# SHINGRIX 

Version : GDSv07/IPIv02

## SHINGRIX

## Herpes zoster (HZ, or shingles) vaccine (non-live recombinant, ASO1 $_{\mathrm{B}}$ adjuvanted)

## Qualitative and Quantitative Composition

After reconstitution, 1 dose ( 0.5 ml ) contains 50 micrograms of gE antigen ${ }^{1}$ adjuvanted with $\mathrm{ASO1}_{\mathrm{B}}{ }^{2}$.
${ }^{1}$ Varicella Zoster Virus (VZV) glycoprotein E (gE) produced by recombinant DNA technology in Chinese Hamster Ovary (CHO) cells
${ }^{2}$ The GlaxoSmithKline proprietary $\mathrm{ASO1}_{\mathrm{B}}$ Adjuvant System is composed of the plant extract Quillaja saponaria Molina, fraction 21 (QS-21) ( 50 micrograms) and 3-O-desacyl-4'-monophosphoryl lipid A (MPL) from Salmonella minnesota ( 50 micrograms)
The powder is white. The suspension is an opalescent, colourless to pale brownish liquid.

## Clinical Information

## Indications

SHINGRIX is indicated for the prevention of herpes zoster (HZ) and HZ-related complications, such as post-herpetic neuralgia (PHN), in:

- adults 50 years of age or older;
- adults 18 years of age or older at increased risk of HZ .

The use of SHINGRIX should be based on official recommendations.

## Dosage and Administration

Pharmaceutical form: powder and suspension for suspension for injection
The immunisation schedules for SHINGRIX should be based on official recommendations.

## Posology

The primary vaccination schedule consists of two doses of 0.5 ml each; an initial dose followed by a second dose 2 to 6 months later

For subjects who are immunodeficient, immunosuppressed or likely to become immunosuppressed due to known disease or therapy, and whom would benefit from a shorter vaccination schedule, the second dose can be given 1 to 2 months after the initial dose (see Pharmacodynamic Effects).

The need for booster doses has not been established.
SHINGRIX can be given with the same schedule in individuals previously vaccinated with live attenuated HZ vaccine (see Pharmacodynamic Effects).
SHINGRIX is not indicated for prevention of primary varicella infection.

## Method of administration

SHINGRIX is for intramuscular injection only, preferably in the deltoid muscle
For instructions on reconstitution of the medicinal product before administration, see Use and Handling.

## Contraindications

Hypersensitivity to the active substances or to any component of the vaccine (see Qualitative and Quantitative Composition and List of Excipients).

## Warnings and Precautions

## Prior to immunization

It is good clinical practice to precede vaccination by a review of the medical history (especially with regard to previous vaccination and possible occurrence of undesirable events) and a clinical examination.
As with all injectable vaccines, appropriate medical treatment and supervision should always be readily available in case of an anaphylactic event following the administration of the vaccine.
As with other vaccines, vaccination with SHINGRIX should be postponed in subjects suffering from an acute severe febrile illness. The presence of a minor infection, such as a cold, should not result in the deferral of vaccination.
As with any vaccine, a protective immune response may not be elicited in all vaccinees.
In a post-marketing observational study in individuals aged 65 years or older, an increased risk of Guillain-Barré syndrome (estimated 3 excess cases per million doses administered) was observed during the 42 days following vaccination with SHINGRIX. Available information is insufficient to determine a causal relationship with SHINGRIX.

## Precautions for use

Do not administer the vaccine intravascularly, intradermally or subcutaneously.
Maladministration via the subcutaneous route may lead to an increase in transient local reactions.
As with other vaccines administered intramuscularly, SHINGRIX should be given with caution to individuals with thrombocytopenia or any coagulation disorder since bleeding may occur following an intramuscular administration to these subjects.
Syncope (fainting) can occur following, or even before, any vaccination as a psychogenic response to the needle injection. It is important that procedures are in place to avoid injury from faints.

## Interactions

## Use with other vaccines.

SHINGRIX can be given concomitantly with unadjuvanted seasonal influenza vaccine, 23 -valent pneumococcal polysaccharide vaccine (PPV23), pneumococcal conjugate vaccine (PCV) or reduced antigen diphtheria-tetanusacellular pertussis vaccine (dTpa) (see Pharmacodynamic Effects).
The adverse reactions of fever and shivering were more frequent when PPV23 vaccine was co-administered with SHINGRIX compared to when SHINGRIX was given alone (see Adverse Reactions).

