A novel nonlive, adjuvanted herpes zoster subunit vaccine: a report on the emerging clinical data and safety profile

Abstract: Herpes zoster (HZ) is an acute vesicular dermatitis with a typical dermatomal distribution, caused by the varicella zoster virus (VZV), often preceded and accompanied by prodromal pain or pruritus. HZ may be related to several complications such as postherpetic neuralgia (PHN). The incidence and severity of the disease increase with aging, due to immunosenescence and in particular to the decline of the specific cell-mediated immunity (CMI). The impact of HZ in terms of morbidity and short- and long-term complications, the availability of suboptimal treatment options to date, and the high costs for the diagnostic and clinical-therapeutic management of patients have motivated the search for a new preventive approach through the development of a vaccine. The vaccine currently in use with live-attenuated virus (ZVL) has been shown to be effective in reducing the incidence of HZ, its impact, and the onset of PHN, although the efficacy is lower in older subjects and tends to decrease some years after immunization. A new adjuvanted recombinant subunit vaccine (HZ/su), containing the VZV glycoprotein E (gE) and the AS01B adjuvant system, is now a very promising alternative to ZVL; in several clinical studies, it showed a good safety profile and was able to elicit high immune humoral and cell-mediated responses, both maintained up to 9 years. Furthermore, HZ/su vaccine was effective both in preventing HZ and in reducing the onset of PHN and other complications. HZ/su has been recommended and preferred over ZVL by the Advisory Committee on Immunization Practices (ACIP) for the prevention of HZ and its complications in immunocompetent adults aged ≥50 years, even if already vaccinated with ZVL, through a two-dose schedule. HZ/su has been approved in Canada, USA, Europe, and Japan and is currently being approved in Australia. The aim of this review was to describe the epidemiological data, HZ and PHN risks and their impact on the social life and common life of infected people, and ZVL and HZ/su vaccine development including various clinical trials and efficacy, safety, and tolerability profiles.

Keywords: herpes zoster, adjuvanted recombinant herpes zoster subunit vaccine, live-attenuated herpes zoster vaccine

Background
Varicella zoster virus (VZV) is an α-herpes virus, with a double-stranded DNA genome, that exclusively infects humans. The primary infection causes chickenpox (varicella), during which the virus becomes latent in the ganglia of the nervous system. The mechanism by which cells become latently infected is unclear; it has been hypothesized for a long time, observing the similar interest of the cutaneous areas involved in varicella and herpes zoster (HZ), that the virus enters from the epidermal nerve endings and undergoes a retrograde axonal migration; however, it seems more plausible that the cause of VZV latentization is the varicella-associated viremia that would explain the
involved in the development of a vaccine.13

The treatment of HZ involves the use of antiviral drugs to be started possibly within the first 3 days after the onset of the disease. This therapy improves pain due to HZ but has usually no effect on reducing the risk of PHN. Treatment of this complication is difficult and is based on symptomatic drugs (lidocaine and capsaicin for local use; gabapentin and pregabalin, and tricyclic antidepressants). Combination therapy or opioid use can be performed; however, it seems that PHN-related pain is alleviated only in half of the cases.1

As a whole, the impact of HZ in terms of morbidity and short- and long-term complications, the availability of suboptimal treatment options, and the high costs for the diagnostic and clinical therapeutic management of patients have motivated the search for a new preventive approach through the development of a vaccine.13

HZ also has a strong impact in terms of health costs. Direct costs due to treatment, hospitalizations, medical and diagnostic examinations, and other health resources have the greatest economic weight. The advanced age, the presence of comorbidity or immunosuppression, the duration of hospitalization, the use of drugs, and the severity of symptoms involve high direct costs. The indirect costs (absent from work and loss of productivity) concern subjects of working age and are therefore associated with the age of onset of the disease.12

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Live-attenuated HZ vaccine (ZVL)

Due to immunosenescence, there is a progressive decline in the VZV-specific CMI, associated with a higher incidence and severity of HZ and PHN in the elderly; otherwise, the levels of VZV-specific antibodies remain relatively constant with advancing age. An episode of HZ causes a reactivation (booster effect) of the long-lasting CMI; therefore, the risk of recurrence of HZ is quite low in immunocompetent subjects. In light of this evidence, it has been hypothesized that, by stimulating the specific cell-mediated response, the frequency and severity of HZ and its complications could be reduced. This target has been pursued developing a live-attenuated vaccine with high antigen content.

