

Vaccine Medical Directive and Delegation Moderna COVID-19 Vaccine

	Delegation of Authority to		
DELEGATED	Delegation of Authority to:		
PROCEDURE	Prescribe a drug		
	Sell a drug		
ORDER TO	Administer Dispense Sell		
AUTHORIZING MD			
AUTHORIZED IMPLEMENTERS	Public Health Sudbury & Districts Public Health Nurses, Registered Nurses, Registered Practical Nurses, graduates of an accredited Nursing Program in Ontario, post-secondary nursing students, medical students of an accredited Medical Program in Ontario, Midwives, Radiation Therapists, Respiratory Therapists, Physician Assistants, Pharmacists and Paramedics who have completed their Certification of Competence Module.		
	Paramedic students from Collège Boréal and Cambrian College who have received formal didactic and practical education in IM, medication administration, and sharp safety, in a formative and summative evaluation process, following the paramedic NOCP's (National Occupational Competency Profile). This was completed in a supervised setting with certified faculty from Collège Boréal and Cambrian College. Second year RPN students from Collège Boréal who have received formal didactic and practical education in IM, medication administration, and sharp safety, in a formative and summative evaluation process, as per Standards of Practice College of Nurses. This was completed in a supervised setting with certified faculty from Collège Boréal and Cambrian College.		
	Pharmacy Technicians who have completed an approved injection course through the College of Pharmacists and who are working with a regulated health professional who can obtain informed consent and provide patient education may perform the act of injection under this medical directive.		
CLINICAL	Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 Vaccine) for the prevention		
INDICATIONS/	of coronavirus disease 2019 (COVID-19) caused by severe acute respiratory		
PURPOSE	syndrome coronavirus 2 (SARS-CoV-2) in individuals 18 years of age and older in whom contraindications are not present.		
SITUATIONAL	Informed consent		
CONDITIONS	Absence of contraindication(s)		
	• The use of Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 Vaccine) is		
	permitted under a Health Canada interim authorization delivered in		
	accordance with section 5 of the COVID-19 Interim order (IO). The interim		
	order is available <u>here</u> . The product monograph is available <u>here</u> .		
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CONTRAINDICATIONS	Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 Vaccine) is contraindicated for use by implementers authorized under this medical directive for the following individuals ^{2,3} :				
	1. Individuals with a history of anaphylaxis after previous administration of the vaccine.				
	2. Individuals who have ever had an anaphylactic reaction to any component of				
	an mRNA vaccine or its packaging. Refer to the vaccine component section for				
	product specific component information.				
	3. Individuals who are under the age of 18 years .				
PRESCRIBED	Moderna COVID-19 Vaccine is permitted for the following individuals under the				
CIRCUMSTANCES	prescribed circumstances described below:				
	1. Individuals who have had an allergic reaction within 4 hours of receiving a previous				
	dose of an mRNA COVID-19, or any components of the mRNA COVID-19 vaccine				
	(including polyethylene glycol [PEG]), tromethamine, or polysorbate, only if they				
	have been evaluated by an allergist/immunologist and it is determined that the				
	person can safely receive the vaccine, and the individual provides documentation as				
	required ⁵ (or as current reference)				
	2. Individuals who are breastfeeding or pregnant should be encouraged to be				
	vaccinated following the same recommendations as the general population. These				
	individuals should be informed of the latest evidence on the safety of mRNA COVID				
	19 vaccines in order to make informed decisions. ⁸				
	3. Individuals with autoimmune conditions and immunocompromised persons				
	should follow the same recommendations for COVID vaccination as for the				
	general adult population. They should be informed of the latest evidence on				
	the safety of mRNA COVID 19 vaccines in order to make decisions. Individuals				
	who are immunosuppressed from disease or treatment should be informed				
	that they may have a reduced immune response to any authorized COVID 19				
	vaccine series. ⁸				
	4. As a precautionary measure, the second dose in the mRNA COVID-19				
	vaccination series should be deferred in individuals who experience myocarditis				
	or pericarditis following the first dose of an mRNA COVID-19 vaccine until more				
	information is available. Individuals that have a history of myocarditis or				
	pericarditis should consult with their primary care provider prior to				
	immunization and be approved. ^(1, 2)				
WARNINGS/	Caution is advised in the administration of intramuscular injections in people with				
PRECAUTIONS	bleeding disorders. Refer to the Injection Techniques Certification Module for				
TRECAUTIONS	further information.				
	Cardiovascular: Myocarditis and Poricarditis				
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	Rare cases of myocarditis and/or pericarditis following vaccination with mRNA COVID-19				
	vaccine have been reported in Canada and internationally. These cases occurred most				
	frequently in adolescents and younger adults under 30 years of age, more frequently in				
	males compared to females, and more commonly after the second dose. When deciding				
	whether to administer an mRNA vaccine to an individual with a history of myocarditis or				
	pericarditis, consider the individual's clinical circumstances. Individuals who experience				
	any of the following symptoms within several days of vaccination should seek medical				

