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**Swiss Summary of the Risk Management Plan (RMP) for  
IMLYGIC® (Talimogene Laherparepvec)**

RMP Summary: Version 2, November 2021  
EU RMP: Version 9.3, 25 January 2021

The Risk Management Plan (RMP) is a comprehensive document submitted as part of the application dossier for market approval of a medicine. The RMP summary contains information on the medicine's safety profile and explains the measures that are taken in order to further investigate and follow the risks as well as to prevent or minimize them.

The RMP summary of IMLYGIC® is a concise document and does not claim to be exhaustive.

As the RMP is an international document, the summary might differ from the “Arzneimittelinformation / Information sur le médicament” approved and published in Switzerland, e.g. by mentioning risks occurring in populations or indications not included in the Swiss authorization.

Please note that the reference document which is valid and relevant for the effective and safe use of IMLYGIC® in Switzerland is the “Arzneimittelinformation/ Information sur le médicament” (see [www.swissmedic.ch](http://www.swissmedic.ch)) approved and authorized by Swissmedic.

AMGEN Switzerland AG is fully responsible for the accuracy and correctness of the content of the published summary RMP of IMLYGIC®.

## The medicine and what it is used for

Imlygic is authorized for treatment of adults with unresectable (cannot be removed by surgery) melanoma (a kind of skin cancer) that is regionally (in the skin or lymph nodes near the original skin tumor) or distantly metastatic (spread to distant areas of skin or lymph nodes) (Stage IIIB, IIIC and IVM1a) with no bone, brain, lung or other visceral (internal organ) disease. It contains talimogene laherparepvec as the active substance and it is given by intralesional injection (injection into the tumor).

Further information about the evaluation of Imlygic's benefits can be found in Imlygic's EPAR, including in its plain-language summary, available on the European Medicines Agency (EMA) website, under the medicine's webpage:

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

## Risks associated with the medicine and activities to minimize or further characterize the risks

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

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

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

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

In the case of Imlygic, these measures are supplemented with *additional risk minimization measures* mentioned under relevant risks, below.

In addition to these measures, information about adverse events is collected continuously and regularly analyzed including periodic safety update report (PSUR) assessment so that immediate action can be taken as necessary. These measures constitute *routine pharmacovigilance activities*.

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

## List of Important Risks and Missing Information

Important risks of Imlygic are risks that need special risk management activities to further investigate or minimize the risk, so that the medicinal product can be safely administered. Important risks can be regarded as identified or potential. Identified risks are concerns for which there is sufficient proof of a link with the use of Imlygic. Potential risks are concerns for which an association with the use of this medicine is possible based on available data, but this association has not been established yet and needs further evaluation. Missing information refers to information on the safety of the medicinal product that is currently missing and needs to be collected (eg, on the long-term use of the medicine).

<b>List of important risks and missing information</b>	
Important Identified Risk	<ul style="list-style-type: none"> <li>Disseminated herpetic infection in severely immunocompromised individuals (those with any severe congenital or acquired cellular and/or humoral immune deficiency)</li> <li>Accidental exposure of healthcare provider to talimogene laherparepvec</li> <li>Immune-mediated adverse reactions</li> </ul>
Important Potential Risk	<ul style="list-style-type: none"> <li>Disseminated herpetic infection in immunocompromised patients (such as those with human immunodeficiency virus/acquired immunodeficiency syndrome (HIV/AIDS), leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)</li> <li>Transmission of talimogene laherparepvec from patient to close contacts or healthcare providers via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</li> <li>Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients</li> <li>Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type herpes simplex virus type 1 in patients</li> <li>Immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection</li> <li>Combination with other therapies like chemotherapy or immunosuppressive agents</li> <li>Talimogene laherparepvec-mediated anti-granulocyte macrophage colony stimulating factor antibody response</li> </ul>
Missing Information	<ul style="list-style-type: none"> <li>Pregnant and lactating women</li> <li>Pediatric patients</li> <li>Long-term safety data</li> <li>Long-term efficacy data</li> <li>Treatment of patients with metastatic lesions greater than 3 cm</li> </ul>

