Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

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In Australia circa 2010, 4.1 million (90% credible interval [CrI] 2.3–6.4 million) episodes of foodborne gastroenteritis occurred, many of which might have resulted in sequelae. We estimated the number of illnesses, hospitalizations, and deaths from Guillain-Barré syndrome, hemolytic uremic syndrome, irritable bowel syndrome, and reactive arthritis that were associated with contaminated food in Australia. Data from published studies, hospital records, and mortality reports were combined with multipliers to adjust for different transmission routes. We used Monte Carlo simulation to estimate median estimates and 90% CrIs. In Australia, circa 2010, we estimated that 35,840 (90% CrI 25,000–54,000) illnesses, 1,080 (90% CrI 700–1,600) hospitalizations, and 10 (90% CrI 5–14) deaths occurred from foodborne gastroenteritis–associated sequelae. Campylobacter spp. infection was responsible for 80% of incident cases. Reducing the incidence of campylobacteriosis and other foodborne diseases would minimize the health effects of sequelae.

Foodborne gastroenteritis is a major source of illness in Australia, causing an estimated 4.1 million (90% credible interval [CrI] 2.3–6.4 million) illnesses, 30,600 (90% CrI 28,000–34,000) hospitalizations, and 60 (90% CrI 53–63) deaths each year (1). In addition to the direct effects of these illnesses, infection with some pathogens can result in sequelae, which can be severe, require multiple hospitalizations, and be costly to society (2). We report on the effects of sequelae associated with Guillain-Barré syndrome (GBS), hemolytic uremic syndrome (HUS), irritable bowel syndrome (IBS), and reactive arthritis (ReA) from 5 pathogens acquired from contaminated food in Australia.

Each of these 4 sequel illnesses are preceded by different gastrointestinal infections and have unique characteristics. GBS, a rare but serious autoimmune illness, affects the nervous system and causes acute flaccid paralysis. GBS can occur as a sequel to Campylobacter spp. infection 10 days–3 weeks after gastroenteritis (3,4). HUS is characterized by acute renal failure, hemolytic anemia, and thrombocytopenia and can result from infection with Shiga toxin–producing Escherichia coli (STEC) ≈4–10 days after onset of gastroenteritis (5,6). IBS is a gastrointestinal disorder that causes abdominal pain and bowel dysfunction. It is not life threatening, but it can cause substantial health effects after illness with Campylobacter spp., nontyphoidal Salmonella enterica serotypes (hereafter referred to as nontyphoidal Salmonella spp.), or Shigella spp. (7,8). ReA is a type of spondyloarthritis that can develop up to 4 weeks after an enteric infection from Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., or Yersinia enterocolitica (9). We estimated the number of illnesses, hospitalizations, and deaths resulting from GBS, HUS, IBS, and ReA from selected foodborne pathogens in Australia in a typical year circa 2010.

Methods

We estimated the effects of foodborne sequelae acquired in Australia circa 2010 using data from multiple sources in Australia and from international peer-reviewed literature. We defined foodborne sequelae as illnesses occurring after bacterial gastroenteritis caused by eating contaminated food. Sequelae were defined as the secondary adverse health outcomes resulting from a previous infection by a microbial pathogen and clearly distinguishable from the initial health event (10). Illness can be acute, such as with HUS, or chronic (lasting for many years), as with IBS. We estimated incidence, hospitalizations, and deaths with uncertainty bounds using Monte Carlo simulation in @Risk version 6 (http://www.palisade.com/), which incorporates uncertainty in both data and inputs. Each stage of our calculation was represented by a probability distribution, and our final estimates of incidence, hospitalizations, and deaths were summarized by the median and 90% CrI. Similar to a recent study in the United States (11), we used empirical distributions for source distributions, such as the number of hospitalizations or deaths, to avoid assumptions about the expected shape of these distributions. All other inputs were modeled by using the PERT (project evaluation and review technique) distribution, which enables the input of a minimum, maximum, and modal value, or 3 percentile points, such as a median value and 95% bounds. We used this distribution widely in our analyses because
it enables asymmetric distributions and can be produced from many data sources, including expert elicitation data. The Australian National University Human Research Ethics Committee approved the study.

Incidence of Sequelae
Several pathogens are associated with the development of sequelae. Community estimates of foodborne illness from Kirk et al. (1) for Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., STEC, and Y. enterocolitica were used for estimating the incidence of foodborne sequelae (Table 1). Although Shigella spp. and nontyphoidal Salmonella spp. have been associated with HUS and STEC has been associated with IBS and ReA, data on which to base estimates are limited. In addition, although other pathogens, such as Chlamydia trachomatis, Clostridium difficile, Giardia lamblia, and norovirus, have been associated with these sequelae (12–15), we assessed only pathogens commonly associated with sequelae, domestically acquired, and with a foodborne transmission pathway. A “sequelae multiplier,” which is the proportion of sequelae cases that develop after enteric infection with a specific bacterial pathogen, was applied to our estimates of domestically acquired foodborne gastrointestinal cases caused by that pathogen (1). For each sequel illness, we reviewed relevant studies published during 1995–2012 using systematic reviews and studies using Australian data where possible to estimate the relevant sequelae multipliers. We reviewed articles about sequelae after infection with Campylobacter spp., E. coli, nontyphoidal Salmonella spp., Shigella spp., and Y. enterocolitica, and we estimated sequelae multipliers for GBS, HUS, IBS, and ReA after bacterial gastrointestinal infection on the basis of these reviews (Table 2). Relevant articles and additional information are documented in online Technical Appendix 1 (http://wwwnc.cdc.gov/EID/article/20/11/13-1316-Techapp1.pdf).

Our sequelae multiplier for GBS was based on 30.4 (range 19.2–94.5) cases of GBS per 100,000 cases of campylobacteriosis using data from studies from the United Kingdom, Sweden, and the United States (16–18). For HUS, the sequelae multiplier used was 3% (95% CI 1.7%–5.4%) from a South Australian study on STEC and HUS notifications during 1997–2009 (19). On the basis of data from Haagsma et al. (20), we assumed that 8.8% (95% CI 7.2%–10.4%) of foodborne disease caused by Campylobacter spp., nontyphoidal Salmonella spp., and Shigella spp. result in IBS. We used a separate sequelae multiplier for each pathogen that resulted in ReA. We assumed that 7% (range 2.8%–16%) of foodborne cases of Campylobacter spp., 8.5% (range 0%–26%) of foodborne cases of nontyphoidal Salmonella spp., 9.7% (range 1.2%–9.8%) of foodborne cases of Shigella spp., and 12% (range 0%–23.1%) of foodborne cases of Y. enterocolitica result in ReA (see full reference list in online Technical Appendix 1). Total foodborne IBS and ReA cases reflect the sum of modeled IBS and ReA cases from these 3 and 4 pathogens, respectively. Details on the sequelae multipliers and incidence estimation methods are in online Technical Appendix 1 and online Technical Appendix 2 (http://wwwnc.cdc.gov/EID/article/20/11/13-1316-Techapp2.pdf).

We compared the incidence of sequelae circa 2010 to that of sequelae circa 2000 by applying the same sequelae multipliers to estimates of the incidence of acute gastroenteritis to specific pathogens in 2006–2010 and 1996–2000, respectively. The estimates of incidence of acute gastroenteritis were based on notification data for Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., STEC, and Y. enterocolitica (19,21,22), (online Technical Appendix 3, http://wwwnc.cdc.gov/EID/article/20/11/13-1316-Techapp3.pdf).

Hospitalizations and Deaths
To estimate hospitalizations associated with IBS from foodborne Campylobacter spp., nontyphoidal Salmonella spp., and Shigella spp. and hospitalizations associated with ReA from foodborne Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., and Y. enterocolitica, we used hospitalization data for 2006–2010 from all Australian states and territories, according to the International Classification of Diseases, Tenth Revision, Australian Modification (ICD-10-AM) codes. To estimate deaths for all 4

| Table 1. Pathogens associated with GBS, HUS, IBS, and ReA included in this study, Australia, circa 2010* |
|--------------------------------------------------|
| Pathogen                                      | GBS | HUS | IBS | ReA |
| Campylobacter spp.                            | X   | X   | X   |     |
| Nontyphoidal Salmonella spp.†                 | X   | X   | X   |     |
| Shigella spp.                                 |     |     |     |     |
| Shiga toxin–producing Escherichia coli         |     |     |     |     |
| Yersinia enterocolitica                       |     |     |     |     |

*GBS, Guillain-Barré syndrome; HUS, hemolytic uremic syndrome; IBS, irritable bowel syndrome; ReA, reactive arthritis.
†Nontyphoidal S. enterica serotypes.

| Table 2. Sequelae multipliers extracted from the literature about domestically acquired foodborne bacterial gastroenteritis* |
|--------------------------------------------------|
| Sequelae                                      | ICD-10-AM code | Incidence after bacterial infection, % |
|------------------------------------------------|
| GBS, median (range)                            | G61.0           | 0.0304 (0.0192–0.0945) |
| HUS, median (95% CI)                           | D59.3           | 3 (1.7–5.1) |
| IBS, median (95% CI)                           | K58.0           | 8.8 (7.2–10.4) |
| ReA, median (range)                            | M02.1           | 7–12 (0–28) |
|                                                | M02.3           | M02.8 |
|                                                | M03.2           |

*CrI, credible interval; GBS, Guillain-Barré syndrome; HUS, hemolytic uremic syndrome; IBS, irritable bowel syndrome; ICD-10-AM, International Classification of Diseases, Tenth Revision, Australian Modification; ReA, reactive arthritis.
sequelae illnesses resulting from the respective foodborne pathogens, we used national death data for 2001–2010 from the Australian Bureau of Statistics, using ICD-10-AM codes (online Technical Appendix 4, http://wwwnc.cdc.gov/EID/article/20/11-1316-Techapp4.pdf). Principal diagnosis and additional diagnoses were included for hospitalizations, and underlying and contributing causes were included for deaths. Because we had only 1 year of hospitalization data for Victoria and 2 years for New South Wales, we extrapolated from these data to derive a distribution of the number of hospitalizations across all states, which was modeled as an empirical distribution. For these states, we assumed the same number of hospitalizations each year to adjust for missing data. Because of the severity of GBS and HUS, hospitalization estimates for these illnesses were not modeled, and all persons with estimated incident cases from contaminated food were considered to have been hospitalized.

