Should Sex Be Considered an Effect Modifier in the Evaluation of Influenza Vaccine Effectiveness?

**BRIEF REPORT**

**Open Forum Infectious Diseases**

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We investigated sex as a potential modifier of influenza vaccine effectiveness (VE) between 2010–2011 and 2016–2017 in Canada. Overall VE was 49% (95% confidence interval [CI], 43% to 55%) for females and 38% (95% CI, 28% to 46%) for males (absolute difference [AD], 11%; P = .03). Sex differences were greater for influenza A(H3N2) (AD, 17%; P = .07) and B(Victoria) (AD, 20%; P = .08) compared with A(H1N1) pdm09 (AD, 10%; P = .19) or B(Yamagata) (AD, –3%; P = .68). They were also more pronounced in older adults ≥50 years (AD, 19%; P = .03) compared with those <20 years (AD, 4%; P = .74) or 20–49 years (AD, –1%; P = .90) but with variation by subtype/lineage. More definitive investigations of VE by sex and age are warranted to elucidate these potential interactions.

**Keywords.** effect modification; gender; influenza vaccine; influenza virus; sex; vaccine effectiveness.

Sex is a potential confounder in the evaluation of influenza vaccine effectiveness (VE) due to its plausible association both with the likelihood of receiving influenza vaccine and with having an influenza exposure [1]. Females tend to have higher influenza vaccination coverage rates than males (at least among young adults in Canada and the United States [2, 3]), likely related to their greater propensity for seeking health care. Women also tend to have greater opportunities for exposure to influenza associated with their traditional gender roles as primary caregivers for children and the elderly or their greater propensity to work in health care or other occupations that may increase the likelihood of exposure [1]. Conversely, morbidity and mortality due to influenza are generally thought to be higher among males (the so-called “man flu” phenomenon [4]). This association may have a biological basis, with greater sex differences in influenza risk reported in the youngest and oldest age groups [5–9].

Multivariable VE analyses are typically adjusted for sex to account for this potential source of bias. Although there may be true biological differences in response to influenza vaccine between males and females, sex is rarely considered a potential effect modifier of influenza VE. Here we investigate the interaction between sex and influenza vaccination for VE against medically attended, laboratory-confirmed influenza illness across 7 seasons (2010–2011 to 2016–2017) in Canada. We further explore whether sex differences in VE vary by influenza subtype/lineage, age, or season.

**METHODS**

The current analysis utilized historical databases of the Canadian Sentinel Practitioner Surveillance Network (SPSN) from 2010–2011 to 2016–2017 according to methods previously described [10–16]. Briefly, respiratory specimens were collected from outpatient 1 or more years old presenting to sentinel practitioners within 7 days of onset of influenza-like illness using a standardized case definition. Specimens were tested for influenza viruses by real-time reverse transcription polymerase chain reaction at public health reference laboratories in each participating province (Alberta, British Columbia, Ontario, and Quebec). Patients testing positive for influenza were considered cases; those testing negative for influenza were considered controls.

Patient data, including sex and vaccination status, were recorded on the laboratory requisition form by the ordering physician at the time of specimen collection before influenza diagnosis. Vaccination status was based on patient self-report but may have also been documented in the physician’s records. Patients were considered vaccinated if they received seasonal influenza vaccine ≥2 weeks before symptom onset; patients who received influenza vaccine <2 weeks before symptom onset were excluded. All patients or their parent/guardian provided verbal consent. Institutional review boards in each province provided approval for this study.

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Differences between male and female patients were compared by chi-square test for categorical variables or the nonparametric Wilcoxon rank-sum test for continuous variables. Odds ratios (ORs) for influenza test positivity (by influenza A subtype or influenza B lineage) comparing vaccinated with unvaccinated patients were derived using logistic regression according to a test-negative study design for all seasons combined. Seasons for which there was minimal circulation of a given subtype/lineage were excluded from the pooled analysis for that outcome. Covariates included in the interaction models were vaccination status, sex, age group, comorbidity, province, specimen collection interval (days from symptom onset to specimen collection), calendar time (based on week of specimen collection, modeled as a cubic B-spline function with 3 equal knots), season, and an interaction term for vaccine*sex. VE was derived as \((1 - \text{OR}) \times 100\%\), where \(\text{OR} = \exp[\beta_{\text{vac}} + \beta_{\text{vac}} \times \text{sex}]\) for males (sex = 1). To verify the results and ensure that our interaction models were adequately specified, we also conducted separate analyses in male and female strata; sex-stratified VE estimates generally differed by ≤5% from the interaction models (data not shown). All analyses were performed in SAS, version 9.4 (SAS Inc., Cary, NC).

