Characterization of Eimeria brunetti Isolated from a Poultry Farm in Japan

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ABSTRACT. None of anticoccidial vaccines (Trivalent TAM™, monovalent Neca™ and imported pentavalent Paracox®-5) contain Eimeria brunetti in Japan, which has not been regarded as a cause of coccidiosis, because of its low prevalence. However, we have recently reported the evidence of a high nationwide prevalence of this species. In this report, we describe the characteristics of E. brunetti infection was almost completely blocked. Consequently, it is suggested that the characteristics of E. brunetti are various among the strains, but the pathogenicity of the Japanese Nb strain is enough strong to cause clinical coccidiosis.

KEY WORDS: anti-drug sensitivity, Eimeria brunetti, Japan, live vaccine, pathogenicity.

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Chicken coccidiosis induced by Eimeria parasites causes huge economic losses to intensive poultry industries worldwide [12]. Seven species of Eimeria have been reported to infect chickens so far. Among these species, E. acervulina, E. brunetti, E. maxima, E. necatrix and E. tenella are highly pathogenetic, causing production losses in industry due to clinical or subclinical coccidiosis.

Generally, the poultry industry uses specific anticoccidial drugs or live anticoccidial vaccines as prophylactic therapies to prevent and control the disease. There are concerns about drug residues in poultry products and a strong consumer desire to ban drugs from animal feeds. As a result, the demand for products derived from organic chickens, without feeding of chemicals or antibiotics, tends to increase in Japan as European countries. It is estimated that live anticoccidial vaccines in Japan are applied to one hundred million chickens per year [13].

Three live coccidiosis vaccines have been registered and sold in Japan. Trivalent TAM™ (Nisseiken Co., Ltd., Tokyo, Japan) contains precociously attenuated strains of E. acervulina, E. maxima and E. tenella. Monovalent Neca™ (Nisseiken Co., Ltd.) contains a precociously attenuated strain of E. necatrix. Pentavalent Paracox®-5 (MSD Animal Health, Milton Keynes, U.K.) which contains precociously attenuated strains of E. acervulina, E. maxima, E. mitis and E. tenella has been imported from the UK. However, none of these products contain E. brunetti, because its occurrence in chicken flocks has been regarded to be less intensive in Japan than in other studies. No oocysts of E. brunetti were detected on several national surveys of broiler flocks in 1970’s [8–10]. In 1990, two strains of E. brunetti were first isolated from poultry farms in Hokkaido and Kumamoto. This suggested a greater prevalence of E. brunetti, as these are the northern-most and southern-most parts of Japan, respectively [7]. No further surveys or cases have been reported since 1991.

E. brunetti infections have been recently diagnosed in many cases from samples submitted to our laboratory from commercial chicken farms [3]. This has forced us to review our concept of E. brunetti as a species capable of causing coccidiosis in Japan also. Characteristics of the Japanese isolate of E. brunetti haven’t been evaluated still yet. For the verification, we describe a Japanese strain of E. brunetti, focusing on its pathogenicity and sensitivity to drugs in this paper.

MATERIALS AND METHODS

Parasite: The isolates of E. brunetti, as Nb strain, was derived from a single oocyst isolated from feces of breeder chickens in Miyazaki prefecture and has been maintained in our laboratory using specific-pathogen-free (SPF) layer chickens. The correct species identification and purity of the strain were confirmed by PCR employing an assay directed towards the ITS-1 [3].

Experimental design: E. brunetti infection was induced in a group of ten 35-day-old SPF chickens by oral inoculation with 1 × 102, 1 × 103, 1 × 104 or 1 × 105 sporulated oocysts/bird. The birds were raised in wire-floored cages for 7 days and then necropsied. Experimental infections were evaluated...
using growth ratio, mortality and intestinal lesion scores. The growth ratio was obtained individually with the following formula: (body weight at termination − body weight at initiation)/ body weight at initiation × 100. Intestinal lesion scores were graded according to Johnson’s method [2]. Values with superscripts indicate significant difference (P<0.05) from uninfected control group.