We estimated incidences of hospitalization and death using a statistical model that incorporates uncertainty in case numbers and in multipliers using probability distributions (Figure), which is adjusted from the hospitalization estimation flow chart in Kirk et al. (1). We assumed that all estimated incident foodborne Campylobacter-associated GBS and STEC-associated HUS case-patients were hospitalized, so those cases were not modeled; however, multipliers were still needed for GBS and HUS to estimate deaths. Sequelae-associated deaths were estimated by using the same methods as for hospitalizations (Figure). Input data arose from the data sources discussed above or from multipliers that are discussed below.

**Domestically Acquired Multiplier**

The “domestically acquired multiplier” adjusted for the proportion of case-patients who acquired their infection in Australia. We estimated domestically applied multipliers for the antecedent bacterial gastrointestinal pathogens using notifiable surveillance data from each state, extrapolated to give national estimates (1). We adopted the domestically acquired multiplier for Campylobacter spp. of 0.97 (90% CrI 0.91–0.99) for GBS and the domestically acquired multiplier for STEC 0.79 (90% CrI 0.73–0.83) for HUS (1). For IBS and ReA, a combined domestically acquired multiplier for Campylobacter spp., nontyphoidal Salmonella spp., and Shigella spp. for IBS and Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp. and Y. enterocolitica for ReA was calculated as a weighted average of the domestically acquired multipliers for each pathogen, weighted by the total number of IBS cases for each pathogen. The ReA bacterial multiplier was then further multiplied by a foodborne multiplier for Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., and Y. enterocolitica by using a weighted average of the foodborne multipliers for each pathogen as was done for IBS (online Technical Appendices 4 and 5).

**Proportion Foodborne Multiplier**

For each of the 4 sequelae, we calculated the proportion of hospitalizations and deaths from foodborne pathogens using 2 multipliers: a “bacterial multiplier” to attribute the proportion of overall cases of each of the sequelae illnesses to specific pathogens and a “foodborne multiplier” to attribute illnesses to foodborne exposure. The bacterial multiplier, which was the proportion of sequel cases attributable to their antecedent bacterial pathogen, was extracted from systematic reviews for GBS and HUS (4,23) and multiplied by the foodborne proportion for Campylobacter spp. and STEC, respectively. For IBS and ReA, from the literature we extracted a midpoint and range of the proportion of cases that resulted from infectious gastroenteritis (12,20,24). The IBS bacterial multiplier was then further multiplied by a foodborne multiplier for Campylobacter spp., nontyphoidal Salmonella spp., and Shigella spp., which was calculated as a weighted average of the foodborne multipliers for each pathogen, weighted by the total number of IBS cases for each pathogen. The ReA bacterial multiplier was then also multiplied by the foodborne multiplier for Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., and Y. enterocolitica by using a weighted average of the foodborne multipliers for each pathogen as was done for IBS (online Technical Appendices 4 and 5).
Results

Incidence

We estimated that, circa 2010 in Australia, 70 (90% CrI 30–150) new cases of Campylobacter-associated GBS, 70 (90% CrI 25–200) new cases of STEC-associated HUS, 19,500 (90% CrI 12,500–30,700) new cases of Campylobacter-, nontyphoidal Salmonella- and Shigella-associated IBS, and 16,200 (90% CrI 8,750–30,450) new cases of Campylobacter-, nontyphoidal Salmonella-, Shigella-, and Y. enterocolitica–associated ReA were domestically acquired and caused by contaminated food (Table 3). We estimated that 35,840 (90% CrI 25,000–54,000) domestically acquired sequel illnesses resulted from foodborne gastroenteritis—an incidence rate of 1,620 (90% CrI 1,150–2,450) sequelae cases per million population annually. Campylobacter spp. infection resulted in the largest number of sequelae cases annually; ≈80% of the 36,000 sequel illnesses were attributable to Campylobacter spp. alone.

Comparison with Estimates Circa 2000

Using data circa 2000, we estimated that 50 GBS cases, 55 HUS cases, 14,800 IBS cases, and 12,500 ReA cases occurred each year. Elsewhere, we estimated that the rate of foodborne campylobacteriosis was approximately 13% higher in 2010 than 2000 (1); this increase led to a 13% decrease in Campylobacter-associated GBS in 2010 over 2000. Similarly, we estimated that the rate of foodborne salmonellosis was 24% higher in 2010 than in 2000 (1). These factors combine to explain much of the increase in IBS and ReA. The rate of STEC-associated HUS remained about the same in 2000 and 2010 (online Technical Appendix 3).

Hospitalizations and Deaths

We estimated that, circa 2010 in Australia, 1,080 (90% CrI 700–1,600) hospitalizations for sequelae illnesses occurred from domestically acquired foodborne gastroenteritis, equating to 50 (90% CrI 30–70) hospitalizations per million population per year (Table 4). We estimated a total of 10 (90% CrI 5–14) deaths from sequelae to domestically acquired foodborne gastroenteritis—a rate of 0.5 (90% CrI 0.2–0.6) deaths per million population per year (Table 4).

Discussion

Our study demonstrates that foodborne gastroenteritis in Australia results in substantial severe and disabling sequelae. We estimated a yearly rate of 1,620 incident cases of sequelae illnesses, 50 hospitalizations, and 0.5 deaths per million population circa 2010. In addition, a comparison with estimates recalculated for 2000 indicates an increase in the rates of GBS, IBS, and ReA since 2000, which is consistent with and directly related to rising levels of antecedent foodborne illnesses caused by Campylobacter spp. and nontyphoidal Salmonella spp. during this period (1). This increase highlights the importance of quantifying sequelae when estimating the effects of foodborne disease and provides further impetus for reducing illness from foodborne bacterial pathogens.

The impact of Campylobacter spp. infection in the community is high. Approximately 179,000 cases of foodborne campylobacteriosis occur in Australia each year (1), and Campylobacter spp. was responsible for 80% of the foodborne sequelae illness estimated in this study. The reported rate of infection from Campylobacter spp. in Australia has increased since 2010 (1) and is higher than in many other industrialized countries. For example, the rate of Campylobacter spp. for Australia was ≈10 times higher than that for the United States (25), double that for the Netherlands (26), and slightly higher than that for the United Kingdom (27). In the Netherlands, a lower rate of acute Campylobacter spp. gastroenteritis has contributed to lower estimates of rates of sequel illnesses than our estimates for GBS, IBS, and ReA (26).

In New Zealand, food safety interventions have been effective in lowering campylobacteriosis rates and sequelae. In 2006, high campylobacteriosis notification rates (>3,800 cases per million population) prompted increased research on Campylobacter spp., which resulted in the introduction of food safety and poultry industry interventions, including Campylobacter spp. performance targets at primary processing plants and promotion of freezing all fresh poultry meat (28). By 2008, the rate of campylobacteriosis notifications decreased by 54% to 1,615 cases per million population (28). In addition, after these interventions in New Zealand, the rate of GBS hospitalizations decreased by 13% (29). The less dramatic decrease in GBS than in campylobacteriosis might be explained by the fact that Campylobacter spp. is not the only cause of GBS. If Australia were to experience decreases similar to those in New Zealand, we would expect the rate of foodborne campylobacteriosis in the community to drop from approximately 8,400 to 3,864 cases per million population. Sequelae would decrease from 1,620 to 870 cases per million population per year. Furthermore, total GBS-associated hospitalizations, including GBS from all causes and readmissions, would decrease from ≈73 to 63 hospitalizations per million population annually.

A comparison of our foodborne Campylobacter-associated GBS incidence estimates with raw hospitalization data showed many more hospitalizations than incident cases. This finding probably is attributable to repeat hospitalizations. We took a conservative approach by basing incidence estimates on community estimates of campylobacteriosis and assuming that all persons with incident cases were hospitalized. A yearly median of 1,536 (range
1,428–1,632) primary and additional GBS diagnoses occurred in Australian hospitals during 2006–2010 (including GBS from all causes and readmissions) and equates to a median rate of 73.1 (range 64.7–77.4) GBS-associated hospitalizations per million population each year. This rate is within the range from a New Zealand study, which found a median rate of 56.3 (range 42.1–75.9) GBS-associated hospitalizations during a 13-year period, with ≈41% of case-patients being readmitted, resulting in 23.2 (range 15.3–29.3) incident GBS hospitalizations per million population each year (29). If we assume that 41% of Australia’s 1,536 GBS hospitalizations are readmissions and apply the domestically acquired multiplier and foodborne proportion multiplier used to estimate GBS-associated deaths (online Technical Appendix 4), we would estimate 170 (90% CrI 60–265) incident foodborne Campylobacter-associated GBS hospitalizations. This point estimate is higher than our current estimate of 70, although the credible interval includes our estimate. A validation study of medical records of persons with GBS would enable us to better characterize readmissions for GBS.

