
6. Contents of the pack and other information

1. What Imlytic is and what it is used for

Imlytic is used to treat adult patients with a type of skin cancer called melanoma that has spread in the skin or to the lymph nodes, when surgery is not an option.

The active ingredient of Imlytic is talimogene laherparepvec. This is a weakened form of herpes simplex virus type-1 (HSV-1), which is commonly called the cold sore virus. To get Imlytic from HSV-1, the virus has been changed so that it multiplies more effectively in tumours than in normal cells. This leads to destruction of infected tumour cells. This medicine also works by helping your immune system to recognise and destroy tumours throughout your body.

2. What you need to know before and during Imlytic treatment

You will not be given Imlytic:

- if you are allergic to talimogene laherparepvec or any of the other ingredients of this medicine (listed in section 6).
- if your healthcare professional has told you that you have a severely weakened immune system.

Warnings and precautions

Talk to your healthcare professional before being given this medicine.

Life-threatening herpes infection

Life-threatening herpes infection including spreading to any part of the body far from the injection site (disseminated herpetic infection) may occur. If you have any new or worsening symptoms, tell your healthcare professional immediately. Tell your healthcare professional if you have or have ever had a

weakened immune system, if you have HIV/AIDS, blood or bone marrow cancer, or if you are taking steroids or other medicines that suppress your immune system because you may be at increased risk of life-threatening herpes infection.

Accidental spread of Imlytic to yourself and others

Imlytic can be spread to other parts of your body or to other people through direct contact with your body fluids or injection sites.

You should do the following to avoid spreading Imlytic to other areas of your body or to your close contacts (close contacts include household members, caregivers, sex partners, or someone you share a bed with):

- Avoid direct contact between your injection sites or body fluids (e.g. blood and urine) and close contacts (e.g. use latex condoms when engaging in sexual activity, avoid kissing close contacts if either of you has an open mouth sore) while you are being treated with this medicine and up to 30 days after your last dose.
- Avoid touching or scratching the injection sites.
- Keep injection sites covered with airtight and watertight dressings at all times. Apply the dressing as instructed by your healthcare professional. If the dressing comes loose or falls off, replace it immediately with a clean dressing.
- Place all used dressings and cleaning materials in a sealed plastic bag and throw them away in your household waste.

You should tell your close contacts to:

- Avoid direct contact with your body fluids or injection sites.
- Wear gloves while changing your dressing.

If your close contacts are accidentally exposed to Imlytic, they should clean the affected area on their body with soap and water and/or a disinfectant. If they develop signs or symptoms of herpes infection, you should ask them to contact their healthcare professional. If herpetic lesions (blisters or sores) are suspected, patients or close contacts have the option of follow-up testing by the Marketing Authorisation Holder for further characterisation of the infection. Please discuss with your healthcare professional.

Close contacts who are pregnant or who have a weakened immune system, and newborns

Ensure that your close contacts who are pregnant or who have a weakened immune system do not touch injection sites, used dressings and cleaning materials. Keep used dressings and cleaning materials away from newborns.

Herpes infection

Cold sores or a more serious herpes infection may occur during or after treatment with Imlytic. Signs and symptoms related to treatment with Imlytic may be the same as for herpes infections, and include but are not limited to pain, burning or tingling in a blister around the mouth, genitals, on the fingers or ears, eye pain, light sensitivity, discharge from the eyes, or blurry vision, weakness in arms or legs, extreme drowsiness (feeling sleepy), and mental confusion. If you have these signs or any new symptoms, you should follow standard hygiene practices to prevent viral transmission to others. If herpetic lesions (blisters or sores) are suspected, patients or close contacts have the option of follow-up testing by the Marketing Authorisation Holder for further characterisation of the infection. Please discuss with your healthcare professional.

Infection and delayed healing at injection site

Imlytic may cause infection at the injection site. Signs and symptoms of infection include pain, redness, warmth, swelling, discharge or a sore (ulcer), fever, and chills. The injection site may take

longer to heal than normal. You should tell your healthcare professional if you notice any of these symptoms.

Autoimmune reactions

Imlytic may cause autoimmune reactions (an over-reaction of the body's immune system). Some people taking this medicine have developed inflammation in the kidneys (glomerulonephritis), narrowing or blockage of blood vessels (vasculitis), swelling of the lungs (pneumonitis), worsening skin scaling (psoriasis), and areas of skin without any colour (vitiligo). Inform your healthcare professional if you have a history of autoimmune disease.

Plasmacytoma

Imlytic may cause cancerous white blood cells to gather at or near the injection site (plasmacytoma). Inform your healthcare professional if you have a history of blood cancer including multiple myeloma.