**RESULTS**

Females were over-represented among SPSN participants (60% vs 40%), both among influenza cases (58% vs 42%) and test-negative controls (61% vs 39%; \(P < .01\)). The age distribution varied by sex, with greater over-representation of females among adults age 20–49 years (62% vs 38%) and older adults age ≥50 years (63% vs 37%) than children and adolescents younger than age 20 years (51% vs 49%; \(P < .01\)). Overall, females were slightly less likely to test positive for influenza compared with males (40% vs 43%; \(P < .01\)), mostly driven by detection of A(H3N2) (17% vs 19%; \(P < .01\)), and less so by A(H1N1)pdm09 (10% vs 10%; \(P = .71\)), B(Yamagata) (6% vs 7%; \(P = .18\)), or B(Victoria) (4% vs 5%; \(P = .04\)). Females also had higher vaccination coverage than males overall (29% vs 23%; \(P < .01\)) and among negative controls (34% vs 27%; \(P < .01\)) but not among influenza cases (21% vs 19%; \(P = .10\)) (Supplementary Tables 1 and 2).

In general, adjusted VE was higher among females than males, although this varied by influenza subtype/lineage, age group, season, and sex. Overall, for any influenza, adjusted VE was 49% (95% confidence interval [CI], 43% to 55%) for females vs 38% (95% CI, 28% to 46%) for males (absolute difference [AD], 11%; \(P\) value for interaction term = .03) (Figure 1). The greatest absolute differences between females and males were for A(H3N2) and B(Victoria). For A(H3N2), adjusted VE in all seasons combined was 34% (95% CI, 22% to 44%) for females vs 17% (95% CI, 1% to 32%) for males (AD, 17%; \(P = .07\)). Excluding the 2014–2015 season, for which overall VE against A(H3N2) was negligible [14], adjusted VE was 49% (95% CI, 37% to 58%) and 29% (95% CI, 9% to 44%) for females and males, respectively (AD, 20%; \(P = .03\)). For B(Victoria), adjusted VE was 65% (95% CI, 50% to 76%) and 45% (95% CI, 18% to 63%) for females and males, respectively (AD, 20%; \(P = .08\)).

The same pattern was observed for A(H1N1)pdm09 but with

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**Figure 1.** Vaccine effectiveness estimates by sex for influenza A subtype and influenza B lineage. \(P\) values indicate significance of vaccine*sex interaction. Covariates included in the interaction model were vaccination status (no, yes), sex (female, male), age group (1–8, 9–19, 20–49, 50–64, ≥65 years), comorbidity (no, yes), province (AB, BC, ON, QC), collection interval (≤4, 5–7 days), calendar time (week of specimen collection based on cubic B-spline with 3 equal knots), season, and vaccine*sex. \(\text{A(H1N1)pdm09 analysis excludes 2014–2015 and 2016–2017 due to low A(H3N2) circulation those seasons. \(\text{A(H3N2) analysis excludes 2013–2014 due to low A(H3N2) circulation that season.}\)
a smaller absolute difference in VE among females (63%; 95% CI, 53% to 70%) vs males (53%; 95% CI, 36% to 65%; AD, 10%; P = .19). No sex differences were seen for B(Yamagata), with a VE of 55% (95% CI, 41% to 65%) for females vs 58% (95% CI, 42% to 70%) for males (AD, −3%; P = .68). When restricted to vaccinated participants, the adjusted odds of influenza diagnosis was significantly higher among males than females overall (OR, 1.31; 95% CI, 1.12 to 1.54) and for A(H3N2) (OR, 1.45; 95% CI, 1.17 to 1.81) and B(Victoria) (OR, 1.83; 95% CI, 1.13 to 2.94; not adjusted for calendar time due to sample size limitations), but not A(H1N1)pdm09 (OR, 1.29; 95% CI, 0.92 to 1.80) or B(Yamagata) (OR, 1.03; 95% CI, 0.71 to 1.50). Conversely, this same effect was not observed among unvaccinated participants overall (OR, 1.06; 95% CI, 0.97 to 1.16) or for any subtype/lineage (data not shown).

