Invasive Group A Streptococcal (iGAS) Disease

CDNA National Guidelines for Public Health Units

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**Summary of revision history**

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| 2.1 | 22 August 2025 | Interim Australian Centre for Disease Control | Amended antimicrobial susceptibility section to avoid misinterpretation. Endorsed by CDNA on 22 August 2025.  |
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**Disclaimer**

These guidelines outline Australia’s national minimum standard for surveillance, laboratory testing, case management and contact management for Invasive Group A *Streptococcal* (iGAS) disease. The intention of these guidelines is to reflect the current available evidence base, with pragmatic guidance provided where evidence is still evolving. Jurisdictions may implement policies that exceed the national minimum standard based on local epidemiological context. CDNA will continue to review and update these guidelines as new information becomes available on iGAS and the situation in Australia.

Readers should not rely solely on the information contained within these guidelines. Guideline information is not intended to be a substitute for advice from other relevant sources including, but not limited to, the advice from a public health specialist or other health professional. Clinical judgement and discretion may be required in the interpretation and application of these guidelines.

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# Summary

## Public health priority

|  |  |
| --- | --- |
| **Priority classification** | **Public health response timeline** |
| High  | For birthing person[[1]](#footnote-1)-neonate pairs and clusters, applicable public health action should be initiated within 1 working day of notification. |
| Routine  | For all other cases, applicable public health action should be initiated within 3 working days of notification.  |

## Case management

Initiate public health response within 1 working day of notification for birthing person-neonate pairs, and within 3 working days of notification for all other cases, both probable and confirmed. Isolate case and practice standard and droplet precautions until 24 hours after initiation of appropriate and effective antibiotic treatment. Exclude case from child-care, school, other educational institutions, or work, until 24 hours after initiation of effective antibiotic treatment.

## Contact management

Provide information to all identified close contacts of both probable and confirmed cases and liaise with treating clinical teams for antibiotics for chemoprophylaxis to be given to eligible close contacts.

# The disease

## Infectious agent

The infectious agent is *Streptococcus pyogenes,* also known as Group A *Streptococcus* (GAS), a Gram positive, ß-haemolytic bacterium.

## Reservoir

Humans.

## Mode of transmission

GAS is spread through direct person-to-person transmission, via droplet spread or direct contact with patients or carriers. Typically, transmission occurs through respiratory droplets but can also occur through contact with secretions (such as saliva, wound discharge, or nasal secretions) from an infected person, or through skin-to-skin contact.

People with GAS disease (e.g., pharyngitis or impetigo) are much more likely to transmit the bacteria to others than asymptomatic carriers. GAS infection is rarely transmitted by indirect contact through objects.

## Incubation period

The incubation period for iGAS is not well defined (1, 2). Cases of iGAS may be preceded by superficial non-invasive GAS infections, such as GAS pharyngitis (incubation period is usually 1 to 3 days) or GAS impetigo (estimated incubation period is 7 to 10 days).

Secondary cases of iGAS infection have been identified up to 30 days after the identification of the initial case, though this is rare (3, 4).

## Infectious period

For contact tracing purposes, iGAS cases are considered infectious from 7 days before onset of GAS-related symptoms until 24 hours after commencement of appropriate antibiotic treatment. The 7 days prior to onset is included to account for a potential period of communicability related to asymptomatic carriage and to capture the source.

## Clinical presentation and outcome

There are a range of clinical presentations related to GAS infections. These include common mild illnesses such as scarlet fever, tonsilitis or pharyngitis (also known as “strep throat”) and skin or soft tissue infections such as impetigo or cellulitis. In rare instances, GAS infections can lead to invasive GAS (iGAS).

Invasive GAS disease is defined by the isolation of GAS from a normally sterile site, such as blood, cerebrospinal fluid, or bone marrow. GAS can enter these sites through a break in the skin (e.g. a cut, puncture, or surgical wound), or via exposure to respiratory or wound secretions from a person carrying the bacteria.

Presentations may include bacteraemia, sepsis, empyema, osteomyelitis, septic arthritis, meningitis, puerperal sepsis, and life-threatening conditions such as streptococcal toxic shock syndrome (STSS) and necrotising fasciitis. Necrotising fasciitis (NF) can lead to life-long complications such as limb-loss and severe scarring, and in approximately 20-30% of cases, death (5). STSS can have similar complications to NF and the case fatality rate is approximately 30% (6). In cases where patients have NF and STSS concurrently, a case-fatality rate of up to 30% has been observed (7).

## People at increased risk of disease

***Birthing person-neonate pairs***

Birthing person-neonate pairs are considered to be the highest risk group for secondary iGAS infection. Though evidence is limited, the relative risk of secondary infections in birthing person-neonate pairs has been estimated at 12-times higher than other close contacts of a case (3).

***Household contacts of a case***

While the available evidence is limited and variable, a range of small studies have found increased risk of secondary iGAS cases in household contacts of cases, ranging from 19 times higher than the general population to over 2,000 times higher (3,8,9).

While an increased risk of infection has been noted in all household contacts of iGAS cases compared to the general population, secondary cases among household contacts are rare (10).

Elderly contacts of a single case in a household or household-like setting have been identified as at higher risk for secondary iGAS infection than other household contacts (3,11).

***Residents or attendees of institutional settings***

Institutions can refer to a range of settings outside the home where people may reside (short-term or long-term) or attend for care or schooling, including, but not limited to:

* Childcare centres
* Aged care or residential care facilities
* Prisons and jails
* Hospitals
* Schools
* Military barracks
* Hostels or shelters

Outbreaks of iGAS infection in aged care facilities, hospitals, and childcare facilities are well-documented in the literature both in Australia and internationally, with aged care facilities being the most frequently reported setting of iGAS outbreaks (12–31).

Other institutional settings where there is likely to be close contact between residents and attendees, high density living with potential for overcrowding, and poor hygiene (such as prisons, military barracks, hostels, and schools) are generally understood to have a higher baseline risk for communicable disease transmission (32–34). However, evidence specific to the increased risk of iGAS transmission in these institutions is not well documented in the literature.

Communities of people experiencing homelessness or utilising safe-injecting rooms or other community-based harm-reduction facilities, can experience increased risk of iGAS infection, and whole genome sequencing has identified connections between apparently sporadic cases in these communities (35–37). However, as these cases are often linked to risk behaviours (such as shared needle use) and environmental exposures rather than facilities, they are referred to in these Guidelines as ‘community outbreaks.’

***Other priority groups***

There are a number of environmental, sociodemographic and health risk factors that can lead groups of people to experience increased risk of iGAS infection, such as:

* Overcrowding in a household setting or other high-density living environments with potential for overcrowding (such as correctional facilities or aged care facilities).
* Receiving wound care in environments or settings with inadequate infection prevention and control practices.
* Frequent skin infections or wounds.
* Shared needle use.
* Poor hygiene environments.
* Chronic disease (particularly diabetes and heart disease).
* Immunocompromising conditions (such as those in receipt of chemotherapy or high-dose steroids).
* Experiencing discrimination in healthcare or barriers to accessing appropriate healthcare services.

Groups who may experience some of these risk factors at a higher rate than the general population include Aboriginal and Torres Strait Islander people, the elderly, people experiencing homelessness, people experiencing poverty, people who inject drugs, and children aged <5 years (15,17,20,27,32,37–44).

Post-surgical, postpartum, and burns patients are also at increased risk of infection; as broken cutaneous or mucosal barriers may facilitate invasive infection after exposure to GAS (8, 10). Acute viral respiratory infections, particularly influenza, are risk factors for developing iGAS (45), and in children, varicella (chickenpox) infection has been noted as a risk factor (46,47).

## Disease occurrence and public health significance

Internationally, the reported incidence in high-resource countries is estimated to be between 2 and 4 cases per 100,000 population, per year (8). At the time of writing, the national incidence of iGAS in Australia is unclear. iGAS became nationally notifiable in Australia from 01 July 2021, and while some jurisdictions had been collecting data on iGAS for up to a decade prior to this time, iGAS did not become notifiable in all jurisdictions until September 2022 (see Table 1).

