Risk factors for infection with *Campylobacter jejuni* flaA genotypes

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SUMMARY

We aimed to explore *Campylobacter* genotype-specific risk factors in Australia. Isolates collected prospectively from cases recruited into a case-control study were genotyped using *flaA* restriction fragment-length polymorphism typing (*flaA* genotyping). Exposure information for cases and controls was collected by telephone interview. Risk factors were examined for major *flaA* genotypes using logistic and multinomial regression. Five *flaA* genotypes accounted for 325 of 590 (55%) cases — *flaA*-6b (*n* = 129), *flaA*-6 (*n* = 70), *flaA*-10 (*n* = 48), *flaA*-2 (*n* = 43), *flaA*-131 (*n* = 35). In Australia, infections due to *flaA*-10 and *flaA*-2 were found to be significantly associated with eating non-poultry meat (beef and ham, respectively) in both case-control and inter-genotype comparisons. All major genotypes apart from *flaA*-10 were associated with chicken consumption in the case-control comparisons. Based on several clinical criteria, infections due to *flaA*-2 were more severe than those due to other genotypes. Thus genotype analysis may reveal genotype-specific niches and differences in virulence and transmission routes.

INTRODUCTION

The disease burden of *Campylobacter* infections is considerable in Australia, where it is the most common notifiable disease [1]. *Campylobacter* infections are largely sporadic and estimated to cause about 223 000 cases of gastroenteritis each year [2]. The need for public health intervention is highlighted by the steady rise in notifications in Australia since the early 1990s [1, 3].

Case-control studies of potential risk factors for campylobacteriosis have been undertaken in several countries [4–11] including Australia [12, 13]. Consumption of chicken is the most commonly identified risk factor for sporadic campylobacteriosis. However, this and other statistically significantly risk factors often do not explain the majority of cases.
Further sources of *Campylobacter* infection may be detected if strains with distinct ecologies (including differential survival characteristics, and therefore specific environmental niches or host preference), with varying virulence (causing disease of variable severity) and different transmission routes could be distinguished. Numerous subtyping methods have been developed and applied to *Campylobacter* isolates [15–19]. These methods, used to determine genomic diversity, generally have been applied to diverse collections of isolates [20, 21]. Methods such as serotyping, multilocus sequence typing (MLST) and pulsed-field gel electrophoresis (PFGE) have been used to distinguish outbreak from sporadic isolates [18, 22, 23]. In addition, a variety of methods have been used to compare animal and human isolates. Some studies have concluded that finding the same subtype in animal and human isolates is evidence of transmission from animals to humans or is evidence of a common source for animals and humans [17, 24–28]. However, the potential to identify genotype-specific risk factors for sporadic infection is yet to be evaluated.

The aim of this study was to determine sources and risk factors of *Campylobacter jejuni* for specific *flaA* genotypes. We hypothesize that the application of *flaA* genotyping to isolates from a case-control study may allow detection of further sources through the examination of genotype-specific risk factors. Study isolates were collected prospectively from sporadic cases recruited into a case-control study and genotyped using *flaA* restriction fragment-length polymorphism (RFLP; *flaA* genotyping) analysis, a moderate throughput, low cost method [29] with good correlation with MLST [30]. Genotyping data were linked to exposure data for the investigation of risk factors.

**METHODS**

**Study base**

Data and *Campylobacter* isolates collected for this study were drawn from an Australian case-control study that was conducted in five states between September 2001 and September 2002. A description of the study and risk factors for subjects aged ≥5 years was reported by Stafford *et al.* [13]. The study recruited cases and controls from five of the eight states and territories in Australia. The largest state, New South Wales, was excluded because campylobacteriosis is not notifiable there and the Northern Territory and Australian Capital Territory were not included in the study because too few cases are notified. As previously described [13], each site aimed to prospectively recruit 200 cases of all ages using systematic sampling from a notifiable disease register; every second case from Tasmania and Victoria, every fourth and sixth case from Queensland and South Australia, respectively and every case from Western Australia. The sampling strategy was based on the expected number of cases notified from participating laboratories and the number required to detect significant associations with hypothesized risk factors [13]. Furthermore, sample size calculations determined that about 550 isolates would probably be collected and that number would prove sufficient to detect significant associations between some genotypes and risk factors.

