Adverse events associated with 2010 CSL and other inactivated influenza vaccines

The 2010 trivalent influenza vaccine (TIV) manufactured in Australia by CSL Biotherapies (CSL) caused an excess of febrile reactions, including febrile convulsions, among Australian children.1 A retrospective cohort study conducted in Western Australia found that 57% of 209 children aged 6–59 months who received CSL TIV experienced a febrile reaction, compared with 17% of 110 children who received another TIV (P < 0.0001). The Australian Technical Advisory Group on Immunisation concluded that the rate of febrile convulsions among children vaccinated with the 2010 CSL TIV may have been as high as 1 per 100.2 The Therapeutic Goods Administration (TGA) stated that the cause of these reactions had still not been determined as of 8 July 2011.3

In this article, we consider a common factor in several clusters of adverse events following immunisation (AEFI) in the past 15 years — the use of deoxycholate as the virus-splitting agent in the manufacture of influenza vaccines.

Split-virus and whole-virus vaccines

In the 1960s, the introduction of a manufacturing process to chemically disrupt or “split” influenza viruses successfully reduced the reactogenicity of seasonal vaccines, generally without compromising immunogenicity.4 Inactivated (killed) virus, split through detergent or solvent solubilisation of the lipid membrane, still forms the basis for currently manufactured split influenza virus vaccines, including CSL influenza vaccines.4,5

Compared with whole-virus vaccines, split-virus vaccines have been shown to have a much improved profile for AEFI among all age groups. For example, a 1977 trial found that only 10% of 68 children who received a split-virus vaccine had rectal temperatures of ≥ 100°F compared with 40% of 65 children who received whole-virus vaccine (P < 0.01).6 Similarly, of 333 hospital staff being vaccinated for the first time and receiving a 1989 seasonal TIV, 13% of those who received a split-virus formulation reported generalised aching during the 48 hours after vaccination, compared with 26% of those who received a whole-virus formulation (P < 0.01).7

Inactivation of the influenza virus is mainly achieved using β-propiolactone or formaldehyde.4 CSL uses β-propiolactone to inactivate the virus but is one of only a few manufacturers globally to use deoxycholate (specifically, sodium taurodeoxycholate) as a splitting agent.5

Summary

- The 2010 trivalent influenza vaccine (TIV) manufactured by CSL Biotherapies (CSL) was associated with increased febrile reactions, including febrile convulsions, among Australian children.
- CSL is one of the few manufacturers that use deoxycholate as the virus-splitting agent in the manufacture of TIV. Clusters of adverse events following immunisation (AEFI) have been previously linked to other deoxycholate-split TIV formulations in Europe and Canada.
- We hypothesise that suboptimal virus splitting or other mechanisms related to the use of deoxycholate may have played a role in adverse events linked to the 2010 CSL TIV.
- This hypothesis garners support from a recent United States Food and Drug Administration warning letter indicating that CSL failed to determine optimal splitting conditions for new virus strains and that assays to assess virus splitting had not been validated.
- While there may be other causes, the use of deoxycholate should be further explored. Comprehensive and timely investigations of AEFI, especially those involving children, are necessary to prevent their recurrence and to maintain public confidence in vaccination programs.

Deoxycholate-related clusters of adverse events following immunisation

During the 2000–2001 influenza immunisation campaign in Canada, a novel adverse event — designated oculo respirator syndrome (ORS) — was identified in association with a domestically manufactured split-virus TIV. Although systemic symptoms were also reported, the syndrome was recognised because of allergic-like ocular and respiratory symptoms. Of 960 reported ORS cases, 96% of the 937 cases where the administered vaccine brand was known followed receipt of a domestically produced deoxycholate-split TIV, while only 1% were reported following receipt of an imported TIV that used Triton X-100 (Sigma-Aldrich, St Louis, Mo, USA) as the splitting agent. The precise number of doses administered during the immunisation campaign was not available but, based on case reports and doses distributed, ORS was reported 150 times more frequently in association with the deoxycholate-split vaccine than with the Triton X-100 split vaccine.

In a review of this cluster, the Global Advisory Committee on Vaccine Safety of the World Health Organization summarised that...
an investigation into the manufacturing process and transmission electron microscopy of the vaccine material found that there was variation in the process of disaggregation of virion particles, resulting in a disproportionate number of unsplit virion aggregates in the implicated product.  

This was thought to particularly apply to the A/ Panama/2007/99 H3N2 component that was introduced as a new vaccine strain in 2000–2001. With appropriate remedial action, including the supplementary use of Triton X-100 as a second splitting agent, the problem was largely resolved the following season.  

At that time, ORS was thought to have first emerged in Canada. However, investigations revealed that a similar cluster of adverse events had occurred at least 5 years earlier in several European countries. The European cluster was associated with an influenza vaccine from another manufacturer that used deoxycholate as the splitting agent. Morphological aberrations similar to those identified in the Canadian vaccine were also reported with this vaccine. In Italy, the vaccine was associated with a 10-fold increase in AEFI reported during 1995–1996. In Czechoslovakia, the experience led to a call for systematic evaluation of AEFI that were possibly due to partially disrupted virions, as well as the routine morphological examination of individual vaccine lots as part of release control.  

