Policy & Procedure Title: Administration of Nonviable or Expired Vaccine

Issuing Date: 11/19/2014
Policy & Procedure Number: 22
Revision Date: 04/01/2017
Policy & Procedure Approval Authority: [Signature]

Policy & Procedure Summary

Refrigerators and freezers used to store publicly provided vaccines must be capable of reaching and maintaining the required temperatures established by the vaccine manufactures, the Centers for Disease Control and Prevention (CDC), and the National Institute of Standards and Technology (NIST) standards. If vaccine shipping or storage temperatures are recorded above or below the required temperature range, do not use the vaccine until the viability (potency) of the vaccine has been established by the vaccine manufacturer.

Vaccines that have been deemed nonviable (to have lost potency) due to temperature excursions or vaccines beyond the expiration date should never be administered. If these vaccines are administered inadvertently, doses are not counted as valid and should be repeated. All patients, not only those who received publicly-funded doses, should be revaccinated. Serologic testing to confirm a vaccine response may be performed for certain vaccinations. This practice; however, is generally not recommended when the viability is in question due to a temperature excursion.

The Indiana State Department of Health (ISDH) Immunization Division will make every effort to work with the enrolled providers to address the administration of the nonviable vaccines while balancing clinic needs, cost to patients, providers and health plans, the risk of illnesses or outbreaks, and overall effect on public health.

In the event that a large number of patients have received doses of nonviable vaccine, providers will be placed on “suspend” status and will be terminated for failure to comply with any corrective action plans that are issued.

Policy Statement

All providers enrolled with either the Vaccine(s) for Children (VFC) and/or the Adult Immunization Program must comply with this policy. Providers must also comply with the other ISDH Immunization Division policies for receipt, storage and proper management and handling of publicly funded vaccine.

Determining the Need for Revaccination

In instances where vaccine potency is lost due to improper storage and handling, the decision to issue a recall notification and/or to revaccinate will be made by the Indiana State Health Commissioner or designee and the Immunization Division Director or designee. All final decisions regarding recall notification and/or revaccination will be sent in writing to the responsible clinician and/or medical director at the facility where the vaccines were administered. Upon receipt of this notification, it will be the responsibility of the provider to determine the identity the names and contact information for all patients in need of revaccination.

In isolated situations where expired doses of vaccine are administered, it will be the responsibility of the responsible clinician and primary vaccine coordinator to revaccinate patients per the vaccine administration guidelines in this policy.

Vaccine Administration

All patient(s) should be revaccinated during the clinic visit, if there is viable vaccine available and the error is caught during the clinic visit. If the error is not caught during the clinic visit, the vaccine doses should be repeated as soon as possible. If the nonviable or expired dose is a live virus vaccine, providers must wait at least 4 weeks after the previous dose was given to repeat vaccination.

Assistance Provided by ISDH
The ISDH Immunization Division will offer assistance to enrolled providers to ensure patients are revaccinated in a timely manner. This includes, but is not limited to:

- Technical assistance using CHIRP for recall purposes. This includes marking doses as “subpotent” in the registry.
- On-site assistance to review proper vaccine storage & handling policies and procedures
- Technical assistance with placing additional orders for vaccines
- Technical assistance with developing protocols in conjunction with the clinic, pharmaceutical companies, and Centers for Disease Control & Prevention (CDC)
- Technical assistance with drafting written or verbal patient correspondence

The ISDH field representative will reach-out to the local health department in the county where the enrolled provider site is located to inform them of the incident. The local health department may provide assistance as needed.

The ISDH Immunization Division will document the event in PEAR, including resolution of the issue.

**Serology Titers to Validate Immunity**

The cost of performing serology testing in lieu of revaccination is the responsibility of the enrolled provider site. The ISDH Immunization Division does not offer any technical assistance to conduct post-vaccination serology testing following known temperature excursions.

The ISDH Immunization Division does not recommend serology testing following known temperature excursions. However, an exception could include vaccines for which a series (more than one dose) is indicated and the potency of all doses in the series is questionable. For instance, if an adolescent or adult received three Hepatitis B vaccines during poor storage times, it could be appropriate to give a first potent dose and draw serology tests at the same time. The additional two doses might be waived depending on the test results.

Serology testing exists for some of the vaccines (many are available through clinical labs, but not all). If the provider opts to choose drawing serology titers instead of revaccinating patients, the provider is required to validate the lab is CLIA certified (test is FDA approved and validated by the lab). Providers also need to consider the following:

1) No level of circulating diphtheria or tetanus antibodies confers absolute protection. Diphtheria has been reported in persons with high antibody levels.
2) An adequate immune response from one component in a combination vaccine is not an indication of the potency of the other vaccine components.
3) No data available on obtaining post-vaccine antibody titers for PCV7, MCV4, HPV or Rotavirus vaccines.

**Provider Responsibility**

If a provider declines or is otherwise incapable to recall patients who received questionable doses, ISDH will request a list of affected patients and, in conjunction with the local health department, will conduct its own recall of these patients. In these instances the provider will be asked to replace the nonviable/expired publicly-purchased vaccine with privately purchased stock. Failure to do so will result in permanent termination from all immunization programs with the Indiana State Department of Health.

If a clinic declines to provide a list of affected patients, ISDH will issue a community notice alerting patients that they have received potentially non-viable vaccine at this clinic, and encouraging patients to contact the local health department to explore revaccination. The ISDH may also send notification to the Indiana Attorney General regarding instances of provider non-compliance to ensure the health and well-being of patients is protected.
References & Resources

https://www.cdc.gov/vaccines/hcp/admin/storage/toolkit/storage-handling-toolkit.pdf


Refrigerator/Freezer Temperature Log

Revision History
11/19/2014, Created
04/01/2017, Revised