**Summary of Important Risks**

Important Identified Risk: Disseminated herpetic infection (herpes infection occurring throughout the body) in severely immunocompromised individuals (those with any severe congenital [present at birth] or acquired cellular and/or humoral immune deficiency [weakness of the immune system])	
Evidence for linking the risk to the medicine	This important identified risk was identified based on nonclinical data.
Risk factors and risk groups	Individuals with any severe congenital or acquired cellular and/or humoral immune deficiency.
Risk minimization measures	<p>Routine risk measures:</p> <ul style="list-style-type: none"> <li>• SmPC Sections 4.3, 4.4, and 5.3</li> <li>• PL Section 2</li> </ul> <p>Additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance activities	<p>Additional pharmacovigilance activities:</p> <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180062</li> <li>• Study 20180099</li> <li>• Quantitative polymerase chain reaction (qPCR) testing for talimogene laherparepvec DNA (a laboratory test to detect the presence of talimogene laherparepvec DNA)</li> </ul> <p>See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan</p>

Important Identified Risk: Accidental exposure of healthcare provider to talimogene laherparepvec	
Evidence for linking the risk to the medicine	This risk was identified based on reports in the clinical study setting.
Risk factors and risk groups	<p>Numerous factors, some modifiable and some not, place healthcare providers at an increased risk for accidental exposure such as sustaining a needle stick injury. These factors include occupation, training, proper disposal of sharps, and medical activity being performed (<a href="#">National Institute for Occupational Safety and Health, DHHS (NIOSH), 1999; Publication No. 2000-2108</a>).</p>

Risk minimization measures	<p>Routine risk communication:</p> <ul style="list-style-type: none"> <li>• SmPC Sections 4.2, 4.4, and 6.6</li> <li>• PL Section 2</li> </ul> <p>Additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> </ul>
Additional pharmacovigilance activities	<p>Additional pharmacovigilance activities:</p> <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> <p>See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan</p>

Important Identified Risk: Immune-mediated adverse reactions	
Evidence for linking the risk to the medicine	This is considered an important identified risk based on reports in the clinical study setting.
Risk factors and risk groups	Risk factors for an immune-mediated adverse reaction include host factors (eg, demographics, other comorbidities), host genotypes ( <a href="#">Thong and Tan, Br J Clin Pharmacol, 2011; 71:684-700</a> ), and pre-existing autoimmune disease.
Risk minimization measures	<p>Routine risk communication:</p> <ul style="list-style-type: none"> <li>• SmPC Sections 4.4 and 4.8</li> <li>• PL Sections 2 and 4</li> </ul> <p>Additional risk minimization measures: None</p>

Important Potential Risk: Disseminated herpetic infection (herpes infection occurring throughout the body) in immunocompromised patients (such as those with HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)	
Evidence for linking the risk to the medicine	This risk is considered an important potential risk based on nonclinical data.
Risk factors and risk groups	<p>Immunosuppression can be due to congenital immunodeficiency, acquired disease (HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, generalized malignancy), pharmacotherapy (immunosuppressive agents, radiation, or large amounts of corticosteroids), or extremes of age (neonates and elderly) (<a href="#">Chinen and Shearer, J Allergy Clin Immunol, 2010; 125(suppl 2):195-203</a>; <a href="#">Notarangelo, J Allergy Clin Immunol, 2010; 125(suppl 2):182-194</a>).</p> <p>The precise risk factors applicable to this risk with talimogene laherparepvec are unknown.</p>

Risk minimization measures	<p>Routine risk communication:</p> <ul style="list-style-type: none"> <li>• SmPC Sections 4.4 and 5.3</li> <li>• PL Section 2</li> </ul> <p>Additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance activities	<p>Additional pharmacovigilance activities:</p> <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180062</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> <p>See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan</p>

<p>Important Potential Risk: Transmission of talimogene laherparepvec from patient to close contacts or healthcare providers via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</p>	
Evidence for linking the risk to the medicine	<p>This risk is considered an important potential risk based on clinical and nonclinical data.</p>
Risk factors and risk groups	<p>Direct contact with injected lesions, protective dressings, or body fluids of treated patients. The likelihood of transfer of talimogene laherparepvec to a close contact or healthcare provider increases if the contact has a break in the skin or mucous membranes.</p>
Risk minimization measures	<p>Routine risk communication:</p> <ul style="list-style-type: none"> <li>• SmPC Sections 4.4 and 6.6</li> <li>• PL Section 2</li> </ul> <p>Additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance activities	<p>Additional pharmacovigilance activities:</p> <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180062</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> <p>See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan</p>

