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**THIS CHAPTER MEETS THE FOLLOWING IMMUNIZATION COMPETENCIES FOR HEALTH
PROFESSIONAL (PHAC, 2008): [HTTP://WWW.PHAC-ASPC.GC.CA/IM/PDF/ICHP-CIPS-ENG.PDF](http://www.phac-aspc.gc.ca/im/pdf/ichp-cips-eng.pdf)**

#8: Administration of Immunizing Agents

- ◆ Competency: Prepares and administers immunization agents correctly.

1.0 PREPARATION FOR ADMINISTRATION OF BIOLOGICAL PRODUCTS

1.1 Client Health Assessment

Each time the client presents for immunization, screen them to ensure that they are well and can safely receive the recommended vaccines based on their **HALO** assessment (**H**ealth, **A**ge, **L**ifestyle, **O**ccupation). Refer to [Chapter 6, Contraindications and Precautions](#) for additional considerations prior to immunization.

1.1.1 General Screening Questions

1. Is your child/are you feeling sick today? Does your child/do you have diarrhea, vomiting or a high fever today?
2. Does your child/do you have allergies to medications, vaccine components, latex or foods?
3. Has your child/have you ever had a life-threatening allergic (anaphylactic) reaction after receiving a vaccine in the past?
4. Has your child/have you had any reactions to vaccines that you were concerned about?
5. Has your child/have you received any vaccines from other providers such as a family physician, Nurse Practitioner, pharmacist or from a travel clinic?
6. Was a parent of this child born outside of Canada?
7. Has your child/have you received any vaccines or a TB test in the past 4 weeks?
8. In the past year, has your child/have you received any blood products or a transfusion, immune globulins (antibodies) or antiviral drugs?
9. Has your infant had an episode of intussusception? Does your infant have an uncorrected congenital gastrointestinal malformation (e.g. Meckel's diverticulum)?
10. Is there a history of severe combined immunodeficiency (SCID) or a history of recurrent, unexplained early deaths in the family?
11. Does your child/do you have any diagnosed medical conditions such as:
 - a. Kidney, liver (hepatitis A, B or C), heart or lung diseases (e.g., asthma)?
 - b. An abnormal or absent spleen?
 - c. Bone marrow problems or a blood disorders (e.g. sickle cell disease or anemia)?
 - d. A bleeding disorder or are on long-term aspirin therapy?
 - e. Any metabolic diseases (e.g., diabetes mellitus)?
 - f. A chronic cerebrospinal fluid leak or a shunt for hydrocephaly?
 - g. Encephalopathy (e.g., coma, decreased level of consciousness, prolonged seizures) not attributable to another identifiable cause within 7 days after receiving a dose of a pertussis-containing vaccine?
 - h. Uncontrolled seizures, progressive encephalopathy or other progressive or neurological disorder that is not stabilized with treatment?
12. Does your child/do you have any form of immune system problems or suppression related to cancer, leukemia, lymphoma, HIV infection or AIDS, or a congenital immune system problem (e.g. severe combined immunodeficiency disorder [SCID], B-lymphocyte [humoral] immunity, T-lymphocyte [cell-mediated] immunity, complement system [properdin, or factor D deficiencies] or phagocytic functions)?
13. Is your child/are you currently taking cortisone, prednisone, and other corticosteroids or immune-suppressing drugs, receiving anticancer drugs (chemotherapy) or having radiation (X-ray) treatments?
 - a. Examples of immune suppressing drugs may include anti-rheumatic drugs, and drugs used for the management of inflammatory bowel disease.
14. Has the mother taken any monoclonal antibody medications during her pregnancy with this child? (Refer to [Appendix 8.2: Monoclonal Antibody Medications](#) for list).
15. Has your child/have you ever had a cochlear, solid organ, islet cell or stem cell transplant, or are recommended to have a transplant in the future?
16. Has your child/have you ever had a nervous system disorder (e.g. Guillain-Barré syndrome)?
17. Is your child/are you pregnant or is there chance she/you could become pregnant during the next month?
18. Has your child/have you ever had a positive TB skin test?
19. Any episode of transient or idiopathic thrombocytopenia following a vaccine?

1.2 Standard Precautions

- Biological products, syringes and needles are considered sterile once manufactured. Therefore, aseptic technique must be practiced when preparing these products.
- Wash hands well using soap and water, or a waterless hand cleanser prior to preparing biological products and between clients.
- The use of non-sterile or sterile gloves during immunization is not recommended and unnecessary unless the immunizer is concerned about contact with a client's body fluids or the immunizer's skin on their hands is not intact. Gloves must be changed between clients.
- Use a separate, sterile safety engineered sharp device (SESSED) for each injection.
- Engage the safety mechanism on needles immediately following administration of the biological product.
- Immediately discard the syringe/needle into a sharps container that is placed safely away from clients (and children).
- Never recap needles or empty used needles/syringes from one sharps container to another.
- Immediately report needle stick injuries to a supervisor for consideration of possible post-exposure immunoprophylaxis. Follow worksite occupational health and safety protocol.

1.3 Product Preparation

If the protective cap on a single-dose vial is removed, or if a manufacturer's pre-filled syringe is opened (e.g., syringe cap removed), the vaccine should be used on that clinic day or discarded CIG).

1.3.1 Pre-Preparation: Pre-Loading of Syringes

Pre-preparation of prefilled syringes and pre-loading of syringes with biological products that come in vial or ampoule presentations is discouraged because of the uncertainty of product stability in syringes, risk of contamination, increased potential for administration errors, and biological product wastage.

1.3.1.1 Filter Needles

Filtration needles are not recommended for vaccine preparation or administration as they may filter out active ingredients such as adjuvants (CIG). In 2010, the Canadian Agency for Drugs and Technologies in Health (CADTH) published a review titled *Filtered Needles for Withdrawing Medication from Glass Ampoules: A Review of the Cost-Effectiveness and Incidence of Complications*. Regarding the use of filtered needles in public health nursing practice, the Ministry of Health endorses the following conclusion statement in this report: "Due to the lack of recent published literature, no conclusions can be drawn on the incidence of complications from glass particle administration, or the cost-effectiveness of using a filtered needle when withdrawing medication from a glass ampoule". This includes diluents in ampoule presentations as well.

1.3.1.2 Combination of Contents of Multi-Dose Vials

Withdrawing contents from multiple multidose vials to make a vaccine dose and prevent wastage is not recommended by the Ministry of Health because of potential contamination risks and lot number discrepancies.

1.3.2 Preparation Instructions

1. When preparing any biological product, consider the “10 Rights” of immunization:
 1. Right client
 2. Right assessment
 3. Right client education
 4. Right to refuse
 5. Right product
 6. Right dose
 7. Right route
 8. Right time
 9. Right documentation
 10. Right evaluation
2. Thoroughly wash hands with soap and water or cleanse with a sanitizer.
3. Prepare necessary materials (e.g., single use, disposable sterile syringe/needle, vaccine, diluent if required, 70% isopropyl alcohol, sharps container, supplies for the management of anaphylaxis).
4. Different vaccines must never be combined in the same syringe.
5. Check the characteristics of the product to be administered:
 - Correct product, form of presentation and expiry date.
 - Check three times that it is the correct product: when removing from fridge/biological cooler, when drawing up and/or reconstituting, and prior to administration.
 - Expected appearance: are there any irregularities (e.g., particulate matter, damage)?
 - Expiry date. If only the month and year are provided for the expiry date, the biological product can be used to the end of that month.
6. If a previously opened multi-dose vial is available, check the date that the vial was opened (as recorded on the label). Most multi-dose vials must be used within 30 days of opening, unless the manufacturer specifies another period (i.e., once punctured, some MDVs are stable to the expiry date noted on the vial).
7. **If there is discoloration, extraneous particulate matter, or obvious lack of re-suspension**, mark the product as “DO NOT USE,” return it to proper storage conditions, complete a [Vaccine Problem Supply report form](#) and fax or email the form with a picture of the product to the Ministry of Health as directed in [SIM Ch. 9](#).

1.3.3 Vials

1. Wash hands or cleanse with a sanitizer.
2. Remove the plastic cap covering the vial.
3. Cleanse the surface of the rubber stopper using a cotton pad/swab moistened with 70% isopropyl alcohol. Allow to air dry.
4. Gently swirl the vial immediately before removing each dose to ensure that the contents are fully dispersed.
 - For a product in a “ready to go” liquid presentation, draw into the syringe a volume of air equal to the quantity of biological product to be removed.
 - For lyophilized, or freeze-dried vaccines requiring reconstitution, the diluent acts as the air in the syringe so there is no need to draw air into the diluent syringe.
5. Hold/place the vial right side up and insert the needle through the centre of the rubber stopper. **Do not insert blunt needles with or without a filter into vials because of coring risk.**
6. Slowly inject the air or diluent from the syringe.
7. If the biological product was reconstituted, gently swirl the vial to ensure the contents are fully dispersed.
 - Single dose vial - withdraw all contents to ensure client receives full concentration of antigens.
 - Multidose vial - withdraw the required quantity of biological product into the syringe.

8. Remove the needle from the vial and expel any air bubbles from the syringe.
9. It is not necessary to change needles between drawing up the biological product into the syringe and immunizing the client. Change the needle only if it is damaged or becomes contaminated.
10. Discard the empty vial into a sharps container.
11. If it is the first entry into a multi-dose vial, record the date (include day, month and year) on the label of the vial.
12. Immediately return multi-dose vials to the refrigerator/biological cooler.

1.3.4 Vaccines with Diluents

1. Before reconstituting, check labels on both the vaccine vial and the diluent vial to verify:
 - That they are the correct products to mix together.
 - That both are within their expiration dates.
2. Reconstitute vaccines just prior to use by:
 - Removing the protective caps and wiping each stopper with an alcohol swab;
 - Inserting needle of syringe into diluent vial and withdrawing entire contents. Do not insert blunt needles with or without a filter into vials because of coring risk;
 - Inject diluent into vaccine vial and gently agitating to thoroughly dissolve the lyophilized powder. Draw up all contents to ensure client receives full concentration of antigens.
 - If reconstituted vaccine is not used immediately or comes in a multi-dose vial:
 - Clearly mark the vial with the date and time that the vaccine was reconstituted.
 - Protect vaccines from light.
3. Changing the needle after the reconstitution is unnecessary unless the needle has been damaged or contaminated.
4. Check the colour and appearance of the reconstituted vaccine to ensure it matches the description on the package insert.

1.3.5 Ampoules

1. Gently swirl the ampoule immediately before removing the contents to ensure that the contents are fully dispersed.
2. Tap the ampoule lightly to ensure that the contents are in the lower part of the ampoule.
3. Using a swab moistened with isopropyl alcohol, wipe the neck area of the ampoule prior to opening to prevent bacterial contamination of ampoule contents.
4. Break the neck of the ampoule using the alcohol swab, cotton ball or cotton gauze. If you cut yourself in breaking the ampoule, discard the ampoule, since the product may be contaminated. Wash your hands and cover the cut before continuing.
5. Draw up all contents to ensure client receives full concentration of antigens using a sterile syringe and needle. It is not necessary to change needles between drawing up the biological product into the syringe and administering it to the client.
6. Discard the ampoule into a sharps container.
7. Expel the air bubbles from the syringe.

1.3.6 Multidose Vials

1. Refer to section 1.3.3 Vials for details.
2. Hold/place the vial right side up and insert the needle through the centre of the rubber stopper. Do not insert blunt needles with or without a filter into multidose vials because of coring risk.