Cases were defined as individuals with diarrhoea (≥3 or more loose stools in a 24 h period), who had culture-confirmed *C. jejuni* infection (and no other pathogen such as *Salmonella*, *Shigella*, or the enteric protozoa detected), whose isolates were *flaA* genotyped, whose stool samples were collected within 10 days of diarrhoea onset and who were interviewed within 30 days of onset. Telephone interviews were conducted when verbal consent was given by the study subject (or carer if the case was aged <16 years).

Controls were drawn (about one per case) from a control bank generated during a national cross-sectional survey of gastroenteritis conducted in 2001; a description of the survey is given by Hall *et al.* [31] and a description of control selection is reported by Stafford *et al.* [13]. In brief, households were selected using random digit dialling during the gastroenteritis survey, a household member was asked to participate in the survey and subsequently consent to be part of a control bank. From this control bank, potential controls were selected and frequency-matched to cases by age bands (0–4, 5–9, 10–19, 20–29, 30–59 and ≥60 years) in each state at a ratio of 1:1. If a person did not wish to participate or was excluded on the criteria (described below), a subsequent person was sought from the control bank. Once a control had been selected from a household, that household was no longer eligible for future selection of controls. Controls were interviewed within 30 days of interview of a notified case.

Cases and controls were excluded if they did not have a phone number, were unable to be contacted...
after at least six attempts, they/their parents were non-English-speakers, they could not answer questions, or if a household member had had diarrhoea or a confirmed Campylobacter infection in the 4 weeks prior to onset. In addition, cases were excluded if they were unable to recall the diarrhoea onset date or they were part of an outbreak.

**Questionnaire**

A standard questionnaire was administered by telephone to obtain information on a range of variables, including host factors (underlying illnesses, prior consumption of antimicrobial agents, antacids and immune suppressive therapies), overseas travel, dining locations, consumption of water and food (fruit and vegetables, meat, poultry, seafood, eggs and dairy products), animal and pet exposures, and demographics. Cases were asked additional questions about their illness and treatment. For cases, all questions related to the 7-day period before their onset of diarrhoea, except for prior use of antibiotics, antacids or immune suppressive therapies, which were based on the preceding 4 weeks. Exposure information was not collected from subjects who had travelled outside Australia during the 7-day exposure period. Controls were asked about the 7 days or 4 weeks prior to interview.

**Laboratory methods**

**Isolates**

Isolates from diarrhoeal stool cultures were stored and subsequently identified to species level by hippurate hydrolysis and PCR as described previously [32, 33]. Those identified as *C. jejuni* using these methods were flaA genotyped.

**flaA genotyping**

Flagellin A RFLP typing (*flaA* genotyping) was performed according to the method described by Nachamkin *et al.* [15]. Briefly, this involved PCR amplification of *flaA*, followed by digestion of products with the restriction enzyme *DdeI* and separation of fragments by agarose gel electrophoresis. Genotyping was performed by five laboratories, using standardized reagents and methods (including DNA extraction, PCR, restriction enzyme digestion, agarose gel electrophoresis and photography), which had been optimized by one laboratory.

**Quality assurance**

A set of eight isolates was distributed to the laboratories that genotyped isolates. Tiff images of *flaA* genotyping gels were loaded onto a BioNumerics database located at one laboratory and examined for comparability. Feedback was provided on accuracy of the patterns and image quality. Reaction or photography conditions were modified, if necessary, to produce images consistent with those from prior testing of the quality assurance set. The positive control, NCTC 11351, was included in each test run. GeneRuler™ 100 bp DNA LadderPlus (MBI Fermentas, Vilnius, Lithuania) was included in lanes 1, 5, 10, and 15 of all gels. If the gel pattern of the positive control was not compatible with those included in the BioNumerics database, gel images were to be rejected, however, this did not occur.

**Assignment of flaA genotypes**

Tiff images of gels were loaded onto a BioNumerics database and patterns were normalized according to molecular-weight standards on each gel. *flaA* genotypes were designated by a number that was assigned arbitrarily; numbered genotypes differed from each other by at least two bands and subtypes of numbered genotypes, designated by a letter (e.g. *flaA*-6 and *flaA*-6b) differed by a single band only. The numbering of genotypes was consistent with a previous Australian study [29, 30] and isolates from the previous study were included in the BioNumerics library used for analysis in this study. Genotypes were grouped together using the Dice band matching coefficient and UPGMA clustering method with a position tolerance of 1% and an optimization of 1% which clustered at >90% similarity [29]. Resultant dendrograms were checked visually by two researchers and about 10% were re-verified by the second researcher in the case of discrepancies.