In the overall age-stratified analysis, the greatest absolute differences in VE between females and males were observed in older adults ≥50 years (Figure 2). Among adults ≥50 years, the adjusted VE was 48% (95% CI, 38% to 57%) vs 29% (95% CI, 10% to 44%) in females and males, respectively (AD, 19%; P = .03), whereas the VE was 49% (95% CI, 31% to 62%) vs 45% (95% CI, 24% to 59%; AD, 4%; P = .74) in those age <20 years and 47% (95% CI, 37% to 56%) vs 48% (95% CI, 33% to 60%; AD, −1%; P = .90) in those age 20–49 years, the latter age group comprising the majority of SPSN participants. The prevalence of comorbidities increased with age but did not significantly differ between females and males (20% vs 19%; P = .23), except in those age 50–64 years (27% vs 33%; P < .01). When an additional interaction term for comorbidity*sex was included in the fully adjusted VE model, including age adjustment, sex differences persisted: 48% (95% CI, 42% to 54%) in females vs 39% (95% CI, 29% to 48%) in males (AD, 9%; P = .08 for vaccine*sex and P = .21 for comorbidity*sex).

By subtype/lineage, larger absolute differences were seen among children and adolescents age <20 years for A(H1N1)pdm09 and B(Victoria) and older adults age ≥50 years for A(H3N2), A(H1N1)pdm09, and B(Victoria) (Supplementary Table 3). Smaller or negative absolute differences were seen across all subtypes/lineages for adults age 20–49 years. The addition of an interaction term for age*sex did not meaningfully change the pattern of higher VE in females (48%; 95% CI, 41% to 54%) vs males (40%; 95% CI, 30% to 49%) in the overall analysis (AD, 8%; P = .16 for vaccine*sex and P < .01 for age*sex). By subtype/lineage, the VE estimates were also similar when adjusted for the age*sex interaction, although the P value for the age*sex interaction term was only statistically significant for B(Yamagata) (Supplementary Table 4).

The finding of higher VE in females was generally consistent across seasons, although the opposite pattern of higher VE in males was observed in 2011–2012 for B(Victoria), 2012–2013 for A(H3N2) and A(H1N1)pdm09, and in 2014–2015 and 2015–2016 for B(Yamagata) (Supplementary Table 5). Few season-specific comparisons reached statistical significance, likely due to the smaller sample size.

DISCUSSION

Our analysis investigated sex as an effect modifier in the association between influenza vaccination and medically attended outpatient illness across 7 influenza seasons in Canada. Sex is often considered a potential confounder in the analysis of influenza VE [1], although it is not consistently included in adjusted VE estimation and the association will likely vary by study

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**Figure 2.** Vaccine effectiveness estimates by sex for patient age groups. P values indicate significance of vaccine*sex interaction. Covariates included in the interaction model were vaccination status (no, yes), sex (female, male), comorbidity (no, yes), province (AB, BC, ON, QC), collection interval (≤4, 5–7 days), calendar time (week of specimen collection based on cubic B-spline with 3 equal knots), season, and vaccine*sex. Abbreviations: AD, absolute difference (Δ female – male); CI, confidence interval; VE, vaccine effectiveness.
population and setting [17]. In our own SPSN VE analyses, sex was included as a covariate for some seasons where a bivariate association was seen for both vaccination status and influenza test positivity [14, 15] but was otherwise omitted.

Few studies have investigated the role of sex as an effect modifier for influenza-related outcomes. In a cohort study of community-dwelling elderly adults from 1990–2000, Nichol et al. found evidence of an interaction between vaccination and sex for all-cause mortality, with lower effectiveness found in males (P = .03) [18]. Vila-Córcoles et al. found a lower risk for all-cause mortality in females compared with males across all age groups in community-dwelling elderly adults in Spain from 2002–2005, but with the difference in mortality risk between females and males narrowing as age increased [19]. However, both of these studies were limited to a population of elderly adults age ≥65 years, non-specific outcomes, and observational study designs; as such, they are likely not directly comparable to our findings and may suffer from other systematic biases.