1.3.7 Prefilled Syringe-Vaccine Vial Format

1. Do not remove any air that is present in the prefilled vaccine- or diluent-containing syringe prior to inserting it into the vaccine-containing vial. This air is required to ensure that all of the liquid is inserted into the vial to attain the correct volume of reconstituted vaccine. If no air is noted, a little amount of air should be withdrawn into the syringe prior to inserting its contents into the vial.
2. Ensure that the vial is sitting on the counter so that the stopper cap is facing upright. Remove the plastic cap and wipe the stopper with an alcohol pad and let dry. Hold the vial steady on the counter. Hold the syringe so that the needle is pointing down and the air bubble (point 1 above) is present under the syringe stopper. Insert the needle straight down into the centre of the vial stopper and then inject this vaccine into the vaccine vial. This will ensure that the air bubble in the syringe will clear all vaccine into the vial. There should be no remaining vaccine in the syringe or needle. (The question and answer below was recently published in the *Immunization Action Coalition* newsletter Volume 17 Issue 1 and is available at: <http://www.immunize.org/va/va38.pdf>.
 - **Q:** Some single-dose pre-loaded vaccines come with an air pocket in the syringe chamber. Do we need to expel the air pocket before vaccinating?
 - **A:** No. You do not need to get rid of the air pocket. The air will be absorbed. This is not true for syringes that you fill yourself; you should expel air bubbles from these syringes prior to vaccination to the extent that you can readily do so).
3. Remove the syringe and gently shake the vial contents for reconstitution. Draw into the syringe a small volume of air. With the vial sitting flat on the counter (stopper facing up), insert the air from the syringe into the vial; this added pressure ensures that all vial content can be withdrawn.
4. Pick up and invert the vial upside down. Pull the needle back until the tip is in the liquid. Pull back on the plunger until you remove the entire liquid contents of the vial; a bit of air from within the vial can also be withdrawn into the syringe to ensure the needle is cleared of vaccine.
5. With the syringe upright so that the needle is at the top, remove the needle from the vial and if there are air bubbles in the syringe, tap them gently so that they move under the needle and then gently push them out of the needle, being very careful not to expel any vaccine. This action primes the needle.
6. Recap the needle and the vaccine is ready to administer.

1.4 Scheduling and Administration of Multiple Injections

There are no contraindications to receiving multiple injections of vaccines at the same clinic visit. There is no increase in side effects or reduced vaccine effectiveness. [CIG](#) Guideline 3 states, “*Vaccine providers should use all clinical opportunities to screen for needed vaccines and to administer all vaccine doses for which a vaccine recipient is eligible at the time of each visit.*” Adherence to this standard of practice will avoid missed immunization opportunities and the possibility of susceptible individuals contracting vaccine-preventable diseases.

1.4.1 Practice Considerations

- When two or more injectable biological products are to be administered, they may be administered in the same limb (except for rabies vaccine and rabies immune globulin), providing the distance between injection sites is a minimum of 2.5 cm (1 inch,) apart so that local reactions can be distinguished for each product administered.
- Give sweet oral vaccines first and then biological products that are known to cause more stinging and/or pain last (e.g., give DTaP-IPV-Hib first, followed by MMRV and/or then pneumococcal conjugate vaccine).
- If multiple injections are to be given, and two health care providers are available, ask the client if they would prefer to have the biological products administered simultaneously in different limbs. The premise is that this procedure allows the client more control in the immunization experience and may decrease anxiety from anticipation of next injection(s).

1.5 Publicly Funded Immunizations Following Non-Conforming Situations

- Publicly funded vaccine doses administered any time before the eligible minimum age or eligible minimum interval as recommended in the [SIM](#) are considered medication administration errors.

4-Day Grace Period Principles

- The US Advisory Committee on Immunization Practices (ACIP) established the “4-day grace period” that may be applied when assessing the validity of historic documented vaccine doses that were administered up to four days before the minimum age or minimum interval.
- The 4-day grace period **may be applied by RHAs and FNJs** when reviewing **historical** client immunization records (such as childhood records for school entry).
- The 4-day grace period **should not** be applied by RHAs and FNJs:
 - To schedule future immunization visits for **publicly funded** vaccines.
 - To allow for the administration of publicly funded vaccines **at the discretion of the PHN** when they encounter a client before the minimum ages and/or intervals for a vaccine have been met.
- In certain situations, like accelerated schedules for travel or other situations such as during communicable disease investigations or when immunizing hard to reach populations, a regional **MHO may discretionarily** determine that doses of selected vaccines administered before the recommended minimum age or minimum intervals supersede the 4-day grace period and are valid doses. This is a regional decision **and** written regional policies (to address such instances should exist regarding the 4-day grace period (for publicly or non-publicly funded vaccines). Where a policy doesn’t exist, the MHO recommendation must be documented in Panorama.
 - PHNs must document in Panorama the rationale regarding the application of the 4-day grace period (outside of historical immunizations) in a client’s immunization record.
- The 4-day grace period does not apply to post-exposure rabies vaccine, or to accelerated vaccine schedules (e.g., HB, HAHB).**

A. Administration of a vaccine 4 or fewer days before the minimum age or minimum interval.

Inactivated vaccines

- Inactivated vaccine doses administered up to 4 days before the minimum age or minimum interval for a vaccine antigen(s) may be assessed as valid.

Live vaccines

- Live injectable and oral vaccine doses administered up to 4 days before the minimum age for a vaccine antigen(s) may be assessed **as valid**.
- Live injectable and oral vaccine doses administered up to 4 days before the minimum interval for a vaccine antigen(s) **are invalid** regardless of circumstances.
 - This applies whether the same or different injectable antigens have been administered (e.g., MMRV and MMRV, or MMR and Var). Invalid doses need to be repeated at the appropriate minimum interval from the invalid dose.
- NOTE: Because live oral and intranasal vaccines do not interfere with viral antibody production from other live (oral, intranasal or injectable) vaccines, they can be given regardless of the interval in relation to other vaccines.

B. Administration of a vaccine 5 or more days before the minimum age or minimum interval.

- Live or inactivated vaccine doses administered 5 days or more before the minimum age or minimum interval **are invalid** and need to be repeated at the appropriate minimum interval from the invalid dose.

1.5.1 Vaccines Given at Less than the Recommended Minimum Interval

- Refer to SIM, [Chapter 5, Section 2.1, Minimum Intervals between Vaccine Series Doses](#).
- Generally, a vaccine dose that was given 5 days or more before the minimum interval is an invalid dose and must be repeated at the correct minimum interval.
- If two live injectable vaccines are not given on the same day and are given at less than the recommended minimum intervals, the second vaccine given is invalid and must be repeated at the correct minimum interval from administration.

1.5.2 Vaccines Given at Less than the Recommended Minimum Age

- Refer to SIM, [Chapter 5, Section 2.1, Minimum Intervals between Vaccine Series Doses](#).
- Generally, a vaccine given at less than the acceptable minimum age is an invalid dose must be repeated once the client is of acceptable age. Refer to section 1.5 above.

1.5.3 Vaccines Given by Incorrect Route

- Refer to SIM, [Chapter 5, Section 4.3, Individuals Who Received a Vaccine by a Route Other than that Recommended](#) for information.

1.5.4 Reduced Doses of Vaccine

- Refer to SIM, [Chapter 5, Section 4.4, Individuals Who Received an Inappropriate Vaccine Dose for information](#).

1.5.5 Expired Vaccines

- A. If there is no urgency to repeat the expired dose and the client is agreeable**, the immunizer may contact the manufacturer's medical information department (do online search for contact information) to request if they have data to support the potency of the administered expired vaccine dose.
 - The inquirer **must** request to receive printed confirmation of these data from the manufacturer for inclusion/uploading in the client's medical/immunization record.
 - NOTE: If A is not feasible by the immunizer, refer to B.**
- B. If an expired live or non-live dose** was inadvertently given; and the client is not agreeable to having the manufacturer contacted; **or** the manufacturer does not have potency data as in **A**, or time is a factor, it is **an invalid dose and** should be repeated.

NOTE: Document a client's refusal for a repeat dose in their record.

- If the error **is detected on the same day** that administration occurred, repeat the dose that same day at a different injection site. The repeat dose is a valid dose.
- If the error **is not detected** on the same day:
 - For a **non-live vaccine**, a repeat dose should be given as soon as possible.
 - However, recombinant zoster vaccine (RZV; SHINGRIX™) should be administered 28 days after the invalid dose, to reduce the burden of adverse reactions which occurs with this vaccine.
 - For a **live vaccine**, a 28-day interval is required, because circulating interferon may interfere with the replication of the second live vaccine.
 - For rotavirus vaccine doses, the repeat dose should be administered after a 28-day interval from the invalid dose or at the maximum age for the vaccine dose (whichever is earlier).

2.0 ADMINISTRATION ROUTES, SITES AND TECHNIQUES

Providers must ensure that biological products are correctly administered using the recommended dose, route, site and schedule to optimize product effectiveness and reduce the risks of local reactions and adverse events. **Always read the product monograph thoroughly for comprehensive administration instructions.**

2.1 Special Considerations

- Most injectable live, attenuated and inactivated vaccines are to be administered by one specific route as stated in the product monograph. However, specific vaccine product monographs indicate two acceptable routes for administration.
- Some vaccines that are indicated only for intranasal or oral administration routes are packaged in devices that resemble injection syringes.
- Immune globulins must be administered in sites distanced from vaccine sites.

2.1.1 Limb Integrity

Injection of a vaccine into an area where lymphatic circulation may be impaired (e.g., local lymphedema, lymphangioma, axillary lymph node dissection, arteriovenous (A-V) fistula, upper limb amputation) could theoretically result in an impaired immune response due to impaired vaccine absorption, although there are no data to support this. Consider an alternative injection site if possible. There is no evidence or theoretical rationale for avoiding injection through a tattoo or superficial birthmark. Vastus lateralis may be used as an alternative site for all ages.

2.1.2 Persons with Bleeding Disorders

Haematoma formation and excessive bruising post IM injection may occur among clients with bleeding disorders or those that are on anticoagulant therapy. The MHO and/or client's physician is best positioned to assess an individual client's ability to safely receive IM injections. A fine gauge needle (23, 25, or 27 G) should be used. Z-track technique may be used to prevent bleeding. Do NOT aspiration. Apply direct pressure (without rubbing) to the injection site for 2 minutes or longer. The client or caregiver should receive information on the risk of developing and managing a haematoma.

2.2 Injection Guidelines

- Before administering any biological products, visually inspect the skin's surface over the injection site for bruises, scars, or inflammation. Sites with established tattoos are safe to use.
- Palpate the site as biological products should not be injected where there is poor muscle mass, and existing inflammation, itching, scars, nodules, sensitivity, induration, or pain.
- For those 4 years and older, moderately rub or stroke the injection site prior to injection.
- Cleanse the site for 5 seconds with isopropyl alcohol and allow for drying before administering the injection.
- Coach the client to relax the limb muscles prior to injection.
- Administer the least painful or stinging products first.
- For IM and SC injections, insert the needle quickly and firmly into the injection site, and stabilize the syringe to prevent it from moving.
- Do not aspirate the syringe for any injections. Aspiration is painful, may traumatize tissues, and affect absorption.
- Rapid injection of vaccines (not immunoglobulins or PPD) reduces pain during injection.

-
- Withdraw IM and SC needles quickly, at the same angle that it was inserted. Promptly activate the safety engineered sharps device.
 - All needles and sharps must be immediately disposed of as a single unit into a sharps container.
 - Apply gentle pressure with a cotton ball to the injection site for 10 seconds after the injection.
 - If the dose leaked out during administration, immediately administer another dose to ensure complete immunization status of the client.
 - Adhesive bandages are not recommended for use because they may:
 - Contain latex;
 - Irritate the injection site; or
 - Become a choking hazard for infants and young children.

2.3 Anatomical Guidelines and Sites

Longer needle lengths reduce the incidence of localized side effects. If the needle strikes bone during insertion, withdraw it slightly before injection of the product into the muscle. Use clinical judgment in selecting appropriate injection sites and needle sizes for IM injections based upon the:

- Client's age;
- Injection site muscle mass;
- Thickness of adipose tissue over the injection site;
- Volume of the product to be administered;
- Number of products to be administered; and
- Viscosity of the biological product.

