SOGC Statement on COVID-19 Vaccination in Pregnancy

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Pre-amble
The SOGC acknowledges the need for guidance related to the COVID-19 vaccine and pregnancy and during lactation. We recognize the difficulty facing women and their health care providers at this time, due to the absence of clinical trials that can support evidence-informed recommendations about the COVID-19 vaccine for pregnant and breastfeeding populations. Information related to COVID-19, the impact of the disease on pregnancy and data related to COVID-19 vaccines in development are rapidly evolving. The information contained herein is subject to change as further evidence becomes available and currently applies to the RNA based COVID-19 vaccines.

Consensus Statement: Women who are pregnant or breastfeeding should be offered vaccination at anytime if they are eligible and no contraindications exist.

This decision is based on the women’s personal values and an understanding that the risk of infection and/or morbidity from COVID-19 outweighs the theorized and undescribed risk of being vaccinated during pregnancy or while breastfeeding. Women should not be precluded from vaccination based on pregnancy status or breastfeeding.

SARS-CoV-2 and the impact on pregnancy
Most pregnant women who become infected with SARS-CoV-2 have mild-to-moderate symptoms and many are asymptomatic. However, both Canadian and international data from large studies spanning multiple jurisdictions demonstrate that approximately 8-11% of pregnant women require hospitalization for COVID-related morbidity and between 2-4% of pregnant women require admission to an intensive care unit. Compared to non-pregnant women with COVID-19, pregnant women are at increased risk of invasive ventilation with an equivalent mortality to age-matched peers. The risk of severe morbidity from COVID-19 in pregnant women appears to be associated with risk factors including age ≥ 35 years old, asthma, obesity, preexisting diabetes, preexisting
hypertension and heart disease.\(^2\)\(^4\) In addition, both Canadian and US data show an increased risk of preterm birth associated with COVID-19 infection in pregnancy, which will cause consequent morbidity to the infant related to prematurity.\(^2\)\(^3\)\(^4\)

**COVID-19 Vaccines**

On December 9\(^{th}\), 2020, Health Canada authorized the first COVID-19 vaccine in Canada: Pfizer-BioNTech COVID-19 vaccine.\(^5\) Shortly thereafter, a second vaccine was approved - the Moderna COVID-19 vaccine. Both vaccines approved to date have a messenger RNA (mRNA) platform. The Pfizer-BioNTech COVID-19 vaccine uses a new mRNA vaccine platform. This model consists of mRNA encapsulated by a lipid nanoparticle, which allows the mRNA entrance into host (human) cells. The mRNA in the vaccine codes for the SARS-CoV-2 spike protein, which is utilized by the virus to bind to human receptors and promote viral replication. The vaccine provides the host cell instructions to manufacture only this spike protein and express it on its surface. Recognizing the spike protein as a foreign antigen, the host immune system is then activated to produce an immune response.\(^6\) The mRNA does not enter the nucleus or alter human DNA and human cells do not have the machinery to allow it to do so.

The Pfizer-BioNTech and Moderna COVID-19 vaccines are administered as a series of two intramuscular injections given 21-28 days apart.\(^7\) The safety and efficacy Pfizer-BioNTech COVID-19 vaccine has been demonstrated for adults 16 years and older in Phase II and Phase III trials involving the randomization of approximately 44,000 individuals.\(^8\) These trials demonstrated a vaccine efficacy of 94.6% for preventing symptomatic COVID-19 cases at least 7 days following the second dose.\(^8\)

The available safety data is based on an interim analysis of 37,586 adults, of whom approximately 9,500 individuals had at least 2 months of follow-up after receiving the vaccine. There were no clinically meaningful differences in adverse events or severe adverse events in the vaccine group compared to control except for lymphadenopathy which occurred in 0.3% of the vaccine group compared to <0.1% of the placebo group. The most commonly reported side effects from the vaccine were pain at the injection site (66-83%), fatigue (51-59%) and headache (39-52%). Fever was reported in 11-16% of patients, particularly following the second dose.\(^8\) Safety monitoring will continue for 2 years following vaccine administration.
Pregnant and breastfeeding women were excluded from the Phase II and Phase III studies for the Pfizer-BioNTech COVID-19 vaccine. However, there were 23 women (12 in the vaccine arm and 11 in the placebo arm) who reported pregnancies during the trial and who are being followed for pregnancy outcomes with no reports of adverse effects to date. Currently, there are no other safety or efficacy data available for pregnant or breastfeeding women. The Developmental and Reproductive Toxicity (DART) animal studies for the Moderna and Pfizer-BioNTech vaccines are ongoing. According to the World Health Organization and the American College of Obstetricians & Gynecologists, no major safety signals have been identified.

