Stigma, separation, sorrow: leprosy in Australia

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Abstract. Leprosy (Hansen’s disease) was introduced to Australia in the mid-1800s and its story reflects the attitudes of the 19th and 20th centuries, with treatment including segregation, paternalism, and racism. The approaches taken within the Australian states were similar and based on isolating people affected by leprosy, as both a measure to assist the patient but, more importantly, to protect the European society. The most devastating effects of this introduced disease and these approaches were on Indigenous Australians. With the advent of effective antimicrobials, isolation practices were slowly replaced with community-based treatment. However, the term ‘leper’ still evokes negative images in Australian society today.

Introduction

Leprosy (Hansen’s disease), a treatable disease affecting peripheral nerves, skin and mucous surfaces, is caused by Mycobacterium leprae and Mycobacterium lepromatosis1. As a clinical illness, it only manifests in a small percentage of people who come in contact with infectious patients; however, immunological tests show that most such contacts process the organism without developing clinical symptoms or signs2. The disfigurement caused by the disease has traditionally incited fear of those diseased and has led to exclusion and stigma.

The term ‘leper’ is offensive to persons affected by leprosy and is only used in this article where historically appropriate. The terms ‘leper colony/station’, ‘lazaret’ and ‘leprosarium’ all broadly refer to areas set aside for the segregation of persons affected by leprosy, with the term ‘leprosarium’ indicating a more medical approach to isolation. Derby Leprosarium is now known as Bungarun, but its previous name is used in this article.

Early records

Following European colonisation, the earliest records of leprosy in Australia date from the 1850s, with no evidence of its presence among Indigenous Australians before this time3–5. The disease pattern varied between the colonies. In Victoria, Chinese immigrants on the goldfields were the earliest reported cases, but the disease did not become established in the European settlers. However, in New South Wales and Queensland, early epidemiological studies suggested that the disease spread to Europeans from both Chinese immigrants and South Sea Islanders brought to Australia as indentured labour3–5. Leprosy was introduced in the northern areas of Western Australia during the 1880s, with the probable sources being Chinese labourers or lugger crews (originating from endemic countries) in the pearling industry6. By 1890, leprosy had been reported in the Northern Territory, presumed to have been introduced by Chinese immigrants working on railway construction and in mines4,7. The earliest cases of leprosy in South Australia and Tasmania were reported in the mid-20th century and notifications remained low in these states8. In the early years, leprosy was difficult to diagnose, and records were incomplete and not accurate4.

Quarantine and isolation

In the late 19th century, all Australian states enacted Public Health Acts and all states except Tasmania included leprosy in these Acts or in succeeding legislation specific for leprosy4. Quarantine stations were established at this time, to protect the population from infectious exotic diseases, including leprosy. The fear associated with the disease meant that facilities for people affected by leprosy were sited away from existing quarantine station buildings or were constructed on islands, removed from centres of population.

In New South Wales, The Coast Hospital, Little Bay, Sydney, replaced the North Head Quarantine Station in 1881. Chinese leprosy-affected patients were initially housed in huts, with a lazaret replacing an isolation ward in 1890. A contemporary report describes the lazaret’s surroundings in glowing terms and the inmates as contented9.

In Victoria, the 1897 description of the ‘leper camp’ some distance from the Point Nepean Quarantine Station was less flattering, with the leprosy-affected inmates ‘calmly awaiting the end of their misery … with philosophical resignation’10. In the
In the 1930s, those remaining were moved to Coode Island, Melbourne\textsuperscript{11,12}. In the 1940s, when definitive treatment became available, leprosy-affected persons were sent to the Exotic Diseases Hospital, a stand-alone institution originally built to house sufferers of such diseases as typhus and smallpox, situated next to the Queen’s Memorial Infectious Diseases Hospital, Fairfield (M. Sandland, pers. comm.).