Table 1: Immunization Route and Site, Needle Length and Gauge and Total Daily Site Volume by Age Group

Intradermal (ID)	5°-15° angle	Needle size & gauge	Site volume
All ages	- inner (volar) forearm - over the deltoid muscle - suprascapular area on the back - over the anterolateral thigh	1 cm (0.39 inches) 26-27 gauge	0.1 mL
Subcutaneous (SC)	45° angle	Needle size & gauge	Site volume
< 1 year	anterolateral thigh	1.6 cm (5/8 inch) 25 gauge	0.5 mL
≥ 1 year:	- upper triceps area - anterolateral thigh		0.5 mL
Intramuscular (IM)	90° angle	Needle length ¹ (22-25 gauge ²)	Site volume
Newborns (< 28 days) & preterm infants	Vastus lateralis	1.6 cm (5/8 inch)	1 mL
Infants (1 to 12 months)	Vastus lateralis	2.2 cm to 2.5 cm (7/8 inch to 1 inch)	1 mL
Young children (1 to 3 years)	Deltoid ³	1.6 cm to 2.5 cm (5/8 inch to 1 inch)	1 mL
	Vastus lateralis	2.5 cm to 3.2 cm (1 inch to 1 ¼ inch)	2 mL
Children (3 to 11 years)	Deltoid	1.6 cm to 2.5 cm (5/8 inch to 1 inch)	1 mL
	Vastus lateralis	2.5 cm to 3.2 cm (1 inch to 1 ¼ inch)	2 mL
Adolescents and adults (12+ years) (weight-based)	Deltoid	For those weighing < 130 lbs (< 60 kg) 1.6 cm to 2.5 cm (5/8 inch to 1 inch)	1 mL
		Males weighing 130 - 260 lbs (60 - 118 kg) Females weighing 130 - 200 lbs (60 - 90 kg) 2.5 cm (1 inch)	2 mL
		Males weighing more than 260lbs (118 kg) Females weighing more than 200lbs (90kg) 3.8 cm (1 ½ inch)	2 mL
Adolescents and adults (12+ years)	Vastus lateralis	2.5 cm to 3.8 cm (1 inch to 1 ½ inch)	3 mL

Adapted from [Vaccine Administration: A Guide to Selecting Needle Gauge and Length](#) (PHAC, 2024)

¹ A range of needle lengths are provided as clinical judgment should be used when selecting needle length for IM injections. Consideration should be given to vaccine recipient's weight, gender and age. **These recommendations are based on the practice of having the skin stretched flat (between thumb and forefinger) at the time of administration.**

- **NOTE:** Ensure staff are aware that 'bunching' a muscle prior to IM injection **is not an evidence-informed practice**; skin must be held flat and an appropriate needle size used for each client.

² A larger gauge needle (e.g., 22 gauge) may be required when administering viscous or larger volume products such as immune globulin.

³ The deltoid site is often selected for toddlers and young children because temporary muscle pain post-vaccination in the anterolateral thigh muscle may affect ambulation.

Table 2: Immune Globulin Preparation Injection Site, Needle Length and Total Daily Site Volume per Age Group

NOTE: Vital signs are not required to be taken before or after IM HBIg/Ig/RabIg/Tig/VarIg administration.

CLIENT AGE	SITE (90° IM) ²	NEEDLE LENGTH	SIZE (Gauge)	MAX. VOLUME ¹
Children				
• Birth to less than 12 months ⁴	Vastus lateralis	7/8" – 1"	25	1 mL
	Ventrogluteal (≥7 months)	7/8" – 1"	25	1 mL
• 12 months up to and including 4 years ⁴	Deltoid	1"	22-25	1 mL
	Vastus lateralis	1"	22-25	2 mL
	Ventrogluteal	1"	22-25	1 mL
• 5 years up to and including 17 years	Deltoid ³	1" – 1½"	22-25	1 mL
	Vastus lateralis	1" – 1½"	20-25	3 mL
	Ventrogluteal	1" – 1½"	20-25	3 mL
	Dorsogluteal ⁵	1" – 1½"	20-25	3 mL
Adults				
• 18 years and older	Deltoid ³	1" – 1½"	20-22	2 mL
	Vastus lateralis	1" – 1½"	20-22	5 mL
	Ventrogluteal	1" – 1½"	20-22	4 mL
	Dorsogluteal ⁵	1" – 1½"	20-22	5 mL

Adapted from BCCDC Communicable Disease Control Manual (2015) [Immune Globulin Preparations](#)

¹ The maximum volumes noted in the table are strictly guidelines. The immunizer must assess the adequacy of the individual's potential immunization sites to accommodate greater product volumes (e.g., [IM Ig administration in an infant](#)) prior to administration.

² IM immune globulin injections **must be** separated by minimum 2.5 cm if given in the same limb (e.g., Tig and RabIg in adult deltoid). **It is recommended to administer in different sites if possible.**

³ One deltoid should be reserved for the administration of rabies vaccine **on day 0** of rabies post-exposure immunoprophylaxis.

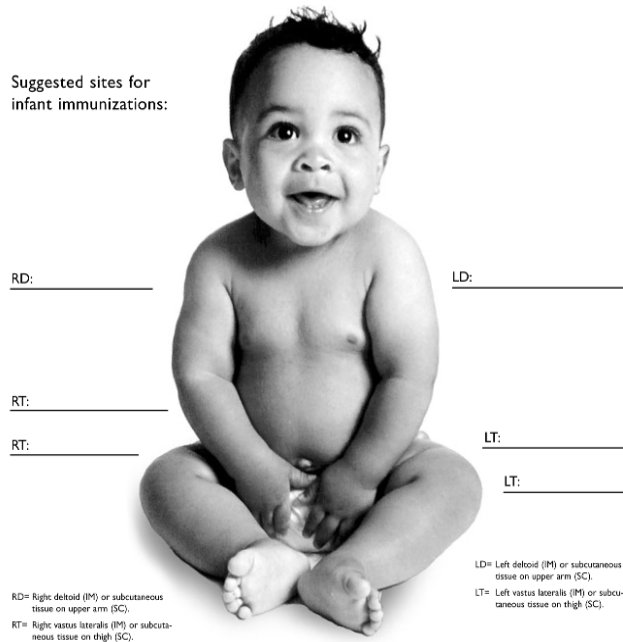
⁴ To facilitate administration of IMIg in children, injection volumes of up to 3 mL could be considered to reduce the number of injections, using clinical judgement. For high volume injections, the anterolateral thigh is generally preferred due to the greater muscle mass. Clinical judgement should be used when selecting the most appropriate site for IMIg administration ([NACI 2025](#)).

⁵ Use of the dorsogluteal site is **only recommended in adolescents and adults** when the ventrogluteal and vastus lateralis sites have had maximum volumes of an immune globulin preparation injected and an additional volume still needs to be administered. This is due to the possibility of sciatic nerve injuries when the injection is done in the dorsogluteal site.

Infant and Child Immunization Site Maps

Immunization Site Map

Suggested sites for
infant immunizations:



IMMUNIZATION
TECHNIQUES
Safe • Effective • Caring

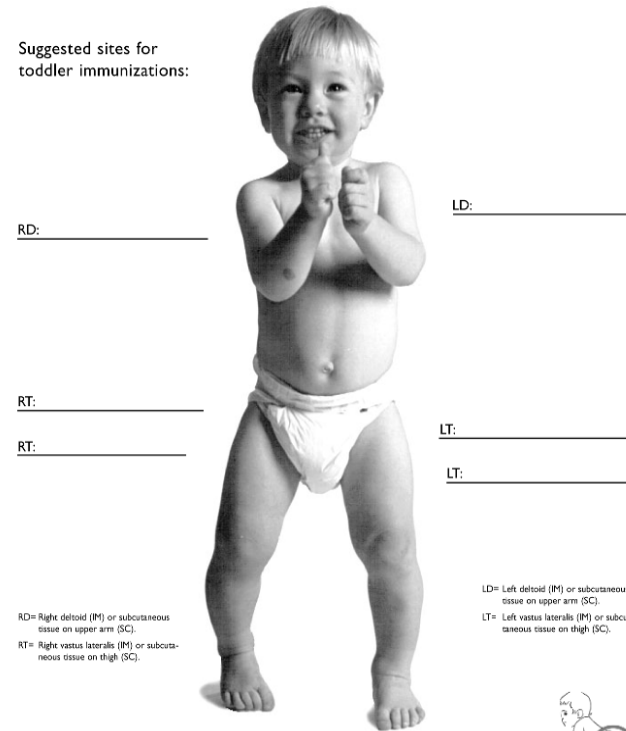
California Department of Health Services • Immunization Branch • 2151 Berkeley Way • Berkeley, CA 94704



IMM-718 (5/01)

Immunization Site Map

Suggested sites for
toddler immunizations:



IMMUNIZATION
TECHNIQUES
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California Department of Health Services • Immunization Branch • 2151 Berkeley Way • Berkeley, CA 94704

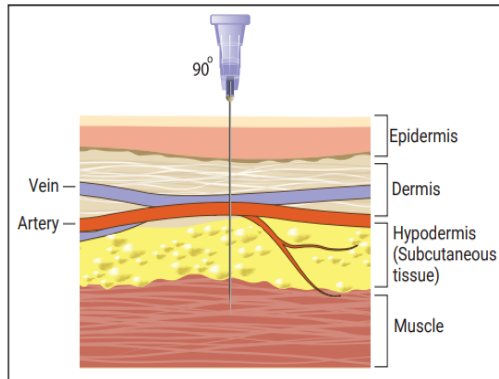


IMM-718 (5/01)

(Source retrieved April 2, 2011 from: <http://www.immunize.org/clinic/administering-vaccines>)

2.4 Intramuscular (IM)

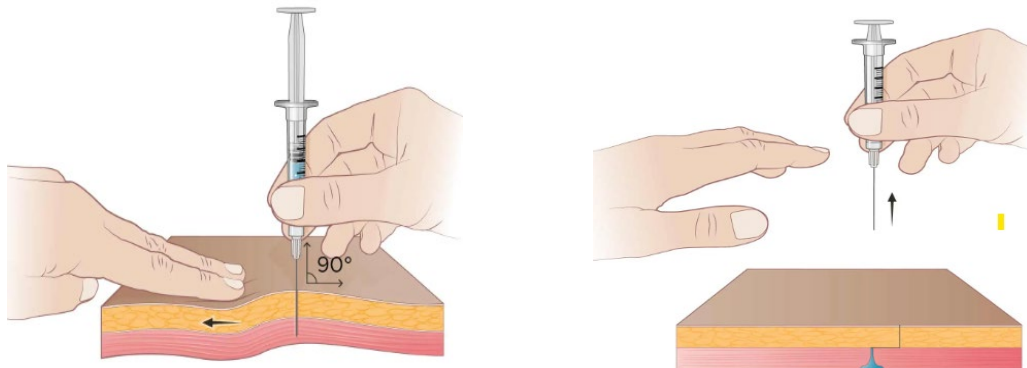
The IM route is used to deposit a biological product deep into a muscle at a 90° angle, where it can elicit the best immune response. Assess the vaccine recipient's weight, gender and age in selecting needle length. If such a product is erroneously injected into the dermis or adipose tissue, irritation, induration, inflammation, or an abscess may form at the injection site. **Do not aspirate before injection.**



Source: [Vaccine Administration: A Guide to Landmarking](#)

Two techniques are acceptable to administer IM injections:

1. Use the **thumb and index finger to gently stretch the skin FLAT over** the site while inserting the needle at a 90° angle to the skin.
2. Use the [Z-track](#) method only for vastus lateralis and ventrogluteal sites. Z-tracking involves using the free hand to displace the skin and adipose tissue at least one inch laterally in relation to the underlying muscle, prior to 90° insertion of the needle into the muscle. The Z-tracking must be kept in place until after the biological product is deposited into muscle and the needle is withdrawn, therefore preventing leaking of the medication into the subcutaneous tissue (see below). This technique seals off the puncture tract and traps the product within the muscle, minimising pain, irritation and bleeding at the site.