If SHINGRIX is to be given at the same time as another injectable vaccine, the vaccines should always be administered at different injection sites.

## Pregnancy and Lactation

Fertility
Animal studies indicate no effects of SHINGRIX on male or female fertility.

## Pregnancy

There are no data on the use of SHINGRIX in pregnant women. Animal studies performed with SHINGRIX administered to female rats do not indicate any harmful effects with respect to pregnancy (see Non-clinical information).

## Lactation

## The effect on breast-fed infants of administration of SHINGRIX to their mothers has not been studied

## Effects on Ability to Drive and Use Machines

No studies on the effects of SHINGRIX on the ability to drive and use machines have been performed.

## Adverse Reactions <br> Clinical trial data

The safety profile presented below is based on a pooled analysis of more than 14,500 adults $\geq 50$ years of age, who have received at least one dose of SHINGRIX. These data were generated in placebo-controlled clinical studies (conducted in Europe, North America, Latin America, Asia and Australia) where SHINGRIX was administered according to a 0,2 -month schedule.
Additionally, in clinical studies, 1,587 subjects $\geq 18$ years of age who are immunodeficient or immunosuppressed due to disease or therapy (referred to as immunocompromised (IC)), were vaccinated with at least 1 dose of SHINGRIX. The reported adverse reactions were consistent with those presented in the Table below.
Adverse reactions reported are listed according to the following frequency:
Very common ( $\geq 1 / 10$ ); Common ( $\geq 1 / 100$ to $<1 / 10$ ); Uncommon ( $\geq 1 / 1,000$ to $<1 / 100$ ); Rare ( $\geq 1 / 10,000$ to $<1 / 1,000$ ); Very rare ( $<1 / 10,000$ )

| System Organ Class | Frequency | Adverse reactions |
| :--- | :--- | :--- |
| Nervous system disorders | Very common | headache |
| Gastrointestinal disorders | Very common | gastrointestinal symptoms (including <br> nausea, vomiting, diarrhoea and/or <br> abdominal pain) |
| Musculoskeletal and connective tissue <br> disorders | Very common | myalgia |
|  | Uncommon | arthralgia |
| General disorders and administration <br> site conditions | Very common | injection site reactions (such as pain, <br> redness, swelling), fatigue, chills, fever |

Overall, there was a higher incidence of some adverse reactions in younger age groups. However, the overall frequency and severity of these events did not indicate a clinically meaningful different reactogenicity profile in the younger age strata. In IC adult studies, there was a higher incidence of pain at the injection site, fatigue, myalgia, headache, shivering and fever in subjects aged 18 to 49 years compared with those aged 50 years and older. In older adult studies, there was a higher incidence of pain and swelling at the injection site, fatigue, myalgia, headache, shivering, fever and gastrointestinal symptoms in subjects aged 50 to 69 years compared with those aged 70 years and older.
In a clinical study where 119 subjects $\geq 50$ years of age were vaccinated with SHINGRIX following a 0, 6-month schedule, the safety profile was similar to that observed in subjects vaccinated with SHINGRIX following a 0,2 month schedule.
In a clinical study including 865 adults $\geq 50$ years of age, fever and shivering were reported more frequently when PPV23 vaccine was co-administered with SHINGRIX ( $16 \%$ and $21 \%$, respectively) compared to when SHINGRIX was given alone ( $7 \%$ for both adverse reactions).

## Post-marketing data

| System Organ Class | Frequency | Adverse reactions |
| :---: | :---: | :---: |
| Immune system disorders | Rare | hypersensitivity reactions including rash, <br> urticaria, angioedema |

## Overdose

Insufficient data are available.

## Pharmacological Properties

## Pharmacodynamics

## ATC Code

Pharmacotherapeutic group: Varicella zoster vaccines, ATC code: Јо7вкоз.

## Mechanism of Action

SHINGRIX is designed to induce antigen-specific cellular and humoral immune responses in individuals with pre existing immunity against VZV.
Non-clinical data show that ASO1s induces a local and transient activation of the innate immune system through specific molecular pathways. This facilitates the recruitment and activation of antigen presenting cells carrying gEderived antigens in the draining lymph node, which in turn leads to the generation of gE-specific CD4+ T cells and
antibodies. The adjuvant effect of $\mathrm{ASO1}_{\mathrm{B}}$ is the result of interactions between MPL and QS-21 formulated in liposomes.