Table 1: Month and year that iGAS became notifiable in each Australian State and Territory

|  |  |
| --- | --- |
| **Jurisdiction** | **Month and year iGAS became notifiable** |
| Australian Capital Territory | February 2022 |
| New South Wales | September 2022 |
| Northern Territory | May 2011 |
| Queensland | December 2005 |
| South Australia | October 2021 |
| Tasmania | July 2022 |
| Victoria | February 2022 |
| Western Australia | August 2021 |

Several cohort studies, linked data studies and analyses of data collected by jurisdictions provide some insight into the estimated incidence of iGAS in Australia:

* NSW hospitalisation data for iGAS bacteraemia for the period 2010 to 2020 found a hospitalisation rate of 4.0 admissions per 100,000 population, per year in 2017 and rates were highest in people in northern and western NSW. Rates for Aboriginal and Torres Strait Islander people were twice as high as rates in non-Indigenous people (48).
* In the Northern Territory, iGAS notification data from 1 May 2011 to 30 April 2021 found an average rate of 28.2 per 100,000 population per year, and that over half of cases over the period were female (55%). The case fatality rate for the period was 6% and outbreaks were detected on 4 occasions. Among the cases notified during this period, 66% had a pre-existing chronic medical condition (commonly type 2 diabetes or kidney disease) and notification rates in Aboriginal and Torres Strait Islander people were 8 times higher than rates in non-Indigenous people. A high incidence of iGAS among hemodialysis patients (2,205 per 100,000 person-years in the dialysis population) has also been observed in the Northern Territory (49).
* In Queensland, iGAS notification data between 2006 and 2015 found an incidence rate of 4.5 notifications per 100,000 per year. Notification rates in Aboriginal and Torres Strait Islander people were 8 times higher than rates in non-Indigenous people (50).
* A linked data study of identified cases of iGAS in Victoria between 1 January 2007 and 31 December 2017 found a median annual incidence of 3.1 cases per 100,000 population. Of the 88% of cases with hospitalisation information in the study, 33% were admitted to an intensive care unit. The case fatality rate was 5.6% over the study period, reaching 13.5% among those aged ≥75 years. The study also found a sharp increase in cases in 2017, where the incidence rate was the highest for the study period (5.2 cases per 100,000 population per year) (51).
* A cohort study of hospital data from seven major Australian paediatric centres for the period 1 July 2016 to 30 June 2018 estimated a mean annual minimum incidence rate of 1.6 per 100,000 children (aged 18 years and younger) across the study period (6).
* A population-based data linkage study of Western Australian hospital, pathology and deaths data found 2,337 cases of iGAS within the study period (1 January 2000 to 31 December 2018). Age-standardised incidence rates increased from 2 per 100,000 population in 2000 to 9.1 per 100,000 in 2017 (per year, adjusted for age group and sex). The study found that more than half of cases were in boys or men, the median age of cases was 44 years and over a third of cases were in Aboriginal and Torres Strait Islander people. Incidence rates among Aboriginal and Torres Strait Islander people were consistently higher than those for non-Indigenous people across the study period, with rates for Aboriginal and Torres Strait Islander people ranging up to 13 times higher than rates for non-Indigenous people. The study did not find any evidence of seasonal trends (52).
* At the time of writing, there were no published studies or guidelines about the incidence rate of iGAS in the ACT, Tasmania, or South Australia. As iGAS has been notifiable in these jurisdictions for less than a year, preliminary incidence rates from national level data have not been included. Where practicable, these states may wish to use the reported incidence rates of neighbouring jurisdictions as an estimate of their baseline iGAS rates.

Seasonality of iGAS disease at the national level in Australia has not been well-established, though some evidence from Victorian studies suggests cases may increase during periods of increased influenza circulation (53). Internationally, studies from other high-resource countries have found seasonal peaks in winter to early spring (54–57) and, in some instances, differing seasonal peaks depending on *emm* type (58).

Public health management of iGAS and associated conditions can be complex and demanding. This is related to the potential for serious complications and death, the fact that incidence is highest among certain populations, including those with potentially complex sociodemographic factors (such as birthing person-neonate pairs, Aboriginal and Torres Strait Islander people, infants, and the elderly) and that clusters of cases may occur.

# Routine prevention activities

* Primordial prevention aimed at reducing the levels of GAS circulating in the population through improved social determinants of health (including housing, health hardware, and education about basic hygiene practices).
* Institutional settings can reduce risk by following infection prevention and control practices and encouraging basic hygiene practices.
* Primary prevention focusing on the early detection and treatment of GAS infections (throat and skin infections) in line with clinical practice guidelines, including the [Sore Throat Clinical Practice Guidelines](https://www.rch.org.au/clinicalguide/guideline_index/Sore_throat/) and the [National Healthy Skin Guideline](https://infectiousdiseases.telethonkids.org.au/resources/skin-guidelines/).
* There is currently no vaccine available for GAS.

# Surveillance objectives

* To monitor the epidemiology of the disease to inform prevention strategies.
* To promptly identify cases and any high-risk close contacts in order that appropriate public health action can be taken.
* To identify clusters of cases in order that appropriate public health action can be taken.
* To monitor the effectiveness of current control measures and to provide an evidence base for further review of national guidelines.

# Data management

In accordance with the assigned public health priority levels, confirmed and probable cases of iGAS disease should be entered onto the notifiable diseases database within:

* 1 working day for birthing-person neonate pairs, and
* 3 working days for all other cases.

Ensure that data on Aboriginal and Torres Strait Islander status is collected and entered into the jurisdictional database. Typing results and case outcome should be added to the database when available.

# 6. Case definition

Both **confirmed cases** and **probable cases** should be notified.

***Confirmed case***

A confirmed case requires **laboratory definitive evidence**only.

***Probable case***

A probable case requires laboratory suggestive AND clinical suggestive evidence.

***Laboratory definitive evidence***

1. Isolation of Group A Streptococci (*Streptococcus pyogenes*) by culture from a **normally sterile site**.1,2,3

OR

1. Detection of Group A Streptococci (*Streptococcus pyogenes*) by nucleic acid testing from a **normally sterile site**.1,2,3

***Laboratory suggestive evidence***

1. Isolation of Group A Streptococci (*Streptococcus pyogenes*) by culture from a **normally** **non-sterile site**, including deep tissue abscess at procedure or post-mortem.

OR

1. Detection of Group A Streptococci (*Streptococcus pyogenes*) by nucleic acid testing from a **normally** **non-sterile site**, including deep tissue abscess at procedure or post-mortem.

***Clinical suggestive evidence***

Clinical presentation consistent with severe invasive GAS infection4 such as:

* streptococcal toxic shock syndrome (STSS) that includes both hypotension and multi-organ failure
* necrotising fasciitis (NF)
* puerperal and/or neonatal sepsis.

***Case definition notes***

1 Where growth of GAS represents invasion into a normally sterile site and not contiguous spread related to tissue degeneration (such as a deep diabetic ulcer leading to adjacent bone infection). Normally sterile sites include:

blood, cerebrospinal fluid, pleural fluid, peritoneal fluid, pericardial fluid, joint fluid, bone, bone marrow.

internal organs; specimens obtained from surgery or aspirate from one of the following: lymph node, brain, heart, liver, spleen, vitreous fluid, kidney, pancreas, ovary, or vascular tissue.

2 Lung tissue is not a normally sterile site.

3 Interpretation of post-mortem specimens from normally sterile sites should be interpreted with caution, preferably in conjunction with a pathologist and/or clinical microbiologist.

4 The following clinical presentation is not considered sufficient to meet the probable case definition:

An abscess that forms above the pretracheal fascia (e.g. peritonsillar abscess, parapharyngeal abscess).

# 7. Laboratory testing

***Test method***

Bacterial culture is the current gold standard for GAS confirmation, however NAAT assays are also available for blood and sterile tissues and fluids.

Traditional agar plate culture methods (combined with Gram stain which is suggestive but non-specific) rely on incubation in a carbon dioxide added atmosphere to enhance growth.

