Novel Borna Virus in Psittacine Birds with Proventricular Dilatation Disease

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Pyrosequencing of cDNA from brains of parrots with proventricular dilatation disease (PDD), an unexplained fatal inflammatory central, autonomic, and peripheral nervous system disease, showed 2 strains of a novel Borna virus. Real-time PCR confirmed virus presence in brain, proventriculus, and adrenal gland of 3 birds with PDD but not in 4 unaffected birds.

Borna disease virus (BDV) is the causative agent of Borna disease, a meningoencephalitis of horses and sheep in central Europe (1). As the prototype and only known member of the family Bornaviridae in the order Mononegavirales (nonsegmented, negative-strand RNA viruses), BDV is atypical in its nuclear localization of transcription, alternative splicing, and differential use of initiation and termination signals. Sequence analysis of isolates obtained from various species over several decades has shown remarkable sequence conservation; only 2 genotypes are known. The virus is highly neurotropic and infects the central, peripheral, and autonomic nervous systems. Although ungulates remain the best known natural host, the introduction of sensitive molecular and serologic assays enabled by subtractive cloning of the BDV genome facilitated surveys of sensitive molecular and serologic assays. Despite the observation that disease can be transferred to naive birds through inoculation with tissue homogenates or fecal material from affected birds (1), no other researches have described particles in tissue consistent in appearance with adenoviruses or paramyxoviruses. Whether any of these agents can be implicated in the pathogenesis of PDD is unknown.

On a quest for the causative agent of PDD, we investigated 3 birds with a PDD diagnosis based on clinical history and histologic criteria (Table 1). RNA was extracted from brains, pooled and randomly amplified for unbiased high-throughput sequencing (8), yielding 96,698 reads, ranging from 40 nt to 353 nt. After implementation of algorithms for vertebrate sequence subtraction and contiguous fragment assembly, GenBank searches using BLAST (http://blast.ncbi.nlm.nih.gov/blast.cgi) indicated a relation to BDV for a total of 11 contigs covering ≈1.1 kb of sequence distributed in 6 clusters throughout the N (230 nt), P (450 nt), G (250 nt), and L (120, 80, and 250 nt) genes (Figure 1). Divergent sequences between multiple overlapping contigs in these 6 regions indicated at least 2 different strains. Analysis of ≈5.5 kb of genomic sequence generated by standard PCR by using primers (sequence available upon request) based on the identified sequence fragments confirmed 2 strains, one from bird 1367, another from birds 1034 and 1322 (GenBank accession nos. FJ169440 and FJ169441, respectively) and indicated conservation of the unique genome organization that is characteristic for the family Bornaviridae (Figure 1). Sequence divergence between the avian strains (86% identity at nucleotide level) is an outbreak of neurologic disease in farmed ostriches in Israel has been attributed to BDV (4). BDV nucleic acids have been reported in feces of wild mallards and jackdaws in Sweden (5).

Proventricular dilatation disease (PDD), also known as proventricular dilatation syndrome or macaw wasting disease, is a disorder of birds wherein inflammation of the central, peripheral, and autonomic nervous systems is associated with gastrointestinal dysfunction and neurologic signs that may include ataxia and seizures (6). Although a presumptive diagnosis can be achieved through imaging studies and biopsy, for most animals, definitive diagnosis is made postmortem, only after detailed histologic analysis indicates the accumulation of lymphocytes in nerves that supply the proventriculus and ventriculus, in their associated ganglia, and in brain. Features consistent with PDD have been reported in >50 avian species; however, PDD is most commonly described in exotic companion birds such as macaws and parrots. Whether this reflects better case ascertainment or factors that influence exposure or susceptibility is unclear. An infectious basis is supported by the observation that disease can be transferred to naive birds through inoculation with tissue homogenates or fecal material from affected birds (7). One electron microscopic study showed the presence of spherical, 83-nm particles in macaw embryo cells after inoculation with feces from a diseased macaw; other researchers have described particles in tissue consistent in appearance with adenoviruses or paramyxoviruses. Whether any of these agents can be implicated in the pathogenesis of PDD is unknown.

