

# 2020-2021 Seasonal Influenza (Flu) Vaccine Consent Form

Section 1: Patient Information				Date (MM/DD/YYYY):	
Last Name:	First Name:	Prov. Health Number:	Gender:		
Main Phone Number:	Alternate Phone Number:	Date of Birth (MM/DD/YYYY):	Age:	Child's weight: (kg / lb)	
Address:	City:	Province:	Postal Code:		
Emergency Contact's Last Name:	Emergency Contact's First Name:	Relationship:	Emergency Contact's Main Phone Number:		
Emergency Contact's Alternate Phone Number:	Ask your pharmacist about age restriction for flu shots in a pharmacy				

Section 2: Screening Questionnaire Refer to <a href="#">Screening Questionnaire Action Guide</a> for recommendations		Yes	No
Are you, or have you been <b>sick within the past 3 days</b> ? (fever greater than 39.5°C, breathing problems, or active infection)			
Have you had difficulty breathing, wheezing or chest tightness within 24 hours of getting an <b>influenza vaccine</b> ?			
Are you <b>allergic</b> to any part of the influenza vaccine, or have you had a severe, life-threatening allergic reaction to a past influenza vaccine?			
Are you allergic (eg. Wheezing, chest tightness, difficulty breathing, hives) to: <b>• Contact lens solution • Egg or egg products • Formaldehyde • Gelatin • Gentamicin • Kanamycin • Neomycin • Thimerosal • Polymyxin B</b>			
Do you have a serious allergy to <b>latex or natural rubber</b> ?			
Have you had a reaction to <b>eggs or egg products</b> but can still eat small amounts of egg? (eg. Stomach ache, skin reaction)			
Have you had <b>Guillian-Barré Syndrome</b> within 6 weeks of getting an influenza vaccine? <b>Oculo-Respiratory Syndrome</b> ?			
Have you ever had a <b>seizure</b> or have an active, new, or changing <b>neurological disorder</b> ?			
Do you have <b>bleeding problems</b> or use <b>blood thinners</b> ? (eg. Warfarin)			
Are you <b>pregnant, nursing, or do you intend to become pregnant</b> ?			
Have you received your pneumonia vaccines? If yes, which vaccine _____ and when: _____			
Have you received your shingles vaccines? If yes, which vaccine _____ and when: _____			
Have you received any <b>vaccines</b> in the last 4 weeks?			
For children under 18 years old: Is the child using, or will be using an <b>aspirin/aspirin-containing therapy</b> in the next 4 weeks?			
Do you have <b>severe asthma</b> (on high dose inhaled or oral corticosteroids) or medically attended <b>wheezing</b> in the past 7 days?			
Have you received in the past 48 hours or do you intend to receive in the next 2 weeks <b>flu antiviral therapy</b> ? (eg. Oseltamivir)?			
Do you have any medical conditions (eg. Cancer, leukemia, HIV/AIDS) or take medications that weaken the <b>immune system</b> ?			
Do you provide health care services to or do you have close contact with persons who are <b>immunocompromised</b> ?			
Are you allergic (eg. Wheezing, chest tightness, difficulty breathing, hives) to Arginine?			

Section 3: Consent Given By Patient/Agent		
<p>I, the undersigned patient, parent or guardian, have read or have had explained to me information about the seasonal influenza vaccine ("Vaccine") as outlined on the Flu Vaccine Fact Sheet. I have had the chance to ask questions, and answers were given to my satisfaction. I understand the risks and benefits of receiving the Vaccine. After getting the Vaccine, I agree to wait in the clinic/pharmacy for 15 minutes (or the time recommended by the pharmacist).</p> <p>I am aware it is possible (yet rare) to have an extreme allergic reaction to any component of the Vaccine. Serious reactions called "anaphylaxis" can be life-threatening medical emergencies. Symptoms of an anaphylactic reaction may include hives, difficulty breathing, swelling of the tongue, throat, and/or lips. If I experience such symptoms following vaccination, I am aware it may require the administration of epinephrine, diphenhydramine, beta-agonists, and/or antihistamines to treat this reaction and 9-1-1 will be called to provide additional assistance. In the event of anaphylaxis, I, my agent, and/or EMS paramedics will receive a copy of this form. I understand the information contained on this form, may be disclosed to the public health authority and to other required parties for the purpose of adverse event and drug safety reporting.</p>		
<input type="checkbox"/> I confirm that I want to receive the seasonal influenza vaccine <b>OR</b> <input type="checkbox"/> I confirm that I want my child to receive the seasonal influenza vaccine		
Patient/Agent Name (& Relationship)	Patient/Agent Signature	Date Signed (MM/DD/YYYY)

