The responses of Aboriginal Canadians to adjuvanted pandemic (H1N1) 2009 influenza vaccine

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Background: Because many Aboriginal Canadians had severe cases of pandemic (H1N1) 2009 influenza, they were given priority access to vaccine. However, it was not known if the single recommended dose would adequately protect people at high risk, prompting our study to assess responses to the vaccine among Aboriginal Canadians.

Methods: We enrolled First Nations and Métis adults aged 20–59 years in our prospective cohort study. Participants were given one 0.5-mL dose of ASO3-adjuvanted pandemic (H1N1) 2009 vaccine (Arepanrix, GlaxoSmithKline Canada). Blood samples were taken at baseline and 21–28 days after vaccination. Paired sera were tested for hemagglutination-inhibiting antibodies at a reference laboratory. To assess vaccine safety, we monitored the injection site symptoms of each participant for seven days. We also monitored patients for general symptoms within 7 days of vaccination and any use of the health care system for 21–28 days after vaccination.

Results: We enrolled 138 participants in the study (95 First Nations, 43 Métis), 137 of whom provided all safety data and 136 of whom provided both blood samples. First Nations and Métis participants had similar characteristics, including high rates of chronic health conditions (74.4%–76.8%). Pre-existing antibody to the virus was detected in 34.3% of the participants, all of whom boosted strongly with vaccination (seroprotection rate [titre ≥ 40] 100%, geometric mean titre 531–667). Participants with no pre-existing antibody also responded well. Fifty-eight of 59 (98.3%) First Nations participants showed seroprotection and a geometric mean titre of 353.6; all 30 Métis participants with no pre-existing antibody showed seroprotection and a geometric mean titre of 376.2. Pain at the injection site and general symptoms frequently occurred but were short-lived and generally not severe, although three participants (2.2%) sought medical attention for general symptoms.

Interpretation: First Nations and Métis adults responded robustly to ASO3-adjuvanted pandemic (H1N1) 2009 vaccine. Virtually all participants showed protective titres, including those with chronic health conditions.

Trial registration: ClinicalTrials.gov trial register no. NCT.01001026.
ASO3-adjuvanted vaccine (Pandemrix, GlaxoSmithKline, Rixensart, Belgium) given to 65 adults aged 18–60 years. The European product was believed to be equivalent to the Canadian-made vaccine, but this had not yet been shown.

We wondered if the recommended single dose would be adequate for Aboriginal Canadian adults given their heightened risk of severe influenza during the first wave. We were unable to identify any previous studies of influenza vaccines involving Aboriginal Canadians to determine if their responses would be similar to other Canadians or to the healthy European study participants on whom the dosing recommendation was based. Consequently, we undertook a study involving First Nations and Métis adults to assess their responses to the pandemic vaccine.

Methods

Eligible participants were self-identified First Nations or Métis adults aged 20–59 years, including people with chronic health conditions not associated with immune dysfunction. Most of the participants recruited to the study were urban residents. Exclusion criteria included pregnancy, allergy to eggs, bleeding disorders and prior receipt of the 2009–10 seasonal trivalent influenza vaccine. The study period lasted from November 2009 to January 2010 during the second wave of the influenza pandemic in Canada.

Blood samples to measure titres of hemagglutination-inhibiting antibodies against the pandemic (H1N1) 2009 influenza virus were obtained at baseline and 21–28 days after vaccination. Participants were given a single 0.5-mL dose of Arepanrix vaccine injected into the deltoid muscle. One commercial lot of vaccine was used containing 3.75 µg (H1N1) 2009 influenza hemagglutinin per dose, with ASO3 adjuvant. To assess vaccine safety, participants kept a daily diary for seven days after vaccination to record any changes at the injection site or general symptoms, including fever. Thereafter, only health events requiring medical attention were documented.

At the first visit, participants were given the vaccination and a nasal test kit (flocked swab, transport medium) to take home and use if they developed an influenza-like illness (with cough and fever) while enrolled in the study. Participants were provided with instructions for using the kit, and reminders were given during each subsequent contact. When a kit was used, arrangements were made to collect the sample and have it tested for influenza virus by polymerase chain reaction at a central laboratory.