Important Potential Risk: Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients	
Evidence for linking the risk to the medicine	This risk is considered an important potential risk based on clinical data.
Risk factors and risk groups	No risk factors have been identified.
Risk minimization measures	Routine risk communication: <ul style="list-style-type: none"> <li>• SmPC Section 4.4</li> <li>• PL Section 2</li> </ul> Additional risk minimization measures: <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance activities	Additional pharmacovigilance activities: <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180062</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

Important Potential Risk: Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type herpes simplex virus type 1 in patients	
Evidence for linking the risk to the medicine	This risk is considered an important potential risk based on nonclinical data.
Risk factors and risk groups	Previous infection with wild-type herpes simplex virus type 1. Fever, stress, and other factors are common triggers of recurrence.
Risk minimization measures	Routine risk communication: <ul style="list-style-type: none"> <li>• SmPC Section 4.4</li> <li>• PL Section 2</li> </ul> Additional risk minimization measures: <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance activities	Additional pharmacovigilance activities: <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180062</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan



Important Potential Risk: Immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection	
Evidence for linking the risk to the medicine	This important potential risk was identified based on theoretical concern and limited data with immunocompromised patients treated with talimogene laherparepvec.
Risk factors and risk groups	Immunosuppression can be due to congenital immunodeficiency, acquired disease (HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, generalized malignancy), pharmacotherapy (immunosuppressive agents, radiation or large amounts of corticosteroids), or extremes of age (neonates and elderly) ( <a href="#">Chinen and Shearer, J Allergy Clin Immunol, 2010; 125(suppl 2):195-203</a> ; <a href="#">Notarangelo, J Allergy Clin Immunol, 2010; 125(suppl 2):182-194</a> ). The precise risk factors applicable to this risk with talimogene laherparepvec are unknown.
Risk minimization measures	Routine risk communication: <ul style="list-style-type: none"> <li>• SmPC Sections 4.3, 4.4, and 5.3</li> <li>• PL Section 2</li> </ul> Additional risk minimization measures: <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> </ul>
Additional pharmacovigilance activities	Additional pharmacovigilance activities: <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20180099</li> <li>• qPCR testing for talimogene laherparepvec DNA</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

Important Potential Risk: Combination with other therapies like chemotherapy or immunosuppressive agents	
Evidence for linking the risk to the medicine	This is considered an important potential risk based on nonclinical data from immunocompromised mice.
Risk factors and risk groups	Patients receiving concomitant chemotherapeutic or immunosuppressive therapies.
Risk minimization measures	Routine risk communication: <ul style="list-style-type: none"> <li>• SmPC Section 4.4</li> <li>• PL Section 2</li> </ul>

Important Potential Risk: Talimogene laherparepvec-mediated anti-granulocyte macrophage colony stimulating factor antibody response (development of antibodies to granulocyte macrophage colony stimulating factor, which is a chemical in the body that increases the production of white blood cells)	
Evidence for linking the risk to the medicine	This risk is considered an important potential risk based on theoretical concerns.
Risk factors and risk groups	Risk factors are unknown for the development of antibodies against granulocyte macrophage colony stimulating factor ( <a href="#">Meager et al, Immunology, 1999; 97:526-532</a> ).
Risk minimization measures	No risk minimization measures
Additional pharmacovigilance activities	Additional pharmacovigilance activities: <ul style="list-style-type: none"> <li>• Testing of anti-granulocyte macrophage colony stimulating factor antibodies</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

### Missing information

Missing Information: Pregnant and lactating women	
Risk minimization measures	Routine risk communication <ul style="list-style-type: none"> <li>• SmPC Sections 4.4, 4.6, and 5.3</li> <li>• PL Section 2</li> </ul> Additional risk minimization measures: <ul style="list-style-type: none"> <li>• Managed Distribution Program</li> <li>• Physician Education Booklet</li> <li>• Patient Safety Brochure</li> <li>• Patient Alert Card</li> </ul>
Additional pharmacovigilance Activities	Additional pharmacovigilance activities <ul style="list-style-type: none"> <li>• Study 20180062</li> <li>• Study 20180099</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

Missing Information: Pediatric patients	
Risk minimization measures	Routine risk communication <ul style="list-style-type: none"> <li>• SmPC Sections 4.2</li> <li>• PL Section none</li> </ul> Additional risk minimization measures: None

Additional pharmacovigilance Activities	Additional pharmacovigilance activities <ul style="list-style-type: none"> <li>• Study 20110261</li> <li>• Study to be determined</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan
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<b>Missing Information:</b> Long-term safety data	
Risk minimization measures	No risk minimization measures
Additional pharmacovigilance Activities	Additional pharmacovigilance activities <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20120139</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