Identification relies on a combination of traditional methods (no single method is perfect) and includes: (i) Gram stain morphology, (ii) beta-haemolysis on blood agar plates, (iii) Latex agglutination (Lancefield) grouping, differentiating from other species, (iv) bacitracin sensitivity, and (v) PYR testing assay, is used for the detection of pyrrolidonyl arylamidase enzyme (also called pyrrolidonyl aminopeptidase) activity in GAS. MALDI-TOF is also a validated method of identification, as are commercial biochemical panels.

NAAT testing can include commercial molecular panels or in-house detections. One molecular target amplified is the highly conserved sdaB gene, which encodes for DNase B, an extracellular antigen of GAS, and the basis for the anti-DNase B antibody test used to substantiate a likely GAS infection. An alternative target is speB, also highly conserved, and it encodes for streptococcal pyrogenic exotoxin B or SPEB, a cysteine protease, major virulence factor, and superantigen whose expression mediates toxic shock, seen with various syndromes of severe acute pyogenic infections due to GAS.

Both of these target genes are specific for GAS, and both are single gene copies in the *Streptococcus* *pyogenes* chromosome.

Serology (ASOT and Anti-DNAase b) is available and may be useful in assisting diagnosis of GAS immune mediated syndromes such as acute rheumatic fever but is not sufficient for iGAS diagnosis.

***Suitable specimen types***

Blood culture (sterile procedure), cerebrospinal fluid (CSF), body fluid from a normally sterile site (such as syringe aspirate), surgical biopsy material (i.e., fasciitis), pus and swabs (with transport media).

Post-mortem specimens may be collected but interpretation requires consideration of specimen type, disease causing death, and time from death to specimen collection, due to the effects of post-mortem bacterial translocation.

***Specimen collection and handling***

Keep specimens moist and collect into sterile containers. Use sterile saline wrapped gauze if necessary to avoid surgical specimens drying out.

***Test sensitivity***

Nucleic acid testing offers an alternative way to improve speed and accuracy in GAS diagnosis and has been shown to have superior sensitivity and specificity compared to conventional throat cultures and pharyngitis and clinical diagnosis. Similar studies have not been conducted for iGAS and culture positivity rates can be influenced by specimen collection, handling during transport and time to processing.

***Strain differentiation***

Various typing systems are available for epidemiological purposes. GAS has been classically subdivided based upon serotyping of surface-expressed M and major pilus subunit protein T typing. M-typing using specific antisera has been largely replaced by *emm*-typing, by DNA sequencing the variable region of the *emm* gene. While useful it may not be specific enough when common *emm* types occur geographically, hence multi-locus sequence typing (MLST) and the even more specific whole genome sequencing (WGS) are also used. *Emm* type of GAS strains can also be determined from the WGS results.

***Antimicrobial susceptibility***

GAS remains universally penicillin sensitive. However alternative agents are also tested due to patient allergies and include vancomycin, macrolides, trimethoprim/sulfamethoxazole, and clindamycin. Resistance to usually effective antimicrobials can develop, such as clindamycin, to which constitutional or inducible reduced susceptibility may develop.

Testing is classically done by phenotypic disc-based methods, but Minimum Inhibitory Concentration (MIC) testing and broth microdilution are also available. Breakpoints and standardised methods are protocol based on either EUCAST (European Committee on Antimicrobial Susceptibility Testing) or CLSI (Clinical and Laboratory Standards Institute).

Mutations in GAS penicillin binding protein genes have been noted in many countries but have not yet caused penicillin resistance but do confer some reduced susceptibility to other beta lactam antibiotics.

# 8. Case management

## Response times

Case investigation and appropriate public health action for probable and confirmed cases should commence within:

* 1 working day of notification for birthing-person neonate pairs, noting that earlier action should be undertaken where possible. As these cases are managed in a hospital setting, the treating clinical team should implement chemoprophylaxis in a timely manner, with PHU follow up confirming appropriate management has occurred.
* 1 working day of notification for two or more epidemiologically linked cases (see Section 11: *Special Situations*).
* 3 working days for all other cases.

## Case investigation

The response to a notification will usually be carried out in collaboration with the case’s treating clinical team. As such, public health response procedures for both confirmed and probable cases should include the following actions:

* Identify whether the case is part of a birthing person-neonate pair, or works, attends, resides, or may have acquired their infection in an institutional setting or facility (i.e. childcare centre, aged or residential care facility, prison, or hospital).
* Liaise with the treating clinician and/or nursing team to obtain demographic details and determine whether antibiotics for chemoprophylaxis or education materials have been provided to the case and applicable close contacts.
* If proceeding to contact the case and/or next of kin, confirm with the treating clinician whether the case or guardian is aware of the diagnosis.
* Endeavour to provide an appropriate fact sheet to close contacts (Appendix 2). Factsheets may be provided to contacts by the treating clinician, the case or their next of kin.
* If it is identified that the case works, attends, or resides in an institutional setting or facility (particularly residential care facilities or childcare):
	+ Provide information to the institution or facility manager for distribution.
	+ Request the institution or facility report any further iGAS cases occurring within 3 months for residential care facilities and 30 days for all other institutional settings.
	+ Consider undertaking a review of notifications for other iGAS cases that may be linked to the facility or institution.
* Liaise with the treating clinician to facilitate the support of the case and/or family with a social worker, Aboriginal or Torres Strait Islander Liaison Officer or interpreter as required.

## Case treatment

Treatment is the responsibility of the treating clinician. For antibiotic treatment recommendations refer to the current edition of [*Therapeutic Guidelines: Antibiotic*](https://tgldcdp.tg.org.au/topicTeaser?guidelinePage=Antibiotic&amp;etgAccess=true).

## Isolation and restriction

Cases should be managed using appropriate standard and droplet transmission-based precautions until completion of 24 hours of treatment with appropriate antibiotics, as per the [*Australian Guidelines for the Prevention and Control of Infection in Healthcare*](https://www.safetyandquality.gov.au/our-work/infection-prevention-and-control/australian-guidelines-prevention-and-control-infection-healthcare). This is applicable to any setting where the case is being treated, including healthcare facilities and residential care facilities.

# 9. Control of environment

None routinely required.

# 10. Contact management: contacts of a single case

***Identification of contacts***

The aim of identifying contacts is to:

* Identify those who may be the potential source of the GAS strain that has led to a case of iGAS and to identify those who may be at increased risk of having been exposed to that GAS strain.
* Provide contacts with information about iGAS infection and their level of risk aimed at both allaying unnecessary anxiety and advising them of what action to take if they develop symptoms.
* Recommend and coordinate provision of antibiotics for chemoprophylaxis if indicated (Table 2).

***Contact definitions***

For a single case of iGAS, public health follow-up focuses on identifying the subsets of close contacts who require information only, and those who may also require antibiotics for chemoprophylaxis.

Contact groups for which information **and** antibiotics for chemoprophylaxis should be routinely provided:

1. Birthing person-neonate pairs: where either the birthing person or neonate develop iGAS disease during the first 28 days after birth.

Contact groups for which information should be routinely provided:

1. Household or household-like contacts:
2. any person who spent at least 24 hours (cumulative) in the same household or household-like setting with the case during the case’s infectious period or,
3. had sexual relations, or other intimate contact with a case during the case’s infectious period.
4. Institutional contacts:
5. Child-care contacts: any child, staff member or child carer (paid or unpaid) who spent at least 8 hours at the same childcare centre as a confirmed case of iGAS during the case’s infectious period.
6. Other institutions: any resident, attendee or staff member who worked at or resided in the same institution or facility (e.g. residential aged care or disability facility, shelter, dormitory, hostel, prison, military barracks, hospitals etc.) as the case during the case’s infectious period, for at least 24 hours. Particularly, residents sharing a room with a case and any staff member who provided airway or wound care for a case where appropriate transmission-based precautions were not used.

