Stroke incidence and subtypes in Aboriginal people in remote Australia: a healthcare network population-based study

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ABSTRACT

Objectives We aimed to compare the incidence, subtypes and aetiology of stroke, and in-hospital death due to stroke, between Aboriginal and non-Aboriginal people in Central Australia, a remote region of Australia where a high proportion Aboriginal people reside (40% of the population). We hypothesised that the rates of stroke, particularly in younger adults, would be greater in the Aboriginal population, compared with the non-Aboriginal population; we aimed to elucidate causes for any identified disparities.

Design A retrospective population-based study of patients hospitalised with stroke within a defined region from 1 January 2011 to 31 December 2014.

Setting Alice Springs Hospital, the only neuroimaging-capable acute hospital in Central Australia, serving a network of 50 healthcare facilities covering 672 000 km².

Participants 161 residents (63.4% Aboriginal) of the catchment area admitted to hospital with stroke.

Primary and secondary outcome measures Rates of first-ever stroke, overall (all events) stroke and in-hospital death.

Results Of 121 residents with first-ever stroke, 61% identified as Aboriginal. Median onset-age (54 years) was 17 years younger in Aboriginal patients (p<0.001), and age-standardised stroke incidence was threefold that of non-Aboriginal patients (153 vs 51 per 100 000, incidence rate ratio 3.0, 95% CI 2 to 4). The rate ratios for the overall rate of stroke (first-ever and recurrent) were similar. In Aboriginal patients aged <55 years, the incidence of ischaemic stroke was 14-fold greater (95% CI 4 to 45), and intracerebral haemorrhage 19-fold greater (95% CI 3 to 142) than in non-Aboriginal patients. Crude prevalence of diabetes mellitus (70.3% vs 34.0%, p<0.001) and hypercholesterolaemia (68.9% vs 51.1%, p=0.049) was greater, and age-standardised in-hospital deaths were fivefold greater (35 vs 7 per 100 000, 95% CI 2 to 11) in Aboriginal patients than in non-Aboriginal patients.

Conclusions Stroke incidence (both subtypes) and in-hospital deaths for remote Aboriginal Australians are dramatically greater than in non-Aboriginal people, especially in patients aged <55 years.

INTRODUCTION

Stroke is the leading cause of disability and the third largest cause of death in Australia. Aboriginal and Torres Strait Islander (hereafter referred to as Aboriginal) Australians comprise 3.3% of the Australian population but have disproportionately greater morbidity and mortality rates. Life expectancy is 8 years less than among non-Aboriginal Australians and in remote and very remote areas the life-expectancy gap is 14 years, reflecting further disadvantage of remote and very remote Aboriginal residents compared with their non-remote counterparts.

These disparities highlight limitations to date in the translation of research into improved health outcomes for Aboriginal people.
Australians. This has been attributed to research being performed ‘on’ rather than in partnership with Aboriginal Australians, as well as a failure to recognise and address the catastrophic effects of colonisation on health outcomes for Aboriginal Australians. There is increasing awareness of the need for research designed in partnership with and in response to the priorities of Aboriginal Australians, to redress this. High-quality evidence, taking into account social, cultural, geographical and institutional factors to inform effective healthcare reform and policy development, is urgently required.

Published data on stroke among Aboriginal Australians are sparse. Stroke-related mortality rates in Aboriginal Australians in 2015 were reported to be 1.5-fold that of non-Aboriginal Australians with a burden of disease (disability-adjusted life years) of 2.3-fold that of non-Aboriginal Australians. Due to limitations of administrative data with potential for inaccuracies in classification of Indigenous status and cause of death, these disparities are likely underestimated. The only prospective population-based stroke incidence study in Aboriginal Australians provided evidence of excess rates in young people. However, estimates of incidence were imprecise, relying on only 15 Aboriginal patients with stroke, and while regional areas were covered, remote populations were not. Retrospective and linked-data studies indicate greater incidence and prevalence of stroke in Aboriginal people, but comprise insufficient detail for in-depth stroke phenotyping required to inform targeted primary and secondary prevention.