The live-attenuated HZ vaccine, approved and currently in use (ZVL; Zostavax; Merck Sharpe & Dohme Corp, USA), consists of the same strain used in the pediatric varicella vaccine (Oka/Merck strain), with a high antigenic content; the latter is at least 14 times higher (19,400 plaque-forming units [PFU]) than the pediatric VZV live-attenuated vaccine.

ZVL must be stored and transported refrigerated (2–8°C) and administered subcutaneously or intramuscularly in a single dose (0.65 mL).

Efficacy, safety, and effectiveness of ZVL have been extensively studied.

The Shingles Prevention Study (SPS), a double-blind multicenter randomized controlled clinical trial, enrolled >38,000 subjects aged ≥60 years, with a history of chickenpox or of living for >30 years in an endemic area. The subjects enrolled received a dose of vaccine or placebo and were followed up ~3 years to evaluate the efficacy against the burden of illness (BOI, which includes incidence, severity, and duration of acute and chronic HZ-related pain during a 6-month follow-up), PHN, and the incidence of HZ.

ZVL showed an efficacy of 61.1, 66.5, and 51.3% in reducing the BOI and the incidence of PHN and of HZ, respectively, with the excellent safety and tolerability profiles.

A lower level of efficacy was found in preventing the occurrence of HZ in older subjects (63.9% in the 60–69 years age group vs 37.6% in the ≥70 years age group), while no differences were found related to the gender. The average duration of pain, discomfort, and severity of the disease were lower in the vaccinated subjects than in the placebo group. The effect of the ZVL in reducing the severity of the disease was greater in older subjects, although the reduction in incidence was, on the contrary, greater in younger age groups.

Another double-blind, randomized, controlled trial that integrated the SPS results was the Zoster Efficacy and Safety Trial (ZEST), in which the same criteria of inclusion and exclusion of the SPS were used enrolling a population of ~22,500 healthy subjects aged between 50 and 59 years. The enrolled subjects received a dose of vaccine or placebo and were followed up for at least 1 year; the efficacy against HZ was 69.8%. Therefore, the results of both studies showed a similar level of efficacy in the 50–59- and 60–69-year-old subjects; in these age groups, the efficacy was higher than in ≥70-year-old subjects (37.6%), probably due to a better immune response in younger subjects (Figure 1). The vaccine was also effective in improving functional status and reducing the interference of HZ on daily activities and quality of life.

The duration of efficacy was further investigated in the short-term persistence substudy (STPS) and the long-term persistence study (LTPS) evaluating a subset of subjects enrolled in the SPS. Recently, the summary of the characteristics of the product (SCP) has been updated with some data on the duration of protection. In detail, the vaccine efficacy against HZ 3 years after vaccination was 60, 55, 50, and 48% in 50–59-, 60–69-, 70–79-, and >80-year-old subjects, respectively. The efficacy against HZ 5 years after vaccination was 49, 46, and 44% in 60–69-, 70–79-, and >80-year-old subjects, respectively.

The efficacy against PHN 3 years after immunization was 98, 74, 73, and 63% in 50–59-, 60–69-, 70–79-, and >80-year-old subjects, respectively. The efficacy for this point 5 years after immunization was 72, 69, and 61% in the 60–69, 70–79, and >80 years age groups, respectively.

During the last years, it has been verified that the results of efficacy as well as those of effectiveness studies are consistent. In particular, the effectiveness data are consistent with each other, although they come from studies carried out with different designs and different characteristics of the enrolled populations (Figure 1).

All studies showed the good safety and tolerability profiles of ZVL.