Our approach has several limitations. First, our comparison of sequelae estimates for 2000–2010 assumes a constant rate of sequelae illness after gastrointestinal infection over time. Although our methods provide an indirect method of assessing changes in sequelae incidence over time, the approach is useful because it enables comparison of the population-level effect of sequelae at these 2 time points. Second, our study measured incidence and not prevalence of sequelae. We estimated the number of new cases every year and did not quantify the long-term effects of these sequelae. Third, our study does not estimate all sequelae illness from foodborne disease pathogens. We did not include sequelae, such as end-stage renal disease, inflammatory bowel disease, and encephalitis, in our estimates. We chose GBS, HUS, IBS, and ReA for this study because they were known, well studied, and well characterized in available data sources. These provide a good basis to begin to understand the effects of foodborne sequelae and the policy implications of reducing illness from preceding bacterial pathogens.

Our estimates for GBS, HUS, IBS, and ReA incidence relied heavily on the quality of the literature we reviewed. We used Australian data and systematic reviews wherever possible. The Australian hospitalization and deaths data we used were of high quality and included both principal and additional diagnoses from all states. However, because data were missing from some states in some years, we extrapolated from these data to the remaining years. Finally, ICD-10 and ICD-10-AM coding can be problematic when co-morbid conditions are present, when hospital transfers occur, or when diagnostic criteria are inconsistent. Therefore, our estimates for sequelae hospitalizations and deaths may be conservative because they do not account for these coding errors.

Table 3. Estimated number of sequelae illnesses resulting from domestically acquired foodborne bacterial gastroenteritis, Australia, circa 2010*

| Sequelae, pathogen | Median no. Illnesses (90% CrI) | Median rate (90% CrI)† |
|--------------------|-------------------------------|------------------------|
| GBS, Campylobacter spp. | 70 (30–150) | 3.1 (2–6) |
| HUS, STEC | 70 (25–200) | 3.3 (1–9) |
| IBS | 915 (550–1,400) | 43 (25–70) |
| ReA | 25 (20–40) | 1 (1–2) |
| Total | 1,080 (700–1,600) | 50 (30–70) |

*CrI, credible interval; GBS, Guillain-Barré syndrome; HUS, hemolytic uremic syndrome; IBS, irritable bowel syndrome; ReA, reactive arthritis; STEC, Shiga toxin–producing Escherichia coli.
†No. cases per million population.
‡i.e., nontyphoidal Campylobacter spp.
§Simulated values, which might not add to total because of rounding and variation over simulations.

1,428–1,632) primary and additional GBS diagnoses occurred in Australian hospitals during 2006–2010 (including GBS from all causes and readmissions) and equates to a median rate of 73.1 (range 64.7–77.4) GBS-associated hospitalizations per million population each year. This rate is within the range from a New Zealand study, which found a median rate of 56.3 (range 42.1–75.9) GBS-associated hospitalizations during a 13-year period, with ≈41% of case-patients being readmitted, resulting in 23.2 (range 15.3–29.3) incident GBS hospitalizations per million population each year (29). If we assume that 41% of Australia’s 1,536 GBS hospitalizations are readmissions and apply the domestically acquired multiplier and foodborne proportion multiplier used to estimate GBS-associated deaths (online Technical Appendix 4), we would estimate 170 (90% CrI 60–265) incident foodborne Campylobacter-associated GBS hospitalizations. This point estimate is higher than our current estimate of 70, although the credible interval includes our estimate. A validation study of medical records of persons with GBS would enable us to better characterize readmissions for GBS.

Our approach has several limitations. First, our comparison of sequelae estimates for 2000–2010 assumes a constant rate of sequelae illness after gastrointestinal infection over time. Although our methods provide an indirect method of assessing changes in sequelae incidence over time, the approach is useful because it enables comparison of the population-level effect of sequelae at these 2 time points. Second, our study measured incidence and not prevalence of sequelae. We estimated the number of new cases every year and did not quantify the long-term effects of these sequelae. Third, our study does not estimate all sequelae illness from foodborne disease pathogens. We did not include sequelae, such as end-stage renal disease, inflammatory bowel disease, and encephalitis, in our estimates. We chose GBS, HUS, IBS, and ReA for this study because they were known, well studied, and well characterized in available data sources. These provide a good basis to begin to understand the effects of foodborne sequelae and the policy implications of reducing illness from preceding bacterial pathogens.

Our estimates for GBS, HUS, IBS, and ReA incidence relied heavily on the quality of the literature we reviewed. We used Australian data and systematic reviews wherever possible. The Australian hospitalization and deaths data we used were of high quality and included both principal and additional diagnoses from all states. However, because data were missing from some states in some years, we extrapolated from these data to the remaining years. Finally, ICD-10 and ICD-10-AM coding can be problematic when co-morbid conditions are present, when hospital transfers occur, or when diagnostic criteria are inconsistent. Therefore, our estimates for sequelae hospitalizations and deaths may be conservative because they do not account for these coding errors.

Table 4. Estimated number of sequelae-associated hospitalizations and deaths caused by domestically acquired foodborne bacterial gastroenteritis, Australia, circa 2010*

| Sequelae | Hospitalizations | Rate (90% CrI)† | Deaths | Rate (90% CrI)† |
|----------|-----------------|----------------|--------|----------------|
| GBS      | 70 (30–150)     | 3.1 (2–6)      | 6 (2–10) | 0.3 (0.1–0.5)  |
| HUS      | 70 (25–200)     | 3.3 (1–9)      | 2 (1–3) | 0.1 (0.03–0.12) |
| IBS      | 915 (550–1,400) | 43 (25–70)     | 2 (1–2) | 0.1 (0.05–0.11) |
| ReA      | 25 (20–40)      | 1 (1–2)        | 0       | 0              |
| Total    | 1,080 (700–1,600) | 50 (30–70)  | 10 (5–14) | 0.5 (0.2–0.6)  |

*CrI, credible interval; GBS, Guillain-Barré syndrome; HUS, hemolytic uremic syndrome; IBS, irritable bowel syndrome; ReA, reactive arthritis; STEC, Shiga toxin–producing Escherichia coli.
†Cases per million population.
The sequelae estimates from this study showed that the impact of foodborne Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., STEC, and Y. enterocolitica was much greater than when consideration is given simply to the initial acute illness. Campylobacter spp. infection, in particular, was highlighted as an increasing problem in Australia. Our estimates provide a basis for costing studies, which can be useful for developing food safety policies and interventions. Finally, this study highlights the need for better data from large population-based studies in Australia to further characterize sequelae, as well as foodborne pathogens.

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Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

Technical Appendix 1

Sequelae Incidence after Bacterial Gastroenteritis: The Sequelae Multiplier

For each sequel, a multiplier was used that estimated the proportion of bacterial gastroenteritis cases that developed into chronic sequelae. This appendix summarizes the relevant studies published during 1995–2012, which we selected for review, as well as the sequelae multipliers that were estimated for Guillain-Barré syndrome (GBS), hemolytic uremic syndrome (HUS), irritable bowel syndrome (IBS), and reactive arthritis (ReA).

GBS

A few studies have quantified the incidence of GBS illness following *Campylobacter* spp. infection by using large cohorts of patients or the literature (online Technical Appendix 1 Table 1). In a population-based cohort study in the United Kingdom, including 2 months of follow-up, 3 cases of GBS occurred among 15,587 *Campylobacter* spp. cases. This yielded a rate of 19.2 cases of GBS per 100,000 cases of campylobacteriosis (1). In Sweden, 0.03% of a cohort of 29,567 persons with laboratory-confirmed *C. jejuni* infection developed GBS illness after 2 months of follow-up, yielding an annual incidence of 30.4 cases of GBS per 100,000 cases of *C. jejuni* infection (95% CI 13.9–57.8) of *C. jejuni* infection (2). In a literature review, Allos (3) estimated that in the United States, GBS develops in 1 of every 1,058 cases, or 94.5 per 100,000 cases, of *C. jejuni* infection. Baker et al. (4) performed a study of hospital records in New Zealand, which found a rate of 414 cases of GBS per 100,000 *Campylobacter* spp. hospitalizations.

For the sequelae multiplier, a midpoint of 30.4 cases of GBS per 1000,000 cases of campylobacteriosis was taken from the study by McCarthy and Gieseke (2) using a minimum value of 19.2 per 100,000 from the UK study and a maximum value of 94.5 per 100,000 from the study by Allos (3). Although the study by Baker et al. (4) is a valuable one, we excluded it from
the calculation of our sequelae multiplier because persons hospitalized with *Campylobacter* spp. infection may not be representative of *Campylobacter* spp. cases in the community.

### Technical Appendix 1 Table 1. Incidence of GBS after infection with *Campylobacter* spp.*

| Reference                  | Study years     | Type of study | Country     | No. GBS cases/*Campylobacter* spp. patients | Incidence per 100,000 (95% CI) |
|----------------------------|-----------------|---------------|-------------|--------------------------------------------|--------------------------------|
| Baker et al. (4)           | 1995–2008       | Cohort        | New Zealand | 35/8,448 hospitalizations                   | 414 (373–459)                  |
| Tam et al. (1)             | 1991–2001       | Cohort        | UK          | 3/15,587 cases                             | 19.2 (17.1–21.5)               |
| McCarthy and Giesecke (2) | 1987–1995       | Cohort        | Sweden      | 9/29,563 cases                             | 30.4 (13.9–57.8)               |
| Allos (3)                  | 1964–1996†      | Review and estimation | Global/USA | 1/1058 cases                              | 94.5 (2.4–525)                 |

*GBS, Guillain-Barré syndrome.†Years of reviewed studies.

