Mycobacteria other than Mycobacterium tuberculosis (TB) cause a significant burden of disease in children. Non tuberculous mycobacteria (NTM) are free living soil and water organisms known to cause a spectrum of diseases including lymphadenitis, pulmonary disease, skin and soft tissue infections, ear infections, skeletal infections and disseminated infection.

The annual incidence of NTM infections in the developed world is believed to be increasing possibly due to increasing awareness, better identification techniques and changing population groups. However, the magnitude of this problem in children is unquantified. Our study aims to expand knowledge recently gained in Australia through laboratory surveillance.

NTM infections are known to be associated with some medical conditions including human immunodeficiency virus (HIV) infection, malignancy, chronic granulomatous disease (CGD) and chronic lung disease including cystic fibrosis and bronchiectasis. However most often NTM infections occur in otherwise healthy children. Emerging data from recent studies show that even in healthy children, subtle underlying immunodeficiency or genotype differences may exist, contributing to susceptibility to NTM infection.

The natural history of NTM infection has not been well described and optimal management remains unclear. There is evidence that a proportion of NTM lymphadenitis will spontaneously resolve. In children requiring intervention (e.g. due to suppurative changes) surgical clearance has been accepted as the therapy of choice. While surgery is curative in most cases, a proportion of children fail initial surgical management and may require repeated surgery or the addition of medical therapy. The role of medical therapy as first line treatment is unconfirmed. Consensus on medical treatment regimens has not been reached. Information from this study will contribute to efforts to improve the detection and the outcome of affected children.

STUDY OBJECTIVES
1. Estimate the incidence of newly diagnosed NTM infection in children seen by child health specialists in Australia
2. Describe the epidemiology and spectrum of disease and document known risk factors.
3. Describe diagnostic investigations used in Australia; frequency of use of skin testing and the clinical utility of the test, including differential skin testing.

CASE DEFINITION
Please report any child under 15 years of age seen in the previous month newly diagnosed with:

1. DEFINITE NTM:
Any child in whom a non-tuberculous mycobacterium species has been identified either by isolation on culture or by polymerase chain reaction (PCR) from a sample from a sterile site.

OR

2. PROBABLE NTM:
A child who presents with any clinical features compatible with NTM (see below)
AND has undergone one or more of the supportive investigations (see below)
AND in whom Mycobacterium tuberculosis (TB) infection is unlikely.

<table>
<thead>
<tr>
<th>Compatible clinical features</th>
<th>Supportive investigations (one or more)</th>
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<tr>
<td>lymphadenopathy (any site)*</td>
<td><strong>Microbiology</strong>: Acid fast bacilli (AFB) seen on sample or biopsy specimens or AFB grown from non-sterile site sample or positive AFB PCR on non-sterile site sample</td>
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<tr>
<td>pulmonary disease with or without constitutional symptoms b</td>
<td><strong>Histopathology</strong>: Granulomatous inflammation or caseous necrosis or AFB seen</td>
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<tr>
<td>skeletal infection</td>
<td><strong>Skin testing</strong>: Tuberculin PPD skin testing &gt; 5mm and less than 15mm and/or Avian PPD ≥10mm</td>
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<tr>
<td>cutaneous infection</td>
<td></td>
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<tr>
<td>ear disease</td>
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* Clinical features of NTM lymphadenitis includes typical firm LN consistency +/- overlying skin changes (e.g. violaceous hue), with no associated constitutional symptoms

b constitutional symptoms referable to NTM infection include unexplained lethargy, fevers and/or anorexia and weight loss, generally only occurring with disseminated infections or pulmonary infections in chronic lung disease

c Avian PPD, manufactured by Commonwealth Serum Laboratories (CSL) Limited. Intradermal dose 10 IU
FOLLOW-UP OF REPORTED CASES
A brief questionnaire requesting further details will be forwarded to clinicians that report a case of NTM to the APSU.

If you have any comments or questions please contact:
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REFERENCES