

CDC 57.133 rev 4, v 6.6

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**Patient Vaccination** 

\* required for saving ^ conditionally required Page 1 of 2 \*Facility ID: \*Event #: \*Patient ID: Social Security #: Secondary ID: Medicare #: Patient Name, Last: Middle: \*Gender: M F Other \*Date of Birth: Ethnicity (Specify): Race (Specify): \*Event Type: FLUVAX \*Influenza subtype: 
Seasonal 
Non-Seasonal \*Date Admitted to Facility: \*Vaccine offered: \( \subseteq \text{Yes} \subseteq \text{No} \) \*Vaccine declined: \( \subseteq \text{Yes} \quad \text{No} \) Reason(s) vaccine declined (Check either section A or B but not both) A. Medical contraindication(s) (check all that apply) B. Personal reason(s) for declining (check all that apply): ☐ Allergy to vaccine components ☐ Fear of needles/injections History of Guillian-Barre syndrome within 6 ☐ Fear of side effects weeks of previous influenza vaccination ☐ Current febrile illness (Temp > 101.5°F) ☐ Perceived ineffectiveness of vaccine Other (specify): Religious or philosophical objections Concern for transmitting vaccine virus to contacts Other (specify): \*Vaccine administered: Yes No ^Date Vaccine Administered: ^Type of influenza vaccine administered: Seasonal: ☐ Afluria® ∐Agriflu® ☐Fluarix® ∐FluLaval® ☐Flumist® Fluzone® Fluzone High-Dose® LOther (Specify): \_\_\_\_\_ ∐Fluvirin<sub>®</sub> Non-seasonal: Other (specify): Live attenuated influenza vaccine (LAIV) e.g., nasal ☐ Inactivated vaccine (TIV) ^Manufacturer: \_\_\_\_\_ ^Lot number: \_\_\_\_\_ ^Route of administration: Intramuscular Subcutaneous Vaccine Information Statement (VIS) Provided to Patient: ☐ Inactivated Influenza VIS ☐ None or unknown Live Attenuated Influenza VIS Edition Date: Person Administering Vaccine: Title: Vaccinator ID: Name: Last: First: Middle: Work Address: State: Zip Code: Assurance of Confidentiality. The voluntarily provided information obtained in this surveillance system that would permit identification of any individual or institution is collected with a guarantee that it will be held in strict confidence, will be used only for the purposes stated, and will not otherwise be disclosed or released without the consent of the individual, or the institution in accordance with Sections 304, 306 and 308(d) of the Public Health Service Act (42 USC 242b, 242k, and 242m(d)). Public reporting burden of this collection of information is estimated to average 10 minutes per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. An agency may not conduct or sponsor, and a person is not required to respond to a collection of information unless it displays a currently valid OMB control number. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to CDC, Reports Clearance Officer, 1600 Clifton Rd., MS D-74, Atlanta, GA 30333, ATTN: PRA (0920-0666).





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Label		Label	
Comments			