## Pharmacodynamic Effects

## Efficacy of SHINGRIX

## Efficacy Against Herpes Zoster (HZ) and Post-Herpetic Neuralgia (PHN)

Two phase III, placebo-controlled, observer-blind efficacy studies of SHINGRIX were conducted in adults $\geq 50$ years with 2 doses administered 2 months apart:

- Zoster-006 (ZOE-50): total vaccinated cohort (TVC) of 15,405 subjects $\geq 50$ years who received at least one dose of either SHINGRIX ( $\mathrm{N}=7,695$ ) or placebo ( $\mathrm{N}=7,710$ ).
- Zoster-022 (ZOE-70): TVC of 13,900 subjects $\geq 70$ years who received at least one dose of either SHINGRIX ( $\mathrm{N}=6,950$ ) or placebo $(\mathrm{N}=6,950)$.
Two phase III, placebo-controlled, observer-blind studies evaluating SHINGRIX efficacy were conducted in IC adults $\geq 18$ years with 2 doses administered 1-2 months apart:
- Zoster-002: TVC of 1,846 autologous hematopoietic stem cell transplants (aHSCT) recipients who received at least one dose of either SHINGRIX ( $\mathrm{N}=922$ ) or placebo ( $\mathrm{N}=924$ ) post-transplant.
- Zoster-039: TVC of 562 subjects with hematologic malignancies who received at least one dose of either SHINGRIX ( $\mathrm{N}=283$ ) or placebo $(\mathrm{N}=279)$ during a cancer therapy course or after the full cancer therapy course.
Incidence of HZ and PHN cases as well as vaccine efficacy were evaluated in the modified Total Vaccinated Cohort (mTVC i.e. excluding subjects who did not receive the second dose of vaccine or who had a confirmed diagnosis of HZ within one month after the second dose).
SHINGRIX significantly decreased the incidence of HZ and PHN compared with placebo in:
- adults $\geq 50$ years (Zoster-006): 6 vs. 210 HZ cases and 0 vs. 18 PHN cases.
- adults $\geq 70$ years (pooled analysis of Zoster- 006 and Zoster-022): 25 vs. 284 HZ cases and 4 vs .36 PHN cases.
- adults $\geq 18$ years with aHSCT (Zoster-002): 49 vs. 135 HZ cases and 1 vs. 9 PHN cases.
- adults $\geq 18$ years with hematologic malignancies (Zoster-039): 2 vs .14 HZ cases (PHN was not assessed as study endpoint). Vaccine efficacy was calculated post-hoc.
Vaccine efficacy results are presented in Table 1.
Table 1: SHINGRIX efficacy against HZ and PHN (mTVC)

| Age (years) | HZ |  |  | PHN |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | N | Efficacy (\%) | 95\% CI | N | Efficacy (\%) | 95\% CI |
| Zoster-006* |  |  |  |  |  |  |
| $\geq 50$ | 7,344 | 97.2 | 93.7; 99.0 | 7,340 | 100.0 | 77.1; 100.0 |
| 50-59 | 3,492 | 96.6 | 89.6; 99.4 | 3,491 | 100.0 | 40.8; 100.0 |
| $\geq 60$ | 3,852 | 97.6 | 92.7; 99.6 | 3,849 | 100.0 | 55.2; 100.0 |
| 60-69 | 2,141 | 97.4 | 90.1; 99.7 | 2,140 | $100.0^{5}$ | < 0; 100.0 |
| Pooled Zoster-006 and Zoster-022** |  |  |  |  |  |  |
| $\geq 70$ | 8,250 | 91.3 | 86.8; 94.5 | 8,250 | 88.8 | 68.7; 97.1 |
| 70-79 | 6,468 | 91.3 | 86.0; 94.9 | 6,468 | 93.0 | 72.4; 99.2 |
| $\geq 80$ | 1,782 | 91.4 | 80.2; 97.0 | 1,782 | $71.2^{5}$ | <0; 97.1 |
| Zoster-002*** (aHSCT recipients ${ }^{\text {\# }}$ ) |  |  |  |  |  |  |
| $\geq 18$ | 870 | 68.2 | 55.5; 77.6 | 870 | 89.3 | 22.5; 99.8 |
| 18-49 | 213 | 71.8 | 38.7; 88.3 | 213 | $100.0^{5}$ | < 0; 100.0 |
| $\geq 50$ | 657 | 67.3 | 52.6; 77.9 | 657 | 88.0 | 10.4; 99.8 |

