Microsporidia of the genus *Encephalitozoon* (*E. cuniculi*, *E. hellem*, and *E. intestinalis*) are intracellular pathogens infecting a wide range of animal species. Because spores can be released to the environment via hosts’ feces, urine, and respiratory secretions, they can be ingested or inhaled, posing a risk for zoonotic infection in humans (1). The primary site of *Encephalitozoon* spp. infection is the small-intestine epithelium, but dissemination and systemic infections are also well known. Infection of a broad spectrum of cell types has been noted, especially for *E. cuniculi* (1); various pathological changes affecting the digestive, urinary, and respiratory tracts and the nervous system may occur. Because encephalitozoons are opportunistic pathogens, the extraintestinal and disseminated infections and severe symptoms they cause are of concern in immunocompromised hosts, such as transplant recipients or persons living with HIV (2,3). Microsporidiosis develops in patients whose immune response has been weakened by diabetes or malignant disease treated with chemotherapy (2,4).

We describe the case of 2 bird owners in Poland who acquired *E. cuniculi*-caused microsporidiosis from their infected pet birds. The Human Research Ethics Committee of Wroclaw Medical University (Wroclaw, Poland) approved this study in accordance with agreement no. KB-549/2012. Patients provided written informed consent before examination.

**The Study**

The 2 patients, a woman and a man, both 41 years of age, had nonspecific symptoms of fatigue, exhaustion, joint and muscle pain, frequent colds, and headaches, progressive and more severe in the woman. We observed fever reaching 38°C and lasting several months in both patients. Moreover, the woman had intense night palpitations, symptoms similar to bronchitis (occasionally treated with antimicrobial drugs), blurred vision, dizziness, and impaired concentration. She had had bipolar disorder and diabetes for years. Magnetic resonance imaging scans of her head revealed single minor demyelinating or vascular changes in the white matter of the frontal lobes. No information about diabetes treatment was available. No abnormalities were shown in abdominal ultrasound or chest radiograph. She tested seronegative for *Borrelia burgdorferi* and *Chlamydia psittaci* infection. Except for a low leukocyte count (3.00 cells/µL), the woman’s basic laboratory tests of blood and urine and her electrocardiograms showed no abnormalities.

Symptoms emerged in the patients after 2 years of breeding exotic birds together. They had 30 birds of various species: budgerigars, canaries, diamond doves, tricolored parrotfinches, Gouldian finches, and diamond firetails. During the 2-year period, 17 birds died from infectious and metabolic diseases, trauma, and management-related issues.

As a part of standard flock management practice, we performed tentative postmortem diagnostic

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**Encephalitozoon cuniculi** and Extraintestinal Microsporidiosis in Bird Owners

Marta Kicia, Żaneta Zajączkowska, Martin Kváč, Kamil Cebulski, Nikola Holubová, Piotr Wencel, Leszek Mayer, Maria Wesolowska, Bohumil Sak
cytology in 1 tricolored parrotfinch; we suspected disseminated microsporidial infection and conducted a detailed investigation of pooled feces collected from 8 aviaries at the patients’ home and tissues from a dead budgerigar. All samples were delivered on ice to the Laboratory of Veterinary and Medical Protistology, Biology Centre of the Czech Academy of Sciences (České Budějovice, Czech Republic). We collected patients’ urine and stool specimens every 2 days for 1 week, 3 samples from each, during diagnosis and follow-up and examined them at the Department of Biology and Medical Parasitology, Wrocław Medical University.

We homogenized all samples by mechanical disruption before genomic DNA extraction as described previously (5). We performed nested PCR protocols amplifying a partial sequence of 16S rRNA gene (130 bp), the entire internal transcribed spacer region, and a partial sequence of 5.8S rRNA gene (137–139 bp) of *Encephalitozoon* spp. (6). In addition, we checked for the presence of *Enterocytozoon bieneusi* (7) and then performed a phylogenetic analysis of PCR products (5). We used standard light microscopy methods for spore detection (8). We deposited the sequences obtained in this study in GenBank (accession nos. OK356650–60).