### Table 1. The pathogenicity of the Japanese Nb strain of *E. brunetti*

| Number of oocysts inoculated per bird | Mortality rate | Mean growth ratio<sup>a)</sup> (relative growth ratio) | Mean lesion score<sup>b)</sup> |
|--------------------------------------|----------------|----------------------------------------------------------|------------------------------|
|                                      |                | Jejunum, ileum                                          | Rectum                       |
| 1 × 10²                              | 0/10           | 38.7 ± 8.3 (0.89)                                       | 1.7<sup>c)</sup>              |
| 1 × 10³                              | 0/10           | 22.2 ± 8.7<sup>c)</sup> (0.51)                           | 1.3<sup>c)</sup>              |
| 1 × 10⁴                              | 0/10           | 9.3 ± 8.2<sup>c)</sup> (0.21)                            | 1.3<sup>c)</sup>              |
| 1 × 10⁵                              | 3/10           | −4.2 ± 4.9<sup>c)</sup> (−0.1)                           | 1.9<sup>c)</sup>              |
| None                                 | 0/10           | 43.7 ± 7.4 (1.0)                                        | 0                            |

<sup>a)</sup> (body weight at termination − body weight at initiation)/ body weight at initiation × 100. <sup>b)</sup> Johnson and Reid [2]. <sup>c)</sup> Values with superscripts indicate significant difference (P<0.05) from uninfected control group.

### Table 2. Oocyst output of the Japanese Nb strain of *E. brunetti*

| Number of oocysts inoculated per bird | Days after inoculation |
|--------------------------------------|------------------------|
|                                      | 4         | 5         | 6         | 7         | 8         | 9         | 10        | 11        |
| 1 × 10²                              | -         | 4.7<sup>a)</sup> | 7.4       | 7.4       | 7.2       | 6.1       | 5.3       | -         |
| 1 × 10³                              | -         | -         | 7.6       | 7.6       | 7.2       | 5.1       | -         | -         |
| 1 × 10⁴                              | -         | 4.2       | 7.7       | 8.1       | 7.2       | 6.1       | 5.8       | -         |
| 1 × 10⁵                              | -         | 4.1       | 7.2       | 7.7       | 7.3       | 5.3       | -         | -         |
| None                                 | -         | -         | -         | -         | -         | -         | -         | -         |

<sup>a)</sup> Oocysts/g feces (OPG) is shown by log. Detection limit is log2.0.

### Table 3. Developing stages of the Japanese Nb strain of *E. brunetti* present in intestinal tissue

| Day after inoculation | Duodenum | Jejunum | Ileum | Cecum | Rectum |
|-----------------------|----------|---------|-------|-------|--------|
| 1                     | 0        | 0.5 (F) | 0     | 0     | 0      |
| 2                     | 0        | 1 (F)   | 0     | 0     | 0      |
| 3                     | 0.5 (F)  | 1.5 (F) | 0     | 0     | 0      |
| 4                     | 0.5 (F>S) | 2 (S)   | 2 (S) | 1.5 (S) | 1 (S) |
| 5                     | 2 (S)    | 2 (S>G) | 2 (S>G) | 2 (S>G) | 2 (S>G) |
| 6                     | 0.5 (G)  | 1 (G>S) | 1 (G>S) | 1.5 (G) | 1.5 (G) |
| 7                     | 0.5 (G)  | 1 (G)   | 1.5 (G) | 1.5 (G) | 1 (G)  |
| 8                     | 0        | 0       | 0.5 (G) | 1 (G)   | 0.5 (G) |

Two birds were sampled daily after inoculation with 1 × 10⁵ oocysts per bird. <sup>a)</sup> 0=no organisms; 1-less than 10% of host cells are infected; 2=more than 10% of host cells are infected. <sup>b)</sup>F=first generation schizonts; S=other generation schizonts or immature gametocytes; G=mature gametocytes.