ZVL is indicated in subjects aged ≥50 years; it can be administered to patients with a history of HZ and also to subjects whose clinical history is unknown. Currently, there are no indications for the administration of ZVL to those who have already received the varicella vaccine. ZVL is contraindicated in subjects with a history of hypersensitivity to any component of the vaccine, suffering from primitive or acquired immunodeficiency, on immunosuppressive therapy (including high doses of corticosteroids), with untreated active tuberculosis and pregnancy. ZVL should be administered 14–30 days before the start of any immunosuppressive treatment.
ZVL can be co-administered with the flu vaccine, provided it is injected into different anatomical sites, but it would not be co-administered with the 23-valent pneumococcal vaccine (PPSV23) because, as shown in a randomized clinical study, it causes a reduced humoral immune response to ZVL compared with administration of the vaccines 4 weeks apart.\textsuperscript{13,33}

However, as protection against HZ is mainly maintained by VZV-specific T CMI, a cohort study did not find any difference in the protection against HZ when ZVL was co-administered with PPSV23 in respect to a 4 weeks separate administration.\textsuperscript{34}

These observations were recently confirmed by a retrospective cohort study, which showed that there is no lack of protection against HZ following the co-administration of the two vaccines compared to their separate administration.\textsuperscript{35}

Several countries in the world have already included or foresee to include HZ vaccination in their immunization calendar. Although the strategies of use of the ZVL vaccine worldwide are heterogeneous, there is a clear convergence on the definition of the vaccination targets based on age group cohorts with greater burden of disease and the presence of risk factors starting from the age of 50 years. Where immunization has been adopted and implemented, the results are very interesting. For example, the UK data for the first 3 years after the introduction of ZVL showed a decrease of ~17,000 and 3,300 cases of HZ and PHN, respectively, in a population of 5.5 million eligible subjects for the vaccination program. The first assessment of vaccine efficacy in UK demonstrated a good impact on both HZ and PHN.\textsuperscript{36,37}

**Adjuvanted recombinant HZ subunit vaccine**

An alternative option to the ZVL vaccine is the adjuvanted recombinant HZ subunit vaccine (HZ/su) containing the glycoprotein E (gE) of the VZV and the adjuvant system AS01B.
The gE has been selected as a vaccinal antigen because it is the most expressed viral protein, is essential for the intercellular viral replication and diffusion, and is one of the targets of the VZV-specific immune response. \textsuperscript{38} gE is able to stimulate the humoral immune response, through the production of neutralizing antibodies, and also the cell-mediated response, through the activation of CD4\textsuperscript{+} T cells. \textsuperscript{39} The AS01B is a liposomal adjuvant that contains 50 µg of 3-O-desacil-4’-monophosphoryl lipid A and 50 µg of the saponin 
\textit{Quillaja saponaria}, fraction 21 (QS21), which stimulates the activation of the innate immune response, the production of cytokines, and the recruitment of antigen-presenting cells. \textsuperscript{39,40} The monophosphoryl lipid A, in synergy with QS21, stimulates the production of γ-interferon and, therefore, enhances the immune response, acting as a toll-like receptor four agonist. \textsuperscript{41}

As demonstrated in preclinical studies, \textsuperscript{42} the AS01B adjuvant also enhances the specific antibody and cell-mediated response to the recombinant protein. \textsuperscript{38} In animal models, gE stimulated a higher activation of CD4\textsuperscript{+} T lymphocytes when associated with the adjuvant AS01B compared to when administered in the absence or with another adjuvant. \textsuperscript{42}

In 2012, a Phase I/II study was conducted in Belgium by Leroux-Roels et al with the aim of investigating the safety and immunogenicity profiles of the subunit vaccine compared to the live-attenuated vaccine (VZV pediatric vaccine). The study showed that HZ/su induces a greater immune response, both cell mediated and humoral, which remains higher than the one obtained using VZV pediatric vaccine up to 42 months both in young adults (18–30 years) and in older subjects (50–70 years).

This trial also showed an acceptable safety profile of the HZ/su vaccine, despite a greater frequency of nonsevere adverse reactions (such as asthenia, myalgia, headache, and pain at the injection site) compared to the VZV pediatric vaccine. \textsuperscript{43}

Even the Phase I study by Lal et al \textsuperscript{44} with the administration of two doses of HZ vaccine (2-month interval) on healthy volunteers in Japan showed an acceptable safety profile.

In 2013, Chlibek et al conducted a multicentric and international Phase II trial with the aim of comparing the efficacy and safety of different adjuvanted subunit vaccines compared to the same vaccine without adjuvant. The study involved ≥50-year-old adults randomly divided into four groups who were given the gE adjuvanted subunit vaccine containing either the liposomal complex AS01B (HZ/su) or the AS01E (gE/AS01E; AS01E contains 500 µg of dioleoyl phosphatidylcholine, 125 µg of cholesterol, 25 µg of Monophosphoryl Lipid A, and 25 µg of QS21), the nonadjuvanted subunit vaccine (gE), and placebo, respectively. The results showed that the immune response is significantly higher with the two adjuvanted vaccines in comparison to the nonadjuvanted vaccine and that the HZ/su vaccine provides a better response than the adjuvanted gE/AS01E vaccine.