<b>Missing Information:</b> Long-term efficacy data	
Risk minimization measures	No risk minimization measures
Additional pharmacovigilance Activities	Additional pharmacovigilance activities <ul style="list-style-type: none"> <li>• Study 20130193</li> <li>• Study 20120139</li> </ul> See <a href="#">Section II.C</a> of this summary for an overview of the postauthorization development plan

<b>Missing Information:</b> Treatment of patients with metastatic lesions greater than 3 cm	
Risk minimization measures	No risk minimization measures

## Postauthorization Development Plan

### Studies Which Are Conditions of the Marketing Authorization

The following studies are conditions of the marketing authorization:

Study Short Name	Purpose of the Study
<p>Study 20110265</p> <p>A phase 1b/3, multicenter trial of talimogene laherparepvec in combination with pembrolizumab (MK-3475) for treatment of unresectable, stage IIIB to IVM1c melanoma</p>	<p><u>Primary Objectives</u></p> <ul style="list-style-type: none"> <li>Phase 1b: To evaluate the safety, as assessed by incidence of dose-limiting toxicity, of talimogene laherparepvec in combination with pembrolizumab in subjects with previously untreated, unresectable, stage IIIB to IVM1c melanoma.</li> <li>Phase 3: To evaluate the efficacy of talimogene laherparepvec with pembrolizumab versus placebo with pembrolizumab, as assessed by progression-free survival (response evaluation by blinded independent central review using modified Response Evaluation Criteria in Solid Tumors 1.1 [RECIST]) and overall survival.</li> </ul> <p><u>Efficacy uncertainties addressed:</u> Preliminary efficacy</p>
<p>Study 20110266</p> <p>A phase 2, multicenter, randomized, open-label trial assessing the efficacy and safety of talimogene laherparepvec neoadjuvant treatment plus surgery versus surgery alone for resectable, stage IIIB to IVM1a melanoma</p>	<p><u>Primary Objective</u></p> <p>To estimate the treatment effect of neoadjuvant talimogene laherparepvec plus surgery compared to surgery alone on recurrence-free survival.</p> <p><u>Efficacy uncertainties addressed:</u> Preliminary efficacy and safety</p>

### Other Studies in Postauthorization Development Plan

Study Short Name	Purpose of the Study
<p>Study 20130193</p> <p>A postmarketing prospective cohort study of melanoma patients treated with IMLYGIC® (talimogene laherparepvec) in clinical practice to characterize the risk of herpetic infection among patients, close contacts, and health care providers; and long-term safety in treated patients.</p>	<p>Estimate the incidence rate of herpetic infection detection of talimogene laherparepvec DNA among patients for up to 5 years after the first IMLYGIC dose.</p> <p><u>Safety concerns addressed:</u></p> <ul style="list-style-type: none"> <li>Disseminated herpetic infection in severely immunocompromised individuals (those with any severe congenital or acquired cellular and/or humoral immune deficiency)</li> <li>Accidental exposure of healthcare provider to talimogene laherparepvec</li> <li>Disseminated herpetic infection in immunocompromised patients (such as those with HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)</li> <li>Transmission of talimogene laherparepvec from patient to close contacts or healthcare providers via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</li> </ul>