**Statistical analyses**

Logistic regression analysis was used to compare demographic characteristics and host factors between study cases and (a) study controls, (b) cases that did not have isolates *flaA* genotyped and (c) cases notified through the national surveillance system. Logistic regression analysis was also used to identify potential risk factors for specific *flaA* genotypes. The exposures reported for cases of each major *flaA* genotype were compared to those for all study
controls combined, in order to increase the power of hypothesis testing. A final model for each genotype was constructed by including all exposure variables with \( P < 0.1 \) (in univariate analyses) and using backwards stepwise elimination, controlling for confounders (demographic and host factors). Models were tested for goodness of fit and compared using the likelihood ratio test. Genotype-specific population-attributable fractions (PAF) were calculated for each risk factor from final models for each of the major \( flaA \) genotypes and for the group comprising ‘other’ \( flaA \) genotypes.

To allow for the possibility that some food and environmental exposures were location-specific, we included terms for the interaction between exposure and state (as a categorical variable) in the logistic regression models. The significance of multiple interaction terms was tested using the likelihood ratio test. Only significant interactions are reported.

Multinomial regression was applied to data on cases to explore differences in exposure variables for the major \( flaA \) genotypes, using ‘other’ \( flaA \) genotypes, comprising the remaining study cases, as the reference group, controlling for confounders. This type of analysis was also used to compare demographic and clinical characteristics between cases infected with each of the major \( flaA \) genotypes. Results are expressed as relative risk ratios (RRRs), as is appropriate for this case-only analysis [34].

Analyses were performed using Stata version 9.1 (Stata Corporation, College Station, TX, USA).

RESULTS

Recruitment of cases and controls

During the study period there were 8847 *Campylobacter jejuni* notifications in the five participating states and, of these, 1019 cases were recruited and interviewed (12%). There were 590 (58%) cases for which an isolate was stored, subsequently found to be *C. jejuni*, and \( flaA \) genotyped, representing 7% of notifications. There were no significant differences in age distribution between 590 study cases and cases notified in 2001 but fewer males were included (52-5% among cases vs. 54-4% for notifications, \( P = 0.03 \)). When cases for whom isolates were genotyped were compared to cases that did not have an isolates genotyped for demographic, host factor and clinical characteristics, a greater number of the former were educated to school level only [37% vs. 28%; odds ratio (OR) 1.5, 95% confidence interval (CI) 1.1–2.0] but no further differences were detected.

A total of 967 controls were recruited and there were statistically significant, but modest, differences between study cases and controls with respect to sex, income, place of residence and use of acid-reducing medications (Table 1).

flaA genotype distribution

Among 590 isolates, there were 61 different \( flaA \) genotypes, of which five accounted for 325 (55%); an image of the electrophoretic patterns of the major genotypes is shown in the Fig.), and 21 (4%) were single isolates. The five major \( flaA \) genotypes comprised \( flaA-6b \) (\( n = 129 \), 22%), \( flaA-6 \) (\( n = 70 \), 12%), \( flaA-10 \) (\( n = 48 \), 8%), \( flaA-2 \) (\( n = 43 \), 7%), \( flaA-131 \) (\( n = 35 \), 6%); the remaining 265 study cases comprised the ‘other’ genotype group used in case-only analyses. Some geographic differences were noted for the major genotypes: \( flaA-10 \), and -131 were identified in all states, \( flaA-6b \) and -2 were found in four of the five states but \( flaA-6b \), the most common genotype, was not found in South Australia and \( flaA-6 \) was found in South Australia and Queensland only. Of the major \( flaA \) genotypes, only \( flaA-2 \), and -10 were detected among the 13 isolates from overseas travellers (data not shown). The full descriptive epidemiology of \( flaA \) genotypes and detection of clusters will be reported separately.

The five major \( flaA \) genotypes (\( flaA-2 \), -6, -6b, -10, and -131) were analysed separately for risk factors.