In contrast, as far as tolerability is concerned and adjuvanted vaccines have shown a greater occurrence of adverse reactions, all of which transient and of mild-to-moderate degree, such as pain at the injection site among local reactions and asthenia among the general ones. \textsuperscript{45}

In 2014, Chlibek et al conducted another Phase II study based on the experience of the previous year. In this study, the investigators wanted to look for the safety and immunogenicity profiles between different formulations and schedules of the gE subunit vaccine adjuvanted with AS01B. The study design enrolled 714 subjects aged ≥60 years randomly divided into five groups. Different formulations/schedules were used, as follows: one single-dose of gE/AS01B vaccine with a 100 µg of gE, two doses of gE/AS01B (2 months apart) with 25, 50, or 100 µg of gE, and two doses of nonadjuvanted gE vaccine (2 months apart) with 100 µg of gE. \textsuperscript{46}

The results, analyzed at 1, 2, 3, 12, 24, and 36 months after the first administration, showed that the two-dose schedule of adjuvanted vaccine leads to a gE-specific clonal expansion of CD4\textsuperscript{+} T lymphocytes. This expansion was registered with all formulations analyzed and was three times higher than the one obtained with the monoadministration of adjuvanted vaccine or the two-dose administration of the nonadjuvanted vaccine. Furthermore, immune response remained high over time. The different formulations showed that the humoral response induced with the administration of two doses of 50 or 100 µg of gE/AS01B is similar and superior to that obtained in the other groups; the nonadjuvanted gE vaccine was found to be able to elicit the humoral response but not the cell-mediated one. Besides, this study confirms the greater reactogenicity of the adjuvanted vaccine compared to the nonadjuvanted vaccine; however, reactogenicity does not change significantly taking into account the different formulations.

In 2016, Chlibek et al \textsuperscript{47} published a further Phase II trial with the aim of assessing the vaccine-induced immune response in the cohort of immunized subjects enrolled in the previous study performed in 2014. In this study, the follow-up involved only those belonging to the group that received the HZ/su vaccine (gE/AS01B adjuvanted vaccine with the 50 µg formulation) and evaluated the duration of cell-mediated and humoral immune responses at 48, 60, and 72 months after the first dose. The results showed that after 72 months both
cell-mediated and humoral responses tend to decrease by 20–25% compared to the 36th month, remaining however satisfactory and significantly higher than the prevaccination level. In detail, the HZ/su vaccine provides a 3.8 times higher cell-mediated response and a 7.3 times higher humoral response than the prevaccination status.

In 2018, Schwarz et al made a further extension of the previous study evaluating humoral and cell-mediated immune responses 9 years after the first dose. The results of the follow-up have confirmed that, even in this case, the immunity persists significantly above the prevaccination level regardless of the age of the analyzed subjects. In particular, the study estimated that this immunity will remain high up to at least 15 years after vaccination.48

A study that provided relevant information in defining the optimal schedule in terms of time interval between the two doses of HZ/su vaccine was conducted by Lal et al in 2018. In this study, the average geometric titers (GMTs) were analyzed 1 month after the second dose of vaccine given accordingly to a 0–2, 0–6, and 0–12 months' schedule. The study demonstrated an excellent immune response in the 0–6 months’ schedule group, comparable to that obtained in the 0–2 months’ schedule group. Besides, GMTs were lower using the 0–12 months’ schedule; this suggests that the second dose of HZ/su should be optimally (ie, without loss of efficacy) administered in a time window between 2 and 6 months after the first dose.49

Another point studied in the approval trials of the HZ/su vaccine has been the evaluation of the efficacy of the vaccination on anamnestically positive subjects for HZ and on subjects already vaccinated with ZVL.