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DOI: 10.3201/eid1412.080984

Emerging Infectious Diseases • www.cdc.gov/eid • Vol. 14, No. 12, December 2008
similar to that observed between the most divergent strains isolated from ungulates (84% identity). Pairwise compari-
son of the avian strains with these 2 ungulate isolates that represent the 2 previously known genotypes of BDV, strain V, NC_001607 (9), and No/98, AJ311524 (10), indicated <70% sequence conservation at the nucleotide level and <80% at the overall amino acid level (Figure 2, Table 2). These data are compatible with the avian strains represent-
ing a new species.

Primers and probes for quantitative real-time PCR were selected in the amino-terminal region of the phos-
phoprotein (P) gene matching P sequences for strain 1367 (set 1367, forward: 5′-AGAAGACCCGCTGACAGCA-3′, reverse: 5′-AAGCTTCTCGACGGGAACAG-3′, probe: 6FAM-5′-TCGTGGGGACCTCGATCTCACTCG-3′-TMR); the diag-
nostic assay has been made available through ProMed mail (www.promedmail.org, archive no. 20080726.2287). Real-
time PCR showed levels of viral RNA exceeding 10^7 copies in all tissues tested from birds with PDD but not in control
birds. In 2 birds (1034 and 1322) virus load was higher in the brain than in the proventriculus and gizzard or adrenal
gland; in 1 bird (1367), the load was lower in the brain than in the adrenals or proventriculi (Table 1).

Western immunoblot and nondenaturing dot blot ex-
periments were pursued by using brain, proventriculus, and
adrenal homogenates of the 3 animals that were quantitated
against recombinant BDV strain V nucleoprotein (N) or
P, as well as immune sera from BDV He/80–infected rats.
Positive and negative controls were BDV He/80–infected
rat brain homogenate and uninfected rat brain homogenate,
respectively. A strong signal was obtained with positive
control material in western immunoblots and dot blots that

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**Table 1. Real-time PCR measurement of viral sequences in birds with PPD**

| Case no. | Species† | Age/sex | PDD/ control | Primer/probe set | Organ | Ct | Virus copies‡ (300 ng total RNA) |
|----------|----------|---------|--------------|-----------------|-------|----|----------------------------------|
| 1034     | Canindae macaw | 30 y/M | PDD | 1034–1322 | Brain | 17.94 | 8.88 × 10^8 |
|          |          |        |              | Adrenal gland | 20.51 | 8.77 × 10^6 |
|          |          |        |              | Proventriculus and gizzard | 25.26 | 1.22 × 10^4 |
| 1322     | Vinaceous Amazon | 1 y/M | PDD | 1034–1322 | Brain | 17.64 | 1.17 × 10^4 |
|          |          |        |              | Proventriculus | 28.47 | 6.76 × 10^3 |
| 1367     | Canindae macaw | 30 y/M | PDD | 1367 | Brain | 27.33 | 8.89 × 10^5 |
|          |          |        |              | Adrenal gland | 18.92 | 3.87 × 10^4 |
|          |          |        |              | Proventriculus | 23.82 | 1.13 × 10^5 |
| 5473     | Leadbetter’s cockatoo | 17 y/M | Control | 1034–1322 and 1367 | Brain | >36§ | Negative |
| 4858     | Long-tailed parakeet | Adult/M | Control | 1034–1322 and 1367 | Brain | >36§ | Negative |
| 2020     | Cockatiel | Young/M | Control | 1034–1322 and 1367 | Brain | >36§ | Negative |
| 3616     | Eclectus parrot | 0.25 y/F | Control | 1034–1322 and 1367 | Brain | >36§ | Negative |

*PDD, poventricular dilatation disease; Ct, cycle threshold. †All birds were captive animals from California, USA. ‡Copy numbers were calculated on the basis of a standard curve generated from cloned target sequences. §A Ct of >36 was rated as negative based on the highest dilution of standard representing 5 copies.

