Post-exposure passive immunisation for preventing measles
(Protocol)

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Post-exposure passive immunisation for preventing measles

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Editorial group: Cochrane Acute Respiratory Infections Group.


ABSTRACT

This is the protocol for a review and there is no abstract. The objectives are as follows:
To assess the effectiveness and safety of intramuscular injection or intravenous infusion of immunoglobulins for preventing measles when administered to exposed susceptible people before the onset of symptoms.
BACKGROUND

Before vaccination against measles was available, annual case numbers were estimated at 130 million (WHO 1999), and the disease caused between five and eight million deaths globally each year (Moss 2009). With the introduction of the vaccine, the worldwide number of cases began to decline (WHO 1999) and this trend continued with increasing vaccination coverage (WHO 2009a). However, the number of measles cases worldwide exceeded 300,000 in 2010 (WHO 2011) and measles is still an important cause of global mortality as identified by the joint World Health Organization (WHO) and United Nations International Children's Emergency Fund (UNICEF) Global Immunization Vision and Strategy 2005-2015 (WHO 2005). One of the strategy’s goals is measles mortality reduction. In 2008, measles caused around 164,000 deaths (WHO 2009b).

Further to mortality reduction, most WHO regions have set measles elimination goals and reported on progress towards these (Castillo-Solorzano 2011; Martin 2011; Sniadack 2011; WHO 2008). Many countries have noted continued reductions in incidence (WHO 2012a) and even elimination of endemic transmission (Parker Fiebelkorn 2010). However, the WHO cautions that failure to maintain high vaccination coverage in all areas of a country results in resurgence of the disease (WHO 2009a). Certainly, there are many recent published reports of measles outbreaks among countries with high vaccination coverage (CDC 2011a; Delaporte 2011; DVD CDC 2011; Hoskins 2011; Parker Fiebelkorn 2010; Smithson 2010; Takimoto 2011; Vainio 2011) and the WHO confirms that the incidence of measles worldwide increased in 2010 because of large outbreaks in some regions (WHO 2012a).

In countries with low incidences of measles, elimination strategies typically include an urgent response to a single reported case, including confirmation of the diagnosis, contact tracing and post-exposure prophylaxis (CDC 1998; CDNA 2009; NZ MoH 2011; UK DoH 2010). Post-exposure prophylaxis may be a vaccination, which seems to be effective at preventing disease onset if administered within 72 hours of exposure (Barrabéig 2011), or may involve passive immunisation with immunoglobulin, particularly if outside this 72-hour window (Heymann 2008).

Description of the intervention

The practice of passive immunisation against measles has been used since the 1930s (Zingher 2005). Polyclonal immunoglobulins are administered parenterally to susceptible individuals, who have been in contact with an infectious case of measles, in an attempt to prevent the onset of disease or modify disease expression (Keller 2000). Immunoglobulin preparations were originally made from animal sera but today are made from pooled donated human plasma (Gonik 2011). Both intramuscular and intravenous preparations are in use. Product names vary from country to country; so too the concentration of disease-specific immunoglobulins in the products will generally reflect circulating antibody levels in the donating populations (Sawyer 2000). Although, in some countries, minimum neutralising antibody concentrations to measles may be regulated (Sawyer 2000). Current recommendations for dose calculations vary by country, although they are all calculated according to body weight (CDC 2011b; CDNA 2009; ID HPA 2009; NZ MoH 2011). Regardless of the dose recommended, passive immunisation is not currently recommended if more than six days have elapsed since exposure to measles (CDC 2011b; CDNA 2009; ID HPA 2009; NZ MoH 2011).

Description of the condition

Measles is a highly communicable viral illness (Heymann 2008). The measles virus is an enveloped, single-stranded RNA Morbillivirus of the family Paramyxoviridae (Heymann 2008; WHO 2009a). The virus is shed from the respiratory tract of infected persons and transmitted by aerosolised droplets or by direct contact with respiratory secretions (WHO 2009a). Someone with measles is contagious from one day before the symptoms start until four days after the rash appears. A susceptible person exposed to measles will usually develop symptoms after around 10 days but this may range from 7 to 18 days after exposure (Heymann 2008). Symptoms of measles include fever, conjunctivitis, runny nose, cough and a red blotchy rash (WHO 1999). The illness is often more severe in infants and adults than in children (Heymann 2008). Complications occur more frequently in cases in low-income than high-income countries (75% or more versus 10% to 15% of cases, respectively) (WHO 1999). Middle ear infection and pneumonia are fairly common complications, occurring in 5% to 15%, and 5% to 10% of children with measles, respectively (WHO 2009a). Encephalitis is a serious, but rarer, complication of measles, occurring in about 1 out of every 1000 cases (WHO 2009a). A slowly progressing neurological disease, subacute sclerosing panencephalitis (SSPE), very rarely (1 out of 100,000 cases) occurs several years after the original measles infection, most often in children infected with measles under the age of two years (Heymann 2008).