	<ul style="list-style-type: none"> <li>• Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients</li> <li>• Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type herpes simplex virus type 1 in patients</li> <li>• Immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection</li> <li>• Long-term safety data</li> <li>• Long-term efficacy data</li> </ul>
<p>Study 20110261 A phase 1 multi center, open label, dose de-escalation study to evaluate the safety and efficacy of talimogene laherparepvec in pediatric subjects with advanced non-central nervous system (outside brain and spinal cord) tumors that are amenable to direct injection.</p>	<p>To evaluate the safety and tolerability of talimogene laherparepvec as assessed by incidence of dose-limiting toxicities, in pediatric subjects with advanced non-central nervous system tumors that are amenable to direct injection. <u>Safety concerns addressed:</u> Pediatric patients</p>
<p>Study 20120139 A registry study to evaluate the survival and long-term safety and subject who previously received talimogene laherparepvec in Amgen or BioVEX-sponsored clinical trials</p>	<ul style="list-style-type: none"> <li>• To evaluate the long-term safety of talimogene laherparepvec</li> <li>• To monitor the subject overall survival</li> </ul> <p><u>Safety concerns addressed:</u></p> <ul style="list-style-type: none"> <li>• Long-term safety data</li> <li>• Long-term efficacy data</li> </ul>
<p>Study to be determined A randomized, controlled study to evaluate the safety and efficacy of talimogene laherparevec in children from birth to &lt; 18 years of age with a pediatric solid malignant tumor as part of a multi-modal treatment approach</p>	<p>To be determined <u>Safety concerns addressed:</u> Pediatric patients</p>
<p>Study 20180062 A cross-sectional survey to evaluate patient knowledge of safety messages included in the Patient Safety Brochure and Patient Alert Card for IMLYGIC</p>	<p><u>Primary Objective</u> To evaluate patients' knowledge levels of the key messages evaluate included in the IMLYGIC Patient Safety Brochure among patients who receive IMLYGIC.</p> <p><u>Secondary Objective</u> To evaluate patients' levels of receipt and reading of the IMLYGIC Patient Safety Brochure and receipt, reading, and use (ie, carrying) of the Patient Alert Card among patients who receive IMLYGIC. Patients' understanding of the purpose of the Patient Alert Card will also be assessed. <u>Safety concerns addressed</u></p> <ul style="list-style-type: none"> <li>• Disseminated herpetic infection in severely immunocompromised individuals (those with any severe congenital or acquired cellular and/or humoral immune deficiency)</li> <li>• Disseminated herpetic infection in immunocompromised patients (such as those with HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other</li> </ul>

	<p>immunosuppressive agents)</p> <ul style="list-style-type: none"> <li>• Transmission of talimogene laherparepvec from patient to close contacts or healthcare providers via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</li> <li>• Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients</li> <li>• Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type herpes simplex virus type 1 in patients</li> <li>• Pregnant and lactating women</li> </ul>
<p>Study 20180099 A cross-sectional survey to evaluate physician knowledge of safety messages included in the Physician Education Booklet (PEB) for IMLYGIC®</p>	<p>Primary Objective To evaluate physicians' knowledge levels of the key messages included in the IMLYGIC Physician Education Booklet among physicians who completed the required IMLYGIC training.</p> <p><u>Secondary Objectives</u></p> <ul style="list-style-type: none"> <li>• To evaluate physicians' levels of receipt and reading of the IMLYGIC Physician Education Booklet among physicians who completed the required IMLYGIC training.</li> <li>• To evaluate physicians' understanding of the requirements to distribute the Patient Safety Brochure and Patient Alert Card.</li> </ul> <p><u>Safety concerns addressed</u></p> <ul style="list-style-type: none"> <li>• Disseminated herpetic infection in severely immunocompromised individuals (those with any severe congenital or acquired cellular and/or humoral immune deficiency)</li> <li>• Accidental exposure of healthcare provider to talimogene laherparepvec</li> <li>• Disseminated herpetic infection in immunocompromised patients (such as those with HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)</li> <li>• Transmission of talimogene laherparepvec from patient to close contacts or healthcare providers via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</li> <li>• Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients</li> <li>• Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type herpes simplex virus type 1 in patients</li> <li>• Immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection</li> <li>• Pregnant and lactating women</li> </ul>

## Summary of changes to the risk management plan over time

### Major changes to the Risk Management Plan over time

Version	Approval Date Procedure	Change
1.2 (W)	At the time of authorization Date of RMP: 04 November 2015 Date of approval: 16 December 2015 EMA/H/C/00002771	<p><b><u>Safety Concerns:</u></b></p> <p><b><u>Important Identified Risks:</u></b></p> <ul style="list-style-type: none"> <li>• Disseminated herpetic infection in severely immunocompromised individuals (those with any severe congenital or acquired cellular and/or humoral immune deficiency)</li> <li>• Accidental exposure of HCP to talimogene laherparepvec</li> <li>• Obstructive airway disorder</li> <li>• Immune-mediated adverse reactions</li> <li>• Plasmacytoma at the injection site</li> <li>• Deep vein thrombosis</li> <li>• Cellulitis at site of injection</li> </ul> <p><b><u>Important Potential Risks:</u></b></p> <ul style="list-style-type: none"> <li>• Disseminated herpetic infection in immunocompromised patients (such as those with HIV/AIDS, leukemia, lymphoma, common variable immunodeficiency, or those who require high-dose steroids or other immunosuppressive agents)</li> <li>• Transmission of talimogene laherparepvec from patient to close contacts or HCPs via direct contact with injected lesions or body fluids resulting in symptomatic infection (primary or reactivation)</li> <li>• Symptomatic talimogene laherparepvec infection in non-tumor tissue in treated patients</li> <li>• Symptomatic herpetic infection due to latency and reactivation of talimogene laherparepvec or wild-type HSV-1 in patients</li> <li>• Immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection</li> <li>• Combination with other therapies like chemotherapy or immunosuppressive agents</li> <li>• Recombination of talimogene laherparepvec with wild-type HSV-1 virus may occur</li> <li>• Impaired wound healing at site of injection</li> <li>• Delayed next line treatment in non-responders</li> <li>• Loss of efficacy in patients treated with systemic acyclovir for complications</li> </ul>