We detected microsporidial DNA in the fecal specimens of all birds and in tissue samples from both tested birds. Genotyping revealed the presence of *E. hellem* genotype 1A in all tested specimens from the lung, liver, duodenum, and jejunum with ileum from the budgerigar, whereas *E. cuniculi* genotype II was found in pooled feces and in all sections from the liver, gizzard with proventriculus, and duodenum with jejunum and ileum from the tricolored parrotfinch. *E. cuniculi* genotype II was present in 2 of the woman’s urine samples and in 1 of the man’s urine samples (Figure). Spores were confirmed in all samples. *E. bieneusi* was not found. We administered 400 mg albendazole daily for 10 days to both patients. In follow-up examination 3 months after treatment, the patients’ urine repeatedly tested negative for *E. cuniculi* in all independent samplings, and the patients gradually improved. Patients’ stools remained negative during the entire diagnostic process. No subsequent follow-up was conducted.

![Figure](image-url). Phylogenetic relationships of *Encephalitozoon cuniculi* genotype II and *E. hellem* genotype 1A obtained from 2 exotic bird breeders and 2 of their birds compared with other *Encephalitozoon* species and genotypes. Bold type indicates sequences obtained in this study, identified by isolate number (e.g., C1.615); black circles indicate isolates from humans; squares indicate isolates from birds. We analyzed a partial sequence of 16S rRNA gene, the entire internal transcribed spacer region, and a partial sequence of 5.8S rRNA gene inferred by neighbor-joining analyses and computed using the Tamura 3-parameter method. We modeled the rate variation among sites with a gamma distribution. Percentages of replicate trees in which the associated taxa clustered together in the bootstrap test (1,000 replicates) are shown next to the branches. The final dataset contained a total of 220 positions. GenBank accession numbers are in parentheses. Scale bar indicates nucleotide substitutions per site.
Conclusions

Birds are a common source in the propagation of encephalitozoons in the environment (prevalence ≤15%), either acting as mechanical vectors or developing active infection (9–11). In both wild-living and captive birds, E. hellem is the most prevalent Encephalitozoon species, whereas E. cuniculi has been detected less frequently (9,10). Whether animal hosts indeed propagate microsporidia or serve as transmission vectors is debatable (9). However, the presence of pathogens in both feces and tissues of birds in our study confirms microsporidial proliferation in these animals rather than the passage of spores through the digestive tract; this finding suggests that encephalitozoons may have been circulating in this breeding group for some time. Although most earlier reports demonstrated asymptomatic infections and low infection intensity among birds, intermittent spore shedding in naturally infected birds contaminates the environment (11). Owners could be in constant contact with spores, which highly increases the risk for infection by ingestion or inhalation of spores. As a slow-growing pathogen, E. cuniculi can lead to chronic infection and microsporidiosis.

In immunocompetent hosts, an immune-controlled balance in the host–parasite relationship is established, and extraintestinal infections remain asymptomatic (5). Symptomatic cases of microsporidial infection usually manifest as self-limiting diarrhea (2). Symptomatic extraintestinal Encephalitozoon infections in immunocompetent humans are uncommon and usually present as keratoconjunctivitis (2). Of note, disseminated microsporidiosis caused by E. cuniculi genotype I with involvement of brain and urinary and intestinal tracts has been described in men with type 2 diabetes (4). However, diabetes is considered a risk factor for opportunistic infections. Experimental infection of diabetic mice with E. cuniculi resulted in more symptoms and a higher pathogen burden than in nondiabetic animals (12); indeed, the woman with diabetes in our study experienced more severe symptoms, which led to serious impairments in everyday functioning.