A risk assessment should be undertaken for institutional settings to determine risk areas and inform where public health action is required. Where there is physical separation between sections of an institution or facility with negligible crossover of residents, attendees, and staff, only the residents, attendees and staff in the affected sub-division (e.g. dormitory, hostel room or prison sub-division) will be considered contacts. Wards in healthcare settings where the case was a longer-term patient and/or had substantial interaction with other patients should be considered, noting that those sharing a room on a hospital ward may meet household or house-like contact definition.

1. Airway exposed healthcare workers: healthcare workers who have had unprotected close exposure of their airway to large particle respiratory droplets of a case during airway management (e.g. suctioning, intubation), or mouth to mouth resuscitation within the infectious period. Provision of information regarding risk exposure for airway exposed healthcare workers is the responsibility of the hospital, however PHUs should ensure appropriate management has been undertaken.

These categories have been developed based on the limited evidence available regarding the public health management of close contacts of a single case of iGAS.

***Populations at increased risk***

Some people who fall into the above close contact categories may have additional risk factors that contribute to iGAS infection risk.

According to the limited evidence available (as outlined in Section 2: *Other priority groups* and Section 2: *Disease occurrence and public health significance*) the following groups may be at higher risk at baseline for iGAS infection:

* Aboriginal and Torres Strait Islander people
* elderly people (particularly those aged >75 years)
* children aged <5 years
* people with a chronic or immunocompromising disease or condition
* haemodialysis recipients
* people who inject drugs
* people experiencing homelessness
* people residing at or attending institutions prone to poor hygiene, contact with bodily fluids or overcrowding (such as prisons, hospitals, military barracks etc.)
* other special risk groups unique to states or territories.

These risk factors may be considered by PHUs and treating clinical teams when determining most appropriate management for contacts of a single case. PHUs may also wish to consider severity of disease in these determinations.

***Education***

When close contacts of a case are identified, provide an information fact sheet to these contacts about the disease and how it is spread as soon as possible, noting this may occur via the treating clinician, case or next of kin. Where the PHU is relying on a clinician, case or next of kin to identify and disseminate information to contacts, they should provide advice on close contact identification. It is acknowledged that depending on the method adopted to disseminate information to contacts, there may be variability in completeness of contact identification.

Information in the fact sheet in Appendix 2 can be adapted for the cultural and literacy needs of recipients, but all fact sheets provided to contacts should include the following information and advice:

* close contacts may be at higher risk of developing iGAS disease for up to a 30-day period following last contact with a case,
* symptoms of GAS and iGAS to monitor for,
* if symptoms of GAS develop (e.g. impetigo, pharyngitis), contacts should seek non-urgent medical review,
* if symptoms of iGAS develop, contacts should seek immediate medical attention.

In instances where a single case has occurred in an institutional setting, appropriate fact sheets should be provided to facility management for circulation as appropriate to staff, residents or, in the case of a childcare centre, to parents of attendee children.

***Chemoprophylaxis***

Routine provision of antibiotics for chemoprophylaxis to all close contacts of a single case is generally not recommended, as evidence of the efficacy of this strategy in preventing secondary cases is limited (3).

Based on this limited evidence, these Guidelines suggest routine provision of antibiotics for chemoprophylaxis by PHUs for birthing parent-neonate pairs *only* (Table 2)*.* PHUs may wish to adopt a risk-assessment based approach for all other contact types on a case-by-case basis in cooperation with the treating clinical team, taking into account factors outlined in the *Other priority groups* and *Populations at increased risk* sections of this document.

Decisions to provide antibiotics for chemoprophylaxis to any close contacts of a single case should take into account the benefits and risks including:

* individual risks factors for developing iGAS
* the specific context of a case and their close contacts
* the risk of side effects, and
* other potential consequences of providing antibiotics to asymptomatic persons (i.e., elimination of protective flora, or potential to develop antibiotic resistance).

Antibiotics for chemoprophylaxis should be given to eligible contacts of a single case of iGAS as soon as possible after the contact is identified, preferably within 48 hours of exposure to the original case or, at least, within 48 hours of the case being notified, noting that the utility of administering antibiotics for chemoprophylaxis beyond 10 days of iGAS diagnosis in the initial case is limited. As of August 2023,[*Therapeutic Guidelines: Antibiotic*](https://tgldcdp.tg.org.au/viewTopic?etgAccess=true&guidelinePage=Antibiotic&topicfile=invasive-group-A-streptococcal-infection&guidelinename=Antibiotic&sectionId=toc_d1e129#toc_d1e129) *– Prophylaxis regimens for invasive iGAS infection* indicates that the optimal antibiotic prophylaxis regimen for iGAS infection has not been determined but suitable regimens include:

1. benzathine benzylpenicillin intramuscularly\*, as a single dose#

* adult: 1.2 million units (2.3 mL)
* child less than 10 kg: 0.45 million units (0.9 mL)
* child 10 kg to less than 20 kg: 0.6 million units (1.2 mL)
* child 20 kg or more: 1.2 million units (2.3 mL)

OR

2. cefalexin 1 g (neonate and child: 25 mg/kg up to 1 g) orally, 12-hourly for 10 days.

\* The ventrogluteal site is preferred for administration of intramuscular benzathine benzylpenicillin because of reduced pain and risk of nerve injury.

For instructions on intramuscular injection at the ventrogluteal site, see [Figure 2.57](https://tgldcdp.tg.org.au/viewTopic?etgAccess=true&guidelinePage=Antibiotic&topicfile=using-antibacterial-drugs&guidelinename=Antibiotic&sectionId=toc_d1e324#toc_d1e324)

# It is unclear if eradication of pharyngeal group A *streptococcus* carriage is required to prevent secondary cases. Limited evidence suggests the

addition of rifampicin to benzathine benzylpenicillin increases the rate of pharyngeal carriage eradication.

However, the role of rifampicin in the prevention of secondary invasive group A *streptococcal* infection is uncertain, and routine combination

prophylaxis is not recommended.

For close contacts with delayed non-severe hypersensitivity to penicillins, cefalexin can be used in most cases^.

For close contacts with immediate (non-severe or severe) or delayed severe hypersensitivity to penicillins, antibiotic choice depends on the susceptibility of the isolate from the index case

(as rates of resistance to non–beta-lactam antibiotics are higher). If susceptibility results are not available, a reasonable regimen is:

* azithromycin 500 mg (child: 12 mg/kg up to 500 mg) orally, daily for 5 days.

^ It is safe to use cefalexin in patients who had a delayed non-severe reaction to a penicillin in the distant past. It is also safe to use cefalexin in

Patients who have had a delayed non-severe reaction recently, unless the reaction involved amoxicillin or ampicillin, because cross-reactivity

between these drugs is possible. For patients who have had a recent delayed non-severe reaction to amoxicillin or ampicillin, use the drug recommended for patients with immediate (non-severe or severe) or delayed severe hypersensitivity.

Recommendations in these Guidelines regarding the provision of antibiotics for chemoprophylaxis to close contacts of a single case are not intended as a substitute for the expert knowledge of treating clinical teams, nor are they intended to override jurisdictional best-practice in accordance with their specific populations and contexts. Decisions about antibiotics for chemoprophylaxis must always take into account the individual and population risks and benefits of this intervention.

Table 2: Recommended public health responses for close contacts of a single case of invasive Group A *Streptococcus* infection, by contact type\*

|  |  |  |
| --- | --- | --- |
| Contact group  | Routine provision of antibiotics for chemoprophylaxis recommended?  | Provision of fact sheet recommended?^ |
| Birthing-parent neonate pairs | YES | YES |
| Other household or household-like contacts | NO# | YES |
| Institutional contacts | NO# | YES |
| Airway-exposed healthcare workers~ | NO  | YES |

\*PHUs should be aware of and implement public health jurisdictional guidelines, where these may be different from the CDNA guidelines. It is not the responsibility of Public Health Units (PHUs) to undertake contact management based on guidelines of individual hospitals if these differ from public health jurisdictional or CDNA guidelines.

# Chemoprophylaxis may be considered by PHU and treating clinicians for contacts with additional individual risk factors on a case-by-case basis.

^Fact sheets may be provided by PHUs for distribution to contacts to the following groups: cases, treating clinical teams of a case, family members/next of kin of a case and facility management for distribution to institutional residents or attendees.

