New drugs

Live Japanese encephalitis vaccine

Approved indication: prevention of Japanese encephalitis

Imojev (Sanofi Pasteur) vials containing one dose of freeze-dried powder for reconstitution

Australian Medicines Handbook section 20.1

The currently approved vaccine for Japanese encephalitis in Australia is an inactivated vaccine given in two doses (Aust Prescr 2009;32:82-6). This new product is a live attenuated vaccine using the yellow fever vaccine virus – strain 17D-204 – as a vector. In this virus, two of the yellow fever genes have been replaced by Japanese encephalitis genes (strain SA14-14-2) encoding a premembrane and an envelope protein. This chimeric virus is propagated in tissue culture before being freeze-dried for use. Vaccination is recommended for people who live in or are travelling to areas where Japanese encephalitis is endemic, such as Papua New Guinea and parts of Asia. It is also recommended for people who work with the virus in laboratories. The vaccine is indicated from 12 months of age as a single subcutaneous injection.

In an early dose-finding trial, most of the 87 adults who were given the live vaccine (1.8–5.8 log10 plaque-forming units subcutaneously) developed neutralising antibodies to the chimeric vaccine strain and, to a lesser degree, to the wild-type Japanese encephalitis strains Beijing-1, Nakama and 902/97. A second dose of the vaccine 30 days later did not boost this immune response.1

In a phase III immunogenicity trial of 820 adults (enrolled from Australia and the USA), one dose (4 log10 plaque-forming units) of the live vaccine was comparable to three doses of an inactivated Japanese encephalitis vaccine. A month after vaccination, 99% of people in the live vaccine group were considered to have protective levels of neutralising antibody to the vaccine strain compared to 74.8% of people in the comparator group.2

In a paediatric trial, almost all children (aged 1–5 years) developed seroprotective antibody levels after a single dose of the vaccine, regardless of whether they had been previously vaccinated with an inactive vaccine. Many of the children developed neutralising antibodies that cross-reacted with other Japanese encephalitis strains.3

It is not currently known if a booster of the live vaccine will be needed, but from longer-term immunogenicity studies it seems antibody responses last for at least four years in adults and six months in children.

In adults, the most common adverse reactions to the vaccine included headache (23.9%), fatigue (21%), malaise (17%), myalgia (14.7%) and injection-site pain (11.8%). These were also common in children as well as irritability (28.5%), loss of appetite (25.9%), fever (20.7%), vomiting (19.2%) and abnormal crying (19.1%).

Vaccination should be postponed in the event of a fever or acute illness. The vaccine is contraindicated in people with impaired cellular immunity, including those on immunosuppressive therapies such as chemotherapy or high-dose steroids (for 14 days or more), and people with symptomatic HIV infection. This vaccine should not be given to women during pregnancy and lactation as there is a theoretical risk that the virus could cross the placenta or be secreted in breast milk.

In children, other vaccines should not be given at the same time. However, in adults concomitant yellow fever vaccine can be administered in the opposite arm or leg. Prescribers should be aware that plasma products such as immunoglobulins could potentially neutralise the live virus so vaccination after receiving blood products should be avoided for at least six weeks.

This live vaccine seems to produce durable immune responses to Japanese encephalitis but its actual efficacy will not be known until after marketing. It may be preferable to the current inactivated vaccine as it is approved for use in children and only one dose is indicated.

Manufacturer provided additional useful information

REFERENCES


The Transparency score (T) is explained in ‘New drugs: T-score for transparency’, Aust Prescr 2011;34:26–7.

* At the time the comment was prepared, information about this drug was available on the website of the Food and Drug Administration in the USA (www.fda.gov).

† At the time the comment was prepared, a scientific discussion about this drug was available on the website of the European Medicines Agency (www.ema.europa.eu).

& At the time the comment was prepared, information about this drug was available on the website of the Therapeutic Goods Administration (www.tga.gov.au/industry/pm-auspar.htm)