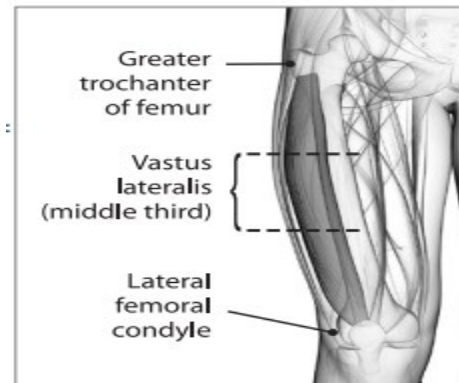
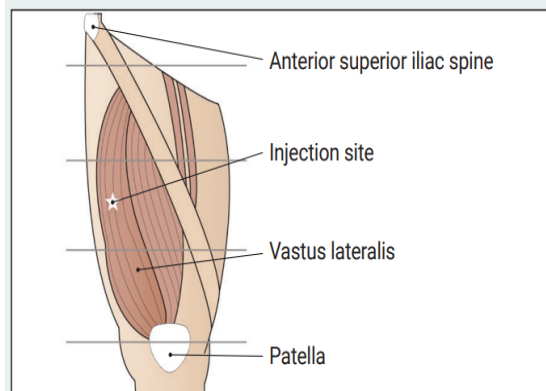
Source: <https://www.healthline.com/health/z-track-injection#how-to>

2.4.1 Vastus lateralis

This is the preferred sites for IM injections in infants less than 12 months of age because they are well developed at birth and are safely distanced from any nerves or major blood vessels. Using these sites can temporarily affect the movement of the child's leg.

To landmark the site:

1. Visually divide the length of the muscle that originates on the greater trochanter of the femur and the lateral border of the kneecap into thirds.
2. The anterolateral aspect of the middle third is where the belly of the vastus lateralis muscle lies, and the injection site is in the middle of this area.
3. In width, the site extends from the midline of the top of the thigh to the midline of the outer side of the thigh.



Source: [Vaccine Administration: A Guide to Landmarking](#)

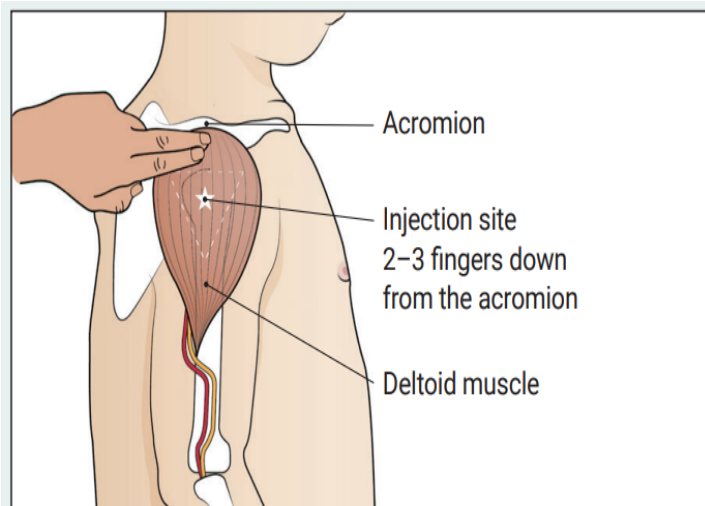
2.4.2 Deltoid

The deltoid muscles are the preferred IM injection sites for adults and children 12 months of age or older. The deltoid muscle is insufficiently developed in younger infants.

To landmark the site:

1. Visualize an inverted triangle with its base at the acromion process and its peak just below the axilla.
2. The injection site is in the center of the triangle.
3. Refer to [Appendix 8.1](#) for illustrations to administer multiple IM injections in one arm for adults

NOTE: Accurate landmarking is very important to prevent a **Shoulder Injury Related to Vaccine Administration (SIRVA)** that can result in damage to tissues and structure in the shoulder area and joint.



Source: [Vaccine Administration: A Guide to Landmarking](#)

2.4.3 Ventrogluteal (For Immunoglobulin Administration Only)

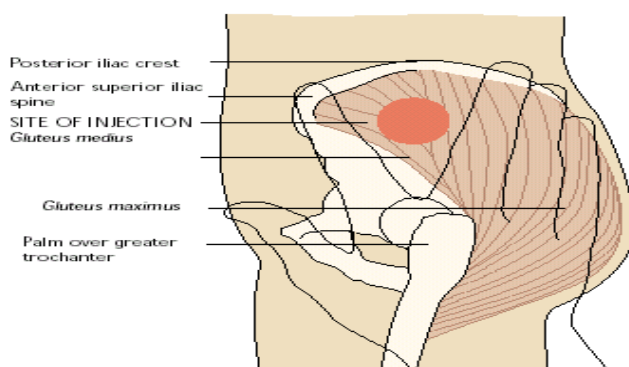
Do not use this site to administer active immunizing agents

The ventrogluteal site accommodates large volumes of immune globulin preparations (i.e., Ig, HBIG, RabIG, TIG, VarIG) by IM injection. Appropriate site selection of the gluteal muscle is necessary to avoid injury to the sciatic nerve. This site can be used in those over 7 months of age.

This muscle is accessible in the supine, prone, and side lying position.

To landmark this site in **children and adults**:

1. Use the right hand to locate the site on the left hip, and the left hand to locate the site on the right hip.
2. Place heel of the hand over the greater trochanter of the client's hip with wrist almost perpendicular to the femur.
3. Point the thumb toward the client's groin and the fingers toward the client's head.
4. Point index finger to the anterior superior iliac spine and extend the middle finger back along the iliac crest toward the buttock. The index finger, the middle finger, and the iliac crest form a V-shaped triangle.
5. The injection site is the centre of the triangle as noted below.



Source: http://www.bccdc.ca/resource-gallery/Documents/Guidelines%20and%20Forms/Guidelines%20and%20Manuals/Epid/CD%20Manual/Chapter%202%20-%20Imms/Appendix_B_Administration.pdf

To landmark this site in **infants 7 months and older**:

1. Lay the infant down over a carer's lap, so the legs are down at about a 45-degree angle. The person restraining the child is holding the infant's feet together with the right hand, and the left hand is positioned on the infant's bottom, with their hand placed to indicate the injection site.
2. The tip of the middle finger is on the iliac crest.
3. The tip of the index finger is on the anterior superior iliac spine.
4. The tip of the thumb is held against the index finger and placed on the greater trochanter.
5. The injection site is in the middle of the triangle formed by the index and middle fingers spread apart.

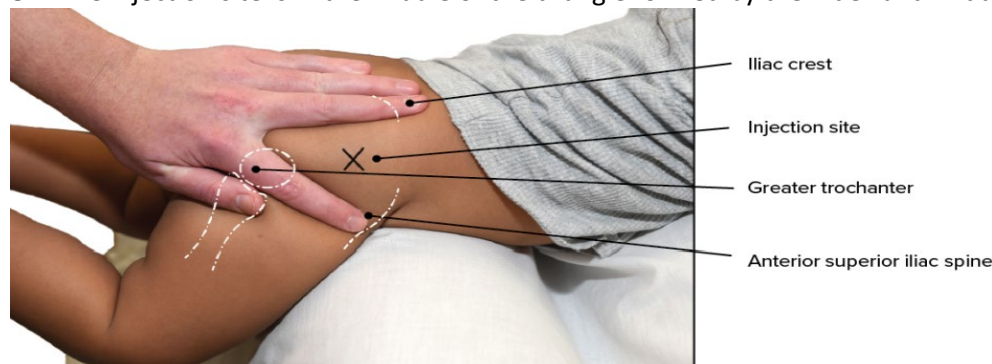


Figure source: [Australian Immunization Handbook](#)

2.4.4 Dorsogluteal (For Immunoglobulin Administration Only)

Do not use this site for administering active immunizing agents into the gluteal muscle.

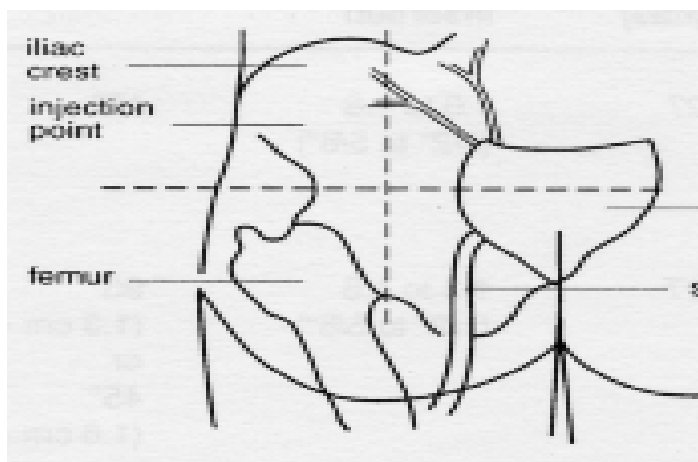
The dorsogluteal site is only to be used for the IM injection of large volumes of immune globulin preparations when the ventrogluteal and vastus lateralis sites have had maximum volumes of an immune globulin preparation injected and an additional volume still needs to be administered. This is due to the possibility of sciatic nerve injuries when the injection is done in the dorsogluteal site.

This site should only be used in individuals over 5 years of age.

Place client in a prone, side lying, or standing position.

To landmark this site:

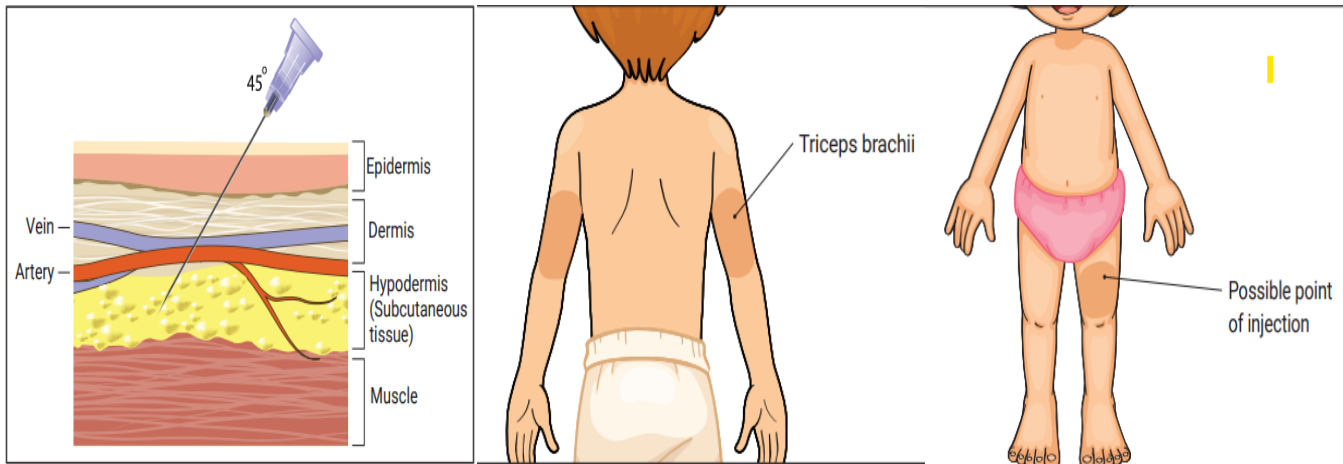
1. Divide the buttock into 4 quadrants.
2. The injection site is the centre of the upper outer quadrant (see below).