Patient characteristics and symptom profile for the major \( flaA \) genotypes

Comparison of cases due to each of the five major \( flaA \) genotypes with those due to all ‘other’ genotypes using multinomial regression showed no differences in the following characteristics (proportions for all study cases are shown in parentheses): proportion of males (53%), those with cramps (89%), persistent diarrhoea (12%), those that were hospitalized (13%), those treated with anti-diarrhoeal medications (48%) or intravenous fluids (12%). A significantly higher proportion of subjects with \( flaA-2 \), compared to ‘other’ genotypes, had fever (RRR 2.3, 95% CI 1.1–5.4), bloody diarrhoea (RRR 2.2, 95%
Sources of C. jejuni and risk factors

Subjects that had travelled internationally were included in the study but exposure information was not collected. When compared to controls, overseas travel was significantly associated with flaA-10 disease (OR 14·5, 95% CI 2·3–85·7). A total of 66 exposure variables were examined in univariate analyses of cases infected with the major flaA genotypes who acquired their infections locally, compared to controls.
Table 2. Final multiple logistic regression models* for exposures associated with major Campylobacter jejuni flaA genotypes

| Exposure† | flaA-2§ | flaA-6 | flaA-6b || flaA-10* | flaA-131# | Other** |
|-----------|---------|--------|---------|---------|----------|---------|
|           | (n = 43) | (n = 129) | (n = 48) | (n = 35) | (n = 265) |
| Chicken   | 3.6 (1.3–10.3) | 6.5% (11.6–86.5%) | | 3.6 (1.1–12.9) | 3.6 (1.9–6.6) | |
| Undercooked | 4.7 (1.7–13.4) | 17.0% (2.2–30.9%) | | 3.6 (1.1–12.9) | 3.6 (1.9–6.6) | |
| Barbecued chicken | 1.9 (1.2–2.9) | 16.8% (3.5–28.3%) | | 2.6 (1.2–5.5) | 17.2% (0.5–31.2) | |
| Undercooked beef | | | | 2.6 (1.2–5.5) | 17.2% (0.5–31.2) | |
| Diced beef | 0.3 (0.1–0.7) | 0.6 (0.3–0.9) | | 0.6 (0.3–0.8) | | |
| Veal | 0.3 (0.1–0.9) | 0.3 (0.1–0.9) | | 0.6 (0.3–0.8) | | |
| Offal | 2.9 (1.1–7.9) | 11.4 (4.1–31.8) | | 2.5 (1.1–6.0) | 2.5 (1.1–6.0) | |
| Paté | 2.2 (1.1–4.6) | 3.3 (1.2–9.6) | | 2.5 (1.1–6.0) | | |
| Ham | 2.4 (1.1–5.2) | 39.3% (10–62.8) | | 0.5 (0.3–0.9) | 2.5 (1.1–6.0) | 6.0% (2.0–10.0) |
| Organic fruit/vegetables | 0.4 (0.2–0.9) | 0.4 (0.2–0.9) | | 0.5 (0.3–0.9) | 2.5 (1.1–6.0) | 6.0% (2.0–10.0) |
| Bottle water | | | | | | |
| Puppy | 4.0 (1.1–14.8) | 9.0% (n.c.) | | 2.5 (1.1–6.0) | | |
| Pet chicken | 2.1 (1.1–4.3) | 5.3% (n.c.) | | 2.5 (1.1–6.0) | | |
| Pet chicken < 6 months | 20.6 (3.0–117.4) | 8.2% (0.4–15.4) | | 10.2 (2.6–40.4) | 13.4% (0.8–24.4) | |
| Farm birds | 11.3 (2.5–51.9) | [5.8%, n.c.] | | | | |
| Poor barbeque technique | 4.4 (1.2–16.3) | 8.7% (n.c.) | | 4.6 (1.2–18.4) | 7.6% (n.c.) | 6.9% (n.c.) |