Difficulty breathing

If you have a tumour in your neck, your healthcare professional may warn you that you might experience compression of your airways during treatment.

Patients with no prior herpes infection

If you have never had herpes infection in the past, you may be more likely to get fever, chills, and flu-like illness within the period of the first 6 treatments.

Children and adolescents

The use of Imlytic has been studied in children and young adults aged 7 to \leq 21 years with advanced non-central nervous system tumours amenable to direct injection. The use of Imlytic has not been studied in children under 7 years of age.

Other medicines and Imlytic

Tell your healthcare professional if you are taking, have recently taken or might take any other medicines, including medicines, such as acyclovir, to treat or prevent herpes infections. Acyclovir and other anti-viral treatments may decrease the effects of Imlytic.

Pregnancy and breast-feeding

Ask your healthcare professional for advice if you:

- think you may be pregnant; or
- are planning to have a baby.

Your healthcare professional will determine if Imlytic is right for you.

If you are pregnant or breast-feeding, ask your healthcare professional for advice before being given this medicine. Imlytic may harm your unborn baby.

Women who are able to become pregnant should use effective contraception to avoid pregnancy during treatment with Imlytic. Talk to your healthcare professional about suitable methods of contraception.

It is not known whether Imlytic passes into breast milk. It is important to tell your healthcare professional if you are breast-feeding or plan to do so. They will then help you decide whether to stop breast-feeding, or whether to stop taking Imlytic, taking into account the benefit of breast-feeding to the baby and the benefit of Imlytic to you.

Driving and using machines

When you are being treated with Imlytic you may experience symptoms such as dizziness or confusion. This may impair your ability to drive or operate machinery. Use caution when driving or operating machinery until you are certain that this medicine does not adversely affect you.

Imlytic contains sodium and sorbitol

This medicine contains 7.7 mg sodium (main component of cooking/table salt) in each 1 mL vial. This is equivalent to 0.4% of the recommended maximum daily dietary intake of sodium for an adult.

This medicine contains 20 mg sorbitol in each 1 mL vial.

3. How Imlytic is given

This medicine is given in a healthcare facility under the supervision of a healthcare professional. The initial recommended dose is up to 4 mL of Imlytic at a concentration of 10^6 (1 million) PFU/mL. Subsequent doses will be up to 4 mL of Imlytic at a concentration of 10^8 (100 million) PFU/mL.

Your healthcare professional will inject this medicine directly into your tumour(s) with a needle and a syringe. Your second injection will be given 3 weeks after the first injection. After that, you will receive injections every 2 weeks for as long as you have the tumour(s).

Your healthcare professional will decide which tumour(s) to inject and may not inject every tumour. Your existing tumour(s) may increase in size and new tumour(s) could appear while you are being treated with Imlytic.

You can expect to be treated with Imlytic for at least 6 months or longer.

If you miss a dose of Imlytic

It is important for you to keep all your appointments to receive this medicine. If you miss an appointment, ask your healthcare professional when to schedule your next dose.

4. Possible side effects

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

Keeping wounds clean and dressed can help prevent infections caused by bacteria (cellulitis) at the injection site.

Flu-like illness, fevers and chills have been seen in patients treated with Imlytic. These symptoms generally resolve within the first 72 hours after treatment.

The following side effects have been reported in patients receiving Imlytic:

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

- Tissue swelling (peripheral oedema)
- Headache
- Cough
- Vomiting, diarrhoea, constipation, nausea
- Muscle pain (myalgia), painful/swollen joints (arthralgia), limb pain
- Flu-like illness, fever (pyrexia), chills, fatigue, pain
- Pain, redness, bleeding, swelling, inflammation, secretion, discharge, and warmth at the injection site

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

- Infection caused by bacteria (cellulitis), cold sores (oral herpes)
- Tumour pain, infected tumour
- Tiredness, headaches, dizziness and looking pale (low red blood cell numbers - anaemia)
- Side effects related to the immune system:
 - fever, fatigue, weight loss, muscle and joint pain (narrowing or blockage of blood vessels - vasculitis)
 - shortness of breath, cough, fatigue, loss of appetite, unintentional weight loss (inflammation of the lungs - pneumonitis)
 - increase in patches of skin which are dry, red and covered in silvery scales (worsening scaling of the skin - worsening psoriasis)
 - pink or cola-coloured urine, frothy urine, high blood pressure, fluid retention (inflammation of kidneys - glomerulonephritis)
- Dehydration
- Confusion, anxiety, depression, dizziness, difficulty sleeping (insomnia)
- Pain in ear, throat, abdomen, groin, back and underarm
- Faster heart rate at rest (tachycardia)
- Pain, swelling, heat, and tenderness in a leg or arm due to a blood clot within a vein (deep vein thrombosis), high blood pressure (hypertension), redness in the face (flushing)
- Shortness of breath (dyspnoea), upper respiratory infection
- Abdominal discomfort
- Areas of skin without any colour (vitiligo), rash, inflamed skin (dermatitis)
- Generally feeling unwell
- Weight loss
- Wound complication, secretion, bruising (contusion), pain after procedure

