Experimental *Campylobacter jejuni* infection in the chicken: an animal model of axonal Guillain-Barré syndrome

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Abstract

**Objective**—To develop and characterise an animal model of paralytic neuropathy after *Campylobacter jejuni* infection. *Campylobacter* infection precedes development of many cases of Guillain-Barré syndrome and is particularly associated with cases having prominent axonal degeneration. Understanding the pathogenesis of Guillain-Barré syndrome after *C jejuni* infection has been slowed by the lack of animal models.

**Methods**—A spontaneous paralytic neuropathy is described that developed in chickens from the farms of four patients with Guillain-Barré syndrome. The production of paralytic neuropathy in chickens experimentally fed *Campylobacter jejuni* isolated from one of these patients is reported. The sciatic nerves of the spontaneously paralysed chickens were examined pathologically in teased fibres, in plastic embedded sections, and by electron microscopy. Two large groups of chickens were then fed cultures of a *C jejuni* (Penner type O:19) isolated from one of these patients.

**Results**—The chickens with spontaneous paralysis had pathologically non-inflammatory neuropathy. Pathology in the sciatic nerves ranged from no detectable changes to severe Wallerian-like degeneration. In the experimentally inoculated groups, an average of 33% of the chickens became paralysed. The median time after inoculation to paralysis was 12 days. The lesions found in the first few days of paralysis included nodal lengthening and paranodal demyelination. In those animals that survived for several days after onset of weakness, the pathology was dominated by extensive Wallerian-like degeneration. Animals that survived for weeks with no clinically apparent neuropathy had paranodal remyelination in some teased nerve fibres, reflecting earlier paranodal demyelination.

**Conclusion**—Experimental inoculation with *C jejuni* may provide a new model for understanding some forms of Guillain-Barré syndrome.

*Campylobacter jejuni* infection is increasingly recognised as an important antecedent of the Guillain-Barré syndromes. 

Guillain-Barré syndrome as clinically diagnosed includes patients with inflammatory demyelinating changes and patients dominated pathologically by Wallerian-like axonal degeneration. Some patients with demyelinating Guillain-Barré syndrome have antecedent *C jejuni* infection, but *C jejuni* seems to be particularly associated with “axonal” forms of Guillain-Barré syndrome.

This association applies to the nearly purely motor syndrome termed acute motor axonal neuropathy (AMAN) that was recognised and characterised in studies of the summer epidemics that occur in northern China.

An important constraint on pathogenetic studies has been the absence of an animal model of neuropathy induced by *C jejuni*. This report describes the occurrence of paralytic neuropathy in chickens in two settings. The first was a spontaneous disorder identified in the flocks of some farm families in which a member had recently developed Guillain-Barré syndrome. The second was produced experimentally by feeding *C jejuni* isolated from a patient with AMAN. These results may provide new models for the dissection of pathogenetic mechanisms in Guillain-Barré syndrome associated with *C jejuni* infection.

**Methods**

**SPONTANEOUS-DISEASE GROUP**

Patients admitted to the Second Teaching Hospital with Guillain-Barré syndrome are routinely questioned about similar disease in their villages or about sick animals on their farms. In the summer of 1993, the families of four patients gave a history of weak chickens developing near the time of onset of paralysis in the patient. Two of the authors travelled to the farms, and examined and purchased the chickens. Five chickens with spontaneously occurring disease were purchased from the farms of the four families, each of which had one member with Guillain-Barré syndrome. In all of the chickens, both the legs and the wings
were involved. The sciatic nerves from these chickens were removed at necropsy and examined as described for the experimental groups below.

EXPERIMENTAL GROUPS
All birds in the experimental groups were inbred Esparpuk chickens, a local white feathered strain that resembles the Leghorn (AK Asbury, personal communication). All of the experimental birds were less than six months old. Half were males and half females. They were housed 15 to a cage; the control birds were caged in a separate region of the animal house. They were routinely raised by a local farm supplier, and were not from specific pathogen free, Marek's disease free, or avian leukosis free flocks. These birds were not cultured for *C. jejuni* before beginning the experimental studies, but subsequent studies have shown that *C. jejuni* can be cultured normally from nearly all such chickens.