In Queensland, Peel Island Quarantine Station (Figure 1) in Moreton Bay was the successor in 1907 to lazarets at Dayman Island and Friday island in the Torres Strait (for non-Europeans) and Stradbroke Island in Moreton Bay (for Europeans). Peel Island housed both Indigenous Australians and Europeans, but in 1940 the Indigenous Australians were transferred to a separate facility at Fantome Island, North Queensland\textsuperscript{13}.

In Western Australia, early isolation facilities were at Woodman Point and Wooroloo Sanatorium in the south, and Bezout Island, Derby, Cossack and Beagle Bay in the north. In the 1930s, Indigenous Australians affected by leprosy were transported to Channel Island, Darwin, Northern Territory. After years of delay and argument between local communities and Government officials, Derby Leprosarium (Figure 2) was opened in 1936\textsuperscript{16}.

In the Northern Territory (part of South Australia until 1911) the ‘leprosy station’ on Mud Island, Darwin (functioning from 1884) (Figure 3) was described as a ‘living hell lazaret’\textsuperscript{11}. The choice of Channel Island as a replacement in 1931 was only marginally better, with scarce water and fuel supplies and inadequate medical care. It was only in 1955 that a leprosarium was built on the mainland at East Arm\textsuperscript{7}.

In South Australia, the Torrens Island Quarantine Station, established in 1877, records three persons with leprosy admitted to the hospital between 1944 and 1968\textsuperscript{15}.

In Tasmania, the Quarantine Station on Bruny Island (1884–1955) does not appear to have hosted any people affected by leprosy\textsuperscript{16}.

**Indefinite detention challenged**

In the interwar years of the 20th century, lifetime segregation of leprosy-affected people was challenged by EH Molesworth and Leonard Rogers\textsuperscript{17}. Molesworth, an Australian leprologist, sharply criticised the inhumanity of this practice\textsuperscript{18}. Rogers, drawing on international experience, argued that indefinite isolation was ineffective because those in the early stages of the disease would not come forward for treatment if threatened with incarceration and because medical practitioners would be reluctant to expose their patients to such a fate. Patients segregated with advanced disease would not benefit from the treatment offered to them (injected chaulmoogra and hydnocarpus oils, reported to arrest symptoms if administered in the early stages), and those with early symptoms would remain in the community, able to spread leprosy to their (untraced) contacts\textsuperscript{19}. However, CE Cook (Chief Medical Officer of the Northern Territory) and Raphael Cilento (Director-General of Health, Queensland) both opposed these arguments and argued persuasively for isolation. Their views prevailed and indefinite detention practices continued\textsuperscript{13,17}.

**Indigenous Australians**

The earliest cases in Indigenous Australians were reported in the 1890s; by the 1920s, these notifications outnumbered all others\textsuperscript{4}.
Unlike Europeans, Indigenous Australians did not traditionally fear leprosy and did not reject those who were affected. However, in the first half of the 20th century, the leprosy control strategies of State and Commonwealth Governments dictated that all cases should be identified and placed in isolation. Rounded up in police-assisted ‘leprosy raids’ in Queensland, or by ‘leprosy patrols’ in northern Western Australia, becoming a ‘leper suspect’ transported in chains in the Northern Territory, facing lifelong separation from their communities and removal of babies at birth – it was entirely reasonable that Indigenous Australians would make every effort to flee and hide from authorities, or escape from custody. If unsuccessful, they were kept in prison-like conditions, with the prospect of painful injections of chaulmoogra oil and poorly funded, inadequate facilities. A telling statistic in 1940 is the allowance per patient on Fantome Island (£100 per annum), compared with the European patient on the Peel Island Lazaret (£1000 per annum).

The first really effective treatment for leprosy, sulphone therapy, was available in Australia in 1947. In 1953, the first report of the
World Health Organization (WHO) Expert Committee on Leprosy called for a reconsideration of compulsory isolation practices. Australia did not change its policies through the 1950s, although treated patients could then be released from isolation under certain conditions, which included access to medical care, separate accommodation, and no domiciliary contact with children. These conditions automatically excluded most Indigenous Australians, with prevailing attitudes being expressed in this 1952 description by Dr AH Humphry, Commonwealth Department of Health, Darwin: ‘his standard of hygiene is poor, he will not sleep apart, nor can he restrain his intense fondness for children.’