Differential disruption of virions treated with various splitting agents was suspected as far back as 1984, when it was observed that possibly the subunits produced by sodium deoxycholate are larger than those produced by other disruptive agents and their antigenic properties may, in consequence, approximate that of whole viruses.  

Based on earlier paediatric experience with whole-virus vaccines, inadequately split virions might be expected to trigger febrile reactions in young children, although the mechanism for this is not understood.  

### Investigations into 2010 CSL trivalent influenza vaccine reactogenicity  

An interim report from the TGA published in October 2010 identified an increase in the neuraminidase concentration of the CSL 2010 seasonal vaccine as a possible cause for the reported AEFI. At the time, a number of other potential causes were excluded, including increased haemagglutinin concentration and the presence of live virus, endotoxins or contaminants. The report also included investigations to detect whole virus particles. None were detected, even when the final vaccine product was concentrated by ultracentrifugation. However, the extent to which incomplete splitting was ruled out appeared to be uncertain, with the report acknowledging that “a significant presence of intact viral particles would have been a concern with regard to potential pyrogenicity from whole virus particles.”  

In the United States, CSL TIV was initially approved by the Food and Drug Administration (FDA) in 2007 for use in adults, and in 2009 for use in children aged 6 months and older. On 15 June 2011, the FDA issued a warning letter to CSL. After reviewing the US regulator’s findings, the Australian regulator (the TGA) concluded that, “The two regulators are in agreement over the problems identified at CSL.” The FDA highlighted potential issues with the splitting procedure used by CSL. According to the FDA, sodium taurodeoxycholate lots that failed identification tests at CSL were nonetheless accepted for use. The FDA stated that CSL “failed to determine optimal splitting conditions for new virus strains before the strains [were] used in production” and that “the tests used to evaluate the completeness of virus splitting were “deficient” as the assays used were not “validated for their ability to discriminate between split and whole virus”. These concerns, together with the previous Canadian and European experiences, suggest that incompletely split virus, and perhaps other factors related to the use of deoxycholate, provide a plausible explanation of the AEFI associated with the 2010 CSL TIV.  

### Increased reactogenicity of CSL trivalent influenza vaccine in children before 2010  

While deoxycholate use in vaccine manufacturing was a common factor in several adverse event clusters between 1995 and 2001, it is important to note that no specific concerns have been raised about the safety profile of deoxycholate-split TIV products between 2001 and 2010. However, limited data from two clinical studies suggest that CSL TIV may have been, at least intermittently, associated with increased rates of febrile reactions in children before 2010. The only paediatric trial of CSL TIV published to date reported that 22.5% of participants under 3 years of age experienced fever after receiving the 2005 formulation, but this figure rose to 39.5% after vaccination with the 2006 formulation (P < 0.005); one of the 272 study participants vaccinated in 2006 had a febrile convulsion. Furthermore, data emerging from a randomised trial conducted in the US in 2009 show that 37% of children aged 6 months to 3 years who received a first dose of 2009–2010 CSL TIV manufactured in Australia for the influenza season in North America experienced a febrile reaction, compared with 14% of those who received a first dose of a comparator brand of 2009–2010 TIV that was manufactured using Triton X-100 as the splitting agent (P < 0.0005).  

Implicating deoxycholate in increased reactogenicity associated with 2010 CSL TIV remains speculative. Its mechanism of action may not be limited to the effects of whole or incompletely split virus; the effect on the immune response of increased aggregation and altered antigenic presentation of surface proteins (including neuraminidase) or internal viral components may also be significant. Lipid remnants, and perhaps even residuals of the splitting agent itself, are less likely to be important. Should suboptimal virus splitting secondary to deoxycholate use ultimately be implicated as a factor in the increased reactogenicity of CSL TIV, the problem may be intermittent, and may potentially be solvable, as the deoxycholate parameters can be adjusted to ensure optimal disruption of specific virus strains.
Conclusion

The benefits of TIV to those at risk of severe influenza complications should not be lost in this analysis. Acceptance of population-based vaccination programs depends on public trust in safe manufacturing, together with open and robust regulatory monitoring. Such trust is gradually earned but more easily eroded. A comprehensive and timely explanation of major episodes of AEFI, and the measures taken to prevent recurrences, is a reasonable public expectation.

Competing interests: Heath Kelly has received funding from CSL Biotherapies to attend a study design meeting. Danuta Skowronski was the principal investigator on a clinical trial for which influenza vaccine was provided free by Sanofi Pasteur. Gaston De Serres has previously received research grants for unrelated studies from GlaxoSmithKline and Sanofi Pasteur. The views expressed are ours and may not represent the views of our respective institutions.

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