Zoster-039 (hematologic malignancy patients")

| $\geq 18$ | 259 | $87.2^{* * * *}$ | $44.2 ; 98.6$ | - | - |
| :--- | :--- | :---: | :---: | :---: | :---: |
| N | Number of evaluable subjects |  |  |  |  |
| Cl | Confidence interval |  |  |  |  |
| $*$ | Over a median follow-up period of 3.1 and 4.1 years for reporting HZ and PHN cases, |  |  |  |  |
| respectively |  |  |  |  |  |
| $* *$ |  |  |  |  |  |
| $* * *$ | Over a median follow-up period of 4.0 years for reporting HZ and PHN cases |  |  |  |  |
| $* * * *$ | Over a median follow-up period of 21 months for reporting HZ and PHN cases |  |  |  |  |
| $\#$ | VE calculation was performed post-hoc; median follow-up period of 11.1 months |  |  |  |  |
| \# | antiviral prophylaxis in line with the local standard of care was permitted |  |  |  |  |
| $\S$ | Not statistically significant |  |  |  |  |

Zoster-006 mTVC: $\mathrm{N}($ SHINGRIX $)=7,344, \mathrm{~N}$ (Placebo) $=7,415$
Pooled analysis of Zoster-006 and Zoster-022 mTVC: $\mathrm{N}($ SHINGRIX $)=8,250, \mathrm{~N}$ (Placebo) $=8,346$
Zoster-002 mTVC: $\mathrm{N}($ SHINGRIX $)=870, \mathrm{~N}$ (Placebo) $=851$
Zoster-039 mTVC: $\mathrm{N}($ SHINGRIX $)=259, \mathrm{~N}$ (Placebo) $=256$
In the fourth year after vaccination, the efficacy against HZ was $93.1 \%$ ( $95 \% \mathrm{Cl}: 81.2 ; 98.2$ ) and $87.9 \%$ ( $95 \%$ CI: 73.3; 95.4) in subjects $\geq 50$ years (Zoster-006) and subjects $\geq 70$ years (pooled Zoster-006 and Zoster-022), respectively.
In Zoster-002, during a follow-up period starting 1 month post-dose 2 (i.e. corresponding to approximately 6 months after aHSCT) until 1 year after aHSCT, when the risk for HZ is the highest, the efficacy against HZ was $76.2 \%$ ( $95 \% \mathrm{CI}: 61.1$; 86.0).

## Efficacy Against Other HZ-Related Complications

The evaluated HZ -related complications (other than PHN) were: HZ vasculitis, disseminated disease, ophthalmic disease, neurologic disease including stroke, and visceral disease.
In the pooled analysis of Zoster-006 and Zoster-022, SHINGRIX significantly reduced HZ-related complications by $93.7 \%$ ( $95 \% \mathrm{CI}: 59.5 ; 99.9$ ) and $91.6 \%$ ( $95 \% \mathrm{CI}: 43.3 ; 99.8$ ) in subjects $\geq 50$ years ( 1 vs. 16 cases) and subjects $\geq 70$ years (1 vs. 12 cases), respectively.
In Zoster-002, SHINGRIX significantly reduced HZ-related complications by $77.8 \%$ ( $95 \% \mathrm{CI}: 19.0 ; 96.0$ ) in aHSCT recipients $\geq 18$ years ( 3 vs 13 cases).
In addition, in Zoster-002, SHINGRIX significantly reduced HZ-related hospitalisations by $84.7 \%$ ( $95 \% \mathrm{CI}$ : 32.1; 96.6) ( 2 vs. 13 cases).

## Effect of SHINGRIX on HZ-Associated Pain

In Zoster-022, SHINGRIX significantly reduced the use and the duration of HZ -associated pain medication by $39.6 \%$ ( $95 \%$ CI: $10.7 ; 64.8$ ) and $49.3 \%$ ( $95 \%$ CI: $2.9 ; 73.5$ ), respectively, in subjects $\geq 70$ years with at least one confirmed HZ episode. The median duration of pain medication use was 30.0 and 38.0 days in the SHINGRIX and placebo group, respectively.
Overall there was a general trend towards less severe HZ-associated pain in subjects vaccinated with SHINGRIX compared to placebo.
In Zoster-002, SHINGRIX significantly reduced the duration of severe 'worst' HZ-associated pain by $38.5 \%$ ( $95 \% \mathrm{Cl}$ : 11.0; 57.6) in aHSCT recipients $\geq 18$ years with at least one confirmed HZ episode.