Control group 1 consisted of four chickens that did not receive *C. jejuni*. Experimental group 1 consisted of 33 chickens that were fed *C. jejuni* by gavage, presumably into the gizzard, as described below. Experimental group 2 duplicated group 1 in a larger study of 66 chickens, all fed *C. jejuni* by the protocol described below.

STOOL CULTURE AND PREPARATION OF *C. JEJUNI* INOCULUM

The *C. jejuni* isolate used in all experiments was obtained from an eight year old boy (HB93–13) with AMAN (patient 4 in results). His stool was cultured by an enrichment method. Fresh stool was cultured in Preston Enriched Broth (Oxoid) in a microaerophilic (5% *O*₂, 3–11% *CO*₂) environment at 41°C. After 24 hours, the culture was inoculated on the selective CCDA plates and incubated under microaerophilic conditions at 15°C for 48 hours. The *C. jejuni* strain was typed as Penner O:19 by the USPHS Centers for Disease Control (Atlanta, GA, USA).

The *C. jejuni* was stored at –70°C in brain-heart infusion broth with 15% glycerol. The inocula were prepared by thawing the stored *C. jejuni* to room temperature. After immediate transfer to a CCDA plate, the sample was incubated in a microaerophilic environment for 48 hours at 37°C. The bacteria were flushed with enrichment broth to a standard turbidity. The chickens in the experimental groups fed *C. jejuni* each received 5 ml of the inoculum by gastric gavage. The gavage tube was of sufficient length to reach the gizzard.

### CLINICAL AND PATHOLOGICAL ANALYSIS

Animals were assessed regularly for weakness of the hind limbs, difficulty standing or walking, wing droop, head drop, and difficulty feeding. In many animals, once weakness began it progressed very rapidly (see results) and they died on the day of onset. When possible, weak animals were allowed to survive for several days before being killed by exsanguination. The sciatic nerves of all of the control chickens and the affected experimental chickens, as well as some clinically unaffected experimental chickens, were removed. No other pathological specimens were obtained. Specimens were processed in three ways: (1) samples were fixed with 4% paraformaldehyde, stained with 4% osmium, softened in graded concentrations of glycerol, stored in 100% glycerol, and examined after teasing; (2) other paraformaldehyde fixed samples were embedded in paraffin for haematoxylin and eosin staining and for combined silver (modified Naumienko-Feigin stain)-luxol fast blue staining; and (3) samples were fixed by 5% glutaraldehyde, postfixed in osmium, embedded in Epon, and sectioned at 1 μm. Thin sections of selected glutaraldehyde fixed samples were examined by electron microscopy.

### Results

#### SPONTANEOUS DISEASE GROUP

The table summarises the histories of the four patients whose families had weak chickens. Three of the four had the AMAN pattern of Guillain–Barré syndrome. All of the chickens that were examined were weak. All had *C. jejuni* cultured from their stools. One was typed (chicken No 3, table) and found to be Penner 019 Lior H36.

The pathological examination of these chickens was restricted to the sciatic nerves. The most severely affected was chicken No 3, obtained from the family of patient HB–93–4, an 11 year old girl with pronounced slowing of her nerve conduction velocities and with a clinical diagnosis of AIDP. This chicken had mild mononuclear infiltration in the sciatic nerve, with occasional fibres undergoing demyelination and numerous fibres undergoing Wallerian-like degeneration. The sciatic nerve from the other weak chicken from this farm was histologically normal.

Chicken No 1 from the farm of a 47 year old woman with AMAN had several teased fibres with definite lengthening of the nodes of Ranvier and mild paranodal demyelination with occasional remyelination (fig 1A and B).