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

- Incision site infection
- A tumour of cancerous white blood cells that grows at or near the injection site (plasmacytoma)
- Eye infection caused by herpes (keratitis herpetic)
- Compressed airways (obstructive airways disorder)
- Allergic reaction (hypersensitivity)

Reporting of side effects

If you get any side effects, talk to your healthcare professional. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in [Appendix V](#). By reporting side effects you can help provide more information on the safety of this medicine.

5. How Imlytic is stored

Imlytic will be stored by the healthcare professionals at your healthcare facility.

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

Store and transport frozen at -90°C to -70°C.

Store in the original carton in order to protect from light.

This medicinal product contains genetically modified cells. Local guidelines should be followed.

6. Contents of the pack and other information

What Imlytic contains

- The active substance is talimogene laherparepvec.
Each vial contains 1 extractable mL of solution at a nominal concentration of 1×10^6 (1 million) plaque forming units (PFU)/mL or 1×10^8 (100 million) PFU/mL.
- The other ingredients are di-sodium phosphate dihydrate, sodium dihydrogen phosphate dihydrate, sodium chloride, myo-inositol, sorbitol (E420), water for injections (see section 2).

What Imlytic looks like and contents of the pack

Imlytic is a clear to semi-translucent (10^6 PFU/mL) or semi-translucent to opaque (10^8 PFU/mL) liquid. It is supplied as a 1 mL preservative-free solution in a single-use vial (cyclic olefin polymer plastic resin) with stopper (chlorobutyl elastomer) and seal (aluminium) with flip-off cap (polypropylene).

The vial cap is colour coded: 10^6 PFU/mL is light green and 10^8 PFU/mL is royal blue.

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The Netherlands

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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 web site:
<https://www.ema.europa.eu>

The following information is intended for healthcare professionals only:

This medicinal product contains genetically modified organisms. Personal protective equipment (e.g. protective gown or laboratory coat, safety glasses, or face shield and gloves) should be worn while preparing or administering talimogene laherparepvec.

After administration, change gloves prior to applying occlusive dressings to injected lesions. Wipe the exterior of occlusive dressing with an alcohol wipe. It is recommended to keep injection sites covered with airtight and watertight dressings at all times, if possible.

Thawing Imlytic vials

- Before use, thaw frozen Imlytic vials at room temperature (20°C to 25°C) until Imlytic is liquid. The time to achieve complete vial thaw is expected to be 30 to 70 minutes, depending on the ambient temperature. Gently swirl. Do NOT shake.
- Vials should be thawed and stored in the original carton until administration in order to protect from light.

After thawing

- After thawing, administer Imlytic as soon as practically feasible.
- Thawed Imlytic is stable when stored at temperatures of 2°C up to 25°C protected from light in its original vial, in a syringe, or in the original vial followed by a syringe. Do not exceed the storage times specified in table 1 and table 2.
- If storing thawed Imlytic in the original vial followed by a syringe:
 - the same temperature range should be maintained throughout the duration of storage until administration.
 - the storage time in the syringe at ambient temperature up to 25°C cannot exceed 2 hours for 10⁶ (1 million) PFU/mL and 4 hours for 10⁸ (100 million) PFU/mL (see table 1).
 - the maximum cumulative storage time (storage time in vial plus storage time in syringe) cannot exceed the durations in table 2.
- Imlytic must not be refrozen once it has thawed. Discard any thawed Imlytic in the vial or syringe stored longer than the specified times below.

Table 1. Maximum storage time for thawed Imlytic in syringe

	10⁶ (1 million) PFU/mL	10⁸ (100 million) PFU/mL
2°C to 8°C	8 hours	8 hours
up to 25°C	2 hours	4 hours

Table 2. Maximum cumulative storage time (storage time in vial plus storage time in syringe) for thawed Imlytic

	10⁶ (1 million) PFU/mL	10⁸ (100 million) PFU/mL
2°C to 8°C	24 hours	1 week (7 days)
up to 25°C	12 hours	24 hours

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