attention immediately: chest pain, shortness of breath, feelings of a fast-beating, fluttering or pounding heart. ^{1,2}

Allergic Reactions

For individuals who have had an allergic reaction within 4 hours and/or anaphylaxis that occurred with a vaccine or injectable medication that does not contain a component or cross-reacting component of the mRNA COVID-19 vaccines, the mRNA COVID-19 vaccine can be given with an extended observation post-vaccination of 30 minutes in the clinic.

Persons with allergy issues like allergic rhinitis, asthma and eczema can receive the vaccine with an extended observation post-vaccination of 15-30 minutes in the clinic.

Adverse Reactions

The most commonly reported adverse drug reactions after administration of Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 Vaccine) are injection site pain, fatigue, headache, muscle pain and stiffness, chills, nausea or vomiting, joint pain and fever. Uncommon reactions include swollen lymph nodes. Reactions are generally mild or moderate in intensity and of limited duration. Some adverse events, including fever, are more frequent after the second dose of vaccine.^{1,2}

Drug: Drug Interactions

Vaccines

COVID-19 vaccines should not be given simultaneously with other live or inactivated vaccines at this time, unless other vaccines are required for post-exposure prophylaxis.²

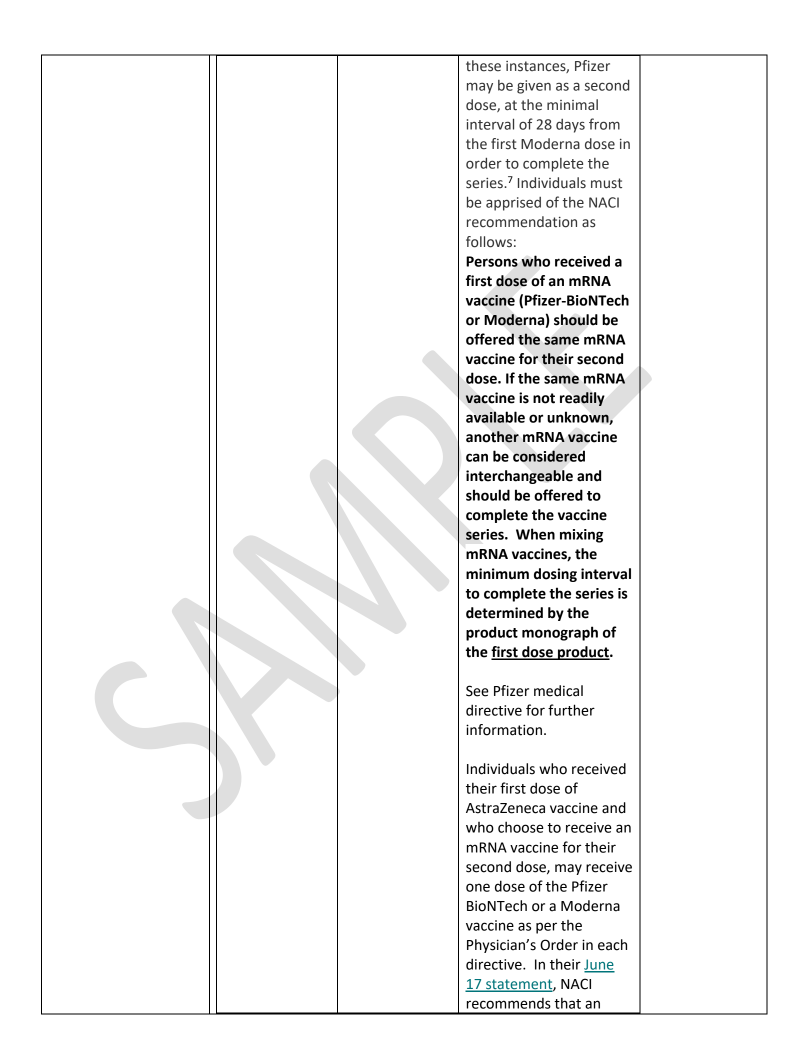
In the absence of evidence, it would be prudent to wait for a period of at least **28 days after each vaccine dose of an mRNA COVID-19 vaccine** before the administration of another vaccine (except in the case where another vaccine is required for post-exposure prophylaxis) due to the elicitation of an inflammatory cytokine response.²