## Immunogenicity of SHINGRIX

An immunological correlate of protection has not been established; therefore the level of immune response that provides protection against HZ is unknown.

In adults $\geq 50$ years, the immune responses to SHINGRIX were evaluated in a subset of subjects from the phase II efficacy studies Zoster-006 [humoral immunity and cell-mediated immunity (CMI)] and Zoster-022 (humora immunity). The gE-specific immune responses (humoral and CMI) elicited by SHINGRIX at 1 month post-dose 2 are presented in Tables 2 and 3, respectively.
Table 2: Humoral immunogenicity of SHINGRIX in adults $\geq \mathbf{5 0}$ years at 1 month post-dose 2 (ATP cohort for immunogenicity)

Anti-gE immune response ${ }^{\wedge}$

| Anti-gE immune response^^ |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: |
| Age <br> group <br> (years) | N | VRR§ (\%) <br> $(95 \% \mathrm{CI})$ | GMC <br> $(95 \% \mathrm{Cl})$ | Median fold increase of <br> concentrations vs pre- <br> vaccination (Q1; Q3) |

Zoster-006

| $\geq 50$ | 1,070 | 98.5 <br> $(97.6 ; 99.1)$ | $52,376.6$ <br> $(50,264.1 ; 54,577.9)$ | 41.9 <br> $(20.8 ; 86.9)$ |
| :---: | :---: | :---: | :---: | :---: |

Pooled Zoster-006 and Zoster-022

| $\geq 70$ | 742 | $\begin{gathered} 96.6 \\ (95.1 ; 97.8) \end{gathered}$ | $\begin{gathered} 49,691.5 \\ (47,250.8 ; 52,258.2) \end{gathered}$ | $\begin{gathered} 34.3 \\ (16.7 ; 68.5) \end{gathered}$ |
| :---: | :---: | :---: | :---: | :---: |
| ATP | According-To-Protocol |  |  |  |
|  | Anti-gE immune response = anti-gE antibody levels, measured by anti-gE enzyme-linked |  |  |  |
| immunosorbent assay (gE ELISA) |  |  |  |  |
| N | Number of evaluable subjects at the specified time point (for the GMC) |  |  |  |
| § | Vaccine response rate (VRR) for anti-gE is defined as the percentage of subjects who have at |  |  |  |
| least a 4 -fold increase in the post-dose 2 anti-gE antibodies concentration as compared to the pre-vaccination anti-gE antibodies (subjects seropositive at baseline), or as compared to the anti-gE antibodies cut-off value for seropositivity (subjects seronegative at baseline) |  |  |  |  |
| Cl | Confidence interval |  |  |  |
| GMC | Geometric Mean Concentration |  |  |  |
| Q1; Q3 | First and third quartiles |  |  |  |

At 3 years post-dose 2, the median fold increase over baseline was 9.3 (Q1: 4.9; Q3: 19.5) in adults $\geq 50$ years (Zoster-006) and 7.2 (Q1: 3.5; Q3: 14.5) in adults $\geq 70$ years (pooled Zoster-006 and Zoster-022).
Table 3: Cell-mediated immunogenicity of SHINGRIX in adults $\geq 50$ years at 1 month post-dose 2 (ATP cohort for immunogenicity)

## gE-specific CD4[2+] T cell response ${ }^{\wedge}$

| Age |
| :---: | :---: | :---: | :---: |
| group |
| (years) |$\quad \mathrm{N} \quad$| Median frequency |
| :---: |
| (Q1; Q3) |

Zoster-006

| $\geq 50$ | 164 | $1,844.1$ <br> $(1,253.6 ; 2,932.3)$ | 24.6 <br> $(9.9 ; 744.2)$ |
| :---: | :---: | :---: | :---: |
| $\geq 70^{*}$ | 52 | $1,494.6$ <br> $(922.9 ; 2,067.1)$ | 33.2 <br> $(10.0 ; 1,052.0)$ |