Alice Springs Hospital (ASH) is the major acute hospital (186-beds) for Central Australia, serving a vast catchment area spanning approximately 1.6 million km². ASH is the only neuroimaging-capable referral centre for all fifty primary healthcare facilities in the region, including the smaller (20-bed) Tennant Creek Hospital, 509 km distant by road. It is geographically isolated from other health services; flight distances are 1319 km from the nearest tertiary stroke centre and 1041 km from another CT scanner (Katherine, Northern Territory). There are no on-site neurologists or stroke specialists. Due to probable near-complete region-wide case ascertainment and a high proportion of Aboriginal people (40% of the resident population), ASH is ideally suited for a quasi-population-based stroke incidence study in remote-living Aboriginal Australians. This study is part of a research programme of stroke epidemiology in the South Australian and Northern Territory regions of Australia, initiated and endorsed by Aboriginal stakeholders.

AIMS

Using rigorous case evaluation based on all available clinical and neuroimaging data, we aim to compare the incidence of stroke and types of stroke, and stroke-related death between Aboriginal and non-Aboriginal Australians hospitalised with acute stroke within a defined region of Central Australia.

METHODS

Study period

The study comprises patients admitted from 1 January 2011 to 31 December 2014, inclusive. This study period was selected as it directly predated the introduction of a Stroke Protocol at ASH, thereby providing a baseline for measuring the efficacy of subsequent stroke interventions.

Study population

All ASH stroke admissions among residents of Alice Springs, Tennant Creek and the Anangu Pitjantjatjara Yankunytjatjara lands (figure 1) comprising a total population of 41562 people (40% Aboriginal) over approximately 672 000 km² (slightly larger than the size of France). We selected this smaller region within the ASH greater catchment area as this area comprises a distinct geographical region. Additionally, we determined that patients residing in this region would be most unlikely to present to other health services, allowing for precision in the calculation of incidence rates. Non-residents were excluded. ASH is the sole hospital serving this area that has a brain imaging facility. All patients from these regions with suspected stroke are routinely transferred from fifty primary healthcare facilities (including public primary healthcare centres, Aboriginal Community Controlled Health Services, Tennant Creek Hospital and privately managed general practices) to ASH.

Definition

WHO clinical definition of stroke was used, inclusive of ischaemic stroke (IS), spontaneous intracerebral haemorrhage (ICH) and non-traumatic subarachnoid
haemorrhage. Transient ischaemic attack, spontaneous extradural haematoma, subdural haematoma and traumatic haemorrhage were excluded.15

Data collection
Stroke admissions were identified using the International Classification of Diseases, 10th revision, Australian Modification (ICD-10-AM) codes I60, I61, I62.9 I63 and I64 (all diagnosis fields) from hospital administrative databases. Case information was extracted from ASH and tertiary hospital medical records for patients aged ≥18 years using a prespecified data extraction template and managed electronically. Risk factors were identified from medical records from ASH and from supplementary records from primary healthcare providers and tertiary referral centres, where available. Records examined included patient self-reported history, documented physician diagnoses prior to or during admission, in-hospital clinical parameters and laboratory results (ie, total cholesterol concentration documented on the electronic health records). When not identified, risk factors were recorded as absent. All neuro-imaging was reviewed and in-hospital deaths recorded. All authors had full access to the study data, including statistical reports and tables.

Case assessment
Clinical and radiological information was examined by the same senior vascular neurologist (TJK) to assign final stroke diagnosis and aetiological classification using the Trial of ORG 10172 in Acute Stroke Treatment14 and structural vascular lesions, medication, amyloid angiopathy, systemic disease, hypertension, or undetermined (SMASH-U) ICH15 criteria. We included only definite (evident on neuroimaging) or probable strokes (rapid-onset focal neurological disturbance lasting >24 hours or leading to death, without a non-stroke cause identified).13 Recurrent stroke was defined as a stroke event in a patient with a history of prior stroke during or before the study period. All others were deemed first-ever (incident) stroke. In-hospital death related to stroke was defined as death during hospital admission as a direct result of, or due to the complications of stroke. These deaths were used as numerators for population-based in-hospital death rates and for case fatality ratios.