In our own study using laboratory-confirmed outcomes and a test-negative design, we observed a pattern of higher VE in females, suggesting that females may respond better to influenza vaccine than their male counterparts. Our finding of increased odds of influenza among vaccinated but not unvaccinated male vs female participants for A(H3N2) and B(Victoria) reinforces our interpretation. A theoretical advantage of the test-negative design for influenza VE evaluation is that it minimizes biases associated with health care–seeking behavior as all participants presented to health care and met a standardized testing indication for influenza-like illness. Together, these findings suggest that biological sex differences in the response to vaccine, rather than gender differences in health care seeking or vaccination status reporting, likely explain the observed differences in influenza VE between males and females.

Although few studies have investigated sex differences in vaccine protection, females have been shown to have stronger innate and adaptive immune responses, including more pronounced antibody response to influenza vaccine, in association with higher rates of local and systemic adverse events following immunization [20–24]. The biological mechanisms underpinning these sex differences are not well understood. Some have attributed these differences to sex steroids that alter the function of immune cells by binding to specific receptors and influencing cell signaling pathways [20, 21]. At certain concentrations, estrogens, particularly estradiol, can function in a pro-inflammatory role, whereas testosterone and progesterone are considered immunosuppressive [20, 21, 25]. Hormone-mediating factors, such as pregnancy, can also modulate the immune response to influenza infection [1, 20, 21]. Although data on pregnancy were not available for this study, sex effects were not apparent among adults during their prime reproductive years, and pregnant women would have comprised only a small proportion of study participants. We observed the greatest absolute differences in VE among older adults (during or after the onset of menopause in females), and to a lesser extent in young children (before the onset of puberty) for certain subtypes/lineages. As such, our age-related findings are consistent with the epidemiological studies cited above but may be contrary to expected patterns based on sex hormone mechanisms alone [20, 21]. Mutations or polymorphisms in genes on the X chromosome that encode immunological proteins can affect activation of cytokine receptors and regulatory processes [20, 21]. Other sex- or gender-dependent factors should also be considered in the interaction between sex and influenza VE.

This study was limited by the available sample size despite pooling across multiple seasons. Although results were generally consistent across outcomes and seasons, the overall effects were small and may be due to chance. In the combined all-season analysis, the interaction term for vaccine*sex was marginally significant at the α < .10 level for A(H3N2) and B(Victoria) outcomes. The interaction terms for A(H1N1)pdm09 and B(Yamagata) were not statistically significant, despite higher VE in females for A(H1N1)pdm09. In our analysis, sex differences were greatest for influenza A(H3N2) and B(Victoria) in older age groups ≥50 years. Influenza A(H3N2) is associated with greater disease burden among elderly adults, whereas A(H1N1)pdm09 is notable for its impact in younger adults [26–27]. Both influenza B(Victoria) and B(Yamagata) disproportionately affect children, with B(Yamagata) exhibiting a bimodal age distribution also affecting older adults [28]. Other factors, such as the higher prevalence of comorbidities in older adults, may also contribute to these findings, although VE analyses were adjusted for age and comorbidity. Ultimately, however, it remains possible that our observational study design did not adequately account for other potential biases, residual confounders, or interactions.

In conclusion, we observed a modest effect of sex on influenza VE across most outcomes and seasons, with higher VE estimates generally seen among females. These sex effects were age dependent, with greater effects in older adults age ≥50 years compared with those age <20 years or 20–49 years, but with some variation by subtype/lineage. Overall, these effects likely represent a complex interplay between birth cohort (ie, immunological prime-boost) effects, hormonal influences, and other contributing agent–host–environment factors [1, 15, 20, 21]. The clinical implications are unclear, although some have argued for sex-based design of influenza vaccination strategies [21]. More definitive investigations of VE by sex and age are ultimately needed to elucidate these potential interactions.

**Supplementary Data**

Supplementary materials are available at *Open Forum Infectious Diseases* online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.
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