OR, Odds ratio, CI, confidence interval, n.c., 95% CI unable to be calculated due to small numbers.
* Logistic regression models controlled for age, sex (and state, when appropriate).
† Logistic regression models controlled for age, sex (and state, when appropriate).
‡ Population-attributable fraction shown in bold within square brackets with 95% CI determined for cases infected with each flaA genotype; blank cells indicate that no statistically significant association was found for the respective flaA genotype and exposure.
§ Exposure period of 7 days.
¶ Model for this flaA genotype included state.
|| Model for this flaA genotype included consumption of acid-reducing medications.
* Model for this flaA genotype included chronic gastrointestinal illness and immunosuppressive therapies.
# Model for this flaA genotype included age and place of residence.
** All remaining flaA genotyped cases. Model for this flaA genotype included chronic gastrointestinal illness.
In final multivariate models, constructed to explain exposures associated with locally acquired disease (Table 2), infection with *flaA*-6b, the most common *flaA* genotype, was independently associated with consumption of barbecued chicken, offal, paté and exposure to pet chickens. The second most common *flaA* genotype, *flaA*-6, was independently associated with consuming chicken and exposure to farm birds (ducks, geese, etc.). Infection with *flaA*-10 was associated with undercooked beef, offal, paté, exposure to young pet chickens (aged <6 months), and poor food handling. *flaA*-131 was associated with chicken (meat and pets) and poor food handling, and *flaA*-2 was associated with consumption of undercooked chicken, ham, exposure to puppies and poor food-handling practices (Table 2). When all ‘other’ *flaA* genotypes combined (comprising 56 *flaA* genotypes) were compared to controls, disease among this group was associated with undercooked chicken, offal, and bottled water (Table 2). Of the seven *flaA*-10 case-patients that consumed offal, three specified lamb, one chicken and the remaining three cases did not indicate a species. Of the seven *flaA*-6b case-patients that consumed offal, two specified lamb, the remaining gave no details on species. Among *flaA*-6b and -10 cases that ate offal, there was no geographical or time clustering. While the odds ratios for some of the risk factors implicated may be of borderline statistical significance, some (e.g. offal and contact with pet chickens and farm birds) were unusually high. In the final multivariate models for each of the major *flaA* genotypes (ducks, geese, etc.) were unusually high. In the significance, some (e.g. offal and contact with pet factors implicated may be of borderline statistical comparison group in their exposure to various exposures associated with *flaA*-6b and *flaA*-131, and reflected exposures associated with disease due to those *flaA* genotypes in the final models (Table 2). *flaA*-2 infected cases were significantly more likely to consume ham (Tables 2 and 3). Poultry meat exposures did not differ between *flaA* types in multinomial regression analyses (Table 3), but were significantly associated with disease due to *flaA*-2, -6, -6b and -131 genotypes when compared to healthy controls (Table 2).

**DISCUSSION**

Case-control studies of campylobacteriosis have consistently identified chicken consumption as the most commonly associated risk factor [7, 9, 11, 13, 35]. Similarly we found chicken associated with disease due to four out of five major *flaA* genotypes, suggesting that chicken harbour a range of *C. jejuni* genotypes. Interventions aimed at minimizing chicken contamination are needed to reduce the burden of disease, and have been initiated in Iceland and Denmark [36, 37]. Here we have attempted to determine whether separate analysis of *C. jejuni* genotypes for risk factors may provide insights into further sources of this important disease.

We demonstrated the benefit of using molecular methods to more specifically define cases of campylobacteriosis to study possible risk factors for infection. We found that illness due to *C. jejuni* genotype *flaA*-10 was independently associated with undercooked beef consumption. In both case-control and case-only comparisons undercooked beef was associated with *flaA*-10 disease (Tables 2 and 3). Non-poultry meat has not previously been identified as a risk factor for campylobacteriosis in Australia [13] ([L. E. Unicomb et al., unpublished results]). Consumption of raw milk and/or contact with calves have been implicated in four Australian outbreaks [38] (OzFoodNet Outbreak Register, M. Kirk, personal communication, July 2006) and exposure to non-poultry meats and bovine husbandry have been associated with *Campylobacter* illness in case-control studies conducted in other countries [5, 8, 9, 11]. By way of comparison, the case-control study from which subjects in this study were drawn had 881 cases and 833 controls aged >5 years. It found undercooked chicken, offal, ownership of domestic chickens aged <6 months, and domestic dogs aged <6 months as risk factors [13].