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**Table: Summaries of patients with paralysed chickens**

<table>
<thead>
<tr>
<th>Designation</th>
<th>Age (y)</th>
<th>Antecedent diarrhoea</th>
<th><em>C. jejuni</em> type</th>
<th>Electro-diagnostic designation*</th>
<th>Paralysed chickens examined</th>
<th>Sciatic nerve pathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>HB93-2</td>
<td>47½</td>
<td>–</td>
<td>–</td>
<td>AMAN</td>
<td>1 (No 1)</td>
<td>Nodal lengthening, paranodal demyelination</td>
</tr>
<tr>
<td>HB93-4</td>
<td>11½</td>
<td>–</td>
<td>–</td>
<td>AIDP</td>
<td>2 (No 2, No 3)</td>
<td>No 3 had mild mononuclear inflammation, mild internodal demyelination, and extensive Wallerian-like degeneration</td>
</tr>
<tr>
<td>HB93-9</td>
<td>6½</td>
<td>–</td>
<td>–</td>
<td>AMAN</td>
<td>1 (No 4)</td>
<td>Normal</td>
</tr>
<tr>
<td>HB93-13</td>
<td>8½</td>
<td>–</td>
<td>+ (Penner O:19)</td>
<td>AMAN</td>
<td>1 (No 5)</td>
<td>Normal</td>
</tr>
</tbody>
</table>

*Criteria as described in Ho et al.*
The weak chickens No 4 and 5 from the farms of children with the AMAN pattern had normal sciatic nerves.

EXPERIMENTAL GROUPS
The control chickens (those that were not fed C jejuni) developed neither diarrhoea nor paralysis. They were killed on day 46, and sciatic nerves were normal both by teased fibre preparations and by silver-LFB and haematoxylin and eosin stained paraffin sections. In particular, there was no paranodal demyelination or remyelination (fig 1A) and no evidence of Wallerian-like degeneration. Occasional mild paranodal changes were interpreted as preparative artifacts.

Group 1
The 33 chickens from group 1 all developed diarrhoea, beginning two to four days after feeding and lasting three to 14 days. C jejuni was cultured from their stools. Penner typing of these cultures was not done.

Eighteen of these 33 (55%) chickens became weak. Affected chickens were unable to stand, and some were paralysed. At early stages the chickens developed wing droop and head drop. Figure 2 shows cumulative plots of the times of onset of paralysis. The earliest onset of weakness (one chicken) was day 5 after inoculation. The median time to development of weakness was 12 days. Eleven of the 18 weak animals in group 1 were killed or died within a day of onset of weakness. The other seven were allowed to survive for up to 11 days after the onset.

Pathologically, five of the sciatic nerves from these 18 weak chickens were abnormal by examination of teased fibres in the mid-sciatic nerve. Three of these five were from animals that survived for 10 or 11 days after onset of weakness. The most severe abnormality was extensive ongoing Wallerian-like degeneration affecting about 25% of total fibres (figs 1D, 3A–D) in a bird in which weakness became evident 18 days after giving C jejuni. This bird died on day 19. The affected chickens also showed varying degrees of lengthening of the nodes of Ranvier and, in rare instances, paranodal demyelination (fig 1E). Mononuclear cells overlying these widened paranodes could occasionally be identified (fig 1E). Several fibres were identified by electron microscopy in which macrophages were present within the periaxial space of affected internodes, separating the inner myelin lamellae from the axon, which was often degenerating (fig 3D).

The 15 clinically unaffected chickens from group 1 were all killed on day 46, and their sciatic nerves were examined by teased fibre studies. Only one nerve from these animals showed a few fibres undergoing late stages of Wallerian-like degeneration. Five of these 15 nerves had paranodal remyelination, with short intercalated internodes on one or both sides of the node. The proportion of fibres affected was never greater than 3%–4%.