How the intervention might work

Whether injected or infused, the administered immunoglobulins distribute throughout the recipient’s body (Birdsall 2009). The mechanism by which the recipient is protected from disease involves interaction between the immunoglobulins, the invading measles virus particles and the cells and molecules of the recipient’s immune system (Reading 2007). The exact mechanisms by which
viral infectivity is mitigated by antibodies within the body are not comprehensively understood, but vary according to the structure and functionality of the particular antibodies as they encounter the particular virus particles (Reading 2007). In general, measles-specific antibodies bind to invading measles virus particles and this may prevent their entry into cells directly, or trigger other immune mechanisms that result in neutralisation or destruction of the virus (Birdsall 2009; Keller 2000; Reading 2007).

Why it is important to do this review

The effectiveness of post-exposure prophylaxis against measles with immunoglobulins is generally accepted (ATAGI 2008; CDC 1998; NZ MoH 2011; Ramsay 2009). However, effectiveness rates vary considerably among identified reports (King 1991; Ordman 1944; Shepeard 2009; Stokes 1944). Further, national recommendations for the use of post-exposure immunoglobulins for measles differ across a number of Western countries (Best 2011; CDC 1998; CDNA 2009; ID HPA 2009; NZ MoH 2011; Ramsay 2009). Disease incidences (WHO 2012b), immunisation schedules (ATAGI 2008; Gustavo 2008; HPA 2011; NZ MoH 2011), measles-containing vaccine coverage (WHO 2012b) and relevant literature are similar. Differences in immunoglobulin dosage recommendations may reflect differences in the minimum levels of measles-specific antibodies in intramuscular preparations (Best 2011; Ramsay 2009; Sawyer 2000).

We could not identify any systematic review evidence of the effectiveness of post-exposure passive immunisation against measles, nor any systematic review evidence of the minimum effective dosage of immunoglobulin for post-exposure prophylaxis against measles. Recent guidance from the United Kingdom on the required dosage of intramuscular immunoglobulin is based on a single study (Endo 2001; Ramsay 2009).

A systematic review of the evidence of effectiveness of passive immunisation for the post-exposure prophylaxis of measles would help to clarify the effectiveness rate and the minimum effective dose and identify differences in benefit or harm across population groups. These outcomes would be valuable to guide public health practice in countries with low incidences of measles.

OBJECTIVES

To assess the effectiveness and safety of intramuscular injection or intravenous infusion of immunoglobulins for preventing measles when administered to exposed susceptible people before the onset of symptoms.

METHODS

Criteria for considering studies for this review

Types of studies

We will include randomised controlled trials (RCTs), quasi-RCTs and prospective non-RCTs (cohort studies), irrespective of blinding, publication status, language or unit of randomisation. We will include prospective non-RCTs given that more recent studies, using current immunoglobulin preparations, are likely to be non-randomised for ethical reasons. The intervention has been part of public health practice since the 1930s and as such, any RCTs are likely to have been conducted at a time when the antibody levels of blood donors were due to infection with measles rather than vaccination. To inform practice appropriately, any evidence of the effectiveness of current immunoglobulin preparations should be included.

Types of participants

People of any age, sex or ethnic origin who were susceptible (no history of measles and not vaccinated against measles and/or measles immunoglobulin G (IgG) negative) and exposed to measles virus or exposed to someone diagnosed with measles, and who were asymptomatic at the time of intervention or control administration. The primary study’s definition of ‘exposed’ will be accepted and any differences explored via subgroup analysis.

Types of interventions

1. Intervention: intramuscular injection of polyclonal immunoglobulins; intravenous infusion of polyclonal immunoglobulins.
2. Control: no intervention or placebo or live attenuated measles virus vaccine.

We will also include studies assessing different brands or preparations of polyclonal immunoglobulins or different dosages of immunoglobulins. We will only include studies where the intervention (and control) are administered to participants after exposure to measles and before the participants develop measles symptoms.