Calculation of rates
Incidence (first-ever), attack (all strokes), recurrent stroke and in-hospital death rates were calculated per 100 000/year, world-population-standardised using the direct method.16 Case-fatality (incident strokes resulting in death as a proportion of all incident stroke events) was also calculated. CIs were calculated using the Poisson distribution. The population-at-risk was identified from the 2011 Census data of the study catchment area (figure 1). Comparisons were performed using Student’s t-test (continuous variables), Mann-Whitney U test (non-parametric variables), χ² test or Fisher’s exact test (binary outcomes).

Waiver of consent was granted.

Patient and public involvement
This study represents one component of the four-part ‘South Australian and Northern Territory Stroke Study: Stroke in Aboriginal and Torres Strait Islander people; incidence, mortality and disease burden’ which was designed in response to an identified need to determine burden of disease, outcomes and areas for service development for Aboriginal and Torres Strait Islander Australians. This research programme was prompted by a senior Aboriginal researcher (AB) and conducted by the research team in partnership with relevant Aboriginal stakeholders. Thus, Aboriginal health councils, Aboriginal research units, medical services, communities and Elders were consulted during the design, conduct and reporting of the study. As waiver of consent was obtained, it was not appropriate or possible to involve patients in the design, conduct or reporting of our research. Data management has been conducted in accordance with the CARE Principles for Indigenous Data Governance, which recommend data governance is designed to ensure Indigenous Peoples’ authority to control, responsibility for, and capacity to derive collective benefit from the data.17

RESULTS
Ascertainment
Of 274 coding-derived events, 43 cases were non-stroke (positive predictive value for coding of 84%, 95% CI 80% to 88%). Thirty cases were duplicates, 34 cases were non-residents, 40 cases were recurrent strokes and 6 case files were unlocatable. The remaining 121 patients with first-ever stroke were included in the analysis.

Incidence
During the study period, 121 residents (61.2% Aboriginal) were admitted with first-ever stroke (table 1). Median age at stroke occurrence in Aboriginal patients (54 years) was 17 years younger than that for non-Aboriginal patients (p<0.001). The sex distribution (~47% male) was similar between populations. Age-standardised stroke incidence in the Aboriginal population was threefold greater than in the non-Aboriginal population (95% CI 2 to 4; table 2). In those aged <55 years, this cut-off approximating the median age of Aboriginal people in our study, the age-standardised incidence of stroke in the Aboriginal population was near-fourteen-fold greater than in the non-Aboriginal population (table 3).

Attack rates
The overall rate of stroke was threefold greater in the Aboriginal population than in the non-Aboriginal population (table 4), a similar ratio to first-ever strokes. Of the total number of patients presenting with stroke aged less than 65 years, 21 (28%) Aboriginal patients presented with recurrent stroke compared with
one (5%) non-Aboriginal patient. Overall, the age-standardised rate of recurrent stroke in the Aboriginal population was almost 2-fold that in the non-Aboriginal population (59 vs 32 per 100 000/year, incidence rate ratio 1.8, 95% CI 1.2 to 2.8).

### Risk factors
All conventional risk factors for stroke, except smoking, appeared more frequent in Aboriginal than non-Aboriginal patients, but estimates were only statistically different for diabetes mellitus (70.3% vs 34.0%, p<0.001) and hypercholesterolaemia (68.9% vs 51.1%, p=0.049; table 1).

### Investigations and stroke subtypes
Neuroimaging was performed in 100% of patients with first-ever stroke; CT brain was performed in 98%, MRI in 11% and both in 10%. Of 88 patients with IS, 