Figure 1  Osmicated teased nerve fibres. Panels A, B, and E photographed with Nomarski optics. (A) Normal node of Ranvier (arrows) from a control chicken; original magnification × 1100. (B) Paranodal demyelination and remyelination (chicken No 1) from the spontaneous disease group. Thin arrows identify the thin (remyelinated) myelin sheath in the former paranode. Thick black arrows identify the nodes; original magnification × 1100. (C) Fibre undergoing Wallerian-like degeneration from the same chicken; original magnification × 240. (D) Teased nerve fibres undergoing Wallerian-like degeneration from the most severely affected chicken in experimental group 1; original magnification × 650. (E) Paranodal demyelination in a chicken from experimental group 1. Demyelinated axon is identified by asterisks. Two mononuclear cells, presumably macrophages, are identified by arrows; original magnification × 1100.

Figure 2  Increasing proportion of paralysed chickens with time in groups 1 and 2. The earliest animal (group 2) became weak on day 5. The median time was day 12.
Group 2
Of the 66 chickens in group 2, 15 (23%) developed paralysis. The earliest onset of weakness was on day 5 in two animals. The others became weak on days 9–26 (fig 2). Five of these 15 animals had sciatic nerve pathology by teased fibre preparations. Wallerian-like degeneration was found in two animals allowed to survive for several days after developing weakness. One of these chickens became weak on day 5 and was killed on day 12 (at a time when the animal was clinically improving). The other severely affected chicken became weak on day 17 and was killed on day 58. In addition to fibre loss and occasional fibres undergoing late stages of Wallerian-like degeneration, this chicken had extensive paranodal remyelination.

As in group 1, most of the animals in group 2 that were killed on the day their weakness developed had normal or nearly normal sciatic nerves. The 41 animals that seemed clinically healthy were not examined pathologically.

In neither group were the mononuclear cell invasion or internodal demyelination characteristic of Marek’s disease identified.

Discussion
This project began with the clinical discovery of paralysis in the chicken flocks of four families each of which had one member with Guillain-Barré syndrome. The temporal association was striking: these families knew of no previous paralytic disease in their chicken flocks, nor of those within their villages. In addition,
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as discussed later, there were pathological similarities in two of these chickens with the pathology identified in necropsies of patients dying with the AMAN pattern of Guillain-Barré syndrome. That C jejuni might be the link between the human and chicken diseases was suggested by the culture of C jejuni from the stools of one of the patients and by the fact that chickens are colonised by and excrete C jejuni normally. We have confirmed that this applies to the flocks in Hebei Province by isolating C jejuni from the culture of chicken faeces from provincial villages in which neither Guillain-Barré syndrome nor weak chickens were found (CY Li, TW Ho, R Liu, and I Nachamkin, unpublished data). Thus in these farm families, the spread of neuritogenic strains of C jejuni from chickens to humans is a real possibility. However, because C jejuni was successfully isolated from only one of these patients, it is not possible to compare the strains obtained from the chickens and patients. In the absence of such data, the simultaneous occurrence of human and spontaneous chicken disease can only provide a tantalising link. The experimental inoculation of chickens with the one successful stool isolation was undertaken to explore that possible link more rigorously.

The pathology in the most severely affected animal with spontaneous disease was dominated by Wallerian-like degeneration of sciatic nerve fibres. In addition, this animal and one other had evidence of nodal lengthening and paranodal demyelination. Strikingly, in three of the chickens there were no identifiable abnormalities in the sciatic nerves. Similarly, very minimal pathology has been noted in some patients dying with the AMAN pattern of Guillain-Barré syndrome, as well as in many of the chickens fed C jejuni. The possible basis for this apparently anomalous finding is discussed later.