**HUS**

A variety of organisms, drugs and conditions can initiate the symptoms of HUS, but the majority of HUS cases are post-diarrheal—usually caused by Shiga toxin–producing *Escherichia coli* (STEC) (5). In developed communities, STEC is the most commonly implicated organism in HUS (6), and in children, 90% of HUS cases are due to STEC (5). HUS is also associated with *Shigella dysenteria* serotype 1, particularly in less developed communities (6); however, a recent systematic review was unable to find an adequate number of studies to quantify the association between *S. dysenteria* serotype 1 and HUS (7). In addition, in a few studies, HUS has been associated with *Clostridium difficile* and *Salmonella enterica* serotype Typhi, but the evidence is limited (8–10). Therefore we estimated food-related HUS cases as a sequel to STEC, which may create an underestimation of HUS if there are food-related HUS cases in Australia from other organisms.

Several sources have reported that 3%-7% of sporadic STEC infections develop into HUS (11–14). Australian studies support this estimate range. Vally et al. (15) examined South Australian surveillance data and identified 14 HUS cases and 460 STEC cases, resulting in an estimate of 3% of STEC cases developing into HUS. Sixty percent of HUS case-patients were ≤15 years of age. In addition, in a case–control study in 6 Australian jurisdictions, 113 STEC case-patients were identified, 44 of whom were infected with O157 and 66 who were infected with non-O157 (14). Eight (7%) of all the STEC cases, 1 (2%) case-patient with O157, and 7 (10%) case-patients infected with non-O157 developed HUS (14). Although STEC O157 is more commonly associated with HUS worldwide (6), data on geographic differences in STEC serotypes suggest that in Australia, “non-O157:H7 STEC strains predominate,” and STEC O157:H7 is not as frequently implicated in “diarrhea-associated HUS” (16).
Overseas studies have reported higher proportions of STEC infections developing into HUS. In a cohort study of Argentinian children, aged \( \leq 15 \) years, 8 (8.6%) of 93 STEC patients developed HUS (17). Through enhanced surveillance in the Netherlands, Van Duynhoven et al. (18) found that HUS developed in 12 of 82 (14.6%) patients. Seventy-five percent of HUS case-patients were \( \leq 15 \) years (18). With the highest proportion from all reviewed studies, a Swiss linkage study found that HUS developed in 13 (29.5%) of 44 STEC patients, all of whom were \( \leq 15 \) years of age (19). Several studies on the incidence of HUS after STEC outbreaks have found that \( \approx 20\% \) of STEC cases develop into HUS (20–23). However, Sigmundsdottir et al. found no HUS cases among 9 STEC outbreak patients in Iceland (24) (Technical Appendix 1 Table 2).

A sequelae multiplier proportion of 3% (95% CI 1.7%–5.4%) was chosen, based on the South Australian study by Vally et al. (15). This study was chosen because STEC surveillance in South Australia is more complete than for other Australian states (11) and would therefore give a more representative estimate for Australia than the other available studies.

### Technical Appendix 1 Table 2. Incidence of HUS after STEC*

| Reference | Study years | Study type | Country | Age of HUS case-patients | No. HUS cases/no. STEC cases | STEC cases developing into HUS, % |
|-----------|-------------|------------|---------|--------------------------|-----------------------------|---------------------------------|
| Bradley et al. (20) | 2008 | Epidemiology investigation and case–control: after an outbreak | USA | Median: 46 y (range 1–88 y), 60% adult | 11/56 | 20 |
| Lopez et al. (17) | 2006 | Prospective cohort | Argentina | \( \leq 15 \) y | 8/93 | 8.6 |
| Neil et al. (21) | 2009 | Case–control: after an outbreak | USA | Not stated | 10/57 | 18 |
| Vally et al. (15) | 1997–2009 | Surveillance | Australia | Range: 5–60+, 60% aged \( \leq 15 \) y | 14/460 | 3 |
| Frank et al. (22) | 2011 | Surveillance: after an outbreak | Germany | Median 42, 88% aged >15 y | 845/3816 | 22 |
| Kappelli et al. (19) | 2000–2009 | Linkage | Switzerland | Median: 3.5 y (range 0–15 y) | 13/44 | 29.5 |
| McPherson et al. (14) | 2003–2007 | Case–control | Australia | Median: 4 y (range 1–62 y) | 8/113 | 7 |
| Sigmundsdottir et al. (24) | 2007 | Cohort: after an outbreak | Iceland | Not stated | 0/9 | 0 |
| Rangel et al. (25) | 1982–2002 | Outbreak surveillance | USA | Not stated | 354/8598 | 4.1 |
| Jay et al. (23) | 1999 | Epidemiology investigation and case–control: after an outbreak | USA | Not stated | 3/13 | 23 |
| Van Duynhoven et al. (18) | 1999–2001 | Enhanced surveillance | The Netherlands | Range: 0–70 y, 75% aged \( \leq 15 \) y | 12/82 | 14.6 |

*HUS, hemolytic uremic syndrome; STEC, Shiga toxin–producing Escherichia coli.*

**IBS**

There have been a few systematic reviews and/or meta-analyses on the association between intestinal infection and post-infectious IBS (PI-IBS). A recent review suggests the proportion of persons developing IBS following gastrointestinal infection is 4%–35% (26).
2010, Haagsma et al. (27) found that 1 year after infection from nontyphoidal *S. enterica* serotypes (hereafter referred to as nontyphoidal *Salmonella* spp.), nontyphoidal *Salmonella* spp., *Shigella* spp., or *Campylobacter* spp., IBS developed in 9% (95% CI 7.2–10.7) of patients. Similarly, in a systematic review of 18 studies, Thabane et al. (28) found a pooled incidence of PI-IBS of 10% (95% CI 9.4–85.6). Comparably, Halvorson et al. (29) reviewed 8 studies on nontyphoidal *Salmonella* spp., *Shigella* spp., bacterial unspecified, or unspecified, and their association with IBS, and calculated a median prevalence of IBS of 9.8% (interquartile range 4.0–13.3) in the exposed group and 1.2% Interquartile rate range 0.04–1.8) in the control group. A review by Smith and Bayles (30) found a mean prevalence of PI-IBS of 15% from 15 studies, with species of *Campylobacter*, nontyphoidal *Salmonella* spp., and/or *Shigella* as the most common agents of infection.

In the United Kingdom, Neal et al. (31) performed a postal survey and found that 25% of subjects had persistently altered bowel habits after bacterial gastroenteritis from nontyphoidal *Salmonella* spp., *Shigella* spp., or *Campylobacter* spp.; however, only 7% met the Rome criteria for new IBS. Also in the United Kingdom, Parry et al. (32) looked at the relationship between IBS and bacterial gastroenteritis from *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., *E. coli* O157, and *Aeromonas sobria*, and calculated an incidence of new IBS of 16.7% in the exposed group and 1.9% in the control group.

Studies looking at singular pathogens have also found an association between infectious gastroenteritis outbreaks and IBS. After an outbreak in 2002 in Spain, Mearin et al. (33) noted that before the outbreak, the prevalence of IBS was similar in case-patients and controls (2.9% vs. 2.3%); however, 3 months after the outbreak, IBS prevalence in case-patients had increased (9.2% vs. 1.7%), and 12 months after the outbreak, prevalence in case-patients remained higher (10.2% vs. 0.7%). The cumulative incidence was 7.4% at 3 months, 10.9% at 6 months, and 11.6% at 12 months. In Korea, 12 months after a *Shigella* spp. outbreak, Ji et al. (34) found that IBS had developed in 15 (14.9%) of 101 case-patients and 6 of 102 (5.9%) controls. In Canada, 2–3 years after an outbreak of *E. coli* O157:H7 and *Campylobacter* spp., 27.5% of 904 subjects with self-reported gastroenteritis reported IBS, and 36.2% of 464 subjects with clinically suspected gastroenteritis reported IBS (35). In a pediatric cohort from the Canadian outbreak, the cumulative incidence of PI-IBS for exposed subjects was 10.5% vs. a cumulative incidence in controls of 2.5% (36).
There have been studies on the association of *G. lamblia* with IBS; however, these have produced inconsistent results. While Wensaas et al. (37) found a high prevalence of IBS in exposed patients 2 years after acute giardiasis, Penrose et al. (38) found no linear association between *G. lamblia* and IBS, and a study by D’Anchino et al. (39) concluded that *G. lamblia* infection is a trigger for exacerbating preexisting IBS but could not conclude that *G. lamblia* causes IBS. PI-IBS has also been shown to develop after norovirus. Marshall et al. (40) performed a 2-year study after a norovirus outbreak; of the 89 respondents who reported an acute enteric illness during the outbreak and did not have preexisting IBS, 23.6% reported symptoms consistent with PI-IBS at 3 months versus 3.4% who reported symptoms but remained well during the outbreak. However, at 6, 12, and 24 months, the prevalence of IBS did not differ statistically among exposed and unexposed individuals, suggesting that PI-IBS might be more transient after viral gastroenteritis than it is after bacterial dysentery (40) (Technical Appendix 1 Table 3).