~Where these occur in a hospital setting, contact management of airway-exposed healthcare workers should be undertaken by the hospital, however PHUs should ensure appropriate management has been undertaken.

# 11. Special situations

## Household clusters

***Definition***

Where two or more cases of iGAS infection occur in the same household or household-like setting within 30 days of symptom onset in the initial case.

***Management***

In the event of a household cluster, the entire household should be offered antibiotics for chemoprophylaxis as per [*Therapeutic Guidelines: Antibiotic*](https://tgldcdp.tg.org.au/viewTopic?etgAccess=true&guidelinePage=Antibiotic&topicfile=invasive-group-A-streptococcal-infection&guidelinename=Antibiotic&sectionId=toc_d1e129#toc_d1e129) *– Prevention of invasive group A streptococcal infection* and provided with the appropriate fact sheet. This includes all persons ordinarily resident in the affected household regardless of whether they were identified as a contact of a case.

Where a secondary case is identified after the initial case has completed a course of antibiotics, the initial case does not require additional antibiotics for chemoprophylaxis.

Specimens from household clusters should be sent to a reference laboratory for molecular typing. The administration of antibiotics for chemoprophylaxis to the household should not be delayed while waiting for molecular typing results.

## Clusters in institutional settings

Where two or more cases of iGAS infection occur in an institution within 3 months of onset in the initial case.

Institutions can include, but are not limited to:

* Childcare centres
* Schools
* Aged care or residential care facilities
* Prisons
* Hospitals
* Hostels
* Military barracks.

For clusters in hospitals, utilise the below guidance in conjunction with hospital guidance or applicable jurisdictional guidance for special risk groups in healthcare (i.e. dialysis patients in the Northern Territory).

***Definition***

Confirmed cluster: 2 or more cases that are epidemiologically linked within an institution that occur within a 3-month period and are identical on molecular typing where cases are not household contacts of each other.

Possible/suspected cluster: 2 or more cases that are epidemiologically linked within an institution that occur within a 3-month period where cases are not household contacts of each other.

Epidemiological link: Where cases occur in a physical or geographical context and a plausible mode of transmission accounts for infection spreading between people.

***Management***

For the purposes of initiating public health management, it is preferable that a cluster be confirmed via molecular typing prior to response. However, if there is strong epidemiological evidence of a cluster or, if staff members, residents or other attendees of an institution belong to a group at increased risk of disease (see *People at increased risk of disease*), a public health response may be initiated prior to receiving results for a suspected cluster or outbreak.

Coordinating the provision of antibiotics for chemoprophylaxis, in a both confirmed and suspected institutional cluster other than in a healthcare facility, is the responsibility of PHUs. PHUs may engage the assistance of any clinicians working within an institution for provision and dissemination of antibiotics. Whilst the PHUs coordinate the overall contact management strategy, PHUs should liaise with staff health or infection control teams (where they exist).

Hospitals should take primary responsibility for implementing prevention and control measures for their facility, including identification of acquisition, and follow-up of staff, patients, and visitors. Depending on respective workloads and resources, PHUs may be able to assist hospitals with follow-up of patients discharged to the community.

In initiating a public health response to a cluster or outbreak, PHUs should:

* Consider convening an Outbreak Management team, including but not limited to infection control advisors, surveillance officers, epidemiologists, public health physicians, and any primary care providers for the facility and facility management staff.
* Identify contacts of cases and provide antibiotics for chemoprophylaxis, where:
	+ a cluster or outbreak is confirmed via molecular typing, or there is clear and strong epidemiological evidence of transmission within the facility, all asymptomatic residents and staff should be offered antibiotics for chemoprophylaxis simultaneously.
	+ linked cases have occurred within a subdivision of the facility and there is negligible crossover of residents and staff between subdivisions, antibiotics for chemoprophylaxis can be restricted to those within the affected subdivision.
* Review public health surveillance databases and information provided by facility management to establish if cluster cases are linked to other previously notified cases.
* Send isolates to reference laboratory for molecular typing.
* Consider communicating information about the cluster to local healthcare providers.

PHUs should engage institutional facility management to:

* Distribute appropriate fact sheets and education materials to staff and residents of the facility, and, in the case of childcare facilities, parents of attending children.
* Review cleaning, hygiene, and infection control practices, including the implementation of appropriate transmission-based precautions for cases and their carers or staff caring for the case in health or residential facilities.
* Establish enhanced surveillance of the facility for further linked cases. This period may vary but should be for at least 3 months after the most recent case was diagnosed. This should include educating facility management on symptoms and signs of iGAS infection and establishing a process by which facility management can notify the appropriate public health service.

Screening asymptomatic people for GAS carriage through testing is not routinely recommended but may be considered as part of a cluster investigation in an institutional setting depending on the risk factors for residents, staff, or attendees.

***Restriction***

Restriction of asymptomatic contacts and screening for GAS carriage is not routinely recommended, including in institutional settings. If a staff member working in an institutional setting has been screened and identified as a GAS carrier during a cluster investigation, the staff member should be excluded from work until 24 hours after initiation of appropriate and effective antibiotic treatment.

## Other community clusters

Clustering of iGAS cases in community settings outside institutions and healthcare settings present distinct challenges for public health response. Community clusters can occur in a range of settings and contexts, including universities, sports clubs (where the sport involves close contact), among people who inject drugs (PWID) and among people experiencing homelessness (PEH).

Stigma, marginalisation, and criminalisation of injecting drug use and homelessness are a challenge to effective engagement with PWID and/or PEH. It is important to keep this in mind when responding to any increase in cases among this population.

Contacts within PEH and PWID populations who have open wounds or lesions are at higher risk for transmission. Contact tracing may be challenging among some PWID as individuals may not be willing to provide contact information for their peers. PHUs are encouraged to engage with any community services used by PWID and/or PEH cases to identify and provide appropriate treatment and information to contacts. This engagement also serves the purpose of determining whether a case has been linked to sheltered accommodation, a drug service or specific injecting network, prison setting, healthcare setting or other institution in the 7 days prior to onset of symptoms.

Where high-risk close contacts of cluster cases are identified in PWID of PEH populations (e.g. close contacts with chronic conditions, current open wounds or with whom the case has shared needles), liaise with community services to organise provision of antibiotics for chemoprophylaxis.

## Elevated rates within the population with no cluster identified

Each state and territory should monitor rates of iGAS and compare against a relevant baseline to determine if iGAS rates are increasing. These rates may be different for different regions, and tolerable thresholds for increase may be lower for populations at increased risk of disease (see *People at increased risk of disease*). The baseline rates can be drawn from jurisdictional surveillance data or research (as per Section 2*: Disease occurrence and public health significance*). PHUs should, after considering available data, establish a threshold for increase from baseline rates appropriate for their population at which public health action should be triggered. If this threshold is met, the following actions should be considered by the PHU:

* Publishing a clinician’s alert
* Releasing a media alert, with links to relevant fact sheets
* Other community messaging, with the aim to both inform and reassure
* Clinician education

All alerts and messaging should serve to both inform the public of the increase in rates but also to alleviate excess anxiety.

## Aboriginal and Torres Strait Islander people

Available Australian research studies and surveillance data have established that the rate of iGAS infections is higher in Aboriginal and Torres Strait Islander people than in the non-Indigenous population (see *Disease occurrence and public health significance*). Aboriginal and Torres Strait Islander people are at increased risk of iGAS transmission and severe outcomes due to a number of intersectional risk factors, including:

* Crowded and inadequate housing,
* Barriers to accessing appropriate and timely healthcare, including limited access to culturally appropriate healthcare, institutional racism, mistrust of mainstream health services and remoteness; and
* Community burden of disease, including higher rates of comorbid and immunocompromising diseases and conditions.

As such, a lower threshold, appropriate to the needs and circumstances of Aboriginal and Torres Strait Islander populations in each jurisdiction, should be used to initiate disease control measures if a cluster is suspected in an Aboriginal and/or Torres Strait Islander household, household-like setting, or community.