Disease caused by *flaA*-2 was associated with exposure to ham in both case-control and case-only comparisons (Tables 2 and 3). This finding was unexpected, since pigs are predominantly (but not exclusively) infected with *C. coli* [39]. Previously, a ham-containing salad has been implicated in a *C. coli* outbreak [40] and it has been detected as a risk factor in a case-control study conducted in the United States [9]. It is unclear how processed meats such as ham could be the source of disease; contamination at retail outlets from raw meats may occur. Alternatively, this finding may reflect cross-contamination...
Table 3. *Multinomial logistic regression analysis* for the major Campylobacter jejuni flaA genotypes compared to ‘other’ flaA genotypes

<table>
<thead>
<tr>
<th>Exposure</th>
<th>flaA-2 (n=43)</th>
<th>flaA-6 (n=70)</th>
<th>flaA-6b (n=129)</th>
<th>flaA-10 (n=48)</th>
<th>flaA-131 (n=35)</th>
<th>Other† (n=265)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Poultry meat</td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chicken</td>
<td>90% 1.5(0.5–4.5)</td>
<td>93% 2.4(0.8–7.2)</td>
<td>82% 0.8(0.4–1.4)</td>
<td>85% 1.0(0.4–2.5)</td>
<td>85% 0.9(0.3–2.8)</td>
<td>85%</td>
</tr>
<tr>
<td>Undercooked</td>
<td>12% 1.0(0.4–2.8)</td>
<td>6% 0.5(0.2–1.5)</td>
<td>6% 0.5(0.2–1.2)</td>
<td>5% 0.4(0.1–1.5)</td>
<td>9% 0.8(0.2–2.9)</td>
<td>12%</td>
</tr>
<tr>
<td>Barbecued</td>
<td>38% 0.6(0.3–1.3)</td>
<td>37% 0.6(0.3–1.0)</td>
<td>47% 0.9(0.6–1.4)</td>
<td>46% 1.0(0.5–1.9)</td>
<td>51% 0.9(0.4–1.8)</td>
<td>49%</td>
</tr>
<tr>
<td>Non-poultry meat</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Undercooked beef</td>
<td>17% 1.1(0.5–2.7)</td>
<td>14% 0.9(0.4–2.1)</td>
<td>30% 1.1(0.6–2.0)</td>
<td>21% 2.5(1.2–5.2)</td>
<td>16% 1.7(0.7–4.3)</td>
<td>15%</td>
</tr>
<tr>
<td>Diced beef</td>
<td>21% 1.7(0.7–4.2)</td>
<td>7% 0.5(0.2–1.5)</td>
<td>22% 1.8(1.0–3.4)</td>
<td>11% 0.8(0.3–2.5)</td>
<td>26% 2.2(0.8–5.7)</td>
<td>13%</td>
</tr>
<tr>
<td>Any sausage</td>
<td>34% 0.3(0.1–0.6)</td>
<td>14% 0.3(0.1–0.6)</td>
<td>13% 0.3(0.1–0.6)</td>
<td>45% 1.4(0.5–3.0)</td>
<td>35% 1.4(0.5–3.0)</td>
<td>35%</td>
</tr>
<tr>
<td>Veal</td>
<td>5% 0.7(0.2–3.3)</td>
<td>12% 1.8(0.7–4.5)</td>
<td>4% 0.5(0.2–1.5)</td>
<td>11% 1.7(0.5–5.3)</td>
<td>4% 0.5(0.1–3.8)</td>
<td>7%</td>
</tr>
<tr>
<td>Offal</td>
<td>0% —</td>
<td>3% 0.6(0.1–2.9)</td>
<td>5% 1.2(0.5–3.2)</td>
<td>16% 3.9(1.4–10.8)</td>
<td>0% —</td>
<td>5%</td>
</tr>
<tr>
<td>Paté</td>
<td>5% 1.0(0.2–5.3)</td>
<td>6% 1.4(0.4–4.6)</td>
<td>9% 2.3(1.1–5.5)</td>
<td>13% 3.4(1.2–9.8)</td>
<td>0% —</td>
<td>4%</td>
</tr>
<tr>
<td>Ham</td>
<td>73% 2.6(1.3–5.5)</td>
<td>5% 1.0(0.6–1.7)</td>
<td>60% 1.5(0.8–2.9)</td>
<td>45% 0.8(0.4–1.6)</td>
<td>51% 0.8(0.4–1.6)</td>
<td>51%</td>
</tr>
<tr>
<td>Other food</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Organic fruit/vegetables</td>
<td>10% 1.6(0.5–5.0)</td>
<td>7% 1.1(0.4–3.3)</td>
<td>6% 0.8(0.3–2.1)</td>
<td>10% 1.5(0.5–4.7)</td>
<td>6% 0.9(0.2–4.2)</td>
<td>7%</td>
</tr>
<tr>
<td>Environmental</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bottled water</td>
<td>12% 1.2(0.4–3.3)</td>
<td>11% 1.1(0.5–2.6)</td>
<td>6% 0.6(0.2–1.3)</td>
<td>16% 1.5(0.6–3.7)</td>
<td>9% 0.9(0.3–3.2)</td>
<td>6%</td>
</tr>
<tr>
<td>Puppy</td>
<td>8% 0.9(0.2–2.4)</td>
<td>5% 0.5(0.2–1.9)</td>
<td>4% 0.4(0.2–1.2)</td>
<td>7% 0.7(0.2–2.7)</td>
<td>3% 0.3(0.0–2.3)</td>
<td>8%</td>
</tr>
<tr>
<td>Pet chicken</td>
<td>7% 0.7(0.2–2.4)</td>
<td>7% 0.7(0.2–2.2)</td>
<td>11% 1.1(0.6–2.2)</td>
<td>9% 0.9(0.3–2.7)</td>
<td>20% 1.9(0.8–4.9)</td>
<td>10%</td>
</tr>
<tr>
<td>Pet chicken &lt;6 months</td>
<td>0% —</td>
<td>4% 0.5(0.3–7.3)</td>
<td>2% 0.7(0.2–2.8)</td>
<td>4% 1.5(0.3–7.3)</td>
<td>11% 3.1(0.8–11.2)</td>
<td>3%</td>
</tr>
<tr>
<td>Farm birds</td>
<td>2% 1.6(0.2–4.8)</td>
<td>4% 2.9(0.6–13.2)</td>
<td>1% 0.5(0.1–4.6)</td>
<td>2% 1.5(0.2–13.4)</td>
<td>0% —</td>
<td>2%</td>
</tr>
<tr>
<td>Poor barbeque technique</td>
<td>9% 2.0(0.6–6.5)</td>
<td>1% 0.3(0.0–2.2)</td>
<td>3% 0.6(0.2–2.0)</td>
<td>6% 1.3(0.4–4.8)</td>
<td>9% 2.2(0.6–8.5)</td>
<td>5%</td>
</tr>
</tbody>
</table>