Types of outcome measures

Primary outcomes

1. Cases of measles. The diagnosis may be made by detection or isolation of measles virus in urine or respiratory secretions; by detection of measles virus antigen in urine or respiratory secretions; by serological detection of immunoglobulin M (IgM) to measles in the absence of vaccination eight days to eight weeks prior to testing; by IgG seroconversion or by a fourfold or greater rise in titre to measles virus in the absence of vaccination eight days to eight weeks prior to testing; or by symptoms consistent
with measles (fever, a red blotchy rash, conjunctivitis, runny nose and cough) or modified measles (prolonged incubation period, milder fever, cough, runny nose, conjunctivitis and sparse discrete rash of short duration).

2. Mortality due to measles.

**Secondary outcomes**

1. Prevention of measles outbreak (higher than expected incidence) as identified by active surveillance.
2. Cessation of measles outbreak (return to expected incidence) as identified by active and/or passive surveillance.
3. Complications due to measles such as otitis media, pneumonia or encephalitis.
4. Occurrence and type of adverse events. We will analyse two types of adverse events: serious adverse events and non-serious adverse events. A serious adverse event is "any untoward medical occurrence that at any dose results in death, is life-threatening, requires inpatient hospitalisation or prolongation of existing hospitalisation, results in persistent or significant disability/incapacity, or is a congenital anomaly/birth defect" (EMEA 1995). We will classify all other events as non-serious. We will specifically extract data on: blood-borne virus infection; anaphylaxis; generalised hypersensitivity; and injection site reactions. We will also include any other adverse event reported as such by study authors.

**Data collection and analysis**

**Selection of studies**

Two review authors (MY, GN) will independently inspect the title and abstract (as available) of each reference identified by the search and determine the potential relevance of each article. If identified by either review author as potentially relevant, we will retrieve the full article. Both review authors will inspect each full article independently, using an eligibility checklist based on the inclusion criteria, to determine inclusion in the review. We will resolve any disagreements through discussion, or by consulting a third review author (AC). We will exclude studies not meeting the eligibility criteria and we will state the reasons for exclusion.

We will list duplicate publications with the main publication for included studies. We will attempt to write to corresponding authors if uncertainties about duplicate publications exist.

**Data extraction and management**

Two review authors (MY, AC) will independently extract data from the included studies using pre-designed data extraction forms. We will resolve disagreement by discussion, or by a third review author (GN), independently extracting the data, as necessary. We will attempt to contact study authors for clarification or further information as necessary.

We will extract the following data.

1. The study
   i) First author, publication year/not published.
   ii) Location.
   iii) Date study undertaken.
   iv) Randomised/quasi-randomised/non-randomised.

2. Participants
   i) Number in each group.
   ii) Age range in each group.
   iii) Proportion of adults, children, infants (aged < one year) in each group.
   iv) Gender distribution in each group.
   v) Proportion of high risk individuals in each group: those with immunodeficiency; pregnancy or age under one year.
   vi) Range of time since exposure in each group.
   vii) Average time since exposure in each group.
   viii) Any measure of baseline comparability and result of this, if calculated.

3. Intervention
   i) Intervention group: product used, concentration of measles antibody if known, volume given, route of administration.
   ii) Control group: placebo/vaccine/product/other, concentration of measles antibody if relevant and known, volume given, route of administration.

4. Outcomes
   i) Primary and secondary (as above).
Assessment of risk of bias in included studies

Two review authors (MY, AC) will independently assess the risk of bias for included studies. We will resolve any disagreements by discussion or by including a third review author (GN) as needed. For randomised and quasi-randomised studies, we will assess: randomisation sequence generation; allocation concealment; blinding of participants, personnel and outcome assessors; incomplete outcome data; drop-out/ selective reporting; and other potential sources of bias. We will report the risk of bias using The Cochrane Collaboration’s tool for assessing ‘Risk of bias’ (Higgins 2011). For non-randomised studies, allocation concealment is not applicable. We will therefore assess: blinding of participants, personnel and outcome assessors; incomplete outcome data; drop-out/ selective reporting; group differences at baseline; management of confounders and other potential sources of bias. We will report the risk of bias using a modified version of the risk of bias tool (Higgins 2011).

Measures of treatment effect

Outcomes, as identified above, are dichotomous. We will express these outcomes as risk ratios (RRs) and calculate 95% confidence intervals (CIs) for each.