**Evaluation of drug resistance:** To evaluate the drug sensitivity, a group of ten 14-day-old SPF chickens were orally inoculated with 1 × 10⁵ sporulated oocysts/bird, kept in wire-floored cages for 7 days and then necropsied. Four medicated infected, one unmedicated infected and one unmedicated uninfected (control) groups were included in this study. As sulfa anticoccidial drugs, diaveridine (19.2 ppm)/sulfaquinoxaline (76.8 ppm) obtained from Sumika Enviro-Science Co., Ltd. (Nishinomiya, Japan) and ormetoprim (75 ppm)/sulfamonomethoxine (225 ppm) obtained from Meiji Seika Pharma Co., Ltd. (Tokyo, Japan) were used in the drinking water on days 0 to 3 after infection. As ionophorous polyether anticoccidial drugs, salinomycin (50 ppm) obtained from Nichiku Yakuhin Kogyo Corporation (Ayase, Japan) and lasalocid (75 ppm) obtained from Scientific Feed...
Laboratory Co., Ltd. (Tokyo, Japan) were used in the feed continuously. The sensitivity against anticoccidial drugs is represented by calculating the anticoccidial index (ACI) according to the method of Merck Sharp & Dohme Co., Ltd. [5] as follows: ACI = (relative growth ratio + survival ratio) − (lesion index + oocyst index). The strain was considered sensitive if the ACI was >161, partially resistant if the ACI was 121 to 160 and resistant if the ACI was <120.

Animal care and use: The experiments presented here were carried out according to protocols pre-approved by the Animal Care and Use Committee of Nippon Institute for Biological Science (Tokyo, Japan) in accordance with Regulation of Animal Experimentation of Nippon Institute for Biological Science.

RESULTS

Experimentally infected chickens showed a great reduction in their body weight gain which correlated with the dose of oocysts (Table 1). There were 3 mortalities in the group that was infected with $1 \times 10^5$ oocysts. Despite showing the great reduction of body weight gains, the intestinal lesions in the infected chickens were rather mild. Typical lesions were found in the rectum, including slight swelling, a color change into whitish-orange and formation of a bellows-like shape with folded circular rings along the tube. In the other intestinal compartment, the only abnormality observed was a slight paling of the serosal surface. There was no correlation between the output patterns and the number of inoculated oocysts. Oocysts were seen in droppings from 5 to 10 DPI among almost groups (Table 2). It indicates that prepatent and patent periods of the strain were about 5 and 6 days, respectively.

Histological observation of the developing stages of the parasite is summarized in Table 3, showing the infection rate of host cells and stage transition of parasites on each day after inoculation. Mature (arrowheads) and immature (arrows) schizonts were observed in the mucosal cells and subepithelial zone of the disrupted villi of the jejunum at 4 days after infection (Fig. 1). Mature female gametocytes (arrowheads) and mature male gametocytes (arrow) were observed in the mucosal cells and subepithelial zone of the villi of the rectum at 6 days after infection (Fig. 2). The number of female gametocytes was much larger than that of male gametocytes. The onset of parasite development occurred in the upper portion of the small intestine, and subsequently parasites were found in the lower tract. The parasite density in the jejunum of E. brunetti-infected SPF chickens was the highest among the locations examined, although the gross lesions were mild (Table 2). The ACIs of medicated and unmedicated infected groups are presented in Table 4. Of the 4 medicated infected groups, only the ACI of the salinomycin group was in the partial resistant. Other drugs were so effective that the E. brunetti infection was almost completely blocked.

DISCUSSION

Severe lesions were often observed in the field chickens infected with E. brunetti(data not shown). However, such lesions did not develop in the intestines of SPF chickens under experimental conditions in this study. This may be due to differences between commercial field chickens and SPF chickens to E. brunetti infection in the susceptibility. Otherwise, this may associate with Clostridium perfringens infection. Because necrotic enteritis and coccidiosis often
The field veterinarians had diagnosed as was detected by PCR in considerable samples which 

E. brunetti E. necatrix infections in breeder pullet flocks have been misdiagnosed as E. brunetti infection process, especially when mixed infections are present. We criteria, leading to confusion in the species-determination species overlap in their measured values or patterns using these prepatent time. It has been well documented that several species could account for this, at least in part. Traditional classification techniques use criteria, such as oocyst size, parasitic site in the intestine, gross lesion figures and variations in the major parasitic site has been generally considered to be the lower small intestine [1, 4].

