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Background. We conducted a phase 1, randomized, double-blind, placebo-controlled trial of a replication-defective HSV-2 vaccine, HSV529 (deleted for UL5 and UL29), in 60 healthy adults aged 18 to 40 years.

Methods. Subjects were enrolled in groups of 20 from 3 serogroups: HSV1+ /HSV2+ (group 1), HSV1+/HSV2- (group 2), and HSV1-/HSV2- (group 3). At months 0, 1, and 6, 15 subjects in each group received HSV529 intramuscularly and 5 subjects received placebo. The primary endpoint was the frequency of solicited injection site and systemic reactions from day 0 to 7 after each vaccination and unsolicited adverse events up to 6 months after the last dose.

Results. 89% of vaccine recipients experienced a mild to moderate solicited injection site reaction vs. 47% of placebo recipients (P = 0.006, 95% CI 0.129, 0.676) that did not preclude additional doses. 64% of vaccine recipients experienced solicited systemic reactions vs. 53% of placebo recipients (P = 0.44, 95% CI -0.179, 0.402). Two serious adverse events occurred in 2 participants and were assessed as unrelated to HSV529 administration. Serum neutralizing antibody titers significantly increased from baseline after 3 doses of HSV529 compared with placebo in group 3 only (P < 0.001). This increase persisted up to 6 months after the third dose of vaccine (P < 0.001). Serum and vaginal antibodies to HSV2 glycoprotein D (gD) also significantly increased after 3 doses of vaccine in group 3 subjects (P < 0.001 and P = 0.012, respectively). The mean vaginal gD titer after 3 doses was about one-third of the mean serum gD titer. In addition, the vaccine induced significant levels of HSV2-specific antibody-dependent cellular cytotoxicity (ADCC) after 3 doses in group 3 subjects compared with placebo (P < 0.001).

Vaccine-induced CD4 T-cell responses were detected in 46%, 27%, and 36% of subjects in groups 1, 2, and 3, respectively, one month after the third dose of vaccine. CD8 T-cell responses were detected in 8%, 18%, and 14% of subjects in groups 1, 2, and 3, respectively, at the same time point.

Conclusion. The HSV529 vaccine was safe, well-tolerated, and immunogenic, eliciting significant neutralizing, gD, and ADCC-mediating antibodies, and modest cellular immune responses in HSV seronegative individuals. NCT01915212

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1346. Results of a Safety Pooled Analysis of an Adjuvanted Herpes Zoster Subunit Vaccine in More than 14,500 Participants Aged 50 Years or Older
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Background. The recombinant herpes zoster (HZ) subunit vaccine (HZ/su) has shown efficacy against HZ in adults ≥50 and ≥70 years of age (YOA), in two pivotal Phase III clinical trials (NCT01165177, NCT01165229). A pooled safety analysis of data from these two efficacy studies was performed, including a comparative analysis on HZ/su vs. placebo groups, to provide a comprehensive understanding of the HZ/su safety profile.

Methods. Two pivotal, randomized, placebo-controlled Phase III studies, assessed the efficacy, reactogenicity and safety of HZ/su, administered immunologically according to a 0, 2-month schedule. Solicited and unsolicited adverse events (AEs) were collected for 7 and 30 days after each dose, respectively; serious AEs (SAEs) for 1 year after last dose; fatal and related SAEs and potential immune-mediated diseases (pIMDs) during the entire study period. Reactogenicity was assessed in a subset of participants; safety was assessed in all vaccinated participants.

Results. 29,305 participants ≥50 YOA (HZ/su: 14,645; placebo: 14,660) were included in the pooled analysis. HZ/su was more reactogenic than placebo. Local reactions were mostly mild to moderate in intensity and transient (median duration = 3 days); the percentages of participants reporting SAEs, fatal SAEs and pIMDs were similar in both groups, at 30 days and 1 year after last dose (Figures 1 and 2). Percentages of fatal SAEs ranged between 4.3% (95% Confidence Interval [CI]: 4.0-4.7) and 4.6% (95% CI: 4.3-5.0) and pIMDs between 1.2% (95% CI: 1.1-1.4) and 1.4% (95% CI: 1.2-1.6), in HZ/su vs. placebo in participants ≥50 YOA.

Conclusion. No safety concern was identified. Together with the high efficacy against HZ (97.2% [95% CI: 93.7–99.0]), ¹¹91.3% [95% CI: 86.8–94.5]¹², the safety data supports a favorable benefit/risk profile of HZ/su in participants ≥50 YOA.