In 2017, Vink et al published a Phase III study conducted on 60 Japanese subjects aged ≥50 years evaluating the safety and immunogenicity of HZ/su vaccine when administered subcutaneously vs intramuscularly. The results of the study showed a comparable immune response between subcutaneous administration and intramuscular administration; however, a greater reactogenicity was registered using the subcutaneous route of administration.52

The development of a nonlive-inactivated vaccine for the prevention of HZ represents an important tool to be available in subjects with impaired immune status. As a matter of fact, these patients, notwithstanding their greater risk of contracting the disease and developing severe complications, until today could not undergo vaccination due to the risk possibly related to the administration of the live-attenuated vaccine (ZVL).53 In this regard, studies conducted by Berkowitz et al54 and by Stadtmauer et al55 on HIV-infected subjects and hematopoietic cell transplant recipients, respectively, are very relevant.

The first study, conducted on ≥18-year-old HIV+ subjects and stratified according to the CD4+ T-cell count in three groups (≥200, 50–199, and ≥500 cells/μl), evaluated the efficacy and safety profiles of the HZ/su compared to placebo group. The results showed that both VZV-specific humoral and cell-mediated immunities reached high values, with a peak after the second dose, persisting for at least 18 months.

In addition, from the clinical point of view, the HZ/su was found to be safe and well tolerated in these patients without any effect on the HIV viral load.54

The second one was a case–control study conducted on 121 patients with multiple myeloma, non-Hodgkin lymphoma (B cell or T cell), Hodgkin lymphoma, or acute myeloid leukemia who had undergone autologous hematopoietic stem-cell transplant 50–70 days earlier. These patients were randomly divided into four groups who, according to the 0–1–3 months’ schedule, received three doses of HZ/su, three doses of gE/AS01E, one dose of placebo, and two doses of HZ/su or three doses of placebo, respectively. A high immune response, both humoral and cell mediated, was achieved with the three-dose schedule of the adjuvanted vaccine independently of the adjuvant used. In addition, the safety profile was judged adequate, since most reactions were classified as mild–moderate.55

A recent clinical trial, not yet published, was conducted in autologous hematopoietic cell transplant recipients aged 18 years and older to evaluate the efficacy of HZ/su in the prevention of HZ.
The results show high efficacy in the prevention of HZ and PHN with a very significant reduction in the duration of “worst” HZ-associated pain calculated in days vs placebo.66

In 2017, Schwarz et al investigated the co-administration of HZ/su and quadrivalent seasonal inactivated influenza vaccine, in order to evaluate the immunogenicity and safety vs the single administration of the vaccines in different times.

The results of the study show no interference in the immune responses to either vaccine when the vaccines were co-administered, and no safety concerns were identified.57

In addition to numerous and relevant studies aimed at investigating the immunogenicity, safety, and tolerability of the HZ/su vaccine, two approval pivotal trials published by Lal et al (ZOE-50)38 and by Cunningham et al (ZOE-70)40 in 2015 and 2016, respectively, have been undertaken in order to evaluate its impact on the reduction in the disease and its complications.

The ZOE-50 is a multicenter randomized study conducted in 18 countries enrolling 15,411 subjects aged ≥50 years divided in two groups receiving two doses of HZ/su or two doses of placebo 2 months apart, respectively. During an average follow-up time of 3.2 years, 210 and six cases of HZ were registered in the placebo group and in the HZ/su group, respectively. Thus, the efficacy of HZ/su against HZ was 97.2%. In addition, the efficacy in subjects aged ≥70 years was found to be comparable to that registered in younger subjects.

As in previous studies, however, the HZ/su group reported more frequently, than the placebo group, the occurrence of solicited local and systemic reactions in the first 7 days, most of them of mild-to-moderate intensity and transient.38

The ZOE-70 study continues the experience of the ZOE-50 study with the enrollment of 13,900 older (aged ≥70 years) subjects randomized to receive two doses of HZ/su or placebo; in both groups, a 2-month interval between doses was used. As in the study conducted on younger patients, also in this case, there was a clear difference in the occurrence of HZ cases (23 and 223 in the vaccinated group and in the placebo group, respectively) with an estimated efficacy of 89.8% in the reduction in the appearance of the disease at a mean follow-up period of 3.7 years.40

In addition to the abovementioned high efficacy of the HZ/su, it is important to highlight the impressive reduction in the risk of developing complications HZ correlated, such as PHN, in vaccinated subjects compared with the placebo group (Figure 2).58

A predefined pooled analysis considering all together (16,596) subjects aged ≥70 years enrolled in both ZOE-50 and ZOE-70 studies estimated the efficacy of the vaccine against HZ (91.3%) and PHN (88.8%) in the older population. In the same analysis, no relevant waning of HZ/su efficacy against HZ was found from year 1 (97.6%) to year 4 (87.9%). Regarding the safety profile of HZ/su vaccine, compared to placebo, this study also confirmed a greater

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**Figure 2** Incidence rate of HZ and PHN per 1,000 Person-Years in HZ/su group and in placebo group.