The meta-analysis by Haagsma et al. (27), which suggests that IBS develops in ≈9% (95% CI 7.2%–10.7%) of *Campylobacter* spp., nontyphoidal *Salmonella* spp., and *Shigella* spp. case-patients at 10–12 months of follow-up was chosen as the sequelae multiplier to simulate the plausible proportion of these bacterial pathogens that cause IBS using an alternate PERT distribution. While studies of multiple pathogens have found different rates of PI-IBS depending on etiology, this proportion was chosen for all 3 pathogens because it is a pooled rate that comes from a recent meta-analysis and is similar to PI-IBS rates after bacterial gastroenteritis that were reported in other studies (28,29,41).

| Reference                  | Year of publication | Study years | Country   | Study type            | Foodborne pathogen                                                                 | IBS patients after infectious gastroenteritis, % |
|----------------------------|---------------------|-------------|-----------|-----------------------|-----------------------------------------------------------------------------------|-----------------------------------------------|
| Koh et al. (41)            | 2012                | 2008–2010   | Korea     | Prospective cohort    | Nontyphoidal *Salmonella* spp., *Shigella* spp., STEC O157, *Vibrio cholerae*     | 9.2% at 3 mo†, 12.3% at 6 mo†                  |
| Wensaas et al. (37)        | 2012                | 2007–2008   | Norway    | Historic cohort       | *Giardia lamblia*                                                                  | 46.1% at 3 y                                  |
| Schwille-Kiuntke et al. (26)| 2011               | -           | Global    | Systematic review     | *Campylobacter* spp., *Escherichia coli*, *G. lamblia*, norovirus, nontyphoidal *Salmonella* spp., *Shigella* spp., *Trichinella britovi*, bacterial, viral, and parasitic gastroenteritis and travelers’ diarrhea | 4%–36% Incidence range                         |
| Thabane et al. (36)        | 2010                | 2002–2008   | Canada    | Outbreak study        | *E. coli* O157:H7, *Campylobacter* spp.                                           | 10.5%†                                        |
| Haagsma et al. (26)        | 2010                | -           | The Netherlands | Meta-analysis         | Campylobacter spp., nontyphoidal *Salmonella* spp.,                                | 9% (95% CI 7.2–10.7)                           |
| Reference                  | Year of publication | Study years | Country                     | Study type                          | Foodborne pathogen                                                                 | IBS patients after infectious gastroenteritis, % |
|----------------------------|---------------------|-------------|-----------------------------|------------------------------------|-----------------------------------------------------------------------------------|-----------------------------------------------|
| Marshall et al. (35)       | 2009                | 2002–2008   | Canada                      | Outbreak study                     | *Shigella* spp., *E. coli* O157:H7, *Campylobacter* spp.                           | 10% (95% CI 9.4–85.6), 4%–32% incidence range |
| Thabane et al. (28)        | 2007                | -           | Canada, China, Israel, Korea, New Zealand, UK, USA | Systematic review and meta-analysis | *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., confirmed bacterial gastroenteritis, and self-reported illness | 10% (95% CI 9.4–85.6), 4%–32% incidence range |
| Marshall et al. (40)       | 2007                | 2002–2004   | Canada                      | Outbreak study                     | Norovirus                                                                         | 23.6% at 3 mo                                  |
| Smith and Bayles (30)      | 2007                | -           | Canada, China, Korea, Spain, UK, USA | Systematic review                  | *Campylobacter* spp., *Cryptosporidium* spp., *E. coli*, *G. lamblia*, nontyphoidal *Salmonella* spp., *Shigella* spp., bacterial, and unspecified | 15% (range 3.4–31.6)‡                           |
| Halvorson et al. (29)      | 2006                | -           | Canada, China, Korea, Spain, UK, USA | Systematic review and meta-analysis | *Shigella* spp.                                                                   | 9.8% (IQR 4.0–13.3)‡                           |
| Ji et al. (34)             | 2005                | 2001–2002   | Korea                       | Outbreak study                     | *Shigella* spp.                                                                   | 14.9% at 1 y                                  |
| Mearin et al. (33)         | 2005                | 2002–2003   | Spain                       | Cohort study after an outbreak     | Nontyphoidal *Salmonella* spp.                                                    | 7.4% at 3 mo†, 10.9% at 6 mo†, 11.6% at 1 y† 16.7% at 6 mo† |
| Parry et al. (32)          | 2003                | 2000–2001   | UK                          | Prospective case–control study     | *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., STEC O157, *Aeromonas sobria* | 7% at 6 mo                                      |
| Neal et al. (31)           | 1997                | 1994        | UK                          | Cross-sectional                    | *Campylobacter* spp., nontyphoidal *Salmonella* spp., and *Shigella* spp.         |                                               |

*IBS, irritable bowel syndrome; IQR, interquartile range; nontyphoidal *Salmonella* spp., nontyphoidal *S. enterica* serotypes; STEC, Shiga toxin–producing *E. coli*.
†Cumulative incidence.
‡Median prevalence.

**ReA**

The causes of ReA are ambiguous because no formal definition or agreed-upon diagnostic criteria exist (42,43). Although the primary focus of the infection is usually through the gut or urogenital track, ReA has also been associated with respiratory pathogens (42). The classical gastrointestinal microbes resulting in ReA are *Yersinia enterocolitica*, nontyphoidal *Salmonella* spp., *Shigella* spp., and *Campylobacter* spp (43), and most agree that the term “ReA” should be applied only to infection caused by these gastrointestinal pathogens and *Chlamydia* spp (43); however, nonclassical ReA forms have been associated by a variety of other bacteria, including *Brucella* and *Staphylococcus*, and many authors have applied the term ReA for arthritis after infection with *C. difficile*, *Cryptosporidium*, *Giardia lamblia*, *E. coli*, and *Strongyloides* spp (43,44). With the majority of the literature focusing on the 4 classical gastrointestinal pathogens as triggers for ReA, we chose to use these to estimate the incidence of ReA due to contaminated food. If other enteric pathogens are in fact associated with ReA, our estimates of foodborne ReA may be conservative.
We were unable to find any published systematic reviews that report a global incidence rate for ReA after infection with the bacterial pathogens *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., and *Y. enterocolitica*. Because there are no diagnostic criteria for ReA, the case definition and the resulting incidences vary (42). The literature suggests that the incidence of ReA as a sequel to bacterial gastroenteritis varies by the enteric pathogen. For each of the bacterial enteric pathogens that precede ReA, we compiled papers that reported the proportion of cases that developed into ReA published in 2000 or later where all enteric cases were confirmed by a laboratory (Technical Appendix 1 Table 4). Because there is still quite a bit of variation in incidence in studies by pathogen, the median and range for *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., and *Y. enterocolitica* from the studies in Technical Appendix 1 Table 4 were calculated for the sequelae multiplier and used to simulate a distribution of the plausible proportion of cases that result in this sequel using an alternate PERT or PERT distribution, respectively. From the literature, we assume that 7% (range 2.8%-16%) of foodborne *Campylobacter* spp., 8.5% (range 0%-26%) of foodborne nontyphoidal *Salmonella* spp., 9.7% (range 1.2%-9.8%) of foodborne *Shigella* spp., and 12% (range 0%-23.1%) of foodborne *Y. enterocolitica* result in ReA. These distributions were then applied to the estimates of domestically acquired foodborne cases for each of the preceding bacterial pathogens.

### Technical Appendix 1 Table 4. ReA incidence* by foodborne pathogen, Australia, 2010

| Reference | Study years | Study type | Country | ReA cases/gastroenteritis cases |
|-----------|-------------|------------|---------|---------------------------------|
| Schonberg-Norio et al. (45) | 2002 | Cross sectional | Finland | 8/201 (4.0%) |
| Doorduyn et al. (46) | 2005 | Case–control | The Netherlands | 20/434 (4.6%) |
| Townes et al. (47) | 2002–2004 | Cohort | USA | 302/2384 (12.7%) |
| Schiellerup et al. (48) | 2002–2003 | Case–case comparison | Denmark | 131/1003 (13.1%) |
| Pope et al. (49) | 1966–2006 | Review | Europe | 1%–5% |
| Rees et al. (50) | 1998–1999 | Cohort | USA | 9/324 (2.8%) |
| Hannu (51) | 1997–1998 | Cohort | Finland | 45/609 (7.4%) |
| Locht and Krogh (52) | 1997–1999 | Cohort | Denmark | 27/173 (15.6%) |
| Arnedo-Pena et al. (53) | 2005 | Outbreak study | Spain | 6/67 (9%) |
| Doorduyn et al. (46) | 2005 | Case–control | The Netherlands | 8/181 (4.4%) |
| Townes et al. (47) | 2002–2004 | Cohort | USA | 204/1356 (15.0%) |
| Schiellerup et al. (48) | 2002–2003 | Case–case comparison | Denmark | 104/619 (16.8%) |
| Lee et al. (54) | 1999 | Outbreak study | Australia | 39/261 (14.6%) |
| Rees et al. (50) | 1998–1999 | Cohort | USA | 2/100 (2.0%) |
| Buxton et al. (55) | 1999–2000 | Case–control | Canada | 17/66 (25.7%) |
| Hannu et al. (56) | 1999 | Outbreak study | Finland | 5/63 (7.9%) |
| Rudwaleit et al. (57) | 1998 | Outbreak study | Germany | 0/286 (0%) (children only) |
| Urfer et al. (58) | 1993 | Outbreak study | Switzerland | 1/156 (0.6%) |
| Townes et al. (47) | 2002–2004 | Cohort | USA | 29/299 (9.7%) |
| Schiellerup et al. (48) | 2002–2003 | Case–case comparison | Denmark | 10/102 (9.8%) |
| Rees et al. (50) | 1998–1999 | Cohort | USA | 1/81 (1.2%) |
| Reference         | Study years | Study type          | Country    | ReA cases/gastroenteritis cases | ReA cases/Yersinia enterocolitica cases |
|-------------------|-------------|---------------------|------------|---------------------------------|----------------------------------------|
| Huovinen et al. (59) | 2006       | Case–control       | Finland    | 11/248 (4.4%)                  |                                        |
| Townes et al. (47)  | 2002–2004   | Cohort              | USA        | 5/35 (14.3%)                   |                                        |
| Schiellerup et al. (48) | 2002–2003  | Case–case comparison | Denmark    | 21/91 (23.1%)                 |                                        |
| Rees et al. (50)    | 1998–1999   | Cohort              | USA        | 0/8 (0%)                        |                                        |
| Hannu et al. (60)   | 1998        | Outbreak study      | Finland    | 4/33 (12.1%)                   |                                        |

*Incidence of ReA after Campylobacter spp. infection: median 7%, range 2.8%–16%; after Salmonella spp. infection: median 8.5%, range 0%–26%; after Shigella spp. infection: median 9.7%, range 1.2%–9.8%; after Yersinia enterocolitica infection: median 12%, range 0%–23.1%. ReA, reactive arthritis. Nontyphoidal Salmonella spp., nontyphoidal S. enterica serotypes.