### Table 1: Demographics, risk factor profile and in-hospital deaths among patients with first-ever stroke (2011–2014)

| Variable | Aboriginal N (%) | Non-Aboriginal N (%) | P value* |
|----------|------------------|----------------------|---------|
| Male     | 35 (47.3)        | 22 (46.8)            | 1.0     |
| Age      |                  |                      |         |
| Median (Q1, Q3), years | 54 (48–66)        | 71 (58–80)           | <0.001  |
| <55 years (n) | 40 (54.1)        | 7 (14.9)             | <0.001  |
| Risk factors † |              |                      |         |
| Atrial fibrillation | 12 (16.2)        | 7 (14.9)             | 0.845   |
| Diabetes mellitus | 52 (70.3)        | 16 (34.0)            | <0.001  |
| Hypertension | 60 (81.1)        | 32 (68.1)            | 0.102   |
| Obesity   | 10 (13.5)        | 6 (12.8)             | 0.906   |
| Smoker    | 14 (18.9)        | 10 (21.3)            | 0.751   |
| Hypercholesterolaemia | 51 (68.9)        | 24 (51.1)            | 0.049   |
| Ischaemic heart disease | 22 (29.7)        | 10 (21.3)            | 0.304   |
| Rheumatic heart disease | 5 (6.8)          | 1 (2.1)              | 0.403   |
| In-hospital death |              |                      |         |
| Case fatality | 17 (23.0)        | 5 (10.6)             | 0.086   |
| Crude rate of death‡ | 25               | 5                    | 0.001   |
| Age-standardised rate of death‡ | 35               | 7                    | <0.001  |

*All p values for comparison are based on crude measurements, unless otherwise stated.
†Risk factors identified from medical records, and includes history, or newly diagnosed from clinical parameters or laboratory results.
‡Rate per 100 000 population/year.

### Table 2: Age-specific incidence rates of first-ever stroke (per 100 000 population/year) in the Alice springs catchment area (2011–2014)

| Age, years | Population at risk | Aboriginal | Non-Aboriginal | Incidence rate ratio (IRR) |
|------------|--------------------|------------|---------------|---------------------------|
|            | n                  | n          | Rate 95% CI   | n            | Rate 95% CI   | IRR 95% CI | P value |
| 0–14*      | 5115               | 4568       | …  …  …      | …  …  …      | …  …  …  …   | 8.0 1 to 94 | 0.035   |
| 15–24*     | 3238               | 3070       | …  …  …      | …  …  …      | …  …  …  …   | 6.8 1 to 32 | 0.005   |
| 25–34      | 2788               | 4453       | 5 45 6 to 84 | 1 6 0 to 17  | 8.0 1 to 94 | 0.035   |
| 35–44      | 2347               | 3995       | 8 85 26 to 144 | 2 13 0 to 30 | 6.8 1 to 32 | 0.005   |
| 45–54      | 1556               | 4117       | 27 434 271 to 597 | 4 24 0 to 48 | 17.9 7 to 48 | <0.001  |
| 55–64      | 964                | 3053       | 14 363 173 to 553 | 12 98 43 to 154 | 3.7 2 to 8 | <0.001  |
| ≥65        | 748                | 1550       | 20 668 376 to 960 | 28 452 285 to 619 | 1.5 1 to 3 | 0.177   |
| Total*     | 16 756             | 24 806     | 74 110 85 to 136 | 47 47 34 to 61 | 2.3 2 to 3 | <0.001  |
| Standardised† | 153               | 129 to 177 | 51 37 to 65 | 3.0 2 to 4 | <0.001   |

*Ethics approval not obtained for those aged <18 years. However, calculated rates include the whole population.
†Age-standardised rate, standardised to WHO world population (rate per 100 000 population/year).
Table 3  Age-standardised incidence rate (per 100 000 population/year) of first-ever stroke, by type of stroke, in Alice springs catchment area (2011–2014)