ATP According-To-Protocol
$\wedge \quad$ gE-specific CD4[2+] T cell response $=\mathrm{gE}$-specific CD4 +T cell activity, measured by intracellular cytokine staining (ICS) assay (CD4[2+] T cells = CD4+ T cells expressing at least 2 of 4 selected immune markers)
$\mathrm{N} \quad$ Number of evaluable subjects at the specified time point for the median frequency.
Q1; Q3 First and third quartiles
The gE-specific CD4[2+] data in the $\geq 70$ YOA group were only generated in Zoster-006 because $\mathrm{CD} 4+\mathrm{T}$ cell activity was not assessed in Zoster-022
At 3 years post-dose 2, in Zoster-006, the median fold increase over baseline was 7.9 (Q1: 2.7; Q3: 31.6) in adults $\geq 50$ years and 7.3 (Q1:1.7; Q3: 31.6) in adults $\geq 70$ years.
Data from a phase II, open-label, single group, follow-up clinical study in adults $\geq 60$ years (Zoster-024) indicate that the vaccine-induced immune response (humoral and CMI) persists up to Month 72 (approximately 6 years postdose 1 i.e. 70 months post-dose 2 ), following a 0,2 -month schedule ( $N=119$ ).
The median anti-gE antibody concentration was greater than 7 -fold above the baseline pre-vaccination median concentration. The median frequency of gE-specific CD4[2+] T cells was greater than 3.7-fold above baseline prevaccination median frequency.
In IC adults $\geq 18$ years, the humoral and CMI responses to SHINGRIX were evaluated in:

- one phase I/II study: Zoster-015 (HIV infected subjects);
- one phase II/III study: Zoster-028 (patients with solid tumors undergoing chemotherapy);
- three phase III studies: Zoster-002 (aHSCT recipients vaccinated post-transplant), Zoster-039 (patients with hematologic malignancies vaccinated during a cancer therapy course or after the full cancer therapy course) and Zoster-041 (renal transplant recipients on chronic immunosuppressive treatment at the time of vaccination).
The gE-specific immune responses (humoral and CMI) elicited by SHINGRIX at 1 month post-dose 2 in all IC populations studied are presented in Tables 4 and 5, respectively.
Table 4: Humoral Immunogenicity of SHINGRIX in IC adults $\geq 18$ years at 1 month post-dose 2 (ATP cohort for immunogenicity)


## Anti-gE immune response^

| N | $\begin{aligned} & \text { VRR }^{\S} \text { (\%) } \\ & \text { (95\% CI) } \end{aligned}$ | $\begin{gathered} \text { GMC } \\ (95 \% \mathrm{CI}) \end{gathered}$ | Median fold increase of concentrations vs prevaccination (Q1; Q3) |
| :---: | :---: | :---: | :---: |
| Zoster-002 (aHSCT recipients) |  |  |  |
| 82 | $\begin{gathered} \hline 67.1 \\ (55.8 ; 77.1) \\ \hline \end{gathered}$ | $\begin{gathered} 12,753.2 \\ (7,973.0 ; 20,399.4) \\ \hline \end{gathered}$ | $\begin{gathered} 14.1 \\ (1.7 ; 137.0) \\ \hline \end{gathered}$ |
| Zoster-028 (solid tumor patients) |  |  |  |
| 87 | $\begin{gathered} 86.2 \\ (77.1 ; 92.7) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 18,291.7 \\ (14,432.1 ; 23,183.5) \\ \hline \end{gathered}$ | $\begin{gathered} 21.5 \\ (7.0 ; 45.2) \\ \hline \end{gathered}$ |


| Zoster-039 (hematologic malignancy patients) |  |  |  |
| :---: | :---: | :---: | :---: |
| 217 | $\begin{gathered} 65.4 \\ (58.7 ; 71.7) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 13,445.6 \\ (10,158.9 ; 17,795.6) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 17.2 \\ (1.4 ; 87.4) \\ \hline \end{gathered}$ |
| Zoster-041 (renal transplant recipients) |  |  |  |
| 121 | $\begin{gathered} \hline 80.2 \\ (71.9 ; 86.9) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 19,163.8 \\ (15,041.5 ; 24,416.0) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 15.1 \\ (6.1 ; 35.0) \\ \hline \end{gathered}$ |
| Zoster-015 (HIV infected subjects) |  |  |  |
| 53 | $\begin{gathered} 98.1 \\ (89.9 ; 100) \\ \hline \end{gathered}$ | $\begin{gathered} \hline 42,723.6 \\ (31,233.0 ; 58,441.6) \\ \hline \end{gathered}$ | $\begin{gathered} 40.9 \\ (18.8 ; 93.0) \\ \hline \end{gathered}$ |
| ATP According-To-Protocol |  |  |  |
| Anti-gE immune response = anti-gE antibody levels, measured by anti-gE enzyme-linked immunosorbent assay (gE ELISA) |  |  |  |
| N Number of evaluable subjects at the specified time point (for the GMC) |  |  |  |
| least a 4 -fold increase in the post-dose 2 anti-gE antibodies concentration as compared to the pre-vaccination anti-gE antibodies (subjects seropositive at baseline), or as compared to the anti-gE antibodies cut-off value for seropositivity (subjects seronegative at baseline) |  |  |  |
| Cl | ence interval |  |  |
| GMC | tric Mean Co |  |  |
| Q1; Q3 | nd third quar |  |  |