**Abbreviations:** HZ, herpes zoster; HZ/su, HZ subunit vaccine; PHN, postherpetic neuralgia.
frequency of occurrence of mild or moderate local or systemic adverse reactions in recipients (79 vs 30% in the placebo group). Regarding severe reactions, deaths, and potential immune-mediated diseases, no statistically significant differences were observed between the two groups in both ZOE studies and the compliance of the subjects enrolled in carrying out the whole two-dose vaccination cycle with HZ/su was very high.\(^{10,59}\)

As reported in the technical data sheet of HZ/su, the effect on the reduction of disease commitment (BOI) was 98.4% in subjects aged ≥50 years (ZOE-50) and 92.1% in subjects aged ≥70 years (ZOE-50 and ZOE-70 together). The BOI score includes the incidence of HZ with the severity and duration of acute pain and chronic pain associated with HZ over a period of 6 months following the onset of rash.\(^{60}\)

In 2018, Kovac et al analyzed data from the ZOE-50 and ZOE-70 studies in order to evaluate further information from the follow-up of enrolled patients such as mortality, number of hospitalizations, and HZ complications (excluding PHN already analyzed by the previous study) defined as HZ-related vasculitis, stroke, and disease with neurological and ophthalmic involvements. The results showed that the efficacy of the vaccine in reducing the impact of the aforementioned HZ complications has been 93.7 and 91.6% in ≥50- and ≥70-year-old subjects, respectively. Furthermore, during the follow-up, five hospitalizations for HZ were registered, all belonging to the group receiving placebo; no HZ-related deaths were recorded in both groups.\(^{61}\)

**Pharmaco-economic evaluations**

An absolutely relevant point coming out from the recent literature on the use of the HZ/su vaccine is its potential impact on public health and cost-effectiveness.

In this regard, the study of Curran et al, conducted on the basis of the results of efficacy studies in the prevention of HZ and PHN, estimated the impact of vaccination with HZ/su on the German population aged ≥50 years, comparing it with the use of ZVL. The results were analyzed dividing the population into three age groups (50–59, 60–69, and ≥70 years) and assuming a 40% coverage for both vaccines and a 70% coverage for the second dose of HZ/su. According to the authors, the HZ/su vaccine would allow, in the three age groups and from the time of vaccination to the end of life, to avoid 725,233, 533,162, and 486,794 cases vs 198,477, 196,000, and 104,640 cases in respect to the ZVL vaccine. The number needed to vaccinate (NNV) to prevent a case of HZ would be between 8 and 11 for HZ/su vs 20–50 for ZVL, and the NNV to prevent a case of PHN would range between 39 and 53 vs 94 and 198 for HZ/su and ZVL, respectively.\(^{62}\)

Also in a recent experience, published by Watanabe et al,\(^{63}\) the potential impact on the Japanese population of HZ/su vaccination compared to ZVL was estimated.

The results of vaccination with HZ/su could prevent ~3.3 million HZ cases, 692,000 cases of PHN, and 281,000 cases of other complications, compared with the prevention of 0.8 million HZ cases, 216,000 PHN cases, and 57,000 other complications with vaccination with ZVL. The NNV in order to prevent one HZ case would range between 6 and 14 using HZ/su and between 21 and 138 using ZVL.\(^{64}\)

Also, from the pharmaco-economic point of view, the HZ/su vaccine guarantees a significant impact in terms of optimization of public health spending, with a high probability of being cost-effective, as demonstrated by the study by Le et al.\(^{64}\) The authors estimated that at a cost of $280 ($140 per dose), HZ/su would be more effective and less expensive than ZVL in all age groups considered (≥60, ≥70, and ≥80 years) while the Incremental Cost-Effectiveness Ratio (ICER) of vaccination with HZ/su compared to nonvaccination would range between 20,038 and $30,084 per QALY, according to the age considered.