|                      | Aboriginal Population† | N   | Rate‡ | 95% CI | Non-Aboriginal Population† | N   | Rate‡ | 95% CI | Incidence rate ratio (IRR)* | 95% CI | P value |
|----------------------|------------------------|-----|-------|--------|-----------------------------|-----|-------|--------|-----------------------------|--------|---------|
| **All stroke**       |                        |     |       |        |                             |     |       |        |                             |        |         |
| <55 years            | 15 044                 | 40  | 66    | 46 to 87 | 20 203                      | 7   | 9     | 2 to 15 |                             |        |         |
| Standardised*        | 68                     | 52  | 84    |        | 5                            | 5   | 1     | 10     | 13.6                         | 5 to 34 | <0.001  |
| All ages             | 16 756                 | 74  | 110   | 85 to 136 | 24 806                      | 47  | 47    | 34 to 61 |                             |        |         |
| Standardised*        | 153                    | 129 | 177   |        | 51                           | 51  | 37    | 65     | 3.0                          | 2 to 4  | <0.001  |
| **Ischaemic stroke** |                        |     |       |        |                             |     |       |        |                             |        |         |
| <55 years            | 15 044                 | 25  | 42    | 25 to 58 | 20 203                      | 4   | 5     | 0 to 10 |                             |        |         |
| Standardised*        | 42                     | 29  | 55    |        | 3                            | 0   | 7     |        | 14.0                         | 4 to 45 | <0.001  |
| All ages             | 16 756                 | 49  | 73    | 53 to 94 | 24 806                      | 39  | 39    | 27 to 52 |                             |        |         |
| Standardised*        | 103                    | 83  | 123   |        | 42                           | 30 | 55    |        | 2.5                          | 2 to 4  | <0.001  |
| **Intracerebral haemorrhage** |            |     |       |        |                             |     |       |        |                             |        |         |
| <55 years            | 15 044                 | 11  | 18    | 7 to 29  | 20 203                      | 1   | 1     | 0 to 4  |                             |        |         |
| Standardised*        | 19                     | 11  | 28    | 11 to 28 | 1                            | 0   | 2     |        | 19.0                         | 3 to 142 | <0.001  |
| All ages             | 16 756                 | 19  | 28    | 16 to 41 | 24 806                      | 4   | 4     | 0 to 8  |                             |        |         |
| Standardised*        | 39                     | 27  | 51    |        | 5                            | 0   | 9     |        | 7.8                          | 3 to 20  | <0.001  |
| **Subarachnoid haemorrhage** |                      |     |       |        |                             |     |       |        |                             |        |         |
| <55 years            | 15 044                 | 4   | 7     | 0 to 13  | 20 203                      | 2   | 2     | 0 to 6  |                             |        |         |
| Standardised*        | 7                      | 2   | 12    | 2 to 12 | 2                            | 0   | 4     |        | 3.5                          | 1 to 17  | 0.109   |
| All ages             | 16 756                 | 6   | 9     | 2 to 16  | 24 806                      | 4   | 4     | 0 to 8  |                             |        |         |
| Standardised*        | 11                     | 5   | 18    |        | 4                            | 0   | 7     |        | 2.8                          | 1 to 9   | 0.119   |

*Age-standardised rate, standardised to WHO world population (rate per 100 000 population/year).
†Population at risk.
‡Ethics approval not obtained for age <18 years. However, calculated rates include the whole population.
Table 4  Age-specific attack rates of stroke (per 100 000 population/year) in Alice springs catchment area (2011–2014)

| Age, year | Aboriginal | Non-Aboriginal | Incidence rate ratio (IRR) |
|-----------|------------|----------------|----------------------------|
|           | Population at risk | Rate | 95% CI | n | Rate | 95% CI | n | Rate | 95% CI | P value |
| 0–14*     | 5115       | 4568 | -     | - | -     | - | - | 1 | 6  | 5 to 17 | - | - | - | - |
| 15–24*    | 3238       | 3070 | -     | - | -     | - | - | 2 | 13 | 5 to 30 | 9.4 | 2 to 42 | 9.4 | <0.001 |<0.001 |<0.001 |
| 25–34     | 2788       | 4453 | 63    | 16 to 109 | 1 | 6  | 5 to 17 | 11.2 | 1 to 90 | 0.007 |<0.001 |<0.001 |
| 35–44     | 2347       | 3995 | 117   | 48 to 186 | 2 | 13 | 5 to 30 | 9.4 | 2 to 42 | 9.4 |<0.001 |<0.001 |
| 45–54     | 1556       | 4117 | 530   | 350 to 711 | 5 | 30 | 4 to 57 | 17.6 | 7 to 45 |<0.001 |<0.001 |<0.001 |
| 55–64     | 964        | 3053 | 622   | 374 to 871 | 12 | 98 | 43 to 154 | 6.4 | 3 to 13 |<0.001 |<0.001 |<0.001 |
| >65       | 748        | 1550 | 902   | 564 to 1241 | 39 | 629 | 432 to 826 | 1.4 | 1 to 2 | 0.146 |<0.001 |<0.001 |
| Total     | 16 756     | 24 806 | 152 | 123 to 182 | 59 | 59 | 44 to 75 | 2.6 | 2 to 4 |<0.001 |<0.001 |<0.001 |
| Standardised† | 212 | 184 to 241 | 66 | 50 to 82 | 3.2 | 2 to 4 |