Source: http://www.bccdc.ca/resource-gallery/Documents/Guidelines%20and%20Forms/Guidelines%20and%20Manuals/Epid/CD%20Manual/Chapter%20%20-%20Imms/Appendix_B_Administration.pdf

2.5 Subcutaneous (SC)

This route is used when less rapid absorption is indicated. Use a 25-gauge $\frac{5}{8}$ inch needle at **45°** angle for subcutaneous injections. The upper outer triceps area is recommended for all individuals 12 months and older because the thickness of adipose tissue is greater than the adipose tissue over the deltoid muscle. The area over the anterolateral thigh is recommended for infants less than 12 months of age. **Do not aspirate before injection.**



Source: [Vaccine Administration: A Guide to Landmarking](#)

2.6 Intradermal Vaccines (ID)

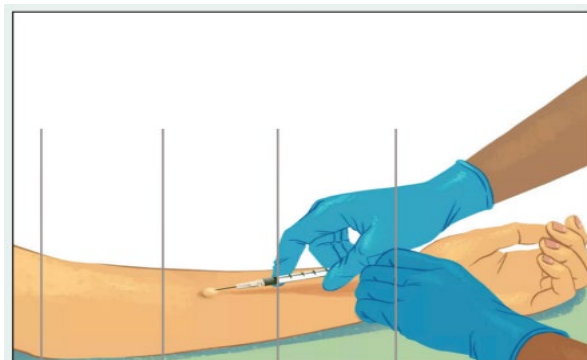
This immunization route is used when slow absorption **of vaccines** into the circulation is desired. Administer at a 5° to 15° angle. Care must be taken when administering a vaccine using the ID route properly, as a suboptimal immune response may occur if the vaccine is administered subcutaneously. **Do not aspirate before injection.**



Source: [Vaccine Administration: A Guide to Landmarking](#)

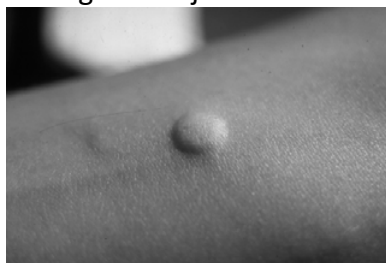
2.7 Intradermal Tuberculosis Skin Test (TST) (to screen for latent TB infection)

- **Schedule TST before, on same day as, or 4 weeks after a live vaccine has been received.**
1. The usual site for a TST is the flexor (anterior) surface of the inner (volar) forearm. Avoid red or swollen areas and visible veins.
 2. Use a 1 mL tuberculin syringe and a 26- or 27-gauge ¼-½ inch needle.
 3. Clean the vial with an alcohol swab and let dry before withdrawing 0.1 mL (5 tuberculin units) of purified protein derivative (PPD).
 4. Clean the injection site with an alcohol swab and let area dry before insertion.
 - Ensure that a topical anesthetic cream was not used prior to TST, as localized edema can occur at the injection site and may be confused with a positive TST.
 5. Gently stretch the skin to ensure that it is taut.
 6. Insert the needle at a 5° to 15° degree angle intradermally with the bevel up, until the needle tip is visible under the skin



Source: [Vaccine Administration: A Guide to Landmarking](#). For a demonstration of ID administration, see the following video: <https://www.youtube.com/watch?v=f3wMIDAdg0>.

- **Do not aspirate before injecting PPD.** Inject the PPD slowly. A defined bleb (wheal) approximately 6-10 mm in diameter should appear, then disappear within 15 minutes after injection (see photos above).
- Some PPD or blood may leak out of the site, but this is not cause for concern.
- Do not cover the site with a bandage or dressing.
- Tell client not to scratch or massage the injection site.



- If a bleb has not formed, the TST must be repeated on another appropriate site, at least 2 inches from the first site, and the second site must be circled to identify that this site result must be read.
7. Read the TST results in 48 - 72 hours.
 - Palpate for induration and mark the transverse edges with a pen.

- Measure the widest transverse diameter between the induration edges only, NOT the redness.
 - Always measure the TST result using a flexible calibrated ruler and **record measurement in millimetres only**.
 - If the result is between whole numbers (e.g., 5.6 mm) **round down to smallest whole number** (e.g., 5 mm). Negative TST results must be recorded as 0 mm.
 - Refer to Table 3: *TB Skin Test Result Interpretations* below.
8. Two-Step TB Skin Testing
- a) Two-step testing to detect a boosting response may be indicated for certain clients (e.g., HCWs or those ≥ 50 years). Refer to the *Saskatchewan TB Program Manual* for more information.
 - Two step testing can identify booster responders who have an anamnestic immune response from the TST. Individuals whose 1st TST is less than 10 mm are retested in 1 to 3 weeks to check for this response.
 - b) Two-step testing can also identify newly infected case contacts whose initial test was negative. They are retested up to 8 weeks later to assess if they have converted to being infected.
9. All TST results shall be document on the appropriate client health record/document and forwarded to TB Control Saskatchewan with the exception of tests performed off-reserve for the purpose of general occupational screening, travel screening and post-secondary education requirements.

Table 3: Interpretation of TST results and cutoff thresholds in various populations ([Canadian Tuberculosis Standards, 8th Edition](#))

TST Result	Situation in which reaction is considered positive
<5 mm	In general, this is considered negative
≥ 5 mm	People living with HIV Known recent (<2years) contact with a patient with infectious TB disease Fibronodular disease on chest x-ray (evidence of healed, untreated TB) Prior to organ transplantation and receipt of immunosuppressive therapy Prior to receipt of biologic drugs, such as tumor necrosis factor alpha inhibitors, or disease-modifying antirheumatic drugs Prior to receipt of other immunosuppressive drugs, such as corticosteroids (equivalent of ≥ 15 mg per day of prednisone for at least one month) Stage 4 or 5 chronic kidney disease (with or without dialysis)
≥ 10 mm	Recent (<2years) conversion of TST from negative to positive Diabetes (controlled or uncontrolled) Malnutrition (<90% of ideal body weight) Current tobacco smoker (any amount) Daily consumption of >3 alcoholic drinks Silicosis Hematologic malignancies (lymphomas and leukemia) and certain carcinomas (such as cancers of head, neck, lung and/or gastrointestinal tract) Any population considered at low risk of disease.

- All adverse events related to the administration of PPD must be documented in the patient's health record and recorded on the Health Canada, *Canada Vigilance Adverse Reaction Reporting Form*, available online at: http://www.hc-sc.gc.ca/dhp-mps/alt_formats/pdf/medeff/report-declaration/ar-ei_form-eng.pdf. These forms must also be forwarded to the regional MHO for review.

2.8 Infiltration of Rabies Immune Globulin (Rablg)

Infiltration of Rablg into a wound incurred by an animal bite protects individuals that have not been previously immunized with rabies vaccine. Two important goals of Rablg infiltration are:

1. To neutralize the rabies virus at the wound site as soon as possible after exposure; and
2. To prevent the migration of the rabies virus from peripheral nerves to the central nervous system.

These regimens are applicable for persons in all age groups, including children.

- Administer 20 IU/kg body weight.
- Because Rablg might partially suppress active production of rabies virus antibodies, no more than the calculated dose should be administered.
- If anatomically feasible, the full calculated dose of Rablg should be infiltrated around and into the depth of the wound(s).
- When the calculated dose is insufficient to infiltrate all the wounds, Rablg can be diluted with saline to obtain the necessary quantity up to a 3:1 dilution (e.g., 30 ml of physiological saline for 10 ml of Rablg for a total of 40 ml).
- Any remaining Rablg (including diluted) should be injected IM in the deltoid or vastus lateralis sites.
- Do not administer Rablg in the same syringe as rabies vaccine.
- Do not administer Rablg in the same anatomical site on the same day that rabies vaccine is given.

2.9 Intranasal (IN)

Live attenuated influenza vaccine (LAIV) is administered intranasally and is not to be injected. .

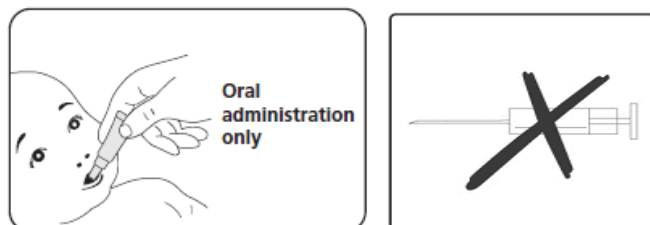
1. Remove the rubber tip protector. Do not remove the dose-divider clip at the other end of the sprayer.
 - a. The administration device is an AccuSpray™ nasal spray syringe that has a dose divider clip that allows for measured 0.1 mL spray into each nostril.
2. With the patient in an upright position, place the tip just inside the nostril to ensure LAIV is delivered into the nose. The patient should breathe normally.
3. With a single motion, depress the plunger as rapidly as possible until the dose-divider clip prevents you from going further.
4. Pinch and remove the dose-divider clip from the plunger.
5. Place the tip just inside the other nostril, and with a single motion, depress plunger as rapidly as possible to deliver the remaining vaccine.
6. The vaccine does not need to be repeated if the client coughs or sneezes after administration of the LAIV.



(Image retrieved February 7, 2012 from: http://www.cdc.gov/vaccines/pubs/pinkbook/downloads/appendices/D/vacc_admin.pdf)

2.10 Oral (PO)

Ready for Use Liquids: RotaTeq® is an oral solution and must never be injected.



1. Tear open the pouch and remove the dosing tube.
2. Clear the fluid from the dispensing tip by holding tube vertically and tapping cap.
3. Open the dosing tube in 2 easy motions:
 - a) Puncture the dispensing tip by screwing cap clockwise until it becomes tight.
 - b) Remove cap by turning it counterclockwise.
4. The infant should be seated in a reclining position.
5. Administer dose by gently squeezing liquid into infant's mouth toward the inner cheek until dosing tube is empty. A residual drop may remain in the tip of the tube.
6. Re-administration of another dose is not recommended if dose is spat out or regurgitated by the infant.

Capsules: Vaccine capsules should not be chewed but swallowed as soon as possible after placing in the mouth. Capsules should be taken with at least 4 oz of cool or lukewarm water as per the recommended schedule.

Sachet and Vaccine Vial Combinations: Medicines, food and drink need to be avoided for 1 hour before and after ingesting the vaccine. Do not use milk, juice or a carbonated beverage to mix the vaccine/sachet in.

1. The effervescent sachet needs to be mixed into a cup of 5 oz. cool water.
2. The vial must be shaken well and then poured into the cup.
3. Stir contents gently for 5 to 10 seconds.
4. The vaccine should be swallowed as soon after mixing as possible.

Double Sachet Combinations: Medicines, food and drink need to be avoided for 1 hour before and after ingesting the vaccine. Do not use milk, juice or a carbonated beverage to mix the sachets in.

1. The sachets must be folded along the solid black line and cut along the dotted line after insuring that the contents have been displaced to the bottom to prevent spillage.
2. The contents of both chambers are to be emptied simultaneously into 100 mL of cold or lukewarm water.
3. Gently mix the sachet contents for 5 to 10 seconds.
4. The vaccine should be swallowed as soon after mixing as possible.

3.0 MANAGEMENT OF PAIN AND ANXIETY

3.1 Positioning Infants and Children

The *Comforting Restraint* methods as presented below, actively involve the caregiver in embracing the child and controlling all four limbs. It avoids “holding down” or overpowering the child, but it helps the immunizer to steady and control the limb of the injection site.

3.1.1 Infants and Toddlers

3.1.1.1 Vastus lateralis

Verbally instruct and physically guide the caregiver to hold the child so that the vastus lateralis site is clearly visible and the child is firmly restrained to prevent movement during the immunization.

1. Ask the caregiver to fully uncover/unclothe the child’s leg and hold the child in a seated or semi-recumbent position on their lap as in the diagrams below.