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**Figure 1. Conservation of genome organization, regulatory sequences, and protein domains of Borna disease virus (BDV) in novel strains from parrots 1034, 1322, and 1367.**
used either antisera to recombinant viral proteins or rat immune sera; no signal was obtained with bird homogenates or uninfected rat brain homogenate.

Conclusions

We have not yet determined whether purified virus induces PDD or an adaptive immune response occurs in association with disease. Nonetheless, given what is known about BDV pathogenesis with other strains in other hosts (11), the bornaviruses identified in these birds must be considered a biologically plausible candidate causative agent. Infection, lymphocyte infiltration, and dysfunction of the central, peripheral, and autonomic nervous system are common to PDD as well as to classical Borna disease in natural disease and experimental models (2,12,13). Proteins and antisera we have used for 2 decades for BDV diagnostics failed to detect this virus in our PCR-positive birds. Thus, it will be important to revisit epidemiologic surveys that we, and others, have undertaken to investigate the role of bornaviruses in human disease (14). From a personal perspective, we are intrigued that whereas molecular discovery of the first BDV in the late 1980s required an investment of 2 years in subtractive cloning (15), high-throughput sequencing, bioinformatics, and sequence databases enabled discovery of these 2 strains in 2 weeks.

This work was supported by National Institutes of Health awards AI062705, AI051292, HL083850, and AI57158 (Northeast Biodefense Center – Lipkin).

During review of this manuscript similar findings were independently reported by Kistler AL, Gancz A, Clubb S, Skewes-Cox P, Fischer K, Sorber K, et al. Recovery of divergent avian bornaviruses from cases of proventricular dilatation disease: identification of a candidate etiologic agent. Virol J. 2008;5:88.

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Table 2. Percent sequence conservation between Borna disease virus strain V(No/98) and novel strains

| Amino acid | Strain V | No/98 | 1034/1322 | 1367 |
|------------|----------|-------|-----------|------|
| Nucleotide|          |       |           |      |
| Strain V  |          |       |           |      |
| P: 99/96* | 84       | 67    | 66        |      |
| X: 84/81 |           |       |           |      |
| M: 100/98 | 66       |       |           |      |
| G: 95/94 |           |       |           |      |
| L: 98/96 |           |       |           |      |
| 1034/1322 |          |       |           |      |
| P: 76/61 | 77/63    |       |           |      |
| X: 55/51 | 49/45    |       |           |      |
| M: 91/85 | 91/85    |       |           | 86 |
| G: 75/66 | 75/66    |       |           |      |
| L: 81/74 | 81/74    |       |           |      |
| 1367     |          | 76/63 |           |      |
| P: 98/95 | 88/88    |       |           |      |
| X: 99/95 | 99/95    |       |           |      |
| M: 95/86 | 94/83    |       |           |      |
| G: 74/66 | 74/66    |       |           |      |
| L: 82/76 | 82/76    | 87/97 |           |      |

* Amino acid conservation is indicated as percent similarity/percent identity.

Figure 2. A) Similarity plot between Borna disease virus (BDV) prototype strain V nucleotide sequence and those of characterized BDV strains He/80fr and No/98 compared with novel strains 1367 and 1034/1322. Gene regions corresponding to the nucleoprotein (N), phosphoprotein (P), X protein (X), matrix protein (M), glycoprotein (G) and L-polymerase protein (L), and nucleotide positions are indicated. B) A tree representing the evolutionary history was inferred by using the neighbor-joining method. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (500 replicates) is shown next to the branches. The tree is drawn to scale with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree (number of base substitutions per site; see scale bar). Evolutionary distances were computed by using a Kimura 2-parameter model; a total of 5,449 positions in the final dataset were analyzed by using MEGA4 software (www.megasoftware.net).
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