To date, Phase III data is available for two other COVID-19 vaccines. The Moderna mRNA-1273 vaccine also utilizes mRNA vaccine technology with the SARS-CoV-2 spike protein as its antigenic target. In Phase III trials randomizing 30,000 individuals, the vaccine was reported to have 94.1% efficacy against symptomatic COVID-19 with no serious safety concerns identified during the initial 2 month follow-up period. For the Moderna vaccine, the most common side effects were fatigue, myalgia and headache, with no difference in rates of serious adverse events in the vaccine compared to placebo group. Although the study excluded pregnant and breastfeeding women, 13 individuals were found to be pregnant during the trial thus far, 6 in the vaccine arm and 7 in the placebo arm. The outcomes of the pregnancies are being monitored and will be reported upon. Of note, reproductive toxicity studies of the Moderna vaccine in rats, administered during mating and in gestation, have revealed no signals of reproductive or developmental concerns.

The Oxford-AstraZeneca ChAdOx1 nCoV-19 vaccine is not an mRNA vaccine. Instead, it utilizes a Chimpanzee adenovirus vector vaccine platform and preliminary Phase III data from 11,636 participants demonstrates an overall vaccine efficacy of 70.4% against symptomatic COVID-19 disease. Unfortunately, all the vaccines for which Phase III results are available excluded pregnant or breastfeeding women from their trials.

Similarly, breastfeeding women were also excluded from the Phase III trials available at present. Therefore, there is no data on the safety of COVID-19 vaccines in lactating women or the effects of mRNA vaccines on the breastfed infant or on milk production. Because mRNA vaccines are not considered live virus vaccines, they are not hypothesized to be a risk to the breastfeeding infant.
Considerations for COVID-19 Vaccination During Pregnancy and Breastfeeding

Decades of experience with other vaccines administered during pregnancy would suggest that we could expect a similar efficacy for the COVID-19 vaccines in pregnant women compared to non-pregnant women. Vaccines in general are immunogenic, safe, and efficacious when delivered to pregnant women. While there have been no red flags or hypothesized mechanisms for potential harm associated with the administration of an mRNA vaccine during pregnancy, until more data is available, the potential risks of vaccination to a pregnant woman and her fetus remain unknown. What is known, however, is that an unvaccinated pregnant woman remains at risk of COVID-19 infection and remains at heightened risk of severe morbidity if infected compared to non-pregnant counterparts. Severe infection with COVID-19 carries risks to both maternal, fetal and neonatal health. While pregnancy itself does not appear to increase the risk of becoming infected with SARS-CoV-2, pregnant individuals may be in work-related (e.g., health-care worker, front line workers etc.) or community situations (e.g., caregiver, indigenous communities, outbreak setting, etc.) where the risk of infection is considerable. Owing to maternal age or underlying comorbidities, some pregnant individuals are at high risk of severe COVID-related morbidity.

NACI has advised “that a complete vaccine series with a COVID-19 vaccine may be offered to pregnant individuals in the authorized age group, without contraindications to the vaccine, if a risk assessment deems that the benefits outweigh the potential risks for the individual and the fetus, and if informed consent includes discussion about the absence of evidence on the use of COVID-19 vaccine in this population (Discretionary NACI Recommendation)”.

We recommend that pregnant and breastfeeding women who are eligible for the COVID-19 vaccine due to exposure risk, medical status, or other circumstances should be able to make an informed decision by having access to up-to-date information about the safety and efficacy of the vaccine (including clear information about the data that is not yet available) and information about the risks of COVID-19 infection for them. The concern around vaccination in the absence of evidence of safety in pregnancy has been debated in the literature. The PREVENT Working Group state, “the absence of evidence and the mere theoretical or even documented risk of fetal harm is generally not sufficient to justify denying pregnant women access to a vaccine in an outbreak or epidemic.” During an epidemic, the default should be to offer vaccines to pregnant women alongside other affected populations.
Universal exclusion of pregnant women from receiving the COVID-19 vaccine based on an undocumented and hypothetical risk to the fetus would leave pregnant women vulnerable to severe morbidity and their infant to preterm birth risk, which would compromise fetal health. Conversely, lack of safety and efficacy data for this population precludes making a recommendation for routine COVID-19 vaccination for all pregnant and breastfeeding individuals.

Pregnant and breastfeeding individuals will likely look to their prenatal care provider to assist in making decisions weighing the risks and benefits so that they might arrive at a well informed and autonomous decision that is right for them as an individual. Such a discussion should prioritize patient autonomy and should include, but not be limited to assessment of:

- Local epidemiology and risk of community acquisition of COVID-19
- Workplace situation and risk of work-related acquisition of COVID-19
- Individual risk for COVID-related morbidity including consideration for comorbidities such as advanced maternal age, immunosuppressive conditions, pre-existing diabetes, pre-existing hypertension, obesity or chronic respiratory and/or cardiac conditions
- Available data related to the safety of the vaccine during pregnancy and lactation
- Data that is not yet available related to the safety and efficacy of the vaccine for pregnant and breastfeeding women
- Individual beliefs and personal risk assessment of the available data

**Individuals who proceed with vaccination**

Individuals should be informed of the expected side effects following vaccination. While pain at the injection site, fatigue and headache are the most commonly reported symptoms following vaccination, fever was reported 16% of the time for young, non-pregnant individuals. Pregnant women can be counselled to treat mild post-vaccination fevers with antipyretics (e.g., acetaminophen). Active surveillance is ongoing for the Pfizer-BioNTech COVID-19 vaccine. As such, prenatal care providers are encouraged to inform themselves on local procedures for active surveillance and to notify the appropriate channels when a pregnant or breastfeeding mother is receiving a dose of the vaccine.
Timing of vaccination during pregnancy and vaccine interval

In theory, immunization of a pregnant woman may confer benefit to a newborn infant through a mechanism of maternal vaccination similar to what is seen for pertussis and influenza vaccination during pregnancy. However, until such time when a newborn benefit is confirmed, the primary indication for administration of a COVID-19 vaccine to a pregnant individual remains for maternal protection. As such, for now, there is no data to guide administration at a particular gestational age and it may be considered at any gestational age including the first trimester.