Unit of analysis issues

Should cluster-randomised trials be included in the review, we will attempt to extract RRs and 95% CIs resulting from analyses that have accounted for the clustering directly from the paper(s). If this is possible, we will proceed to meta-analyse the data using the inverse variance method. If this is not possible, we will extract the number of clusters, the average size of each cluster, the outcome data at the level of the individuals and an estimate of the intra-cluster correlation coefficient and proceed to reduce the trial(s) to their ‘effective sample size’ for meta-analysis (Higgins 2011). Should studies with multiple intervention groups, for example different doses of immunoglobulins compared to control, be included in the review, we will calculate a weighted average of the relevant pair-wise comparisons from the study and a variance for the study, taking into account the correlation between the comparisons.

Dealing with missing data

We will attempt to contact the trial authors for any missing data. Where missing data exceeds 20%, or where data are clearly missing in different proportions in the treatment groups (P < 0.05), we will exclude the study from further assessment of the relevant outcome. For smaller amounts of missing data, we will undertake sensitivity analysis after assuming worst case and best case scenarios.

Assessment of heterogeneity

We will explore the presence of heterogeneity firstly by comparing studies’ population groups and interventions. If heterogeneity is clear, we will not proceed to meta-analysis. If there is no obvious heterogeneity, we will proceed to meta-analysis and consider the forest plot for each primary outcome and the secondary outcome, “serious adverse events”. If statistical heterogeneity is clear visually, we will proceed to subgroup and sensitivity analyses and re-examine the heterogeneity of these results separately. If heterogeneity is not obvious in the initial forest plots, we will consider the Cochrane Chi² test and I² statistic for each outcome. We will consider an I² estimate of 60% or more, alongside a Chi² P value of 0.1 or less important heterogeneity and again proceed to subgroup and sensitivity analyses. If heterogeneity is not explained by subgroup or methodological differences, we will report the results of a random-effects model for the relevant outcome(s).

Assessment of reporting biases

In the event of multiple publications of the same study, we will list the subsequent papers with the main paper and only enter data once. If uncertainty exists in this respect, we will attempt to contact the study authors. We will assess publication bias by examining funnel plots if we have sufficient studies (at least 10).

Data synthesis

We will calculate the RR and 95% CI for each outcome measured in each study. We will use the fixed-effect model for each primary outcome and the secondary outcome, “serious adverse events” and examine the forest plots to assess heterogeneity. We will then explore possible reasons for apparent heterogeneity via subgroup and sensitivity analyses. If these do not explain the apparent heterogeneity, we will use a random-effects model to analyse the pooled studies and report this result. If heterogeneity is minimal, or explained by subgroup or sensitivity analyses, we will report these using a fixed-effect model.

Even in the absence of apparent significant heterogeneity for the primary outcome of “cases of measles”, we will proceed to subgroup and sensitivity analysis if we have a sufficient number of included studies. We will report the results of the other secondary outcomes descriptively.

Subgroup analysis and investigation of heterogeneity

For subgroup analysis we will include:

- study type;
- age of participants (infants/children/adults/combinations);
- proportion of high risk individuals (pregnant plus immunodeficient plus aged under one year);
- dose of immunoglobulins;
- dose of measles-specific immunoglobulins;
• route of administration of immunoglobulins;
• timing of administration of intervention in relation to exposure; and
• differences in primary study definitions of ‘exposed’ if necessary.

Sensitivity analysis
For sensitivity analysis we will include:
• risk of bias of included studies; and
• studies with imputed missing data.

ACKNOWLEDGEMENTS
We would like to thank Liz Dooley and Clare Dooley for their support and assistance. We also wish to thank the following people for commenting on the draft protocol: Theresa Wangham, Sushil Kabra, Segun Bello, Viviana Rodriguez and Taixiang Wu.