In the late 1950s, European patients in Queensland and Western Australia were beginning to move from Peel Island and Wooloroo to hospital and then home isolation. Yet the facilities for Indigenous Australians did not close until the 1970s (Fantome Island) or the 1980s (Derby, East Arm), with many Fantome Island patients simply transferred to Palm Island. This treatment reflected both official attitudes (that Indigenous Australians were irresponsible with their health) and structural shortfalls in Government health and welfare services for Indigenous Australians.

### Leprosy in Australia today

Triple antibiotic therapy (dapsone, rifampicin and clofazimine) was introduced in the 1980s, and the treatment of leprosy changed to outpatient consultation and monitoring of antimicrobial therapy and any adverse reactions. Responsibilities for leprosy diagnosis and treatment shifted to specific infectious diseases hospitals (e.g. Fairfield Infectious Diseases Hospital, Melbourne) or to major hospitals. Leprosy-affected persons today are treated in outpatient clinics, unless there is a clinical indication for hospitalisation, such as planned corrective surgery or treatment of immune reactions (M. Sandland, pers. comm.).

Since 1925, leprosy notifications have had peaks in 1940, 1944 and 1957 (dominated by Western Australia and the Northern Territory numbers), with occasional reports today, predominantly in those from endemic countries.

From the harrowing descriptions in 1867 of Victorian Chinese immigrants with a ‘loathsome disease’ to the appropriate outpatient treatment in the Northern Territory today, it is clear that Australian approaches to leprosy have undergone radical improvement. Nevertheless, within Australian society in 2020, the term ‘leper colony’ is still used as a description of shame and isolation, an attitude that reflects the ignorance, paternalism and racism in our all too recent past.

### Conflicts of interest

The author declares no conflicts of interest.

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COVID-19 microbiology experience with a difference

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While many microbiologists (and microbiologists-at-heart like myself) turned to nurturing sourdough starters to keep their hands busy and buoy their spirits during the periods of COVID-19-related lockdown in 2020, I turned to the crochet hook. The rhythmic nature of crochet was not only therapeutic, but the feeling of productivity was the perfect accompaniment to morning shifts supervising primary school for my three children. What started as teaching myself to crochet ended up as a collection of the most abundant genus of bacteria found in the oral cavity, plus a couple of disease-associated species. The beady eyes and smiling mouths may be artistic license, but each microbe is crocheted to scale (1 μM = 10 cm). I teach oral microbiology at the University of Melbourne Melbourne Dental School, and these creations will be making an appearance in class next year as my students explore the incredible diversity of microorganisms that call the mouth home.

Quiz

Who am I?

(1) We are considered part of the core microbiome of the mouth, and are some of the earliest colonisers of the teeth. However, some of our genus can cause nasty infections such as meningitis and gonorrhea.
(2) We are some of the earliest bacteria to colonise the mouth after birth, and make up a large proportion of bacteria found at different oral sites. We are a heterogenous genus – some of us are associated with tooth decay by turning the sugars you eat into acid, whereas others can produce alkaline substances that balance this acid out. Under a microscope we are often found in pairs or short chains.
(3) One of my relations is responsible for syphilis, while I am associated with gum disease (periodontitis). I use my periplasmic flagella to get around.
(4) Members of my genus are found in many places in the human body including the gastrointestinal tract and the vagina. Some of us can also be used in the production of fermented dairy products such as cheese and yoghurt.
(5) My extracellular vesicles might look cute, but I am considered one of the major aetiologic agents of gum disease. I may also be related to systemic conditions such as Alzheimer’s disease and cardiovascular disease when I escape the mouth.

Answers available on page 216.