* Multinomial models controlled for age and sex.
† All cases with flaA genotypes apart from the five major flaA genotypes.
‡ Relative risk ratio (RRR) with 95% confidence intervals (CI) given in parentheses. RRR and 95% CIs given in bold indicate that a statistically significant difference was found for the respective flaA genotype when compared to ‘other’ flaA genotypes.
of ham and other foods during preparation in the home.

Consumption of offal (from a variety of species) was associated with disease due to flaA-6b, -10 and ‘other’ genotypes, and was also reported for all Campylobacter species and genotypes in an Australian case-control study [13]. Furthermore, duck liver consumption has been implicated in one Australian case-control study [13]. Furthermore, duck liver consumption has been implicated in one Australian case-control study [13]. Furthermore, duck liver consumption has been implicated in one Australian case-control study [13].

Gel patterns for flaA-6 and flaA-6b genotypes differed by the size of one band (Fig.), probably from an insertion or deletion in the flaA gene. flaA-6 was detected in a previous study from January 1999 to July 2001 in New South Wales [29, 30] and has been detected in a subsequent study in South Australia from November 2005 to March 2006 (B. Coombs, personal communication, December 2006). Geographic and temporal differences in flaA-6 and -6b distribution suggest that one may be a variant of the other. When flaA-6 and -6b were analysed as a single genotype (controlling for the potential confounders age and sex) similar variables were significantly associated with disease, in univariate models, as those for the more common flaA-6b alone (data not shown). The two genotypes were analysed separately as we could not assume that they are variants without further laboratory investigations.