PHARMACY USE ONLY Section 4: Prescription Templates Influenza Vaccine Used					
<b>HEALTH CARE PROVIDER'S DECLARATION:</b>					
<input type="checkbox"/> I confirm the above named patient is capable of providing consent for the seasonal influenza vaccine and that the seasonal influenza vaccine should be given to the patient. I am administering the seasonal influenza vaccine no more than <u>21 days</u> after the consent was signed by the Guardian or Committee, Representative, or Temporary Substitute Decision Maker of the patient.					
<b>Trivalent Influenza Vaccine (TIV):</b> <input type="checkbox"/> <b>AGRIFLU®</b> 0.5 mL IM DIN 02346850	<input type="checkbox"/> <b>FLUAD®</b> 0.5 mL IM DIN 02362384	<input type="checkbox"/> <b>FLUAD Pediatric®</b> 0.25 mL IM DIN 02434881	<input type="checkbox"/> <b>FLUVIRAL®</b> 0.5 mL IM DIN 02420686	<input type="checkbox"/> <b>FLUZONE High-Dose®</b> 0.5 mL IM DIN 02445646	<input type="checkbox"/> <b>INFLUVAC®</b> 0.5 mL IM DIN 02269562
<input type="checkbox"/> <b>FLULAVAL™ TETRA</b> 0.5mL IM DIN 02420783	<input type="checkbox"/> <b>AFLURIA® TETRA</b> <input type="checkbox"/> 0.5mL IM pre-filled syringe DIN 02473283 <input type="checkbox"/> 5mL IM multi-dose vial DIN 02473313	<input type="checkbox"/> <b>FLUZONE® QUADRIVALENT</b> <input type="checkbox"/> 0.5mL IM single-dose vial DIN 02420643 <input type="checkbox"/> 5mL IM multi-dose vial DIN 02432730	<input type="checkbox"/> <b>INFLUVAC® TETRA</b> 0.5mL IM DIN 02484854	<input type="checkbox"/> <b>Live Attenuated Influenza Vaccine (LAIV): FLUMIST®</b> 0.1mL per nostril (0.2mL total dose intra-nasally) DIN 02426544	
Date of Immunization (MM/DD/YYYY):	Time of Immunization:	Vaccine Lot #:	Vaccine Expiry (MM/YYYY):	Health Care Provider's Name & License #:	Signature:
Site of Administration: <input type="checkbox"/> Left Arm <input type="checkbox"/> Right Arm <input type="checkbox"/> Intranasal			Contacted Primary Prescriber: <input type="checkbox"/> Yes <input type="checkbox"/> No		Emergency Treatment: <input type="checkbox"/> Yes (see attached) <input type="checkbox"/> No
NS Only	Patient condition before:	Response during:		Response immediately after:	

## Epinephrine Emergency Treatment

Patient's Last Name:	Patient's First Name:	Patient's Date of Birth (MM/DD/YYYY):		
<table style="width: 100%; border: none;"><tr><td style="width: 50%; border: none; vertical-align: top;"><input type="checkbox"/> EpiPen® 0.3mg/0.3mL DIN 00509558 If weight is &gt;30kg or 66 lbs</td><td style="width: 50%; border: none; vertical-align: top;"><input type="checkbox"/> EpiPen® Junior 0.15mg/0.3mL DIN 00578657 If weight is between 15-30kg or 33-66 lbs</td></tr></table>			<input type="checkbox"/> EpiPen® 0.3mg/0.3mL DIN 00509558 If weight is >30kg or 66 lbs	<input type="checkbox"/> EpiPen® Junior 0.15mg/0.3mL DIN 00578657 If weight is between 15-30kg or 33-66 lbs
<input type="checkbox"/> EpiPen® 0.3mg/0.3mL DIN 00509558 If weight is >30kg or 66 lbs	<input type="checkbox"/> EpiPen® Junior 0.15mg/0.3mL DIN 00578657 If weight is between 15-30kg or 33-66 lbs			
Date of Administration (MM/DD/YYYY):	Times of Administration			
Number of Doses Administered:	1.	(if applicable)		
Health Care Provider's Name & License #:	2.	(if applicable)		
Additional Notes (including other emergency measures taken or treatments administered):	3.	(if applicable)		
	Signature:			
	Date (MM/DD/YYYY) & Time of Follow-up with Patient/Agent:			