*Ethics approval not obtained for age <18 years. However, calculated rates include the whole population.
†Age-standardised rate, standardised to WHO world population (rate per 100 000 population/year).

In-hospital case fatality and rate of death related to stroke

In-hospital case fatality appeared to be higher in Aboriginal people (23.0%) than non-Aboriginal people (10.6%, p=0.086), although not statistically significant at conventional levels. However, the crude and age-standardised rates were 21.2 and 6.6 per 100 000 population/year, respectively (95% CI 2.1 to 3.8 and 2.1 to 3.8, respectively, p=0.023). Other causes of death were infective (14.2%), heart failure (8.0%), and chronic respiratory disease (4.9%).

The age-standardised incidence of IS was over twofold greater in the entire Aboriginal population than in the entire non-Aboriginal population and 14-fold greater in the Aboriginal population aged <55 years compared with the non-Aboriginal population aged ≥55 years (table 3).

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| 15–24*    | 3238       | 3070 | -     | - | -     | - | - | 2 | 13 | 5 to 30 | 9.4 | 2 to 42 | 9.4 |<0.001 |<0.001 |<0.001 |
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rates of incident stroke resulting in death were significantly greater; fivefold greater in the Aboriginal than non-Aboriginal samples (table 1).

**DISCUSSION**

We present the first detailed stroke-neurologist curated data on stroke incidence and subtypes in remote Aboriginal Australians within a well-defined healthcare network system. Our results demonstrate significant disparities in age-standardised stroke incidence rates and in-hospital death rates between Aboriginal and non-Aboriginal populations. Disparities were especially marked in patients aged <55 years. Furthermore, Aboriginal patients appeared to more commonly have ICH than non-Aboriginal patients; this form of stroke has much poorer outcomes than for IS.

Aboriginal patients had stroke at an earlier age than non-Aboriginal people, and this disparity appeared greater than that reported previously, although CIs are wide. The disparities in the younger Aboriginal patients is consistent with earlier onset of cardiovascular comorbidity in the Aboriginal population, which is reported to occur on average 10–20 years earlier than in the non-Aboriginal population. Indeed, a recent study showed that the stroke risk in Aboriginal patients with atrial fibrillation aged under 60 years was three times higher than their non-Aboriginal counterparts, with an even higher disparity for fatal stroke. This is reflected in our study's findings; although only diabetes mellitus and hypercholesterolaemia were shown to have a significantly greater crude prevalence in Aboriginal patients, we found similar if not greater prevalence of all cardiovascular risk factors in Aboriginal patients, despite the 17-year age gap between the median age of first-ever stroke in Aboriginal and non-Aboriginal patients. Of note is that 6.8% of Aboriginal stroke patients had a history of rheumatic heart disease, a condition that is endemic in this population generally and in this region specifically.

Notably, our incidence rates for Aboriginal people are generally greater than those found in other studies where the composition of the population is less remote. This finding might be explained by greater rates of cardiovascular comorbidities in Aboriginal people residing in remote areas, compared with those in non-remote areas, although we acknowledge the relatively small sample size and overlapping confidence intervals. Furthermore, it appears that Aboriginal people are not only at greater risk of first stroke, but also of recurrent stroke. This may be a consequence of marked inequalities in access to and appropriateness of secondary prevention, particularly in remote Aboriginal communities.