***Household cases and clusters***

For Aboriginal and Torres Strait Islander people, disease risk needs to be communicated in a culturally safe manner so individuals and families understand the recommended management plan and contribute to how this, and contact tracing, can be actioned appropriately.

Consider referring household and household-like contacts to their Aboriginal Community Controlled Health Service, Aboriginal Medical Service or local health providers for ongoing assessment and follow up if GAS symptoms develop for culturally appropriate follow-up. If iGAS symptoms develop in a household contact of a case, culturally appropriate medical attention should be sought immediately.

Culturally appropriate educational resources should be used where needed.

***Community clusters***

Further investigation should be undertaken by the local PHU if increases in notification numbers and/or rates are identified for an Aboriginal and/or Torres Strait Islander community. The trigger for investigation should be guided by the community size, composition, and underlying risk factors. The nature of any action will depend on a number of factors, including the size of the community.

Local Aboriginal Health Workers and Practitioners and local Aboriginal Community Controlled Health Organisations are key stakeholders and should be included in the development of community cluster responses.

# References

1. Communicable Disease Control Invasive Group A *Streptococcal* Disease 2017. Available from: <http://www.bccdc.ca/resource-gallery/Documents/Guidelines%20and%20Forms/Guidelines%20and%20Manuals/Epid/CD%20Manual/Chapter%201%20-%20CDC/iGAS.pdf>
2. Group A *Streptococcal* Infections, Invasive (iGAS). Northwest Territories, Canada 2021 Available from: <https://www.hss.gov.nt.ca/professionals/sites/professionals/files/resources/cdc-invasive-group-a-streptococcal.pdf>
3. Mearkle R, Saavedra-Campos M, Lamagni T, Usdin M, Coelho J, Chalker V, et al. Household transmission of Invasive Group A *Streptococcus* infections in England: a population-based study, 2009, 2011 to 2013. Eurosurveillance. 2017 May 11;22(19). Doi: 10.2807/1560-7917.ES.2017.22.19.30532
4. Heymann DL, American Public Health Association. Control of communicable diseases manual: an official report of the American Public Health Association. Washington, DC: American Public Health Association; 2008.
5. Nawijn F, Smeeing DPJ, Houwert RM, Leenen LPH, Hietbrink F. Time is of the essence when treating necrotizing soft tissue infections: a systematic review and meta-analysis. World Journal of Emergency Surgery. 2020 Jan 8;15(1). Doi: 10.1186/s13017-019-0286-6
6. Oliver J, Thielemans E, McMinn A, Baker C, Britton PN, Clark JE, et al. Invasive Group A *Streptococcus* disease in Australian children: 2016 to 2018 – a descriptive cohort study. BMC Public Health. 2019 Dec 30;19(1). Doi: 10.1186/s12889-019-8085-2.
7. CDC. Necrotizing Fasciitis: Acting Fast Is Key. Centers for Disease Control and Prevention. 2019 [viewed 17 March 2023]. Available from: <https://www.cdc.gov/groupastrep/diseases-public/necrotizing-fasciitis.html>
8. Steer AC, Lamagni T, Curtis N, Carapetis JR. Invasive Group A *Streptococcal* Disease. Drugs. 2012 Jun;72(9):1213–27. Doi: 10.2165/11634180-000000000-00000
9. Carapetis JR, Jacoby P, Carville K, Ang SJ., J, Curtis N, Andrews R. Effectiveness of Clindamycin and Intravenous Immunoglobulin, and Risk of Disease in Contacts, in Invasive Group A *Streptococcal* Infections. Clinical Infectious Diseases. 2014 Apr 29;59(3):358–65. Doi: 10.1093/cid/ciu304.
10. Prevention of Invasive Group A *Streptococcal* Disease among Household Contacts of Case Patients and among Postpartum and Postsurgical Patients: Recommendations from the Centers for Disease Control and Prevention. Clinical Infectious Diseases. 2002 Oct 15;35(8):950–9. Doi: 10.1086/342692
11. Adebanjo T, Apostol M, Alden N, Petit S, Tunali A, Torres S, et al. Evaluating Household Transmission of Invasive Group A *Streptococcus* Disease in the United States Using Population-based Surveillance Data, 2013–2016. Clinical Infectious Diseases. 2019 Apr 1; 70(7). Doi: 10.1093/cid/ciz716.
12. Steer JA, Lamagni T, Healy B, Morgan M, Dryden M, Rao B, et al. Guidelines for prevention and control of Group A *Streptococcal* infection in acute healthcare and maternity settings in the UK. Journal of Infection . 2012 Jan;64(1):1–18. Doi: 10.1016/j.jinf.2011.11.001
13. Saavedra-Campos M, Simone B, Balasegaram S, Wright A, Usdin M, Lamagni T Estimating the risk of Invasive Group A *Streptococcus* infection in care home residents in England, 2009–2010. Epidemiology and Infection. 2017 Aug 14;145(13):2759–65. Doi: 10.1017/s0950268817001674
14. Cummins A, Millership S, Lamagni T, Foster K. Control measures for Invasive Group A *Streptococci* (iGAS) outbreaks in care homes. Journal of Infection. 2012 Feb;64(2):156–61. Doi: 10.1016/j.jinf.2011.11.017
15. Nanduri SA, Metcalf BJ, Arwady MA, Edens C, Lavin MA, Morgan J, et al. Prolonged and large outbreak of Invasive Group A *Streptococcus* disease within a nursing home: repeated intrafacility transmission of a single strain. Clinical Microbiology and Infection. 2019 Feb;25(2): 248.e1–7. Doi: 10.1016/j.cmi.2018.04.034
16. Worthing KA, Werno A, Pink R, McIntyre L, Carter GP, Williamson DA, et al. Biphasic Outbreak of Invasive Group A *Streptococcus* Disease in Eldercare Facility, New Zealand. Emerging Infectious Diseases. 2020 May 1;26(5):841–8. Doi: 10.3201/eid2605.190131
17. ‌ Ahmed S, Diebold K, Brandvold JM, Ewaidah S, Black S, Ogundimu A, et al. The Role of Wound Care in two Group A *Streptococcal* Outbreaks in a Chicago Skilled Nursing Facility, 2015‒2016. Open Forum Infectious Diseases. 2018 Jun 26; 5(7):ofy145. Doi: 10.1093/ofid/ofy145
18. Auerbach SB. Outbreak of Invasive Group A *Streptococcal* Infections in a Nursing Home. Archives of Internal Medicine. 1992 May 1;152(5):1017.
19. Thigpen MC, Thomas DM, Gloss D, Park SY, Khan AJ, Fogelman VL, et al. Nursing Home Outbreak of Invasive Group A *Streptococca*l Infections Caused by 2 Distinct Strains. Infection Control & Hospital Epidemiology. 2007 Jan;28(1):68–74. Doi: 10.1017/S0195941700045677
20. Greene CM, Chris Van Beneden, Javadi M, Skoff TH, Beall B, Facklam RR, et al. Cluster of deaths from Group A *Streptococcus* in a long-term care facility, Georgia, 2001. American Journal of Infection Control. 2005 Mar 1;33(2):108–13. Doi: 10.1016/j.ajic.2004.07.009
21. Daneman N, Green KA, Low DE, Simor AE, Willey B, Schwartz B, et al. Surveillance for Hospital Outbreaks of Invasive Group A *Streptococcal* Infections in Ontario, Canada, 1992 to 2000. Annals of Internal Medicine. 2007 Aug 21;147(4):234. Doi: 10.7326/0003-4819-147-4-200708210-00004
22. Chalker VJ, Smith A, Al-Shahib A, Botchway S, Macdonald E, Daniel R, et al. Integration of Genomic and Other Epidemiologic Data to Investigate and Control a Cross-Institutional Outbreak of *Streptococcus* pyogenes. Emerging Infectious Diseases. 2016 Jun;22(6):973-980. Doi: 10.3201/eid2206.142050
23. Rainbow J, Jewell B, Danila R, Boxrud D, Beall B, Chris Van Beneden, et al. Invasive Group A *Streptococcal* disease in nursing homes, Minnesota, 1995-2006. Emerging Infectious Diseases. 2008 May 1; 14(5):772-777. Doi: 10.3201/eid1405.0704072.
24. Vasant BR, Jarvinen KAJ, Fang NX, Smith HV, Jennison AV. Mass prophylaxis in an outbreak of Invasive Group A *Streptococcal* disease in a residential aged care facility. Communicable Diseases Intelligence. 2019 May 15;43.
25. ‌Mahida N, Beal A, Trigg D, Vaughan N, Boswell T. Outbreak of Invasive Group A *Streptococcus* infection: contaminated patient curtains and cross-infection on an ear, nose and throat ward. Journal of Hospital Infection. 2014 Jul;87(3):141–4. Doi: 10.1016/j.jhin.2014.04.007
26. Dickinson H, Reacher M, Nazareth B, Eagle H, Fowler D, Underwood A, et al. Whole-genome sequencing in the investigation of recurrent invasive Group A *Streptococcus* outbreaks in a maternity unit. Journal of Hospital Infection. 2019 Mar;101(3):320–6. Doi: 10.1016/j.jhin.2018.03.018