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Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

Technical Appendix 2

Methods to Estimate Sequelae Incidence

For all 4 sequelae illnesses, we used data from notifiable surveillance (either national or state notifications) to estimate incidence of acute gastroenteritis due to relevant pathogens and then adjusted this using a sequelae multiplier, which is the proportion of bacterial infections that lead to sequelae illnesses (online Technical Appendix 1, http://wwwnc.cdc.gov/EID/article/20/11/13-1316-Techapp1.pdf). This approach is displayed in the Technical Appendix 2 Figure, where the left-hand column describes each input or output distribution, the central column illustrates the distribution, and the right-hand column describes the type and source of data underlying each input distribution. The final estimate is produced from a statistical model that incorporates uncertainty in case numbers in multipliers using probability distributions. That is, at each stage of the calculation, the estimate is represented by a probability distribution, and our final estimates and credible intervals are computed from this distribution. Further details on the estimation of incidence of acute illness due to each of the causal pathogens can be found in Kirk et al. (1).

The sequelae multiplier was modelled by using the PERT (Project Evaluation and Review Techniques) distribution, which is widely used for expert elicitation and risk assessment studies. It is based on the beta distribution and allows the input of minimum, maximum, and modal values. The alternate PERT distribution can be specified by 3 percentile points, such as a median value and 95% credible intervals (CrIs). Alternate PERT was used for the hemolytic uremic syndrome and irritable bowel syndrome sequelae multiplier, as the multiplier used was from another study that used median and 95% CIs. Alternate PERT was also used for reactive arthritis sequelae multipliers to enable a median value to be input, except in the case of the Shigella-associated reactive arthritis, where an alternate PERT distribution would not fit the data, and a PERT distribution was used instead. PERT allows for asymmetric distributions and can be easily produced from many data sources.
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Technical Appendix 2 Figure. Flowchart for the approach used to calculate the estimated number of sequelae cases in the community, Australia, circa 2010. PERT, project evaluation and review technique.
Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

Technical Appendix 3

Comparison with Estimates from 2000

Hall et al. estimated incidence, hospitalizations, and deaths for these 4 sequelae illnesses in Australia circa 2000 (1). Because methods and data sources have changed since the 2000 estimation effort, we recalculated incidence estimates for the sequelae in 2000 using our current methods and equivalent data from that earlier time period to validly compare rates over time. We used National Notifiable Disease Surveillance System data from 1996 to 2000 to recalculate the estimates for the incidence of all cases of gastroenteritis due to foodborne Campylobacter spp., nontyphoidal Salmonella enterica serotypes (hereafter referred to as nontyphoidal Salmonella spp.), Shigella spp., and Yersinia enterocolitica (2,3), and South Australian data from 1998–2000 (3) to recalculate the 2000 estimate for the incidence of gastroenteritis due to Shiga toxin–producing Escherichia coli (STEC). Further details on the method and recalculated circa 2000 estimates for Campylobacter spp., nontyphoidal Salmonella spp., and Shigella spp. can be found in the methods section and Table 3 of Kirk et al. (4). The estimates of foodborne illness from STEC and Y. enterocolitica for circa 2000 were calculated solely for this paper, using the same methods described in Kirk et al. (4) and the data described above.

Sequelae multipliers for the 2010 estimates were then applied to the recalculated 2000 estimates of incidence of acute gastroenteritis. The Technical Appendix 3 Table presents a comparison of the recalculated incidence estimates of sequelae of Guillain-Barré syndrome, hemolytic uremic syndrome, irritable bowel syndrome, and reactive arthritis for 2000 and 2010. Changes in sequelae illness from 2000 to 2010 reflect changes in the incidence of the preceding bacterial pathogen because the rate of sequelae after foodborne gastroenteritis, otherwise referred to as the sequelae multiplier, is assumed to be constant over this time period.
### Technical Appendix 3 Table. Comparison of incidence estimates and rates of 4 sequelae, Australia, circa 2000 and 2010*

| Illness | 2000 Incidence (90% CrI) | 2000 Rate per million (90% CrI) | 2010 Incidence (90% CrI) | 2010 Rate per million (90% CrI) | Rate ratio (90% CrI) |
|---------|-------------------------|-------------------------------|-------------------------|-------------------------------|----------------------|
| GBS     | 50 (25–100)             | 2.8 (1–6)                     | 70 (30–150)             | 3.1 (2–6)                     | 1.13 (0.5–3.6)       |
| HUS     | 55 (15–175)             | 3 (1–9)                       | 70 (25–200)             | 3 (1–9)                       | 1 (0.3–3.5)          |
| IBS     | 14,800 (9,500–23,500)   | 850 (550–1,350)               | 19,500 (12,500–30,700)  | 915 (570–1,440)               | 1.07 (0.5–2.0)       |
| ReA     | 12,500 (6,700–23,000)   | 730 (380–1,325)               | 16,200 (8,750–30,400)   | 765 (415–1,375)               | 1.06 (0.4–2.5)       |

*GBS, Guillain-Barré syndrome; HUS, hemolytic uremic syndrome; IBS, irritable bowel syndrome; ReA, reactive arthritis; CrI, credible interval.

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Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

Technical Appendix 4

Methods to Estimate Sequelae Hospitalizations and Deaths

To estimate hospitalizations due to irritable bowel syndrome (IBS) and reactive arthritis (ReA), we used hospitalization data for 2006–2010 from all Australian states and territories, using International Classification of Disease, Tenth Revision, Australian Modification (ICD-10-AM) codes. All estimated incident foodborne Campylobacter-associated Guillain-Barré syndrome (GBS) and Shiga toxin–producing Escherichia coli (STEC)–associated hemolytic uremic syndrome (HUS) cases were considered hospitalized, so were not modeled. The estimate for hospitalizations due to GBS and HUS is the estimate for GBS and HUS incidence. To estimate deaths for all 4 sequelae illnesses, we used national deaths data for 2001–2010 from the Australian Bureau of Statistics, using ICD-10 codes (Technical Appendix 4 Table 1). The final estimate included 2 multipliers, which are discussed below.

| Sequelae                | Mortality ICD-10 code and description | Hospitalization ICD-10-AM code and description |
|-------------------------|---------------------------------------|-----------------------------------------------|
| Guillain-Barré syndrome | G610: Guillain-Barré syndrome          |                                                |
| Hemolytic uremic syndrome | D593: Hemolytic uremic syndrome      |                                                |
| Irritable bowel syndrome | K58: Irritable bowel syndrome         | K58.0: Irritable bowel syndrome with diarrhea  |
|                         |                                       | K58.9: Irritable bowel syndrome without diarrhea|
| Reactive arthritis     | M021: Postdysenteric arthropathy      | M02.1: Postdysenteric arthropathy              |
|                         | M028: Other reactive arthropathies    | M02.3: Reiter’s disease                       |
|                         |                                       | M02.8: Other reactive arthropathies           |
|                         |                                       | M03.2: Other postinfectious arthropathies in diseases classified elsewhere|

*ICD-10-AM, International Classification of Diseases, Tenth Revision; AM, Australian Modification; --, all patients with incident cases are assumed to have been hospitalized so hospitalization data not used for this pathogen.

Domestically Acquired Multiplier

This multiplier adjusts for the proportion of case-patients who acquired infection in Australia with values for each sequelae in Technical Appendix 4 Table 2. For GBS, we adopted the domestically acquired multiplier for Campylobacter spp. (I). Given the relatively small numbers of notified cases of HUS, we adopted the domestically acquired multiplier for STEC (I). The domestically acquired multiplier for IBS was calculated as a weighted average of the
domestically acquired multipliers for *Campylobacter* spp., nontyphoidal *Salmonella enterica* serotypes (hereafter referred to as nontyphoidal *Salmonella* spp.), and *Shigella* spp., weighted by the total number of IBS cases for each pathogen. Similarly, the domestically acquired multiplier for ReA was calculated as a weighted average of the domestically acquired multipliers for *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp., and *Yersinia enterocolitica*, weighted by the total number of ReA cases for each pathogen.

### Technical Appendix 4 Table 2. Domestically acquired multipliers*

| Sequelae                          | Domestically acquired multiplier |
|-----------------------------------|----------------------------------|
| Guillain-Barré syndrome           | 0.97 (range 0.91–0.99)           |
| Hemolytic uremic syndrome         | 0.99 (range 0.93–1.00)           |
| Irritable bowel syndrome          | 0.91 (90% CrI 0.88–0.94)         |
| Reactive arthritis                | 0.91 (90% CrI 0.86–0.95)         |

*Crl, credible interval.