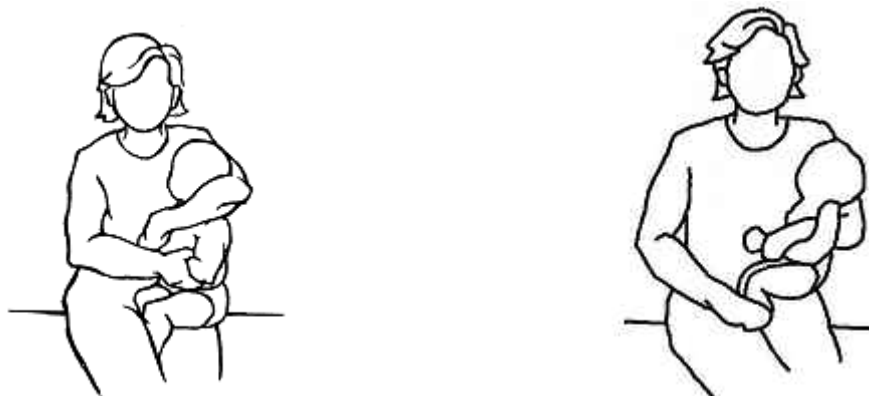
2. Ensure the child’s arm that is positioned closest to the caregiver, is tucked into the caregiver’s side, or placed behind the caregiver’s back. The child’s other arm is controlled with the caregiver’s arm and hand placed over it. In the case of children under 1 year of age, the caregiver can control both arms with one hand.
3. Instruct and guide the caregiver to firmly hold the child’s legs and feet between his or her thighs and control them with their free hand. The caregiver’s hand may be placed over the child’s knee to prevent the leg from being raised by the child during the immunization as shown below.



3.1.1.2 Deltoid

Verbally instruct and physically guide the caregiver to hold the child so that the deltoid site is clearly visible and the child is firmly restrained to prevent movement during the immunization.

1. Ask the caregiver to fully uncover/unclothe the child's arm and hold the child in a seated or semi-recumbent position on their lap as in the diagrams below.



2. Ensure the child's arm that is positioned closest to the caregiver is tucked into the caregiver's side or placed behind the caregiver's back. The child's other arm is controlled with the caregiver's arm and hand placed over it as pictured above.
3. Instruct and guide the caregiver to firmly hold the child's legs and feet between his or her thighs, and control them with their free hand, if necessary.

3.1.2 Kindergarten-Aged and Older Children

3.1.2.1 Deltoid

Verbally instruct and physically guide the caregiver to hold the child so that the deltoid site is clearly visible and the child is firmly restrained to prevent movement during the immunization.

1. Ask the caregiver to fully uncover/unclothe the child's arm and seat the child on parent's lap or have the child stand in front of the seated caregiver.



2. Ensure the child's arm that is positioned closest to the caregiver is placed behind the caregiver's back. The child's other arm is held close to the child's body with the caregiver's arm and hand placed over it as pictured above.

3.2 Recommendations for a More Successful Immunization Experience

All healthcare providers are encouraged to learn about the [CARD System](#) (Comfort, Ask, Relax, Distract) framework that helps people cope with pain, fear, and stress during vaccination procedures, offering strategies for a more positive experience. There are four components to the [CARD System](#):

1. **Comfort:** Focuses on creating a comfortable environment and providing physical comfort.
2. **Ask:** Encourages open communication and addressing concerns or questions.
3. **Relax:** Emphasizes relaxation techniques to reduce anxiety and tension.
4. **Distract:** Uses distractions to shift attention away from the procedure.

Benefits include:

- Reduced stress-related reactions like fear, pain, dizziness, and fainting during vaccination.
- Improved vaccination experience for the person receiving the vaccine and those supporting them.
- Teaching of lifelong coping skills that can be used in other stressful situations.

3.2.1 Foster a Culture of Empathy and Respect

- Ask about the child's previous experiences with needles. Individual responses to stress are influenced by temperament, environment and past experience.
- Acknowledge the child's feelings. Give permission to cry.
- Do not give false reassurance (i.e., "it won't hurt"). Honest reassurance is "it may hurt a bit, but I think you can handle it."
- Do not tolerate threats, shaming, or manipulation from the child's caregiver. When a caregiver threatens a child, the most helpful response is to offer empathy to the caregiver, state a neutral fact or principle and offer hope (e.g., "This must be frustrating for you. Immunizations are never emergencies. I think we can work out something we can all live with").
- Discourage the use of bribes and encourage their efforts – no matter how small.
- Remain firm as you manage the process. At the same time, show respect for the child.

3.2.2 Structure the Clinical Environment

- When a caregiver presents with more than one child, immunize the most anxious one first (usually the eldest), even if the caregiver thinks otherwise. The needle is the focus of the child's fear and watching while someone else is immunized may increase fear and anxiety.
- Provide privacy and prepare the immunization ahead, if possible, always out of sight of the child. If the child asks to see the needle, explain you will show it after the procedure.
- Describe what you plan to do, thereby displaying respect for a child's right to know, confidence in their ability to manage, and interest in addressing concerns. The child may wonder how long the needle will be in their arm or how quickly it will go in. Threatened loss of control is a factor in needle fear.
- Consider the use of practice dolls with children under 6. Offer to immunize a stuffed toy or doll, and invite the child to hold the "patient". Use a syringe without a needle and go through all the steps, explaining each one as you proceed.
- Provide limited, realistic choices and let the child decide (e.g., "Would you like to use your right or left arm?" "Would you prefer to sit or stand?"). Offering realistic choices creates a setting where the child can maintain some personal control and contributes to an atmosphere of mutual respect. Supportive, secure positioning can be achieved with a child (depending upon age) either standing or sitting.
- Do not have the caregiver restrain the child before you are ready to administer the vaccine. The longer the child is restrained the greater the loss of personal control and hence increased anxiety. The goal of restraint is not to overpower the child, but to assist the child to remain as still as possible for the procedure.
- Manage the time and set limits. If the child cannot calm him or herself, acknowledge their effort and offer a rest period. If there is no other alternative, reschedule the immunization.

3.3 Evidence-Based Interventions for Pain and Anxiety

The physical discomfort associated with injections is short-lasting but may cause anxiety and a life-long fear of needles, medical procedures and health care providers. Evidence-based management interventions enable caregiver and immunizers to comfort and reassure children, and may reduce potential psychological effects (e.g., trauma) of such procedures. It is recommended that PHNs:

- a) Refer to *CIG Vaccine Administration Practices Table 4: Immunization* pain management strategies for children, by age groups (as Appendix 8.3 in this chapter).
- b) Watch the *Reduce the Pain of Vaccination in Babies* video at:
<http://www.youtube.com/watch?v=dZcBc9UnMtw>;
- c) Read *Reducing pain during vaccine injections: clinical practice guidelines* (Taddio et al, 2015)
<http://www.cmaj.ca/content/early/2015/08/24/cmaj.150391>; and
- d) Read *Pain Management during Vaccine Injections A Clinician's Guide: Children < 3 years*
(http://immunize.ca/uploads/pain/5p_less_than_3_web_e.pdf)

3.3.1 Infants and Toddlers

- A child, who is dry, fed, and emotionally content, may react less strongly to pain during immunization.
- Ask caregivers to position and cuddle their baby on their lap during the immunization. Allow the baby to suck before, during and after the immunization:
 - If mothers are breastfeeding, encourage them to nurse their baby before, during and after the immunizations. Breastfeeding calms and comforts both the baby and mother and can reduce baby's pain.
 - There is some evidence that giving a sweet tasting solution to non-breastfeeding infants just prior to injections may provide analgesia. Honey is not to be given to infants under 12 months of age.
 - Encourage non-breastfeeding mothers and other care providers to use a pacifier or formula to comfort and calm the baby.

3.3.2 Older Toddlers and Children

Children may “pick up” on their caregiver’s anxiety. By coaching caregivers to adopt a calm affect, caregivers and children may have a better immunization experience. Distraction techniques and behaviour modification measures are effective in decreasing pain response during and following immunization. Regardless of the type of distraction, the more the child is involved in the distraction, the lower their pain.

- Coach children to count or squeeze hard on the hand of the caregiver.
- Work with the caregiver to use distraction techniques such as singing, reading, blowing soap bubbles or using toys.
- Slow, deep breathing has a physiologic calming effect and can, at minimum, limit anxiety escalation. Coach children to engage in slow deep breathing or blowing (away the pain) during the each immunization.
- Briefly describe to the child what will happen during each immunization and how it will feel. Do not say that it won't hurt. Instead say the pain lasts a short time and feels like a “sting”, “poke” or “squeeze”.
- Answer children's questions:
 - Why do I need a needle? (“To help you stay healthy.”)
 - What will happen? (“The medicine will be put in your arm with a needle.”)
 - How will it feel? (“You may feel a poke or small sting that will last a few seconds.”)

3.4 Topical Anaesthetics

Clients who are concerned about the pain associated with immunization may be interested in using topical anaesthetics. Children and adults may feel a sense of control when using these products, and distress and anxiety may be prevented or lessened. This is preferable, instead of delaying or avoiding immunization because of a fear of needles. Pain relief occurs at the insertion point of the needle, but pressure felt within the muscle upon injection of the product is generally unaffected.

Topical anaesthetics should be applied to intact skin over a suitable injection site for the minimum duration of time required (see [Table 4](#)) and removed prior to immunization. When used correctly, they cause superficial localized anaesthesia at the injection site. While the risk of serious adverse reactions such as methemoglobinemia and seizures is low, risk increases if a greater than recommended amount is applied, if the product is left on longer than recommended and/or the product is applied to non-intact skin.

Non-prescription topical anaesthetics are available at pharmacies and clients are encouraged to consult a pharmacist for further product information. It is the responsibility of parents/clients who wish to use these products to follow the manufacturer's instructions and check the contraindications, precautions, and possible side effects prior to use.

3.4.1 Cream and Gel Formulations

Refer to [Table 4: Non-Prescription Topical Anaesthetics Available in Canada](#)

Generally, products should not be used by individuals who are sensitive to any of the components in these products. Lidocaine and prilocaine are amides, while tetracaine is an ester. True allergies to topical anesthetics are rare, with most reactions likely caused by other ingredients, such as preservatives. If an allergic reaction occurs, it is advisable to try a product from a different class, preferably one that is preservative-free or uses a different preservative.

3.4.1.1 MAXILENE™ (RGR Pharma)

MAXILENE™ is the brand name of a topical anaesthetic consisting of liposomal lidocaine 4% or 5% and is indicated to relieve the pain and/or itching associated with minor burns, sunburn, minor cuts, scrapes, insect bites or minor skin irritations in individuals 2 years of age and older. It needs to be used with caution in individuals with heart block or severe hepatic impairment. A physician must be consulted prior to use in children under the age of 2 years.

3.4.1.2 EMLA® Cream (Aspen Pharmacare Canada)

EMLA® is the brand name for a topical anaesthetic consisting of 2.5% lidocaine and 2.5% prilocaine. It comes in a multidose cream and a single dose patch. The product requires a longer application time than other topical anaesthetics. EMLA® is contraindicated in individuals with congenital or idiopathic methemoglobinemia; in pre-term infants (gestation less than 37 weeks); in infants less than 3 months of age (young infants have a higher risk of developing methemoglobinemia); and in infants less than 12 months of age who are taking methemoglobinemia-inducing agents (e.g., sulfonamides). It needs to be used with caution in individuals with heart block or severe hepatic impairment. Refer to the product monograph for detailed information (in references).

3.4.1.3 AMETOP® GEL 4% (Valeo Pharma)

AMETOP® GEL 4% is the brand name for a topical anaesthetic consisting of 4% tetracaine that is indicated for use in individuals one month of age and older. AMETOP® GEL 4% achieves anaesthesia within 30 to 45 minutes and may last for four to six hours after removal. When compared to EMLA®, AMETOP® GEL 4% has been found to provide better anaesthesia when used before needle insertion in children. It is contraindicated in premature babies or full-term infants less than one month of age, in whom the metabolic pathway for tetracaine may not be fully developed.

