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Acute gastritis associated with invading *Helicobacter heilmannii*
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To the Editor:

*H. heilmannii* are unculturable, and gram-negative rod, apparently larger than *H. pylori*, and are not usually (frequency 0.1-6.2%) observed in routine gastric biopsy.\(^1\,^2\) Most infected stomachs show only mild gastritis but rarely is it associated with acute gastritis, peptic ulcer, gastric carcinoma or mucosa-associated lymphoid tissue lymphoma.\(^2\,^3\) Although *H. heilmannii* exclusively floats in the mucous layer above the surface and foveolar epithelial cells, Heilmann and Borchard using transmission electron microscopy showed that the organisms could attach to and penetrate into gastric parietal cells.\(^1\) Using giemsa stain and immunostaining, we first report that *H. heilmannii* can penetrate to gastric epithelial cells and further invade into the stroma.

A previously healthy 58-year-old man presented with epigastric pain. Gastric endoscopy showed severe hemorrhage and erosion in the body and the antrum of his stomach. A biopsy taken from body mucosa showed erosion, hemorrhage, marked infiltration of polymorphonuclear leukocytes (neutrophils), and glandular destruction (Fig. 1a). Giemsa stain revealed clusters of corkscrew-shaped bacilli in the surface mucous layer and some foveolae. The organisms were positive for polyclonal anti-*Helicobacter pylori* antibody (DakoCytomation, Kyoto, Japan), consistent with *Helicobacter (H.) heilmannii*. Spiral bacilli were also present within foveolar epithelial cells and in the stroma (Fig. 1a inset). He underwent antimicrobial treatment by oral administration of lansoprazole, amoxillin and clarithromycin; simulating the standard treatment for *H. pylori* gastritis. Re-endoscopy two weeks after the initial gastroscopy showed an improvement of his acute gastritis and the disappearance of the organisms; however, two healing
linear ulcers were retained in his antral mucosa. Gastroscopy and biopsy six months after the initial endoscopy showed no evidence of abnormal mucosal lesions or *H. heilmannii* infection.

For over three years he had kept in his house three healthy pet dogs with good pedigrees obtained from breeders. Several months before the initial gastroscopy, he picked up a healthy but then homeless cat and kept it in his restaurant that was separated from his house. The dogs and the cat were anesthetized and underwent gastroscopy with the consent of the patient. Biopsy from all the animals revealed minimal to moderate gastritis with the specimens from the cat showing spiral bacilli presenting in the mucous layer, gastric foveolae, and cytoplasm of parietal cells with intracytoplasmic vacuolation around the organisms (Fig. 1b). They also cross immunostained with anti-*H. pylori* antibody.

The frequency of *H. heilmannii* infection in dogs is 67 to 100% and in cats it is 42 to 100%, while its infection in humans has been postulated to be an example of zoonosis. The patient’s dogs that lived in his house were not infected with the organism because they had been kept apart from the infected cat kept that lived in his restaurant. This suggests that *H. heilmannii* infection between animals is connected with poor sanitary conditions in which infected animals live together. Owners of pet animals that have grown up in a clean environment are not always oversensitive to *H. heilmannii* infection. This case of acute *H. heilmannii* gastritis was almost certainly caused by organisms derived from patient’s cat; if so, the incubation period of the disease may be presumed to have been within several months. The mechanism by which indolent organisms in a cat could cause acute gastritis in a human stomach is unknown; however, a possible explanation may
include that a potentially aggressive strain of *H. heilmannii* penetrated into the epithelial cells and the stroma resulting in neutrophilic infiltration and subsequent glandular destruction. We concluded that invading *H. heilmannii* organisms could cause acute gastritis in humans when animals with the organisms are kept as a pet.

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**References**


**Figure legend**

FIG. 1 (a) Gastric histology of the patient (magnification×100); the inset highlights spiral *H. heilmannii* organisms (arrows) in the destructed foveolae and in the stroma (×1000, anti-*H. pylori* immunostaining). (b) Histology of the previously homeless cat showing colonized organisms (arrows) in the cytoplasm of parietal cells (×1000, Giemsa).