### Proportion Foodborne Multiplier

This multiplier adjusts for the proportion of illness that is acquired from food and was required only to estimate hospitalizations and deaths. Sequelae can arise from a source other than a bacterial pathogen, from a bacterial pathogen that was not foodborne, or from a foodborne pathogen. Only this latter category is considered a foodborne source. The proportion foodborne multiplier is the simulated product of the bacterial multiplier and the weighted foodborne multiplier and can be found in Technical Appendix 4 Table 3. The approach for calculating the proportion foodborne multiplier for each sequel is described as follows:

### Technical Appendix 4 Table 3. Proportion foodborne multiplier*

| Sequelae                          | Foodborne multiplier |
|-----------------------------------|----------------------|
| Guillain-Barré syndrome           | 0.25 (90% CrI 0.1–0.43) |
| Hemolytic uremic syndrome         | 0.33 (90% CrI 0.17–0.53) |
| Irritable bowel syndrome          | 0.13 (90% CrI 0.08–0.20) |
| Reactive arthritis                | 0.48 (90% CrI 0.36–0.62) |

*Crl, credible interval.

**GBS**

There have been several reviews, as well as many case–control and cross-sectional studies, that estimated the percentage of GBS cases attributable to *Campylobacter* spp. (Technical Appendix 4 Table 3). Poropatich et al. (8) performed a systematic review of 30 case–control studies and concluded that 31.0% of GBS cases might be attributable to a previous infection due to *Campylobacter* spp. (8). The other global systematic review of GBS incidence does not look at *Campylobacter* spp. specifically or perform a meta-analysis (9). Other (nonsystematic) reviews have found that 13%–72% (10) and 8%–50% (11) of GBS occurs as a sequel to campylobacteriosis. We assume that 31% (range 4.8%–72%) of cases of GBS arise
from *Campylobacter* spp. (2). Multiplied together with the *Campylobacter* spp. foodborne multiplier of 0.77 (90% CrI 0.62–0.89) (1) led to a foodborne multiplier for GBS of 0.25 (90% CrI 0.11–0.43).

**Technical Appendix 4 Table 4. Proportion of Guillain-Barré syndrome attributable to *Campylobacter* spp.*

| Reference          | Study years | Country  | Study type       | No. GBS cases | No. *Campylobacter* spp. cases based on GBS cases attributable to campylobacteriosis |
|--------------------|-------------|----------|------------------|---------------|--------------------------------------------------------------------------------------|
| Poropatich et al. (8) | 1982–2010  | Global   | Systematic review | 2,502         | Stool samples or serology 31% (range 4.8%–71.7%)                                      |
| McGrogan et al. (9) | 2006–2007   | Bangladesh | Systematic case-control | 100         | Stool samples and serology                                                       |
| Islam et al. (12)  | 2006–2007   | Global   | Prospective case-control | 237         | Stool samples and serology                                                       |
| Sivadon-Tardy et al. (13) | 1999–2005 | France   | Cross sectional | 553          | Corrected community incidence estimate Serology                                    |
| Tam et al. (14)    | 1991–2001   | UK       | Nested case-control | 263          | Serology 22%                                                                      |
| Sivadon-Tardy et al. (15) | 1996–2001 | France   | Cross sectional | 1049         | Stool samples and serology                                                       |
| Takahashi et al. (16) | 1990–2003 | Japan    | Case-control | 1146         | Community incidence estimate Serology                                                |
| Tam et al. (17)    | 2000–2001   | UK       | Estimation       | 88            | Stool samples or serology                                                        |
| Hadden and Gregson (10) | –           | Global   | Review           | 273/470      | Best estimate 30%–40% (range 8%–50%)                                               |
| Nachamkin et al. (11) | –           | USA      | Review           | 36/70        | Best estimate 30%–40% (range 8%–50%)                                               |

**HUS**

Technical Appendix 4 Table 5 presents the percentage of cases of HUS that arise from STEC estimated in 4 different papers, including a global systematic review. From this, we assumed that 61% (range 30%–85%) of HUS cases arise from STEC, modelled as a PERT distribution. Multiplied with the STEC foodborne multiplier of 0.56 (90% credible interval [CrI] 0.32–0.83) (1) led to a foodborne multiplier for HUS of 0.33 (90% CrI 0.18–0.54).

**Technical Appendix 4 Table 5. Proportion of HUS attributable to STEC**

| Reference          | Study years | Study type       | Country    | No. STEC isolations/no. STEC cases that develop into HUS |
|--------------------|-------------|------------------|------------|--------------------------------------------------------|
| Walker et al. (18) | 1980–2011   | Systematic review | Global     | 60.8% (range 30%–85.2%)                                  |
| Askar et al. (19)  | 2011        | Surveillance     | Germany    | 58%                                                     |
| Elliot et al. (20) | 1994–1998   | Surveillance     | Australia  | 51%                                                     |
| Van de Kar (21)    | 1989–1993   | Case control     | The Netherlands | 77.8%                                                 |

**IBS**

We estimated the proportion of IBS cases from *Campylobacter* spp., nontyphoidal *Salmonella* spp., or *Shigella* spp. based on the proportion of IBS considered to be postinfectious in the literature. In 1962, Chaudhary and Truelove (22) reported IBS occurring from infective dysentery, with 34 (26.2%) of 130 patients dating symptoms back to an attack of gastroenteritis.
More recently, review studies have estimated that 6%-17% (23) and 7%–33% of IBS is postinfectious (24). In the meta-analysis and estimation by Haagsma et al. (25), the authors considered that 17% of IBS is due to campylobacteriosis, salmonellosis, or shigellosis from the top end of the range of 6%-17% by Spiller and Garsed (23). We assumed 17% of IBS to be triggered by a gastrointestinal infection (25), with a range of 7%–33% from the review by Schwille-Kiuntke et al. (24). Because more than just Campylobacter spp., nontyphoidal Salmonella spp. and Shigella spp. can cause postinfectious IBS, this may be an overestimate.

A foodborne multiplier for the combined 3 pathogens of 73% (90% CrI 64%–82%) was calculated as a weighted average of the foodborne multipliers for each pathogen, weighted by the total number of IBS cases for each pathogen. Multiplied by the above PERT distribution of 17% (range 6%–33%), gave a foodborne multiplier for IBS of 13% (90% CrI 8%–20%).

### Technical Appendix 4 Table 6. Proportion of IBS attributable to infectious gastroenteritis*

| Reference              | Publication year | Study type           | Country         | No. postinfectious IBS cases/IBS cases | IBS that is postinfectious, % |
|------------------------|------------------|----------------------|-----------------|----------------------------------------|------------------------------|
| Chaudhary and Truelove (22) | 1962             | Epidemiologic report | UK              | 34/130                                  | 26.2                         |
| Spiller and Garsed (23)  | 2009             | Review               | Global          | –                                       | 6–17                         |
| Haagsma et al. (25)     | 2010             | Meta-analysis and estimation | The Netherlands | –                                       | 17                           |
| Schwille-Kiuntke et al. (24) | 2013             | Review               | Global          | –                                       | 7–33                         |

*IBS, irritable bowel syndrome. Boldface indicates chosen proportion for foodborne multiplier calculation.

ReA

In a review of ReA, Hannu et al. (4) compiled population-based studies on the annual incidence of ReA—both from enteric and urogenital infection. We used this compilation and calculated the proportion of ReA due to enteric infection by dividing the enteric incidence by the total incidence found in each study (Technical Appendix 4 Table 7). We used the midpoint and range of the proportions from these studies for the bacterial multiplier. We therefore assumed a median of 66.7% of ReA is due to an enteric infection, with a range of 50%–94.7%. If enteric infections preceding ReA are from other infections besides campylobacteriosis, salmonellosis, shigellosis, or yersiniosis, using this distribution to estimate ReA cases from these infections may cause an overestimation.

We adjusted for the proportion foodborne using a weighted average of the foodborne multipliers for Campylobacter spp., nontyphoidal Salmonella spp., Shigella spp., and Y. enterocolitica, weighted by the total number of ReA cases for each pathogen. This gave a foodborne multiplier of 72% (90% CrI 60%–82%). Multiplied by the above alternate PERT
distribution of median 66.7% (range 50%–94.7%), gave a foodborne multiplier for reactive arthritis of 48% (90% CrI 36%–61%).

| Reference            | Country        | Year | Incidence per 100,000 | Enteric | Urogenital | Total | No. ReA due to enteric infection/total no. enteric infections |
|----------------------|----------------|------|-----------------------|---------|------------|-------|-----------------------------------------------------------------|
| Isomaki et al. (26)  | Finland        | 1978 | 14                    | 13      | 27         | 14/27 (51.9%)       |
| Kvien et al. (27)    | Norway         | 1994 | 5                     | 5       | 10         | 5/10 (50%)          |
| Savolainen et al. (28)| Finland       | 2000 | 7                     | 3       | 10         | 7/10 (70%)          |
| Soderlin et al. (29) | Sweden         | 2002 | 18                    | 1       | 19         | 18/19 (94.7%)       |
| Townes et al. (30)   | USA            | 2008 | 0.6–3.1               | NA      | NA         | NA                |
| Hanova et al. (31)   | Czech Republic | 2010 | 6                     | 3       | ≈9         | 6/9 (66.7%)         |

*Adapted from the table of annual incidence of reactive arthritis based on population studies in Hannu et al. (4). NA, not applicable.