Study staff reviewed the diaries and influenza-like illnesses with participants by telephone on day 7 and in person 21–28 days after vaccination. Paired sera were tested for hemagglutination-inhibiting antibodies at the National Microbiology Laboratory in Winnipeg, Manitoba. Standard criteria were used to assess antibody responses, including seroprotection rates, geometric mean titres, geometric mean fold rises and seroconversion rates. Antibody titres were expressed as the reciprocal value (e.g., 40 in place of 1:40). Because protocol adherence was excellent, the same participants were included in the immunogenicity and safety analyses.

We had intended to enrol 200 participants in the study. If a minimum of 110 evaluable participants could be enrolled, the study would have at least 80% power to detect a 15% or greater departure from the 100% seroprotection rate reported in the earlier study of ASO3-adjuvanted pandemic (H1N1) 2009 influenza vaccine.

Fisher exact test was used to compare proportional data, and analysis of variance (ANOVA) was used to compare continuous data, including geometric mean titre and geometric mean fold rise.

This study was undertaken by the Public Health Agency of Canada/Canadian Institutes of Health Research Influenza Research Network (PCIRN) centres in Winnipeg; Edmonton, Alberta; and Vancouver, British Columbia. The funders played no role in the analysis of the data or in the design and conduct of the study. Aboriginal community leaders were consulted to obtain their approval and advice for implementing the protocol in a culturally appropriate manner. The protocol was approved by the research ethics boards of Health Canada and the participating institutions, including the University of British Columbia Clinical Research Ethics Board as the lead institution. The trial was registered with ClinicalTrials.gov as trial number NCT.01001026. Written informed consent was obtained from each participant upon enrollment in the study.

Results

We enrolled 138 participants in the study (100 from Winnipeg, 27 from Edmonton and 11 from Vancouver), and 137 participants provided symptom summaries at both interviews. Blood samples were obtained from 137 participants at baseline and from 136 participants after vaccination, for a protocol completion rate of 98.6%.

The demographics of the participants from the First Nations (n = 95) and Métis (n = 43) groups were similar (Table 1). First Nations participants gave 25 Nation/Band affiliations, with Ojibway (33 people) and Cree (23 people) predominating. Underlying health conditions were reported by 76.8% of First Nations participants...
and by 74.4% of Métis participants. The most commonly reported conditions were respiratory (17 participants, 8 of whom reported asthma), cardiovascular (17 participants, 13 of whom reported hypertension), endocrine (15 participants, 8 of whom reported diabetes mellitus and 7 of whom reported thyroid disorders), gastrointestinal (12 participants, 3 of whom reported chronic hepatitis), dermatologic (12 participants, 6 of whom reported eczema and 3 of whom reported psoriasis), psychological (11 participants, 10 of whom reported depression) and hematologic disorders (7 participants, 6 of whom reported anemia). Eleven participants had body mass indices greater than 40 kg/m². Thirty-eight participants (27.5%) had conditions that posed high risk for influenza complications as per Canadian guidelines. None of the participants were receiving immunosuppressive medications, including one participant with sarcoidosis and one participant with rheumatoid arthritis.

Hemagglutination-inhibiting antibody responses are summarized in Table 2. Pre-existing antibody to the pandemic (H1N1) 2009 influenza virus was detectable in about one-third of participants (47, 34.3%). The responses of primed (baseline titre ≥ 10) and naive participants (i.e., people with no detectable antibody at baseline) were assessed separately. After vaccination, at least 99.3% (95% confidence interval [CI] 96.0–100.0) of First Nations and Métis participants had protective titres (≥ 40). Primed and naive participants in both groups mounted robust responses as measured by geometric mean titres (Table 2). However, postvaccination geometric mean titres were significantly higher in primed participants than in naive participants (ANOVA, p < 0.01).