Contrary to previous reports [8–10], our recent study revealed that E. brunetti is very prevalent in Japan [3]. In this study, we showed that E. brunetti infections with a Japanese isolate cause similar mortalities and suppression of body weight gain as other tested isolates [1, 4]. So, the Japanese Nb strain has similar high pathogenicity to oversea strains. Furthermore, the involvement of E. brunetti in field coccidiosis cases must be evaluated. The drug sensitivity was also found to be normal, although our strain was slightly resistant to salinomycin. This coccidiostatic drug is generally thought to be slightly ineffective against E. brunetti [6].

The reasons for the recent increase in the diagnosis of E. brunetti infections in Japan remain unclear. Improvements of the techniques for the detection of E. brunetti in field samples based on PCR, rather than morphological diagnosis, could account for this, at least in part. Traditional classification techniques use criteria, such as oocyst size, parasitic site in the intestine, gross lesion figures and variations in prepatent time. It has been well documented that several species overlap in their measured values or patterns using these criteria, leading to confusion in the species-determination process, especially when mixed infections are present. We suspect that the clinical signs attributed to E. brunetti infections in breeder pullet flocks have been misdiagnosed as those of E. necatrix. Actually, we experienced that only E. brunetti was detected by PCR in considerable samples which the field veterinarians had diagnosed as E. necatrix infection (data not shown).

Opinions on the prevalence of E. brunetti were basically derived from information supplied from field surveys [8–10]. These were performed mainly on broiler flocks, while recent data for detection came from breeder pullet and layer chicken flocks [3]. E. brunetti was commonly thought to infect both younger and older chicken flocks [4]. Therefore, the epidemiology of E. brunetti should be evaluated using surveys from not only broiler flocks, but also breeder pullet and layer chicken flocks.

From the recent and present findings, the broad distribution and pathogenicity of E. brunetti in Japan has emerged. This indicates that an appropriate plan to confront the risk associated with outbreaks of this species is required. The fact that E. brunetti outbreaks are usually found in older chicken flocks means that using coccidiostatic drugs that are normally applied extensively to broilers would not be suitable for this species. Vaccination would be the best method to mitigate this risk, but appropriate vaccines effective against this species are not available in Japan. Therefore, development of an effective E. brunetti vaccine that is safe for use in the poultry industry in Japan is urgently required.

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### Table 4. Sensitivity of the Japanese Nb strain of *E. brunetti* against anticoccidial drugs

| Drug                        | Relative growth ratio | Survival ratio | Lesion index b) | Oocyst index c) | Antibiotic index d) (ACI) |
|-----------------------------|-----------------------|----------------|------------------|-----------------|--------------------------|
| Diaveridine/sulfamethoxine   | 94.10                 | 100            | 0                | 0               | 194                      |
| (in water)                  |                       |                |                  |                 |                          |
| Ormetoprim/sulfadimethoxine | 96.51                 | 100            | 0                | 0               | 197                      |
| (in water)                  |                       |                |                  |                 |                          |
| Salinomycin (in feed)       | 49.18                 | 100            | 1                | 5               | 143                      |
| Lasalocid (in feed)         | 96.88                 | 100            | 0                | 0               | 197                      |
| None                        | -4.3                  | 100            | 16               | 40              | 40                       |

a) Each group contains ten birds which were inoculated with 1 × 10⁵ oocysts per bird. b) Lesion index: the total amount of lesion score of ten birds based on Johnson and Reid [2]. c) Oocyst index: based on the ratio (%) of OPG to the none-drug group. 0–1%=0; 1–25%=5; 26–50%=10; 51–75%=20; and 76–100%=40. d) Antibiotic index = (relative growth ratio + survival ratio) – (lesion index + oocyst index). ACI: 200–161 = sensitive; 121–160 = partially resistant; and <120 = resistant.
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