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Sequelae of Foodborne Illness Caused by 5 Pathogens, Australia, Circa 2010

Technical Appendix 5

Model Inputs for 4 Sequelae Illnesses Due to Contaminated Food

Incidence

Technical Appendix 5 Table 1. Guillain-Barré Syndrome

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Antecedent bacterial gastroenteritis cases: estimated number of foodborne *Campylobacter* spp. cases (1) | Outcome | 5%, median, 95% values: 108500, 179000, 290000 (circca 2010) 82500, 139000, 227000 (circca 2000) |
| Sequelae multiplier: this proportion was a midpoint between estimates from the literature reported in Tam et al. (2), McCarthy and Gieseke (3), and Allos et al. (4) | PERT | Minimum, modal, maximum values: 0.000192, 0.000304, 0.000945 |
| Total foodborne illness: foodborne *Campylobacter* spp. cases × Sequelae multiplier | Outcome | 5%, median, 95% values: 30, 75, 150 (circca 2010) 25, 50, 100 (circca 2000) |
| Rate of foodborne illness from *Campylobacter* spp. per million population | Outcome | 5%, median, 95% values: 2, 3.1, 6 (circca 2010) 1, 2.8, 6 (circca 2000) |

Technical Appendix 5 Table 2. Hemolytic uremic syndrome

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Antecedent bacterial gastroenteritis cases: estimated number of foodborne STEC cases (1) | Outcome | 5%, median, 95% values: 950, 2350, 5850 (circca 2010) 550, 1900, 5000 (circca 2000) |
| Sequelae multiplier: this proportion is from Vally et al. (5) | Alternate PERT | 2.5%, Median, 97.5% values: 0.017, 0.03, 0.051 |
| Total foodborne illness: foodborne STEC cases × Sequelae multiplier | Outcome | 5%, median, 95% values: 25, 70, 200 (circca 2010) 15, 55, 175 (circca 2000) |
| Rate of foodborne illness from STEC per million | Outcome | 5%, median, 95% values: 1, 3.3, 9 (circca 2010) 1, 3.0, 9 (circca 2000) |

*STEC, Shiga toxin–producing *Escherichia coli*.

Technical Appendix 5 Table 3. Irritable bowel syndrome

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Antecedent bacterial gastroenteritis cases: Estimated number of foodborne *Campylobacter* spp. cases (1) | Outcome | 5%, median, 95% values: 108500, 179000, 290000 (circca 2010) 82500, 139000, 227000 (circca 2000) |
| Estimated number of foodborne nontyphoidal *Salmonella* spp. cases (1) | Outcome | 5%, median, 95% values: 21200, 39600, 73400 (circca 2010) 15000, 28000, 50000 (circca 2000) |
| Estimated number of foodborne *Shigella* spp. cases (1) | Outcome | 5%, median, 95% values: 150, 350, 850 (circca 2010) 150, 515, 1300 (circca 2000) |
| Sequelae multiplier: This proportion was from Haagsmsa et al. (6) | Alternate PERT | 2.5%, Median, 97.5% values: 0.072, 0.088, 0.104 |
| Total foodborne illness Foodborne *Campylobacter* spp. cases × Sequelae multiplier | Outcome | 5%, median, 95% values: 12500, 19500, 30700 (circca 2010) |
### Hospitalizations and Deaths

**Technical Appendix 5 Table 5. Guillain-Barré syndrome**

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Average number of deaths per year: Australian Bureau of Statistics death data | Empirical | 2001–2010: 24.5 |
| Population adjustment: Australian resident population June quarter, [http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0De c%202011?OpenDocument](http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0Dec%202011?OpenDocument) [cited 2012 Aug 16] | Empirical | By year (2001–2010): 19413240, 19651438, 19885435, 20127363, 20394791, 20697880, 21015936, 21384427, 21778845, 22065317 |
| Domestically acquired multiplier: *Campylobacter* spp. domestic acquired multiplier | PERT | Minimum, modal, maximum values: 0.91, 0.97, 0.99 |
| Foodborne multiplier: derived from: | Outcome | 5%, median, 95% values: 0.1, 0.25, 0.43 |
| Bacterial multiplier—the proportion of Guillain-Barré syndrome that is attributable to *Campylobacter* spp. from Poropatich et al. (7) × *Campylobacter* spp. foodborne proportion (1) | PERT | Minimum, modal, maximum values: 0.048, 0.31, 0.717 |
| | Alternate | 5%, median, 95% values: 0.62, 0.77, 0.89 |
| Total foodborne deaths: circa 2010 | Outcome | 5%, median, 95% values: 2, 6, 10 |
| Rate of foodborne deaths per million: circa 2010 | Outcome | 5%, median, 95% values: 0.1, 0.3, 0.5 |

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### Campylobacter attributable to multipliers + Foodborne Model

**Technical Appendix 5 Table 6. Hemolytic uremic syndrome**

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Rate of foodborne illness per million | Outcome | 5%, median, 95% values: 570, 915, 1440 (circa 2010) 550, 850, 1350 (circa 2000) |

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### Technical Appendix 5 Table 4. Reactive arthritis

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Antecedent bacterial gastroenteritis cases: | | |
| Estimated number of foodborne *Campylobacter* spp. cases (1) | Outcome | 5%, median, 95% values: 108500, 179000, 290000 (circa 2010) 82500, 139000, 227000 (circa 2000) |
| Estimated number of foodborne nontyphoidal *Salmonella* spp. cases (1) | Outcome | 5%, median, 95% values: 21200, 39600, 73400 (circa 2010) 15000, 28000, 50000 (circa 2000) |
| Estimated number of foodborne *Shigella* spp. cases (1) | Outcome | 5%, median, 95% values: 150, 350, 850 (circa 2010) 175, 515, 1300 (circa 2000) |
| Estimated number of foodborne *Yersinia enterocolitica* cases (1) | Outcome | 5%, median, 95% values: 650, 1150, 1950 (circa 2010) 300, 800, 1650 (circa 2000) |

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Sequela multipliers: The proportion for each of the 4 pathogens was calculated from the literature. See Technical Appendix 1 for further explanation.

- **Campylobacter** spp. sequelae multiplier
  - Alternate
  - PERT
  - Minimum, median, maximum values: 0.028, 0.07, 0.16

- **Nontyphoidal *Salmonella*** spp. sequelae multiplier
  - Alternate
  - PERT
  - Minimum, median, maximum values: 0, 0.085, 0.26

- **Shigella** spp. sequelae multiplier
  - PERT
  - Minimum, modal, maximum values: 0.012, 0.097, 0.098

- **Yersinia enterocolitica** sequelae multiplier
  - Alternate
  - PERT
  - Minimum, median, maximum values: 0, 0.12, 0.231

**Total foodborne illness:**

- Foodborne *Campylobacter* spp. cases × Sequelae multiplier + Foodborne nontyphoidal *Salmonella* spp. cases × Sequelae multiplier + Foodborne *Shigella* spp. cases × Sequelae multiplier + Foodborne *Y. enterocolitica* cases × Sequelae multiplier
  - Outcome
  - 5%, median, 95% values: 8750, 16200, 30400 (circa 2010) 6700, 12500, 23000 (circa 2000)

**Rate of foodborne illness from *Campylobacter* spp., nontyphoidal *Salmonella* spp., *Shigella* spp. and *Y. enterocolitica* per million:

- Outcome
  - 5%, median, 95% values: 415, 765, 1375 (circa 2010) 380, 730, 1325 (circa 2000)
### Page 3 of 4

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**Technical Appendix 5 Table 7. Irritable bowel syndrome**

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Average number of deaths per year: Australian Bureau of Statistics death data | Empirical | By year (2001–2010): 4,2 |
| Population adjustment: Australian resident population June quarter, [http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0De c%202011?OpenDocument](http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0Dec%202011?OpenDocument) [cited 2012 Aug 16] | Empirical | By year (2001–2010): 19413240, 19651438, 19895435, 20127363, 20394791, 20697880, 21015936, 21384427, 21778845, 22065317 |
| Domestically acquired multiplier: STEC domestically acquired multiplier | PERT | Minimum, modal, maximum values: 0.93, 0.9, 1 |
| Foodborne multiplier: derived from: Bacterial multiplier—the proportion of HUS that is attributable to STEC from Walker et al. (8) x STEC foodborne proportion (1) | Outcome | 5%, median, 95% values: 0.17, 0.33, 0.53 |
| Total foodborne deaths: circa 2010 | Outcome | 5% | median, 95% values: 0.03, 0.1, 0.12 |

*HUS, hemolytic uremic syndrome; STEC, Shiga toxin–producing Escherichia coli.*

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**Technical Appendix 5 Table 8. Reactive arthritis**

| Model input, source, and comments | Distribution | Data for model input |
|-----------------------------------|--------------|----------------------|
| Average number of deaths per year: Australian Bureau of Statistics death data | Empirical | By year (2006–2010): 63, 50, 50, 70, 70 |
| Population adjustment: Australian resident population June quarter, [http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0De c%202011?OpenDocument](http://www.abs.gov.au/AUSSTATS/abs@.nsf/DetailsPage/3101.0Dec%202011?OpenDocument) [cited 2012 Aug 16] | Empirical | By year (2001–2010): 19413240, 19651438, 19895435, 20127363, 20394791, 20697880, 21015936, 21384427, 21778845, 22065317 |
| Domestically acquired multiplier: a weighted multiplier from *Campylobacter* spp., nontyphoidal *Salmonella* spp., and *Shigella* spp. domestically multipliers | Alternate | 5%, median, 95% values: 0.88, 0.91, 0.94 |
| Foodborne multiplier: derived from: Bacterial multiplier—proportion of IBS that is post-infectious extracted from the literature (6,9) x weighted *Campylobacter* spp., nontyphoidal *Salmonella* spp., and *Shigella* spp. foodborne proportion (1) | Alternate | 5%, median, 95% values: 0.06, 0.17, 0.33 |
| Total foodborne hospitalizations: circa 2010 | Outcome | 5%, median, 95% values: 25, 43, 70 |
| Rate of foodborne hospitalizations per million: circa 2010 | Outcome | 5%, median, 95% values: 0.05, 0.1, 0.11 |

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