EXPERIMENTAL FEEDING OF C. JEEJUNI
The spontaneous disease led us to feed C. jejuni isolated from a patient with AMAN (patient 4) to groups of experimental chickens. The two major findings in this experiment were, firstly, that most chickens fed C. jejuni by this protocol developed diarrhoea; and, secondly, that a proportion of these chickens developed paralytic neuropathy. Considering the two experimental groups together 33% of the animals developed weakness. Pathological studies confirmed a neuropathic basis for the weakness and indicated that some apparently healthy animals also had subclinical neuropathy, but also pointed out that very minimal or no changes could be present in the early stages of paralysis.

Several aspects of the experimental design will need to be altered in future studies. The strains of C jejuni that the chickens were almost certainly excreting before inoculation should be determined. Because both the broth and the organisms contributed to the turbidity of the inoculum, we could not determine the number of organisms inoculated; determination of the size of the inoculum will need to be done by culture. Inoculation of a wider range of C jejuni isolates is necessary. Finally, more extensive sampling of the PNS and CNS should be planned, including spinal roots and motor nerve terminals.

PATHOLOGICAL CHANGES
The pathological changes in paralysed chickens ranged from extensive Wallerian-like degeneration to no detectible abnormalities. Overall, the changes in these experimental birds had similarities to those seen in the AMAN pattern of Guillain-Barré syndrome in humans. As in AMAN, lymphocytic infiltration was uncommon and scant (in the present study, the only chicken with identifiable lymphocytic infiltration of the sciatic nerve was No 3 in the spontaneous disease group). In AMAN, the mildest and probably the earliest change is nodal lengthening, which can occur with or without evidence of associated paranodal myelin damage. Lengthening and paranodal demyelination were seen in some of the experimentally fed animals killed early in the course of the disease, as well as in chickens No 1 and No 3 with spontaneous disease. In addition, among the experimental animals fed C jejuni that did not develop clinical neuropathy and that were allowed to survive for nearly two months, some had evidence of paranodal remyelination. This change, never seen in control animals, presumably reflects remyelination in segments affected by earlier paranodal demyelination.

Wallerian-like degeneration was the dominant change in those animals that survived several days while weak. Wallerian-like degeneration is similarly the major finding in severe cases of AMAN. A provocative link between the disorders is the finding in the chickens of occasional fibres that had macrophages in the space normally occupied by the axon and inside otherwise normal appearing myelin sheaths. This is a very prominent change in the axon patterns of Guillain-Barré syndrome. As described below, in AMAN this pattern has been linked to the presence of antibody and complement activation products in the periaxonal space.

A final pathological similarity is that, as in AMAN, many animals with paralysis had no detectable changes in the sciatic nerves. In some cases of AMAN, the changes in the spinal roots and sciatic nerves can be surprisingly mild in view of the severe (and fatal) paralysis. Similarly, AMAN paralysis can be rapidly reversible, indicating that some lesion milder than Wallerian-like degeneration must be present to allow early recovery. In AMAN, the possibilities of selective motor nerve terminal degeneration (TW Ho et al, unpublished data) or of antibody induced failure of impulse conduction (M Roberts, A Vincent, J Newsom-Davis, H Willson, TW Ho, unpublished data) being considered. Similar considerations could apply to the chicken model described here. Motor nerve terminals were not examined. Alternatively, disease could be concentrated within the spinal roots,
which were also unexamined in the present material.

Finally, the present data do not indicate the mechanism by which paralysis develops. Because the axonal patterns of Guillain-Barré syndrome are presumed to be immune mediated, an allergic reaction to some constituent of the organism, possibly the lipopolysaccharide, is attractive. However, the present data do not exclude the possibility of a neurotoxin produced by the organism. We have recently produced paralytic disease in one of eight rabbits immunised using complete Freund's adjuvant containing a lipopolysaccharide preparation from the same C. jejuni isolate used in these studies (CY Li, WQ Tian, P Xue, RC Liu, C Yang, unpublished data). The pathology was similar to that in axonal Guillain-Barré syndrome and in the present study, including the presence of periaxonal macrophages. These results support an immune mediated mechanism.

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