Symptoms during the week after vaccination were frequent but generally mild (Appendix 1, available at www.cmaj.ca/lookup/suppl/doi:10.1503/cmaj.110196/-/DC1). Pain at the injection site was noted by 75.9% of participants, but only 5.8% rated it as being severe (i.e., limiting the participant’s activities). Redness at the injection site was reported by 12.4% of participants, and swelling was reported by 18.2% of participants. The reported rates of general symptoms during the week after vaccination were as follows: myalgia, 54.0%; tiredness, 52.6%; headache, 40.1%; malaise, 32.8%; and arthralgia, 26.3%. No fevers of 38.0°C or higher were recorded. Most symptoms resolved by day 6 postvaccination. Three participants (2.2%) sought medical attention for general symptoms, and no one sought medical attention for local symptoms. Three participants required admission to hospital during the study period for illnesses unrelated to vaccination. Three participants described having an influenza-like illness before their final visit, but the results of polymerase chain reaction tests for influenza were negative.

**Interpretation**

Both First Nations and Métis adults responded robustly to the adjuvanted vaccine. Participants with evidence of prior infection with the pandemic (H1N1) 2009 influenza virus boosted well with vaccination (Table 2), and those with no pre-existing antibodies also had strong responses to vaccination. The observed seroprotection rate of 98.8% (95% CI 94.2–100.0) among naive participants was remarkable, exceeding the 89.2% (95% CI 81.1–94.7) seroprotection rate of naive, non-Aboriginal adults in a concurrent PCIRN study of the same vaccine involving people in the same age group. The sizes of response (geometric mean titres) were significantly higher among First Nations and Métis participants (Table 2) than among non-Aboriginal Canadian adults (geometric mean titre 141, 95% CI 109–183) in the concurrent study, attesting to the adequacy of responses of both First Nations and Métis participants. However, the “seroprotection rate” is a correlate of protection based on seasonal influenza vaccines, which might not apply to a pandemic virus. With seasonal vaccines, a hemagglutination-inhibiting antibody titre of 1:40 in young adults correlated to 50% protective efficacy, with higher titres affording greater protection. The mean titre in the current study was 10-fold higher than the threshold value, thus

| Characteristic                             | First Nations n = 95 | Métis n = 43 | p value† |
|--------------------------------------------|----------------------|--------------|----------|
| Age, yr, mean (SD)                         | 36.2 (10.1)          | 39.7 (12.2)  | 0.08‡    |
| Sex                                         |                      |              |          |
| Men                                        | 34 (35.8)            | 13 (30.2)    | 0.57§    |
| Women                                      | 61 (64.2)            | 30 (69.8)    | —        |
| Body mass index, kg/m², mean (SD)          | 29.6 (5.9)           | 30.2 (6.5)   | 0.58‡    |
| One or more chronic health conditions      | 73 (76.8)            | 32 (74.4)    | 0.83§    |
| Past influenza vaccination                  | 46 (48.4)            | 14 (32.6)    | 0.10§    |

Note: SD = standard deviation.  
*Unless otherwise indicated.  
†No statistically significant differences existed between groups.  
‡Analysis of variance.  
§Fisher exact test.
increasing the likelihood of protection.

The robust nature of the immune response of Aboriginal adults to influenza vaccine was unanticipated. An analogous situation might be hepatitis C infection, which North American Aboriginal people have an enhanced capacity to resolve. Studies of innate responses of First Nations volunteers in Manitoba showed a greater tendency toward a proinflammatory cytokine milieu in response to hepatitis C virus, potentially aiding clearance of infection. A similar proinflammatory innate response to adjuvanted influenza vaccine could plausibly result in stronger antibody responses. A greater pro-inflammatory response to influenza infection might be the basis for the severe disease that affected some First Nations people during the pandemic.

Aboriginal participants reported having adverse events during the week after immunization at rates similar to those seen in other studies that evaluated ASO3-adjuvanted pandemic (H1N1) 2009 vaccines in adults. No significant differences in these rates were seen between the Aboriginal participants in our study and the non-Aboriginal participants in the concurrent PCIRN study. Pain at the injection site and general symptoms of tiredness or myalgia were described by over half of the participants in this study during the week after vaccination. These and other symptoms were generally mild and short-lived, having little impact on the ability to perform daily activities. No serious adverse events were attributed to vaccination.