It would be prudent to wait for a period of at least **14 days after the** administration of another vaccine before administrating a COVID-19 vaccine.²

Blood Products and Human Immunoglobulin

COVID-19 vaccines should not be given simultaneously with monoclonal antibodies or convalescent plasma.² In the post-exposure setting, expert clinical opinion should be sought on a case-by-case basis when deciding whether anti-SARS-CoV-2 monoclonal antibodies would be appropriate to administer after receipt of COVID-

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	19 vaccine, taking into consideration the risk of exposure and the risk of severe COVID-19 disease in the individual. ²					
	To date, there is also insufficient evidence on the receipt of both a COVID-19 vaccine and any monoclonal antibodies or convalescent plasma for treatment or					
	prevention of non-COVID-19 disease. Therefore, timing of administration and					
	potential interference between these two products are currently unknown and					
	expert clinical opinion should be sought on a case-by-case basis. ²					
	Oral Analgesics and Antipyretics					
	NACI recommends that prophylactic oral analgesics or antipyretics (e.g.,					
	acetaminophen or ibuprofen) should not be routinely used before or at the time					
	of vaccination, but their use is not a contraindication to vaccination. Oral analgesics					
	or antipyretics may be considered for the management of adverse events (e.g.,					
	pain or fever, respectively), if they occur after vaccination. ²					
	-		d in clinical trials of COVID-1			
			er vaccination. There is curr			
			f oral analgesics for the prev	ention of		
	immunization injed	ction pain or syster	nic reactions. ²			
	Drug: Food Interac	rtions				
	Drug. i obu intera					
	None listed					
PHYSICIAN'S ORDER			273 SARS-CoV-2 Vaccine) in	accordance with		
	the following table:					
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	Age	First dose	Second dose			
	Age 18 years of age		0.5 mL IM no sooner than			
	Age	First dose	0.5 mL IM no sooner than 28 days after the			
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	mRNA vaccine is now		
	preferred as the second		
	dose for individuals who		
	received their first dose		
	of		
	AstraZeneca/COVISHIELD		
	vaccine. ⁽¹¹⁾ This second		
	dose is given at an		
	interval between 8 to 12		
	weeks or more following		
	the AstraZeneca		
	vaccine. ⁹ If eligible for an		
	exception to the		
	extension of the interval,		
	the individual should		
	consult with their health		
	care provider to		
	determine the		
	recommended interval		
	between four and 12		
	weeks. Upon receiving		
	proof of this		
	recommendation, this		
	medical directive		
	provides authority to the		
	immunizer to administer		
	vaccine at the prescribed		
	interval if it is between 4		
	to 12 weeks following		
	the first dose dose. ⁷		
	*If administration of the second dose is delayed it should be given as soon as		
	possible. Every effort should be made to vaccinate with the second dose as		
	outlined above.		
ADDITIONAL DOSES	It may be possible to withdraw an additional 0.5 ml dose of vaccine i.e. an 11 th (5ml		
FROM VACCINE VIALS	vial) or 15 th dose(8ml vial). ¹⁰		
(6)			
	As an interim measure, an additional dose of COVID-19 vaccine may be extracted		
	from up to 3 vials of the same vaccine using aseptic technique as follows:		
	 Prepare vaccine in a clean, designated medication area away from where 		
	vaccination is occurring.		
	• Ensure that all of the vaccine vials accessed to extract an additional dose of		
	vaccine are from the same vaccine lot.		
	Combine vaccine from vials with residual volume only (i.e. not full vials) and		
	do not save up vials until the end of clinic before combining for extra dose.		
	• The different vials accessed have been under the same vaccine storage and		
	handling conditions – do not combine vials that have been thawed and		
	stored at $+2^{c}$ to $+8^{c}$ with those that have just been removed from a freezer.		
			