REFERENCES

Additional references

ATAGI 2008

Barrabeig 2011

Best 2011

Birdsall 2009

Castillo-Solorzano 2011

CDC 1998

CDC 2011a

CDC 2011b

CDNA 2009

Delaporte 2011

DVD CDC 2011

EMEA 1995

Endo 2001

Gonik 2011

Gustavo 2008
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Heymann 2008

Higgins 2011

Hoskins 2011

HPA 2011

ID HPA 2009

Keller 2000

King 1991

Martin 2011

Moss 2009

NZ MoH 2011

Ordman 1944

Parker Fiebelkorn 2010

Ramsay 2009

Reading 2007

Sawyer 2000

Sheppeard 2009

Smithson 2010

Sniadack 2011

Stokes 1944

Takimoto 2011

UK DoH 2010

Vainio 2011
WHO 1999

WHO 2005

WHO 2008

WHO 2009a

WHO 2009b

WHO 2011

WHO 2012a

WHO 2012b

Zingher 2005

* Indicates the major publication for the study

**APPENDICES**

Appendix 1. CENTRAL and MEDLINE search strategy

1 exp Measles/
2 exp Measles virus/
3 measles.tw.
4 (rubeola or rubeolla).tw.
5 or/1-4
6 exp Immunoglobulins/
7 (immunoglobulin* or immuno-globulin* or immun* globulin*).tw, nm.
8 (gammaglobulin* or gamma-globulin* or gamma globulin*).tw, nm.
9 exp Immunization, Passive/
10 (passiv* adj2 (immunotherap* or immuni* or antibody transfer*)).tw.
11 Post-Exposure Prophylaxis/
12 ((post exposur* or post-exposur* or postexposur*) adj2 (prophyla* or prevent*)).tw.
13 or/6-12
14 5 and 13
Appendix 2. CENTRAL and MEDLINE filter for study type

We will combine the following filter for non-randomised prospective intervention studies (not before and after and not time-series studies) with the Cochrane Highly Sensitive Search Strategy for identifying randomized trials (Higgins 2011).

1. exp Cohort Studies/
2. Epidemiologic Studies/
3. Intervention Studies/
4. Evaluation Studies/
5. Program Evaluation/
6. Random Allocation/
7. Clinical Trial/
8. Single-Blind Method/
9. Double-Blind Method/
10. Control Groups/
11. Pilot Projects/
12. controlled clinical trial.pt.
13. clinical trial.pt.
15. multicenter study.pt.
16. evaluation studies.pt.
17. Comparative Study/
18. Multicenter Study/
19. Follow-Up Studies/
20. Prospective Studies/
21. (cohort adj (study or studies)).tw.
22. cohort analy*.tw.
23. cohort*.tw.
24. (("follow up" or follow-up) adj (study or studies or assessment)).tw.
25. (observational adj (study or studies)).tw.
26. longitudinal.tw.
27. prospective.tw.
28. ((single or double* or triple* or treb*) and (blind* or mask*)).tw.
29. trial*.tw.
30. placebo.tw.
31. groups.tw.
32. ("pre test" or pretest or pre-intervention or preintervention or "pre intervention" or "post test" or posttest or post-intervention or postintervention or "post intervention").tw.
33. (pre adj5 post).tw.
34. ((evaluat* or intervention or interventional or treatment) and (control or controlled or study or studies or program* or comparison or comparative or "usual care")).tw.
35. ((intervention or interventional or process or program) adj8 (evaluat* or effect* or outcome*)).tw.
36. (program or programme or secondary analyse*).tw.
37. (quasi-experiment* or Quasiexperiment* or "quasi random*" or quasirandom* or "quasi control*" or quasi control* or ((quasi* or experimental) adj3 (method* or study or studies or trial or design*))).tw.
38. random*.tw.
39. (study adj3 aim*).ab.
40. "our study".ab.
41. multivariate.ab.
42. compared.ab.
43. intervention*.ti.
44. pilot.ti.
45. (multicentre or multicenter or multi-centre or multi-center).ti.
46. controlled.ti.
47. (rat or rats or cow or cows or chicken* or horse or horses or mice or mouse or bovine or animal*).ti.
48. exp animals/ not humans.sh.
49. (or/1-46) not (47 or 48)

**HISTORY**


**CONTRIBUTIONS OF AUTHORS**

Dr Megan Young (MY) and Prof Graeme Nimmo (GN) will obtain copies of the studies and select studies for inclusion in the review.

MY and Prof Allan Cripps (AC) will extract the data and assess the risk of bias in the studies.

MY and Dr Mark Jones (MJ) will enter the data.

MJ will analyse the data and together with MY will interpret the analysis.

All authors will complete the final review.

**DECLARATIONS OF INTEREST**

Dr Megan Young is a public health physician in Queensland, Australia who is involved in the public health management of measles.

**SOURCES OF SUPPORT**

**Internal sources**

- Griffith University, Australia.  
In-kind employee time
- University of Queensland, Australia.  
In-kind employee time
- Queensland Health, Australia.  
In-kind employee time

**External sources**

- No sources of support supplied