Always considering the price of $280 per vaccination cycle, HZ/su would have 73–91% probability, depending on the age groups considered, of being cost-effective at $50,000 per QALY and 78–93% probability at $100,000 per QALY. In comparison, at the same threshold of $100,000 per QALY, ZVL vaccine given to 60-year-old subjects would be cost-effective with the probability of <5%.\(^{64}\)

**HZ/su: approvals and recommendations**

Based on the excellent and promising results of the clinical trials, the HZ/su vaccine, named now Zoster Vaccine Recombinant, Adjuvanted (RZV, Shingrix; GlaxoSmithKline plc, London, UK), has been approved for the first time worldwide in Canada on October 13, 2017, and a few days later (on October 20, 2017) in the USA by the Food and Drug Administration (FDA).

With the approval of the vaccine, the Advisory Committee on Immunization Practices (ACIP) of the Centers for Disease Control (CDC) has made public its indications on the basis of a review and analysis of literature and epidemiological data started in March 2015.

The recommendations of the ACIP can be summarized in three points:

1. RZV is recommended for the prevention of HZ and its complications in immunocompetent adults aged ≥50 years.
2. RZV is recommended for the prevention of HZ and its complications in immunocompetent adults aged ≥50 years previously vaccinated with ZVL.

3. RZV is preferable to ZVL for the prevention of HZ and related complications.

The optimal route of administration is intramuscular, and it is possible to co-administer RZV with other vaccines in another anatomical site in order to improve patient compliance to vaccination; more specifically, RZV can be administered together with the quadrivalent influenza vaccine without any immunogenicity and safety concern.65

Besides, RZV can be co-administered with PPSV23 vaccine as showed in a recent study: immunogenicity of both vaccines is not negatively influenced and co-administration was well tolerated and safe.66

The ideal schedule must be with two doses with a 2–6 months’ interval; it is important to underline that there is no indication to repeat the vaccination cycle if the second dose is administered ≥6 months after the first one, while if the interval is <4 weeks, it would be necessary to consider the first dose null.

The vaccine should not be administered in subjects with active HZ, since it has only preventive and nontherapeutic value, and in pregnant women due to the lack of clinical studies. The only contraindication to vaccination is related to a documented history of a severe allergic reaction to one or more components of the vaccine.

ACIP strongly recommends to administer RZV in subjects with a history of previous HZ, to prevent a possible recurrence of disease, and in patients with comorbidities such as chronic renal failure, diabetes mellitus, rheumatoid arthritis, and COPD for which HZ can have a great and severe impact on their health condition. RZV is not indicated for the prevention of varicella and only for HZ.

Regarding immunocompromised subjects, ACIP currently recommends vaccination only in patients in remission from the immunosuppressive state, in anticipation of a future immunosuppression, or in subjects who take a low dose of immunosuppressive therapy. As in immunosuppressed patients, the HZ disease has a greater incidence and impact and the evaluation of the efficacy and safety profiles of the vaccine should be based on more data in addition to those previously reported;65 in this regard, some clinical trials are currently underway on immunocompromised subjects in renal transplant recipients57 and in chemotherapy for solid68 and hematological tumors.69

RZV has also been approved for marketing in Europe, thanks to the favorable opinion of the European Medicines Agency (EMA), and in Japan on March, 23, 2018, and at present, it is pending approval in Australia.

Given the very recent approval of RZV, at the moment, there are no Phase IV trials related to commercialization, which, when available, will provide further, large-scale and relevant information on effectiveness and safety of the new vaccine.

Conclusion

All evidence available in literature underlines the importance of the new adjuvanted recombinant vaccine (RZV) that seems very promising in the fight against HZ and its complications. RZV has indeed proved very effective in eliciting a strong immune response, regardless of the age of the immunized subject.

The new recombinant vaccine, moreover, could bring important benefits in immunocompromised subjects for which, despite a higher incidence of HZ, vaccination with live-attenuated virus is contraindicated.

It is very important to underline the strong impact that vaccination with RZV can offer in the context of public health in terms of containing the impact that HZ and its complications have on the social, economic, and welfare levels. The only critical points of this new vaccine could be related to its reactogenicity and to the need of a two-dose schedule. Therefore, a well-balanced communication will be necessary in order to guarantee a high compliance to this new preventive intervention.

Disclosure

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