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Long-term proton pump inhibitor administration, *H pylori* and gastric cancer: lessons from the gerbil

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The association between chronic active gastritis and pre-neoplastic conditions as well as invasive cancer of the stomach was established several decades ago. The risk of progression depended on the severity and distribution of gastritis, with cancer, in particular, occurring in subjects with pan-gastritis. Subsequently, *Helicobacter pylori* was recognised as the primary cause of chronic active gastritis, and it was demonstrated that the pattern of gastritis corresponded with the colonisation pattern of *H pylori*. We and others then showed both in animals and in humans that this pattern of colonisation and associated gastritis primarily depended on the level of acid output. Although this hypothesis was widely accepted, it led to intense debate when dealing with the safety of long-term treatment with profound acid suppressors. An elegant, long-awaited study from Japan published in this issue of *Gut* (see page 624) provides compelling evidence that the pattern of *H pylori* colonisation depends on acid output and that this influences the long-term progression to neoplasia.

The current study was based on experiments in gerbils, one of the well-established animal models for the study gastric disease induced by *Helicobacter* spp. Gerbils, used sparingly for biomedical research, were first reported as a model for experimental *H pylori* infection in 1991. Interestingly, the gerbil has been shown to have particular relevant features that can be used to address whether *H pylori* can induce gastric cancer. Japanese investigators have noted intestinal metaplasia, atrophy and gastric ulcers in gerbils experimentally infected with *H pylori*. Following these findings, others noted that gerbils infected with *H pylori* from periods ranging from 15 to 18 months develop gastric adenocarcinoma. The gastric cancers were clearly documented histologically. Vascular invasion and metastases were not observed in either study.

The histological progression of *H pylori*-associated disease in the gerbil closely resembles that observed in humans, including the early appearance of intestinal metaplasia, well-differentiated histological patterns of gastric malignancy, and antral location of the gastric cancers. The development of cancer in the gerbil is also preceded by invagination of atypical glands (cystica profunda) into the submucosa. The association with gastric ulcers in this model continues to be of interest, given clinical studies in human patients indicating a link between gastric ulcer disease and gastric cancer.

Although most of the tumours in the *H pylori* gerbil model originate in the pyloric region of the stomach, significant changes in the oxyntic mucosa consistent with chronic atrophic gastritis are also observed. Glandular tissues in the gastric body and fundus are replaced by atrophy and hyperplastic epithelium of the pseudopyloric type. A similar type of gastric atrophy (loss of oxyntic glands and neck cell hyperplasia) has also been reported in *Helicobacter felis* or *H pylori*-infected C57BL mice.

The gerbil appears to be uniquely susceptible to *H pylori*-induced gastric neoplasia, and further characterisation of the gerbil model has provided important clues on gastric cancer progression and host–bacteria interaction. The unusual susceptibility of this animal species to gastric cancer using fairly standard *H pylori* strains underscores once again the over-riding importance of host factors in determining the outcome from gastric *Helicobacter* spp. infection. Further, a possible role for altered gastrin physiology in the pathogenesis of gastric cancer has been raised by several recent studies. In Mongolian gerbils, *H pylori* leads to marked elevation of serum gastrin levels which coincide temporally with increases in gastric mucosal proliferation rates. This is consistent with experiments in beagle (FVB/N) mice that were rendered moderately hypergastrinaemic through an insulin–gastrin (INS/GAS) transgene. In this model there is increased mucosal proliferation and progressive atrophy, and gastric carcinomas develop in 20-month-old INS/GAS mice. Infection of these hypergastrinaemic mice with *H felis* and *H pylori* accelerates tumorigenesis, with the majority (85%) of infected male mice developing gastric cancer within 8 months after colonisation. We recently reported that this chronic, progressive process is enhanced by co-colonisation with enteric flora.

Using this well-described and reproducible gerbil model, Japanese investigators have conducted a very important study, examining the potential effect of proton pump inhibitor (PPI) therapy on progression of *H pylori*-associated disease. The male gerbils were divided into four groups: *H pylori* (ATCC43504) positives and negatives, with and without administration of a PPI (omeprazole 100 mg/kg body weight/day). At the end point of the 6-month experiment, the authors provided detailed analysis of gastric pathology including morphometric severity of parietal cell loss. The authors made a concerted effort to address confusion in the literature with respect to what constitutes a ‘gastric adenocarcinoma’ in a Mongolian gerbil model and how to differentiate an ‘invasive carcinoma’ from ‘heterotropic glands’ that are frequently encountered in these models during the early course of the disease. The ability to differentiate glandular herniation into the submucosa from the true invasion is an important consideration in evaluating rodent models of gastric cancer induced by *Helicobacter* spp. Indeed with these histopathological criteria in mind the results were remarkable and provide considerable support for previously published literature in humans. *H pylori*-negative male gerbils did not develop gastritis, metaplasia, or cancer irrespective of PPI treatment. *H pylori*-positive gerbils all developed gastritis, and most also developed metaplasia during the later stages of disease. Treatment with omeprazole in *H pylori*-positive animals accelerated progression to atrophy and enhanced hypergastrinaemia,
resulting in a significantly increased incidence of gastric cancer. The authors emphasise that hypergastrinaemia might be promoting the development of H pylori gastric cancer. Given gender susceptibility to H pylori-associated gastric cancer in humans and rodent models, additional studies using H pylori female gerbils on chronic PPI therapy should provide additional interesting insights.

Overall, the findings in the current paper support previous findings by us and others showing the increased risk of gastric atrophy in H pylori-infected patients on PPI therapy. In humans, it remains to be shown whether chronic PPI therapy in H pylori-infected individuals also increases the further progression to metaplasia, dysplasia and neoplasia. These studies will require longer-term follow-up of this subset of patients. In the meantime, the current findings in this paper support the argument regarding the importance of considering H pylori eradication in patients on long-term PPI treatment to cure gastritis, lower gastrin levels, and prevent progression to atrophy, dysplasia and gastric cancer. This recommendation is consistent with European guidelines on H pylori treatment in humans.

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