Table 5: Cell-mediated immunogenicity of SHINGRIX in IC adults $\geq 18$ years at 1 month postdose 2 (ATP cohort for immunogenicity)
gE-specific CD4[2+] T cell response^

| N | Median frequency (Q1; Q3) | Median fold increase of frequency vs. pre-vaccination (Q1; Q3) |
| :---: | :---: | :---: |
| Zoster-002 (aHSCT recipients) |  |  |
| 51 | $\begin{aligned} & 6,644.9 \\ & (1,438.3 ; 13,298.6) \\ & \hline \end{aligned}$ | $\begin{aligned} & 109.0 \\ & (34.4 ; 2,716.4) \end{aligned}$ |
| Zoster-028* (solid tumor patients) |  |  |
| 22 | $\begin{aligned} & \hline 778.8 \\ & (393.1 ; 1,098.2) \\ & \hline \end{aligned}$ | $\begin{aligned} & \hline 4.9 \\ & (1.7 ; 33.0) \end{aligned}$ |
| Zoster-039 (hematologic malignancy patients) |  |  |
| 53 | $\begin{aligned} & \hline 3,081.9 \\ & (1,766.2 ; 7,413.6) \end{aligned}$ | $\begin{aligned} & \hline 45.9 \\ & (16.4 ; 2,221.9) \\ & \hline \end{aligned}$ |
| Zoster-041 (renal transplant recipients) |  |  |
| 32 | $\begin{aligned} & \hline 2,149.0 \\ & (569.4 ; 3,695.1) \\ & \hline \end{aligned}$ | $\begin{aligned} & \hline 47.7 \\ & (14.7 ; 439.6) \end{aligned}$ |
| Zoster-015 (HIV infected subjects) |  |  |
| 41 | $\begin{aligned} & \hline 2,809.7 \\ & (1,554.5 ; 4,663.7) \end{aligned}$ | $\begin{aligned} & \hline 23.4 \\ & (8.5 ; 604.1) \\ & \hline \end{aligned}$ |
|  |  |  |
|  |  |  |
|  |  |  |
|  |  |  |
|  |  |  |
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## Immunogenicity Following Concomitant Vaccination

In four phase III, controlled, open-label clinical studies, adults $\geq 50$ years of age were randomized to receive 2 doses of SHINGRIX 2 months apart administered either concomitantly at the first dose or non-concomitantly with unadjuvanted seasonal influenza vaccine ( $\mathrm{N}=828$; Zoster-004), PPV23 vaccine ( $\mathrm{N}=865$; Zoster-035), PCV13 vaccine ( $\mathrm{N}=912$; Zoster-059) or dTpa vaccine formulated with 0.3 milligrams $\mathrm{Al}^{3+}(\mathrm{N}=830$; Zoster-042). The vaccine response rate (in terms of anti-gE antibodies) was $95.8 \%$ ( $95 \% \mathrm{Cl}$ : 93.3; 97.6), $98.3 \%$ ( $95 \% \mathrm{Cl}: 96.4 ; 99.3$ ), $99.1 \%$ ( $95 \% \mathrm{Cl}: 97.6$; 99.7) and $97.8 \%$ ( $95 \%$ CI: 95.8; 99.1) following co-administration of SHINGRIX with the influenza, PPV23, PCV13 and dTpa vaccine respectively. The immune responses of the co-administered vaccines were unaffected, with the exception of lower geometric mean concentrations (GMCs) for one of the pertussis antigens (pertactin) when SHINGRIX is coadministered with the dTpa vaccine. However, these data do not suggest clinically relevant interference.

Immunogenicity in Subjects with a History of HZ Prior to Vaccination
In a phase III, uncontrolled, open-label clinical study (Zoster-033), 96 adults $\geq 50$ years of age, with a history of HZ , received 2 doses of SHINGRIX 2 months apart. The vaccine response rate (anti-gE antibodies) at 1 month postvaccination was $90.2 \%$ (95\% CI: 81.7; 95.7).