Even though we were able to test only urine and feces and confirmed E. cuniculi in the urinary tract, the symptoms we observed in our patients were complex, indicating disseminated infection. We cannot be confident in the extent to which E. cuniculi infection contributed to these symptoms. However, symptom relief coincided with pathogen clearance after albendazole treatment, which convinced us that microsporidia may have been at least partially involved in symptom development. The severity of symptoms despite the lack of lifelong immunosuppression is puzzling and may arise from high doses of spores acquired as a result of patients’ everyday contact with birds.

In summary, our study documents the risk for bird-to-human transmission of E. cuniculi parasites. Exotic-bird breeders should be aware of the risk for infection with this opportunistic pathogen.

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About the Author

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June 2020

Prions

- Identifying and Interrupting Superspreading Events—Implications for Control of Severe Acute Respiratory Syndrome Coronavirus 2
- Risks Related to Chikungunya Infections among European Union Travelers, 2012–2018
- Manifestations of Toxic Shock Syndrome in Children, Columbus, Ohio, USA, 2010–2017
- Genomic Epidemiology of 2015–2016 Zika Virus Outbreak in Cape Verde
- Epidemiologic Changes of Scrub Typhus in China, 1952–2016
- Pharmacologic Treatments and Supportive Care for Middle East Respiratory Syndrome
- Distribution of Streptococcal Pharyngitis and Acute Rheumatic Fever, Auckland, New Zealand, 2010–2016
- Temporary Fertility Decline after Large Rubella Outbreak, Japan
- Radical Change in Zoonotic Abilities of Atypical BSE Prion Strains as Evidenced by Crossing of Sheep Species Barrier in Transgenic Mice
- Characterization of Sporadic Creutzfeldt-Jakob Disease and History of Neurosurgery to Identify Potential Iatrogenic Cases
- Failures of 13-Valent Conjugated Pneumococcal Vaccine in Age-Appropriately Vaccinated Children 2–59 Months of Age, Spain
- Increased Risk for Carbapenem-Resistant Enterobacteriaceae Colonization in Intensive Care Units after Hospitalization in Emergency Department
- Antimicrobial Resistance in Salmonella enterica Serovar Paratyphi B Variant Java in Poultry from Europe and Latin America
- Invasive Group B Streptococcus Infections in Adults, England, 2015–2016
- Zoonotic Alphaviruses in Fatal and Neurologic Infections in Wildlife and Nonequine Domestic Animals, South Africa
- Effectiveness and Tolerability of Oral Amoxicillin in Pregnant Women with Active Syphilis, Japan, 2010–2018
- Endemic Chromoblastomycosis Caused by Fonsecaea nubica, Madagascar
- Emergence of New Non–Clonal Group 258 High-Risk Clones among Klebsiella pneumoniae Carbapenemase–Producing K. pneumoniae Isolates, France
- Zoonotic Vectorborne Pathogens and Ectoparasites of Dogs and Cats in Eastern and Southeast Asia
- Multihost Transmission of Schistosoma mansoni in Senegal, 2015–2018
- Statin Use and Influenza Vaccine Effectiveness in Persons ≥ 65 Years of Age, Taiwan
- Estimating Risk for Death from Coronavirus Disease, China, January–February 2020
- Epidemiology of Coronavirus Disease in Gansu Province, China, 2020
- Severe Acute Respiratory Syndrome Coronavirus 2 from Patient with Coronavirus Disease, United States
- Syphilis in Maria Salvati (1499–1543), Wife of Giovanni de’ Medici of the Black Bands
- Yaws Disease Caused by Treponema pallidum subspecies pertenue in Wild Chimpanzee, Guinea, 2019
- Fatal Encephalitis Caused by Cristovirus, an Emerging Orthobunyavirus, France
- Increased Community-Associated Clostridioides difficile Infections in Quebec, Canada, 2008–2015
- Melioidosis in a Resident of Texas with No Recent Travel History, United States
- No Adaptation of the Prion Strain in a Heterozygous Case of Variant Creutzfeldt-Jakob Disease
- Prevalence of Escherichia albertii in Raccoons (Procyon lotor), Japan
- Cannabis Use and Fungal Infections in a Commercially Insured Population, United States, 2016

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