There is no clear evidence to direct if vaccine spacing is required. In the absence of evidence, NACI recommends spacing the administration of any other vaccines for 28 days from completion of the COVID-19 vaccines, however other jurisdictions do not recommend any specific spacing and simultaneous administration of other vaccines can occur. COVID-19 vaccines, out of prudence, can be spaced 14 days from any other vaccines. The spacing recommendation is due to the theoretical risk of an increased inflammatory response, and the potential of vaccine adverse events between different vaccines, and not from data that shows a direct effect on efficacy of adverse events. In addition, administration of immunoglobulins is thought to interfere with vaccine efficacy due to live attenuated vaccine and to circulating levels of antibody within the population. However, the rates of circulating antibodies to COVID-19 are low therefore, the impact on vaccine efficacy of COVID-19 is unclear.

Given this, the following can be recommended:

- Wait 14 days after any other vaccine before receiving a COVID-19 vaccine. However, given the context of the global pandemic, simultaneous or closer interval of administration may be considered.
- After receiving a COVID-19 vaccine dose, where possible wait 28 days before receiving any other vaccine, unless a vaccine is required urgently due to an exposure to a virus such as Hepatitis B. Again, given the global pandemic and condensed timelines of pregnancy this may not be possible.
- Time-sensitive interventions such as administration of anti-D immunoglobulin and blood products should not be delayed on account of recent COVID-19 vaccination.
**Vaccination of the pregnant patient in the context of limited vaccine supply**

Certain jurisdictions may manage interruptions of vaccine supply chain by delaying administration of the second dose of a COVID-19 vaccine. There are no physiologic reasons to anticipate that the effect of delaying the second dose of the COVID-19 vaccine would be different for a pregnant individual compared to a non-pregnant individual. Pregnant individuals may resume their vaccine series akin to the non-pregnant population in situations of supply chain interruptions.

In the context of limited vaccine supply, distribution of vaccination will be prioritized differently in each jurisdiction depending on local epidemiology and public health priorities. Decisions made regarding prioritization of pregnant women should reflect that pregnancy, appears to carry increased risk for COVID-related hospitalization, intensive care unit admission and mechanical ventilation. Other additional factors in pregnancy such as advanced maternal age, immunosuppressive conditions, pre-existing diabetes, pre-existing hypertension, obesity or chronic cardiac and respiratory conditions may provide additional risk to pregnant women and could be considered for further prioritization in the context of limited vaccine supply.

**Inadvertent pregnancy following vaccination**

Individuals who are discovered to be pregnant during their vaccine series or shortly afterward should not be counselled to terminate pregnancy based on having received the vaccine. If conception is presumed to predate the first dose, it is recommended to follow the same procedures for active surveillance (as available) as would be activated if the pregnancy was known at the time of vaccination.

Where pregnancy is detected during the vaccine series (i.e. following the first dose, but ahead of the second dose), the decision of whether to complete the vaccine series during pregnancy should be based on an assessment of the potential risks of not being completely vaccinated during pregnancy versus the potential risks of receiving the vaccine during pregnancy (as discussed above). Women should not be precluded or forced to delay the vaccine series. There is no current evidence of teratogenicity within the limited data to date, however this has not been studied as none of the vaccines have been studied in pregnant women. The risk remains of teratogenicity remains theoretical and should not preclude vaccination.
A registry to track pregnancy outcomes for those women that receive any vaccine doses in pregnancy is planned for Canada.

**Individuals contemplating pregnancy**

For an individual planning a pregnancy, it is recommended to complete the entire COVID-19 vaccination series (where possible) to achieve maximal vaccine efficacy ahead of pregnancy. It is not known whether an individual should delay pregnancy following receipt of the vaccine and a risk-benefit discussion for those planning pregnancy should occur similar to the discussion for pregnant and breastfeeding women.

**Future research**

As the evidence evolves, it is becoming clear that pregnant and postpartum women represent a population at increased risk of COVID-related morbidity. Severe COVID-19 infection during pregnancy has important implications for both maternal and fetal health. NACI acknowledges that people of reproductive age constitute a substantial proportion of the Canadian population, yet no data on the use of COVID-19 vaccine in pregnancy are available. We support NACI’s recommendation for the inclusion of pregnant women in clinical trials of COVID-19 vaccines to ensure that this population has equitable access to COVID-19 vaccine options informed by robust safety, immunogenicity, and efficacy data.}\(^5\)
References