Table 2: Hemagglutination-inhibiting antibody responses to adjuvanted pandemic (H1N1) 2009 influenza vaccine among First Nations and Métis adults enrolled in the study

| Antibody response          | No. (%)* | First Nations | Métis |
|----------------------------|----------|---------------|-------|
|                            |          | n = 94        | n = 43|
| Baseline                   |          |               |       |
| Titre ≥ 10                 |          | 34 (36.2)     | 13 (30.2) |
| Seroprotective titre ≥ 40 |          | 16 (17.0)     | 7 (16.3) |
| Geometric mean titre (95% CI) | 10.1 (8.0–12.8) | 8.6 (6.4–11.7) |
| Postvaccination             |          |               |       |
| Seroconversion ≥ 4-fold titre increase | n = 93 | 86 (92.5) | 42 (97.7) |
| Naive                      |          | n = 59        | n = 30 |
|                           |          | 59 (98.3)†    | 30 (100.0) |
| Primed                     |          | n = 34        | n = 13 |
|                           |          | 28 (82.4)†    | 12 (92.3) |
| Seroprotective titre ≥ 40, no. (%; 95% CI) | n = 93 | 92 (98.9; 94.2–100.0) | 43 (100.0; 91.8–100.0) |
| Naive                      |          | n = 59        | n = 30 |
|                           |          | 58 (98.3; 90.9–100.0) | 30 (100.0; 88.4–100.0) |
| Primed                     |          | n = 34        | n = 13 |
|                           |          | 34 (100.0; 89.7–100.0) | 13 (100.0; 75.3–100.0) |
| Geometric mean titre (95% CI) | 445.8 (349.3–569.1) | 417.5 (288.6–604.0) |
| Naive                      |          | 353.6 (261.9–477.3)† | 376.2 (234.1–604.4) |
| Primed                     |          | 666.6 (446.3–995.8)† | 531.0 (285.6–987.4) |
| Geometric mean fold rise (95% CI) | 43.8 (32.1–59.6) | 48.3 (30.6–76.2) |
| Naive                      |          | 69.5 (51.5–93.8)† | 73.5 (45.9–117.8)† |
| Primed                     |          | 19.6 (10.8–35.6)† | 18.3 (7.2–46.6)† |

Note: CI = confidence interval.
*Unless otherwise indicated.
†p < 0.01, comparing primed (titre ≥ 10 at baseline) and naive subjects. No significant differences existed between responses of First Nations and Métis subjects. Seroconversion rates were compared using Fisher exact test, and geometric mean titre and geometric mean fold rise were compared using analysis of variance.
25 different bands and First Nations, as well as Métis. Studying selected reserve communities would not have resulted in as broad a sample population. One-third of the participants had evidence of prior infection with pandemic (H1N1) 2009 influenza upon entry into the study, confirming that participants were drawn from populations at risk of exposure during the first wave. Including participants with chronic health conditions made the study more representative of the Aboriginal population, in contrast to typical vaccine studies that limit enrolment to healthy volunteers.

Because many potential participants had received the pandemic vaccine before our study began, our sample was smaller than intended but still within the desired range. Our results might not apply to all First Nations and Métis people, nor might they apply to Inuit adults, who were not included in the study but who had high rates of severe infection during the pandemic. Antibody titres might have been increased by subsequent exposures to the pandemic virus as the study was conducted during the second wave of infections. However, most participants were enrolled in Winnipeg, which had a milder second wave than the other sites.

There was a high prevalence of underlying medical conditions among the Aboriginal adults in this study. Many of these conditions pose an increased risk for severe influenza infection thus warranting annual vaccination. It would seem prudent to determine if satisfactory responses also follow the administration of unadjuvanted seasonal influenza vaccines among people at high risk.

Conclusion
Adjuvanted pandemic (H1N1) 2009 influenza vaccine resulted in high antibody titres in vulnerable First Nations and Métis people.

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