Table 4: Non-Prescription Topical Anaesthetics Available in Canada

Active Ingredient	Liposomal Lidocaine	Lidocaine/Prilocaine	Tetracaine
Brand	MAXILENE 4™ MAXILENE 5™	EMLA® Cream	AMETOP® GEL 4%
Time to effect	30+ min	60+ min	30+ min
Occlusion recommended?	No*	Yes (for the cream)	Yes
Duration of effect after removal	1-2 hours	≥ 2 hours	4-6 hours
Concentration/ Dosage Form	4% cream	2.5%/2.5% cream	4% gel
	5% cream	2.5%/2.5% patch	
Availability	5 g, 30 g	No dressing 30 g, w/ dressings 5 g	1.5 g
	15 g, 30 g	1 g	
Comments	- Contraindications: allergy, application on mucosae or an open wound or in eyes. - Preserved with benzyl alcohol and propylene glycol.	- Contraindications: allergy, application on mucosae or an open wound or in eyes, methemoglobinemia, G6PD. - Preservative-free	- Contraindications: allergy (including PABA and sulfonamides), application on mucosae or an open wound or in eye - Preserved with parabens. The manufacturer does not advise using it prior to immunization.

*Unless area needs to be covered to prevent child from ingesting anaesthetic.

Source: Saskatchewan Drug Information Services; updated 2025, medSask

3.4.2 Vapocoolants

Evidence has been equivocal, though a systematic review has concluded vapocoolants are no more effective than placebo and are associated with some pain from application. [Gebauer](#) Company's [Ethyl Chloride®](#) and [Pain Ease®](#) sprays provide immediate but time-limited pain numbness but are not recommended for those younger than 4 years old and for children with sickle cell disease, and must not be sprayed on mucosae or an open wound.

3.4.3 Handheld Portable Devices

There is evidence that such devices, such as the [Buzzy®](#), can provide drug-free pain prevention for injections, including vaccines.

3.5 Post-Immunization Client Care

1. Provide current information about the management of common vaccine side effects.
2. Prophylactic administration of acetaminophen prior to or immediately post-immunization for pain management is ineffective and is not recommended because of interference with vaccine induced immune responses. The March 2018 *Paediatrics and Child Health* article **Fever prophylaxis can reduce vaccine responses: A caution** states:

“Prophylactic use of antipyretic/analgesic drugs can reduce immune responses to some infant vaccines, warranting judicious use. The clinical significance of such reduced responses is uncertain but stronger responses are obtained in the absence of prophylaxis. In contrast, using these drugs to treat symptoms once they appear is unlikely to interfere with immune responses and would reduce the number of asymptomatic children exposed to other potential drug adverse effects. The above observations that anti-inflammatory drugs only interfere with antibody responses if present during the first 6 to 8 hours after immunization serve as a reminder that injection site inflammation is an essential first step in initiating responses to vaccines, activating dendritic cells and recruiting macrophages that rapidly transport vaccine antigens to regional lymph nodes where antibody responses begin. Acetaminophen and ibuprofen target different parts of the inflammatory response cascade, likely explaining their differing effects on immune responses.”
3. As of this date, there is no supporting evidence for effective prophylactic use of acetaminophen in children prone to febrile seizures.
4. When fever occurs, acetaminophen (e.g., Tylenol™, Tempra™) is recommended for infants and young children because it controls fevers better than other anti-pyretics. Always counsel caregivers to provide anti-pyretic doses based on their child’s weight, not age. Provide caregivers a copy of the *Caring for Your Child’s Fever* fact sheet, available at: <http://www.saskatchewan.ca/residents/health/accessing-health-care-services/immunization-services#immunization-forms-and-fact-sheets>
5. Ibuprofen (e.g., Motrin™, Advil™ or other brands) can be used for children over 6 months if acetaminophen does not relieve the fever and the child is drinking well. Read the Ibuprofen box for the correct dose. Do not alternate acetaminophen and ibuprofen, as there is a risk of overdose.
 - **Acetylsalicylic acid (Aspirin) is not recommended for children with fever because of its association with a neurological condition known as Reye's syndrome.**
6. Review expected side effect with caregivers or clients, and counsel them when to seek medical attention post-immunization
7. It is recommended that all immunized clients remain in the clinic for 15 minutes post-immunization. This may facilitate the management of any adverse reactions.
 - Those with a previous adverse event should remain at the clinic site for 30 minutes or longer, or according to MHO recommendations.

4.0 REFERENCES

Barron, C., & Cocoman, A. (2008). Administering intramuscular injections to children: What does the evidence say? *Journal of Children's and Young People's Nursing*. 2(3), pp. 138-144.

Canadian Agency for Drugs and Technologies in Health (2010). Filtered Needles for Withdrawing Medication from Glass Ampoules: A Review of the Cost-Effectiveness and Incidence of Complications. Available at: http://www.cadth.ca/media/pdf/L0158_Filtered_needles_final_no_abs.pdf

Canadian Pediatric Society: Position Statement: [Managing pain and distress in children undergoing brief diagnostic and therapeutic procedures](#).

Hogan ME, Smart S, Shah V, Taddio A. A systematic review of vapocoolants for reducing pain from venipuncture and venous cannulation in children and adults. *J Emerg Med*. 2014 Dec;47(6):736-49.

Immunize.org [How to Administer Intranasal and Oral Vaccinations](#)

PHAC (2024) Vaccine Administration: A Guide to Selecting Needle Gauge and Length https://publications.gc.ca/collections/collection_2024/aspc-phac/HP40-353-2024-eng.pdf

PHAC (2024) Vaccine Administration :A Guide to Landmarking: https://publications.gc.ca/collections/collection_2024/aspc-phac/HP40-354-2024-eng.pdf

PHAC. *Canadian Immunization Guide* (Evergreen Ed.). Available at: <http://www.phac-aspc.gc.ca/publicat/cig-gci/index-eng.php>.

PHAC *Canadian Tuberculosis Standards* (8th Ed.) Ottawa, ON. Available at: <https://www.tandfonline.com/toc/ucts20/6/s>

David Scheifele, Brian Ward; Fever prophylaxis can reduce vaccine responses: A caution, *Paediatrics & Child Health*, pxy011, <https://doi.org/10.1093/pch/pxy011>

Taddio, A., Appleton, M., Bortolussi, R., Chambers, C., Dubey, V., Halperin, S., et.al. (2010). Reducing pain in childhood vaccination: an evidence-based clinical practice guideline. *CMAJ*, 182(18), pp. E843-E855. Available at: <http://www.cmaj.ca/cgi/content/full/182/18/E843?maxto=show=&hits=10&RESULTFORMAT=1&andorexacttitle=and&andorexacttitleabs=and&fulltext=emla&andorexactfulltext=and&searchid=1&FIRSTINDEX=0&sortspec=date&resourcetype=HWCIT,HWELTR>

Taddio, A. et al (2015) Reducing pain during vaccine injections: clinical practice guidelines. *CMAJ*. <http://www.cmaj.ca/content/early/2015/08/24/cmaj.150391>

Toronto Public Health. "How to reduce your child's pain from immunization". Available at: www.toronto.ca/health/immunization_children/howtoreducepain.htm
http://www.toronto.ca/health/immunization_children/pdf/howtoreducepain.pdf

Weniger, B. G., & Papnia, M. J. (2008). Alternative vaccines delivery methods. In Plotkin, S., Orenstein, W. & Offit, P (Eds), *Vaccines* (5th Ed). Saunders Elsevier: China.

World Health Organization (1996). Laboratory techniques in rabies (4th Ed.). Geneva: Author. Available at:
http://libdoc.who.int/publications/1996/9241544791_eng.pdf

5.0 APPENDICES

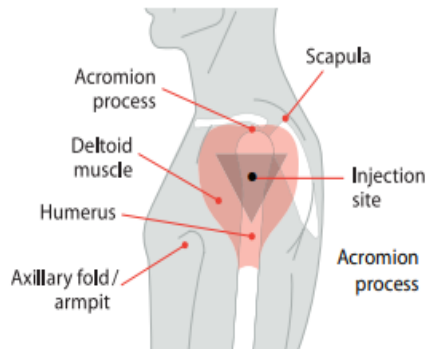
Appendix 8.1: Administration of Multiple IM Injections to the Deltoid Site

It is important to assess the adolescent or adult recipient's muscle mass to accommodate multiple injections into one deltoid site.

The diagrams below illustrate options for administering one, two, or three vaccinations in a single arm, spaced at least 1" apart. Additional injections can also be administered in the opposite arm.

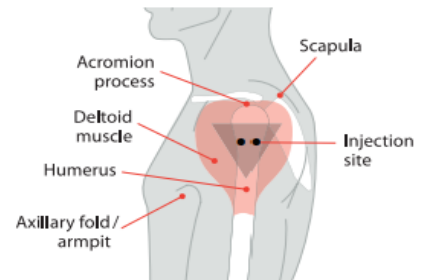
Use anatomical landmarks to determine the injection site in the deltoid muscle (a large, rounded, triangular shape). Find the acromion process, which is the bony point at the end of the shoulder. Then, locate the injection site which will be approximately 2" below the bone and above the axillary fold/armpit.

Single IM injection in deltoid



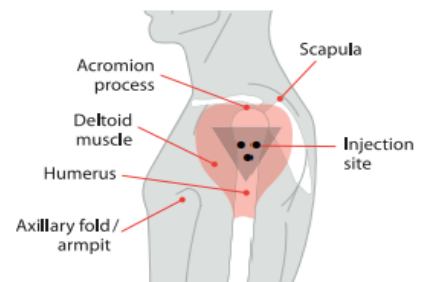
Two IM injections in deltoid

Space injections at least 1" apart.



Three IM injections in deltoid

Space injections at least 1" apart.



Source: [How to Administer Multiple Intramuscular Vaccines to Adults During One Visit](#)

Appendix 8.2: Potentially Immunosuppressive Biologic Agents

- Infants who were exposed during pregnancy to immunosuppressive biologic agents should be referred to the Pediatric Infectious Diseases Clinic for assessment and vaccine counselling, especially regarding live vaccines, such as rotavirus.
- Immunosuppressive biologic agents include those:
 - used for transplant recipients or for cancer treatment, or
 - found in the list below (not an exhaustive list)

Immunosuppressive Agent*	Brand Name(s) (Reference biologic agent listed first, followed by biosimilars in alphabetical order)
Abatacept	Orencia®
Adalimumab	Humira®; Abrilada®; Amgevita™; Hadlima®; Hulio®; Hyrimoz®; Idacio®; Simlandi™; Yuflyma™
Alemtuzumab	Lemtrada®; MabCampath®*
Anakinra	Kineret®
Anifrolumab	Saphnelo
Belimumab	Benlysta™
Bimekizumab	Bimzelx®
Brodalumab	Siliq™
Etanercept	Enbrel®; Erelzi®; Brenzys®; Rymti
Golimumab	Simponi®
Guselkumab	Tremfya®
Infliximab	Remicade®; Avsola®; Ixifi®; Remdantry™; Remsima™; Renflexis®;
Ixekizumab	Taltz®
Lebrikizumab	Eblyss™
Mirikizumab	Omvoh™
Natalizumab	Tysabri®
Ocrelizumab	Ocrevus®
Ofatumumab	Kesimpta™
Risankizumab	Skyrizi®
Rituximab	Rituxan®; Riximyo®; Ruxience™; Truxima™
Sarilumab	Kevzara®
Secukinumab	Cosentyx®
Tildrakizumab	Ilumya™
Tocilizumab	Actemra®; Tyenne®
Ustekinumab	Stelara®; Finlius® ; Jamteki™; Otulfi™; Pyzchiva; Stegeyma®; Wezlana™
Vedolizumab	Entyvio®

* Certolizumab pegol is not included based on data that it does not cross the placenta.