Version	Approval Date Procedure	Change
1.2 (W) continued		<p><u>Important Potential Risks (continued):</u></p> <ul style="list-style-type: none"> <li>• Talimogene laherparepvec-mediated anti-GM-CSF antibody response</li> </ul> <p><u>Missing Information:</u></p> <ul style="list-style-type: none"> <li>• Additional clinical biodistribution and shedding data in melanoma</li> <li>• Pregnant and lactating women</li> <li>• Pediatric patients</li> <li>• Patients below the age of 40 years</li> <li>• Patients with renal or hepatic impairment</li> <li>• Treatment of patients with cardiac impairment</li> <li>• Patients of race or ethnic origin other than white</li> <li>• Long-term safety data</li> <li>• Long-term efficacy data</li> <li>• Treatment of patients with bone metastases</li> <li>• Treatment of patients with cerebral metastases</li> <li>• Treatment of patients with more than 3 visceral lesions</li> <li>• Treatment of patients with metastatic lesions greater than 3 cm</li> <li>• Treatment of patients with ocular melanoma</li> <li>• Treatment of patients with mucosal melanoma</li> </ul> <p><b><u>Pharmacovigilance plan</u></b></p> <p>Specific Adverse Drug Reaction Follow.up Forms:</p> <ul style="list-style-type: none"> <li>• Suspected IMLYGIC (talimogene laherparepvec) or Herpes Virus Associated Adverse Event</li> <li>• Clinical Trial or Postmarket Report of Suspected talimogene laherparepvec Associated Adverse Event for HCP or Close Contact</li> <li>• Suspected IMLYGIC Autoimmune Adverse Event</li> <li>• Pregnancy and lactation follow-up forms</li> </ul>



Version	Approval Date Procedure	Change
1.2 (W) continued		<p><b><u>Category 1 to 3 Studies:</u></b></p> <ul style="list-style-type: none"> <li>• Study 20120139 A registry study to evaluate the survival and long-term safety of subjects with melanoma who previously received talimogene laherparepvec.</li> <li>• Study 20130193 A postmarketing, prospective cohort study of patients treated with talimogene laherparepvec in clinical practice to characterize the</li> <li>• Study 20120324 A phase 2, multicenter, single-arm trial to evaluate the biodistribution and shedding of talimogene laherparepvec in subjects with unresected, stage IIIB to IVM1c melanoma.</li> <li>• Study 20110261 A phase 1, open-label, dose de-escalation study to evaluate the tolerability, safety, and activity of talimogene laherparepvec in children from birth to &lt; 18 years of age with melanoma or with advanced non-central nervous system tumors that are amenable to direct injection and for which no effective treatment is known.</li> <li>• Study Number: To be determined. A Randomized, controlled study to evaluate the safety and efficacy of talimogene laherparepvec in children from birth to &lt; 18 years of age with a pediatric solid malignant tumor as part of a multi-modal treatment approach.</li> </ul> <p><b><u>Postauthorization Efficacy Plan:</u></b></p> <ul style="list-style-type: none"> <li>• Study 20120139 A registry study to evaluate the survival and long-term safety of subjects with melanoma who previously received talimogene laherparepvec.</li> </ul> <p><b><u>Risk Minimization Measures:</u></b></p> <ul style="list-style-type: none"> <li>• Physician Education Booklet</li> <li>• Managed distribution program</li> <li>• Patient safety brochure and patient alert card</li> </ul>
2.0	<p>Date of RMP: 16 August 2016</p> <p>Date of approval: 07 October 2016 EMA/H/C/002771/ IB/0007</p>	<p><b><u>Safety Concerns:</u></b> No changes</p> <p><b><u>Pharmacovigilance Plan:</u></b> Due dates of Studies 20120324 and 20110261 were updated.</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p>