The proportion of stroke attributable to ICH was substantially greater in Aboriginal patients, comprising over 25% of all incident strokes, approximately threefold that of non-Aboriginal patients. The reason for this is unclear, but may be partly attributable to earlier-onset
cardiovascular risk factors for ICH, along with dietary and lifestyle factors and the social determinants of health. Of note, the proportion of ICH in Aboriginal patients is similar to that observed in East Asian populations. This greater proportion of ICH, a type of stroke with greater mortality, also provides an explanation for the greater age-standardised rate of in-hospital death in Aboriginal patients than non-Aboriginal patients with incident stroke. This high stroke mortality is likely a major contributor to the life expectancy ‘gap’ between Aboriginal and non-Aboriginal Australians.

An aetiological classification could not be determined for more than half of patients with IS; this reflects the high proportion of both Aboriginal and non-Aboriginal patients having incomplete investigation in remote Australia. Importantly, this highlights unmet need and inequity in access, which is particularly significant given the younger age profile of Aboriginal patients, which would generally be an indication for more extensive investigation to determine the aetiology of stroke. It is also notable that no patient received thrombolysis, as thrombolytic therapy for stroke was not available at ASH during the time period of the study; this has subsequently been introduced along with a region-specific stroke protocol.

Our study had several limitations. Given its retrospective nature, our results may be affected by incomplete data records, with six records unlocatable, and is subject to bias. Therefore, our findings can only be indirectly compared with prospective incidence studies. Sample sizes in some age categories, including for older Aboriginal patients, are small, thereby reducing the precision of the incidence rates calculated. Suboptimal sensitivity of stroke coding may have also resulted in ascertainment bias (through false negatives), although our methods ensured that no false positives were included in our counts. In addition, underascertainment from remote communities may have occurred, either due to mild stroke syndromes or out-of-hospital fatal strokes. Although in an urban Australian stroke study, 96% of all incident cases presented to hospital, the proportion of patients not presenting to hospital in this remote catchment is unknown and may have been higher than in urban settings. However, the well-defined nature of the ‘ub and spoke’ network, with the requirement that patients with stroke are transferred to ASH for imaging diagnosis and more comprehensive management makes this number likely to be small. In further support for the view that substantial underascertainment in the non-Aboriginal population is unlikely, we found similar age-adjusted incidence rates in the non-Aboriginal population (47 per 100 000) to those in a recent ‘ideal’ rural Australian stroke study (50 per 100 000). If there was underascertainment in Aboriginal patients, the disparities we identified are likely to be underestimated. Incomplete documentation and diagnostic evaluation meant that in a large proportion of cases, stroke aetiology could not be determined. Furthermore, the relatively small sample size limited the ability to determine whether conventional risk factors explained the disparities in incidence.

This study also has a number of strengths. It comprises the largest population-based dataset of stroke in Aboriginal people utilising all available clinical and radiological data for case ascertainment, enabling all strokes to be adjudicated. While our findings of disparities in stroke incidence rates are similar to previously published data, we combine a larger cohort with detailed case analysis. Well-defined population-based methodology, with rigorous case analysis, allowed more accurate stroke ascertainment and phenotyping than previous coding-based studies. Previous studies have been unable to provide these insights, either due to low case numbers, imprecise stroke phenotyping due to the use of administrative data without accompanying clinical evaluation, or datasets almost 20 years old. Lastly, to our knowledge, we have provided the first validation for the positive predictive value of ICD-10-AM codes for stroke in a remote setting.

We have identified areas of unmet need in healthcare services, and provide a foundation on which to explore the efficacy of health intervention on stroke incidence and outcomes in Aboriginal Australians. Partly as a result of this study, in the 5 years following the study period, a stroke protocol has been introduced, including Telestroke-supported intravenous thrombolytic therapy for IS, as well as Telestroke-supported retrieval 1319 km distance to Adelaide in South Australia for endovascular thrombectomy or surgical intervention where appropriate. These interventions have facilitated improved access to reperfusion therapy, investigation, aetiological classification and secondary prevention of stroke in Central Australia. In partnership with Aboriginal researchers, governing bodies, medical services and community groups, these results are being used to inform the development of culturally responsive stroke prevention and management tools for Aboriginal communities in Central Australia. Further research will be undertaken to evaluate the efficacy of these interventions.

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