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1347. A Cost-Effectiveness Analysis of an Adjuvanted Subunit Vaccine for the Prevention of Herpes Zoster and Post-Herpetic Neuralgia
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SAEs, serious adverse events; pIMDs, potential immune-mediated diseases; Error bars depict 95% confidence intervals (CI)
Background. Herpes zoster (HZ) develops in up to 50% of unvaccinated individuals aged ≥ 80 years, accounting for more than 1 million cases of HZ annually in the United States. A live attenuated vaccine (LAV) for HZ is U.S. FDA approved for persons 50 years or older, though CDC Advisory Committee on Immunization Practices (ACIP) recommendations are only for persons beginning at age 60 years. LAV efficacy at preventing HZ is ~70% for persons 50–59 years of age, with lower efficacy in older adults, and it is efficacious in preventing post-herpetic neuralgia (PHN) beyond the HZ prevention. The efficacy of LAV after vaccination wanes over time. A new adjuvanted HZ subunit vaccine (SUV), administered as a two-dose series, has greater than 95% efficacy against HZ in persons 50–69 years of age. SUV efficacy remains greater than 90% in persons vaccinated at age 70 years and older, including the subgroup older than 80 years of age. Overall efficacy of SUV against PHN approaches 90%. The waning rate of efficacy after SUV vaccination is unknown.

Methods. To estimate the relative cost-effectiveness of SUV, LAV and no vaccination (NV) strategies, a Markov model was developed based on published trials and data on vaccine efficacy persistence, quality of life, resource utilization, costs and disease epidemiology. The perspective was U.S. societal, and the cycle length was one year with a time horizon of 20 years. SUV vaccine efficacy was estimated for the base case at the same rate as LAV, all persons were assumed to receive both doses of SUV, and the cost of SUV included both doses.

Results. For individuals vaccinated at age 50 years the incremental cost-effectiveness ratio (ICER) for SUV vs. LAV was $142.81 per quality-adjusted life-year (QALY); at age 60 years the ICER dropped to $59.48 per QALY. The cost-effectiveness ratio of SUV approached that of LAV when the cost of SUV was $500 for persons vaccinated at age 50 years and when the cost was $400 for those vaccinated at age 60 years. The SUV cost that would result in achieving an ICER target of $100,000 per QALY for SUV vaccination vs. NV at age 50 years was $316; at age 60 years the cost of SUV included both doses.

Conclusion. Vaccination at age 60 years with SUV was more cost-effective than LAV when SUV cost was $450 or less. Vaccination with SUV at age 50 years appeared to be cost-effective if SUV cost was ~$15 or less.

Disclosures. All authors: No reported disclosures.

1349. Immunogenicity and Safety of a Candidate Subunit Adjuvanted Herpes Zoster Vaccine in Adults with Solid Tumors Vaccinated Before or During Immunosuppressive Chemotherapy Treatment: A Phase II/III, Randomized Clinical Trial
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Background. No herpes zoster (HZ) vaccine for immunosuppressed individuals is currently available. GSK’s candidate HZ vaccine containing recombinant varicella zoster virus glycoprotein E (gE) subunit and AS01, Adjuvant System (HZ/su) showed >90% efficacy for HZ prevention in immunocompetent adults aged ≥50 years. The HZ incidence in individuals with solid tumors (ST) receiving immunosuppressive chemotherapy (chemo) is estimated as 3–4 times higher than the overall US population aged ≥50 years. We present HZ/su immunogenicity and safety in ST adults aged ≥18 years.

Methods. In this phase II/III, observer-blind, multicenter study (NCT01798056), ST adults received 2 doses of HZ/su or placebo intramuscularly 1–2 months apart and were randomized 4:4:1:1 to receive a first dose 8–30 days (D) pre-chemo (HZ/su – HZ/su-PreC group, placebo – Pl-PreC) or on chemo start (≤ 1 D) (HZ/su-OnC, Pl-OnC). Vaccine response rates (VRRs) and geometric means (GMs) were evaluated for gE humoral immune and gE-specific CD4+ cell-mediated immune (CMI) responses 1 month post dose 2 (M2). Solicited adverse events (AEs) were recorded for 7 days and unsolicited AEs and medically-attended AEs (MAEs) for 30 days after each dose. Solicited general and unsolicited AEs were also collected for 7 days prior to dose 1. Potential immune-mediated diseases (pIMDs) and serious AEs (SAEs) were recorded until 1 year post dose 2. From data dose 1 through M2 is presented.

Results. At M2, 240 subjects (121 HZ/su; 119 placebo) were included in the humoral immunogenicity according-to-protocol (ATP) cohort. All immunogenicity success criteria were met at M2 (Table 1). gE VRRs for ATP humoral immune and CMI sub-cohort (72 subjects: 36 HZ/su; 36 placebo) were higher in HZ/su groups. Humoral GM concentrations and CMI GM frequencies were significantly higher in HZ/su compared with placebo groups. The frequency of AEs was higher in HZ/su vs. placebo groups for solicited local AEs, but similar for solicited general AEs, unsolicited AEs, MAEs and SAEs. No pIMDs, vaccine-related SAEs or transplant rejections were reported (Table 2).

Conclusion. HZ/su was highly immunogenic in adults with RT at M2. No safety concerns were identified.

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