Version	Approval Date Procedure	Version
3.0	Date of RMP: 03 October 2017  Date of approval: 13 November 2017 EMEA/H/C/002771/ IB/0017	<p><b><u>Safety Concerns:</u></b> No changes</p> <p><b><u>Pharmacovigilance Plan:</u></b> Due dates of final analysis clinical study report for Study 20120324 was updated.</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p>
4.0	Date of RMP: 10 September 2018 EMEA/H/C/002771/ II/0028	<p><b><u>Safety Concerns:</u></b> The following important identified risks were reclassified as not important and removed from the RMP:</p> <ul style="list-style-type: none"> <li>• Obstructive airway disorder</li> <li>• Plasmacytoma at the injection site</li> <li>• Deep vein thrombosis</li> <li>• Cellulitis at site of injection</li> </ul> <p>The following important potential risks were reclassified as not important and removed from the RMP:</p> <ul style="list-style-type: none"> <li>• Combination with other therapies like chemotherapy or immunosuppressive agents</li> <li>• Recombination of talimogene laherparepvec with wild-type HSV-1 virus may occur</li> <li>• Impaired wound healing at site of injection</li> <li>• Delayed next line treatment in non-responders</li> <li>• Loss of efficacy in patients treated with systemic acyclovir for complications</li> </ul> <p>The following missing information was removed from the RMP:</p> <ul style="list-style-type: none"> <li>• Use in patients below the age of 40 years</li> <li>• Use in patients with renal or hepatic impairment</li> <li>• Treatment of patients with cardiac impairment</li> <li>• Use in patients of race or ethnic origin other than white</li> <li>• Treatment of patients with bone metastases</li> <li>• Treatment of patients with active cerebral metastases</li> <li>• Treatment of patients with more than 3 visceral lesions</li> <li>• Treatment of patients with metastatic lesions greater than 3 cm</li> <li>• Treatment of patients with ocular melanoma</li> <li>• Treatment of patients with mucosal melanoma</li> </ul> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b></p> <ul style="list-style-type: none"> <li>• Study 20120139 was removed as a postauthorization efficacy study.</li> </ul> <p><b><u>Risk Minimization Measures:</u></b> No change</p>

Version	Approval Date Procedure	Version
5.0	Date of RMP: 15 November 2018 EMA/H/C/002771/ II/0029	<p><b><u>Safety Concerns:</u></b> The following missing information was removed from the RMP:</p> <ul style="list-style-type: none"> <li>Additional clinical biodistribution and shedding data in melanoma</li> </ul> <p><b><u>Pharmacovigilance Plan:</u></b> • Study 20120324 removed as study complete</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p>
4.1	Date of RMP: 29 January 2019 EMA/H/C/002771/ II/0028	<p><b><u>Safety Concerns:</u></b> The following safety concern was reclassified as an important potential risk and added to the RMP:</p> <ul style="list-style-type: none"> <li>Combination with other therapies like chemotherapy or immunosuppressive agents</li> </ul> <p>The following safety concern was reclassified as missing information and added to the RMP:</p> <ul style="list-style-type: none"> <li>Treatment of patients with metastatic lesions greater than 3 cm</li> </ul> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> The following postauthorization efficacy studies were added to the RMP:</p> <ul style="list-style-type: none"> <li>Study 20110265</li> <li>Study 20110266</li> </ul> <p><b><u>Risk Minimization Measures:</u></b> No change</p> <p><b><u>Annexes:</u></b></p> <ul style="list-style-type: none"> <li><a href="#">Annex 5</a>: Protocols for Studies 20110265 and 20110266 were appended</li> </ul>
5.1	Date of RMP: 15 February 2019 EMA/H/C/002771/ II/0029	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p> <p><b><u>Annexes:</u></b> No change</p> <p><b><u>Other Changes:</u></b> Information on detectable DNA from swabs of the exterior of occlusive dressings added to justification text for removal of the missing information 'Additional clinical biodistribution and shedding data in melanoma.'</p>