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Vials cannot be placed into a refrigerator beyond the permitted 24 hours post
puncture in order to have enough vaccine to make up a full extra dose.
This vaccine can be stored in refrigerator or frozen (needs to be thawed before
use), however DO NOT store on dry ice.
Refrigerator
Vials can be stored refrigerated between 2-8 degrees Celsius for up to 30 days prior
to first use.
Once the vial has been entered (needle-punctured), it can be stored at room
temperature or refrigerated (between $+2^{\circ}C$ to $+25^{\circ}C$), but must be discarded after
24 hours. Do not refreeze. ¹ Remember to time and date when vial is first
punctured.
The data in the surface should be used as seen as fare; bla and no later then 24
The dose in the syringe should be used as soon as feasible and no later than 24
hours after the vial was first entered (needle-punctured). ¹
Freezer
Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 vaccine) is stored frozen
between -25 degrees to -15 degree Celsius and should be stored in the original
carton to protect from light.
Vials must be thawed before use by removing the required number of vial(s) from
storage and thaw in the refrigerated conditions between 2 degrees to 8 degrees
Celsius for 2 hours and 30 minutes. Then, let each vial stand at room temperature
for 15 minutes before administering.
Vials alternatively can be thawed at room temperature between 15 degrees to 25
degrees for 1 hour.
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After thawing, DO NOT refreeze. This vaccine is preservative-free.
Any unused vaccine should be disposed of in accordance with local requirements.
While not recommended as routine practice, in exceptional circumstances a single
dose of Moderna vaccine may be transported in a syringe whilst careful attention is
taken to adhere to the parameters as outlined in the following documents
referenced below. ¹² Available data support transportation of one or more thawed
vials in liquid state for up to 12 hours at 2° to 8°C (36° to 46°F) when shipped using
shipping containers which have been qualified to maintain 2° to 8°C (36° to 46°F)
and under routine road and air transport conditions with shaking and vibration
minimized . ⁽¹²⁾
Drawn up vaccine must be administered within 24 hours from the time the vial was
first punctured. ⁽¹¹⁾
This process will only be enacted for exceptional circumstances only with the
approval and support of the Medical Officer of Health.