27. ‌Hancock-Allen JB, Janelle SJ, Lujan K, Bamberg W. Outbreak of Group A *Streptococcus* infections in an outpatient wound clinic—Colorado, 2014. American Journal of Infection Control. 2016 Oct 1; 44(10):1133-1138. Doi: 10.1016/j.ajic.2016.03.058
28. Outbreak of Invasive Group A *Streptococcus* associated with varicella in a childcare center -- Boston, Massachusetts, 1997. PubMed. 1997 Oct 10;46(40):944–8.
29. Agüero J, Ortega-Mendi M, Eliecer Cano M, González de Aledo A, Calvo J, Viloria L, et al. Outbreak of Invasive Group A *Streptococcal* Disease Among Children Attending a Day-Care Center. Pediatric Infectious Disease Journal. 2008 Jul;27(7):602–4. Doi: 10.1097/INF.0b013e31816a0e0a.
30. Collins JP, Shane AL. Infections Associated with Group Childcare. Principles and Practice of Pediatric Infectious Diseases . 2018;25-32.e3.
31. Ortega-Mendi M, Martínez-Martínez L, González de Aledo-Linos A, Agüero-Balbín J, Viloria-Raymundo L, Cano-García ME et al. Outbreak of *streptococcal* toxic shock syndrome in a day care center in Cantabria, Spain, 2006. Revista Espanola de Salud Pubica. 2008 Jan-Feb; 82(1):81-89. Doi: 10.1590/s1135-57272008000100007
32. Riccardo F, Suk J, Espinosa L, Bella A, Giambi C, Del Manso M, et al. Key Dimensions for the Prevention and Control of Communicable Diseases in Institutional Settings: A Scoping Review to Guide the Development of a Tool to Strengthen Preparedness at Migrant Holding Centres in the EU/EEA. International Journal of Environmental Research and Public Health. 2018 May 30;15(6):1120. Doi: 10.3390/ijerph15061120
33. Enggist S, Organisation Mondiale De La Santé. Bureau Régional De L'europe, Al E. Prisons and health. Copenhagen: WHO Regional Office For Europe; 2014.
34. Simpson PL, Simpson M, Adily A, Grant L, Butler T. Prison cell spatial density and infectious and communicable diseases: a systematic review. BMJ Open. 2019 Jul 1;9(7):e026806 Doi: . 10.1136/bmjopen-2018-026806.
35. Dickson C, Pham M, Nguyen V, Brubacher C, Silverman M, Khaled K, et al. Community outbreak of Invasive Group A *Streptococcus* infection in Ontario, Canada. Canada Communicable Disease Report. 2018 Jul 5;44(7/8):182–8.
36. Bundle N, Bubba L, Coelho J, Kwiatkowska R, Cloke R, King S, et al. Ongoing outbreak of invasive and non-invasive disease due to Group A *Streptococcus* (GAS) type *emm*66 among homeless and people who inject drugs in England and Wales, January to December 2016. Eurosurveillance. 2017 Jan 19;22(3). Doi: 10.2807/1560-7917.ES.2017.22.3.30446
37. Vasylyeva TI, Smyrnov P, Strathdee S, Friedman SR. Challenges posed by COVID‐19 to people who inject drugs and lessons from other outbreaks. Journal of the International AIDS Society. 2020 Jul;23(7):e25583. Doi: 10.1002/jia2.25583
38. ‌ Valenciano SJ, Onukwube J, Spiller MW, Thomas A, Como-Sabetti K, Schaffner W, et al. Invasive Group A *Streptococcal* Infections Among People Who Inject Drugs and People Experiencing Homelessness in the United States, 2010-2017. Clinical Infectious Diseases . 2021 Dec 6 ;73(11):e3718–26. Doi: 10.1093/cid/ciaa787.
39. Langley G, Hao Y, Pondo T, Miller L, Petit S, Thomas A, et al. The Impact of Obesity and Diabetes on the Risk of Disease and Death due to Invasive Group A *Streptococcus* Infections in Adults. Clinical Infectious Diseases. 2015 Dec 23;62(7):845–52. Doi: 10.1093/cid/civ1032
40. Boyd R, Patel M, Currie BJ, Holt DC, Harris T, Krause V. High burden of Invasive Group A *Streptococcal* disease in the Northern Territory of Australia. Epidemiology and Infection. 2015 Sep 14;144(5):1018–27. Doi: 10.1017/S0950268815002010
41. Factor SH, Levine OS, Schwartz B, Harrison LH, Farley MM, McGeer A, et al. Invasive Group A *Streptococca*l Disease: Risk Factors for Adults. Emerging Infectious Diseases. 2003 Aug;9(8):970–7. Doi: 10.3201/eid0908.020745
42. ‌ Olufon O, Iyanger N, Cleary V, Lamagni T. An outbreak of Invasive Group A *Streptococcal* infection among elderly patients receiving care from a district nursing team, October 2013 – May 2014. Journal of Infection Prevention. 2015 Mar 6;16(4):174–7. Doi: 10.1177/1757177415572646
43. Moses AE. Invasive Group A *Streptococcal* Infections, Israel. Emerging Infectious Diseases. 2002 Apr;8(4):421–6. Doi: 10.3201/eid0804.010278
44. Levitt A, Mermin J, Jones CM, See I, Butler JC. Infectious Diseases and Injection Drug Use: Public Health Burden and Response. The Journal of Infectious Diseases . 2020 Oct 1; 222(Supp5): S213–7. Doi: 10.1093/infdis/jiaa432
45. Herrera AL, Huber VC, Chaussee MS. The Association between Invasive Group A *Streptococcal* Diseases and Viral Respiratory Tract Infections. Frontiers in Microbiology. 2016 Mar 21;7:342. Doi: 10.3389/fmicb.2016.00342
46. ‌ Tyrrell G, Lovgren M, Kress B, Grimsrud 2005. Varicella-associated Invasive Group A *Streptococcal* disease in Alberta, Canada, 2000–2002. Clinical Infectious Diseases 40(7):1055-1057. Doi: 10.1086/428614.
47. Vugia DJ, Peterson CB, Meyers H, Kim KS, Arrieta A, Schlievert PM, et al. Invasive group A *streptococcal* infections in children with varicella in Southern California. 1996 Feb 1;15(2):146–50. Doi: 10.1097/00006454-199602000-00011
48. Invasive Group A *Streptococcus* control guideline - Control guidelines [Internet]. www.health.nsw.gov.au. Available from: <https://www.health.nsw.gov.au/Infectious/controlguideline/Pages/invasive-group-a-strep.aspx#references>
49. Krause V, Merridew N, Birrell J. Public Health Management of Invasive Group A *Streptococcal* Disease in the Northern Territory Guideline . 2022. Available from: <https://health.nt.gov.au/__data/assets/pdf_file/0009/1111122/invasive-group-A-streptococcal-disease-northern-territory-guidelines.pdf>
50. Invasive Group A *Streptococcal* Disease | Queensland Health . www.health.qld.gov.au. Available from: <https://www.health.qld.gov.au/cdcg/index/igas>
51. Thomson TN, Campbell PT, Gibney KB. The epidemiology of invasive group A *streptococcal* disease in Victoria, 2007–2017: an analysis of linked datasets. Australian and New Zealand Journal of Public Health. 2022 Aug 18; 46(6):878-883. Doi: 10.1111/1753-6405.13290.
52. Wright CM, Moorin R, Pearson G, Dyer JR, Carapetis JR, Manning L. Increasing incidence of invasive group A *streptococcal* disease in Western Australia, particularly among Indigenous people. Medical Journal of Australia. 2021 Jun 5;215(1):36–41. Doi: 10.5694/mja2.51117.
53. Wong NX, Crawford N, Oliver J, McMinn A, Ching NS, Baker C, et al. A Cluster of Pediatric Invasive Group A *Streptococcus* Disease in Melbourne, Australia, Coinciding with a High-Burden Influenza Season. Journal of Pediatric infectious diseases. 2019 Mar 7;14(04):213–8. Doi: 10.1186/s12889-019-8085-2
54. Olafsdottir LB, Erlendsdóttir H, Melo-Cristino J, Weinberger DM, Ramirez M, Kristinsson KG, et al. Invasive infections due to *Streptococcus* pyogenes: seasonal variation of severity and clinical characteristics, Iceland, 1975 to 2012. Eurosurveillance. 2014 May 1;19(17)5-14. Doi: 10.2807/1560-7917.ES.2016.21.10.30158.
55. Nelson GE, Pondo T, Toews KA, Farley MM, Lindegren ML, Lynfield R, et al. Epidemiology of Invasive Group A *Streptococcal* Infections in the United States, 2005–2012. Clinical Infectious Diseases. 2016 Apr 22;63(4):478–86. Doi: 10.1093/cid/ciw248
56. Public Health Scotland 2023. Group A *Streptococcal* infections. Available from: <https://www.hps.scot.nhs.uk/a-to-z-of-topics/streptococcal-infections/group-a-streptococcal-infections/>>
57. Dunne EM, Hutton S, Peterson E, Blackstock AJ, Hahn CG, Turner K, et al. Increasing Incidence of Invasive Group A *Streptococcus* Disease, Idaho, USA, 2008–2019. Emerging Infectious Diseases. 2022 Sep;28(9):1785–95.Doi: 10.3201/eid2809.212129
58. Smeesters PR, Laho D, Beall B, Steer AC, Van Beneden CA. Seasonal, Geographic, and Temporal Trends of *emm* Clusters Associated with Invasive Group A *Streptococcal* Infections in US Multistate Surveillance. Clinical Infectious Diseases. 2017 Feb 10;64(5):694–5