Version	Approval Date Procedure	Version
6.0	Date of RMP: 27 March 2019 EMA/H/C/002771/ II/0028 EMA/H/C/002771/ II/0029	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p> <p><b><u>Annexes:</u></b> No change</p> <p><b><u>Other Changes:</u></b> Consolidation of EU RMP versions 4.1 and 5.1.</p>
7.0	Date of RMP: 26 April 2019 To be confirmed by EMA	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> The following studies were added to evaluate the effectiveness of additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Study 20180062</li> <li>• Study 20180099</li> </ul> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b></p> <ul style="list-style-type: none"> <li>• Plans to evaluate the effectiveness of the additional risk minimization measures were updated as follows: <ul style="list-style-type: none"> <li>- Effectiveness of the managed distribution program will be measured by conducting an internal evaluation of managed distribution process metrics</li> <li>- Effectiveness of the Physician Education Booklet will be measured using a cross-sectional survey (Study 20180099)</li> <li>- Effectiveness of the patient safety brochure and patient alert card will be measured using a cross-sectional survey (Study 20180062)</li> </ul> </li> <li>• Patient safety brochure and patient alert card removed as additional risk minimization measures for the important identified risk of accidental exposure of healthcare provider to talimogene laherparepvec and the important potential risk of immunocompromised patients treated with talimogene laherparepvec and suffering from concomitant infection as these measures are not relevant for these risks.</li> </ul> <p><b><u>Annexes:</u></b></p> <ul style="list-style-type: none"> <li>• <a href="#">Annex 2:</a> Updated to include Studies 20180062 and 20180099</li> <li>• <a href="#">Annex 3:</a> Protocols for Studies 20180062 and 20180099 were appended</li> </ul>

Version	Approval Date Procedure	Version
8.0	Date of RMP: 13 June 2019 EMA/H/C/002771/ IB/0035	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> Clinical study report due date updated for Study 20110265</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p> <p><b><u>Annexes:</u></b> No change</p>
8.1	Date of RMP: 15 July 2019 EMA/H/C/002771/ IB/0035	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> No change</p> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b> No change</p> <p><b><u>Annexes:</u></b> No change</p> <p><b><u>Other changes:</u></b> Removal of all of EU RMP v7.0 changes (procedure EMA/H/C/002771/II/0034) so that only v8.0/v8.1 changes (procedure EMA/H/C/002771/IB/0035) are contained within the current EU RMP.</p>
9.0	Date of RMP: 06 August 2019 To be confirmed by EMA	<p><b><u>Safety Concerns:</u></b> No change</p> <p><b><u>Pharmacovigilance Plan:</u></b> The following studies were added to evaluate the effectiveness of additional risk minimization measures:</p> <ul style="list-style-type: none"> <li>• Study 20180062</li> <li>• Study 20180099</li> </ul> <p><b><u>Postauthorization Efficacy Plan:</u></b> No change</p> <p><b><u>Risk Minimization Measures:</u></b></p> <ul style="list-style-type: none"> <li>• Plans to evaluate the effectiveness of the additional risk minimization measures were updated as follows: <ul style="list-style-type: none"> <li>- Effectiveness of the managed distribution program will be measured by conducting an internal evaluation of managed distribution process metrics</li> <li>- Effectiveness of the Physician Education Booklet will be measured using a cross-sectional survey (Study 20180099)</li> <li>- Effectiveness of the patient safety brochure and patient alert card will be measured using a cross-sectional survey (Study 20180062)</li> </ul> </li> </ul> <p>Patient safety brochure and patient alert card removed as additional risk minimization measures for the important identified risk of accidental exposure of healthcare provider to talimogene laherparepvec and the important potential risk of immunocompromised patients treated with talimogene laherparepvec and suffering from</p>

		concomitant infection as these measures are not relevant for these risks. <b>Annexes:</b> <ul style="list-style-type: none"> <li>• <a href="#">Annex 2</a>: Updated to include Studies 20180062 and 20180099</li> <li>• <a href="#">Annex 3</a>: Protocols for Studies 20180062 and 20180099 were appended</li> </ul>
Version	Approval Date Procedure	Version
9.1	Date of RMP: 12 June 2020	<b><u>Other Changes:</u></b> <b><u>To extend the final report date for the category 3 Study 20180099 from 31 August 2020 to 28 February 2021</u></b>
9.2	Date of RMP: 14 December 2020 Procedure: EMA/H/C/002771/ IB/0042	<b>Other Changes:</b> To extend the final report date for the category 3 Study 20180062 from 31 March 2021 to March 2022
9.3	Date of RMP: 25 January 2021 Procedure: EMA/H/C/002771/ IB/0042	<b>Other Changes:</b> To correct the status of the category 3 Studies 20130193, 20180062, and 20180099 from planned to ongoing.

This summary was created in November 2021.