VACCINE	Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 Vaccine) presents as a white		
PRESENTATION	to off-white frozen suspension for intramuscular injection. It may contain white or		
	translucent product –related particulates. Inspect the vials visually for foreign		
	particulate matter and/or discoloration prior to administration. If either of these		
	conditions exists, the vaccine should not be administered.		
VACCINE	Moderna COVID-19 Vaccine (mRNA-1273 SARS-CoV-2 vaccine) contains a lipid		
COMPONENTS	nanoparticle (LNP) comprised of messenger Ribonucleic acid (mRNA) encoding the		
	viral spike glycoprotein (S) of SARS-CoV-2 and four lipids. Moderna COVID-19		
	vaccine does not contain any preservatives, antibiotics, adjuvants or human or		
	animal derived materials.		
	Non-medicinal ingredients:		
	 1,2-distearoyl-sn-glycero-3-phosphocholine (DSPC) 		
	Acetic acid		
	Cholesterol		
	Lipid SM-102		
	PEG2000 DMG 1,2-dimyristoyl-racglycerol,		
	methoxy-polyethylene glycol*		
	Sodium acetate		
	Sucrose		
	Tromethamine		
	Tromethamine hydrochloride		
	Water for injection		
	• Water for injection		
	Madarna COVID 10 vaccing is supplied in a multi dass 100 type 1 glass vial (sither		
	Moderna COVID-19 vaccine is supplied in a multi-dose 10R type 1 glass vial (either		
	5ml or 8ml) with a 20mm FluroTec [®] coated chlorobutyl elastomer stopper, 20mm		
	flip off aluminum seal. The 5ml vials contain 10 (maximum 11 doses) while the 8 n		
	vials each contains 14 doses (maximum 15 doses).		
	*Polyethylene glycol (PEG) is found in bowl preparation products for colonoscopy,		
	laxatives, cough syrup, cosmetics, skin care products and some food and drinks,		
	however this list is not exhaustive.		
	The vial stopper does not contain natural rubber latex.		
	Vials are packaged in a secondary carton containing a total of ten mRNA-1273		
	vaccine vials per carton.		
EXPIRY DATE AND LOT	The expiration date is not printed on the USA cartons or vials. This will be available		
NUMBER	on Moderna's Canadian Website: <u>https://www.modernacovid19global.com/ca</u> . ¹⁰		
	A list of expiry dates and lot numbers for USA products can be found in Moderna		
	lots expiry.		
	The Lead PHN shall check each lot number utilized prior to preparation and		
	distribution at each shift and clinic. For US products they will check the website		
	linked above for the expiration date and for other producers they will find the		
	expiration date on the box or vials.		

REFERENCES	1. Moderna	Therapeutics Inc. COVID	19 vaccine Moderna COVID-19 Vaccine		
	xp(mRN/	A-1273 SARS-CoV-2 vaccine	e) Product Monograph. June 30, 2021		
	2. National	Advisory Committee on In	nmunization (NACI): Recommendations on		
	the Use	of COVID-19 Vaccine(s). Ma	arch 1, 2021; July 2, 2021		
			Moderna Second Dose Interval & Firefighter		
	• •	Email dated February 23,			
		<i>,</i>	CAG). Recommendations on Exceptions to		
			-19 Vaccines. May 25, 2021 (or as current).		
	5. COVID 19 2021.	9 Vaccination Recommend	ations for Special Populations. March 11,		
		,	nmunization (NACI). An Advisory Committee Committee on Immunization (NACI)- NACI		
	Rapid Re 2021.	sponse: Interchangeability	of Authorized COVID-19 Vaccines. June 1,		
			e of COVID-19 Vaccines-update on Special ervals. Media Lines. May 28, 2021.		
	8. Ministry of Health. June 12, 2021. Ontario Accelerates Second Doses of AstraZeneca COVID-19 Vaccine: Second Dose of mRNA or AstraZeneca can be Administered at an Eight Week Interval with Informed Consent.				
	9. Moderna COVID-19 Vaccine US Supply Communication June 14 2021				
	10. Summary of National Advisory Committee on Immunization statement June 17 2021.				
	-	9 Handling and Storage Gu	idance. Version 6.1. June 24 2021.		
SIGNATURE AND DATE					
	Signature:	Original Signed By	Date: July 3, 2021		
			R: July 2021		