## Immunogenicity in Subjects Receiving 2 Doses of SHINGRIX 6 Months Apart

In a phase III, open-label clinical study (Zoster-026) where 238 subjects $\geq 50$ years of age were equally randomised to receive 2 doses of SHINGRIX 2 or 6 months apart, the vaccine response rate (anti-gE antibodies) at 1 month postvaccination following the 0,6 -month schedule was $96.5 \%$ ( $95 \% \mathrm{CI}$ : 90.4; 99.2).
The humoral immune response (anti-gE antibodies concentration) following the 0,6 -month schedule was not inferior to the humoral immune response following the 0, 2-month schedule, as the $97.5 \% \mathrm{Cl}$ upper limit of the antibodies concentration ratio was below 1.50 [1.16 (97.5\% CI: 0.98; 1.39)].

Immunogenicity in Individuals Previously Vaccinated with Live Attenuated Herpes Zoster (HZ) Vaccine

In a phase III, open-label, multicentre clinical study (Zoster-048), 430 adults $\geq 65$ years of age with or without a previous history of vaccination with live attenuated HZ vaccine $\geq 5$ years earlier were group-matched at a 1:1 ratio to receive 2 doses of SHINGRIX 2 months apart. The immune response to SHINGRIX was unaffected by prior vaccination with live attenuated HZ vaccine.

## Pharmacokinetics

Evaluation of pharmacokinetic properties is not required for vaccines.

## Clinical Studies

See Pharmacodynamic Effects.

## Non-Clinical Information

## Reproductive Toxicology

Administration of $\mathrm{VZV} \mathrm{gE} \mathrm{ASO} 1_{\mathrm{B}}$ to female rats did not indicate any harmful effects with respect to fertility, pregnancy, embryo-foetal development, parturition or postnatal development.
Treatment of male rats did not affect mating performance, fertility or early embryonic development.

## Animal Toxicology and/or Pharmacology

Non-clinical data reveal no special hazard for humans based on conventional studies of acute and repeated dose toxicity, local tolerance and cardiovascular/respiratory safety pharmacology.

## Pharmaceutical Information

## List of Excipients

Powder (gE Antigen):
Sucrose, polysorbate 80, sodium dihydrogen phosphate dihydrate, dipotassium phosphate
Suspension (ASO1 ${ }_{B}$ Adjuvant System):
Dioleoyl phosphatidylcholine, cholesterol, sodium chloride, disodium phosphate anhydrous, potassium dihydrogen phosphate, water for injections

## Shelf Life

The expiry date is indicated on the packaging.
For shelf-life after reconstitution of the medicinal product, see Use and Handling

## Storage

Store in a refrigerator $\left(2^{\circ} \mathrm{C}-8^{\circ} \mathrm{C}\right)$. Do not freeze. Store in the original package in order to protect from light. The storage conditions are detailed on the packaging.
For storage conditions after reconstitution of the medicinal product, see Use and Handling.

## Nature and Contents of Container

- Powder for 1 dose in a vial (type I glass) with a stopper (butyl rubber)
- $\quad$ Suspension for 1 dose in a vial (type I glass) with a stopper (butyl rubber).

SHINGRIX is available in a pack size of 1 vial of powder plus 1 vial of suspension or in a pack size of 10 vials of powder plus 10 vials of suspension.

## Not all pack sizes may be marketed.

## Incompatibilities

This medicinal product must not be mixed with other medicinal products

## Use and Handling

The powder and suspension should be inspected visually for any foreign particulate matter and/or variation of appearance. If either is observed, do not reconstitute the vaccine.

## How to Prepare SHINGRIX

SHINGRIX must be reconstituted prior to administration.

1. Withdraw the entire contents of the vial containing the suspension into the syringe.
2. Add the entire contents of the syringe into the vial containing the powder.

Shake gently until the powder is completely dissolved.
The reconstituted vaccine is an opalescent, colourless to pale brownish liquid.
The reconstituted vaccine should be inspected visually for any foreign particulate matter and/or variation of appearance. If either is observed, do not administer the vaccine.
After reconstitution, the vaccine should be used promptly; if this is not possible, the vaccine should be stored in a refrigerator $\left(2^{\circ} \mathrm{C}-8^{\circ} \mathrm{C}\right)$. If not used within 6 hours it should be discarded.

## Before Administration

1. Withdraw the entire contents of the vial containing the reconstituted vaccine into the syringe
2. Change the needle so that you are using a new needle to administer the vaccine.

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

## Version

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