Clinical manifestations differed slightly among those infected with the major flaA genotypes with flaA-2 infections apparently more severe than those due to other genotypes; this suggests that there may be differences in virulence between flaA genotypes. While the differences between flaA-2 study cases and others were small, they were consistent for symptoms that resulted in missing school, work, recreation or other activities. We have previously reported that flaA genotypes closely predicted MLST; 88% of flaA-2 isolates, when tested by MLST, belonged to clonal complex (CC) 48 [30]. Among the CC 48 human isolates on the international MLST database (http://pubmlst.org/campylobacter; accessed 11 July, 2007), those included had caused Guillain–Barré syndrome, Miller–Fisher syndrome and systemic disease in addition to gastroenteritis. Genotype-specific differences in symptoms by age could not be explored in this study as numbers in each age group were small.

Our findings should be considered in the light of study limitations. Selection bias in recruitment of controls is possible, since people who spend more time at home would be more easily contactable, but several factors were controlled for in analyses. Measurement biases may have occurred as we relied on information from participants that was not validated. However, this applied to both cases and controls. Interviewer bias may have occurred as interviewers knew which interviewees were cases and controls; and recall bias was possible since cases potentially would have better recall than controls. Study cases were selected from notified cases, which include those with more severe disease. Study cases for whom isolates were not genotyped, had similar characteristics to those that were genotyped, a greater number of the former were educated to school level only; this small difference may have had an impact on exposures. We conducted many hypothesis tests, so it is plausible that some statistically significant differences may have arisen by chance; of the 66 exposure variables examined for each subtype, an average of three are expected due to chance alone (using the 5% level of significance). Our findings therefore need to be replicated by further studies to confirm our results.

Detecting genotype-specific risk factors may be better determined by enrolling cases based on their isolate subtype results, focusing on the more common subtypes, thereby including sufficient sample size to test hypotheses. This would further be enhanced by studies of the distribution of flaA genotypes of non-human C. jejuni isolates; such data on Australian non-human isolates are not available. Information on genotypes for other countries is available from the international MLST database. As stated above, flaA genotypes can closely predict MLST CCs; 96% of Australian flaA-6 isolates belong to CC 257, 91% of flaA-10 to CC 354 and 88% of flaA-2 to CC 48 [30]. Among the data from various countries on the MLST database, these clonal complexes have been detected in a variety of non-human samples and countries: CC 257 (flaA-6) was detected from bovine, avian (poultry and other avian), ovine and porcine samples, CC 354 (flaA-10) from bovine, poultry (including environmental) and ovine samples and CC 48 (flaA-2) from bovine, avian (poultry, poultry environment and other avian), ovine, water, sand and domestic pet samples. While these data could potentially be useful in supporting findings from the case-control and case-case analyses, only one of the 124 non-human isolates from the CC 257, 354 and 48 isolates on the database was from Australia. Determining flaA genotypes from systematically collected, non-human sources in Australia may be useful to identify potential reservoirs of genotypes.
Campylobacter jejuni flaA genotype risk factors

flaA genotyping is a gel-based method which has limitations; it requires standardization to achieve comparable results across laboratories but is rapid and cheap. Ideally, this method would be automated to achieve ‘high throughput’ status. There are reports of instability in the flaA locus [41] but the majority of C. jejuni isolates are apparently genetically stable in this region over time [26, 42, 43] and that instability may be strain specific. Sequence-based methods such as MLST remove dependence on visual and potentially subjective, genotype assignment but require expensive equipment to handle medium to high numbers of isolates and reagents are costly [29, 44]. Isolates from this study were genotyped retrospectively, but ‘real time’, genotyping of notified case isolates would be preferable, to enable rapid detection of temporal clusters using a library of common flaA genotypes and should be feasible.

This is the first study to suggest the value of flaA genotyping for identifying strain-specific risk factors for Campylobacter. Case-control analyses using logistic regression and case-only analyses using multinomial regression were employed to detect risk factors for C. jejuni flaA genotypes among a selection of notified cases drawn from five Australian states over a 12-month period. Differences were detected for symptom profile, geographic distribution and exposures between flaA genotypes. The value of flaA genotyping is therefore worthy of further investigations in studies with a larger sample size and in other settings, and particularly in the course of outbreak investigations. The ability of flaA genotyping to detect clusters and outbreaks among apparently sporadic notified cases will be assessed in a separate report (L. E. Unicomb et al., unpublished observations).

APPENDIX. Australian Campylobacter Subtyping Study Group (listed in alphabetical order):

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products and food animals. *Epidemiology and Infection* 2006; 134: 758–767.