Appendix 8.3: Immunization pain management strategies, by age group ¹ (CIG)

Age group	Pain management strategies
All ages	• Injection of vaccines without aspiration • Injecting vaccines that cause the most injection site pain after other vaccines
Infants and young children (3 years of age and under) ²	• Education of parent/caregiver about pain management before and on the day of immunization • Topical anesthetics prior to vaccine injection • Presence of parent/caregiver during vaccine injection • Breastfeeding during vaccine injection (≤ 2 years of age) • If the infant/young child is not breastfed during vaccine injection, a combination of other strategies may be used, such as: ○ Skin-to-skin contact during vaccine injection (≤ 1 month of age) ○ Holding during vaccine injection or holding and rocking/patting after vaccine injection ○ Administration of a sweet-tasting (sucrose or glucose) solution prior to vaccine injection (≤ 2 years of age) ³
Children (3 to 12 years) ²	• Education of parent/caregiver about pain management before and on the day of immunization • Education of individual about pain management for vaccine injection on the day of immunization ⁴ • Topical anesthetics prior to vaccine injection • Presence of parent/caregiver during vaccine injection (≤ 10 years of age) • Sitting up during vaccine injection
Adolescents (12 to 17 years) ²	• Education of parent/caregiver about pain management before and on the day of immunization • Education of individual about pain management for vaccine injection on the day of immunization ⁴ • Sitting up during vaccine injection
Adults (≥ 18 years)	• Education of individual about pain management for vaccine injection on the day of immunization ⁴ • Sitting up during vaccine injection

1 The strategies noted in the table are strong recommendations for reducing pain during vaccine injections from Taddio A, McMurtry CM, Shah V et al. (HELPinKIDS & Adults Team) Reducing pain during vaccine injections: clinical practice guideline. CMAJ 2015; 187(13): 975-982. For additional recommendations on pain management strategies, refer to the article.

2 There is some overlap in ages across these categories (i.e., children aged 3 and 12 years are included in two separate categories) owing to the need to balance over-simplification in creating age categories with appropriate guidance, overlap in the underlying literature base as well as substantial differences in developmental trajectories of individual children.

3 If an infant is scheduled to receive a dose of rotavirus vaccine, provide it first as it is sweet-tasting. In this scenario, no additional sweet-tasting solution is required prior to vaccine administration.

4 For example, give information on what to expect (the procedure and how it will feel) as well as suggestions on how to cope.

Source: <https://www.canada.ca/en/public-health/services/publications/healthy-living/canadian-immunization-guide-part-1-key-immunization-information/page-8-vaccine-administration-practices.html#t4>

Appendix 8.4: Oral Vaccine Administration via Enteral Tube

Saskatchewan Immunization Manual STANDARD WORK	Name of Activity: Procedure – Oral Vaccine Administration via Enteral Tube Role performing Activity: Public Health Nurse		
	Location: Saskatchewan Immunization Manual		Department: Chapter 8
	Date Prepared: 2019-04-16	Last Revision:	Date Approved:

Standard Work Summary: Steps to administer oral vaccines to infants who are unable to take vaccines by mouth and who have an enteral feeding tube.

Anatomical location of enteral feeding tube	A. Nasogastric (NG) B. Orogastric (OG) C. Nasojejunal (NJ) D. Gastrostomy (G-tube) E. Jejunostomy (J-tube) F. Gastro-jejunostomy (G-J tube) G. Percutaneous enterogastrostomy (PEG) H. Percutaneous gastrojejunostomy (PGJ or PEJ)
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Sequence	Task Definition
1.	Check with the parent/ guardian to ensure that they have the necessary supplies for the procedure (e.g., two appropriate syringes at the immunization appointment). - NOTE - If the parent/guardian presents without the necessary supplies, provide the other vaccine(s) that are due. Follow up with a home visit or a clinic visit to administer the oral vaccine. Med cups do not need to be supplied by parents as they will be available at clinics. - Two appropriate syringes (3 mL syringe or larger) **NOTE largest amount of volume need to prime a NG line is 1.8 mL so a 3 mL syringe is adequate. - One appropriate 3 mL syringe for placement confirmation & flushing. - One appropriate 3 mL syringe for administering medication. - Sterile water is recommended for enteral tube flushing when oral vaccines are being administered. Sterile water ampoules can be on hand at clinics and therefore not needed for parents to provide.
2.	Perform hand hygiene. Maintain clean technique when accessing tube& administering medication.

3.	Check placement and patency of the enteral tube. - Do not proceed with vaccine administration by enteral feeding tube unless tube placement and patency has been confirmed. Nasogastric or Orogastric A. Confirm placement of gastric tube in nares or mouth with 2 step process: - Confirm with parent that the centimeter marking visible at a nare or the mouth is correct. - Confirm by visualizing stomach content. - Kink off gastric tube, remove end cap of enteral tube port. Attach empty appropriate syringe to gastric tube. - Gently aspirate 0.5-1.0 mL of stomach content into syringe and then return visualized gastric contents with push pause technique back into stomach. - Kink off gastric tube, remove syringe and recap enteral tube port B. Nasojejunal - Confirm with parent the external baseline length of tube measurement. C. Gastrostomy (button/balloon enteral tube) - Confirm correct placement by ensuring the flange is flush to the skin. D. Jejunostomy - Confirm correct placement by ensuring the flange is flush to the skin. E. Gastro-jejunostomy - Confirm correct placement by ensuring the flange is flush to the skin. F. Percutaneous enterogastrostomy - Confirm correct placement by ensuring the flange is flush to the skin. G. Percutaneous gastrojejunostomy (PGJ or PEJ) - Confirm with parent the external baseline length of tube measurement.
4.	Flushing of gastric tube pre-vaccine administration: - Using same appropriate syringe for placement check, draw up 3mL of sterile water for the flush. - The reason to use the same syringe is to help remove the acidic stomach content residual from sticking to the inside of the syringe. - Clamp off gastric tube, remove cap and attach appropriate syringe with sterile water. - Gently flush with a pause-push technique. - Kink off gastric tube, by folding the tube over on itself, remove the syringe and re cap the enteral tube port.
5.	Administration of vaccine: - Squirt the oral vaccine content into a medicine cup. - Using the second 3 mL syringe, draw up all vaccine from the medicine cup. Once medication drawn up, ensure all air is removed from the syringe prior to connecting to the gastric tube. - Kink off the gastric tube, uncup the enteral tube port and attach it to medication syringe. - Release the kink and use a push-pause technique to administer the medication through the tube. - Kink off tube, remove the medication syringe and recap the enteral tube port. - Dispose of syringe if syringe does not belong to client or have family rinse syringe if syringe belongs to client and they do not want to dispose of it.
6.	Flushing of gastric tube post-vaccine administration: - Draw up 3 mL of sterile water into the designated “flush” syringe. - Kink the tube and uncup the tube port and attach flush syringe. - Unkink tube and gently flush with a pause-push technique. - Kink off tube, remove syringe, and recap tube port. - Dispose of remaining water.
7.	Perform hand hygiene.
8.	Ensure client has all appropriate syringes if provided by family. If syringes provided by health care, dispose of syringes in garbage.
9.	Documentation: - Document vaccine as administered orally (PO). - Add a note in comments section stating the oral vaccine was administered via enteral tube.

Appendix 8.5: Mass Immunization Clinic Infection Control Recommendations

Purpose

- To protect the health of clients and public health employees.
- To prevent the transmission of infectious diseases from person to person.
- To maintain public confidence in the immunization services and delivery.

Clinic advertisement

- Inform clients if physical distancing measures and mask use are in effect in mass immunization clinics.
- Clients are to be made aware that health screening may occur prior to immunization.
- Special population clinics (e.g., for seniors, or those who are immune compromised) should be clearly stated.

Space requirements

- Clinic location can accommodate clients without overcrowding, to accommodate physical distancing measures.
- Proper waste management/disposal.
- Tables, counters, chairs and mats that are easily cleaned and disinfected.
- Separate waiting area for individuals who have been immunized, taking into account physical distancing
- Food, beverages, toys, etc. are not used by clients or staff in the clinic area.

Supplies

- Posters and factsheets on hand hygiene, cough etiquette and infection prevention messages.
- Sufficient alcohol-based hand sanitizer (ABHS) gel, foam or wipes and lotion for clinic staff and clients.
- Surgical masks, face shields, goggles.
- Disposable gloves.
- Surface disinfectant wipes, disinfectant spray, paper towels, disposable cloths, hands-free garbage cans with plastic liners.
- Disposable tissues.

Immunizers

- Shall apply physical distancing measures between immunization stations.
- Shall disinfect chair and table surfaces between all clients.
- Screen clients for symptoms before immunizing them.
- Hand hygiene shall be practiced according to policy including:
 - ✓ Before entering and leaving the work area.
 - ✓ Before preparing or handling sterile products or medications.
 - ✓ Before and after contact with a client.
 - ✓ After removing disposable gloves.
- Food and beverage consumption, including water bottles, is not permitted in clinical areas.

Clients

- Screen selves for illness.
- Signage shall be posted at the entrance to the facility to advise the public that they are required to use ABHS upon entering and leaving the facility and touching door handles; to practice cough etiquette, and if they have disease symptoms that they will not be immunized and should leave the facility immediately.
- All clients shall use ABHS upon sitting at the immunization station.
- Apply physical distancing practices to reduce crowds in queues and post-immunization observation areas to prevent potential transmission.

Appendix 8.6: Immunization Clinic Supply Recommendations

OFFICE SUPPLIES		
Paper 8.5" x 11"	Tape	File boxes/lock boxes/file folder system
Pen, pencils, highlighters, sharpies	Rubber bands to bundle vaccines	Date stamps, stamp pads
Envelopes	Sticky tack for posters	Black clips 2" and 1" for papers
Clipboards	Sticky notes and large wall post for staff updates	Secure folders
Scissors, staplers, staples, paperclips	Stickers	Transparent page protectors
IT REQUIREMENTS		
Laptops	Portable printers/scanners	Power bars
Computer/TV for screening/information videos	Telephones, cell phones, chargers	Extension cords
Thumb drives, MiFi drives	2-way radios	Portable projector
SECURITY REQUIREMENTS		
Organization approved ID badges	Volunteer confidentiality forms	Staff temperature screening list and station
Sign-in/sign-out station for staff, volunteers	Master schedules	
OTHER ITEMS		
Ear protection or back of head strap holders	Rolls of clear packing film or bungee cords	Locking plastic bags
CVA vehicles SUV or van when possible	Tape measures for 6-foot physical distancing	Transportation of dangerous goods forms
Maintenance support/assistance	Totes or duffle bags to transport supplies	Poster for HCW to self-identify to nurses
Plexiglas partitions to set on tables	Bright coloured duct tape	Hand sanitizer dispenser stations or bottles
Sit-stand chairs – local need for injury prevention	Juice boxes, water bottles	Electronic signage
IMMUNIZATION SUPPLIES/EQUIPMENT		
Vaccine (and diluent/adjuvant if applicable)	Face shield, mask with visor	Bandages
3 mL syringes with safety needles, 25 gauge 1"	Biohazard yellow bags (for gowns)	Vaccine insulated cooler bags/coolers
3 mL syringes	Biohazard waste boxes and liners	Ice/cold packs
25 G 1" safety needles	Hand Sanitizer	Latex-free gloves – assorted sizes
25 G 1½" safety needles	Anaphylaxis kits	Min-Max thermometers, coolers
1 mL 25 G tuberculin syringes	Stethoscope	Patient thermometers
18 G 1" needles	Brown paper bags	Mats, blankets
Cotton balls	Alcohol swabs, chlorhexidine swabs	Disposable gowns
Sharps containers (4 L or larger, small)	Procedure towels	Surgical masks - disposable
Blood pressure cuffs (child and adult reg. and large)	Goggles intruder safety, clear, anti-fog lenses with cleaner	Pocket masks with one way valve – child and adult
DOCUMENTATION AND SUPPORT FORMS		
Program document	AEFI report forms	Team lists/Team Schedule
Consent forms	Blood and body fluid exposure incident form	Vaccine Count sheets
Line list form	Emergency phone lists	Stat forms
Vaccine fact sheets	Immunization Wallet cards	Vaccine temperature monitoring sheets
Vaccine product monographs	Incident Report form	Freeze Watch/Warm Mark indicators
GENERAL SUPPLIES AND EQUIPMENT		
Tables, table covers	Signage	Paper towels
Chairs	Garbage bags	Facial tissues
Screening drapes (portable partitions) for privacy	Disinfectant wipes, spray cleaning solution	Flashlight
Traffic ropes and stands	Cart for wheeling supplies	Soap, toilet paper, paper towelling