# Appendices

Appendix 1: PHU Checklist for invasive Group A *Streptococcal* (iGAS) cases

Appendix 2: Invasive Group A *Streptococcal* (iGAS) Infection: Information for the Public

# Appendix 1: PHU checklist for Invasive Group A *Streptococcal* cases (single case)

**Confirm case:**

□ Assess information on case against case definition

**Contact the patient’s doctor to:**

□ Obtain patient’s history

□ Confirm results of relevant pathology tests

□ Determine whether antibiotics have been provided to the case

□ Confirm onset date and symptoms of the illness (if possible)

□ Identify any known likely source of infection, including possible acquisition in a hospital or care facility

**Provide information to patient’s doctor, case or contacts:**

□ Recommend exclusions and restrictions

□ Identify contacts and assess risk of iGAS transmission. Determine whether contact is eligible for antibiotics for chemoprophylaxis and arrange, where applicable, and advise that medical attention be sought if not already obtained for any contacts with current symptoms.

□ Provide with appropriate iGAS Fact Sheet for case and contacts

**Other actions:**

□ Enter case data onto notifiable diseases database

□ For a death, report details to state/territory communicable disease agency

□ Where defined groups of people have been exposed (e.g. residential aged care, childcare), provide facility management with fact sheets for circulation to staff, and residents and, in the case of childcare or schools, for parents or guardians of attendee children.

# Appendix 2: Invasive Group A *Streptococcal* (iGAS) Infection: Information for Contacts

(The content of this sheet can be adapted for different settings and audiences)

You have been given this information sheet because you have been identified as a close contact of a person with Invasive Group A *Streptococca*l (iGAS) infection. Although your risk of developing iGAS infection is small, it is important that you are aware of symptoms to look out for, and that you seek medical attention should you have or develop those symptoms.

***What is Group A Streptococcus (GAS)?***

Group A *Streptococcus* (GAS) – also known as *Streptococcus pyogenes* - is a bacterium commonly found in the throat and on the skin. People may carry GAS and have no symptoms of illness.

***How is GAS spread?***

GAS bacteria are usually spread between people through coughing, sneezing, kissing, or direct skin to skin contact. Both people who carry GAS and people who are unwell with a GAS infection can pass these bacteria on to others.

***What is Invasive Group A Streptococcal*** ***(iGAS)?***

Invasive GAS infection, or iGAS, may occur when GAS bacteria get into parts of the body where these bacteria are not usually found and cause a severe infection, such as the blood, joints, or the birth canal after childbirth.

Although iGAS is uncommon, it is a very serious disease. The infection can develop very quickly and, depending how the disease presents and progresses, can be fatal.

Two severe, but rare, forms of iGAS are necrotising fasciitis (a deep tissue infection often requiring surgery) and Streptococcal Toxic Shock Syndrome (an illness that can cause high fever, low blood pressure, body rash, diarrhea and vomiting, difficulty breathing, and kidney or liver damage).

***Who is at risk of iGAS infections?***

Most people who have close contact with a person with iGAS infection remain well and symptom-free or develop mild throat or skin infections. These infections can easily be treated by your GP. Invasive GAS infections are rare, however there is some evidence that close contacts of a case (e.g. household members, birthing person-neonate pairs within 28 days of birth, sexual partners, childcare attendees and nursing home residents) are at higher risk of infection with GAS due to direct contact with secretions from infected persons.

***What are the symptoms of iGAS infection?***

Early signs and symptoms of iGAS may include:

* fever
* sore throat
* skin rash or redness around a wound site
* severe headache
* shortness of breath
* severe limb pain or muscle aches.

A person with iGAS infection can become very ill within 12 – 24 hours so it is important to be aware of these early signs and symptoms and seek medical attention immediately if you develop these symptoms within 30 days of contact with a person sick with iGAS.

***Do contacts of a person with iGAS infection require treatment?***

Contacts of a person with iGAS infection do not usually require any treatment if they remain well, and secondary cases are rare. However, if you or someone in the household develop any of the symptoms of iGAS infection in the 30 days following contact with someone with an iGAS infection, seek medical advice immediately and tell the doctor you have been in contact with someone recently diagnosed with iGAS infection.

***Infection prevention strategies***

Practicing good hand hygiene (including washing hands with soap and water and using an alcohol-based hand rub) can help reduce the risk of transmission. Any wounds that are draining or have pus must be kept covered with a clean, dry bandage or dressing. Used tapes, dressings or band aids must be discarded in the bin. Bacteria can spread on linen, towels and wash clothes, so launder these items after use with a regular laundry detergent.

***More information***

The following websites provide further information:

[Health Direct – Group A Streptococcal Disease](https://www.healthdirect.gov.au/group-a-streptococcal) (https://www.healthdirect.gov.au)

[Australian Government Department of Health and Aged Care](https://www.health.gov.au/diseases/group-a-streptococcal-disease-invasive-igas) (www.health.gov.au)

[Telethon Kids Institute](https://www.telethonkids.org.au/our-research/research-topics/invasive-strep-a-disease/?s=29757) (telethonkids.org.au)

1. “Birthing-person” refers to someone who gives birth, regardless of their gender identity, which may be female, male, nonbinary, or other, and regardless of their relationship with the neonate (e.g., surrogate pregnancy). [↑](#footnote-ref-1)